800 year ice-core record of nitrogen deposition in Svalbard linked to ocean productivity and biogenic emissions

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13 Abstract

14 We present the records of the two nitrogen species nitrate (NO_3) and ammonium (NH_4) 15 analysed in a new ice core from Lomonosovfonna, Svalbard, in the Eurasian Arctic covering 16 the period 1222-2009. Changes in melt at the Lomonosovfonna glacier are assumed to have a negligible effect on the decadal variations of the investigated compounds. Accordingly, we 17 use decadal records to investigate the major emission sources of NO_3^- and NH_4^+ precursors 18 and find that during the 20th century both records are influenced by anthropogenic pollution 19 20 from Eurasia. In pre-industrial times NO_3^- is highly correlated with methane-sulfonate 21 (MSA), which we explain by a fertilising effect. We propose that enhanced atmospheric NO_3^{-1} 22 concentrations and the corresponding nitrogen input to the ocean trigger the growth of 23 dimethyl-sulfide-(DMS)-producing phytoplankton. Increased DMS production results in 24 elevated fluxes to the atmosphere where it is oxidised to MSA. Eurasia was presumably the 25 main source area also for pre-industrial NO₃, but a more exact source apportionment could not be performed based on our data. This is different for NH₄⁺, where biogenic ammonia 26 27 (NH₃) emissions from Siberian boreal forests were identified as the dominant source of preindustrial NH₄⁺. 28

2 **1** Introduction

3 The Arctic is generally a nutrient limited region (Dickerson, 1985). Nutrients originate from 4 lower latitudes and reach the remote polar areas via long-range transport, local sources are 5 sparse. The major source for bio-available nitrogen in the Arctic is the deposition of reactive atmospheric nitrogen that is present primarily as nitrate (NO_3) and ammonium (NH_4) 6 7 (Björkman et al., 2013; Kühnel et al., 2011). Those species are predominantly removed from 8 the atmosphere by wet deposition (Bergin et al., 1995). NO_3^- is the oxidation product of 9 emitted NO_x (NO and NO₂). At a global scale, major NO₃⁻ sources include biomass burning, 10 emissions from microbial processes in soils, ammonia oxidation, stratospheric injection, 11 lightning, as well as fossil fuel and biofuel combustion, and aircraft emissions (Fibiger et al., 2013; Galloway et al., 2004; Hastings et al., 2004; Wolff, 2013). NH₄⁺ derives from biogenic 12 emissions of ammonia (NH₃) from terrestrial and marine sources, biomass burning, 13 14 agriculture, and livestock breeding (Fuhrer et al., 1996; Galloway et al., 2004; Wolff, 2013). Both NO_3^- and NH_4^+ concentrations in the atmosphere have varied greatly with time and 15 16 space due to changing emissions and the short atmospheric lifetimes of a few days (Adams et 17 al., 1999; Feng and Penner, 2007). Generally, concentrations were low in pre-industrial times 18 and increased due to stronger emissions with beginning of the industrialisation and 19 intensification of agricultural activities (Galloway et al., 2004). The deposition of NO₃⁻ and NH_4^+ in the Arctic is an important nutrient source. Varying concentrations thus greatly affect 20 the nitrogen budget in the Arctic where nutrient supply is limited. 21

22 Ice cores represent an invaluable archive of past atmospheric composition. Ice core studies 23 from the Arctic clearly reveal an anthropogenic influence on the concentrations of NO₃⁻ and NH4⁺ approximately during the last 150 years (Fischer et al., 1998; Fuhrer et al., 1996; Goto-24 Azuma and Koerner, 2001; Kekonen et al., 2002, 2005; Legrand and Mayewski, 1997; 25 26 Matoba et al., 2002; Simões and Zagorodnov, 2001). North America was identified as major 27 pollutant source for south Greenland, both North America and Eurasia for central and north Greenland, and Eurasia for Svalbard (Goto-Azuma and Koerner, 2001; Hicks and Isaksson, 28 2006). However, the pre-industrial sources of NO_3^- and NH_4^+ are still fairly unknown 29 30 (Legrand and Mayewski, 1997; Wolff, 2013). Eichler et al. (2011) identified forest fires as 31 major source of NO₃⁻ in a Siberian Altai ice core from the mid-latitudes. In studies on 32 Greenland ice NO₃⁻ was also associated with forest fires (Whitlow et al., 1994; Wolff et al., 2008). Pre-industrial NH₄⁺ in ice cores from the mid-latitudes was attributed to biogenic
 emissions (Eichler et al., 2009; Kellerhals et al., 2010). Similarly, long-term trends in
 Greenland ice cores have been attributed to changing biogenic emission from North America,
 whereas short-term NH₄⁺ changes were found to correlate with forest fires (Fuhrer et al.,
 1996; Whitlow et al., 1994; Zennaro et al., 2014).

6 Whereas a few records exist from Greenland, there is less information available from the 7 Eurasian Arctic. The NO_3^- and NH_4^+ records of a previous ice core from Lomonosovfonna, Svalbard, retrieved in 1997 (Lomo97, for location see Fig. 1), cover the last 1000 years 8 9 (Divine et al., 2011; Kekonen et al., 2002, 2005). For both species a clear anthropogenic impact is observed in the second half of the 20th century, but the pre-industrial sources remain 10 largely unidentified due to potential runoff that biased the ion records before the mid-16th 11 century (Kekonen et al., 2002, 2005). Nevertheless, the fairly stable concentrations in the 12 NO₃⁻ record from the mid-16th to the mid-19th century are interpreted as input from natural 13 NO₃⁻ sources (Kekonen et al., 2002). An anthropogenic influence in the 20th century is also 14 visible in the NO_3^- and NH_4^+ records of other Eurasian Arctic ice cores (see Fig. 1 for 15 locations) from Holtedahlfonna (Holte05), Svalbard (Beaudon et al., 2013), Snøfjellafonna, 16 17 Svalbard (Goto-Azuma and Koerner, 2001), and Severnava Zemlya (Weiler et al., 2005). The industrial records from these cores are discussed in detail, but pre-industrial sources and 18 19 concentration changes of the inorganic nitrogen species remain unexplained.

The interpretation of NO_3^- and NH_4^+ as paleo-environmental proxies may be hampered by the 20 fact that both undergo post-depositional processes leading to loss from or relocation within 21 22 the snow pack even at temperatures well below the melting point (Pohjola et al., 2002). NO_3^{-1} can be relocated or lost by photolysis and/or evaporation of nitric acid (HNO₃) (Honrath et al., 23 24 1999; Röthlisberger et al., 2002). This loss can be severe at low accumulation sites such as Dome C, Antarctica (Röthlisberger et al., 2000, 2002). At sites with higher accumulation rates 25 26 such as Summit in Greenland or Weissfluhjoch in the European Alps the majority of NO₃⁻ is preserved (Baltensperger et al., 1993; Fibiger et al., 2013). Many studies reveal that NH₄⁺ and 27 NO_3^{-} are preserved in snow and firn cores with respect to percolating melt water (Eichler et 28 al., 2001; Ginot et al., 2010; Moore and Grinsted, 2009; Pohjola et al., 2002), but others report 29 30 a preferential elution of these species compared to other major ions (Brimblecombe et al., 1985; Moore and Grinsted, 2009; Pohjola et al., 2002). The underlying mechanism is not well 31 32 understood, except from the fact that it depends on the overall ion composition.

In this paper we discuss the records of the two nitrogen species NO_3^- and NH_4^+ analysed in a new ice core drilled on Lomonosovfonna, Svalbard, in 2009. The study focuses on the investigation of the major sources of NO_3^- and NH_4^+ deposited in the Eurasian Arctic which highly affects the nutrient budget in the region, along with the effect of melt on the geochemical records of these nitrogen species which will gain importance due to the ongoing global warming.

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8 2 Methods

9 2.1 Drilling site and meteorological setting

10 In 2009, a 149.5 m long ice core was drilled on Lomonosovfonna, Svalbard (1202 m asl; 78°49`24``N, 17°25`59``E, Lomo09), using the Fast Electromechanical Lightweight Ice 11 12 Coring System (FELICS) (Ginot et al., 2002). The 2009 drilling site is 4.6 km south of that in 1997 (Lomo97) (Isaksson et al., 2001). Bedrock was not reached but a radar survey suggested 13 14 it to be at around 200 m (Pettersson, unpublished data). Measured borehole temperatures in the upper 42 m (between -1.7°C and -4.3°C) at the Lomo09 drill site are in good agreement 15 with the average borehole temperature at the Lomo97 site of -2.8°C with a nearly isothermal 16 profile (Van de Wal et al. 2002). Previous studies indicate that summer melt water in the 17 study area is refrozen mostly within the previous winter's snow, and the remainder within the 18 next two to three lower annual layers (Samuelson, 2001). Percolation length was found to be 19 up to 8 years only in the warmest years during the 20th century (Kekonen et al., 2005; Moore 20 21 et al., 2005).

22 Svalbard is located at a climatically sensitive area being surrounded by the Arctic Ocean, the 23 Barents Sea and the Atlantic Ocean, and situated at the southerly edge of the permanent 24 Arctic sea ice and close to the over-turning point of the North Atlantic thermohaline 25 circulation. Further, it is relatively close to the industrialised areas of Eurasia which were 26 found to highly affect the chemical composition of air reaching the archipelago, especially in 27 spring during the Arctic Haze (Eleftheriadis et al., 2009; Eneroth et al., 2003; Forsström et al., 28 2009; Goto-Azuma and Koerner, 2001; Law and Stohl, 2007; Stohl et al., 2007). The Arctic 29 Haze describes a phenomenon of increased aerosol concentration in the end of winter to early 30 spring (Greenaway, 1950; Quinn et al., 2007; Shaw, 1995). At that time of the year 31 temperatures in the Arctic become very low which leads to a thermally very stable

stratification with strong surface inversions (Shaw, 1995; Stohl, 2006). This cold stratified air 1 2 forms a dome over the Arctic that hinders warm air masses from lower latitudes to enter. The boundary of this dome that acts as a transport barrier is called Arctic or Polar Front whose 3 position shifts between summer and winter due to temperature. In summer only the more 4 5 northern parts of the Northern Hemisphere are cold enough to cause a stable stratification of the atmosphere, whereas in winter temperatures in more southern parts are cold enough so 6 7 that the Arctic Front is located as far south as 40°N. Then large areas of Eurasia and partly 8 North America are included in the Arctic dome, facilitating transport of pollution from those 9 regions. In addition, since both dry and wet deposition is reduced within the Arctic dome in 10 winter, aerosol related species have very long lifetimes once within the Arctic dome (Stohl, 11 2006).

12 **2.2 Sampling and analyses**

The Lomonosovfonna 2009 ice core was processed in the cold room (-20°C) at Paul Scherrer Institut, Switzerland, resulting in 3997 samples with a depth resolution of 3-4 cm (details on the method in Eichler et al., 2000). The resolution was adapted to layer thinning with depth, so that even in the deepest and oldest part of the core each year is at least represented by one sample. The inner part of the core was sampled for the analysis of water soluble major ions and the water stable isotopes δ^{18} O and δ D. Outer core sections were analysed for ³H and ²¹⁰Pb used for dating purposes (Eichler et al., 2000).

Concentrations of water soluble major ions, including NO_3^- and NH_4^+ , were determined using ion chromatography (Metrohm 850 Professional IC combined with a 872 Extension Module and a 858 Professional Sample Processor autosampler). Prior to analyses ice samples were melted in a N₂ atmosphere to reduce contamination from laboratory air. A list of the measured ionic species, their detection limits and median concentrations are given in Table 1.

25 2.3 Ice core dating

The Lomo09 ice core covers the time period of 1222 to 2009 (Fig. 2). It was dated with a combination of reference horizons, annual layer counting (ALC), ²¹⁰Pb decay, and a simple glacier flow model. The reference horizons include the tritium peak indicating the year 1963, and the major volcanic eruptions of Bezymianny (1956), Katmai (1912), Tambora (1815), Laki (1783), Hekla (1766), Kuwae (1458/59; Sigl et al., 2013), and Samalas (1257/58;

Lavigne et al., 2013) marked by high non-sea-salt sulphate concentrations and high values for 1 2 the sulphate-residual of the multiple linear regression of all measured ions, a method previously described in Moore et al. (2012). Annual layer counting was performed down to a 3 depth of ~79.7 m weq (= 1750) using the pronounced seasonality of $\delta^{18}O$ and Na^+ 4 (Supplement Fig. S1). A simple glacier flow model (Thompson et al., 1998) was fitted 5 through the volcanic reference horizons. This was used to date the core below ~79.7 m weg 6 7 where ALC was limited due to strong layer thinning. The dating uncertainty for the core down 8 to a depth of ~68 m weg is estimated to be ± 1 year within ± 10 years of the reference horizons 9 and increases to ± 3 years in between. Down to a depth of ~80 m weg the dating uncertainty enlarges to ± 3 years also in proximity of the reference horizons, and below ~80 m weq it 10 increases to ± 10 years. This was calculated using the difference of the year of the volcanic 11 12 eruptions and the modelled date. The average annual accumulation rate is 0.58 ± 0.13 m weq.

13 **2.4 Calculation of annual melt percent**

14 Melt features are formed when surface snow melts and the melt water percolates into deeper 15 layers where it fills the pores and refreezes under the formation of a layer of ice poor or free 16 of air bubbles. The percentage of annual melt in the Lomo09 core was calculated from the 17 thickness of melt features observed during processing of the core (similar to Henderson et al., 18 2006). Clear and bubbly ice appears as transparent area when the core is backlit. If the melt 19 did not affect the whole core diameter, this was accounted for by multiplying the length of the 20 melt feature with the percentage of the core diameter it covered. If for example a melt feature 21 was 20 cm long but only affected one fifth of the core diameter, this melt feature would count 22 the same as a four centimetre long melt feature affecting the whole core diameter. The 23 observed melt features were then summed up per year to calculate the annual melt percent 24 (Fig. 3).

25

26 3 Results and discussion

The records of NO_3^- and NH_4^+ of the Lomo09 core both show the highest concentrations during the period of approximately 1940 to 1980 (Figs. 3 and 4), similar to findings from other Arctic sites (Goto-Azuma and Koerner, 2001). This clearly indicates a strong influence of anthropogenic emissions in recent decades on the chemical composition of aerosols reaching Lomonosovfonna. Both records show a significant decrease after 1980, a trend

similarly observed in the NO_3^- and NH_4^+ records of ice cores from the Siberian Altai (Eichler 1 et al., 2009, 2011) (Fig. 5) and Severnaya Zemlya (Opel et al., 2013; Weiler et al., 2005) 2 influenced mainly by Eurasian pollution. The Siberian Altai core ends in 2001, explaining 3 4 why the continuation of the decrease seen at Lomonosovfonna is not recorded there. The 5 nitrate trend in the Siberian Altai fits well with NO_x emissions as discussed in detail in Eichler et al. (2009) where the Eastern European NO_x emissions are shown. In contrast, NO_3^- 6 7 concentrations in records from Summit, Greenland, and Colle Gnifetti, Swiss Alps (see Fig. 1 8 for locations), affected by Northern American and Western European air masses, respectively, kept rising into the 21st century (Fig. 5). This suggests that the major sources for the increased 9 concentrations of NO_3^- and NH_4^+ in the Lomo09 core are similar to those for the Siberian 10 11 Altai and Severnaya Zemlya, whereas the influence of emissions in North America and 12 Europe is of minor importance. Our finding is in agreement with results from transport 13 modelling and trajectory analysis, identifying Eurasia as source region for Svalbard (Hirdman 14 et al., 2010a and b). Eurasian pollution dominates especially during the autumn and winter period characterised by Artic Haze episodes, in contrast to summer months when North 15 16 Atlantic air masses prevail (Tunved et al., 2013). We thus attribute the observed trend after 17 1940 in NO_3^- to higher NO_x emissions from traffic, energy production, and industrial activities, and in NH₄⁺ to enhanced NH₃ emissions from agriculture and livestock mainly in 18 19 Eurasia (Eichler et al., 2009; Weiler et al., 2005). This is in contrast to studies from Northern America and Western Europe, where anthropogenic pollution started already rising in the 20 second half of the 19th century due to an earlier industrialization and intensification of 21 agriculture (see e.g. Döscher et al., 1996). The anthropogenic impact is also seen in the NO_3^{-1} 22 and- less pronounced- in the NH₄⁺ record of the Lomo97 core (Divine et al., 2011; Kekonen 23 et al., 2005) (Fig. 4), which underlines the spatial representativeness of the Lomo09 ice core 24 data. The NO₃⁻ records of the Lomo09 and Lomo97 cores agree well. This is not the case for 25 the NH₄⁺ records, where the Lomo97 shows higher concentrations, especially before 1900 26 (Fig. 4). We cannot explain this difference, but NH_4^+ is known to be prone to contamination 27 during analysis (Jauhiainen et al., 1999; Kaufmann et al., 2010; Legrand et al., 1984, 1993, 28 1999; Udisti et al., 1994). Three observations indicate that the Lomo09 NH_4^+ concentrations 29 are robust: 1) The preindustrial Lomo09 values are generally lower than those of Lomo97 and 30 31 therefore contamination seems unlikely, 2) the Lomo09 preindustrial ion balance is close to zero (Fig. S2), and 3) the 300 year record of NH_4^+ from Holtedahlfonna (Beaudon et al., 32

2013) is in reasonable agreement with the Lomo09 data, except the period around 1750 (Fig.
 4).

In order to investigate sources of NH_4^+ and NO_3^- and other ionic species in the Lomo09 ice 3 4 core we performed a principal component analysis (PCA). We used 10-year-averages to 5 account for dating uncertainties and smoothing effects by melt-water relocation. Previous studies on the Lomo97 core suggested that the percolation lengths at the site can reach two to 6 7 eight annual layers in the warmest years (Moore et al., 2005; Pohjola et al., 2002). The 8 average annual melt percent of the Lomo97 core was 41% (Pohjola et al., 2002) compared to 9 31% of the Lomo09 core. We thus assume that the maximum percolation lengths in the Lomo09 core do not exceed the eight annual layers determined for the Lomo97 core. As a 10 conservative estimate we used the 10-year-average record of melt percent in the PCA to 11 examine the influence of melt on the NH_4^+ and NO_3^- records. The PCA was performed only 12 for pre-industrial times (1222-1859) to exclude anthropogenic influences on the ion 13 concentrations. Sulphate (SO_4^{2-}) from anthropogenic sources has been shown to increase 14 already during the second half of the 19th century (Moore et al., 2006). 15

16 We obtained six principal components (PCs) from the PCA (Table 2). PC1 has high loadings of sodium (Na⁺), potassium (K⁺), magnesium (Mg²⁺), and chloride (Cl⁻). This component 17 explains 38% of the total variance and contains species that are directly emitted by sea spray. 18 19 PC2 has high loadings of methane-sulfonate (MSA = CH_3SO_3) and NO_3 . MSA has a strictly 20 marine biogenic source. It results from the oxidation of gaseous dimethyl-sulphide (DMS) 21 which is produced by phytoplankton and emitted from the ocean to the atmosphere. This gas 22 release across the sea-air interface differs distinctly from the way sea salt species are emitted to the atmosphere via sea spray because no droplets are involved (Stefels et al., 2007; Vogt 23 24 and Liss, 2009). PC3 has a high loading of NH₄⁺, representing biogenic emissions. Calcium (Ca^{2+}) is the only species that has a high loading in PC4. This suggests that PC4 represents a 25 26 mineral dust component. The melt percent is the only parameter that has a high loading in PC5. Although we do not expect a simple correlation between melt in a particular layer and 27 28 net chemical flux, this result suggests that there is at least not a net loss of the investigated major ions from layers of extreme melt, which together with the findings of Pohjola et al. 29 30 (2002) and Moore et al. (2005) supports our conclusion that melt has not been of major influence on the ion concentrations on the considered decadal time scale.. PC6 has a high 31 loading of SO_4^{2-} , indicating a volcanic source. We can exclude a dominant sea spray and 32

marine biogenic influence on $SO_4^{2^-}$, due to the low loadings within PC1 and PC2, respectively. The results of the PCA are in good correspondence with those of a correlation analysis of the 10-year-averaged records of the ionic species and the melt percent for the preindustrial period (Table 3). Strong correlation is observed for the sea spray related ions Na⁺, K^+ , Mg²⁺, and Cl⁻ (0.59<r²<0.98). Furthermore, MSA and NO₃⁻ are highly correlated and share 60% of data variability. NH₄⁺, Ca²⁺, melt percent and SO₄²⁻ are not significantly correlated with any other species.

8 3.1 Nitrate and methane-sulfonate (NO₃⁻ and MSA)

9 In the Arctic MSA is well known to originate from marine biogenic sources (Legrand, 1997).
10 However, major pre-industrial NO₃⁻ sources in this region are still not fully understood (e.g.,
11 Wolff et al., 2008).

12 Varying atmospheric MSA concentrations have been related to changing sea ice conditions. Studies from Arctic and Antarctic ice cores found positive (Becagli et al., 2009; Legrand et 13 al., 1997), but also negative correlations of MSA and sea ice extent (Rhodes et al., 2009; 14 15 Sharma et al., 2012), see also the review on MSA and sea ice in Antarctica by Abram et al. (2013). After 1920 the Lomo97 core MSA correlates negatively with summer (August) sea-16 17 ice extent and sea surface temperature in the Barents Sea (O'Dwyer et al., 2000) and 18 positively with the instrumental summer temperature record from Svalbard (Isaksson et al., 19 2005). During the period 1600-1920 Isaksson et al. (2005) detected a positive correlation of 20 the Lomo97 MSA and winter (April) sea ice extent in the Barents Sea (Divine and Dick, 2006; Vinje, 2001). The Lomo97 MSA record reveals a pattern with twice as high values 21 prior to about 1920 compared to those of the later 20th century (Isaksson et al., 2005). They 22 23 suggest that it results from a change of source and/or more favourable growing conditions for 24 the DMS-producing phytoplankton in a more extensive sea ice environment before 1920.

In the MSA record of the Lomo09 core we find a similar pattern as in the Lomo97 core with higher concentrations prior to the 20th century and a decreasing trend since around 1900 (Figs. 4 and 6). Hence, we investigate if a coupling of MSA with sea ice conditions around Svalbard exists, using three long-term reconstructions of sea ice extent. These reconstructions include the winter (April) ice extent in the Western Nordic Seas covering the last 800 years (Macias Fauria et al., 2010), the summer (August) location of the sea ice edge in the Barents Sea (BS) that covers the last 400 years (Kinnard et al., 2011), and the summer sea ice extent in the

Arctic Seas extending back to the year 563 (Kinnard et al., 2011). The best agreement was 1 2 observed between the 40-year-lowpass-filtered records of Lomo09 MSA and reconstructed winter (April) Western Nordic Seas ice extent (Macias Fauria et al., 2010). However, less 3 4 than 50% of the variability in the MSA record is explained with changing sea ice conditions 5 (Fig. 6; r=0.56, p<0.001). The most striking feature in both records is the pronounced decrease starting around 1890. This remarkable drop was also observed in the time series of 6 7 the Western Nordic Sea ice extent (period 1860-2000; Vinje et al., 2001), used to calibrate the 8 proxy records (Macias Fauria et al., 2010). The strong decrease is not seen in any of the 9 summer (August) ice records before around 1910 (Fig. 6). Furthermore, the pronounced 10 minimum around 1710 and the peak around 1640 in the BS ice record are not reflected in the 11 Lomo09 MSA record. Thus, our data do not support the connection of MSA at 12 Lomonosovfonna and the BS ice extent stated in O'Dwyer et al. (2000) for the period 1920-13 1997, nor the assumption of Isaksson et al. (2005) that the MSA sources prior and after 1920 14 were the same, i.e. the BS. We explain the positive correlation of Lomo09 MSA and Western Nordic Sea ice extent as follows. The marginal ice zone is known to be the area of highest 15 16 DMS production (Perrette et al., 2011). The larger the sea ice area, the more ice edge area is 17 available for phytoplankton growth and thus DMS production. Furthermore, more ice leads to higher freshwater inflow by melting ice. This results in a stronger stratification of the ocean 18 19 water (Perrette et al., 2011) which keeps the phytoplankton in the euphotic zone. The good 20 correspondence of the Lomo09 MSA record with the Western Nordic Sea ice extent but not 21 with that of the BS is well supported by the findings of Beaudon et al. (2013) pointing to the 22 Greenland Sea as the main source for biogenic related MSA in Svalbard.

In the Lomo09 ice core MSA is highly correlated with NO_3^- before around 1900. The records (Figs. 4 and 6) are similar with shared peaks around 1395, 1475, 1560, 1645, 1695, and 1795. The only exception is the period at about 1350 when the correlation breaks down. After around 1900 there is a decoupling of both species with enhanced NO_3^- concentrations from anthropogenic Eurasian NO_x emissions (see above) and strongly decreased MSA concentrations.

In previous studies NO_3^- was found to correlate with non-sea-salt- Ca^{2+} (nss- Ca^{2+}) (Legrand et al., 1999; Röthlisberger et al., 2000, 2002), suggesting that nss- Ca^{2+} prevents NO_3^- from being re-emitted from the snowpack. However, those studies are from Greenland, consider glacial timescales, and include e.g. the last glacial maximum (LGM) with much higher nss- Ca^{2+}

concentrations. Other studies observed a correlation of NO_3^- and Ca^{2+} in summer and with sea 1 2 salt in winter but they considered only industrial times (Beine et al., 2003; Geng et al., 2010; 3 Teinilä et al., 2003). The empirical orthogonal function (EOF) analysis performed on the ion data of the Lomo97 core suggests in general no correlation between Ca^{2+} and NO_3^{-} , but in 4 some parts of the last 200 years the two species are clearly associated (Kekonen et al., 2002). 5 Kekonen et al. (2002) found NO_3^- and NH_4^+ to covariate during the last 100 years. However, 6 the EOF of the whole core did not show a clear association of NH_4^+ and NO_3^- . Nevertheless, 7 8 they suggested that before 1920 and after 1960 ammonium nitrate (NH₄NO₃) has been 9 common at Lomonosovfonna. They explain this in recent years to be due to Arctic Haze and significant natural sources of NH₄NO₃ during the earlier period. At Holtedahlfonna, Svalbard, 10 NH_4^+ was also associated with NO_3^- before 1880 which Beaudon et al. (2013) interpreted as 11 evidence for NH_4NO_3 to be present. Teinilä et al. (2003) also discovered a correlation of NO_3^- 12 13 and NH_4^+ in recent times which they concluded to result from anthropogenic emissions. Our data neither support a correlation of NO₃⁻ and Ca²⁺, nor of NO₃⁻ and the sea salt species Na⁺, 14 15 nor of NO₃⁻ and NH₄⁺ in pre-industrial times. Instead, they clearly suggest an association of 16 NO_3^- with MSA. Three hypotheses for the high correlation are discussed: (1) post-17 depositional processes caused by melt water percolation affecting NO₃⁻ and MSA in the same way, (2) a common source of NO_3^- and MSA, and (3) NO_3^- fertilisation of the ocean which 18 19 triggers phytoplankton growth and thus DMS and MSA formation.

- 1. The pre-industrial record of the melt percent does share some features with NO_3^- and MSA but there is no significant correlation with NO_3^- or MSA ($r^2=0.1$ with either NO_3^- or MSA) (Table 3, Fig. 4). This is also seen in the PCA where the melt percent and the two ionic species have their highest loadings in different PCs (Table 2). Thus, the correlation of NO_3^- and MSA is not a result of similar relocation during melt events on the decadal time scales considered here.
- 26 2. If both species have a common source this would have to be the ocean because MSA results only from marine DMS production and its oxidation in the atmosphere. NO_3^{-1} is 27 28 only a minor component in sea water with concentrations in the micro-molar range 29 (Chester and Jickells, 2012; Codispoti et al., 2013). The ice core NO₃⁻/Na⁺ ratio of ~0.066 in the Lomo09 core is up to a factor of ten higher than the sea water ratio of 30 0.006 to 0.038 (Keene et al., 1986). Additionally, we can exclude NO_3^{-1} to be derived 31 from sea spray because NO_3^- and the major sea spray components Na^+ , K^+ , Mg^{2+} , and 32 33 Cl⁻ (PC1) do not correlate as seen in the PCA and the correlation analysis (Tables 2

1 and 3). Because of the high solubility of nitrate in water, outgassing of HNO_3 from the 2 ocean is unlikely. Thus, the major NO_3^- source is not the ocean which excludes a 3 common source to cause the strong correlation of NO_3^- and MSA.

3. Elevated atmospheric NO_3^- concentrations due to high NO_x emissions and/or 4 5 enhanced transport to the Arctic in the end of winter lead to an increased amount of NO_3^- dissolved in the ocean surface water. Nutrient supply in the Arctic is known to 6 7 be limited and nitrate depletion is common during the vegetative season (Codispoti et 8 al., 2013). Hence, an increased nitrogen input by dissolved NO₃⁻ leads to a fertilisation 9 of the phytoplankton (Duce et al., 2008). As soon as light becomes available this 10 results in an enhanced production of DMS and finally higher MSA concentrations in 11 the atmosphere. This process takes weeks to months (Codispoti et al., 2013; Sharma et 12 al., 2012). However, such a potential short time lag cannot be resolved from our data.

We therefore propose the fertilising effect to be the dominant cause for the high correlation of NO₃⁻ and MSA in pre-industrial times. In industrial times the records of NO₃⁻ and MSA diverge with increasing NO₃⁻ and decreasing MSA concentrations. This reveals that during the 20th century the effect of decreasing MSA concentrations following reduction in ice extent in the Western Nordic Seas predominates compared to an expected MSA increase caused by enhanced anthropogenic NO₃⁻ levels.

19 The major NO₃⁻ source region for the industrial time is Eurasia indicated by the similarity of 20 the NO₃⁻ records observed in the last 30-40 years in the ice cores from Lomo09, the Siberian Altai, and Severnaya Zemlya (Eichler et al., 2009; Weiler et al., 2005) (Fig. 5). We assume 21 22 that the source region has not changed from pre-industrial to industrial times. In the period 23 1250-1940 NO₃⁻ in the Siberian Altai ice core was ascribed to forest fires and mineral dust as 24 main pre-industrial sources (Eichler et al., 2011). That NO₃⁻ record shows a maximum between 1540 and 1680 (see Fig. 5), attributed to an increased mineral dust input from 25 26 Central Asian deserts (1540-1600) and enhanced fire activity from Siberian boreal forests (1600-1680). This distinct peak in the 16th and 17th century is not observed in the Lomo09 27 NO₃⁻ record and also the general pre-industrial records do not correspond well. We cannot 28 exclude that other regional scale NO_3^- sources in Eurasia had a significant impact on the low 29 30 pre-industrial concentration level. From our data we can therefore not identify major pre-31 industrial NO_3^- sources for the Lomo09 core.

1 3.2 Ammonium (NH₄⁺)

The Lomo09 NH_4^+ record shows very low concentrations between the 13th and 18th century 2 and an increasing trend from around 1750 onwards (Fig. 7). The values are on the same order 3 4 of magnitude as those from other Arctic sites and factor 2-3 lower compared to the Lomo97 5 ice core (Beaudon et al., 2013; Fuhrer et al., 1996; Kehrwald et al., 2012; Kekonen et al., 6 2005; Legrand and De Angelis, 1996; Legrand et al., 1992; Whitlow et al., 1994; Zennaro et 7 al., 2014). The NH₄⁺ record of another Svalbard core from Holtedahlfonna, spanning the last 300 years, shows a differing pattern prior to 1800, but a similar strong increasing trend as the 8 Lomo09 record from the 18th century on (Beaudon et al., 2013) (Fig. 4). The authors interpret 9 the rising concentrations from 1880 as result of anthropogenic mid-latitude pollution reaching 10 the Arctic. However, the earlier increase in NH₄⁺ concentrations in the Lomo09 and Holte05 11 ice core from the 18th century on cannot be related to anthropogenic emissions. As discussed 12 above, anthropogenic NH₃ emissions from Eurasia influence precipitation chemistry in 13 Svalbard only after around 1940. 14

Pre-industrial NH_4^+ was not studied in details in the Lomo97 core but Kekonen et al. (2002) 15 16 suggested NH₄NO₃ to have been common at Lomonosovfonna before 1920. Similarly, Beaudon et al. (2013) postulated that at Holtedahlfonna natural NH₄NO₃ was a common 17 aerosol. Our data do not support this hypothesis since NH_4^+ and NO_3^- are not significantly 18 correlated in pre-industrial times (Tables 2 and 3). In other studies pre-industrial NH_4^+ was 19 20 attributed mainly to biomass burning (e.g., Fuhrer et al., 1996; Kehrwald et al., 2010; Legrand 21 et al., 1992; Whitlow et al., 1994). North America and Canada were identified as major 22 sources for NH₄⁺ in Greenland ice (Fuhrer et al., 1996), whereas Legrand and De Angelis 23 (1996) and Zennaro et al. (2014) suggest an additional Eurasian source. A period of 24 exceptional high fire activity around 1600-1680 in Siberian boreal forests of Eurasia was 25 detected in the ice core fire tracer records from the Siberian Altai and Greenland (Eichler et al., 2011; Zennaro et al., 2014). This unique period did not lead to a maximum in the Lomo09 26 NH_4^+ record. Therefore, we conclude that biomass burning is not a major source for NH_4^+ 27 arriving at Svalbard. 28

The trend in the Lomo09 NH_4^+ record is similar to that in the ice core from Belukha glacier in the Siberian Altai with increasing concentrations already from around 1750 and very low concentrations between 1680 and 1750 (Eichler et al., 2009) (Fig. 7). Before ~1500 NH_4^+ concentration records agree less, which could be explained by increasing dating uncertainty.

At the Belukha site long-term NH_4^+ variations were related to temperature-induced changes of 1 biogenic NH₃ emissions from extended Siberian boreal forests (Eichler et al., 2009). The 2 strong increase after the 18th century was caused by a rise of Siberian temperatures since that 3 4 time. Hence, from the similarity in the Lomo09 and Siberian Altai NH₄⁺ concentration records we conclude that biogenic NH₃ emissions from Siberian boreal forests are the dominant 5 source for NH_4^+ at Lomonosovfonna. Due to the larger distance to the emission sources the 6 NH_4^+ concentrations in the Lomo09 core are about one order of magnitude lower than in the 7 core from Belukha glacier. The NH₄⁺ concentrations in a Greenland ice core (NEEM, for 8 location see Fig. 1) do not show the increase after the 18th century (Zennaro et al., 2014) (Fig. 9 7), implying that biogenic emission trends in Northern America and Eurasia differ. 10

11

12 **4** Summary

We presented the 800 year records of the two nitrogen species NO_3^- and NH_4^+ analysed in a 13 new ice core collected from Lomonosovfonna, Svalbard, in 2009. In general, the NO₃⁻ record 14 of the 2009 ice core agrees reasonably with published data from two previous Svalbard ice 15 cores, Lomonosovfonna 1997 (Kekonen et al., 2005) and Holtedahlfonna 2005 (Beaudon et 16 al., 2013), whereas there is less correspondence between the different NH_4^+ records. On the 17 decadal time scale considered here melt related effects did not significantly alter the 18 19 concentrations of the nitrogen compounds. Both species show a clear impact of anthropogenic pollution in the 20th century, with peak concentrations in the 1970s/1980s. This temporal 20 21 trend points to source regions in Eurasia and the Siberian Arctic, since emissions in Northern America and Western Europe kept rising into the 21st century. In pre-industrial times, i.e. 22 prior to the 20th century, the dominant source of NH₄⁺ was biogenic NH₃ emissions from 23 Siberian boreal forests. During the same period NO_3^- was highly correlated to MSA on a 24 25 decadal time scale. We explained this by a fertilising mechanism where higher atmospheric NO₃⁻ concentrations yield higher nitrogen input to the ocean, triggering the growth of DMS-26 27 producing phytoplankton. Elevated DMS concentrations then result in enhanced concentrations of MSA in the atmosphere. Based on our data it was not possible to resolve 28 major pre-industrial NO₃⁻ sources for Svalbard. 29

30

31 Supplementary material

Here, an example for the annual layer counting (ALC) for the core section between 0 and
20 m weq (Fig. S1) is displayed, followed by the raw data of the ionic species and the ion
balance along depth in m weq (Fig. S2) and the annual melt percent (Fig. S3) of the Lomo09
ice core versus age.

5

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- 1 Table 1 Detection limits and median values $[\mu eq/L]$ for the ions analysed with the
- 2 Metrohm 850 Professional IC. Pre-ind. = pre-industrial time from 1222-1859;

| Anions | | | | Cations | ions | | | | |
|------------------------------|--------------------|----------|------|-------------------|-----------|----------|------|--|--|
| | Detection limit | Median | | | Detection | Median | | | |
| | | Pre-ind. | Ind. | - | limit | Pre-ind. | Ind. | | |
| MSA | 0.005 | 0.09 | 0.05 | Na ⁺ | 0.02 | 8.77 | 7.18 | | |
| Cl. | 0.02 | 10.48 | 8.92 | $\mathbf{NH_4}^+$ | 0.02 | 0.50 | 0.74 | | |
| NO ₃ ⁻ | 0.01 | 0.54 | 0.65 | \mathbf{K}^{+} | 0.02 | 0.25 | 0.19 | | |
| SO4 ²⁻ | 0.02 | 2.08 | 2.63 | Mg^{2+} | 0.03 | 2.10 | 1.32 | | |
| | | | | Ca ²⁺ | 0.04 | 1.43 | 1.02 | | |
| | | | | | | | | | |

3 Ind. = industrial time from 1860-2009; $MSA = CH_3SO_3^-$.

4

1 Table 2 Results of the principal component analysis (PCA) after VARIMAX rotation. Time

2 period: 1222-1859; data: 10-year averages; $MSA = CH_3SO_3$; melt% = melt percent.

3 Values >0.8 marked in bold.

| | PC1 | PC2 | PC3 | PC4 | PC5 | PC6 |
|--------------------------------------|-------|------|-------|-------|-------|------|
| Na ⁺ | 0.97 | 0.06 | 0.05 | 0.11 | -0.03 | 0.08 |
| \mathbf{K}^+ | 0.88 | 0.18 | 0.00 | -0.04 | -0.07 | 0.16 |
| Mg^{2+} | 0.82 | 0.37 | 0.02 | 0.27 | 0.07 | 0.19 |
| СГ | 0.97 | 0.08 | 0.06 | 0.12 | 0.01 | 0.08 |
| MSA | 0.33 | 0.80 | 0.13 | 0.22 | 0.23 | 0.11 |
| NO ₃ | 0.11 | 0.89 | 0.22 | 0.16 | 0.09 | 0.22 |
| NH_4^+ | 0.06 | 0.23 | 0.96 | -0.02 | 0.17 | 0.02 |
| Ca ²⁺ | 0.18 | 0.27 | -0.02 | 0.92 | 0.07 | 0.19 |
| Melt% | -0.05 | 0.19 | 0.16 | 0.07 | 0.96 | 0.07 |
| SO ₄ ²⁻ | 0.29 | 0.28 | 0.02 | 0.21 | 0.08 | 0.88 |
| Variance explained [%] | 38 | 19 | 11 | 11 | 11 | 10 |

4

| 1 Table 3 R^2 values of the correlation | ion analysis of the | ionic species and | the melt percent |
|---|---------------------|-------------------|------------------|
|---|---------------------|-------------------|------------------|

2 (Melt%). Time period: 1222-1859; data: 10-year averages; $MSA = CH_3SO_3^-$; $0.5 < r^2 < 1$

| r^2 | Na^+ | \mathbf{K}^{+} | Mg^{2+} | Cl | MSA | NO ₃ | $\mathbf{NH_4}^+$ | Ca ²⁺ | Melt% | SO ₄ ²⁻ |
|--------------------------------------|--------|------------------|--------------------|------|------|-----------------|-------------------|------------------|-------|--------------------------------------|
| Na ⁺ | 1 | | | | | | | | | |
| \mathbf{K}^{+} | 0.71 | 1 | | | | | | | | |
| Mg^{2+} | 0.71 | 0.59 | 1 | | | | | | | |
| Cl | 0.98 | 0.67 | 0.78 | 1 | | | | | | |
| MSA | 0.17 | 0.16 | 0.41 | 0.20 | 1 | | | | | |
| NO ₃ - | 0.04 | 0.08 | 0.27 | 0.06 | 0.60 | 1 | | | | |
| $\mathbf{NH_4}^+$ | 0.01 | 0.01 | 0.03 | 0.02 | 0.14 | 0.19 | 1 | | | |
| Ca ²⁺ | 0.09 | 0.06 | 0.27 | 0.10 | 0.26 | 0.21 | 0.00 | 1 | | |
| Melt% | 0.00 | 0.00 | 0.02 | 0.00 | 0.15 | 0.11 | 0.13 | 0.04 | 1 | |
| SO ₄ ²⁻ | 0.16 | 0.18 | 0.33 | 0.17 | 0.26 | 0.26 | 0.02 | 0.24 | 0.04 | 1 |



2 Figure 1 (A) Map of Svalbard with the locations of Lomonosovfonna and Holtedahlfonna. (B)

- 3 Map with all ice core locations discussed in the text: Lomo = Lomonosovfonna (red triangle);
- 4 NEEM, Sum = Summit, SZ = Severnaya Zemlya, Bel = Belukha, and CG = Colle Gnifetti
- 5 (black triangles). Satellite image in (B) © PlanetObserver, extracted from DVD-ROM "Der
- 6 Große 3D-Globus 4.0 Premium,"#2008 United Soft Media Verlag GmbH, Munich.
- 7



2 Figure 2 Depth-age relationship of the Lomo09 ice core showing all dating methods applied.

- 3 Depth is given in m weq to account for density variation.
- 4



and annual melt percent of the Lomo09 ice core.



Figure 4 Records of NH_4^+ (green), NO_3^- (red), melt percent (light blue), Na^+ (yellow), and MSA (dark blue) of the Lomo09 ice core. Bold lines are 40-year-lowpass-filtered (40-yr-LP); dashed lines are 10-year averages (10-yr-avg). Raw data are available in the supplementary material (Figs. S2 and S3). Grey dashed lines are 40-year-lowpass-filtered records of NH_4^+ , NO_3^- , Na^+ , and MSA of the Lomo97 ice core (Kekonen et al., 2005) calculated with the updated chronology of Divine et al. (2011). Black dashed lines are 40-year-lowpass-filtered records of NH_4^+ and NO_3^- of the Holte05 ice core (Beaudon et al., 2013).



Figure 5 NO₃⁻ records from Lomo09 (red), Belukha (dark red; Eichler et al., 2009), Summit,
Greenland (grey; Geng et al., 2014), and Colle Gnifetti, Swiss Alps (black; Sigl, 2009). Bold
lines are 100-year-lowpass-filtered (100-yr-LP); dashed lines are 10-year averages (10-yravg).



Figure 6 Records of Lomo09 MSA (dark blue), pre-industrial NO₃⁻ (red), Western Nordic
Seas winter (April) ice extent (turquoise; Macias Fauria et al., 2010), Arctic summer (August)
sea ice extent (light purple; Kinnard et al., 2011), and August sea ice edge position in the
Barents Sea (BS; dark purple; Kinnard et al., 2011). Bold lines are 100-year-lowpass-filtered
(100-yr-LP); dashed lines are 40-year-lowpass-filtered (40-yr-LP).



Figure 7 NH₄⁺ records of the Lomo09 (green), Belukha (light green; Eichler et al., 2009), and
the NEEM (grey; Zennaro et al., 2014) ice cores. Bold lines are 100-year-lowpass-filtered
(100-yr-LP); dashed lines are the 10-year-averages (10-yr-avg).