RESEARCH ARTICLE

Post-depositional loss of nitrate and chloride in Antarctic snow by photolysis and sublimation: a field investigation

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Abstract

Nitrate in snow is subject to post-depositional processing, which leads to a net loss and redistribution within the snowpack. The relative importance of post-depositional loss processes such as the volatilization of nitric acid (HNO₃) and photolysis of nitrate has long been debated. Changes in nitrate and chloride concentrations in the snowpack were investigated at H128 (69°23.584′S, 41°33.712′E), an Antarctic coastal site approximately 100 km from Syowa Station in East Antarctica from December 2015 to February 2016. Results indicate that chloride migrated to deeper sites within the snowpack under the influence of water vapour movement. Moreover, 50% of the nitrate on surface snow was lost to photolysis, and approximately 20% of the nitrate was absent at a depth of 40 cm. To enhance our knowledge of the Antarctic geochemical cycle, this study is the first to suggest chloride ion movement in snowpacks or significant nitrate loss for any Antarctic coastal site.

Introduction

Nitrate in snow is subject to post-depositional processing, which leads to net loss and redistribution within the snowpack (Röthlisberger et al. 2000; Jacobi & Hilker 2007; Frey et al. 2009). The relative importance of the volatilization of nitric acid (HNO₃) and the photolysis of nitrate in the post-depositional loss process have long been debated (Röthlisberger et al. 2000; Blunier et al. 2005). Nitrate photolysis in/on the snowpack is a source of reactive nitrogen oxides (NO₃⁻) in the atmosphere (Honrath et al. 1999; Jones et al. 2001; Shi et al. 2018). The photochemistry of nitrate in the snowpack has significant implications because its photoproducts—NO, NO₂, and radicals—are intimately linked to reactions involving ozone, hydrocarbons and halogens.

Nitrate undergoes photolysis in the quasi-liquid layer of ice and either NO₂⁻ (Eqn. 1) or NO₃⁻ (Eqn. 2) is produced (Mack & Bolton 1999; Grannas et al. 2007; Meusinger et al. 2014).

\[
\text{NO}_3^- + h\nu + H^+ \rightarrow NO_2^- + OH (1)
\]

\[
\text{NO}_3^- + h\nu \rightarrow NO_2^- + O(^3P) \quad (2)
\]

Nitrate in ice cores is a potential proxy for historic atmospheric NO₃ levels (Dibb et al. 1998). Solar variability on centennial to millennial time scales may imprint on the long-term NO₃ record, as suggested by Traversi et al. (2012) in their work on the Talos Dome core record. They reported that traces of solar variability are preserved as nitrate in ice cores drilled in high-accumulation sites (Traversi et al. 2012). However, nitrate concentration in ice cores, which Traversi et al. used as a tracer of solar variability, would have been reduced by the photochemical reactions of nitrate (Noro et al. 2018). Hence, investigation of the nitrate loss in snowpacks at high-accumulation sites is important. In addition, the volatilization of weak acid from the ice surface was reported by Sato et al. (2008). They determined that HNO₃ in snow was not volatilized, and hydrochloric acid was volatilized slowly (Sato et al. 2008). It is possible that if snow is investigated for longer than in the study by Sato et al. (2008), HNO₃ and more hydrochloric acid may be volatilized. Recently, Shi et al. (2019) reported on their observations of nitrate in an Antarctic snowpack that had undergone isotopic
fractionation due to the volatilization process. They also reported that the extent of nitrate evaporation was very small at temperatures lower than −24 °C; however, significant evaporation possibly occurred at −4°C for a few weeks. Therefore, both the photolysis and the evaporation of nitrate have been observed in snowpack extracted from Antarctica.

With the aim of (1) quantifying the chloride and nitrate post-depositional loss in snowpack from Antarctica and (2) determining the snow depth at which nitrate loss occurs, we carried out controlled experiments at an Antarctic coastal site.

Materials and methods

Location

Post-depositional process experiments were conducted at H128 (69°23.584’S, 41°33.712’E), approximately 100 km from Syowa Station in eastern Dronning Maud Land, East Antarctica, during JARE57, from December 2015 to February 2016.

Nitrate volatilization from the ice surface in the laboratory experiment

The pH values of 100 cm³ of the sample solutions, including 100 μmol dm⁻³ of NaNO₃, were adjusted using H₂SO₄ at a pH of 2–5 and then placed in Petri dishes with inner diameters of 95 mm (approximately 1.4 cm height) for analysis. The dishes were loosely covered with aluminium foil and kept in a freezer at −20 and −60 °C for three weeks. Afterwards, the samples were melted at 25 °C for ion chromatography analysis.

Application of quartation for the mixed snow

A quartation method was employed for sample homogenization. A polyethylene sheet cleaned with ultra-pure water (resistivity ≥18.2 MΩ cm) was placed on a metal frame (90 × 90 × 90 cm³) to make a container. The container was then half-filled with surface snow, which was smoothed and divided into four parts. Unmixed blank samples were collected from the centres of the whole sample and the divided parts (a total of five samples). Subsequently, the divided samples were mixed diagonally or orthogonally three times (mixed sample), as shown in Fig. 1. This mixing process is referred to as “quartation” in soil science. The mixed sample was then again divided into four parts to collect mixed blank samples from the centres of the whole sample and the divided parts. The chemical concentrations before and after the homogenization conducted in Antarctica and Japan are shown in Supplementary Table S1.

Natural sample preparation and field sampling

The sample mixed by the quartation method was packed into Teflon-coated glass tubes with inner diameters of 5 cm and lengths of 40 cm, as shown in Fig. 2. The glass tubes were then buried in snowpacks under three varying conditions for four weeks. The first condition is the most natural condition: sample tubes were simply buried in a snowpack (OPEN condition). In the second condition, glass beakers were placed on top of the glass tubes to prevent deposition on the sample (COVERED condition), with a gap left to allow air to pass through. Under the third condition, aluminium tape was wrapped around the glass tubes to shield the sunlight, and glass beakers

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Fig. 1 Quartation mixing step. Black dots show sampling points.

Fig. 2 The Pyrex glass tubes used for the investigation. The inner walls of the tubes were treated with PTFE. The bottoms were covered with mesh to allow air to pass through and to prevent snow from moving down. (a) In the OPEN experimental condition, the top was open and directly in contact with air. (b) In the COVERED condition, the top was covered with a Pyrex beaker and three PTFE spacers allowed air to pass through and prevented the deposition of new snow and particles. (c) The DARK condition was the same as the COVERED condition, except that the tube and beaker were covered with an aluminium sheet to prevent sunlight irradiation.
wrapped with aluminium tape were placed on the glass tubes with gaps to allow for airflow (DARK condition). Every week, three glass tubes were collected from each of the three scenarios (OPEN, COVERED and DARK), and the snow cores from the collected tubes were divided into 5-cm sections. Before and after sampling, the samples in the glass tubes were weighed to calculate the sample density (ca. 0.50 g cm\(^{-3}\)). The cut samples were stored in clean bags in a frozen state prior to laboratory analysis in Japan.

**Amended sample preparation and field sampling**

The containers were set up, and natural snow samples were prepared. Thereafter, 100 cm\(^3\) of NaCl, NH\(_4\)Cl and NaNO\(_3\) (100 mmol dm\(^{-3}\) each) solution was sprayed onto the snow sample. Subsequent mixing, glass tube packing and sampling procedures were conducted as for the natural snow sample (amended sample).

**Observation of sunlight intensity in the snowpack**

Light intensity in the snowpack was observed with a 1.3 m depth pit. An optical fibre spectrophotometer system with a UV/Vis probe (Ocean Optics TP300-UV/VIS probe and USB2000+UV-VIS-ES) was used for the observation. The tip of the probe in a right-angle prism was inserted up to approximately 90 cm from the side of the pit wall at depths of 10, 20, 30, 50, 75 and 100 cm (Fig. 3). The probe was rotated to 0° (upward in vertical direction), 45°, 90° (side), 135° and 180° (downward in vertical direction) to measure the light intensity in the snow pit from these selected directions. In addition, the reference was measured by placing the probe, which was set to 30 cm above the reflector (SRT-99-050, Labsphere), at a fixed height of 150 cm.

We calculated the e-folding depth of actinic flux in the snowpack according to the Beer–Bouguer–Lambert Law (Eqn. 3; France et al. 2011). The e-folding depth was defined as the depth where the actinic flux would be 1/e times (ca. 37%; France et al. 2011).

\[
I_z = I_{z'} \times \exp(e^{-1} \times (z - z')), \tag{3}
\]

where \(z\) (cm) is the depth, \(z'\) (cm) is the initial depth \((z > z')\), \(I_z\) is the light intensity (a.u.) at depth \(z\), \(I_{z'}\) (a.u.) is the light intensity at depth \(z'\) and \(e\) (cm) is the e-folding depth. The average of solar intensities at 90° and 270° was used to calculate \(e\), in the same manner as in previous reports (France et al. 2011). Additionally, the solar intensity in the snowpack was normalized by the sunlight intensity measured by an automatic weather station.

**Temperature observation of the snow**

To determine the temperature of the snow samples, type-k thermocouples were connected to a portable temperature and humidity data logger (TR-72Ui-type Thermo Recorder, T&D Corporation). The thermocouples were inserted into the snowpack at 0, 10, 20, 30 and 40 cm depths to measure the temperature gradient in the snowpack continuously (Fig. 4). After setting the thermocouples, the snow was returned to the pit and the temperature at each depth was measured continuously.

**Ion chromatography analysis**

Ion concentrations in the samples were analysed by ion chromatography in the laboratory. Amended samples were diluted 20 times with ultra-pure water. The anion chromatograph system (883 basic IC plus, Metrohm) consisted of a guard column (SI-90G, Shodex), separation column (SI-90 4E, Shodex) and suppressor for anion analysis, and the eluent was a mixed solution of 1.8 mmol dm\(^{-3}\) Na\(_2\)CO\(_3\) and 1.7 mmol dm\(^{-3}\) NaHCO\(_3\), with a flow rate at 1 cm\(^3\) min\(^{-1}\). The cation

![Fig. 3 Schematic diagram of how light intensity was observed in the snowpack: (a) overall picture; (b) enlarged illustration of the light-receiving unit.](image-url)
chromatograph system (IC7000, Yokogawa) consisted of a separation column (YS-50, Shodex) and an eluent made up of 4 mmol dm$^{-3}$ methanesulfonic acid at 1 cm$^3$ min$^{-1}$ flow rate.

Results

Nitrate volatilization from the ice surface

Figure 5 shows the decrease in the nitrate concentration from the ice surface during the three-week study period at temperatures of −20 and −60 °C. Because the apparent ion concentration changes with the evaporation of water, nitrate concentrations were standardized by [Na$^+$]. At −60 °C, the nitrate concentration decreased by approximately 3 μmol dm$^{-3}$ at a pH of between 2 and 5. At −60 °C, there is no liquid phase on the ice; therefore, nitrate ions cannot bond to protons, restricting nitrate volatilization. Therefore, we did not observe a pH dependence of nitrogen volatilization in our experiments at −60 °C. At −20 °C, nitrate volatilization showed a peak at a pH of 3. The volatilization of hydrochloric acid has already been reported (Sato et al. 2008) and tends to proceed at lower pH levels. Due to the effects of the freezing concentration effect, the volatilization of acid from the ice surface occurred by combining the concentrated H$^+$ ions with the counter anions. However, at a pH of 2, the high concentration of sulfuric acid (5 mM) inhibited the freeze concentration effect, decreasing nitrate volatilization. This suggests that the volatilization of nitrate as a post-depositional process may occur in environments like Antarctica where nitrate is present on the ice surface for long periods.

Homogenization of snow sample by quartation

Supplementary Table S1 shows the ion concentrations and relative SDs of the unmixed and mixed snow samples. Except for the chloride concentration of one sample collected at H128, variations in all the samples reduced after quartation. The unchanged chloride concentration in the H128 sample may be due to limitations in the accuracy of the ion chromatograph rather than ineffectiveness of the quartation process, which produced good results for snow equalization.
Temperature in the snowpack

Supplementary Fig. S1a shows the temperature change in the snowpack at the depths of 0, 10, 20, 30 and 40 cm, as well as the atmospheric temperature measured by an automatic weather station on 2–22 January 2016. Supplementary Fig. S1b shows the atmospheric temperatures recorded on 4 January 2016. The surface temperature fluctuated significantly between 3.9 and −22.8 °C on the account of sunlight heating and radiative cooling. Note that temperatures above 0 °C were caused by the direct solar heating of the sensor; snowmelt was not observed during these experiments. The maximum and minimum daily temperatures were observed at approximately 13:30 and 01:30, respectively. Deeper in the snowpack, the amplitude of the temperature variations decreased. In addition, owing to the adiabatic effect of snow, the time of the temperature peaks at the deeper sites gradually lagged behind that of the surface temperature peaks. The width shift and lag of temperature peaks should change the thermal gradient in the snowpack, controlling the direction of the water vapour movement in a complex manner.

Change in concentrations of ions in natural snow in Antarctica

Concentrations of Na⁺, Cl⁻, NO₃⁻ and SO₄²⁻ in the unmixed blank samples were 8.4 ± 4.3, 5.9 ± 0.2,
1.7 ± 0.5 and 2.0 ± 0.6 μmol dm⁻³, respectively (n = 5, error = 1 SD), while those in mixed blank samples were 4.2 ± 0.9, 5.6 ± 0.3, 1.0 ± 0.1 and 1.3 ± 0.04 μmol dm⁻³, respectively (n = 5, error = 1 SD; Supplementary Table S1). Figure 6 shows the results of these samples. As the apparent ion concentrations change with the evaporation of water, we could not determine whether changes observed in the ion concentrations of the natural samples were due to water vapour movement or increases and decreases in the absolute concentrations of the ions. Therefore, the concentrations of Cl⁻ and NO₃⁻ were compared with that of Na⁺, for which the absolute amount did not change. Note that dry deposition of Na⁺ could increase the Na⁺ concentration instead of water evaporation. However, the Na⁺ concentration increment was observed under both OPEN and COVERED conditions. We therefore concluded that the reason of the Na⁺ concentration change was the water evaporation. The stars in Fig. 6 show the [Cl⁻]/[Na⁺] and [NO₃⁻]/[Na⁺] of the mixed blank sample. If the ion concentration change was due to water vapour movement only, all the plots would be on the straight lines, as shown in the figures. Additionally, if a plot occurs on the right side of this line, it indicates that the concentration of Cl⁻ or NO₃⁻ decreased after the post-depositional processes occurred. Conversely, if the plot is on the left side of this line, the concentration of Cl⁻ or NO₃⁻ increased after the occurrence of post-depositional processes or because of deposition from the atmosphere. Under the conditions of OPEN and COVERED, Cl⁻ was lost in the surface snow (0–5 and 5–10 cm depth). We observed a significant difference between the [Cl⁻]/[Na⁺] of the surface snow sample (0–5 and 5–10 cm depth; p = 0.05). However, the [Cl⁻]/[Na⁺] under the DARK condition was stable. Similar to the Cl⁻ results, NO₃⁻ was lost in the surface snow (0–5 and 5–10 cm depth) under the OPEN and COVERED conditions. There was also a significant difference between the [NO₃⁻]/[Na⁺] of the surface snow sample (0–5 and 5–10 cm depth; p = 0.05). Because most of the data plotted on the right side of the blank under all three conditions, we can conclude that water evaporation occurred in the glass tube. As a result of this water vapour movement, the surface snow became drier, which may have led to the volatilization of HCl and other organic acids. Even under the DARK condition, water vapour movement dried the sample, and the data plotted on the blank line. In other words, HCl emission into the atmosphere does not progress under DARK conditions. A possible reason for this observation is the lower water evaporation in the DARK condition than in the OPEN and COVERED conditions (Fig. 6). Compared to the OPEN and COVERED conditions, the [Na⁺] at 0–5 cm depth was relatively low, which could be attributed to lower water evaporation. The concentration gradient under the DARK condition would therefore be leaner than in other conditions. While nitrate ions behave similarly to chloride ions, the reasons for their behaviour are different, as discussed later. The main cause of nitrate ion loss would be photolysis. Because the photolysis reaction could not proceed, nitrate concentrations were stable under the DARK condition. Under the OPEN and COVERED conditions, data collected at sample depths greater than 10 cm plotted on the right side of the line. This indicates that nitrate photolysis may proceed even deeper than 10 cm in the snowpack. Moreover, under the OPEN and COVERED conditions, the glass tubes prevent the penetration of UV from the side, suggesting that more nitrate can be photolysed in natural conditions than shown in the experiment. Although nitrate photolysis may produce NO₂⁻, this ion was not detected. Therefore, all the NO₃⁻ generated by nitrate photolysis are released into the atmosphere as gases.

The Cl⁻ and NO₃⁻ behaviour in the mixed amended samples was similar to that of the natural samples (Supplementary Fig. S2). This implies that an amended sample could be suitable to conduct an experiment to estimate the post-depositional process in a laboratory when natural Antarctic snow is unavailable. Note that the Cl⁻ values of the amended samples in the lower layers are on the straight lines, which means that the Cl⁻ concentration changes were due to the water evaporation effect. Cl⁻ did not volatilize in this condition, presumably because there

![Fig. 7 Depth profile of normalized light intensity, measured at 12:00–16:00 on 14 January 2015.](image-url)
was insufficient H⁺ required for bonding with the Cl⁻ to volatilize as HCl gas. An excessive amount of Cl⁻ had been added to the amended sample.

Discussion

e-folding depth

The normalized solar intensities and depth dependence of the solar spectrum are shown in Fig. 7 and Supplementary Fig. S3. Several studies have reported the e-folding depth at 305 nm in Antarctica as 2–5, 8–12, 15–31, 10–21 and 24–69 cm, which are dependent on the snow impurity (France et al. 2011; Zatko et al. 2013; Zatko et al. 2016; Winton et al. 2020). Our result for ε (45 cm) was within a range of the ε (24–69 cm) calculated by a model (Zatko et al. 2016; Fig. 7). However, our result was relatively deeper than that of the field observations of 2–5, 8–12, 15–31 and 10–20 cm (France et al. 2011; Zatko et al. 2013; Winton et al. 2020). A shallower ε (8–12 cm) and deeper ε (15–31 cm) were observed near a station and about 11 km away, respectively, and these differences were attributed to the greater snow impurity near the station, as a result of human activities there (Zatko et al. 2013). This suggests that the relatively deeper ε (45 cm) observed in this study would be due to the lower snow impurity more than 100 km away from Syowa Station.

Estimation of the post-depositional loss of chloride and nitrate

The density of snow in the glass tube was approximately 0.5 g cm⁻³ and, from this, chloride and nitrate losses at each depth were calculated. Figure 8 shows the changes in the chloride, nitrate and sodium concentrations under OPEN conditions after four weeks. The [Na⁺] at a depth of 2.5 cm was relatively high, and the water at this depth evaporated. This shows that first-order reactions (i.e., volatilization and photolysis) are affected by changes in concentration of chloride and nitrate caused by the water evaporation. The change in the [Cl⁻] was small at this depth; however, the concentration increased with increasing depth: HCl deposited on the surface of the snowpack gradually migrated downwards. Cl⁻ initially moved through the snowpack as HCl gas or in the grain boundary of the ice as Cl⁻. The reason for the migration was the concentration gradient from the higher [Cl⁻] at the surface to the lower [Cl⁻] deeper down, caused by water evaporation. Additionally, convection within the snowpack would influence the migration. Either way, all the Cl⁻ deposited on the surface migrated through the snowpack, which suggests that other chemical species may also move in the snowpack.

Nitrate near the surface decreased by more than 50%. Previous investigations have suggested that the degree of post-depositional nitrate loss depends largely on the rate of snow accumulation, and photolytic nitrate loss occurs significantly at low snow accumulation sites (Erbland et al. 2013; Erbland et al. 2015; Zatko et al. 2016; Noro et al. 2018; Shi et al. 2018; Winton et al. 2020). Since the surface snow is buried after a short time at high-accumulation sites, the nitrate loss by photolysis should be nominal there. However, Savarino et al. (2007) observed that the increase in the particulate nitrate concentration in the atmosphere at a coastal site was a result of the re-emission of NO₃⁻ species from snow. Snow precipitation was not observed during the present investigation at the site (H128), which was a high-accumulation site, which implies that the nitrate remained on the snow surface during the experiments (Noro et al. 2018). In other words, nitrate photolysis can occur under a low-accumulation condition at coastal sites where post-depositional loss of nitrate was not believed to have occurred (Erbland et al. 2013; Noro et al. 2018; Shi et al. 2018). Moreover, the loss of chloride in the snow, which has a higher volatilization rate than nitrate, was not observed in the glass tube experiments. Therefore, photolysis dominated the post-depositional process of nitrate in the present study, and volatilization proceeded as a minor process.

Nearly 20% of the nitrate disappears completely, even at a depth of 40 cm. This observation can be explained by...
light of 305 nm penetrating deeper sites in the snowpack, as previously mentioned. Most of the nitrate remained at 25 cm because the glass wall obstructed the light at this depth. The nitrate loss was therefore underestimated. In the lower part of the glass tube, photolysis was enabled using light penetrating from the shallower sites.

Conclusion

Our study reveals new and significant information about the Antarctic geochemical cycle. We found that chloride migrated down into the snowpack due to the influence of water vapour movement. Moreover, 50% of the nitrate on surface snow was lost to photolysis, and approximately 20% of the nitrate was absent at a depth of 40 cm. This is the first report of chloride ion movement and a notable nitrate loss at any Antarctic coastal site. Our results show that the post-depositional process should be taken into account when interpreting nitrate records from coastal sites, as the amount of nitrate may be underestimated if this process is not considered at high-accumulation sites.

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Disclosure statement

The authors report no conflict of interest.

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