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Research article

A 60-year atmospheric nitrate isotope record from a southeastern Greenland ice core with minimal postdepositional alteration

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Abstract. Stable isotopes of atmospheric nitrate (NO_3^-) are valuable tools for tracing nitrogen sources and processes; however, their signals in ice core records are often disrupted by postdepositional processes. The ice core from the southeastern Dome (SE-Dome) in Greenland is a potential record of variations in atmospheric chemistry that has experienced a lower postdepositional effect owing to a high accumulation rate ($\sim 1 \text{ m}$ water equivalent per year). Herein, we report 60-year (1959–2014) $\delta^{15}N(NO_3^-)$ and $\Delta^{17}O(NO_3^-)$ records from the SE-Dome ice core. The $\delta^{15}N(NO_3^-)$ decreased from 1960 to 1974 and exhibited clear seasonal changes (high in summer and low in winter). The $\Delta^{17}O(NO_3^-)$ did not exhibit any significant long-term trends, but it did contain seasonal patterns. The mass-weighted annual average of $\delta^{15}N(NO_2^-)$ values in the SE-Dome core were 4.2 \pm 2.8% lower than those in the Greenland Summit ice core between 1959 and 2006. The TRansfer of Atmospheric Nitrate Stable Isotopes To the Snow (TRANSITS) model under the SE-Dome condition estimated changes of only 0.9% for $\delta^{15}N(NO_3^-)$ and -0.2% for $\Delta^{17}O(NO_3^-)$ from the initial deposition. Although differences in the source of NO₃⁻ cannot be discounted, the lower $\delta^{15}N(NO_3^-)$ values observed at SE-Dome compared to Summit were likely due to reduced postdepositional alteration. Therefore, the SE-Dome ice core NO_3^- record offers a precise reconstruction of nitrogen oxides (NO_x) emissions from both North America and western Europe, as well as atmospheric oxidation chemistry and transport, thereby providing reliable insight into atmospheric nitrogen cycling.

1 Introduction

Nitrate (NO_3^-) and its precursors $(NO_x = NO + NO_2)$ play important roles in the atmosphere. Tropospheric NO_x cycling produces ozone (O₃), a key component of the atmospheric oxidative capacity (Finlayson-Pitts and Pitts, 2000). NO_x emitted from various sources undergoes oxidation to form HNO₃, which contributes to acid rain (Shammas et al., 2020) and particulate matter (as NO_3^- ; Zhai et al., 2021); moreover, upon deposition, nitrate alters the nutrient balance in ecosystems (Duce et al., 2008). Owing to increasing fossil fuel and chemical fertilizer use since the beginning of the industrial revolution, NO_x levels in the atmosphere have increased, which is reflected in the elevated NO_3^- concentrations in ice cores, including those collected from Greenland (Neftel et al., 1985; Mayewski et al., 1986). Despite efforts to curb NO_x emissions through pollution mitigation measures and NO_x removal techniques, including three-way catalytic converters, the decline in the ice core NO_3^- concentrations has been relatively gradual, as observed in Greenland (Iizuka et al., 2017) and in an Alpine ice core (Eichler et al., 2023). This underscores the necessity to understand atmospheric NO_3^- dynamics beyond just precursor NO_x emissions, for which ice core NO_3^- can offer invaluable historical perspectives.

In addition to NO_3^- concentrations, NO_3^- stable isotopic compositions provide valuable information. Nitrogen isotopes (δ^{15} N) differ among NO_x sources and can be used to identify the origin of the NO_3^- (Hastings, 2010; Hastings et al., 2013). Previous studies of Greenland ice cores have identified decreases in the $\delta^{15}N$ values of NO₃⁻ ($\delta^{15}N(NO_3^-)$) as early as 1850 CE, which subsequently accelerated after 1950 CE (Hastings et al., 2009; Geng et al., 2014). These decreases in δ^{15} N have been interpreted as a change in source: increased anthropogenic emissions of NO_x from fossil fuel combustion (Hastings et al., 2009) and/or NO_x derived from soil amended with fertilizer (Felix and Elliott, 2013). In addition, complex factors control $\delta^{15}N(NO_3^{-})$. Previous studies have shown that isotopic fractionation can occur during gas-particle partitioning and washout (Freyer, 1991), kinetic NO₂ oxidation (Walters and Michalski, 2015a), and NO_x cycle equilibrium in the atmosphere (Walters et al., 2015b; Walters and Michalski, 2016). Another interpretation of the decrease in δ^{15} N in ice cores is related to changes in isotopic fractionation between gaseous HNO3 and particulate NO_3^- that resulted from acidity changes (Geng et al., 2014). The mass-independent oxygen isotope fractionation signals ($\Delta^{17}O = \delta^{17}O - 0.52 \times \delta^{18}O$) of NO_x and NO₃⁻ can reflect the oxygen source during oxidation (Michalski et al., 2003; Alexander et al., 2009). Positive Δ^{17} O values of NO₃⁻ $(\Delta^{17}O(NO_3^-))$ occur as a result of excess ¹⁷O (i.e., deviation from mass-dependent fractionation) transferred from O₃ to NO₃⁻ during photochemical cycling of NO and NO₂ and the

oxidation of NO₂ into HNO₃. Thus, middle to high latitudes have $\Delta^{17}O(NO_3^-)$ values of 22%-34‰ for atmospheric deposition (Michalski et al., 2003, 2012). The $\Delta^{17}O(NO_3^-)$ measurements in ice cores have also been used to investigate historical atmospheric processes (Geng et al., 2017).

However, postdepositional NO_3^- loss in snow or ice can reduce NO₃⁻ concentrations and change its isotopic compositions (Röthlisberger et al., 2000, 2002; Frey et al., 2009; Akers et al., 2022). Indeed, NO_3^- in snow can undergo photolysis by ultraviolet (UV) light ($\lambda = 290-350$ nm; Berhanu et al., 2014), which produces NO_2 that is released into the atmosphere via diffusion or wind pumping. Although NO₂ can partially reoxidize into NO_3^- in the atmosphere, postdepositional processes can lead to decreases in the $NO_3^$ concentrations in ice cores (Meusinger et al., 2014; Erbland et al., 2015). In addition, postdepositional processes can also cause significant isotopic fractionation (from -47.9% to -55.8%), and the remaining NO₃⁻ becomes enriched in ¹⁵N (Berhanu et al., 2015). In contrast, $\Delta^{17}O(NO_3^-)$ is not directly affected by photolysis but, rather, by the cage effect, in which the intermediate photoproducts (NO₂, NO₂, or ONOO⁻) undergo recombination reactions within snow grains to reform nitrate with exchange with water oxygen or react with radicals (e.g., OH) to regenerate NO₃⁻ after being emitted to the atmosphere (McCabe et al., 2005; Jiang et al., 2021). The alteration of $\Delta^{17}O(NO_3^-)$ can also occur through the reoxidation of NO₂ sourced from snow, which leads to nitrate formation in the overlying atmosphere (Erbland et al., 2013). Because NO_3^- photolysis in snow only occurs in the photic zone, the degree of postdepositional alteration is mostly controlled by the snow accumulation rate, as demonstrated in Antarctica (Akers et al., 2022). Even at the Greenland Summit ice core site, the $\delta^{15}N(NO_3^-)$ values of the snowpack are higher than those of the surface snow and the overlying atmosphere (Jarvis et al., 2009; Geng et al., 2014; Fibiger et al., 2016). The Greenland Ice Sheet Project 2 (GISP2) ice core exhibits decreasing $\delta^{15}N(NO_3^{-})$ values from glacial to interglacial periods (Hastings et al., 2005; Geng et al., 2015), which has been interpreted as the result of two potential causes: (1) changes in the NO_x source or (2) postdepositional effects related to the snow accumulation rate and dust concentrations. Overall, given the impacts of postdepositional processes, ice core records of NO_3^- and its isotopes require careful consideration.

Accurate understanding of atmospheric NO₃⁻ and its related nitrogen cycles can be more reliably obtained from ice core records with minimal postdepositional alterations. In this context, the southeastern Dome (SE-Dome) site in Greenland has distinct characteristics, including a snow accumulation rate of 1.01 ± 0.22 m water equivalent per year (m w.e. a⁻¹; for the period from 1960 to 2014; Iizuka et al., 2017), which is approximately 4 times greater than that at the Summit site (0.22 ± 0.05 m w.e. a⁻¹; Fig. 1c; Geng et al., 2014). The relatively high accumulation rate is expected to reduce the impact of postdepositional loss, preserving a more representative record of atmospheric NO₃⁻ deposition. Furthermore, statistical analysis using the Mann-Kendall test for monotonic trends (Kendall, 1975; Mann, 1945) revealed no significant decadal trends in snow accumulation at either the SE-Dome (Kawakami et al., 2023) or the Summit (p > 0.05;data from Geng et al., 2014) site, suggesting that within-site accumulation variability does not strongly influence nitrate trends at these locations. The Summit and SE-Dome sites receive atmospheric inputs from similar source regions in North America and western Europe, based on Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) 7 d backward-trajectory modeling (Fig. 1b and c). However, the extent to which postdepositional processes affect NO₃⁻ isotopic compositions at these sites has not been systematically assessed. In this study, we present the $\delta^{15}N(NO_3^-)$ and $\Delta^{17}O(NO_3^-)$ records obtained from a 90.45 m ice core drilled at the SE-Dome site. By applying the TRansfer of Atmospheric Nitrate Stable Isotopes To the Snow (TRANSITS) model (Erbland et al., 2015; Jiang et al., 2021) with sitespecific modifications, we evaluate the extent to which postdepositional processes influence NO₃⁻ at SE-Dome and assess its suitability for reconstructing past atmospheric NO_x emissions. Our results demonstrate that SE-Dome preserves a robust NO_3^- isotopic signal with minimal postdepositional modification, making it a valuable archive for investigating anthropogenic changes in atmospheric NO_3^- over the Northern Hemisphere, particularly from eastern North America and western Europe.

2 Materials and methods

2.1 Samples

This study was based on a 90.45 m ice core drilled at the SE-Dome site (67.18° N, 36.37° W; 3170 m a.s.l., meters above sea level) in 2015 (Iizuka et al., 2016). The age-depth scale was determined using the oxygen isotope matching method, which matches the δ^{18} O variations between ice core records and isotope-enabled climate model estimates, and indicated that this ice core covers the period from 1959 to 2014 (Furukawa et al., 2017). The reliability of this dating method generally falls within the 95 % confidence interval (typically around an average of ± 0.9 months). The greatest uncertainty was reported at 2 months in some years (Furukawa et al., 2017). We divided the ice core samples into four seasons: spring (21 March-20 June), summer (21 June-20 September), fall (21 September-20 December), and winter (21 December-20 March). For samples analyzed at a twoseason resolution (1959-1980 and 1995-2014), spring and summer were combined into summer, whereas fall and winter were combined into winter. This approach was adopted to ensure a consistent sample size and minimize analytical uncertainty for periods with lower temporal resolution, while still capturing seasonal variations relevant to NO_3^- deposition and atmospheric conditions.

All SE-Dome ice core samples used in this study were stored in a refrigerated room $(-50 \,^{\circ}\text{C})$ at the Institute of Low Temperature Science (Hokkaido University, Sapporo, Japan). Each ice sample $(3 \,\text{cm} \times 4 \,\text{cm} \text{ cross-dimension})$ was cut using a band saw in a refrigerated room $(-20 \,^{\circ}\text{C})$ and decontaminated by removing the outermost $\sim 5 \,\text{mm}$ of ice with a ceramic knife in a class 10 000 clean booth, resulting in a loss of approximately 30% of the original sample weight. The remaining 70% of the cleaned samples was shipped frozen ($\sim -20 \,^{\circ}\text{C}$) to the Tokyo Institute of Technology (Tokyo Tech, Yokohama, Japan). The samples were then stored in a freezer at $-30 \,^{\circ}\text{C}$ until analysis.

2.2 Sample analysis

The NO₃⁻ in each sample (n = 136) was separated from other ions using ion chromatography (IC; Dionex Integrion, Thermo Fisher Scientific) according to the methods described by Noro et al. (2018). Changes in the isotopic compositions of NO_3^- during ion chromatographic separation were $< 0.4\,\%$ for $\delta^{\dot{1}5}N$ values and within the analytical error range for Δ^{17} O values (Noro et al., 2018). After ion separation, NO_3^- in solution was converted and neutralized to the Na⁺ form by passing it through an ion exchange column. The isotopic compositions of NO₃⁻ were measured using a bacterial method that converts NO_3^- to N₂O (Sigman et al., 2001; McIlvin et al., 2011), followed by N₂O decomposition via microwave-induced plasma (MIP), a technique developed at Tokyo Tech (Hattori et al., 2016). Isotopic reference materials, as well as United States Geological Survey (USGS) standards 32, 34, 35, and their mixtures (prepared in $18.2 \text{ M}\Omega \text{ cm}$ water), were also analyzed with the samples using the same analytical processes.

Stable isotopic compositions are reported as $\delta X = R_{\text{sample}}/R_{\text{reference}} - 1$, where *X* denotes ¹⁵N, ¹⁷O, or ¹⁸O, while *R* denotes the isotope ratios, such as ¹⁵N/¹⁴N, ¹⁷O/¹⁶O, and ¹⁸O/¹⁶O, determined for both sample and standard materials. The δ^{15} N, δ^{18} O, and Δ^{17} O values are reported in per mil (‰) notation. The δ^{15} N values are relative to atmospheric N₂ (air), while the δ^{18} O and Δ^{17} O values are relative to Vienna Standard Mean Ocean Water. By propagating the analytical uncertainties for the IC separation and replicating isotopic measurements of USGS standards 34, 35, and 32, the estimated combined uncertainties were ±0.4‰ for both δ^{15} N(NO₃⁻) and Δ^{17} O(NO₃⁻).

2.3 TRANSITS modeling

The TRANSITS model (Erbland et al., 2015), a multilayer one-dimensional isotopic model, was used to simulate NO_3^- recycling across the air–snow interface (i.e., UV photolysis of NO_3^- , NO_x emission, local NO_2 oxidation, and NO_3^- deposition) and its associated isotopic effects. The model is op-



Figure 1. (a) Map of the southeastern Dome (SE-Dome, star) and Summit (circle) sites in Greenland (annual precipitation from ERA5 reanalysis climate data; Hersbach et al., 2020). Probability distributions for the air masses overlying the (b) SE-Dome and (c) Summit sites from 7 d three-dimensional back-trajectory analysis based on HYSPLIT modeling (1960–2019). Detailed back-trajectory analysis procedures were the same as those in Iizuka et al. (2018). (d) Annual accumulation rate at the SE-Dome (Furukawa et al., 2017) and Summit (Geng et al., 2014) sites.

erated at a weekly resolution (52 time steps per year), and the default snow depth resolution is 1 mm. In each step, $NO_3^$ photolysis is calculated according to the depth-dependent photochemical flux and NO₃⁻ concentration. All generated NO_2 enters the atmosphere and is reoxidized to NO_3^- , which is deposited on the surface snow with the primary $NO_3^$ from long-range transport in the next step. The original snow moves downward as snowfall continues, and newly deposited snow is divided into 1 mm layers. Once the NO_3^- is buried beneath the light transmission band, the layer is regarded as an archive. We adapted the parameters of the TRANSITS model, originally developed for the Summit site by Jiang et al. (2021). This model reproduces the seasonal variation pattern of $\delta^{15}N(NO_3^-)$ in the surface snow at the Summit site reported by Jarvis et al. (2009), highlighting the importance of postdepositional processes at site. In addition, this model estimated the net loss of NO_3^- (4.1%) and associated changes in $\delta^{15}N(NO_3^-)$ (+2.6%) and $\Delta^{17}O(NO_3^-)$ (-0.9%) between primary deposition and NO₃⁻ archived in the ice, under an estimate of the horizontal export fraction of locally reoxidized NO₃⁻ (f_{exp}) of 35 % (Jiang et al., 2021). In this study, we applied the same model under the SE-Dome condition by adjusting the parameters to examine the effects of snow NO_3^- photolysis on the NO_3^- concentration and its isotopes.

The snow accumulation rate was set at $1.01 \text{ m w.e. a}^{-1}$, based on the 1960–2014 average from SE-Dome ice core data (Iizuka et al., 2017), with additional tests conducted at rates of 0.25, 0.6, and $1.4 \text{ m w.e. a}^{-1}$. The mass balance of NO₃⁻ between the snow and atmosphere depends on NO_3^- influxes and outfluxes. We expressed the NO_3^- flux as F_Y , which includes the primary NO₃⁻ flux from long-range transport (F_{pri}), NO₃⁻ flux from NO₃⁻ photolysis (F_P), atmospheric NO₃⁻ deposition flux (F_D), and ice core NO₃⁻ flux buried beneath the light band (F_A) . These fluxes reflect changes in NO_3^- and its isotopic compositions in the snow and atmosphere. The TRANSITS model considers that two processes can change $\delta^{15}N(NO_3^-)$ as a result of isotope fractionation from UV photolysis and NO_3^- deposition (i.e., co-condensation and dry deposition). The nitrogen isotope fractionation constant during photolysis ($^{15}\varepsilon_{p}$) was calculated using a ratio of the ¹⁴NO₃⁻ and ¹⁵NO₃⁻ photolysis rates (${}^{15}\varepsilon_{p} = {}^{15}J/{}^{14}J - 1$, where J represents the photolysis rate constant) in each snow layer (Erbland et al., 2013). The J at different depths (z) (J(z)) was calculated according to Eq. (1):

$$I(z) = \int_{280 \text{ nm}}^{350 \text{ nm}} \Phi(\lambda) \times \sigma_{\text{NO}_3^-}(\lambda) \times I(z,\lambda) d\lambda, \qquad (1)$$

where I is the actinic flux, while Φ and σ are the quantum yield and absorption cross-section of NO₃⁻ photolysis, respectively. The quantum yield of NO₂⁻ photolysis has significant uncertainties (Meusinger et al., 2014). However, it is unlikely that the quantum yield of NO_3^- photolysis would differ substantially between the SE-Dome and Summit sites. As this study compares the differences between these sites, which have notably different snow accumulation rates, the quantum yield of NO_3^- photolysis at the SE-Dome site was set to the same value (0.002) as that estimated for the Greenland Summit site by Jiang et al. (2021). The absorption crosssections of ${}^{14}\text{NO}_3^-$ (${}^{14}\sigma_{\text{NO}_2^-}$) and ${}^{15}\text{NO}_3^-$ (${}^{15}\sigma_{\text{NO}_2^-}$) were derived from Berhanu et al. (2014). The nitrogen isotope fractionation constant during deposition ($^{15}\varepsilon_{d}$) was set to +10%(Erbland et al., 2015). For the oxygen isotopes, only the mass-independent fractionation signal (Δ^{17} O) was modeled. The cage effect (i.e., decrease in Δ^{17} O of the snow NO₃ owing to secondary chemistry during NO₃ photolysis; Mc-Cabe et al., 2005) was set to 15% according to Erbland et al. (2015), and the TRANSITS model calculated the exchange of oxygen atoms with water during UV photolysis and atmospheric NO-NO₂ cycling, both of which alter Δ^{17} O. The Δ^{17} O(NO₂⁻) fractionation mechanisms in the TRANSITS model during these processes are explained in

detail in Jiang et al. (2021).

The atmospheric boundary layer at the SE-Dome site was assumed to be a zero-dimensional well-mixed box, and the snowpack was assumed to be a stack of snow layers deposited at different times. Weekly air temperatures (T), pressures (P), and average boundary layer heights (h) from 1950 to 2020 were obtained from the second-generation European Centre for Medium-Range Weather Forecasts atmospheric analysis of global climate (ERA5) (Hersbach et al., 2020; Khalzan et al., 2022). O₃, OH, peroxyl radical (RO₂ and HO₂), and BrO concentrations were used to calculate the rates of NO-NO2 cycling and NO2 oxidation to HNO3. However, because these records were not available for the SE-Dome, they were extracted from the outputs of the v.12.9.3 (https://doi.org/10.5281/zenodo.3974569, The International GEOS-Chem User Community, 2020) GEOS-Chem atmospheric chemical transport model (http://www.geos-chem. org, last access: 7 January 2024) using the Modern-Era Retrospective analysis for Research and Applications, Version 2 (MERRA-2) meteorological field, with 4° latitudinal and 5° longitudinal resolutions. The GEOS-Chem model was run for the year 2017 after a 1-year spin-up run, and the monthly averages for the O₃, OH, HO₂, and BrO concentrations in the planetary boundary layer in the SE-Dome grid were used. We selected 2017 as the representative period, which should not vary significantly from other recent years and should ensure robust outcomes. Given that tropospheric O₃ concentrations were comparable between the SE-Dome and Summit grids in the GEOS-Chem model, the total O₃ column (TCO) was set to the same value (266-408 DU, Dobson units) as that used in a previous study of the Summit site (Jiang et al., 2021). F_{pri} was estimated to be 16.4, 23.6, 13.3, and 11.5 mg N m⁻² a⁻¹ for spring, summer, fall, and winter, respectively, based on the seasonal NO₃⁻ fluxes at the SE-Dome site from 1960 to 2014 (Iizuka et al., 2018).

An *e*-folding depth, which is the depth to which light enters the snow layer and attenuates to an initial intensity of 1/e (owing to absorption and scattering), for the SE-Dome site was calculated using the snow density (ρ_{snow}), the calculated specific surface area (SSA), and fixed light-absorbing impurity concentrations (Jiang et al., 2021). The ρ_{snow} of 400 kg m⁻³ for the SE-Dome site was obtained from an observation at the SE-Dome (Oyabu et al., 2016). The SSA for the SE-Dome site was determined to be 47.0 m² kg⁻¹ using the relationship between the SSA and ρ_{snow} , according to a previous study (Domine et al., 2007):

$$SSA = -174.13 \times \ln(\rho_{snow}) + 306.4, \tag{2}$$

where SSA is in units of square centimeters per gram $(cm^2 g^{-1})$ and the units for ρ_{snow} were changed to grams per cubic centimeter ($g cm^{-3}$). For the light-absorbing impurity concentrations, we established constant concentrations of the three main light-absorbing impurities in snow: dust, soot (BC), and organic humic-like substances (HULISs). The dust concentration was set to 33.94 ng g^{-1} according to an average concentration for the SE-Dome ice core from 1960 to 2014 (Amino et al., 2021). Owing to a lack of direct observations for BC and HULISs at the SE-Dome, we assumed these concentrations based on the Ca²⁺ concentration ratio between the Summit and SE-Dome sites. Here, [BC]_{Summit} and [HULIS]_{Summit} were set to 1.4 and 31 ng g^{-1} , respectively, according to Jiang et al. (2021). Furthermore, $[Ca^{2+}]$ at the Summit site was set to 6.5 ng g^{-1} according to an average of the 2 m shallow snowpack observation (Geng et al., 2014); $[Ca^{2+}]$ at the SE-Dome site was set to 11.6 ng g⁻¹ according to an average from the SE-Dome ice core from 1960 to 2014 (Iizuka et al., 2018). Thus, [BC]SE-Dome and [HULIS]_{SE-Dome} were calculated as 2.2 and 47.6 ng g^{-1} , respectively, and used for the model calculation. An e-folding depth of 10 cm was obtained based on the above inputs. The calculated e-folding depth for the SE-Dome site was consistent with previous estimates from the GEOS-Chem model investigating the impact of postdepositional effect in snow (Zatko et al., 2016).

The horizontal export fraction of locally reoxidized NO_3^- (f_{exp}) under the SE-Dome condition was calculated with the same scheme described for the Antarctic Plateau (Erbland et al., 2015) and Greenland Summit (Jiang et al., 2021), as outlined in the following equations:

$$f_{\exp} = \frac{\frac{1}{\tau_1}}{\frac{1}{\tau_1} + \frac{1}{\tau_2}} \times \left(1 + \frac{\frac{1}{\tau_2}}{\frac{1}{\tau_3} + \frac{1}{\tau_1}}\right), \text{ where}$$
(3)

$$\tau_1 = \frac{L}{V_{\rm h}},\tag{4}$$

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$$\tau_2 = \frac{1}{k \, \text{[OH]}},\tag{5}$$

$$\tau_3 = \frac{H}{V_{\rm d}}.\tag{6}$$

In these equations, τ_1 , τ_2 , and τ_3 represent the lifetimes of horizontal transport, oxidation of NO2 by OH radicals, and vertical deposition, respectively. L and H denote the respective summer boundary layer height and horizontal characteristic, while $V_{\rm h}$ is the mean horizontal wind speed, k is the rate constant for the NO₂+OH reaction, and V_d is the dry deposition velocity of HNO₃ (Jiang et al., 2021). The values used for the calculation are summarized in Table S2, and the same physicochemical values as those of Summit were used, while parameters such as temperature and boundary layer height were incorporated from ERA5 data. We obtained f_{exp} to be 47 % under the SE-Dome condition, but the calculated f_{exp} may oversimplify the processes governing NO_3^- deposition and the chemical loss pathways of NO_x , as discussed previously (Jiang et al., 2021). Therefore, we considered the sensitivity of postdepositional alteration to variations in f_{exp} between initial deposition and the point at which NO_3^- becomes archived in the ice.

At the initial time (t = 0) in the TRANSITS model, the NO₃⁻ concentration was set to 71.12 ng g⁻¹ based on the average NO₃⁻ concentration in the SE-Dome I ice core (Iizuka et al., 2018), while the δ^{15} N(NO₃⁻) and Δ^{17} O(NO₃⁻) values in the snowpack were set to 0% and 30%, respectively, according to previous TRANSITS settings (Erbland et al., 2015; Jiang et al., 2021). This initial isotopic parameter does not affect the model interpretation of changes in δ^{15} N(NO₃⁻) and Δ^{17} O(NO₃⁻) due to postdepositional processing. The 3-year distributions of NO₃⁻ and its isotopes were simulated with and without NO₃⁻ photolysis scenarios under the SE-Dome condition, from which profiles of NO₃⁻ concentrations, δ^{15} N(NO₃⁻), and Δ^{17} O(NO₃⁻) were output. The parameters used in the TRANSITS model are summarized in Table 1 and Supplement data file 1.

2.4 Statistical analysis

XLSTAT 2023 (Addinsoft, Paris, France) was used for the Mann–Kendall trend analysis over 1959–2014. SPSS Statistics 25 (IBM SPSS, Armonk, NY, USA) was used to perform the *t* tests of annual changes in the $\delta^{15}N(NO_3^-)$ and NO_3^- concentrations. Statistical significance was set at p < 0.05.

3 Results

3.1 Nitrate isotope records from the SE-Dome ice core

The NO_3^- isotope data and fluxes obtained from the SE-Dome ice core are shown in Figs. 2 and S1. The seasonal variations were larger in the samples analyzed at a fourseason resolution (1981–1994), which may have been caused by age errors (Furukawa et al., 2017). Accordingly, massweighted averages were calculated for the summer and winter fractions from the seasonal samples during 1981–1994. From 1959 to 2014, the $\delta^{15}N(NO_3^-)$ values were generally higher in summer (-2.9 ± 2.6%) than in winter (-6.9 ± 2.9%) (Figs. 2a and S1a). To assess annual changes in $\delta^{15}N(NO_3^-)$ over this period, we calculated the annual massweighted average $\delta^{15}N(NO_3^-)$ values and found that they decreased from 1959 to 1974 and exhibited no significant (p > 0.05) trends after 1975 (mean value of $-4.8 \pm 1.3\%$) (Fig. 2a). No clear relationship was observed between the annual variations in the $\delta^{15}N(NO_3^-)$ and NO_3^- concentrations (p = 0.37).

Using a similar method to that employed for $\delta^{15}N(NO_3^-)$, we also calculated the mass-weighted average and annual mass-weighted average for $\Delta^{17}O(NO_3^-)$. The $\Delta^{17}O(NO_3^-)$ also exhibited a seasonal pattern, with lower values in the summer ($27.8 \pm 1.3\%$) than those in the winter ($31.3 \pm$ 1.9%), yielding a mass-weighted average of $29.3 \pm 1.2\%$ over the entire period (Figs. 2b and S1). The average annual $\Delta^{17}O(NO_3^-)$ values were relatively high ($\sim 33\%$) in 1988, whereas low values were observed in 2013 and 2014 (Fig. 2b). Excluding these particular years, no significant annual increases or decreases (p > 0.05) were observed in the $\Delta^{17}O(NO_3^-)$ values.

3.2 TRANSITS model results

We aimed to know the changes in NO_3^- from primary deposition to the ice core archive. However, these changes primarily depend on the f_{exp} value – the fraction of NO₃⁻ exported from the site of photolysis. Therefore, we calculated the dependency of the $\delta^{15}N(NO_3^-)$ and $\Delta^{17}O(NO_3^-)$ values on f_{exp} at the SE-Dome using the same approach as Jiang et al. (2021) (Fig. 3). The postdepositional alterations in $\delta^{15}N(NO_3^-)$ and $\Delta^{17}O(NO_3^-)$ between initial deposition and the ice core NO_3^- concentration at the SE-Dome were dependent on f_{exp} . As shown in Fig. 3, an inverse relationship was observed between f_{exp} and $\delta^{15}N(NO_3^-)$ and $\Delta^{17}O(NO_3^-)$. In the case of δ^{15} N, when f_{exp} is high, a larger proportion of isotopically light NO_3^- is removed from the snowpack due to strong isotopic fractionation during photolysis. As a result, the remaining NO_3^- in the snow becomes isotopically enriched in ¹⁵N. In contrast, for $\Delta^{17}O(NO_3^-)$, when f_{exp} is high, the impact of photolysis-induced isotopic fractionation on $\Delta^{17}O(NO_3^-)$ is minimal; thus, its value remains largely unchanged. Conversely, when f_{exp} is low (i.e., a significant portion of the nitrogen species is emitted and subsequently redeposited), the NO₃⁻ with lower $\Delta^{17}O(NO_3^-)$, originating from NO₂+OH reaction, dominates the signal, leading to a decrease in $\Delta^{17}O(NO_3^{-})$. These distinct mechanisms explain why $\delta^{15}N(NO_3^-)$ and $\tilde{\Delta}^{17}O(NO_3^-)$ exhibit opposite trends.

However, the degree to which the changes in $\delta^{15}N(NO_3^-)$ and $\Delta^{17}O(NO_3^-)$ were dependent on f_{exp} was less evident at the SE-Dome site than at the Summit site (Fig. 3). Notably,

Parameters	Description	Value	Unit*	Data origin
ρ	Snow density	400	${\rm kg}{\rm m}^{-3}$	Oyabu et al. (2016)
SSA	Snow-specific surface area	47.0	$m^2 kg^{-1}$	Domine et al. (2007)
[BC] _{SE-Dome}	BC concentration	2.2	$\rm ngg^{-1}$	See text
[Dust]	Dust concentration	33.94	$ng g^{-1}$	Amino et al. (2021)
[HULIS] _{SE-Dome}	HULIS concentration	47.6	$\rm ngg^{-1}$	See text
ТСО	Total column ozone	See Supplement data file 1	DU	Jiang et al. (2021)
h	Boundary layer height	See Supplement data file 1	m	ERA5 (Hersbach et al., 2020; Khalzan et al., 2022)
Т	Temperature		°C	
Р	Pressure		hPa	
[O ₃]	O ₃ concentration	26.66–32.30	ррb	GEOS-Chem v12.9.3 (https://doi.org/10.5281/ zenodo.3974569, The International GEOS-Chem User Community, 2020)
[BrO]	BrO concentration	0.06–0.76	ppt	
[OH] and [HO ₂]	[OH] and [HO ₂] concentrations	See Supplement data file 1	molec. cm^{-3}	
A	Snow accumulation	101	$\mathrm{cm}\mathrm{a}^{-1}$	Iizuka et al. (2017, 2018)
F _{pri}	Primary nitrate flux	16.28	$kg N m^{-2} a^{-1}$	
fexp	Export fraction	47 %	_	See Table S2
15 _{ep}	N isotope fractionation constant during photolysis	${}^{15}\varepsilon_{\rm p} = J^{15}/J^{14} - 1$	%0	Erbland et al. (2013)
15 _{<i>e</i>d}	N isotope fractionation constant during deposition	10	%0	
fcage	Cage effect	15	%	Erbland et al. (2015)

 Table 1. Parameters used in the TRansfer of Atmospheric Nitrate Stable Isotopes To the Snow (TRANSITS) model for the SE-Dome ice core.

* Note that ppt denotes parts per trillion.

even when the snow accumulation rate for SE-Dome is adjusted from the minimum (0.6 m w.e. a^{-1}) to the maximum (1.4 m w.e. a^{-1}) values, as shown in Fig. 1d, the results indicate that changes in $\delta^{15}N(NO_3^-)$ and $\Delta^{17}O(NO_3^-)$ are less sensitive to f_{exp} compared to Summit. Furthermore, when a snow accumulation rate of 0.25 m w.e. a^{-1} , equivalent to that used in the Summit study (Jiang et al., 2021), was applied, the variations were nearly identical to those observed at Summit (Fig. S2). This suggests that the differences in postdepositional alterations for $\delta^{15}N(NO_3^-)$ and $\Delta^{17}O(NO_3^-)$ are primarily caused by differences in accumulation rates.

Using the method of Erbland et al. (2015), f_{exp} was calculated as 47 % at SE-Dome, reflecting an estimated net loss of 1.4 % due to postdepositional alteration in the NO₃⁻

concentration at the site, with corresponding changes in $\delta^{15}N(NO_3^-)$ and $\Delta^{17}O(NO_3^-)$ of +0.9% and -0.2%, respectively (Fig. 3). In contrast, the estimated net loss of NO_3^- under the Summit condition showed higher values (4.1%) and associated changes in $\delta^{15}N(NO_3^-)$ (+2.6%) and $\Delta^{17}O(NO_3^-)$ (-0.9%) when the f_{exp} value was 35\%, as estimated previously (Jiang et al., 2021). Thus, even when using the same evaluation criteria, the postdepositional alteration in NO_3^- and its isotopic compositions at the SE-Dome site were smaller than those at the Summit site. We note that, under an extreme condition of $f_{exp} = 100\%$, the estimated changes in $\delta^{15}N(NO_3^-)$ from initial deposition due to postdepositional processing under the SE-Dome condition



Figure 2. The NO₃⁻ isotope data and fluxes from the SE-Dome ice core: (a) $\delta^{15}N(NO_3^-)$, (b) $\Delta^{17}O(NO_3^-)$, and (c) NO₃⁻ flux (mmol m⁻² a⁻¹).

were $+1.8\%_0$, which is significantly lower than that under the Summit condition of $+6.8\%_0$ (Fig. 3).

Figure 4 shows the results obtained from the TRANSITS model for NO₃⁻ and its isotopic compositions for the SE-Dome site when considering an f_{exp} value of 47%. The model considering photolysis showed a maximum 6% decrease in the annual NO_3^- concentrations during spring and early summer compared to the scenario without photolysis (Fig. 4a). The postdepositional effects (primarily due to photolytic isotopic fractionation) caused a fluctuation in $\delta^{15}N(NO_3^-)$ of -1% to +2%, with higher values in summer $(1.3 \pm 0.7\%)$ and lower values in winter $(0.2 \pm 0.2\%)$ (Fig. 4b). The variation in the $\Delta^{17}O(NO_3^-)$ value, which was initially set at 30 %, is attributed to a slight decrease in the atmospheric NO_3^- concentration owing to its reoxidization during spring to summer. Thus, when photolysis is minimal in the fall and winter, the $\Delta^{17}O(NO_3^-)$ values remained close to the initial value (30%) (Fig. 4c). Conversely, during spring and summer, when $\delta^{15}N(NO_3^-)$ values increase, a decreasing $\Delta^{17}O(NO_3^-)$ trend was observed (Fig. 4c). However, the extent of this change is minimal, with values reaching a minimum of $\sim 29.6 \%$. The seasonality in the NO₃⁻ concentration and postdepositional alteration in its isotopic composition were less obvious at SE-Dome than that at Sum-



Figure 3. Sensitivity of the changes in $\delta^{15}N(NO_3^-)$ and $\Delta^{17}O(NO_3^-)$ of the ice core nitrate to f_{exp} . Positive or negative values indicate deviations from initial deposition. The shaded area in the SE-Dome calculations represents results obtained using snow accumulation rates of 0.6 and 1.4 m w.e. a⁻¹.

mit, where $\delta^{15}N(NO_3^-)$ varied by > 5 ‰ and $\Delta^{17}O(NO_3^-)$ by $\sim 2\%$ (Jiang et al., 2021).

4 Discussion

4.1 The $\delta^{15}N(NO_3^-)$ values from the SE-Dome and Summit sites

Figure 5 shows the annual average $\delta^{15}N(NO_3^-)$ values obtained from the SE-Dome ice core and the previously published $\delta^{15}N(NO_3^-)$ values from the Summit site (Hastings et al., 2009; Geng et al., 2014). Decreasing trends in $\delta^{15}N(NO_3^-)$ were observed in both the Summit and SE-Dome ice cores until approximately 1974, after which no clear changes occurred (Fig. 5). Notably, based on the overlapping analysis period from 1959 to 2006 (n = 44), the annual $\delta^{15}N(NO_3^-)$ values in the SE-Dome ice core were found to be $4.2 \pm 2.8 \%$ lower than those in the Summit ice core (Fig. 5). The observed differences in $\delta^{15}N(NO_3^-)$ values between the SE-Dome and Summit cores may be attributed to (1) variations in the δ^{15} N values of NO₃⁻ deposited at the two sites and/or (2) variations in the degree of postdepositional alterations between the two sites. These two points are discussed in detail below.

First, regarding the differences in the δ^{15} N values of NO₃⁻ deposited at the two sites, there are two main sources of the NO₃⁻ deposited in the Greenland ice core. One long-range source is derived primarily from anthropogenic sources outside of Greenland. The other source involves NO_x released from NO₃⁻ photolysis within the snowpack, which is then re-oxidized and redeposited. Although the air masses at the SE-



Figure 4. TRANSITS model results for the SE-Dome site: (a) NO₃⁻ concentration, (b) $\delta^{15}N(NO_3^-)$, and (c) $\Delta^{17}O(NO_3^-)$. The black and red lines represent the variations calculated with and without NO₃⁻ photolysis, respectively.



Figure 5. The $\delta^{15}N(NO_3^-)$ values obtained from the SE-Dome (this study) and Summit (Hastings et al., 2009; Geng et al., 2014) ice cores. The shaded area for $\delta^{15}N(NO_3^-)$ values in SE-Dome represents the propagated errors of the annual average, based on the seasonal concentration and $\delta^{15}N$ variations.

Dome and Summit sites have similar source regions – North America and western Europe (Fig. 1b and c) – the degree of influence from reactive nitrogen sources differed between sites (Fig. S3). At the SE-Dome site, the contributions from outside Greenland were relatively high, with nearly equal influence from North America and European Union (EU) countries (Fig. S3b). In contrast, the contributions from EU countries were relatively low at the Summit site, while the North American region (mostly Eastern Canada) and inner Greenland had greater contributions (Fig. S3c). The NO_x sources from western Europe and North America are not necessarily similar; for example, differences in the relative contri-

butions of various NO_x sources are reflected in their $\delta^{15}N$ values, with NO_x from coal and biomass tending to have higher $\delta^{15}N$ values, while NO_x emissions from oil, natural gas, and soil tend to have lower $\delta^{15}N$ values (e.g., Elliott et al., 2019). To date, there have been no studies comprehensively comparing the isotopic composition of atmospheric NO_3^- between Europe and North America. The limited available data show that $\delta^{15}N$ values in total atmospheric NO_3^- (sum of gaseous HNO₃ and particulate NO_3^-) in the northeastern USA range from -10% to +5% (Bekker et al., 2023), while those in rainwater NO_3^- (including both gaseous HNO₃ and particulate NO_3^-) in Switzerland range

from -12 to +6% (Freyer, 1991), which are indistinguishable. A recent study (Song et al., 2021) compiled the δ^{15} N values of precipitated NO₃⁻ between urban and non-urban areas in Europe (n = 8 and n = 15, respectively) and North America (n = 10 and n = 73, respectively), showing no clear distinction between the two regions, although Europe exhibited slightly higher values. Hence, there is no clear evidence that the long-term δ^{15} N trends in European countries consistently remain lower than those in the USA or Canada; thus, the $\sim 4\%$ lower δ^{15} N(NO₃⁻) values observed at SE-Dome, which is relatively more influenced by air masses from Europe, cannot be explained solely by air mass origin differences.

Considering the potential impact of snow-sourced NO_x and reoxidized NO_3^- , it is important to note that the extent of recycled NO_x from NO₃⁻ photolysis in the Greenland ice sheet differed between these two sites. As modeled by Zatko et al. (2016), recycled NO_x is typically more important at inland sites such as the Summit than coastal sites such as the SE-Dome. Additionally, the contribution of air masses from inside Greenland was higher at the Summit site than at the SE-Dome (Fig. S3). Nevertheless, the δ^{15} N values of NO_x and reoxidized NO₃⁻ are typically low due to isotopic fractionation during snow NO3 photolysis in the snow and ice, with $\delta^{15}N(NO_3^-)$ in high-latitude air masses attributed to photochemical NO_x production in snow, resulting in $\delta^{15}N(NO_3^-)$ values of -10% to -43% in polar regions (e.g., Savarino et al., 2007; Morin et al., 2009; Shi et al., 2021). Thus, the contribution of locally recycled NO_3^- , which was greater at the Summit, cannot explain why $\delta^{15}N(NO_3^-)$ values were lower at the SE-Dome than at the Summit site.

Finally, regarding the differences in postdepositional alterations between two sites, we applied parameters specific to the SE-Dome in the TRANSITS model (Figs. 3 and 4). The model results for the SE-Dome, accounting for postdepositional NO₃ photolysis, showed a net NO₃ loss of 1.3 % and a $\delta^{15}N(NO_3^-)$ increase of +0.9% (see Sect. 3.2). In comparison, the Summit condition resulted in a $\sim 4\%$ net NO₃ loss and a +2.6 $\% \delta^{15}$ N(NO₃⁻) increase (Jiang et al., 2021). Although the lower $\delta^{15}N(NO_3^-)$ values at the SE-Dome can be partially explained by the model, they cannot be fully accounted for quantitatively. However, the estimated +2.6%increase in $\delta^{15}N(NO_3^-)$ at the Summit site may be underestimated due to an underestimation of f_{exp} (Jiang et al., 2021). Indeed, an observational study (Honrath et al., 2002) indicates that most of the NO_x and/or HNO_3 emitted from the snow at Summit is largely exported from the local boundary layer if no wet deposition occurs, suggesting that the f_{exp} value can reach ~ 1 under Summit conditions. Therefore, the actual net NO₃⁻ loss and $\delta^{15}N(NO_3^-)$ variation at the Summit site may have been larger than the 4% estimated by Jiang et al. (2021). Thus, when considering higher f_{exp} values, the difference due to postdepositional alterations could be higher than the modeled difference between the SE-Dome (+0.9%), this study) and Summit (+2.6%), Jiang et al., 2021) sites. Indeed, when considering an extreme condition of $f_{exp} = 100 \%$, the difference between the Summit and SE-Dome sites becomes ~ 5% (Fig. 3). Overall, although there is some uncertainty in the model, it is likely that the SE-Dome experienced less postdepositional alteration, thus preserving the atmospheric $\delta^{15}N(NO_3^-)$ values more effectively than at the Summit site.

Although the contribution of different NO₃⁻ sources cannot be entirely ruled out, our analysis shows that the observed $\delta^{15}N(NO_3^-)$ value at SE-Dome, which is $4.2 \pm 2.8 \%$ lower than that at Summit, can largely be attributed to differences in postdepositional alterations.

4.2 Seasonal variations in NO₃⁻ isotopes

The postdepositional effect, as estimated from the TRAN-SITS model, yielded a summer–winter difference in $\delta^{15}N(NO_3^-)$ of $1.1 \pm 0.7\%$ and a difference smaller than 0.5% in $\Delta^{17}O(NO_3^-)$, as described in Sect. 3.2 (Fig. 4b and c). In contrast, except for the anomalous years (1959– 1961, 1972, 1995, 2005, and 2013), the observed summer– winter differences for respective years in the SE-Dome ice core were $5.3 \pm 2.4\%$ (0.4%–9.8%) for $\delta^{15}N(NO_3^-)$ and $-4.2 \pm 1.5\%$ (from -8.2% to -0.4%) for $\Delta^{17}O(NO_3^-)$, respectively (Fig. 2a and b), which were larger than the differences estimated by the TRANSITS model. Thus, the observed differences between the summer and winter NO_3^- isotopes were not solely explained by postdepositional alteration. Consequently, seasonal differences in $\delta^{15}N(NO_3^-)$ and $\Delta^{17}O(NO_3^-)$ likely reflect atmospheric changes.

The observed seasonal $\delta^{15}N(NO_3^-)$ trend (high in summer and low in winter) at the SE-Dome site was consistent with observations made at two coastal Arctic sites (Morin et al., 2008, 2012) but inconsistent with typical seasonal $\delta^{15}N(NO_3^-)$ values of aerosols in midlatitude regions that are high in winter and low in summer (Freyer, 1991; Freyer et al., 1996; Lim et al., 2022). Although the specific process has yet to be identified, the factors controlling high $\delta^{15}N(NO_3^-)$ values in the summer have been comprehensively reviewed (Jiang et al., 2024, and references therein). One possibility is that physicochemical transformations of NO₃⁻ related to temperature influence $\delta^{15}N(NO_3^-)$ values, as suggested by a strong correlation between high $\delta^{15}N(NO_3^-)$ values and summer air temperatures (Morin et al., 2008). Another possibility is the incursion of anthropogenic sources, as proposed by Morin et al. (2009), which is supported by indications that air parcels originating from regions with greater anthropogenic influence carry higher $\delta^{15}N(NO_3^-)$ values. This is supported by observational studies on atmospheric $\delta^{15}N(NO_3^-)$ (e.g., Vicars and Savarino, 2014) and the increased frequency of air masses originating from North America during summer compared to winter (Kahl et al., 1997). While definitive conclusions regarding these observations have not yet been determined, it is hypothesized that the observations may be influenced by a combination of factors, including NO_x sources,

gas–particle partitioning variability influenced by temperature (Freyer, 1991) and acidity (Geng et al., 2014), oxidation pathways (Walters et al., 2015a), and differences in transport efficiency and removal processes (Heaton, 1987; Beyn et al., 2014). Future studies should examine the differences in $\delta^{15}N(NO_3^-)$ variations between both the source and remote regions. Such comparative analyses could enhance the current understanding of the underlying processes that influence isotopic compositions in different geographical contexts.

The observed seasonal changes in $\Delta^{17}O(NO_3^-)$ (high in winter and low in summer; Fig. 2b) were consistent with typical seasonal variations in $\Delta^{17}O(NO_2^-)$ (e.g., Michalski et al., 2003). In summer, increased sunlight promotes the formation of HNO₃ via NO₂+OH reactions, leading to lower $\Delta^{17}O(NO_3^-)$ values. Conversely, in winter, N₂O₅ hydrolytic or NO3 radical pathways forming HNO3 in the presence of O_3 predominate and result in increased $\Delta^{17}O(NO_3^-)$ levels. Although this kind of seasonal variation in $\Delta^{17}O(NO_3^-)$ is well known, we confirmed the historic occurrence of similar seasonal variations in the atmosphere. Although the scope of the current study limits further discussion in this regard, future research should explore the differences in $\Delta^{17}O(NO_3^{-})$ between summer and winter during the preindustrial period, when anthropogenic contributions of NO_3^- were significantly lower.

4.3 Decadal variations in NO₃⁻ isotopes

As discussed in Sect. 4.1, the SE-Dome ice core recorded atmospheric NO₃⁻ deposition with minimal postdepositional effects. The decadal $\delta^{15}N(NO_3^-)$ trend obtained from ice cores in Greenland has been interpreted to indicate changes in the NO_x source (Hastings et al., 2009) and/or atmospheric acidity from the beginning of the industrial revolution to the present (Geng et al., 2014). In response to emission controls since 1975, reasonable changes in dominant NO_x emissions and the adoption of NO_x removal technology (such as threeway catalytic converters) are expected, which can affect the $\delta^{15}N(NO_3^-)$ values (e.g., Walters et al., 2015b). As for acidity, if $\delta^{15}N(NO_3^-)$ is primarily controlled by atmospheric acidity, as proposed by Geng et al. (2014), $\delta^{15}N(NO_3^{-})$ should have increased after approximately 1975 when the atmospheric acidity decreased (owing to SO2 emission controls; Hattori et al., 2021). However, the $\delta^{15}N(NO_3^-)$ values obtained herein did not increase until 2014, indicating that there are multiple factors controlling ice core $\delta^{15}N(NO_3^-)$ values. As this study only covers a relatively limited period (60 years), future studies should address and compare longer ice core records from different regions. Such comparisons would be beneficial for understanding the factors behind isotopic variations, thereby enabling more accurate interpretations of isotopic records reconstructed from ice cores.

The unusually high NO_3^- fluxes observed in the summer of 1987 (1.97 mmol $m^{-2} a^{-1}$) and the spring of 1992 $(1.38 \text{ mmol m}^{-2} \text{ a}^{-1})$ were also notable (Fig. 2c). The extent of forest fires in North America could be the primary driver of this phenomenon, based on coincident high NH_4^+ fluxes during these periods (Iizuka et al., 2018). In 1992, the Mount Pinatubo eruption may have influenced the observations, considering the high SO_4^{2-} concentration $(13.7 \,\mu\text{mol}\,\text{L}^{-1}; \text{ lizuka et al., 2018}).$ The $\delta^{15}\text{N}(\text{NO}_3^{-})$ values during the summer of 1987 (1.3%) and the spring of 1992 (4.8%) were relatively high compared with other years during which $\delta^{15}N(NO_2^-)$ was less than 0%. These higher $\delta^{15}N(NO_3^-)$ values may be related to biomass burning associated with forest fires (-4.3% to +7.0%; Chai et al., 2019). Stratospheric NO₃⁻ inputs may also have high δ^{15} N values, as observed in Antarctic aerosols (Savarino et al., 2007). However, the $\Delta^{17}O(NO_3^-)$ values in the summer of 1987 (28.2%) and the spring of 1992 (29.5%) were not clearly different from other years, which is not consistent with the high $\Delta^{17}O(NO_3^-)$ trend during preindustrial biomass burning (i.e., forest fires) reported in the previous study (Alexander et al., 2004). We also note that no biomass-burning tracers were detected in 1987 or 1992 (Parvin et al., 2019). Further research is therefore required to link nitrate isotopes with specific events such as biomass burning.

The $\Delta^{17}O(NO_3^-)$ record from the SE-Dome core did not exhibit clear trends over the past 60 years. During this period, changes in atmospheric oxidants have occurred, such as increases in tropospheric O₃ over Arctic regions (Law et al., 2023). It is reasonable to estimate that higher O_3 can induce increases in $\Delta^{17}O(NO_3^-)$ by (1) promoting NO₂ formation from $NO + O_3$ reactions and (2) promoting NO_2 oxidation to NO₃ (and subsequently to HNO₃) by O₃. However, such changes were not recorded in the $\Delta^{17}O(NO_3^-)$ data from the SE-Dome ice core. During this period, atmospheric sulfate formation was changed by the promotion of in-cloud $S(IV) + O_3$ reactions, based on increases in $\Delta^{17}O(SO_4^{2-})$ from the same SE-Dome ice core (Hattori et al., 2021). Thus, further research is required to determine the mechanism(s) behind the observed constant $\Delta^{17}O(NO_3^{-})$ values in ice cores after emission controls by comparing $\Delta^{17}O(NO_3^{-})$ values estimated using chemical transport models such as GEOS-Chem (Alexander et al., 2009, 2020). Based on the $\Delta^{17}O(NO_3^{-})$ values recorded in the GISP2 ice core, the variations have been attributed to the intricate BrONO₂ hydrolysis mechanism, which extends beyond the small fluctuations in the $O_3/(HO_2 + RO_2)$ ratio in the relatively colder climate of a glacial period (Geng et al., 2017). Thus, reactive-halogen chemistry may also be a factor that impacts changes in the atmospheric oxidation capacity, specifically in high-latitude regions in the Northern Hemisphere.

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4.4 Comparison with other ice core data

The $\delta^{15}N(NO_3^-)$ value of ice cores collected in Lomonosovfonna, Svalbard, was $-6.9 \pm 1.9 \,\%$ after 1950 (Vega et al., 2015), which is lower than that at the SE-Dome and Summit sites. Given that snow accumulation at Lomonosovfonna $(0.55 \pm 0.1 \text{ m w.e. a}^{-1}; \text{ Vega et al., 2015})$ was higher than at Summit, these low $\delta^{15} \text{N(NO}_3^{-})$ values may reflect less postdepositional alteration. Notably, the relatively low $\delta^{15}N(NO_3^-)$ values at Lomonosovfonna and the SE-Dome site were consistent with the low $\delta^{15}N(NO_3^-)$ values in aerosols observed at two Arctic stations (Morin et al., 2008, 2012). Notably, the SE-Dome site and Svalbard both had lower $\delta^{15}N(NO_3^-)$ values and higher accumulation rates than the Summit site. Additionally, Svalbard is closer to Europe than Greenland, which may indicate a regional source difference. Ice core $\delta^{15}N(NO_3^-)$ records reported from Lomonosovfonna also exhibited decreasing trends until the 1970s, whereas an increase in $\delta^{15}N(NO_3^-)$ was only observed at Lomonosovfonna after the 1990s (Vega et al., 2015). Such differences may be attributed to differences in NO_x sources and spatial chemistries in the Arctic, although it is unclear whether this difference was caused by anthropogenic sources, natural sources, transport, or a combination of these factors.

The $\delta^{15}N(NO_3^-)$ records in ice cores from the Tibetan Plateau also exhibit decreasing trends from 1955 to 2011 (Li et al., 2020). The $\delta^{15}N(NO_3^-)$ of this Tibetan Plateau ice core $(4.2 \pm 3.1\%$ in 1951–2011) is also substantially higher than those of Arctic ice cores, indicating a different regional context. Comparing $\delta^{15}N(NO_3^-)$ records from different locations would be beneficial for determining the regional physical or chemical behaviors of NO_3^- from emission to deposition. This would allow us to better assess the impacts of human activity on nitrogen cycling and take corresponding measures to reduce the adverse effects of NO_3^- on climate and biogeochemical cycles. However, in Antarctica (Akers et al., 2022), the ice core $\delta^{15}N(NO_3^-)$ values varied significantly depending on the snow accumulation rate. It is therefore important to estimate postdepositional alteration for each ice core, refine models with recent information (Shi et al., 2023), and perform reverse calculations for atmospheric $\delta^{15}N(NO_2^-)$ (Jiang et al., 2024). The TRANSITS model used in this study is effective for assessing the sensitivity of nitrate preservation and isotopic compositions to snow accumulation within the same environment. However, as discussed, the archived isotopic composition depends on the f_{exp} value, which varies spatiotemporally. Therefore, efforts should be made to update models that account for the photolysis of snow $NO_3^$ and the recycling and redistribution of reactive nitrogen in boundary layer chemistry within a global chemical transport model (Zatko et al., 2016) by incorporating δ^{15} N information. Additionally, a recent study emphasized the potential impact of microbial alterations on both NO₃⁻ concentrations and its isotopic compositions in an Asian glacier (Hattori et al., 2023); thus, interpretations of NO₃⁻ concentrations and $\delta^{15}N(NO_3^-)$ records in ice cores should proceed with caution. We recommend that interpretations of NO₃⁻ concentrations and $\delta^{15}N(NO_3^-)$ records in ice cores should be accompanied by $\Delta^{17}O(NO_3^-)$ or $\delta^{18}O(NO_3^-)$ records whenever possible to verify atmospheric NO₃⁻ preservation without postdepositional biological alteration.

5 Conclusions

In this study, we reported \sim 60-year (1959–2014) records of NO_3^- isotopic compositions from the SE-Dome ice core in Greenland. The observed $\delta^{15}N(NO_3^-)$ values in the SE-Dome ice core were consistently $\sim 4\%$ lower than those in the Summit ice core record. The high snow accumulation rate at the SE-Dome site reduces the sensitivity of NO_3^- to postdepositional processes, which was supported by outputs from the TRANSITS model. Therefore, we concluded that the SE-Dome ice core, which exhibits superior NO_3^- preservation, is a promising tool for reconstructing changes in atmospheric nitrogen cycling driven by anthropogenic activity. This study was based on results from the SE-Dome I ice core $(\sim 90 \,\mathrm{m})$, which covers the past 60 years. The SE-Dome II core (drilled in 2021) preserves records that extend back to 1800 CE (Iizuka et al., 2021; Kawakami et al., 2023). Thus, there is considerable potential for future research aimed at reconstructing NO₃⁻ aerosol dynamics from the beginning of the industrial revolution to the present. Additionally, while regional comparisons of ice core $\delta^{15}N(NO_3^-)$ records are beneficial for describing the regional physicochemical behaviors of NO₃⁻, it is necessary to account for regional differences in postdepositional alteration when analyzing the spatiotemporal variations in atmospheric NO_3^- isotopes.

Data availability. The data used in this study are available from https://eprints.lib.hokudai.ac.jp/dspace/handle/2115/94240 (last access: 18 March 2025) or https://doi.org/10.14943/hu94240 (Wei et al., 2025).

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