Nitrate in Greenland and Antarctic ice cores: a detailed description of post-depositional processes

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ABSTRACT. A compilation of nitrate (NO₃⁻) data from Greenland has shown that recent NO₃⁻ concentrations reveal a temperature dependence similar to that seen in Antarctica. Except for sites with very low accumulation rates, lower temperatures tend to lead to higher NO₃⁻ concentrations preserved in the ice. Accumulation rate, which is closely linked to temperature, might influence the concentrations preserved in snow as well, but its effect cannot be separated from the temperature imprint. Processes involved in NO₃⁻ deposition are discussed and shown to be temperature- and/or accumulation-rate-dependent. Apart from scavenging of nitric acid (HNO₃) during formation of precipitation, uptake of HNO₃ onto the ice crystal's surface during and after precipitation seems to contribute further to the NO₃⁻ concentrations found in surface snow. Post-depositional loss of NO₃⁻ from the top snow layers is caused by release of HNO₃ and by photolysis of NO₃⁻. It is suggested that photolysis accounts for considerable losses at sites with very low accumulation-rate dependence is quantified, it should be possible to infer changes in atmospheric HNO₃ concentrations.

INTRODUCTION

The nitrate (NO_3^{-}) record in polar ice cores is expected to contain information about past atmospheric concentrations of nitrogen oxides $(NO_x = NO + NO_2)$ due to the close link between NO_3^- and NO_x . The increase in atmospheric NO_x concentrations in the Northern Hemisphere caused by rising fossil-fuel combustion since approximately 1940, for example, is reflected in higher NO3⁻ concentrations in Greenland snow (Neftel and others, 1985). However, past studies have shown that factors other than atmospheric NO_x concentrations also influence NO₃⁻ records (Wolff, 1995). In Greenland as well as in Antarctica, reversible deposition of NO3⁻ and net losses in the top snow layers have been observed. It has been suggested that either re-evaporation of nitric acid (HNO₃) (Dibb and others, 1998; Mulvaney and others, 1998; Röthlisberger and others, 2000a) or photolysis of NO_3^- in the top few centimetres of the snowpack (Honrath and others, 2000; Jones and others, 2000) causes such post-depositional alterations.

In earlier studies, relationships between NO_3^- and accumulation rate have been proposed (Herron, 1982; Legrand and Kirchner, 1990; Yang and others, 1995), with generally higher accumulation rates associated with lower NO_3^- concentrations

and higher NO_3^- depositional fluxes. In a more recent study based on a macroscopic deposition model, Fischer and others (1998) found a second-order polynomial dependence of average firm concentration and inverse snow accumulation.

Based on a compilation of NO_3^- data from more than 50 Antarctic sites covering various temperature and accumulation regimes, it has recently been suggested that temperature is also a key parameter in defining NO_3^- concentrations in Antarctic snow and ice, with lower temperatures leading to higher NO_3^- concentrations preserved in the snow (Röthlisberger and others, 2000a). Elevation has also been linked to NO_3^- concentrations, for both Greenland and Antarctica (Mulvaney and Wolff, 1994; Yang and others, 1996), but no statistically significant relationship has been found in a more recent study (Kreutz and Mayewski, 1999). The inherent connection between temperature, accumulation rate and elevation makes it difficult to distinguish between the separate effects, and no firm conclusions about their relative importance have yet been reached.

Here, we provide a detailed description of the processes involved in NO_3^- re-emission and a discussion of how temperature and accumulation rate affect those processes. In analogy to the compilation of NO_3^- data from Antarctic

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Table 1. Temperature, accumulation rate and NO₃ concentration for Greenland locations used in this study

Site (source)	Location	Altitude	Temperature	Accumulation	NO ₃ ⁻ pre-1940		NO ₃ post-1970	NO ₃ ⁻ post-1970	
		m	°C	$g \mathrm{cm}^{-2} \mathrm{a}^{-1}$	Concentration	Number of years	Concentration	Number of years	
		III	u	gem a	ppo		pp		
1988-8 ¹	$64.6^\circ\mathrm{N}, 43.7^\circ\mathrm{W}$	2550	-19.9	57			94	2	
$1988-7^{1}$	$64.8^{\circ}\mathrm{N}, 44.1^{\circ}\mathrm{W}$	2745	-22.0	38			90	3	
Dye $318C^{2}, 20D^{3}$	$65.0^{\circ}\mathrm{N}, 44.9^{\circ}\mathrm{W}$	2617	-22.3	41	53,55	$174^{\rm a}$, $30^{\rm c}$	100,122,132	15,5,15	
Dye 3 ^{2,4}	65.2° N, 43.8° W	2491	-20.0	49	44,52,52	10 ^b ,11 ^b ,220 ^h	86,114,137	6,7,2	
$1988-4^{1}$	$66.6^{\circ}\mathrm{N}, 45.5^{\circ}\mathrm{W}$	2130	-18.6	30			117	2	
1988-5 ¹	67.0° N, 44.5° W	2250	-20.3	33			115	4	
1988-6 ¹	67.2° N, 43.8° W	2450	-21.6	36			108	6	
EGIG $T05^5$	69.9° N, 47.3° W	1910	-18.0	46			118	5	
$EGIG T09^5$	70.0° N, 46.3° W	2170	-20.0	41			134	9	
EGIG T13 ⁵	70.2° N, 45.0° W	2380	-22.6	46			107	8	
Milcent ²	70.3° N, 44.6° W	2410	-22.0	49	61	7^{g}			
EGIG T 17^5	70.4° N, 44.1° W	2530	-23.8	44			117	9	
EGIG T 21^5	70.6° N, 43.0° W	2700	-24.8	44			112	9	
Site A ^{2, 6}	70.6° N, 35.8° W	3092	-29.4	29	45,64,70	$3^{d}, 2^{c}, 19^{b}$			
Site D^6	70.6° N, 39.6° W	3018	-28.3	34	73	19 ^b			
EGIG T 27^5	70.8° N, 41.6° W	2870	-26.6	39			113	9	
EGIG T31 ⁵	70.9° N. 40.6° W	2970	-27.5	34			133	10	
EGIG T41 ⁵	71.1° N, 37.9° W	3150	-29.5	25			144	12	
Crête ²	71.1° N. 37.3° W	3172	-30.0	28	60.80	$60^{\rm e}.2^{\rm g}$			
EGIG T43 ⁵	71.1° N. 37.3° W	3172	-30.1	23)		144,146	8.14	
EGIG T47 ⁵	71.2° N. 36.0° W	3099	-29.9	22			143	8	
Renland ^{7,8}	71.3° N 267° W	2340	-18.0	42	64.88	$5^{\mathrm{g}} 9^{\mathrm{c}}$	110	Ũ	
FGIG T53 ⁵	71.4° N 32.5° W	2864	-28.0	23	01,00	0,0	134	9	
1987-9 ¹	71.6° N 381° W	3157	-30.4	25			128	5	
EGIG NST085	71.9° N 378° W	3220	-31.0	23			153	5	
1987-3 ¹	71.9° N 39.8° W	3145	-30.3	30			123	2	
1987-5 ¹	72.0° N 37.5° W	3187	-31.5	22			130	6	
FGIG T61 ⁵	72.0° N 39.3° W	9819	-28.8	10			144	8	
1987.1 ¹	72.2 IN, 52.5 W	2012	-31.3	19			194	10	
1097 1	72.5 IN, 57.5 W	2146	-21.5	22			124	5	
CISP9 ATM ¹	72.4° N 20.0° M	2200	22.0	27			190 120 125 129	2502	
FCIC T66 ⁵	72.F IN, 50.0 W	9679	-95.7	17			150	5,5,5,5	
CISP9 ^{1,9,10}	72.5 IN, 50.6 W	2070	-21.0	17	6170	$410^{a} 650^{a}$	109 195 196 125 126 140 140 167	207266994	
CP 1P ^{5,11,12}	72.0 IN, 30.5 W	3200	-21.6	22	69	410,050	120,129,130,139,149,149,149	50,7,5,0,0,2,2,4	
10.97 7 ^l	72.0 IN, 57.0 W	2100	-22.1	19	00	50	157	9,5,21	
1907-7 1000 9 ¹	72.0 IN, 35.5 W	3150	_22.9	10			137	1	
1990-2 1097 6 ¹	72.0 IN, 50.5 W	2004	22.0	17			14.9	0	
1967-0 NCT01 ¹³	73.0 IN, 37.7 W	2002	-33.6	17			140	0	
NCT02 B16 ¹³	73.0° N 276° W	2040	-34.6	10	95	40^{a}	129	13	
North Control ²	73.5 IN, 57.0 W	2020	22.0	12	03	of orb	152	23	
NCT05 ¹³	74.0 IN, 59.0 W	2930	-32.0	13	65,112	9,21	199	19	
NCP 10 ^{8,14}	75.1° N 49.1° W	2075	30.0	17	01	156^{a}	140161	5.11	
NGKIF NCTI0 ¹³	75.1 IN, 42.1 W	2970	-30.9	17	01	150	149,101	J,11	
NG112 NCT14 P10 ¹³	75.7 IN, 50.4 W	2071	-34.2	10	72	40^{a}	101	14	
$\Gamma G I I 4 D I 6$	70.0 IN, 30.4 W	2008	-55.0	10	/ 3 46 65 71	40	120	20	
Camp Century	77.2 IN, 01.1 W	1000	-24.4	30	40,03,71	10°,15',100	120	2	
1988-1 1000 02 ¹	77.2 N, 60.7 W	1000	-24.6	29			101	/	
1988-2,5	77.2 IN, 39.2 W	1/00	-23./	31 95	5.0	10b	90,107	4,2	
Camp Century II ⁺	77.2° N, 60.8° W	1910	-24./	35	52	13 ⁻	110	0.4	
NGT37 B26	77.3° N, 49.2° W	2598	-30.0	18	69	440	110	24	
NGTI8"	77.5° N, 36.4° W	2319	-32.6	11	6.2	110.03	158	14	
NGT23 B20 ¹⁰	/8.8° N, 36.5° W	2147	-31.5	10	83	1130"	155	24	
NGT27 B21 ¹⁵	80.0° N, 41.1° W	2185	-29.6	11	80	40 ^a	126	24	
Hans lausen"	80.5° N, 37.5° W	1271	-21.0	10	/4	30°	91	3	

Notes: ¹Yang and others (1996), data obtained from http://nsidc.org. ²Clausen and Langway (1989). ³Mayewski and others (1990). ⁴Neftel and others (1985). ⁵Fischer and Wagenbach (1996). ⁶Steffensen (1988). ⁷Hansson (1994). ⁸H. B. Clausen, personal communication (2001). ⁹Yang and others (1995). ¹⁰University of Arizona, unpublished data. ¹¹Clausen and others (1997). ¹²Steffensen and others (1996). ¹³Fischer and others (1998). ¹⁴L. B. Larsen, personal communication (2001). ¹⁵Hausbrand (1998). ¹⁶Bigler (2000). ^aContinuous record up to 1940. ^b20th century, before 1940. ^c19th century. ^d18th century. ^e11th to 18th century. ^f15th century. ^g13th century. ^hHolocene.

sites, NO₃⁻ data from many Greenland sites have been gathered in order to illustrate the effect of temperature. Also, we outline how calcium (Ca²⁺) can inhibit NO₃⁻ re-emission.

Another aspect that has been discussed lately is the effect on $\mathrm{NO_3^-}$ concentrations of snow layers containing large amounts of sulphuric acid (H₂SO₄) of volcanic origin. Most studies focused on a few well-known volcanic eruptions during the Holocene which showed post-depositional dis-

placement of NO_3^- away from the H_2SO_4 peak. This behaviour has been found in single events in Greenland and Antarctica (Legrand and Kirchner, 1990; Clausen and others, 1997; Röthlisberger and others, 2000a). However, the mechanisms leading to this effect are only vaguely understood, and hypotheses have not been tested on a large number of cases, because of a lack of sufficient high-resolution data.

In this paper, new high-resolution data from the North-



Fig. 1. Spatial distribution of NO_3^- across Greenland. Concentrations in snow deposited before 1940 are shown in parentheses; the other values correspond to concentrations in snow deposited after 1970. All concentrations are in ppb.

GRIP (North Greenland Ice Core Project) ice core are used to investigate in more detail the effect of volcanic H_2SO_4 on NO_3^- . Volcanic events from the early Holocene, the Last Glacial Maximum (LGM) and some earlier glacial periods are compared and a hypothesis of the mechanism is given.

DATA

Many of the NO₃⁻ data used in this paper are compiled from earlier studies (see Table 1 for sources). Furthermore, data from selected sections of the NorthGRIP ice core (75.1° N, 42.05° W; 2978 m a.s.l.) are presented. These sections were analyzed during the NorthGRIP 2000 field season with a continuous flow analysis (CFA) system, as described in Röthlisberger and others (2000b). Among other compounds, nitrate (NO₃⁻), sulphate (SO₄²⁻) and calcium (Ca²⁺) have been measured at a resolution of approximately 1 cm.



Fig. 2. NO_3^- concentrations vs average temperatures in Greenland and Antarctica with linear trends (triangles: post-1970; squares: pre-1940; solid lines: Greenland; dots and dashed line: Antarctica). The data points at temperatures below $-52^{\circ}C$ correspond to sites with very low accumulation rates and are not used for the calculation of the linear fit shown.



Fig. 3. NO_3^- concentrations vs accumulation rate in Greenland and Antarctica (triangles: post-1970; squares: pre-1940; dots: Antarctica).

$\mathrm{NO}_3^-,$ TEMPERATURE AND ACCUMULATION RATE

In Figure l, average NO_3^- concentrations for different Greenland locations are indicated. In order to account for the anthropogenic increase in NO_3^- concentrations in Greenland, the data have been split into two separate sets, one indicating concentrations from before 1940, which are unaffected by the anthropogenic emissions, and one from after 1970. The data and their sources are listed in Table 1. For both pre-1940 and post-1970, a decreasing trend in NO_3^- concentration with increasing temperature is found (Fig. 2). A similar trend has been found in Antarctica (Röthlisberger and others, 2000a),



Fig. 4. Accumulation rate vs average temperatures in Greenland and Antarctica (squares: Greenland; dots: Antarctica).

where also generally higher concentrations are found at sites with lower temperatures, except for very low-accumulation sites (e.g. Dome C). At Dome C, the concentrations in the top few centimetres can be very high (up to 1000 ppb), but at greater depths, only 15 ppb are preserved, despite the low temperatures (annual mean temperature $\approx -54^{\circ}$ C).

When NO₃⁻ concentrations are plotted against accumulation rate, higher NO₃⁻ concentrations are found at sites with lower accumulation rates (Fig. 3), but again, NO₃⁻ concentrations at very low-accumulation sites do not agree with the general trend. Considering the close relationship between temperature and accumulation rate (Fig. 4), it seems difficult to separate their effects on NO₃⁻ concentrations. Only a detailed consideration of the underlying microphysical processes allows for assigning a temperature or accumulationrate dependence.

In the following discussion, we analyze the temperature and accumulation-rate dependence of processes involved in NO_3^- deposition and re-emission, aiming for a better understanding of:

the cause of the relationship between NO_3^- concentrations and mean annual temperature and accumulation rate at a site

the generally higher $\mathrm{NO_3}^-$ levels in summer snow than in winter snow

the net loss of NO_3^- from snow after deposition.

 $\rm NO_3^-$ can either be predominantly incorporated in the bulk or be adsorbed to the surface of a snow crystal, depending on the deposition pathway. In a cloud with a liquid-water content of >0.01 g m⁻³ and pH > 1, HNO₃ would be completely dissolved in water droplets due to its high solubility, leaving virtually no HNO₃ in the gas phase (Seinfeld and Pandis, 1998). Thus, in the case of liquid or mixed clouds, essentially all HNO₃ is removed from the gas phase independent of the cloud temperature. While there is no specific information about the conditions at cloud level, typical liquid-water contents of $0.1 \, {\rm g m^{-3}}$ and initial HNO₃ concentrations of 20 pptv in the air would lead to NO₃⁻ concentrations of

Table 2. Estimates of $NO_{\mathcal{S}}^{-}$ concentrations in snow due to different deposition mechanisms

	Sur Summer	nmit Winter	Neur Summer	nayer Winter	Sout Summer	h Pole Winter
Pressure (Pa)	67 0 00		99 000		68 000	
Accumulation rate $(\mathbf{g} \mathbf{cm}^{-2} \mathbf{a}^{-1})$	20		34		8.5	
Temperature (K)	259	230	269	247	246	208
Relative humidity	0.75	0.75	0.8	0.8	0.75	0.6
HNO ₃ atm. (pptv)	20	1	6	1	20	1
NO ₃ snow (ppbw)	120		50		100	
Co-condensation (ppbw)	7	9	1	2	26	160
Solubility in ice (ppbw)	19	29	8	12	40	150
Surface uptake (ppbw)		1900		780	840	3300
Dry deposition (ppbw)	30	2	8	1	77	5

Note: The meteorological data are from automatic weather stations; the atmospheric HNO3 summer concentrations are from Dibb and others (1994) and Jones and others (1999). For South Pole, atmospheric concentrations of the same order of magnitude as in Summit have been assumed. The HNO3 winter concentration has only been measured at Neumayer and has been found to be approximately 1 pptv (personal communication from R. Weller, 2001). In the absence of measurements for the other sites, we assume the winter concentrations at Summit and South Pole are the same as in Neumayer. The calculations for co-condensation and solubility in ice are based on Thibert and Dominé (1998). Surface uptake was estimated using a linear regression through the values for temperaturedependent uptake found by Abbatt (1997) and assuming a typical surface area of $4000 \text{ m}^2 \text{ m}^{-3}$ (Narita, 1971). Temperatures higher than 248 K have not been included in the study of surface uptake, so the estimates for surface uptake in Summit and Neumayer during summer are missing. Dry deposition was calculated assuming a dry-deposition velocity of 0.5 cm s (Hauglustaine and others, 1994).

approximately 350 ppbw in fresh snow. On the other hand, the co-condensation of HNO_3 and water (H_2O) molecules on ice crystals (Thibert and Dominé, 1998) would lead to a bulk concentration of 20 ppbw only. In the absence of liquid water, i.e. in ice clouds, the high NO₃⁻ concentrations found in surface snow could not be explained. However, Abbatt (1997) observed a temperature dependence of adsorption of HNO3 on ice surfaces with higher uptake at lower temperatures. For typical summer temperatures at South Pole (246 K; data obtained from http://www.cmdl.noaa.gov), the uptake capacity on fresh snow crystals exceeds the amount of HNO3 available in the cloud, implying that at very cold temperatures where ice clouds predominate, essentially all HNO3 in a cloud is bound to the surface of the snow crystal. While co-condensation, riming and adsorption of HNO₃ determine the distribution of NO_3^- within the ice crystal, its concentration is defined by the initial atmospheric concentration of HNO₃ and the amount of condensed water in the cloud. An imprint of temperature is expected for surface uptake and co-condensation, but not for the HNO₃ taken up in liquid cloud droplets. Based on the results of Abbatt (1997), the snow crystals are expected to efficiently scavenge atmospheric HNO3 on their way to the surface, potentially further increasing the NO_3^{-} concentration of the fresh snow.

Once on the ground, the formation of surface hoar frost (co-condensation), rime (deposition of supercooled fog droplets) as well as dry deposition (adsorption of HNO₃ onto the crystal's surface) leads to additional NO₃⁻ deposition to surface snow. For a given atmospheric HNO₃ concentration, the hoar-frost NO₃⁻ concentration is determined by the water-



Fig. 5. Examples of the influence of volcanic H_2SO_4 on NO_3^- in the NorthGRIP ice core. H^+ concentrations have been inferred from electrical conductivity measurements on the solid ice. Due to very low H^+ concentrations, the scale of the y axis had to be adjusted for the section shown in (b). Of 28 events selected with SO_4^{-2} concentrations exceeding 20 μeqL^- , 14 showed a pattern similar to (a), 12 similar to (b), and 2 similar to (c). The data are shown against depth, as the absolute age is not critical to the illustration of the effect and an absolute time-scale is not yet available.

vapour concentration (Thibert and Dominé, 1998), which is mainly a function of temperature T. Assuming that the relative humidity at the site is similar throughout the year, the NO₃⁻ concentration should depend linearly on l/T in a first-order approach. However, assuming that the HNO3 concentrations in the atmosphere as in Table 2 are representative for the site, the estimated concentrations resulting from co-condensation are lower than observed surface snow concentrations, thus leading to dilution of the surface snow (Table 2). Rime deposition, on the other hand, which is likely to remove all HNO₃ from the air, shows concentrations similar to those in fresh snow, provided that the atmospheric HNO₃ concentration is similar to that at cloud level. A net dry deposition of HNO3 (adsorption of HNO3 onto snow crystals) has the potential to increase the NO_3^{-} concentration in snow. However, it will only contribute where the surface is undersaturated, i.e. at very cold sites and during winter. The contribution in winter is small, due to low atmospheric HNO₃ concentrations. During the summer, the contribution of dry deposition to the NO₃⁻ concentrations in snow at South Pole might be considerable. However, according to Hauglustaine and others, (1994), the dry-deposition velocity of 0.5 cm s⁻¹ for HNO₃ on snow has to be considered as an upper limit. At sites with higher accumulation rates, a specific surface snow layer is buried more rapidly, leaving less time to adsorb additional HNO3 from the atmosphere, given that no saturation has been reached. Therefore, a tendency to higher concentrations at lower accumulation rates is expected.

Besides the diluting effect of co-deposition, processes capable of reducing the NO_3^- concentration in snow are desorp-

tion of HNO₃ from the snow crystal (Dibb and others, 1998; Mulvaney and others, 1998; Röthlisberger and others, 2000a) and photolysis (Honrath and others, 2000; Jones and others, 2000). In both cases, the NO_3^- ion has to be at the surface of a snow crystal, since photolysis of NO₃⁻ in the bulk is not effective (Dubowski and others, 2001). If a NO_3^- ion recombines with a H^+ ion to form HNO₃, it then may desorb into the firm air and eventually diffuse into the air above the snow. Diffusion of NO₃⁻ in ice has been investigated by Thibert and Dominé (1998), who found that diffusion of NO_3^{-} in ice is slower at colder temperatures, with the diffusion coefficient D given by $D = 1.37 \times 10^{-2610/T}$ cm² s⁻¹, with T being the temperature in K. During the summer, the typical time a NO₃⁻ molecule needs to reach the ice surface (diffusion length of $40\,\mu\text{m}$, corresponding to an average crystal radius (Harder and others, 1996)) is of the order of a couple of hours (e.g. Neumayer) to a few days (e.g. South Pole).

The solubility of NO_3^- in ice has been determined for various temperatures (Thibert and Dominé, 1998). For summer conditions, not only the surface snow but also deeper layers are supersaturated with respect to the solubility of NO_3^- in ice (Table 2). In the case of South Pole, the surface remains undersaturated, suggesting that NO_3^- that is expelled from the bulk is simply transferred to the surface without affecting the NO_3^- concentration. At warmer sites (e.g. Neumayer or Summit), the ice surface also might become supersaturated during the summer, leading to release of HNO₃ from the snow into the interstitial air. From a thermodynamic point of view, warmer temperatures during the summer should facilitate the release of HNO₃ from the ice surface. But according to the uptake experiment of Abbatt (1997), only up to 25% of the

HNO₃ that has been taken up is released again afterwards. However, his study emphasized the uptake rather than the release of HNO₃, and a temperature dependence of the release has not been discussed. Once released from the snow crystal's surface, the HNO₃ molecule might make its way out of the snowpack, leading to a net loss of NO₃⁻⁻ in the snow. The molecular diffusion of HNO₃ in the interstitial air is temperature-dependent as well, but probably this is not the limiting factor controlling the transfer out of the snowpack. It is conceivable that temperature-dependent, repeated adsorption and subsequent desorption of a HNO₃ molecule on ice crystals will determine the removal from the snowpack.

Photolysis of NO₃⁻ in the top snow layers results in the production of nitrogen dioxide (NO_2) and hydroxyl radical (OH). NO₂ is expected not to interact with the surrounding snow but to be mixed into the boundary layer rather quickly, leading to a NO_3^{-} depletion in surface snow. The influence is presumably largest at low-accumulation sites, where surface snow is exposed to sunlight for a long time. Furthermore, photolysis should become more efficient at lower latitudes due to more incoming ultraviolet radiation. At Dome C, the top few centimetres of snow seem to reach saturated surface coverage (concentrations in the range of several hundred ppbw) (Röthlisberger and others, 2000b). Deeper layers are then dramatically undersaturated considering surface coverage and solubility in the bulk. No quantitative estimate has yet been made of how much NO_3^- can be lost by photolysis. It therefore remains unclear whether photolysis alone can account for the NO_3^- profile seen in the snow at Dome C. The estimate of the maximum surface uptake relies very much on the surface area in snow and might change due to recrystallization, which has not yet been taken into account.

Although elevation may affect the atmospheric HNO₃ concentration, there is no obvious mechanism by which it can have a direct physical influence on post-depositional processes. A minor influence is expected on the photolysis rate due to changes in the irradiance with altitude and on the gas-phase diffusion due to lower pressure, but both effects might only slightly modulate the changes in NO₃⁻ concentrations.

INTERACTIONS OF NO3⁻ WITH DUST

Recent studies have reported that NO_3^- and Ca^{2+} concentrations are correlated in ice from the last glacial period from Vostok and Dome C, Antarctica (Legrand and others, 1999; Röthlisberger and others, 2000a). During the last glacial period, the Ca^{2+} deposited on the East Antarctic plateau is mainly of terrestrial origin and can thus be used as a proxy for dust. It has been suggested that the reaction of HNO_3 and $CaCO_3$ to form $Ca(NO_3)_2$ prevents NO_3^- from being released from the snow into the gas phase.

In principle, the reaction could take place in the atmosphere or in the snow. For it to occur in the atmosphere, high dust concentrations would need to be in the same season as the maximum nitrate concentrations. A recent paper that studied the reaction of CaCO₃ with HNO₃ derived a formula for the lifetime for removal of HNO₃ by dust (Hanisch and Crowley, 2001). Based on this, and using very rough estimates for the surface area of dust (assuming spherical particles of 1 μ m diameter, a typical density of 2 g cm⁻³, and atmospheric dust concentrations of 10 ng m⁻³), we can estimate a HNO₃ lifetime vs removal by dust in the present-day Antarctic

atmosphere of around 50 days. This is unlikely to be important relative to other removal processes. However, this could be reduced to 2 days under the dustier conditions of the LGM, and in Greenland under LGM conditions one could estimate a lifetime for this removal of only a few hours. It seems possible therefore that under LGM conditions much of the atmospheric HNO₃ could be converted to aerosol calcium nitrate in the atmosphere. In addition, the reaction to form $Ca(NO_3)_2$ might take place in the snow, in which case HNO₃ has to make its way to the snow layer where the Ca^{2+} has been deposited and, if Ca^{2+} is inside the snow grain rather than on its surface, diffuse through it.

INFLUENCE OF VOLCANIC H₂SO₄ ON NO₃⁻

As shown lately in an Antarctic high-resolution record from Dome C (Röthlisberger and others, 2000a), H_2SO_4 of volcanic origin can cause NO_3^- to move in the ice. Several examples of very low NO_3^- concentrations coinciding with H_2SO_4 peaks and increased NO_3^- concentrations above and below this layer have been found in the Dome C as well as the North-GRIP record (Fig. 5a). At Dome C, the effect is first seen at 12 m depth, where the deposits of the Tambora (Indonesia) eruption (AD 1815) are located. This indicates that the processes involved take place or at least start in the firn.

Our hypothesis of the mechanism for NO_3^- displacement in the firn relies on processes similar to those described above, as it includes diffusion of HNO₃ in the firn air. The high concentration of H_2SO_4 in a volcanic layer causes the equilibrium of $H^+ + NO_3^- \rightleftharpoons HNO_3$ to shift towards the righthand side, as a large amount of H^+ from the H_2SO_4 is present. Therefore, a locally elevated concentration of HNO₃ in the firn air is caused, which prompts gas-phase diffusion of the HNO₃ away from the volcanic layer towards either side. In an adjacent layer with no excessive H_2SO_4 , HNO₃ favours dissociation into H^+ and NO_3^- , thus maintaining the gradient in the firn air by removing HNO₃ from the gas phase. It is possible that the movement of NO_3^- progresses via a different mechanism in the ice after pore close-off, possibly by diffusion of ions in the veins.

Figure 5b shows an event where no NO_3^- displacement occurred. In this section, large amounts of alkaline material are present in the ice, as seen by the high Ca^{2+} and negligible H^+ concentrations. In the context of the above hypothesis, the high concentrations of alkaline material (presumably $CaCO_3$) neutralize the H_2SO_4 and prevent the formation of HNO_3 . The assumption that in this case H_2SO_4 undergoes some reactions is supported by the significantly narrower SO_4^{2-} peak compared to the ones seen in acid ice (personal communication from P. Barnes, 2001).

Our hypothesis is challenged by the example shown in Figure 5c. Although much excess H_2SO_4 is present and far too little Ca²⁺ to compensate, no marked NO₃⁻ displacement is seen. It is possible that the Ca²⁺ was unable to neutralize the H_2SO_4 , but that it managed to bind NO₃⁻, thus preventing it from being transferred into the gas-phase. The occurrence of ice layers limiting HNO₃ diffusion in the interstitial air is rather unlikely in NorthGRIP.

CONCLUSIONS

The spatial distribution of nitrate concentrations in Greenland is shown to be strongly related to site temperature, just as it is for Antarctica. Because temperature and snow-accumulation rate are so closely linked, we cannot determine which of these factors is the one exerting physical control on the concentrations seen. In either case, the relationship changes at the very lowest accumulation rates, where it is clear that post-depositional losses are the dominant control on the subsurface concentration.

By examining the individual processes that could contribute to the nitrate concentration in snow, we find that many of them are indeed temperature-dependent, with higher concentrations predicted at lower temperatures, as observed. Some processes could also depend on the accumulation rate, if a longer exposure time at the surface allows additional uptake. Of the processes identified, either uptake by liquid droplets in cloud, if present, or uptake onto the ice surface in the cloud or after deposition can lead to concentrations in fresh snow that are as high as or higher than those observed. However, the role of surface uptake at higher temperatures, as encountered in coastal Antarctica and in Greenland in summer, needs to be quantified. Co-condensation of nitric acid and water, and dissolution of nitric acid within the ice lattice appear to give concentrations that are too low compared to those observed. This suggests tentatively that surface uptake and retention might be rather important in determining the concentrations we see. This process has a temperature dependence (approximately threefold greater uptake at -55°C compared to -25° C (Abbatt, 1997)) similar to that seen in Figure 2. However, according to Abbatt (1997), the uptake is not dependent on the nitric acid concentration in air. The higher nitrate concentrations in post-1970 Greenland snow compared to pre-1940 snow suggest that the snow concentration is somehow related to atmospheric concentrations, and we suggest that this is simply a question of limited supply to the ice surface, because nitric acid is scavenged so efficiently. In that case, a temperature- and accumulation-rate-corrected Holocene nitrate ice-core record of a site with adequate snow-accumulation rate should reflect the flux of nitrate to the surface, which should in turn be related to atmospheric NO_x input.

For sites with very low accumulation rate, losses, possibly due to photolysis, control the concentration seen in the Holocene in such a dominant way that it is unlikely that information about atmospheric nitrate or NO_x can be extracted. Once the ice becomes less acidic (in the last glacial period), the concentration of nitrate seems to be strongly controlled by the calcium or dust content of the atmosphere, and the deposition processes might be significantly altered. The ratio of nitrate to dust might give clues to the past nitrate content of the atmosphere, although in this case it is probable that the nitrate uptake is determined by the content of the atmosphere over the whole transport route of dust from its source to the deposition site, and is not closely related to the local nitrate concentration at the ice-core site.

A number of laboratory and field experiments would help to test the above hypotheses. Laboratory uptake experiments, similar to those carried out at 248 K and below (Abbatt, 1997), are needed at higher temperatures, appropriate to summer temperatures at coastal Antarctic or Greenland sites. Laboratory experiments would also allow an assessment of whether photolysis can account for the magnitude of nitrate losses at low-accumulation sites. Field measurements of the nitric acid content of the atmosphere are lacking for most sites, and particularly for winter. Experiments that follow the evolution of concentration from fresh snow to depth in individual layers are also required. In summary, the factors that control nitrate concentration in ice cores are complex, and interpretation is likely to involve different factors for different locations and time periods. However, if the factors controlling deposition and loss can be better understood, it may still be possible, in some cases, to reconstruct information about the important NO_x cycle in the past.

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