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Concentrations and snow-atmosphere fluxes of reactive nitrogen at Summit, Greenland

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Abstract. Concentrations and fluxes of NO_v (total reactive nitrogen), ozone concentrations and fluxes of sensible heat, water vapor, and momentum were measured from May 1 to July 20, 1995 at Summit, Greenland. Median NO_v concentrations declined from 947 ppt in May to 444 ppt by July. NO, fluxes were observed into and out of the snow, but the magnitudes were usually below 1 µmol m⁻² h⁻¹ because of the low HNO₃ concentration and weak turbulence over the snow surface. Some of the highest observed fluxes may be due to temporary storage by equilibrium sorption of peroxyacetylnitrate (PAN) or other organic nitrogen species on ice surfaces in the upper snowpack. Sublimation of snow at the surface or during blowing snow events is associated with efflux of NO_y from the snowpack. Because the NO_y fluxes during summer at Summit are bidirectional and small in magnitude, the net result of turbulent NO_v exchange is insignificant compared to the 2 µmol m⁻² d⁻¹ mean input from fresh snow during the summer months. If the arctic NO_y reservoir is predominantly PAN (or compounds with similar properties), thermal dissociation of this NO_v is sufficient to support the observed flux of nitrate in fresh snow. Very low HNO₃ concentrations in the surface layer (1% of total NO_v) reflect the poor ventilation of the surface layer over the snowpack combined with the relatively rapid uptake of HNO₃ by fog, falling snow, and direct deposition to the snowpack.

1. Introduction

Concentration records of NO₃ in snow and ice cores have been used to infer variations of atmospheric nitrogen loading and inputs from anthropogenic NO_r emissions, biomass burning, solar activity, and stratospheric denitrification [Legrand and Delmas, 1986; Mayewski et al., 1990; Wolff, 1995]. Inferences from the ice-core nitrate record, however, have been called into question because the chemical and physical processes controlling transport and speciation of N in the atmosphere, deposition to snow, and ultimate preservation in glacial ice are not fully understood [Wolff, 1995]. Our ability to invert the ice-core nitrate record to infer global atmospheric nitrogen concentrations and emission rates depends on a quantitative understanding of which nitrogen species are transported to the Arctic, what processes control conversion and deposition of N, and how extensively post-depositional processes alter the record. The lifetimes for NO, species (we define $NO_x = NO + NO_2$; $NO_y = NO_x + HNO_3 + NO_3 + 2N_2O_5$ + peroxyacetylnitrate (PAN) + other organic nitrates + particulate nitrate) vary from a few days to deposit HNO₃ from

the boundary layer to several months to decompose PAN in the upper troposphere. The amount of NO, that reaches the polar regions depends strongly on the form of NO, that is transported, but the long-lived species, such as PAN, that are transported effectively may not deposit efficiently. Dibb et al. [1994] point out the dilemma that HNO₃ concentrations measured above Summit, Greenland, are very low, yet NO3 is the dominant anion in the snow. Furthermore, the ratio of NO₃ to SO₄² in snow at Summit exceeds the ratio of those ions (as HNO₃ and aerosol SO₄²) in air. They suggest that rapid dry deposition of HNO3 from a shallow boundary layer depletes the gas-phase HNO₃ or that species other than HNO₃ are deposited and converted to nitrate in the snowpack. Concentration profiles in the surface snow from Antarctica and Greenland indicate that NO₃ may migrate within the snowpack (distorting seasonal and interannual variations) or be returned to the atmosphere [Fischer et al., 1998; Fischer and Wagenbach, 1998; Mayewski and Legrand, 1990; Mulvaney et al., 1998; Neubauer and Heumann, 1988], particularly at low-accumulation sites. Volatilization of HNO₃ from snow is an apparent contradiction to laboratory studies showing irreversible uptake of HNO₃ on ice surfaces [Abbatt, 1997; Laird and Sommerfeld, 1995; Zondlo et al., 1997]. Better understanding of the processes controlling HNO₃ removal is needed to objectively distinguish long-term trends in NO₃ deposition [Mayewski et al., 1986; 1990] from postdepositional loss where both processes are important.

This paper presents results from the first measurements of total nitrogen oxides (NO_y) mixing ratios and eddy fluxes above the surface of a polar glacier at Summit, Greenland.

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The goals of this work were to determine the ambient mixing ratios of NO_v, to quantify the direction and magnitude of NO_v fluxes at the snow-atmosphere interface, and to identify processes that control exchange of reactive nitrogen with the snow surface. A previous paper [Dibb et al., 1998] presented a small subset of the 1995 NO_v data focusing only on periods with simultaneous measurement of HNO3 concentration gradients above the snow surface and NO_v. Nitric acid was 1% of the total NO_v. Measurable gradients of HNO₃ were observed and were frequently in opposition to the concurrent NO_v eddy flux, suggesting that N exchange with the snow was more complex than a simple adsorption/desorption of HNO₃. Large apparent deposition velocities for HNO₃ suggest that NO₃ species other than HNO₃ may be exchanging with the snowpack. In this paper we present the complete NO, concentration and flux results for the May-July 1995 summer field season, along with an examination of momentum, heat, and water vapor exchanges over the snowpack. The results in this paper overlap those presented by Dibb et al. [1998] but show different features due to periods that were not included in the previous analysis. We consider what nitrogen species are present, how they are transformed and deposited to the snowpack, and whether they are stable in the snow.

2. Methods

During the 1994 field season an 18 m tower was installed 270 m south of the main camp at Summit (elevation 3200 m, 38.4°E, 72.55°N). Measurements were made during the summer season in both 1994 and 1995; however, we will not discuss the 1994 data here because contamination by combustion sources in and near the camp affected too many of the data. Winds from the clean sector were more frequent and polluting activities were more effectively restricted during the 1995 field season. A three-axis sonic anemometer (Solent Research Anemometer, Gill Instruments) and sampling inlets for NO_v , H_2O , and O_3 measurements were mounted at 17.5 m facing the clean-air sector to the south and east. Instruments and data acquisition and control systems were housed in a covered snow trench at the tower base. Ventilation at the trench ceiling was required to discharge waste heat from the instrumentation and prevent the walls from melting. The instrument rack was enclosed to maintain a moderate operating temperature despite the room temperature being < 0°C. NO. was measured using reduction by H2 on heated gold followed by O₃-chemiluminescent detection [Bakwin et al., 1994; Munger et al., 1996]. The hot gold catalyst quantitatively converts all species of NO, to NO. The catalyst was mounted on the tower with no additional inlet to avoid inlet retention problems for species such as HNO₃. The sample line was attached to a heating cable and sheathed in foam insulation to prevent ice formation downstream from the catalyst. Temperature of the tubing was controlled by manually adjusting the voltage applied to the heating cable. The system was calibrated several times daily by addition of NO in N2; periodic addition of isopropylnitrate (IPN) indicated that conversion efficiency near 100% (for IPN) was maintained throughout the season. The range of NO sensitivity was 0.3 to 0.5 ppt photon⁻¹ s⁻¹; after day 141, the sensitivity only varied between 0.4 and 0.475 ppt photon⁻¹ s⁻¹. Laboratory tests subsequent to the field season were conducted to determine the conversion efficiency for HCN. The configuration did not duplicate field

conditions, but the tests suggested that the catalyst also converted HCN with efficiency >50%, which adds a degree of uncertainty to the true level of the lowest NO_y concentrations but should not affect the fluxes. A fast-response CO₂/H₂O analyzer (LiCor 6262) was used to measure H₂O mixing ratios. The instrument response was checked using a dewpoint calibrator before and after the sampling season and found to be constant. Ozone was determined by UV absorbance (Dasibi 1008). Dry zero air was supplied to the inlet several times a day to determine the zero offsets for both H₂O and O₃.

The anemometer and concentration signals were recorded at 4 Hz, and auxiliary data on pressures and flows used to calibrate the sensors were recorded at 0.5 Hz by a PC-based data acquisition and control system. The data were downloaded to a separate computer daily and processed to generate initial concentration, flux and quality assurance data. Air temperature was computed from the speed of sound reported by the sonic anemometer. Fluxes of NO_y, H₂O, heat, and momentum were calculated for 10 min intervals from the covariance of fluctuations in vertical wind velocity (w') and tracegas concentration (C'), temperature (T') or horizontal wind velocity (u') after removing the linear trend and offsetting the data to account for delays in the sample tubing [McMillen, 1988].

Because the source of contamination at Summit was so close to the sensor, data rejection based on wind sector alone may not be reliable, particularly when winds are light and variable. In order to objectively exclude periods with contamination from camp activities, the NO_y mixing ratio data were rejected if the signal variance was too large. A few additional observations were rejected because the signal changed too much between successive intervals (see Table 1).

Table 1. Statistics of NO_y Data Rejection for the 1995 Summer Field Season at Summit, Greenland

Criteria	Hours		
Overall duration of campaign			
(May 1 - July 20, 1995)	1923		
NO _v Mixing Ratios			
Instrument in sampling mode	1553		
Instruments on	1535		
Variance indicates clean conditions ^a	1320		
ΔNO, indicates clean conditions ^b	1311		
After exclusion of warm-up time and			
miscellaneous failures	1280		
Mixing ratios acceptable	1280		
Wind Velocities			
Instrument operational	1574		
Sonic data valid - frost free (T<0°C)	1255		
In-sector winds ^c	1187		
Valid friction velocity ^d	1168		
Downward momentum flux	1164		
Wind data acceptable	1164		
Eddy flux (w'NO _y ') acceptable	944		

 $^{^{}a}$ 21 < σ NO_v < 1250 and σ NO_v²/NO_v < 2^o0.

blΔNO₃l < 1450. NO₃ thresholds are based on the raw photon count data sampled at 4 Hz in units of counts (or counts² for the variance).

Tower not upwind of the sensors.

 $^{^{}d}$ u* < 0.0046 + 0.045 × \bar{u} (m s⁻¹), u* = (-1*<u'w'>)^{1/2}.

The sonic anemometer failed when rime ice covered the transducers. The computed eddy fluxes were rejected if icing was indicated by anomalous temperatures or if the wind was blowing through the tower. A small number of periods with in-sector winds and valid temperatures still had anomalous wind data and were rejected (Table 1). Between May 1 and July 19, the data system was operational 82% of the time. The wind velocity and NO_y mixing ratio data were acceptable for 66% and 60% of the time, respectively; the NO_y fluxes were acceptable for 50% of the possible observations (Table 1).

3. Results

3.1. Meteorological Conditions

During the summer months the Sun never sets at Summit; at the solstice, the noon Sun is $\approx 41^{\circ}$ above the horizon and the midnight Sun is 5° above the horizon. Air temperatures exhibited diel variation in response to Sun angle superimposed on seasonal and synoptic trends (Figure 1). Tethered-balloon temperature soundings (not shown) revealed that a strong temperature inversion (temperature increases of up to 10° C) existed at or near (within tens of meters) the surface on

many days. Soundings were restricted to times and altitudes with wind speeds below ≈10 m s⁻¹. Maximum temperatures on the tower during the measurement period never exceeded 0°C.

Except for wind storms on June 10-11 and July 3-4, the momentum flux was generally between 0 and -0.1 m² s⁻²; these low values demonstrate the weak turbulence above the cold, aerodynamically smooth snow surface (Figure 2). Grelle and Lindroth [1996] reported problems with heat fluxes calculated from a Solent sonic anemometer at wind speeds >10 m s⁻¹ due to correlated noise in the temperature and wind speed channels. Less than 10% of the observations at Summit were in this range, and the heat fluxes showed no apparent anomalies. If this problem is due to vibration of the sensor, it may depend on specific properties of the mounting and tower configuration. Sensible heat fluxes above the snow were small; 90% of observations had absolute values <20 W m⁻². Negative sensible heat fluxes and decreasing air temperatures were usually observed between the hours of 1600 and 0800, which we refer to here as "night." Peak values of latent heat (evaporation) flux during midday were <15 W m⁻² (0.3 mmol-H₂O m⁻² s⁻¹; 5.4 mg m⁻² s⁻¹). Mean water vapor fluxes during the afternoon were 5 W m⁻² (0.1 mmol H₂O m⁻² s⁻¹). Measur-

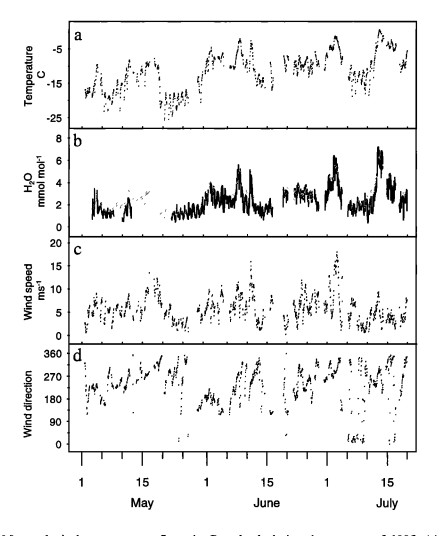


Figure 1. Meteorological parameters at Summit, Greenland, during the summer of 1995: (a) virtual temperature measured by the sonic anemometer, (b) H_2O concentrations shown as black lines superimposed on the saturation vapor pressure for the observed air temperature, (c) wind speed, (d) wind direction.

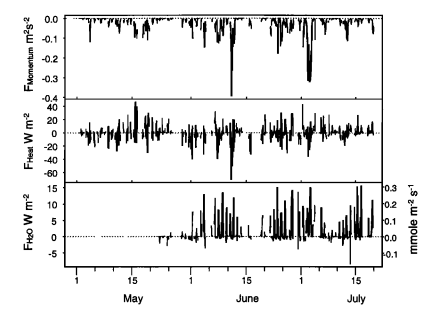


Figure 2. Momentum and sensible and latent heat (water vapor) fluxes measured at Summit, Greenland, during May - July 1995.

able negative values (condensation) were consistently observed at night, due to formation of frost and ground fog. Note that periods of water vapor deposition may be underrepresented because fog was generally accompanied by rime formation on the sonic anemometer. Water vapor concentrations were generally near the saturation vapor pressure (Figure 1a). The small magnitude of water vapor fluxes and rapid diel variations in H_2O concentration imply that the surface layer was fairly shallow and, if air above the inversion was dry, that the entrainment rate was small. Vertical profiles of water vapor concentration that would allow quantification of the water budget and entrainment rates could not be measured accurately in this subzero environment with the available sensors.

3.2. NO_v and O₃ Mixing Ratios

The variance-based criteria used to reject periods of local contamination remove the high NO_y concentration outliers from the data, leaving a population that fits a lognormal distribution (Figure 3). From May 3 to July 19, 1995, NO_y mixing ratios at Summit ranged from <100 ppt to 4900 ppt (ppt = 10^{-12} mol NO_y /mol air). Mixing ratios of NO_y were highest in May (Figure 4). Median mixing ratios decreased by a factor of 2 from May to July, and other statistical measures of the concentration distribution decreased similarly (Table 2).

Ozone concentrations declined from values near 70 ppb in early May to a minimum about 30 ppb in mid-June, then rose

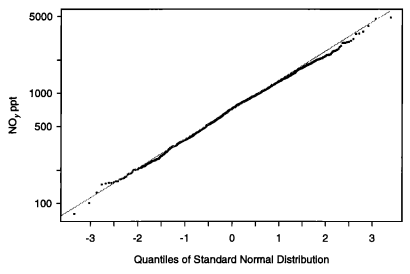


Figure 3. Probability distribution for NO_y concentrations measured at Summit, Greenland, during the 1995 field season, May - July. The line is fit to the 25% and 75% quantiles. Lognormally distributed data would follow a straight line in this plot.

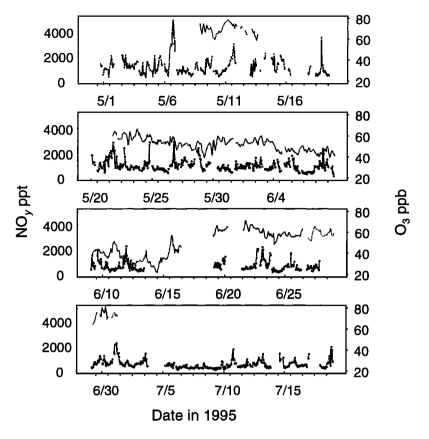


Figure 4. NO_y (black line with dots) and O_3 (gray lines) concentrations at Summit, Greenland, during May-July 1995 are shown for 20-day periods. Downward pointing tick marks indicate noon of each day (local time). Gaps in the O_3 data in the beginning of May and in July are due to instrument malfunctions.

again to 70 ppb (Figure 4). The high O₃ concentrations are characteristic of high-latitude free tropospheric air [Browell et al., 1992; Gregory et al., 1992]. In general, O₃ concentrations had a maximum in the late afternoon, as is typical of sites

where surface deposition is balanced by entrainment of high O₃ from aloft, but the diel variations at Summit were quite small implying a weak deposition velocity for O₃ to snow surfaces [Weseley et al., 1981].

Table 2. NO₂, HNO₃, and O₃ Mixing Ratios and NO₂ Fluxes Between May 3 and July 19, 1995, at Summit, Greenland

Period (N)	Percentile							
	10%	25%	33%	Median	67%	75%	90%	Mean
				NO _y ,	nnt			
Overall (1280)	330	469	547	740	939	1072	1508	850
May (507)	574	751	828	985	1252	1363	1849	1145
June (429)	329	448	518	683	858	965	1242	748
July (344)	233	331	363	444	571	651	947	544
, (,				O_3 ,	ppb			
Overall (943)	37	46	50	55	58	61	68	54
May (318)	49	53	54	58	63	65	72	59
June (601)	33	43	45	51	56	58	63	50
` '				HNO3 at	8m", ppt			
April 25-July 8 (225)	2.7	4.2	4.9	6.4	9.9	11.2	18.3	9.6
May (69)	3.4	5.5	7.4	10.4	13.4	15.7	24.1	12.8
June (134)	2.5	4	4.4	5.5	7.1	8.8	12.6	8.2
July (22)	3.7	4.4	4.6	8,3	10.2	12	14.5	8.3
, ,				NO _ν Flux, μ	$mol m^{-2} h^{-1}$			
Overall (944)	-0.705	-0.151	-0.053	0.002	0.066	0.146	0.717	0.008
May (316)	-0.468	-0.095	-0.041	0.001	0.063	0.178	0.973	0.140
June (339)	-1.154	-0.409	-0.270	0.007	0.126	0.305	0.898	-0.070
July (289)	-0.318	-0.051	-0.022	0.000	0.030	0.066	0.206	-0.045

^a Dibb et al. [1998]

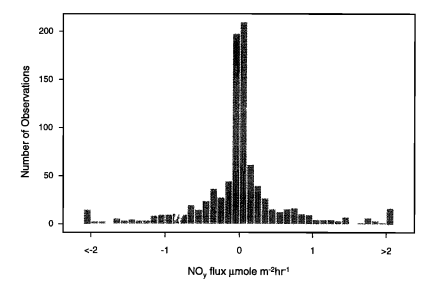


Figure 5. Distribution of NO_y fluxes measured at Summit, Greenland, during the summer of 1995.

3.3. Fluxes

The NO₂ flux observations averaged over hourly intervals are about equally divided between periods of net deposition $(F_{NOy} < 0)$ and net emission $(F_{NOy} > 0)$ (Figure 5, Table 2). We estimate that the absolute value for the smallest observable flux (defined as twice the standard deviation during periods of very small flux) is 0.03 μ mol m⁻² h⁻¹. The absolute values in general are small; 25% were below the 0.03 μ mol m⁻² h⁻¹ (Figure 6). The frequency of fluxes with larger absolute value is highest in June. The NO₂ fluxes are generally weak in July, resulting in the minimum range between 10% and 90% quantiles. In comparison to NO₂ flux, snow deposition of NO₃ averages 3.8 μ mol m⁻² per event [Bergin et al., 1995]; total snow deposition of NO₃ for the period April 28-July 12, 1995, was 149 μ mol m⁻² (J. Dibb, unpublished data, 1995).

4. Discussion

4.1. Nature and Origin of NO, at Summit

Previous observations of nitrogen oxides in the Arctic boundary layer show a spring peak in NO_y concentration that is dominated by PAN [Bottenheim and Gallant, 1989; Bottenheim et al., 1986,1993; Dickerson, 1985; Honrath and Jaffe, 1992; Honrath et al., 1996; Solberg et al., 1997] (Table 3). Enhanced PAN background concentrations are also observed in the northern midlatitudes during the winter months [Brice et al., 1988]. The data from Summit, which is at 3 km, are best compared to measurements in the free troposphere. Aircraft measurements in the Arctic during July and August indicate an increase of PAN concentrations with altitude [Singh et al., 1992a,b] (see Table 3).

Results from a three-dimensional chemical transport model that includes a detailed chemical mechanism for reactive N [Wang et al., 1998] (Table 4) summarize our current understanding of chemical and physical processes that affect longrange transport of reactive nitrogen to the Arctic troposphere. The model predicts that PAN will be the dominant NO₂ species in the Arctic middle troposphere. In the model, PAN over

Summit is mostly derived from long-range transport in the free troposphere of PAN produced at northern midlatitudes from anthropogenic NO, Even a small yield of PAN can lead to a substantial accumulation during the late winter and early spring because the lifetime for PAN is very long at cold temperatures. Emissions from forest and tundra fires during summer may contribute to additional PAN at high latitudes [Singh et al., 1992a]. The spring maximum in the model most likely results from a combination of high emissions of biogenic hydrocarbons (precursors of PAN) and relatively low temperatures (suppressing PAN decomposition) [Liang et al., 1998]. Because natural sources of NO_x (soils, lightning, forest fires) are at a minimum in winter, the NO, peak in spring is most likely of anthropogenic origin. Simulated PAN concentrations exceed 500 ppt in April and decline to levels <200 ppt by July. A vertical gradient of increasing PAN concentrations with altitude develops over the summer, reflecting the dominance of high-altitude transport of pollution as was observed in the Arctic Boundary Layer Expedition (ABLE 3A) aircraft mission over the Arctic [Harriss et al., 1992] and colder temperatures aloft. HNO3 concentrations are depressed in the bottom model layer by deposition. Both the model and aircraft observations have higher HNO₃ concentrations and HNO₃:NO_y ratios than are observed in the surface layer at Summit. However, the model predictions are averaged through a layer that is thicker than the surface inversion at Summit, and the aircraft observations are made in the free troposphere. The lifetime for HNO₃ in a 100 m surface layer with a deposition velocity of 0.7 cm s⁻¹ (the median V_d computed from aerodynamic resistance) is <4 hours and decreases linearly as mixing height decreases. Modeled NO_x concentrations are low (10-20 ppt) throughout the spring and summer due to the ~1 day lifetime for NO_x oxidation.

The NO_v observations from this study (Table 2) exceed the model predictions by a factor of 2-4 but show similar trends. The median NO_v concentration observed at Summit during July (444 ppt) is within the range of NO_v concentrations measured in the free troposphere during ABLE 3A, and a factor of 2 higher than the corresponding PAN concentration (see Table 3). Concentrations of NO_v and PAN in the bound-

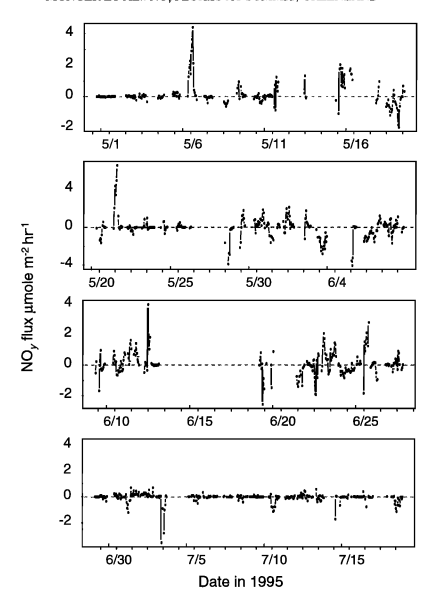


Figure 6. Time series of NO_y fluxes at Summit, Greenland. Each dot represents a 1-hour average flux. Downward pointing tick marks indicate noon of each day (local time).

ary layer (Table 3) also decrease from spring to summer, but the concentrations are considerably less than at Summit or aircraft observations above the boundary layer, consistent with the predicted vertical gradients. The results of this and previous studies taken together with the model predictions point to the accumulation of a reactive N reservoir during late winter and early spring in the Arctic. The dominant identified species is PAN [Singh et al., 1992b], but other compounds such as alkyl nitrates are also present at small concentrations [Muthuramu et al., 1994]. Based on these considerations, we assume that PAN is the NO_v species that is transported to and accumulates in the Arctic midtroposphere and that it probably is the dominant NO_v species at Summit. Concentrations of NO_v, however, were lower at Summit during a 3-week measurement campaign in the summer of 1998 than in 1995, and the contribution from PAN was less than expected [Campbell et al., 1998]. A detailed analysis of air mass histories for the two seasons has not been done to try and understand these

differences. PAN is destroyed by heating during subsidence and would be less abundant in marine air masses uplifted to Summit. The NO_y concentrations measured during ABLE 3A were a factor of 2 higher than the sum of concurrently measured NO_x, PAN, and HNO₃ concentrations; issues about measurement artifacts [Bradshaw et al., 1998] or unidentified species have not been resolved. As temperatures in the middle and lower troposphere warm, PAN (and other organic nitrogen species) thermally decomposes, and the resulting NO₂ is quickly converted to HNO₃ [Fan et al., 1994]. The reservoir concentration in the middle troposphere declines over summer through reaction and dilution by mixing with NO_y-depleted air. Measurements made in the boundary layer, where mean temperatures reach 0°C or above, indicate that NO_y is depleted to low concentrations.

We next examine whether the distinct episodes of elevated mixing ratio lasting for hours at a time (Figure 4) can be attributed to specific sources or transport patterns. The magni-

Table 3. Reactive Nitrogen and Ozone Concentrations at Selected Sites in the Arctic Free Troposphere and
Boundary Layer

Month	Altitude, km	NO _x , ppt	HNO ₃ , ppt	PAN, ppt	NO _y , ppt	O ₃ , ppb
		Free-Troposph	eric, Arctic and Sul	parctic Flights ^a		
July-August	2-4	26	78	140	518	36
July-August	4-6	34	70	285	860	57
			Spitsbergen ^b			
March	0.5	25		304	426	37
April	0.5	35		290	363	33
May	0.5	23		164	180	29
•			Barrow, Alaska ^t			
March					575	
May	0				255	
June	0				89	
July	0				86	
•			Bethel, Alaska ^d			
July-August	0	14			202	32

^aSandholm et al. [1992], Talbot et al. [1992], Singh et al. [1992a], and Gregory et al. [1992].

tude of these episodes decreases from May to July. Analysis of isentropic back trajectories indicates that about 50% of the air parcels arriving at Summit during the May-July 1995 period had passed over northern Canada or the Arctic Ocean within the previous 5 days [Kahl et al., 1997] (J. Kahl, University of Wisconsin, Milwaukee, unpublished trajectory analysis for 1995). Parcels that had been over the Atlantic Ocean or stagnated over Greenland for the previous 5 days accounted for an additional 23% and 12%, respectively. Trajectories from midlatitude regions of North America, Europe, or Asia that could have been significantly influenced by anthropogenic emissions or forest fires within 5 days had a combined frequency of 10% during the summer of 1995. NO, events that occur on timescales of a few hours were not clearly associated with the back trajectories calculated at 12hour intervals. Large day-to-day variability in aerosol ⁷Be and ²¹⁰Pb at Summit has been attributed to short-term fluctuations in vertical mixing between the surface layer and free troposphere [Dibb, 1990]. Because vertical mixing at Summit is intermittent, the surface layer is decoupled from the free troposphere. Variations in NO, concentrations aloft may be delayed in the surface layer or not observed at all.

4.2. Factors Contributing to NO, Flux

4.2.1. Storage. The vertical flux of NO_y measured at 18 m (F_{18}) is equal to the flux at the snow surface, F_{0} , plus any change in concentration (storage) in the 0-18 m column:

$$F_{18} = F_0 + \frac{\partial}{\partial t} \int_0^h C(z) dz$$
 (1)

where C(z) is the number density of NO_y at height z and h = 18 m. Because snow is porous, the lower boundary should be below the surface, but we ignore this contribution because only the upper 0.1 to 1 m is rapidly ventilated [Colbeck, 1989; Waddington et al., 1996], and air space in the snowpack is reduced by the volume of ice. We estimate the magnitude of the storage term by assuming a uniform concentration below the sensor. For the range of observed hourly concentration changes the storage term would be between -1 and 1.5 µmol m⁻² h⁻¹; 50% of the data, however, have an absolute value <0.05 µmol m⁻² h⁻¹. Although the storage term can have a large magnitude at times, the surface exchange, F₀ is not consistently different from the flux at 18 m. More important, the storage term does not bias the results; the least squares fit of F_0 versus. F_{18} has a slope of 0.99 and an intercept not different from 0.

4.2.2. PAN adsorption. Laboratory measurements (R. Friedl, Jet Propulsion Laboratory, personal communication, 1997) indicate that PAN is reversibly adsorbed on ice at 193 K and shows no evidence of chemical reaction. The adsorption coefficient, K_{ads} , is $K_{ads} = 30$ cm (defined as the ratio of PAN adsorbed (molecules cm⁻²) to the concentration in the adjacent air (molecules cm⁻³)). For our application the mass of adsorbed PAN in a volume of snow is determined by the gas phase concentration, adsorption constant, and the bulk surface

Table 4. Reactive N Mixing Ratios (ppt) Simulated Over Summit, Greenland, by a Global Three-Dimensional Model

Month	NO _x		PAN		HNO ₃		NO _y	
	0-0.4km	1.0-2.3 km	0-0.4 km	1.0-2.3 km	0-0.4 km	1.0-2.3 km	0-0.4 km	1.0-2.3 km
April	8	11	547	547	80	162	688	769
May	14	19	257	306	78	187	381	543
June	15	18	191	233	77	197	300	469
July	14	20	133	180	94	207	253	421
August	14	19	132	176	96	199	258	411

Mean model results from Wang et al., [1998] for the atmosphere over Summit. The model was sampled in vertical levels 1 and 3, corresponding to 0-0.4 km and 1-2.3 km above the surface at Summit, which is at 3 km above sea level.

^hSolberg et al. [1997].

^cHonrath and Jaffe [1992].

^dBakwin et al. [1992].

area of the snow. The unadsorbed PAN in the pore space is the gas phase concentration adjusted for the volume of snow-pack filled by ice. Hence the ratio of adsorbed PAN on the grain surfaces (PAN_{sfc}) to free PAN in the pores (PAN_{pare}) for a volume of snow is given by

$$\frac{PAN_{sfc}}{PAN_{pore}} = \frac{K_{ads} \times SV}{\left(1 - \frac{\rho_s}{\rho_i}\right)}$$
 (2)

where SV is the bulk surface area of the snowpack (typically 30 mm⁻¹ for wind-packed surface snow [Davis et al., 1996] (B. Davis, Cold Regions Research and Engineering Laboratory, unpublished data, 1998)), p_e is the bulk density of snow (typically 0.4 g cm⁻³ for wind-packed surface snow [Davis et al., 1996] (B. Davis, CRREL, unpublished data, 1998)), and p, is the density of ice (0.9 g cm⁻³). For the conditions given, which correspond to a layer of small, closely packed grains, adsorbed PAN exceeds PAN in the pores by a factor of 16,000. Equilibrium should be established quickly within the snowpack, but depletion of PAN from the much larger volume of overlying air is limited by the ventilation rate across the snow-air interface. Because this is an equilibrium process, it does not present an infinite sink for PAN. Reversible adsorption on ice surfaces can be thought of in the same way as the storage term described in equation (1). For the snow morphology given above and the adsorption coefficient determined at 193 K, a 1 cm layer of snow would hold 2.85 nmol of PAN per m² at equilibrium with a 1 ppt gas phase concentration. Temporal variations in NO_v concentration range from -400 ppt h⁻¹ to 400 ppt h⁻¹ (90% of data), which translates to upper limit fluxes of -1.1 µmol m⁻² h⁻¹ to 1.1 µmol m⁻² h⁻¹ if ventilation of the snowpack and PAN equilibration at the ice surfaces are rapid and all NO, is (or behaves like) PAN. This range is comparable to the measured fluxes; however, it is probably an overestimate because concentration changes in the snowpack may be damped relative to variations in the overlying air, and the adsorption coefficient for PAN at temperatures observed at Summit is probably less than the value for 193°K. The adsorption constant and reactivity for PAN on ice at temperatures relevant to polar snow need to be determined. This analysis suggests, however, that the porous snowpack provides a mechanism to at least temporarily sequester a relatively large mass of PAN. Adsorption and desorption of PAN in response to changes in ambient concentration provide a mechanism to explain NO_v eddy fluxes that are in opposition to observed HNO3 concentration gradients [Dibb et al., 1998]. The NO, flux measurement does not distinguish the species involved and would not differentiate PAN exchange from release of a different compound (e.g., NO_x [Honrath et al., 1999]). As pointed out by Dibb et al. [1998], conversion of only a small fraction of the NO, passing through the snow could be a large term in the snow nitrate budget.

4.2.3. HNO₃ uptake. Laboratory studies [Abbatt, 1997; Laird and Sommerfeld, 1995; Zondlo et al., 1997] show very efficient uptake of HNO₃ by ice up to monolayer coverage. Slower adsorption corresponding to formation of a multilayer or surface rearrangement is observed as well. Diffusion of HNO₃ through ice crystals is discounted by Sommerfeld et al. [1998], however. Adsorption of a HNO₃ monolayer is largely irreversible [Abbatt, 1997; Laird and Sommerfeld, 1995; Zondlo et al., 1997]. The capacity for monolayer HNO₃ adsorption is weakly temperature dependent and decreases by

only a factor of 4 as temperature increases from 208 K to 248 K [Abbatt, 1997]. As seen below, this apparent contradiction between laboratory measurements of HNO₃ adsorption and HNO₃ volatilization in the field can be reconciled by considering the surface-to-volume relationship of ice grains and the temperature dependence of HNO₃ uptake.

Sommerfeld et al. [1998] report that HNO₃ will be predominantly on the surface of ice grains. The concentration (mol g⁻¹) and surface density (mol cm⁻²) of HNO₃ for an ice grain are related by the S/V ratio, and density of ice. As snow ages its, S/V ratio decreases from >100 mm⁻¹ for new snow, to <5 mm⁻¹ for fully developed depth hoar in summer (B. Davis, CRREL, personal communication, 1998) [Davis et al., 1996]. For the mean snow concentration of 3 nmol g⁻¹ [Dibb et al., 1998] the surface coverage increases from 0.016 x 10¹⁴ molecules cm⁻² to 0.33 x10¹⁴ molecules cm⁻² as the grains increase in size. Conversely, the calculated concentration corresponding to an HNO₃ monolayer at 248 K (0.7 x 10¹⁴ molecules cm⁻²) [Abbatt 1997] decreases from 130 nmol g⁻¹ to 6 nmol g⁻¹ as the snow ages. New snow has a capacity for HNO₃ adsorption well in excess of the observed ambient concentrations, but as long as the snow remains acidic, older coarsegrained snow approaches the monolayer adsorption limit (for ice at 248 K) and may exceed that threshold if the HNO₃ uptake coefficient is even smaller at the warmer temperatures in the summer snowpack. Temperature profiles in the snow can be complex and vary over diel and seasonal timescales. Ice evaporates in the warmer layers, and the vapor is either redeposited in colder layers or escapes entirely [Davis et al., 1996]. The fate of HNO₃ liberated from evaporated ice grains depends on whether the vapor encounters other ice surfaces or is quickly ventilated to the surrounding air. Snow metamorphism could redistribute HNO₃ within the snowpack and modify seasonal layers. This mechanism could account for the large enhancement of NO₃ in the surface snow and the damping of seasonal cycles in older snow [Fischer and Wagenbach, 1996; Mulvaney et al., 1998]. Increasing NO₃ inventories of surface snow layers over the course of several days that have been interpreted as evidence of dry deposition [Bergin et al., 1995; Dibb et al., 1998] may actually be redeposition of HNO₃ migrating from deeper layers. Evaporation at the surface of the snowpack, or sublimation from blowing snow [Pomeroy and Jones, 1996], however, would release HNO₃ to the atmosphere (see section 4.3). The extent of NO₃ loss from the snowpack would depend on the length of time a layer was near the surface, where it is subject to thermal gradients and ventilation, and on the frequency of high winds.

4.3. Illustrative Case Studies

The processes of HNO₃ and PAN adsorption noted above suggest there should be correlation between NO_y fluxes and physical parameters such as temperature, evaporation rate, or changes in ambient concentration. The dominant feature of most of the data is very low NO_y flux associated with weak turbulence, as demonstrated by the small momentum fluxes. The periods with the largest momentum fluxes, however, are consistently associated with positive NO_y and water vapor flux and negative sensible heat flux (Figure 7) expected from sublimation of blowing snow [*Pomeroy and Jones*, 1996]. The ratio of NO_y to H₂O fluxes corresponds to evaporation of snow with a NO₃ concentration of about 20 nmol g⁻¹, which exceeds the observed mean NO₃ concentration in snow at Summit. HNO₃ may be preferentially lost by partial evapora-

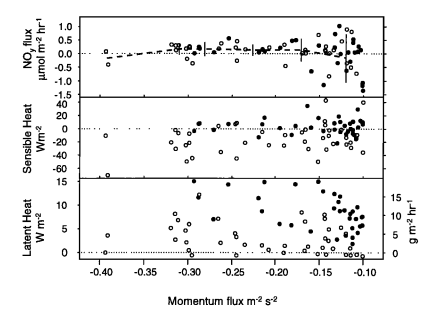


Figure 7. Fluxes of NO_y, sensible heat, and water vapor (latent heat) plotted versus the momentum flux ($F_{MOM} = \langle u'w' \rangle$) when the magnitude of F_{MOM} exceeds 0.1 m⁻² s⁻². Daytime hours are indicated by solid symbols. The mean NO_y fluxes and standard deviations are computed over F_{MOM} intervals of 0.05 m⁻² s⁻² and shown as vertical segments connected by the long-dashed line in the top panel.

tion if it is concentrated on the surface of ice grains rather than in the ice matrix [Sommerfeld et al., 1998].

The interactions between controlling variables at Summit are complex, and correlation plots between NO_v fluxes and any single factor other than momentum flux were generally confounded by the effects of other factors. Physical factors such as snow density, porosity, morphology, or surface temperature that may affect the NO_v flux are difficult to quantify routinely over the large upwind area that contributes to the eddy flux. Except as noted above, defining quantitative relationships between NO_v flux and any single variable is difficult because the data set is too limited to identify trends by binning of the data to selectively hold the other factors constant. Instead we illustrate processes that affect NO, exchange by examining selected periods of the data and pose physical and chemical mechanisms that can be tested in controlled experiments. Periods of enhanced NO_v deposition or efflux are not quantitatively important, and the overall net flux is near zero for the period covered by this data set. However, these infrequent events may illustrate processes that are more significant at other locations or in different seasons.

A pulse of NO_y deposition coincides with a 1500 ppt increase in NO_y concentration the afternoon of June 22 (Figure 8) and is followed by a period of positive NO_y flux during the night. This pattern of alternating deposition and efflux is consistent with temporary storage by reversible surface adsorption. Positive NO_y fluxes continue during the afternoon of June 23 accompanied by wind speeds that peak at 12 m s⁻¹ and observations of snow and blowing snow. Latent heat fluxes that exceed the sensible heat flux suggest strong evaporation and liberation of HNO₃. Another period of high wind on June 24, which only stirred up a layer of blowing snow just above the surface, is associated with a consistent small negative NO_y flux. Evaporation rates are smaller during this period, and the sensible heat flux is larger. The increased sen-

sible heat flux and warmer surface temperatures would weaken the surface inversion and allow mixing over a deeper layer and allow HNO₃ from aloft to reach the surface. Subtle differences in the consolidation of surface snow or shifts in the energy balance can affect the direction of NO_y fluxes.

Several hours of NO_y uptake, with a peak of 1 µmol m⁻² h⁻¹ is associated with a 1500 ppt increase in NO_v concentration on July 1 (Figure 9). Consistent small NO, efflux is observed from midmorning on July 1 to the night of July 2 that is again associated with high wind speed, warmer temperatures, increasing water vapor concentrations, and low ambient NO, concentration, suggesting evaporation of snow into a relatively clean, well-mixed surface layer. A short pulse of NO, deposition occurs on July 3 along with rising NO, concentration that suggests equilibrium surface adsorption. The temperature is colder during this period than during the NO_v concentration increase on July 1, and the surface snow has been densely packed by the wind storm on the previous day. In addition, the slight positive heat flux may enhance vertical mixing and entrainment of air that has a higher proportion of HNO₃.

4.4. Nitrate Scavenging by Precipitating Snow

In this section we consider whether the levels of nitrate observed in snow at Summit are consistent with the NO_y concentrations and fluxes. Based on the recent laboratory investigations of HNO₃ adsorption and evaluation of HNO₃ uptake by cirrus clouds [Abbatt, 1997; Zondlo et al., 1997] (see section 4.2) we expect that HNO₃ is completely scavenged by the ice surface during formation of clouds and snowflakes. Even if a quasi-liquid layer [Conklin and Bales, 1993] were present, HNO₃ uptake would be efficient because of its large solubility and dissociation constants. Although some PAN could adsorb on the fresh snowflakes, the surface area of ice in the cloud is too small for this process to significantly alter

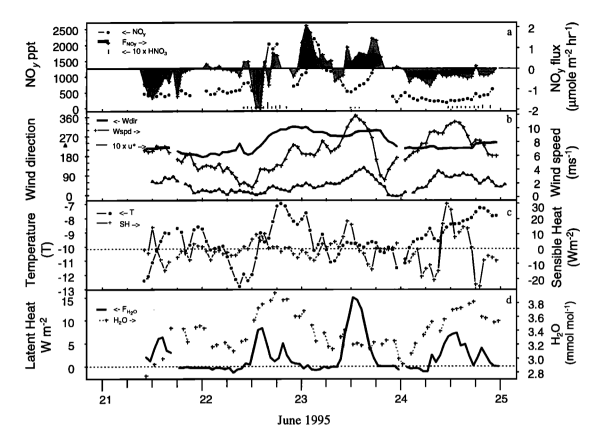


Figure 8. (a) Concentration of NO_y (dots) and NO_y flux (solid shading) shown for a 4 day period beginning on June 21, 1995. Vertical bars indicate the HNO₃ concentration (HNO₃ has been scaled by 10 for greater visibility on the plot). (b) Wind direction (solid line), wind speed (crosses connected by dashed line), and u*, the friction velocity, x10 (triangles) are plotted against time. Blowing snow was generally observed with wind speed >10 m s⁻¹. (c) Temperature (dots), sensible heat flux (crosses), and (d) latent heat flux and H₂O concentration are shown.

the gas phase concentrations, and because the adsorption is reversible, PAN will desorb in response to decreased ambient concentrations. The final concentration of NO₃ in fresh snow depends on the initial concentration of HNO₃ in the air and the mass of water that freezes, which can be computed from the temperatures before and after cloud formation. This mechanism differs from the co-deposition process considered by Silvente and Legrand [1995] because all the HNO₃ present in an air parcel is partitioned into the small fraction of water vapor that freezes. For an HNO₃ concentration of 6 ppt, cooling saturated air from -12°C to -12.5°C is sufficient to produce nitrate concentrations of 3.5 nmol g⁻¹ in snow (Figure 10). Even the low HNO₃ observed in the surface layer could account for the mean NO₃ concentrations in fresh snow, without any additional input from dry deposition.

4.5. Budget Considerations

We next evaluate whether an NO_y reservoir composed of PAN can account for the observed total deposition of NO₃ during summer. The lifetime (1/k) for PAN decomposition below -10°C exceeds 4 weeks (decay rate \leq 4% d⁻¹), but it decreases rapidly to 3 days as the temperature warms above 0°C (decay rate \geq 31% d⁻¹) [Roberts and Bertman, 1992]. Using midday mean air temperatures measured at the surface, the monthly median NO_y concentrations (assuming that NO_y is PAN), and temperature coefficients for PAN decomposition

from Roberts and Bertman [1992], we compute a maximum source of NO, from PAN of 13 ppt d⁻¹, 34 ppt d⁻¹, and 31 ppt d⁻¹ in the surface layer for May, June, and July, respectively. The lifetime for OH oxidation of NO2 to HNO3 at pressure and temperature of Summit and an OH concentration of 1x 10⁶ molecules cm⁻³ is 1 day. Fan et al. [1994] conclude that PAN is extensively decomposed and converted to HNO₃ in subsiding air parcels. Thermal decomposition of PAN followed by oxidation of NO_x to HNO₃ in a 3 km layer above Summit could thus yield 153 µmol m⁻² of deposited NO₃ for the period May 1 to July 15 (1 µmol m⁻² d⁻¹, 2.7 µmol m⁻² d⁻¹, and 2.5 µmol m⁻² d⁻¹ for the months May, June, and July, respectively). From April 28 to July 12, 1995, the nitrate accumulation in snow was 149 µmol m⁻² (J Dibb, unpublished data, 1996). Such close agreement between the NO₃ accumulation rate in fresh snow and PAN decomposition is fortuitous, considering the gross simplification in our assumptions about vertical structure of temperature and PAN concentration. The surface temperatures lie between the inversion temperature and the temperatures in the free troposphere above. Nevertheless, the calculation demonstrates that production of HNO₃ from PAN can potentially account for the NO₃ deposited in snow. As noted in section 4.1, PAN transport and thermal decomposition account for HNO3 concentrations of 80-90 ppt just above the surface layer. Furthermore, the seasonal pattern for PAN decomposition can ex-

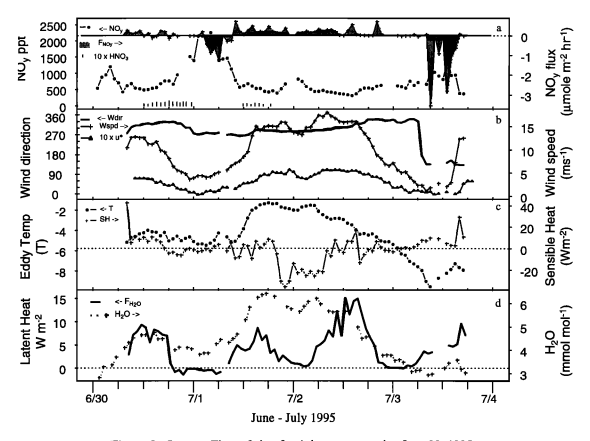


Figure 9. Same as Figure 8, but for 4 days commencing June 30, 1995.

plain the observed summertime peak in snow NO₃ [Davidson et al., 1989; Whitlow et al., 1992] despite decreasing ambient NO₃ concentrations over the summer.

Our observations of small offsetting fluxes of NO_y from the snowpack are consistent with the observations by *Bergin*

et al. [1995] that measured inputs in fresh snow and fog accounted for 99% of the NO₃ present in the snowpack at the end of the season, leaving little room for significant dry deposition or revolatilization of HNO₃. Although the flux observations from Summit do not indicate a net loss of nitrate from

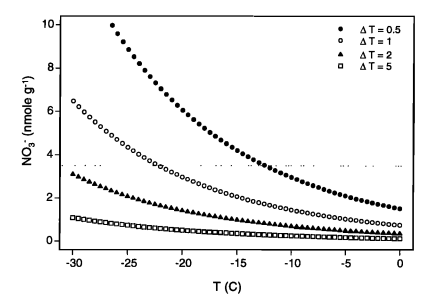


Figure 10. Nitrate concentration in snow that formed by cooling a specified ΔT below the saturation temperature in air with an ambient HNO₃ concentration of 0.27 nmol m³ (6 ppt) [Dibb et al., 1998] is calculated assuming 100% scavenging and is plotted on the vertical axis against the saturation temperature on the horizontal axis. The symbols indicate temperature differences associated with snow formation. The ice formed by cooling saturated air 0.5°C from -12°C would have a nitrate concentration of 3.5 nmol g⁻¹. The dashed horizontal line shows the mean nitrate concentration in snow [Dibb et al., 1998].

the snowpack over the summer, they do not address the potential redistribution of NO₃ within the snowpack or losses in other seasons or at other sites.

5. Conclusions

The concentration of NO_y at Summit, Greenland, declined over the summer from a median of 950 ppt in May to 440 ppt in July. Nitric acid in the surface layer was typically only 1% of the observed NO_y. Predictions based on current understanding of reactive N chemistry indicate that PAN should accumulate in early spring in the Arctic troposphere. Thermal decomposition rates for PAN as temperatures warm in the summer are adequate to supply fresh NO_x that is oxidized to HNO₃ and scavenged by precipitating snow. Variations in the ice-core nitrate record should be interpreted with respect to variations in the long-range transport of PAN to the Arctic, which depends on the supply of precursor hydrocarbons, photochemical activity, temperatures, as well as variations in NO_x emissions.

The turbulent fluxes of NO_y at Summit were bidirectional, but the magnitudes were small. Most of the time, NO_y fluxes were <1 µmol m⁻² h⁻¹. Neither emission nor dry deposition dominated the NO_y flux, and the overall net exchange was insignificant compared to the flux of NO₃ via snow deposition, which contributed 149 µmol m⁻² over 75 days. Water vapor concentrations were close to the saturation vapor pressure and responded quickly to variations in temperature, which demonstrates that the boundary layer is shallow and infrequently mixed with the overlying atmosphere. If the process that limits HNO₃ dry deposition is its entrainment from aloft, the small fluxes that we observed would be sufficient to maintain very low concentrations in the surface layer.

Occasional pulses of NO, exchange into and out of the snow that follow ambient concentration changes suggest that reversible adsorption of compounds such as PAN on the ice surfaces may temporarily store reactive N in the snowpack. However, we cannot rule out adsorption of one species followed by conversion and release as a different compound. Upward fluxes of NO, and H₂O during the highest wind events point to sublimation as a mechanism for releasing NO₃ from the snowpack. Nitric acid is tightly bound to ice surfaces up to a monolayer coverage but may be liberated as the snow ages and the individual grains increase in size. Once liberated, this HNO3 may redeposit on fine-grained layers near the surface or be lost from the snowpack if it is released near the surface and is quickly ventilated. The tendency for NO_v efflux during windy periods that evaporate the snow indicates that postdepositional losses of NO₃ from the snowpack could be more important in drier or windier seasons and sites.

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References

- Abbatt, J. P. D., Interaction of HNO₃ with water-ice surfaces at temperatures of the free troposphere, *Geophys. Res. Lett.*, 24, 1479-1482, 1907.
- Bakwin, P. S., S. C. Wofsy, S.-M. Fan, and D. R. Fitzjarrald, Measurements of NO_x and NO_y concentrations and fluxes over Arctic tundra, *J. Geophys. Res.*, 97, 16,545-16,557, 1992.
- Bakwin, P. S., et al., Reactive nitrogen oxides and ozone above a taiga woodland, J. Geophys. Res., 99, 1927-1936, 1994.
- Bergin, M. H., J.L. Jaffrezo, C. I. Davidson, J. E. Dibb, S. N. Pandis, R. Hillamo, W. Maenhaut, H. D. Kuhns, and T. Makela, The contributions of snow, fog, and dry deposition to the summer flux of anions and cations at Summit, Greenland, J. Geophys. Res, 100, 16,275-16,288, 1995.
- Bottenheim, J. W., and A. J. Gallant, PAN over the Arctic: Observations during AGASP-2 in April 1986, J. Atmos. Chem., 9, 301-316, 1989.
- Bottenheim, J. W., A. G. Gallant, and K. A. Brice, Measurements of NO_y species and O₃ at 82°N latitude, *Geophys. Res. Lett.*, 13, 113-116, 1986.
- Bottenheim, J. W., L. A. Barrie, and E. Atlas, The partitioning of nitrogen oxides in the lower Arctic troposphere during spring 1988, *J. Atmos. Chem.*, 17, 15-27, 1993.
- Bradshaw, J., S. Sandholm, and R. Talbot, An update on reactive odd-nitrogen measurements made during recent NASA Global Tropospheric Experiment programs, *J. Geophys. Res.*, 103, 19,129-19,148, 1998.
- Brice, K. A., J. W. Bottenheim, K. G. Anlauf, and H. A. Wiebe, Long-term measurements of atmospheric peroxyacetylnitrate (PAN) at rural sites in Ontario and Nova Scotia: Seasonal variations and long-range transport, *Tellus*, 40, 408-425, 1988.
- Browell, E. V., C. F. Butler, S. A. Kool, M. A. Fenn, R. C. Harriss, and G. L. Gregory, Large-scale variability of ozone and aerosols in the summertime arctic and sub-arctic troposphere, *J. Geophys. Res.*, 97, 16,433-16,450, 1992.
- Campbell, B. M., P. B. Shepson, S. Bertman, R. Honrath, M. Peterson, and S. Guo, Ambient peroxyacetyl nitrate concentrations at Summit, Greenland, *Eos Trans. AGU*, 79(45), Fall Meet. Suppl., F97, 1998.
- Colbeck, S. C., Air movement in snow due to windpumping, J. Glaciol., 35, 209-213, 1989.
- Conklin, M. H. and R. C. Bales, SO₂ uptake on ice spheres: Liquid nature of the ice-air interface, *J. Geophys. Res.*, 98, 16,851-16,855, 1993.
- Davidson, C. I., J. R. Harrington, M. J. Stephenson, M. J. Small, F. P. Boscoe, and R. E. Gandley, Seasonal variations in sulfate nitrate and chloride in the Greenland ice sheet: Relation to atmospheric concentrations, Atmos. Environ., 23, 2483-2493, 1989.
- Davis, R. E., E. M. Arons, and M. R. Albert, Metamorphism of polar firn: Significance of microstructure in energy, mass and chemical species transfer, in *Chemical Exchange Between the Atmosphere* and Polar Snow, edited by E. W. Wolff and R. G. Bales, pp. 379-401, Springer-Verlag, New York, 1996.
- Dibb, J. É., Beryllium-7 and lead-210 in the atmosphere and surface snow over the Greenland ice sheet in summer of 1989, J. Geophys. Res., 95, 22,407-22,415, 1990.
- Dibb, J. E., R. W. Talbot, and M. H. Bergin, Soluble acidic species in air and snow at Summit, Greenland, Geophys. Res. Lett., 21, 1627-1630, 1994.
- Dibb, J. E., R. W. Talbot, J. W. Munger, D. J. Jacob, and S.-M. Fan, Air-snow exchange of HNO₃ and NO_y at Summit, Greenland, J. Geophys. Res., 103, 3475-3486, 1998.
- Dickerson, R. R., Reactive nitrogen compounds in the Arctic, J. Geophys. Res., 90, 10,739-10,743, 1985.
- Fan, S.-M., D. J. Jacob, D. L. V. Auzerall, J. D. Bradshaw, S. T. Sandholm, D. R. Blake, H. B. Singh, R. W. Talbot, G. L. Gregory, and G. W. Sachse, Origin of tropospheric NO_x over subarctic eastern Canada in summer, J. Geophys. Res., 99, 16,867-16,877, 1994.
- Fischer, H., and D. Wagenbach, Large-scale spatial trends in recent firn chemistry along an east-west transect through central Greenland, Atmos. Environ., 30, 3227-3238, 1996.
- Fischer, H., and D. Wagenbach, Sulfate and nitrate firn concentrations on the Greenland ice sheet, 2, Temporal anthropogenic deposition changes, J. Geophys. Res., 103, 21,935-21,942, 1998.
- Fischer, H., D. Wagenbach, and J. Kipfstuhl, Sulfate and nitrate firn concentrations on the Greenland ice sheet, 1, Large-scale ger-

- graphical deposition changes, J. Geophys. Res., 103, 21,927-21,934, 1998.
- Gregory, G. L., B. E. Anderson, L. S. Warren, E. V. Browell, D. R. Bagwell, and C. H. Hudgins, Tropospheric ozone and aerosol observations: The Alaskan arctic, J. Geophys. Res., 97, 16,451-16,471, 1992.
- Grelle, A. and A. Lindroth, Eddy-correlation system for long-term monitoring of fluxes of heat, water vapour and CO₂, Global Change Biol., 2, 297-307, 1996.
- Harriss, R. C., et al., The Arctic boundary layer expedition (ABLE 3A): July-August 1988, J. Geophys. Res., 97, 16,383-16,394, 1992.
- Honrath, R. E., and D. A. Jaffe, The seasonal cycle of nitrogen oxides in the arctic troposphere at Barrow, Alaska, J. Geophys. Res., 97, 20,615-20,630, 1992.
- Honrath, R. E., A. J. Hamlin, and J. T. Merrill, Transport of ozone precursors from the arctic troposphere to the North Atlantic region, J. Geophys. Res., 101, 29,335-29,351, 1996.
- Honrath, R. E., M. C. Peterson, S. Guo, J. E. Dibb, P. B. Shepson, and B. Campbell, Evidence of NO_x production within or upon ice particles in the Greenland snowpack, *Geophys. Res. Lett.*, 26, 695-698, 1999.
- Kahl, J. D., W. D. Martinez, H. Kuhns, C. I. Davidson, J.-L. Jaffrezo, and J. M. Harris, Air mass trajectories to Summit, Greenland: A 44-year climatology and some episodic events, J. Geophys. Res., 102, 26,861-26,875, 1997.
- Laird, S. K., and R. A. Sommerfeld, Nitric acid adsorption on ice: A preliminary study, Geophys. Res. Lett., 22, 921-923, 1995.
- Legrand, M. R., and R. J. Delmas, Relative contributions of tropospheric and stratospheric sources to nitrate in Antarctic snow, *Tellus*, 38, 236-249, 1986.
- Liang, J., L. W. Horowitz, D. J. Jacob, Y. Wang, A. M. Fiore, J. A. Logan, G. M. Gardner, and J. W. Munger, Seasonal budgets of reactive nitrogen species and ozone over the United States, and export fluxes to the global atmosphere, J. Geophys. Res., 103, 13,435-13,450, 1998.
- Mayewski, P. A., and M. R. Legrand, Recent increase in nitrate concentration of Antarctic snow, *Nature*, 346, 258-260, 1990.
- Mayewski, P. A., W. B. Lyons, M. J. Spencer, M. Twickler, W. Dansgaard, B. Koci, C. I. Davidson, and R. E. Honrath, Sulfate and nitrate concentrations from a south Greenland ice core, Science, 232, 975-977, 1986.
- Mayewski, P. A., W. B. Lyons, M. J. Spencer, M. S. Twickler, C. F. Buck, and S. Whitlow, An ice-core record of atmospheric response to anthropogenic sulphate and nitrate, *Nature*, 346, 554-556, 1990.
- McMillen, R. T., An eddy correlation technique with extended applicability to non-simple terrain, *Boundary Layer Meteorol.*, 43, 231-245, 1988.
- Mulvaney, R., D. Wagenbach, and E. W. Wolff, Postdepositional change in snowpack nitrate from observation of year-round nearsurface snow in coastal Antarctica, J. Geophys. Res., 103, 11,021-11,031, 1998.
- Munger, J. W., S. C. Wofsy, P. S. Bakwin, S.-M. Fan, M. L. Goulden, B. C. Daube, A H. Goldstein, K. E. Moore, and D. R. Fitzjarrald, Atmospheric deposition of reactive nitrogen oxides and ozone in a temperate deciduous forest and a sub-arctic woodland, 1, Measurements and mechanisms, J. Geophys. Res., 101, 12,639-12,657, 1996.
- Muthuramu, K., P. B. Shepson, J. W. Bottenheim, B. T. Jobson, H. Niki, and K. G. Anlauf, Relationships between organic nitrates and surface ozone destruction during Polar Sunrise Experiment 1992, J. Geophys. Res., 99, 25,369-25,378, 1994.
- Neubauer, J. and K. G. Heumann, Nitrate trace determinations in snow and firn core samples of ice shelves at the Weddell Sea, Antarctica, Atmos. Environ., 22, 537-545, 1988.
- Pomeroy, J. W. and H. G. Jones, Wind-blown snow: Sublimation,

- transport, and changes to polar snow, in *Chemical Exchange Between the Atmosphere and Polar Snow*, edited by E. W. Wolff and R. G. Bales, pp. 453-489, Springer-Verlag, New York, 1996.
- Roberts, J. M. and S. B. Bertman, The thermal decomposition of peroxyacetic nitric anhydride (PAN) and peroxymethacrylic nitric anhydride (MPAN), Int. J Chem. Kinet., 24, 297-307, 1992.
- Sandholm, S. T., et al., Summertime tropospheric observations related to N_xO_y distributions and partitioning over Alaska: Arctic Boundary Layer Expedition 3A, *J. Geophys. Res.*, 97, 16,481-16,509, 1992.
- Silvente, E. and M. Legrand, A preliminary study of the air-snow relationship for nitric acid in Greenland, in *Ice Core Studies of Global Biogeochemical Cycles*, edited by R. J. Delmas, pp. 225-240, Springer-Verlag, New York, 1995.
- Singh, H. B., et al., Atmospheric measurements of peroxyacetyl nitrate and other organic nitrates at high latitudes: Possible sources and sinks, J. Geophys. Res., 97, 16,511-16,522, 1992a.
- Singh, H. B., D. Herlth, D. O'Hara, K. Zahnle, J. D. Bradshaw, S. T. Sandholm, R. Talbot, P. J. Crutzen, and M. Kanakidou, Relationship of peroxyacetyl nitrate to active and total odd nitrogen at northern high latitudes: Influence of reservoir species on NO_x and O₃, J. Geophys. Res., 97, 16,523-16,530, 1992b.
- Solberg, S., T. Krognes, F. Stordal, Ø. Hov, H. J. Beine, D. A. Jaffe, K. C. Clemitshaw, and S. A. Penkett, Reactive nitrogen compounds at Spitsbergen in the Norwegian Arctic, J. Atmos. Chem., 28, 209-225, 1997.
- Sommerfeld, R.A., C. A. Knight, and S. K. Laird, Diffusion of HNO₃ in ice, *Geophys. Res. Lett*, 25, 935-938, 1998.
- Talbot, R. W., A. S. Vijgen, and R. C. Harriss, Soluble species in the arctic summer troposphere: Acidic gases, aerosols, and precipitation, J. Geophys. Res., 97, 16,531-16,543, 1992.
- Waddington, E. D., J. Cunningham, and S. L. Harder, The effects of snow ventilation on chemical concentrations, in *Chemical Ex*change Between the Atmosphere and Polar Snow, edited by E. W. Wolff and R. G. Bales, pp. 404-451, Springer-Verlag, New York, 1996.
- Wang, Y., D. J. Jacob, and J. A. Logan, Global simulation of tropospheric O₃-NO_x-hydrocarbon chemistry, 1, Model formulation, J. Geophys. Res., 103, 10,713-10,725, 1998.
- Weseley, M. L., D. R. Cook, and R. M. Williams, Field measurement of small ozone fluxes to snow, wet bare soil and lake water, *Boundary Layer Meteorol.*, 20, 459-471, 1981.
- Whitlow, S., P. A. Mayewski, and J. E. Dibb, A comparison of major chemical species seasonal concentrations and accumulation at the South Pole and Summit, Greenland, Atmos. Environ., 26A, 2045-2054, 1992.
- Wolff, E. W., Nitrate in polar ice, in *Ice Core Studies of Global Biogeochemical Cycles*, edited by R. J. Delmas, pp. 195-224, Springer-Verlag, New York, 1995.
- Zondio, M. A., S. B. Barone, and M. A. Tolbert, Uptake of HNO₃ on ice under upper tropospheric conditions, *Geophys. Res. Lett.*, 24, 1391-1394, 1997.
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