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Surprisingly small HONO emissions from snow surfaces at Browning Pass, Antarctica

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Abstract. Measured fluxes of nitrous acid at Browning Pass, Antarctica were very low, despite conditions that are generally understood as favorable for HONO emissions, including: acidic snow surfaces, an abundance of NO₃⁻ anions in the snow surface, and abundant UV light for NO₃⁻ photolysis. Photochemical modeling suggests noon time HONO fluxes of 5–10 nmol m⁻² h⁻¹; the measured fluxes, however, were close to zero throughout the campaign. The location and state of NO₃⁻ in snow is crucial to its reactivity. The analysis of soluble mineral ions in snow reveals that the NO₃⁻ ion is probably present in aged snows as NaNO₃. This is peculiar to our study site, and we suggest that this may affect the photochemical reactivity of NO₃⁻, by preventing the release of products, or providing a reactive medium for newly formed HONO. In fresh snow, the NO₃⁻ ion is probably present as dissolved or adsorbed HNO₃ and yet, no HONO emissions were observed. We speculate that HONO formation from NO₃⁻ photolysis may involve electron transfer reactions of NO²⁻ from photosensitized organics and that fresh snows at our site had insufficient concentrations of adequate organic compounds to favor this reaction.

1 Introduction

The production of gas phase HONO from snow surfaces is generally understood to proceed through a mechanism similar to that of NO₃ by the photolysis of NO₃⁻ in the snow surface (Honrath et al., 2000), analogous to reactions in liquid phase (Mack and Bolton, 1999). Fluxes were quantified above snow surfaces at Alert, Nunavut as ca. 40 nmol m⁻² h⁻¹ (Zhou et al., 2001), and 5–10 nmol m⁻² h⁻¹ at Summit Greenland (Honrath et al., 2002) (both noon-time maximum values). The absence of measurable HONO fluxes in the marine Arctic at Ny-Alesund, Svalbard was assigned to an alkaline snow surface, which prevented the emission of HONO (Beine et al., 2003). However, in fresh, acidic snow in ozone depleted airmasses, HONO emissions of up to 60 nmol m⁻² h⁻¹ were found at Ny-Alesund (Amoroso et al., 2005). The importance of the surface snow pH was shown again in the Italian Apennines, where up to 120 nmol m⁻² h⁻¹ HONO were deposited on to snow surfaces that were rendered alkaline by Saharan dust deposition (Beine et al., 2005).

We measured HONO fluxes, snow chemical composition and snow optical properties at Browning Pass, Antarctica; a site close to the Ross Sea. All observed conditions were generally favorable for HONO emissions, yet we observed fluxes larger than 5 nmol m⁻² h⁻¹ only on two short occasions under very specific local air flow conditions. This paper discusses the surprisingly low HONO emissions.

2 Experimental

Measurements were carried out at Browning Pass (74°36.915′ S, 163°56.487′ E) (Fig. 1), which is located 10.1 km from the Italian coastal Antarctic station “Mario Zucchelli” (formerly Terra Nova Bay). The site does not receive direct sea spray, and is, during the rare katabatic flow from the Boomerang and Campbell glaciers somewhat removed meteorologically from marine influences by the Northern Foothills (up to 1000 m altitude); however, the marine influence is prevailing. At this field site, which was accessed by helicopter daily, we measured HONO fluxes, chemical and optical snow properties between 9 November (doy 314) and 28 (doy 333), 2004.

2.1 HONO fluxes

Fluxes of HONO were derived from independent chemical measurements of HONO at two sampling heights above the
snow surface (25 and 150 cm) and simultaneous temperature and wind speed measurements at the same heights. The glacier was reasonably flat for a radius of hundreds of meters in all directions. The snow surface showed alternating outcropping hard windpacks and softer layers, and steps between these two types of snow could reach 30 cm. The large snowfall of 17–18 November temporarily smoothed the snow surface. The only obstruction to the local windfield were our instrument containers, which were located ca. 20 m from the sampling site, perpendicular to the prevailing wind directions, which were SSW and NE, roughly parallel to the glacier.

We sampled HONO at 25 and 150 cm above the snow surface, using two independent 2.5 cm (I.D.) light-shielded inlet lines of 20 m length at flow rates of 38 L min\(^{-1}\) to feed the sample into the container where the instrument was placed. The samples were taken from this flow through ca. 50 cm of 1.58 mm (I.D.) tubing at 3 L min\(^{-1}\). The total residence time in the inlet lines was ca. 17 s. Both inlet lines were identical, and no null-gradients between the two inlets were detected.

The details of our measurement technique, including possible interferences, are discussed in Beine et al. (2005) and Amoroso et al. (2005); briefly; gaseous HONO was trapped quantitatively in a 10-turn glass coil sampler using 1-mM phosphate buffer (pH 7). The scrubbing solution was then derivatized with sulfanilamide (SA)/N-(1-naphthyl)ethyldiamine (NED), subsequently analyzed using high-performance liquid chromatography (HPLC), and detected by vis absorption. Typical operation conditions were: sample flow: 3 L min\(^{-1}\), solution flow: 0.2 mL min\(^{-1}\), derivatization conditions: 5 min at 45°C; HPLC: loop: 300 mL C18 reverse phase column (Varian), eluent: 20% acetonitrile in 15 mM HCl.

To characterize the surface-atmosphere interaction and to determine the turbulent fluxes we used a UVW tri-propeller anemometer (Gill, model 200-27005), which measured the three orthogonal wind vectors at 1 Hz sampling frequency. Fluxes were computed using the eddy covariance technique in the post processing. Additionally, profile measurements of air temperatures and wind speeds were performed to be able to compute fluxes of chemical species. The instrument was set up ca. 10 m from the chemical measurements. The derivation of HONO fluxes from this mixed eddy covariance and gradient technique is fully described in Beine et al. (2003, 2005).

The mechanical tri-propeller was calibrated against 3-D sonic anemometers under various field conditions; offsets are taken into account in the computational procedures. The use of a mechanical device introduces a threshold wind speed (0.25 m s\(^{-1}\)). Consequently a detection limit for the observed HONO flux does not depend on the detection limit of the chemical HONO measurements alone. The \(\sigma\) detection limit for the individual HONO measurement was <0.5 pmol mol\(^{-1}\). In the simplest approach we use this value also for the minimum detectable \(\Delta_{\text{HONO}} (=[\text{HONO}_{\text{down}}] - [\text{HONO}_{\text{up}}])\). Two chemical samples were taken at the same time, and analyzed over a 10 min period; the fluxes were calculated for 10 min envelopes around this time of sampling. At the threshold wind speed the minimum \(\Delta_{\text{HONO}}\) translates into a minimum observable flux of ca. 0.3 nmol m\(^{-2}\) h\(^{-1}\) (0.33 nmol m\(^{-2}\) h\(^{-1}\) to 0.28 nmol m\(^{-2}\) h\(^{-1}\) under neutral conditions between –40°C and 0°C).

At mixing ratios above the 10\(\sigma\) quantification level (1.6 pmol mol\(^{-1}\)) the error is ca. 30%. The error attached to the flux gradient method was estimated through a comparison of sensible heat fluxes measured by the tri-propeller and a fast temperature sensor and analysis by eddy covariance and gradient flux calculations. The mean relative error between these two techniques was around 15%. Because both air temperatures and HONO mixing ratios are scalars, we can assume that the flux of a generic scalar variable in similar atmospheric conditions shows similar errors. Using the HONO data as if they were available at 1 Hz, we can calculate an error using eddy covariance theory; such a (hypothetical) error would come to 35%. Taking into account the difference between the eddy covariance and gradient flux techniques, our HONO fluxes generated by the gradient flux theory have a (mean) total error on the order of 50%.

Fig. 1. Map of the area surrounding our measurement site at Brown- ing Pass, near the Italian Mario Zucchelli (formerly Terra Nova Bay) station.
2.2 Snow optical properties

The optical properties (scattering and absorption cross sections) of the Antarctica snowpacks found at Browning Pass were determined by an experimental and modeling method we have used previously in mid-latitude snowpacks (Fisher et al., 2005), and sea ice (King et al., 2005), where details can be found. The modeling was originally described in Lee-Taylor and Madronich (2002). Briefly, the optical properties of a snowpack can be defined and constrained by measuring the wavelength-resolved albedo and the wavelength-resolved transmission of light through snowpack. The transmission measurements are made at a series of depths below a starting depth a few centimeters below the surface of the snowpack where the optical e-folding depth, $\varepsilon$, is constant and only very weakly dependent on the solar zenith angle. (The optical properties of snowpacks are described well by Lee-Taylor and Madronich (2002), Warren (1982), King and Simpson (2001), and Simpson et al. (2002). The albedos of the snowpack were measured using a GER1500 spectrometer ($\lambda=300$–1000 nm) and measured relative to a reflectance standard under identical light conditions. The e-folding depth of the snowpack (the depth of snowpack to reduce the light intensity by 1/e of its initial value) was measured by probing the snow with a flat plate irradiance probe connected to a spectrometer ($\lambda=280$–449 nm) at different depths within the snowpack. During the transmission measurement any changes in the downwelling irradiance were measured by the GER1500 fitted with a cosine collector. The downwelling atmospheric flat-plate irradiance was measured throughout the campaign by the GER1500 fitted with a cosine corrector every 1–5 min (more technical information on the GER1500 and the spectrometer used to sample irradiance in the snow can be found in Fisher et al., 2005).

2.3 Snow sampling and analysis

Snow was sampled and analyzed in a manner essentially similar to that described by Dominé et al. (2002; 2004) for physical measurements. Polyethylene gloves were used to avoid contamination, and snow was sampled into polyethylene sampling vials. For each snow layer, two sets of triplicate samples were taken for anions and cations, respectively. Samples were usually melted and analyzed on site by ion chromatography within 12 h of sampling, and in a few cases were stored frozen for up to 48 h before analysis.

Melted snow was analyzed by ion chromatography (Dionex IC mod. DX120 and DX100) using Dionex AS12 and CS12 columns for anions and cations, respectively, as detailed in Allegrini et al. (1999) and Domine et al. (2004). The 3 $\sigma$ limit of detection was between 0.01 $\mu$eq (for NO$_3^-$) and 0.50 $\mu$eq (for Ca$^{2+}$). It was 0.04 $\mu$eq for Na$^+$.

3 Photochemistry and Photochemical Model

3.1 Photochemistry of the Nitrate Ion in the Snowpack

The aim of the optical measurements of the snowpack was to calculate the photolysis rate, $J(NO_3^-)$ of nitrate in the snowpack and to estimate the expected molecular fluxes of gas-phase NO$_2$ and HONO from the snowpack. The photolysis of nitrate in the snowpack can lead to gas phase nitrogen dioxide production or formation of nitrite, NO$_2^-$, (Chu and Anastasio, 2003; Cotter et al., 2003; Domine and Shepson, 2002; Dubowski et al., 2001; 2002; Couch et al., 2000; Dubowski and Hoffmann, 2000; Honrath et al., 2000):

$$NO_3^- + h\nu \rightarrow NO_2^- + O^-$$ (1)

$$NO_3^- + h\nu \rightarrow NO_2^- + O$$ (2)

The resulting radical oxygen anion will form the hydroxyl radical,

$$O^- + H_2O^+ \rightarrow OH + H_2O$$ (3)

The source of HONO in the snowpack is probably from one or more of the following three sources:

1) Photo production of nitrite, NO$_3^-$, from nitrate photolysis in snowpack. The nitrite is protonated and desorbs from the snow grains as HONO.

$$NO_2^- + H_2O^+ \rightarrow HONO + H_2O$$ (4)

2) Recombination of OH and NO radical within the snowpack, (Mack and Bolton, 1999).

$$NO + OH \rightarrow HONO$$ (5)

The sources of NO may be the photolysis of nitrate that is more photolabile than nitrate under natural sunlight. The source of OH radicals can be the photolysis of nitrate and hydrogen peroxide. The photolysis of H$_2$O$_2$ may be a more efficient production route to produce OH radicals than nitrate depending on the concentrations of H$_2$O$_2$ and nitrate within the snowpack, (Chu and Anastasio, 2005).

$$NO_2^- + h\nu \rightarrow NO + O^-$$ (6)

3) Electron transfer reactions from photosensitized phenoxy groups in organic matter to NO$_2$. This mechanism has been demonstrated by George et al., (2005) on the surface of organic bulk substrates during laboratory experiments, but not on ice surfaces. A recent study (Stemmler et al., 2006) shows that HONO is produced in photo-induced reactions on natural humic acids from NO$_3^-$, possibly on sea salt or ice surfaces with adsorbed organic photosensitizers.
Table 1. Observed HONO fluxes [nmol m\(^{-2}\) h\(^{-1}\)].

|                  | All data | Detected >-0.3 | Detected ≤-0.3 | Below method detection limits |
|------------------|----------|----------------|----------------|-----------------------------|
| N                | 1440     | 675            | 413            | 352                         |
| Minimum          | –26.111  | 0.300          | –26.111        | –0.300                      |
| Maximum          | 29.566   | 29.566         | –0.303         | 0.298                       |
| Mean             | 0.229    | 1.750          | –1.831         | 0.022                       |
| Median           | 0.579    | 3.140          | –3.131         | 0.023                       |
| Upper 95% CI\(^{(a)}\) | 0.801    | 3.439          | –2.776         | 0.039                       |
| Lower 95% CI\(^{(a)}\) | 0.358    | 2.841          | –3.487         | 0.007                       |
| Std. Dev.        | 4.280    | 3.961          | 3.674          | 0.154                       |

(a) CI=Confidence Interval.

If mechanism 3 is significant a fraction of NO\(_2\) formed in Reaction (1) might be converted to HONO. A flux for that case, however, cannot presently be estimated because of the lack of photo-physical and choromophore abundance data.

These mechanisms (1–3) all require nitrogen oxides and the sources of these nitrogen oxides is thought to be the photolysis of nitrate. Thus, we consider the photolysis of nitrate to produce nitrogen dioxide as a measure of snowpack phototochemistry, and photolysis of nitrate to produce nitrite as an estimation of the upper limit of HONO production. The estimate of the flux of HONO from the snowpack is an upper limit because the conversion of nitrite on the snowpack to gaseous HONO is probably not 100% efficient and because HONO may undergo reactions in the snowpack before being emitted to the atmosphere.

3.2 Calculating photolysis rates

The photolysis frequency for Reaction (1) is estimated from a coupled atmosphere-snow radiative-transfer model, TUV-snow, (Lee-Taylor and Madronich, 2002). The model requires knowledge of the optical properties of the snowpack (the scattering and absorption cross-sections), the concentration of nitrate in the snowpack and downwelling irradiance on the snowpack. The cross sections for scattering and absorption are derived from the measured light transmission (e-folding depth, \(\varepsilon\)) and albedo of the snowpack measured at Browning pass. The model is described fully in Lee-Taylor and Madronich (2002) and will only be described very briefly here. The model treats the optical properties of the snowpack empirically with two constants \(\sigma_{\text{scatt}}\) and \(\sigma_{\text{abs}}\), which describe the scattering and absorption properties of the snowpack. With these data the wavelength-resolved actinic flux, \(F(\lambda)\), at each point in a 1 m thick snowbank can be calculated using the DISORT code in the TUV-snow model. The absorption spectrum of nitrate in a thin film of ice, \(\sigma(\lambda)\), of Burley and Johnson (1992) was used. For Reaction (1) to produce NO\(_2\) (and thus HONO), the quantum yield data for ice of Dubowski et al. (2002) was used. A temperature-independent value of \(1.5 \times 10^{-3}\) was taken from Dubowski et al. (2002) and assumed to supersede the older value of \(4.5 \times 10^{-3}\) (Dubowski et al., 2001). With these quantities one can calculate the photolysis frequency, \(J\), according to Eq. (7):

\[
J = \int \sigma(\lambda) \Phi(T) F(\lambda) d\lambda
\]

The photochemical predictions of the model have been shown to be consistent with laboratory measurement of nitrate photochemistry, (Phillips and Simpson, 2005).

4 Results

4.1 HONO Measurements

Table 1 lists the statistics of the observed HONO fluxes during our campaign split for data below and above the method detection limit. To identify periods during which HONO fluxes were actually significantly different from 0, we performed statistical analysis and t-tests for data above the detection limits in bins of varying length in time. Consecutive data above (or below) the detection limits of \(\pm 0.3\) nmol m\(^{-2}\) h\(^{-1}\) fall into an individual bin. The length in time of these bins is shown in Fig. 2c by the size of the symbols, with the smallest size representing a single 10 min measurement, and the largest symbol representing a 200 min (20 measurements) period. Figure 2 shows the averages of the detected data in these bins. The median positive flux was 1.66 nmol m\(^{-2}\) h\(^{-1}\) (mean 2.86; 340 bins); the median negative flux was –1.74 (mean –2.91, 237 bins). Compared to typical Arctic values of HONO and NOx emissions of around 40 nmol m\(^{-2}\) h\(^{-1}\) (Beine et al., 2002, Honrath et al., 2002) these values are very small.

Figure 3 shows a detailed view of our measured mixing ratios for DOY 325-330. A diurnal noon time maximum, correlated with the insolation can be discerned in the HONO
mixing ratios at the two heights with some goodwill. The maximum diurnal variation observed here is 1 pmol/mol.

Dibb et al. (2004) measured HONO mixing ratios in the atmosphere (of up to 70 pmol mol$^{-1}$) and in snowpack interstitial air at South Pole. Since values were 5 times higher in snowpack air than in the atmosphere, they concluded that the snowpack must be a strong source of HONO but they did not actually quantify fluxes. Our mixing ratios were mostly below 5 pmol mol$^{-1}$; the difference in mixing ratios is expected comparing the stable and stratified atmosphere at South Pole (Davis et al., 2001) with a windy site near the coast. Still, Oncley et al. (2004) measured average NOx fluxes at South Pole of ca. 23 nmol m$^{-2}$ h$^{-1}$ (NO $\sim$15 nmol m$^{-2}$ h$^{-1}$). If the chemistry is comparable to what was observed at Alert (Beine et al., 2002), NOx and HONO fluxes should be of similar magnitude.

4.2 Snow structure

The snow surface at Browning pass is shaped by wind. In late October, it consisted of hard windpacks with a positive relief and of softer layers in hollows, each type covering roughly half of the surface. As observed during our stay, snow layers are formed by wind events that remobilize recent precipitation and exposed layers, and are always discontinuous. As a result, the apparently similar-looking windpacks that outcropped were in fact outcrops of different layers (perhaps 2 to 5) formed at different times, so that spatially variable snow chemistry and physics can be expected. During our stay, snowfalls took place on DOY 311/312 (6–7 November) (<1 mm water equivalent), DOY 322/323 (17–18 November) (12 mm) and 25 November (DOY 330) (2.5 mm). The last 2 snowfalls took place under low winds and formed continuous layers that were eventually windblown to form discontinuous layers. Other thin layers with a very small spatial coverage were formed in hollows by wind in the absence of precipitation. Often, these layers were dark, presumably because they had incorporated terrigenous material. The main four surface layers observed were thus (a) hard windpack, (b) soft windpack, (c) recent windblown snow that has been advected in and that had not yet sintered, (d) fresh precipitation, i.e. recent snowfall before it was windblown. The variations of the surface coverage of these 4 snow types are shown in Fig. 4. The snow stratigraphy is composed of alternating hard windpacks formed of small rounded grains, including an incredibly hard layer of density 0.61 (no melting visible) 40 cm down in the deep pit we dug, and softer layers composed of larger crystals showing some variable degrees of faceting, of density as low as 0.3. Numerous ice layers, ice lenses, melt-freeze crusts and layers where some frozen percolating water was present were seen throughout the snowpack, suggesting frequent melting events, and not just in summer. Signs of frozen water were more frequent in softer layers. 120 cm down, and for a depth of at least 70 cm, all the snow layers showed signs of frozen water, and their density was as high as 0.7. This was certainly the summer melt layer. A 3 m probe from the bottom of our 1.9 m pit could not reach solid ice, indicating that our location was in the lower part of the accumulation zone of the glacier.

![Fig. 2. (a) Timeseries of HONO mixing ratio measured at the lower inlet (red symbols) and the upper inlet (blue symbols). (b) Time-series of the friction velocity $u_*$ [m s$^{-1}$]. (c) Timeseries of averaged HONO fluxes above the detection limits of $\pm$ 0.3 nmol m$^{-2}$ h$^{-1}$. The data are averaged in bins of variable length; the length in time of these bins is shown by the size of the symbols. The smallest size representing a single 10 min measurement, and the largest symbol representing a 200 min (20 measurements) period.](www.atmos-chem-phys.net/6/2569/2006/Atmos. Chem. Phys., 6, 2569–2580, 2006)
4.3 General trends in snow chemistry

We analyzed 148 triplicate snow samples from Browning Pass, almost all taken within 2 km from our air sampling site, and >70% taken within 100 m. Table 2 sums up correlations and ratios between major soluble mineral ions. It is clear that sea salt has a major influence on snow composition, as most major ions are correlated to [Na$^+$]. Cl$^-$ and Mg$^{2+}$ show ion/Na$^+$ ratios very close to those of sea water. [K$^+$] shows the best correlation with [Na$^+$], but surprisingly the ratio, [K$^+$]/[Na$^+$] is 50% greater than that of sea water. Furthermore, Cl$^-$ and Na$^+$ were usually the most abundant ions on a molar basis, sometimes exceeding 1000 µeq, while Ca$^{2+}$, SO$_4^{2-}$ and Mg$^{2+}$ were also abundant, reaching 200 µeq. Concentrations of NO$_3^-$ and K$^+$ reached 50 and 40 µeq, respectively. The sea is just a few km from our site, so the trends of Table 2, and the predominance of Na$^+$ and Cl$^-$ were expected. Table 2 shows that SO$_4^{2-}$ has a source other than sea salt, and this is presumably the oxidation of dimethyl sulfide, with a possible volcanic contribution from Mount Melbourne, some 40 km away. The influence of marine biogenic products is expected, as marine organic compounds have been identified in snow in this area (Cincinelli et al., 2001). Ca$^{2+}$ also has an extra source, most likely terrigeneous particles, as a lot of rock outcrops are present nearby. These are mostly granite, but also micashists, and volcanic rocks are present near Mt. Melbourne. Some snow that was windblown during our campaign had a yellow to brownish color, indicating episodes of fairly high concentrations of mineral aerosols. All the snow samples with a high concentration of ions were from the snow that was remobilized during our presence. From their ion balance, most of the snows were acidic. A few samples (20%) were alkaline. All samples from the hard windpacks and from the 25 November (DOY 330) fall were acidic (pH 5.0 to 5.5). 30% of recently windblown samples were alkaline. There was no clear correlation between acidity and total ion concentrations, although alkaline samples showed a slight tendency to be more concentrated that acidic ones. The range of concentrations was large; however, that just one highly concentrated sample could change the trend. Regarding NO$_3^-$ in snow, it is necessary to discuss separately aged and freshly precipitated snows.

4.4 Origins of nitrate in snow

Figure 5a shows the excellent correlation between the concentrations of NO$_3^-$ and Na$^+$ in aged snows. The most likely explanation is that gas phase nitric acid reacted with sea salt to release HCl and form NaNO$_3$ (Fenter et al., 1994). Na$^+$ concentrations are a function of sea salt mass, while the rate of sea salt reaction depends on aerosol surface area. The existence of this excellent correlation suggests that the size distribution of sea salt aerosol varies little at Browning pass. This seems reasonable, given the existence of the Ross sea
Table 2. Correlations between the concentrations of ions and those of Na+, and comparison of the ion/Na+ ratio to that of sea salt.

| Ion               | Cl− | SO4^{2−} | NO3− | Ca^{2+} | Mg^{2+} | K+  |
|-------------------|-----|----------|------|---------|---------|-----|
| Correlation with Na+(a) | 0.88 (0.95)(b) | 0.915 | 0.972 | 0.889   | 0.992   | 0.996|
| Molar ratio [ion]/[Na+]   | 0.951 (1.22)(b) | 0.108 | 0.052(c) | 0.090 | 0.110 | 0.033 |
| Sea salt ratio [ion]/[Na+] | 1.165 | 0.060 | 0.00003 | 0.022 | 0.113 | 0.022 |

(a)=Linear correlation coefficient for least square fit.
(b)=Value if four outliers, presumably caused by highly fractionated sea salt, are removed.
(c)=aged snows only.

Polynya, and among the possible sources (Domine et al., 2004) sea salt aerosol probably comes mostly from nearby sea spray. There is also a good correlation between [NO3−] and [Ca^{2+}] (R²= 0.918), and it could be argued that NO3− in aged snow comes from the reaction of mineral aerosol with gas phase HNO3. This reaction is fast, with a reaction probability γ >0.01 on many minerals (Hanisch and Crowley, 2001). However, since [Ca^{2+}] and [Na+] are well correlated, a [NO3−]−[Na+] correlation results also in a [Ca^{2+}]−[NO3−] correlation, without necessarily a chemical reason. Furthermore, since the correlation with Ca^{2+} is not as good, our preferred interpretation is that NO3− in aged snows is present as NaNO3, after acid attack of sea salt by HNO3.

Figure 5b shows that there is little correlation between [NO3−] and [Na+] for fresh snow samples, i.e. in precipitated snows before they were remobilized by wind. The ion balance of almost all fresh samples is acidic, with corresponding pH values in the range 5–6. The ionic concentrations are also much lower than for many aged snows in the preceding figure. Our interpretation is that the source of NO3− in these snows is gas phase HNO3 that dissolved in the ice to form a solid solution (Thibert and Domine, 1998). Adsorbed HNO3 may also contribute to the NO3− signal (Sokolov and Abbatt, 2002). Back trajectory calculations (10 day isentropic cluster back trajectories at three different potential temperatures), using the British Atmospheric Data Center model, indicate that air masses that generated the precipitations had spent the past few days over the Antarctic continent. The back trajectory is consistent with the low Na+ concentration, and explains why there is no correlation between [Na+] and [NO3−].

In summary, in fresh snows, our data suggest strongly that NO3− is present as dissolved or adsorbed nitric acid taken up from the gas phase, while in aged snows NO3− is present as NaNO3, presumably because of the reaction of atmospheric nitric acid with sea salt.

4.5 Optical properties of the snow

Since photochemistry takes place mostly in the top few cm or tens of cm in the snowpack (Simpson et al., 2002; Leetaylor, 2002; Warren, 1982) our chemical and optical studies focused on the surface snow layers. Snowpits were dug for optical measurements under each one of the main 4 snow types discussed above and in Fig. 4. Seven snowpits were studied in detail, the objective being to obtain optical properties averaged over several snow layers found under and including each type of surface layer considered. The thin snow
layers found at Browning Pass did not allow the optical properties of just the surface layer to be obtained in isolation. In each pit, the stratigraphy had to be simplified to allow analysis of the optical data and modeling of the fluxes of nitrogen oxides leaving the snowpack. The detailed stratigraphy, calculation of $\sigma_{\text{scatt}}$, $\sigma_{\text{abs}}^+$ and $\varepsilon$, and analysis of these data are described in a separate future paper, and details of the method can be found in (Fisher et al., 2005). Table 3 lists the optical and physical properties of typical snowpacks under and including each type of surface layer, together with the nitrate concentrations of these surface layers. The values of the physical parameters in Table 3 reflect in part the structure and the grain size and shape of the surface layers, and in part the effect of heterogeneities in these layers that include the presence of sub-layers and of frozen water. They also reflect the properties of the underlying layers, as no snow layer (except the fresh windblown) was thick enough to be measured in isolation (i.e. to be treated as semi-infinite). Thus values of $\sigma_{\text{scatt}}$, $\sigma_{\text{abs}}^+$, and $\varepsilon$ presented in Table 3 describe the measured optical properties of the surface snow at Browning Pass, but are not to be interpreted as characteristic of the surface layer in isolation.

4.6 Calculating molecular fluxes of NO$_2$ and HONO from the Snowpack

As shown in Fig. 4, the temporal changes in the proportion of outcropping layers and the resulting changes to the snowpack in Browning pass are complex. The total calculated flux of NO$_2$ (or HONO) sourced from the snowpacks at Browning Pass depended on the proportions of snowpack present. The molecular flux from each surface snow layer listed in Table 3 was calculated and the weighted sum of the molecular fluxes from these snowpacks according to Fig. 4 was calculated, in order to compare with the measured molecular flux.

The snow-atmosphere radiation transfer model was run with daily measured ozone columns, each one of the 4 snow types in Table 3 and varying optical depths of cloud (asymmetry factor, $g=0.85$ and altitude $\sim2$ km) to coarsely replicate the measured solar downwelling irradiance (Fig. 6).

When a coarse agreement between modeled and measured values had been achieved the values of $J(\text{NO}_3)$ were calculated in a 1 m deep slab under each type of surface layers. Coarse agreement between the modeled and measured irradiance was refined by multiplying the calculated photolysis rates, $J(\text{NO}_3)$, by the ratio of the measured and modeled irradiance, $I$, at 350 nm, $I_{\text{measured}}/I_{\text{model}}$. While this is a crude method the adjustment is small ($\sim20\%$) and allows molecular fluxes of NO$_2$ and HONO to be estimated.

The depth, $z$, integrated photolysis rate (transfer velocity), $v$ (NO$_2$), is calculated by

$$v (\text{NO}_2) = \int J (\text{NO}_3 \rightarrow \text{NO}_2) dz$$ \hspace{1cm} (8)

The molecular flux, $F$, is the product of the nitrate concentration in snow, $[\text{NO}_3^-]$ and the transfer velocity

$$F (\text{NO}_2) = v (\text{NO}_2) \times [\text{NO}_3^-]$$ \hspace{1cm} (9)

The analysis assumes that (I) $[\text{NO}_3^-]$ is depth independent, and (II) all the photo-produced NO$_2$ and HONO can exit the snowpack with 100% efficiency and do not undergo further photolysis or reaction within the snowpack. Either assumption may not always be true. Assumption (II) implies the molecular fluxes calculated in this paper are upper limits. Using Eqs. (8) and (9) the molecular flux for NO$_2$, $F$ (NO$_2$) was calculated and is shown in Fig. 7. The flux of HONO, $F$ (HONO), was estimated as an upper limit from the reaction to produce NO$_2$ (Fig. 7).

Comparing the flux estimates from the photochemical modeling with our actual measurements a significant discrepancy is obvious. While the maximum observed HONO fluxes are of the same order of magnitude as the modeled estimates, the majority ($>90\%$) of measured HONO fluxes were below 5 nmol m$^{-2}$ h$^{-1}$.
Fig. 6. The measured downwelling surface irradiance recorded throughout the campaign at 350 nm (blue, units of W m$^{-2}$ nm$^{-1}$, right scale). Also plotted is the calculated quantum yield for nitrate photolysis to yield NO$_2$ in ice (Reaction 1, units of 10$^{-3}$, red, left scale). The variation of the quantum yields is due to the temperature of the snowpack recorded at 12.5 cm depth throughout the campaign.

5 Discussion

5.1 The case of aged snows

The particularity of this study is the quasi non-existent HONO fluxes, despite the normal to high NO$_3^-$ concentrations in snow. The currently hypothesized mechanism of HONO formation from NO$_3^-$ photolysis is discussed above, though the details are not well known. From this mechanism alone we would expect daily noontime maximum HONO fluxes of ca. 10 nmol m$^{-2}$ h$^{-1}$, which were not found. Other studies have reported or inferred HONO fluxes associated with NO$_3^-$ at Alert, Summit, and South Pole (Beine et al., 2002; Honrath et al., 2002; Dibb et al., 2004). We argued that NO$_3^-$ is present mostly as NaNO$_3$ in aged snows at our study site, and this seems to be peculiar. During this campaign, we also sampled snow at other nearby coastal sites such as ice tongues and inland on the plateau, all within 200 km from our site, and the correlation between Na$^+$ and NO$_3^-$ was not observed (R$^2$=0.02). Neither did we observe a good correlation between both ions during previous campaigns at Alert and Ny-Ålesund (Domine et al., unpublished results) and it is therefore tempting to relate both characteristics: no HONO fluxes and NO$_3^-$ present as a sodium salt in aged snows.

The state of salt particles in snow (i.e. solid or liquid) is not clear. From the NaCl – H$_2$O phase diagram (e.g. Hall et al., 1988), given the ionic concentrations present and the temperatures encountered, usually between −5 and −15°C, NaCl could not induce melting, and the concentrations of the other ions measured cannot modify this conclusion.

However, ions present in snow may favor the existence of a quasi-liquid or brine layer at the surface of snow crystals, as evidenced in laboratory experiments by Cho et al. (2002). The experiments of Cho et al. (2002) used NaCl concentrations much higher than those present in our snow samples, and the phase behavior for low salt concentrations still has to be established, especially since a basic extrapolation of the data of Cho et al. (2002) to the concentrations found in Browning pass suggests that it may be unimportant here. This conclusion is reached by considering bulk concentrations. Locally, a sea salt particle may induce very high ionic concentrations on a snow crystal surface, causing the formation of a quasi-liquid layer, or actual melting. To our knowledge, the impact of a salt particle on an ice surface has not been investigated and we are not able to conclude on the state of sea salt ions, and of NaNO$_3$ on the surface of snow crystals. It is still reasonable to suggest that NO$_3^-$ in aged snows remains trapped in salt particles or that, even if a liquid phase is formed, NO$_3^-$ reactivity differs from that of NO$_3^-$ present as nitric acid dissolved or adsorbed in/on snow crystals (Beine et al., 2002; 2003).

Perhaps because of matrix effects in the solid state, or solvent "cage effects" in the liquid state, the products of NO$_3^-$ photolysis cannot escape. Another possibility is more simply that HONO, if formed in a salt or brine medium, reacts rapidly with sea salt, presumably to form NaNO$_2$. This reaction is plausible, although available laboratory studies are inconclusive (Vogt and Finlayson-Pitts, 1994).

5.2 The case of fresh snows

The case for aged snows does not explain why no fluxes were observed out of fresh snows, especially after the heavy fall of...
DOY 322/323 (17–18 November), when the NO$_3^-$ concentration was larger than the Cl$^-$ concentration. In fact, the snowpack was particularly inactive with respect to HONO uptake/release when that snowfall covered the surface (Fig. 2), until it was partially windblown by a katabatic wind on DOY 326 (21 November). The mechanism of HONO formation from NO$_3^-$ photolysis is not well known. It is possible that it involves the reaction of NO$_2$ (produced from NO$_3^-$ photolysis) with specific photosensitized organics (George et al., 2005; Stemmler et al., 2006). Species that are expected to efficiently transfer electrons to NO$_2$ include electron donors such as phenols as photosensitizers in reaction cycles with aromatic ketones (George et al., 2005). The organic surface photochemistry was found in soils and other surfaces containing humic acids (Stemmler et al., 2006). If snow surfaces contain these organic chemicals, the HONO flux may depend on the concentration of photosensitizer in the snow; with low concentrations of organic photosensitizers the fluxes of HONO will be low and perhaps non-detectable.

We did not measure organic compounds in the snow or in the air, and we are therefore left to speculate on their concentrations, based on other studies. In general, Antarctic snow may be suspected of having lower concentrations of organic compounds than Arctic snow, because many of the Arctic sources such as soil dust (e.g. at Alert or Ny-Ålesund, (Domine et al., 2002) or forest fire plumes (e.g. at Summit, (Legrand and De Angelis, 1996)) are absent or reduced. Nevertheless, the sea and in particular the nearby Ross sea polynya can be a source of organic compounds, as found in snow samples near our site by Cincinelli et al. (2001). Dibb and Arsenault (2002) measured formic and acetic acids at Summit, Greenland, and at South Pole in the atmosphere and in snowpack interstitial air. Both sites are at comparable elevations and distances from the sea. They found concentrations about twice as high at Summit, supporting the idea that the concentrations of organic compounds in the Antarctic are lower than in the Arctic. However, Grannas et al. (2004) analyzed total organic carbon in snow and found similar values at South Pole and Summit. More importantly, the complex humic style material that Grannas et al. (2004) observed is capable of the type of photosensitization reactions that George et al. (2005) and Stemmler et al. (2006) describe. In summary, the data available on organic compounds in polar snow do not seem to support the idea that the lack of HONO fluxes could be due to a lack of organic photosensitizers. Furthermore, Dibb et al. (2004) inferred strong HONO fluxes out of the snow at South Pole, strengthening our impression that our site is peculiar.

The above considerations on the composition of Antarctic snow are general and do not allow any conclusion on the reactivity of a specific snow layer at a given site. Our monitoring of snow layers showed that fresh snow always had low ionic concentrations. Most of the mineral ion loading came from dry deposition due to wind pumping or while snow was airborne. The same mechanisms that deposit soluble mineral ions probably also deposit organic compounds. Both Arctic and Antarctic measurements cited above involved aged snows and the conclusion that they had a significant organic loading may thus not apply to our fresh snows, that may well have been depleted in organic compounds. We therefore suggest that the concentration of organic compounds was too low to support production of detectable HONO fluxes. This is consistent with our back trajectory calculations that showed that air masses generating fresh precipitation came from the continent. The highest HONO flux out of the snow was measured on DOY 327 (22 November), just after a katabatic wind partially remobilized the thick 17–18 November snowfall that was particularly unreactive (Fig. 2). The
katabatic wind increased the sea salt content of the snowfall snowlayer throughout, with Na⁺ concentrations increasing from 1–5 µeq to 10–20 µeq. It is reasonable, although speculative at this stage, to suggest that this dry deposition of sea salt was accompanied by the deposition of organic compounds of marine origin that allowed HONO formation. The HONO flux out of the snow was short-lived, possibly because the salt subsequently trapped HNO₃ as NaNO₃. More field and laboratory experiments will be necessary to test these suggestions.

6 Conclusions

We have attempted to sum up some of the possible (photo) chemical and physical reaction pathways for HONO production suggested by this study in Fig. 8. Our somewhat speculative suggestions to explain the near-absence of detectable HONO fluxes at Browning pass is that:

1. In aged snow samples, i.e. snows that have been remobilized by wind, NO₃⁻ is present as NaNO₃, and there is a significant concentration of sea salt. The absence of HONO emissions may be due to the fact that the NO₃⁻ is present as NaNO₃, which does not allow photolysis products to escape, possibly because of matrix or solvent cage effects in the phase in which NaNO₃ is contained, probably NaCl, or because HONO is taken up by sea salt.

2. In fresh snow samples, i.e. in recent snow falls before their remobilization by wind, NO₃⁻ is present as dissolved and/or adsorbed HNO₃, as also suggested from studies at Alert and Ny-Ålesund (Beine et al., 2002; 2003). Products of the photolysis of NO₃⁻, including HONO, are thus expected. A possibility is that HONO is produced from NO₂ via electron transfer from photosensitized organic compounds, and that fresh snows had concentrations of organic photosensitizers too low for this photochemistry to be efficient. When wind deposited organic compounds to the snow, the accompanying sea salt trapped all the NO₃⁻ as NaNO₃, preventing HONO production as described above.

In summary, HONO production from nitrate does strongly depend on its physical (surface or volume) and chemical (ice or salt) environment. It is clear that HONO production could follow several pathways, the prevalence of one over the other depending on several aspects of snow chemistry, such as the chemical form of NO₃⁻, the concentration of organic photosensitizer, etc. Well designed laboratory experiments, with different chemistry of the (natural) ice substrate and ideally a control over the location of NO₃⁻ in the substrate, seem necessary to solve the puzzle.

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