Atmospheric nitric oxide and ozone at the WAIS Divide deep coring site: a discussion of local sources and transport in West Antarctica

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Abstract. The first measurements of atmospheric nitric oxide (NO) along with observations of ozone (O3), hydroperoxides (H2O2 and MHP) and snow nitrate (NO3-) on the West Antarctic Ice Sheet (WAIS) were carried out at the WAIS Divide deep ice-corining site between 10 December 2008 and 11 January 2009. Average ±1σ mixing ratios of NO were 19 ± 3 pptv and confirmed prior model estimates for the summer boundary layer above WAIS. Mean ±1σ mixing ratios of O3 of 14 ± 4 ppbv were in the range of previous measurements from overland traverses across WAIS during summer, while average ±1σ concentrations of H2O2 and MHP revealed higher levels with mixing ratios of 743 ± 362 and 519 ± 238 pptv, respectively.

An upper limit for daily average NO2 and NO emission fluxes from snow of 8.6 × 108 and 33.9 × 108 molecule cm−2 s−1, respectively, were estimated based on photolysis of measured NO3− and nitrite (NO2−) in the surface snowpack. The resulting high NOx emission flux may explain the little preservation of NO3− in snow (~30 %) when compared to Summit, Greenland (75–93 %). Assuming rapid and complete mixing into the overlying atmosphere, and steady state of NOx, these snow emissions are equivalent to an average (range) production of atmospheric NOx of 30 (21–566) pptv h−1 for a typical atmospheric boundary-layer depth of 250 (354–13) m. These upper bounds indicate that local emissions from the snowpack are a significant source of short-lived nitrogen oxides above the inner WAIS.

The net O3 production of 0.8 ppbv day−1 triggered with NO higher than 2 pptv is too small to explain the observed O3 variability. Thus, the origins of the air masses reaching WAIS Divide during this campaign were investigated with a 4-day back-trajectory analysis every 4 h. The resulting 168 back trajectories revealed that in 75 % of all runs air originated from the Antarctic coastal slopes (58 %) and the inner WAIS (17 %). For these air sources O3 levels were on average 13 ± 3 ppbv. The remaining 25 % are katabatic outflows from the East Antarctic Plateau above 2500 m. When near-surface air from the East Antarctic Plateau reaches WAIS Divide through a rapid transport of less than 3 days, O3 levels are on average 19 ± 4 ppbv with maximum mixing ratios of 30 ppbv. Episodes of elevated ozone at WAIS Divide are therefore linked to air mass export off the East Antarctic Plateau, demonstrating that outflows from the highly oxidizing summer atmospheric boundary layer in the interior of the continent can episodically raise the mixing ratios of long-lived atmospheric chemical species such as O3 and enhance the oxidative capacity of the atmosphere above WAIS.

1 Introduction

Over the last decade, a large number of field and lab studies have provided evidence of the importance of snow photochemistry on the chemical composition of air above snow-covered surfaces in the polar and mid-latitudes (Grannas et al., 2007, and references therein). The upper snowpack is seen not only as a chemical reservoir but as a chemical reactor. Trace gases emitted by the snowpack include hydrogen peroxide (H2O2), formaldehyde (CH2O) and nitrogen oxides (NOx = NO + NO2), and increase the oxidizing potential of the atmospheric boundary layer through production of
the hydroxyl radical (OH) and ozone (O₃) (Grannas et al., 2007; Thomas et al., 2012). Measurements of atmospheric and snow concentrations and also fluxes from snow surfaces indicate that the polar snowpack emits NOₓ mainly through the photolysis of nitrate (NO₃) in near-surface snow (Grannas et al., 2007; Bauguitte et al., 2012; Frey et al., 2013). Snowpack emissions of NOₓ can contribute significantly to the NOₓ (NOₓ + other oxidized nitrogen species) budget above snow, as observed in continental and coastal Antarctica (Jones et al., 2000; Frey et al., 2013). Bauguitte et al. (2012) reached a similar conclusion at Halley station by highlighting that halogen chemistry over the Antarctic coast controls the lifetime of NOₓ species and reduces the nitric oxide (NO) mixing ratios. When atmospheric turbulence is low or atmospheric boundary layer depths are shallow, these emissions contribute to high NOₓ levels, several hundred pptv, as observed at South Pole and over the East Antarctic Plateau (Davis et al., 2004, 2008; Slusher et al., 2010; Frey et al., 2013). Chen et al. (2004) and Mauldin et al. (2004) showed that at South Pole these high levels of NOₓ shift the HOₓ (OH + HO₂) partitioning towards OH:

\[ \text{NO} + \text{HO}_2 \rightarrow \text{NO}_2 + \text{OH}. \]  

(R1)

NOₓ emissions from snow can also lead to net production of O₃ over the East Antarctic Plateau (Helmg et al., 2008a, and references therein), and thereby shift HOₓ partitioning of the overlying atmosphere.

Above the West Antarctic Ice Sheet (WAIS), enhanced snowpack emissions of NOₓ associated with events of stratospheric O₃ depletion may lower the formation rate of atmospheric H₂O₂ (Frey et al., 2005). Photochemical modeling suggests that atmospheric H₂O₂ is sensitive to the NO background, opening up the possibility of constraining past NOₓ and OH levels using the H₂O₂ ice-core record (Frey et al., 2005, 2006). Therefore, investigation of the current atmospheric boundary-layer photochemistry at a WAIS ice-core drilling site is essential to interpret ice-core records of photochemically active species such as H₂O₂ and NO₃⁻. This can also provide information regarding the contribution of air advection from the East Antarctic Plateau to levels of oxidants above WAIS, similar to Legrand et al. (2009) who observed outflows from the Antarctic Plateau increasing O₃ concentrations at the East Antarctic coastal site Dumont D’Urville.

So far, atmospheric-sampling campaigns in the Antarctic have taken place at existing stations on the coast (Dumont D’Urville, Halley, Neumayer) and on the East Antarctic Plateau (South Pole, Dome C). Only recent airborne campaigns (Eisele and Davis, 2008; Slusher et al., 2010) or scientific overland traverses (Frey et al., 2005) provided information on the composition of the lower atmosphere across the interior of the continent. Thus, spatial data coverage of the lower atmosphere in Antarctica is still sparse and little is known about the variability of O₃ and its precursors above WAIS. For example, the first ground-based measurements across WAIS lasting several days included atmospheric records of H₂O₂, methyl hydroperoxide (MHP), CH₂O, O₃ but not NOₓ (Frey et al., 2005).

The current study includes multi-week first observations of NO and complementary measurements of atmospheric O₁, H₂O₂, MHP and of surface-snow H₂O₂, NO₃⁻ and nitrate (NO₃⁻) in the WAIS region. The aims were to determine the summer composition of the lower atmosphere in the interior of West Antarctica and the relative importance of local production versus air mass transport on the local atmospheric budgets of NOₓ and O₃.

### 2 Methods

From 10 December 2008 to 11 January 2009, atmospheric concentrations of NO and O₃ were continuously measured at WAIS Divide (local time: LT = UTC − 07:30). Mixing ratios of ROOH (H₂O₂ and MHP) were recorded between 31 December 2008 to 7 January 2009. Snow samples were collected daily from the surface and weekly from 30 cm snow pits for chemical analysis of NO₃⁻, NO₂⁻ and H₂O₂.

Atmospheric sampling took place 5 km NW of the WAIS Divide drilling camp (79.467° S, 112.085° W, 1766 m a.m.s.l., http://www.waisdivide.unh.edu). All instruments were run out of a Polarhaven tent heated by a preway heater. Atmospheric measurements were made 1 m above the snow, 10 m upwind (prevailing winds from NE) from the tent, with ambient air drawn through an insulated and heated PFA (1/4″ I.D.) intake line (typically 1.4 STP – L min⁻¹) of 12 m for ROOH, and of 20 m for NO and O₃. In an attempt to minimize artifacts in our atmospheric records, the two generators (3.5 and 5 KW) that provided electricity to the lab were located about 30 m downwind from the sampling lines, and all activities around the site were restricted. However, the heater exhaust was located on the top of the Polarhaven tent.

#### 2.1 Atmospheric sampling

NO was measured using a modified chemiluminescence instrument used previously at South Pole (Davis et al., 2001, 2004). NO mixing ratios recorded at 1 Hz were averaged to 1 min averages. The limit of detection (LOD), defined as 2-σ of the background count rate, was 5 pptv. A two-minute background signal was monitored every 20 min and an automatic 4 min calibration was performed every 2 h by addition of a 2 ppmv NO standard. Due to late delivery of this NO gas standard to the site, calibration was only run during the last 3 days of the campaign. The instrument sensitivity remained fairly constant over the three days, with an average over 16 calibrations of 7.10 ± 0.18 Hz pptv⁻¹, similar to the preseasong value of 7.00 Hz pptv⁻¹ determined in the lab. We therefore used the 3-day average value of these calibrations to process the overall dataset. NO spikes related to pollution from generators and heater exhaust were removed using a moving standard deviation filter with a maximum standard deviation...
of 30 (1.5 times the interquartile range of the dataset). This led to the removal of 25% from the raw NO record.

Surface O$_3$ was monitored at 1 min resolution using a 2B Technologies (Golden, Colorado) O$_3$ monitor, model 205, similar to those previously used in the remote Antarctic such as on the ITASE traverses (Frey et al., 2005) and in an O$_3$ monitoring network (Bauguitte et al., 2011). LOD was 1 ppbv.

Atmospheric ROOH were measured based on continuous scrubbing of sample air followed by separation in an HPLC column and fluorescence detection, described in detail by Frey et al. (2005, 2009a). The detector was calibrated 1–2 times per day with H$_2$O$_2$ solution and MHP standards synthesized in our lab following the protocol described by Frey et al. (2009a). The LOD, 2-$\sigma$ of the baseline, were 87 ppptv for H$_2$O$_2$ and 167 ppptv for MHP. Unexpected variations of the coil-scrubber temperatures may have caused higher LOD than those reported by Frey et al. (2005, 2009a).

### 2.2 Snow sampling

All surface snow and snow pits were sampled in a 7200 m$^2$ clean area upwind from the Polarhaven tent. The top 1 cm of the non-cohesive surface snow, referred to as the skin layer (Frey et al., 2009b; Erbland et al., 2013), was collected daily with a 10 mL glass test tube to assess temporal changes in snow chemistry. Twice during the campaign, the skin layer was sampled simultaneously at five different spots inside the clean area to assess possible local spatial variability of NO$_3^-$, NO$_2^-$ and H$_2$O$_2$.

Weekly snow pits were sampled at 2 cm resolution to a depth of 30 cm, covering the snowpack zone where 85% of NO$_3^-$ photolysis is expected to occur (France et al., 2011). Snowflakes were collected on aluminum foil during the only snow precipitation event observed during the campaign, on 12 December 2008.

All snow samples were collected in 100 mL SCHOTT bottles and kept frozen during storage and transport until analysis 14 months later. The analysis involved melting the snow 1 hour before injecting the sample into a self-built continuous flow analysis (CFA) system, as described by Frey et al. (2006). The LOD, defined as 3-$\sigma$ of the baseline, was 0.4 ppbw for NO$_3^-$, NO$_2^-$ and H$_2$O$_2$. Only values above LOD were used for further calculations. Some loss of NO$_3^-$ in the samples may have occurred between the time of collection and analysis, as Takenaka and Bandow (2007) and O’Driscoll et al. (2012) showed that NO$_2^-$ may be oxidized during freezing and storage.

### 3 Results

#### 3.1 Atmospheric concentrations

The average ±1$\sigma$ (median) of NO over the campaign was 19 ± 31 (10) pptv (Table 1). Some noise remained in the NO dataset after filtering; however, the 4h running median (Fig. 1a) shows very little change in the overall trend of the data due to filtering, with the median value after filtering similar to that of the raw dataset (6 pptv). One-minute data did not exhibit any clear diel cycle (Fig. 1), but 1 h binned data centered on each hour for the measurement period revealed a diel cycle that can be interpreted with the variations of the average solar elevation angle (Fig. 2b). NO mixing ratios increased at 07:00–08:00 LT with a maximum rise of 36% from the daily median of 10 pptv. A decrease was observed afterwards and followed by a second increase of 20% above the median value at 19:00 LT. These peaks in NO occurred as solar elevation angle increased and decreased, with lower values of NO at the maxima and minima of solar elevation angle.

Average ±1$\sigma$ (median) mixing ratios of O$_3$ at WAIS Divide were 14 ± 4 (13) ppbv (Table 1). The mean is two thirds of the 20 ± 2 ppbv average mixing ratio observed at Byrd Station in summer 2002, but is in the range of values from previous measurements between 79.06° S and 85.00° S above the WAIS (Frey et al., 2005) (Fig. 3). Two events of elevated O$_3$ levels were recorded between 24 and 25 December, and between 27 and 29 December, with concentrations in the range of 20 to 30 ppbv (Fig. 1). Concentrations above 25 ppbv were only observed for winds blowing from ENE to SWS. This 135° sector represents 67% of all the wind directions observed during the field campaign (Fig. 4). The hourly binned O$_3$ data (Fig. 2b) show a small diel cycle in phase with solar elevation angle and wind speed. The mixing ratios rose by 5% of the median value (13 ppbv) in the morning, reaching a maximum at 14:00 LT and dropping thereafter in the afternoon.

Concentrations of H$_2$O$_2$ and MHP were measured between 31 December 2008 and 5 January 2009 (Fig. 1). Averages ±1$\sigma$ (medians) were 743 ± 362 (695) and 519 ± 238 (464) pptv for H$_2$O$_2$ and MHP, respectively. Our records are closer to values observed at West Antarctic sites below 1500 m a.m.s.l. and higher than measurements made in the surrounding area (Fig. 3), with mixing ratios of H$_2$O$_2$ that were twice those observed at Byrd station in late November 2002 (Table 1) (Frey et al., 2005). Average ±1$\sigma$ (range) of the MHP: (H$_2$O$_2$ + MHP) ratios were 0.42 ± 0.10 (0.12–0.76). These values are in the range of those previously recorded over WAIS (Frey et al., 2005). Binned values suggest, for both H$_2$O$_2$ and MHP, a diel cycle with respective maximum 44 and 37% above their medians (695 and 464 pptv) observed in the morning (Fig. 2c).
Table 1. Averages ±1σ (medians) of atmospheric and snow concentrations of the chemical species observed at WAIS Divide and nearby sites.

<table>
<thead>
<tr>
<th>Site</th>
<th>Site</th>
<th>atmospheric</th>
<th>skin layer</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>NO (pptv)</td>
<td>O3 (ppbv)</td>
</tr>
<tr>
<td>WAIS Divide</td>
<td></td>
<td>19 ± 31</td>
<td>14 ± 4</td>
</tr>
<tr>
<td>ITASE 00-1a</td>
<td></td>
<td>– –</td>
<td>303 ± 159</td>
</tr>
<tr>
<td>Byrdb</td>
<td></td>
<td>10 ± 2</td>
<td>20 ± 2</td>
</tr>
</tbody>
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*79.38° S, 111.23° W, 1791 m a.m.s.l. (from Frey et al., 2005)
*b 80.02° S, 119.60° W, 1537 m a.m.s.l. (from Frey et al., 2005); NO based on optimum model runs; atmospheric H2O2 from sampling at site RIDS-C (80.00° S, 119.53° W, 1575 m a.m.s.l.); H2O2 in snow from sampling at site RIDS-B in 1996 (79.46° S, 118.04° W, 1650 m a.m.s.l.).

Fig. 1. Atmospheric mixing ratios of NO after filtering and O3 (1 min averages), and of H2O2 and MHP (10-minute averages) during austral summer 2008–2009 at WAIS Divide. The 4 h running median of NO is also shown (red symbols).

3.2 Concentrations in snow

Average ±1σ (median) concentrations of NO2, NO3 and H2O2 in the skin layer at WAIS Divide were 0.6 ± 0.4 (0.5), 137 ± 37 (142) and 238 ± 37 (238) ppbw, respectively (Table 1). Daily concentrations of NO2 in the skin layer showed a decrease of 30 pptw per day ($R^2 = 0.36$) over the campaign (Fig. 5). This decrease represents a rate of 5 % per day of the average concentration of NO2 measured in all of the snow-surface samples. Unlike NO2, NO3 and H2O2 exhibited some variation but no trend was observed for these species.

Fig. 2. Diel variation of (a) average solar elevation angle (SEA) and wind speed, (b) median NO (left axis) and O3 (right axis), and (c) median H2O2 and MHP. Symbols and lines are 1 h binned values centered on each hour; shaded area and error bars indicate the range between the first and third quartiles; local time is UTC – 07:30.
The coefficients of variation of concentrations of NO$_2^\text{−}$, NO$_3^\text{−}$ and H$_2$O$_2$ in the skin layer are 49 %, 26 % and 17 %, respectively. The coefficients of variation for samples collected simultaneously on 1 and 8 January 2009 (Fig. 5) are 17 %, 31 % and 7 % for NO$_2^\text{−}$, NO$_3^\text{−}$ and H$_2$O$_2$, respectively. The similar coefficients of variation for NO$_3^\text{−}$ concentrations in snow imply that spatial variability contributes significantly to the overall variability and thus a temporal trend may be difficult to detect. For NO$_2^\text{−}$ and H$_2$O$_2$, the coefficients of temporal variability (49 % and 17 %, respectively) are more than double those calculated from spatial variability (17 % and 7 %, respectively). The variations of daily concentrations of NO$_2^\text{−}$ and H$_2$O$_2$ in near-surface snow may then be interpreted as temporal trends. Concentrations of H$_2$O$_2$ in the top 5–15 cm of the profile (Fig. 6) may also indicate a temporal trend; seasonal increase in concentrations measured over this period was apparent not only in the top 5 cm of snow, but also down to at least 15 cm. Although there was no new snow accumulation during this period, there was wind redistribution and atmosphere–snow exchange of H$_2$O$_2$ and other atmospheric gas species, or nighttime deposition of fog.

The 30 cm deep profiles of NO$_2^\text{−}$, NO$_3^\text{−}$ and H$_2$O$_2$ illustrated in Fig. 6 represent concentration changes of these species in snow over the last 6 months of 2008, based on local mean annual snow accumulation rate of 0.20 m$_{\text{w eq}}$ yr$^{-1}$ (Banta et al., 2008) and an average snow density of 0.37. Total concentrations of NO$_2^\text{−}$ in the snow column decreased by about 19 % between the first and last snow-pit samplings. NO$_3^\text{−}$ concentrations decrease by 94–188 ppbw over the top 5 cm of snow, reaching ∼30 ppbw below. Total NO$_2^\text{−}$ stored in the 30 cm column decreased by about 65 % across the three snow-pit samplings. Unlike NO$_2^\text{−}$ and NO$_3^\text{−}$, total
Fig. 4. The windrose for WAIS Divide for period of 10 December 2008 to 5 January 2009 and O₃ concentrations from each direction for the same period.

Fig. 5. Surface-snow concentrations and snow concentrations from the 12 December 2008 precipitation of (a) NO₃⁻, (b) NO₂⁻, (c) H₂O₂. The shaded area in (b) represents the NO₂ LOD.

concentrations of H₂O₂ in the top 30 cm of snowpack doubled over the 18 days of sampling. A 233–298 ppbw decrease of H₂O₂ concentrations in the first 10 cm of each snow pit was generally observed.

4 Discussion

4.1 Local photochemistry

4.1.1 Factors controlling atmospheric levels of ROOH, NO, O₃ and snow content of NO₃⁻, NO₂⁻ and H₂O₂

Levels of NO and O₃ measured at WAIS Divide during summer 2008–2009 are slightly higher than coastal values but lower than observed concentrations on the Antarctic Plateau (Fig. 3). The observed mean of NO mixing rates are close to the 10 pptv modeled by Frey et al. (2005) at Byrd, about 160 km from WAIS Divide. Their potential NO concentrations were computed with the NASA Goddard Flight Center (GSFC) point photochemical model that included a snowpack source of H₂O₂ and CH₂O.

The diel cycle of NO (Fig. 2b) does not compare well with either measurements from the East Antarctic Plateau (Dome C) (Frey et al., 2013) or from the West Antarctic coast (Halley) (Bauguitte et al., 2012). Data show a symmetry of the NO diel cycle with respect to local noon that is similar to predictions and observations from Summit/Greenland (Thomas et al., 2012), but with lower mixing ratios and a diel amplitude of 6.5 pptv. The NO variability is mainly in phase with solar radiations but presents two minima around local
noon and midnight and two daily maxima around 07:00 and 19:00 LT. The 36 % rise from the NO median of 10 pptv observed in the morning along with the increase of solar elevation angle suggests that the increase of solar elevation angle enhances the photolytic production of NO from the snowpack. The 2 h binned data of NO and wind speed (data not shown) suggest that higher NO mixing ratios were observed for wind speeds less than 3 m s$^{-1}$, indicating a potential local production of NO from surface snow. The decrease observed after 09:00 LT at high solar elevation angle may result from the decrease of the atmospheric boundary layer height and of the wind speed. Higher wind speeds would result in an increase of turbulent mixing and therefore in efficient vertical upward transport and dilution of surface emissions into a growing volume given by the developing boundary layer (Frey et al., 2013). The increase of NO levels by 20 % of the median value around 19:00 LT may result from the decrease of the atmospheric boundary layer height and wind speed along with solar elevation angle. However, no measurements of the diel variability of the atmospheric boundary layer height were performed during the campaign to confirm this point. These data suggest that both snow photochemistry and physical mixing control the diel cycle of NO at WAIS Divide. Through its different sinks and sources, the NO$_3$ photochemistry may also drive the NO variability but this contribution cannot be defined here as it requires a more detailed analysis using a 1-D model (Thomas et al., 2012).

The interpretation of the 1 month record of O$_3$ in terms of seasonal trend is made difficult given the short period of measurement. However, its comparison with the annual cycles of O$_3$ observed at other Antarctic stations, with concentrations between those observed at Halley and South Pole, suggests a possible influence of both marine and continental air on the WAIS Divide level, and shows characteristic low mixing ratios around the Austral solstice.

The O$_3$ diel cycle (Fig. 2b) is similar to observations from Dome C (Legrand et al., 2009) but with lower amplitude and concentrations. The increase of 5 % of the median value is consistent with O$_3$ produced locally from the snowpack NO$_x$ emissions and confined into a shallow atmospheric boundary layer. Similarly to the NO diel cycle, the following decrease of O$_3$ in the afternoon can be attributed to a deeper atmospheric boundary layer and higher wind speeds that dilute all the produced O$_3$. Because the chemical lifetime of O$_3$ in polar regions is in the order of days (Grannas et al., 2007), it is not expected that photochemical loss is a main driver of the diel cycle of O$_3$. Lack of higher mixing ratios during periods with wind speed less than 3 m s$^{-1}$ (data not shown) suggests that local production of O$_3$ over the surface snow is not the predominant process at WAIS Divide during the austral summer. The amplitude of the diel cycle is relatively low in comparison with the average daily O$_3$ which indicates that another source may be considered to explain the change of O$_3$ over the campaign. The ENE to SWS wind directions associated with high O$_3$ values (Fig. 4) point to a possible contribution of air mass transport of this chemical species at WAIS Divide, e.g. outflow from the East Antarctic Plateau (see Sect. 4.2). A similar conclusion was reached by Helmig et al. (2007a) from the review of other Antarctic station data.

H$_2$O$_2$ concentrations at WAIS Divide are similar to those observed above WAIS at lower latitudes (below 1500 m a.m.s.l.) but 3 times the mixing ratios measured at South Pole (Frey et al., 2009a). As shown in Table 1 and in Fig. 3, the concentrations of H$_2$O$_2$ measured at WAIS Divide are higher than those measured at nearby sites (Frey et al., 2005). A similar comparison between our average MHP mixing ratios and prior measurements across WAIS (Frey et al., 2005, 2009a) is also observed. The average specific humidity of 1.1 g kg$^{-1}$ calculated for the period of ROOH measurement is higher than previous records from nearby sites (Frey et al., 2005). This high value may explain the elevated mixing ratios of H$_2$O$_2$ and MHP measured at WAIS Divide, as water vapor is an important precursor for both species.

H$_2$O$_2$ and MHP exhibited a simultaneous increase with solar elevation angle between 03:00–13:00 LT, followed by a decrease (Fig. 2). The maximum amplitude of H$_2$O$_2$ is observed at 05:00 LT, with a rise of 43 % above the median value of 695 pptv. For MHP, mixing ratios rose to a maximum of 37 % above the median (464 pptv) at 05:00 and 13:00 LT. These results also suggest that both H$_2$O$_2$ and MHP may be affected by photochemistry and an increase of the atmospheric boundary layer depth during the day. Unlike NO, ROOH did not show an increase in the second part of the day but the levels remained low, consistent with an increased uptake by the snowpack when the temperature decreased in

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**Fig. 6.** Snow concentrations of (a) NO$_3^-$, (b) NO$_2^-$ and (c) H$_2$O$_2$ measured in the 30 cm-depth snow pits dug on 18 December 2008 (red squares), 28 December 2008 (blue triangles) and 04 January 2009 (green crosses). The shaded area in (b) represents the NO$_2^-$ LOD.
the evening (Hutterli, 2003, and references therein). The temperature record at the WAIS Divide camp was, however, too intermittent to further investigate the influence of temperature on atmospheric H$_2$O$_2$. A source from the snowpack is not considered significant for MHP since it has a solubility only 0.1% that of H$_2$O$_2$ (Lind and Kok, 1994) and because no MHP has been detected in snow and ice above the current LOD (Frey et al., 2005, 2009a). The change of MHP may depend on the photochemistry of its precursors such as CH$_4$, NMHCs and water vapor, which may decrease with solar elevation angle. Higher H$_2$O$_2$ and MHP levels at low wind speeds (less than 4 m s$^{-1}$) (data not shown) suggest that ROOH are produced locally, coherent with a local photochemical production of MHP and a physical snow-air exchange controlling H$_2$O$_2$ (Frey et al., 2009a). They are then diluted through turbulent transport after production.

NO$_3^-$ values are closer to those measured at coastal sites such as Halley or Neumayer stations (Mulvaney et al., 1998), than to the values observed at higher-altitude sites such as South Pole and Concordia stations (Dibb et al., 2004; France et al., 2011). Concentrations of skin-layer NO$_3^-$ present short-term and local spatial variations as Jarvis et al. (2009) and Wolff et al. (2008) also observed at Summit and Halley. These variations cannot be explained with a scenario of fresh snowfall since only one snow precipitation occurred during the campaign (Fig. 5). Part of these variations of NO$_3^-$ in the skin layer as H$_2$O$_2$ may be caused by processes such as nighttime deposition of NO$_3^-$, in the form of nitric oxide (HNO$_3$) and H$_2$O$_2$ or HNO$_3$ and H$_2$O$_2$ gas exchange. Other part can be attributed to some events of snow removed by the wind since this snow removal would also impact on chemical species other than NO$_3^-$ (such as H$_2$O$_2$), which presents similar variations to NO$_3^-$ on 26–27 December 2008 and on 1–2 January 2009. However, these events are too episodic to explain the overall variations of skin-layer NO$_3^-$.

It is suggested that part of NO$_3^-$ in the top snowpack undergoes some possible post-depositional processes such as photolysis. The contents of NO$_3^-$ and H$_2$O$_2$ in fresh snowfall are in the range of skin-layer concentrations, while previous observations showed that fresh snowfall contains higher concentrations of these species than does aged snow (e.g., Mulvaney et al., 1998). In addition to post-depositional equilibrium with the atmosphere, it is possible that either the snow may have degassed during its collection or prior to analysis. Anastasio and Robles (2007) showed that NO$_3^-$ and H$_2$O$_2$ contribute to half the light absorption in polar snow for wavelengths of 280 nm and above. Since the snowflakes from this diamond-dust-like precipitation stayed exposed to the sun under a clear sky until their collection at the end of the event – over 5 h – it is possible that photochemical loss of NO$_3^-$ and H$_2$O$_2$ may have occurred during the collection process.

The NO$_3^-$ and H$_2$O$_2$ profiles measured in the 30 cm snow pits show a summer peak, as observed in earlier studies (Kreutz et al., 1999, and references therein). The increase over the campaign of H$_2$O$_2$ concentrations in the top 15 cm of snow (Fig. 6) may point to a significant deposition of H$_2$O$_2$ during summer, and may also reflect changes in the overlying atmospheric H$_2$O$_2$ levels, as discussed in Hutterli (2003). NO$_3^-$ in the skin layer does not show a constant increase with time, unlike H$_2$O$_2$. However, a 70% decrease of NO$_3^-$ concentrations with increasing depth in the snow is observed, unlike reported snow-pit measurements from polar sites with similar surface temperature and snow accumulation rate, such as Summit, Greenland (e.g., Burkhart et al., 2004). These NO$_3^-$ profiles are similar to those observed at Dome C (e.g., Frey et al., 2009b; France et al., 2011). Based on the interpretation of these authors, this sharp decrease may therefore indicate that NO$_3^-$ in the top snowpack can be significantly reduced by photolysis. A similar photochemical depletion can be suggested to explain the slight decrease of NO$_3^-$ concentration observed in the 30 cm deep profiles over the campaign.

### 4.1.2 Post-depositional loss of NO$_3^-$ in surface snowpack

WAIS Divide and Summit, Greenland, have similar latitudes, surface temperature and snow accumulation rate (Table 2 and Fig. 7a). Concentrations of NO$_3^-$ in the skin layer are also in the same range for both sites, between 24–206 ppbw for WAIS Divide and 61–207 ppbw for Summit (Fig. 8a, b).

However, the median NO$_3^-$ concentrations in the top 1 m of snow at WAIS Divide is lower than at Summit, 38 vs. 138 ppbw. At very dry and cold sites on the East Antarctic Plateau such as Dome C, mean NO$_3^-$ concentrations are as low as 10–20 ppbw if one excludes the surface layer (Fig. 8c). Note that at Dome C most NO$_3^-$ is concentrated in the top layer with levels 1–2 orders of magnitude larger than at depth (Frey et al., 2009b; Erbland et al., 2013). Comparison of concentrations at the surface and at depth allows estimating net preservation of NO$_3^-$ in snow, which is about 30% at WAIS Divide, 75–93% at Summit (Burkhart et al., 2004; Dibb et al., 2007) and less than 10% at Dome C (Rothlisberger et al., 2002; Frey et al., 2009b).

Differences in average NO$_3^-$ concentrations in the skin layer and preserved at depth between WAIS Divide and Summit can be attributed to (a) a lower background of atmospheric reactive nitrogen in Antarcrica compared to the Arctic due to a larger distance from anthropogenic pollution sources, and (b) differences in processes occurring during and after deposition.

Regarding (b), the relative contributions of the processes involved may vary significantly between different locations, i.e., both physical (evaporation and adsorption of HNO$_3$) and photochemical (photolysis of NO$_3^-$) processes have been put forward to explain NO$_3^-$ net preservation (sum of gain and loss) in polar snow (Rothlisberger et al., 2002; Frey et al., 2009b). Both processes will be sensitive to accumulation rate and timing. For example, little snowfall in the sunlit season

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[S. Masclen et al.: WAIS snowpack emission and air mass transport source](http://www.atmos-chem-phys.net/13/8857/2013/)
Table 2. Concentrations of major ions (in ppbw) measured in surface snow at WAIS Divide and Summit, Greenland, and the respective estimated pH and alkalinity.

<table>
<thead>
<tr>
<th>Sampling year</th>
<th>Na⁺</th>
<th>NH₄⁺</th>
<th>K⁺</th>
<th>Mg²⁺</th>
<th>Ca²⁺</th>
<th>Cl⁻</th>
<th>NO₃⁻</th>
<th>SO₄²⁻</th>
<th>pH</th>
<th>Alk.</th>
</tr>
</thead>
<tbody>
<tr>
<td>WAIS Divide (T.air = −28.5°C; Acc = 20 cm weq yr⁻¹)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2000³</td>
<td>13</td>
<td>–</td>
<td>16</td>
<td>1</td>
<td>24</td>
<td>24</td>
<td>24</td>
<td>6.1</td>
<td>0.37</td>
<td></td>
</tr>
<tr>
<td>2006⁴</td>
<td>6</td>
<td>4</td>
<td>2</td>
<td>1</td>
<td>33</td>
<td>61</td>
<td>43</td>
<td>5.6</td>
<td>0.34</td>
<td></td>
</tr>
<tr>
<td>2006⁵</td>
<td>4</td>
<td>2</td>
<td>0.4</td>
<td>1</td>
<td>2</td>
<td>31</td>
<td>62</td>
<td>5.6</td>
<td>0.27</td>
<td></td>
</tr>
<tr>
<td>2008⁶</td>
<td>31</td>
<td>–</td>
<td>2</td>
<td>5</td>
<td>5</td>
<td>54</td>
<td>67</td>
<td>5.8</td>
<td>0.29</td>
<td></td>
</tr>
<tr>
<td>Average</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>5.8</td>
<td>0.32</td>
<td></td>
</tr>
<tr>
<td>Summit (T.air = −29.5°C; Acc = 24 cm weq yr⁻¹)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1987⁷</td>
<td>2</td>
<td>6</td>
<td>1</td>
<td>1</td>
<td>15</td>
<td>8</td>
<td>40</td>
<td>5.6</td>
<td>1.2</td>
<td></td>
</tr>
<tr>
<td>2003–2009⁸</td>
<td>8</td>
<td>21</td>
<td>10</td>
<td>2</td>
<td>18</td>
<td>42</td>
<td>234</td>
<td>107</td>
<td>5.2</td>
<td>2.4</td>
</tr>
<tr>
<td>2007⁹</td>
<td>4</td>
<td>10</td>
<td>3</td>
<td>2</td>
<td>19</td>
<td>23</td>
<td>195</td>
<td>72</td>
<td>5.3</td>
<td>1.6</td>
</tr>
<tr>
<td>2008⁸</td>
<td>3</td>
<td>5</td>
<td>3</td>
<td>1</td>
<td>6</td>
<td>19</td>
<td>239</td>
<td>54</td>
<td>5.3</td>
<td>0.65</td>
</tr>
<tr>
<td>Average</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>5.3</td>
<td>1.5</td>
<td></td>
</tr>
</tbody>
</table>

³ Based on [H⁺] = ([SO₄²⁻] × 0.12 × [Na⁺]) + [NO₃⁻] + ([Cl⁻] × 1.17 × [Na⁺]) (Legrand and Delmas, 1988).
⁴ In µeq L⁻¹, derived from the ionic balance and attributing the missing part of anions to dissolved carbonate.
⁵ Site ITASE 00-1, top 2 cm of surface snow; unpublished data (Mayewski and Dixon, 2005).
⁶ Top 3 cm of surface snow with the first cm removed; unpublished data from http://www.waisdivide.unh.edu.
⁷ Top 3 cm of surface snow; unpublished data from Kreutz and Koffman, 2011.
⁸ Top 3 cm of surface snow (Mayewski et al., 1990).
⁹ Top 1 cm of surface snow; unpublished data from http://niflheim.nilu.no/geosummit/.
¹⁰ Top 0.5 cm of surface snow (range 0.1–6 cm, Dibb et al., 2010).

Fig. 7. Updated Figs. 2, 3 and 4 from Rothlisberger et al. (2002) with additional measurements from WAIS Divide, Summit, and Dome C (median and range). Data reported from Greenland are NO₃⁻ concentrations in snow deposited either before 1940 (squares) or after 1970 (triangles). The NO₃⁻ levels observed in the snow deposited after 1970 over Greenland are influenced by anthropogenic emissions that do not reach the Antarctic continent, whereas the difference between the pre-1940s records and the Antarctic data is likely due to a difference in alkalinity.

implies longer exposure of surface snow to UV radiation, leading to significant loss of NO₃⁻ at some sites from photolysis (Frey et al., 2009b; Erbland et al., 2013).

In general, NO₃⁻ concentration preserved in polar snow scale with mean annual temperature and accumulation rate (Rothlisberger et al., 2002, Fig. 7). Comparison shows that NO₃⁻ concentrations at WAIS Divide are not unusual at all since they do fall into the range expected in Antarctica (Fig. 7b, c). They are, however, still lower than those in pre-1940 snow at Greenlandic sites of similar accumulation rate and temperature. The slightly lower accumulation and thus slower burial rate of annual snow deposition at WAIS Divide vs. Summit may contribute to this difference in NO₃⁻ preservation.

But, as suggested by the recent literature, variables linked to (post)depositional processes other than accumulation rate may contribute as well to the observed difference and include micro-physical properties and chemical heterogeneity in the snow matrix. Regarding the latter, observations of atmospheric and snow NO₃⁻ in coastal Antarctica showed that deposition of NO₃⁻ spikes is linked to sea salt aerosol, i.e., by conversion of gas-phase NO₃⁻ to NO₃⁻ aerosol and enhanced trapping of gas-phase NO₃⁻ on salty surfaces (Wolff et al., 2008). This enhanced deposition efficiency will be less important further inland. However, higher NO₃⁻ concentrations were also associated with higher dust content (i.e., calcium (Ca²⁺)), suggesting reduced post-depositional loss of NO₃⁻ by photolysis or evaporation (Rothlisberger et al., 2000).
Fig. 8. The 1 m NO$_3^-$ profiles from snowpits reported from WAIS Divide, Summit, and Dome C (Frey et al., 2013; France et al., 2011; Traversi et al., 2009; Burkhart et al., 2004; Hastings, 2004; Rothlisberger et al., 2000; Mayewski et al., 1990, Jarvis (unpublished data)).

comparison with Summit, the lower dust content in WAIS Divide snow, ∼20% as alkaline and ∼30% as much Ca$^{2+}$ (Table 2), may reduce the preservation of NO$_3^-$ in snow and enhance the NO$_x$ emission flux in summer. Systematic differences of NO$_x$ emissions from surface snow could indeed further support increased loss rates and thus less preservation of NO$_3^-$ in surface snow at WAIS Divide. At Summit, Honrath et al. (2002) measured 24 h average NO$_x$ fluxes of $2.5 \times 10^8$ molecule cm$^{-2}$ s$^{-1}$, significantly less than what we calculated for WAIS Divide (see Sec. 4.1.5). However, based on a similar approach to our study, Zatko et al. (2013) recently modeled an NO$_x$ flux at Summit 5–10 times that reported by Honrath et al. (2002). These results point to significant NO$_x$ emissions from NO$_3^-$ in the snowpack at both sites. Our calculations suggest high loss rates, higher than at Summit (see Sect. 4.1.5), but uncertainties in available measurements and model estimates prevent concluding that emission rates are significantly higher at WAIS Divide than at Summit.

Further, the stable isotopic composition of NO$_3^-$ in snow can provide information on the amount of post-depositional NO$_3^-$ loss (e.g., Frey et al., 2009b; Erbland et al., 2013). At Summit, the N and O isotopes of NO$_3^-$ showed that most of the NO$_x$ emitted from the surface snow is recycled back to the snow as NO$_3^-$, explaining high NO$_3^-$ preservation (Hastings, 2004). It can be speculated that if the NO$_3^-$ loss rate was similar at WAIS Divide, then some of the emitted gas phase species may be lost through lateral export, implying a lower recycling efficiency and therefore lower NO$_3^-$ preservation. Thus, NO$_3^-$ post-depositional loss is a nonlinear combination of both accumulation and temperature, but other parameters need also to be considered including snow chemistry to get a more complete process understanding.

4.1.3 Steady-state estimation of atmospheric NO$_2$

Considering the NO–NO$_2$–O$_3$ system, it is reasonable to assume a photo-stationary steady state between NO and NO$_2$ at 1 m above the snowpack (Frey et al., 2013) to infer the potential atmospheric NO$_2$ concentrations from reactions:

\[
\text{NO} + \text{O}_3 \rightarrow \text{NO}_2 + \text{O}_2 \quad \text{(R4)}
\]

The conversion of NO to NO$_2$ described in Reaction (R4) is also achieved through different channels (Reactions R1, R5–R6) with the presence of oxidants such as HO$_x$, peroxy (RO$_2$) or halogen (XO = ClO, BrO, IO) radicals:

\]

NO$_2$ mixing ratios can therefore be estimated from the extended Leighton ratio as derived in Ridley et al. (2000):

\[
[\text{NO}_2] = \frac{[\text{NO}]}{[\text{NO}]} \frac{k_{R4}[\text{O}_3] + k_{R1}[\text{XO}]}{j_{R2}}
\]

(R1)
with $\text{OX}$ the total radical concentration ($[\text{OX}] = [\text{HO}_2] + [\text{RO}_2] + 2[\text{OX}]$) as $\text{OX}$ represents the halogen radicals (Ridley et al., 2000). The photolytic rate constant $k_R$ was calculated from the NCAR/ACD radiative-transfer model TUV version 5.0 (Lee-Taylor and Madronich, 2002) with $O_3$ columns measured by total ozone mapping spectrometer (TOMS) (http://ozoneaq.gsfc.nasa.gov/ozone_overhead_current_v8.md) and assuming clear-sky conditions. $k_{R4}$ and $k_{R1}$ were estimated through the temperature-dependent expressions from Sander et al. (2006). An NO$_2$ concentration of 5 pptv was computed based on the averages of NO and $O_3$ observed at WAIS Divide, and on $4.9\times10^7$ molecule cm$^{-3}$ HO$_2$ + RO$_2$ mixing ratios derived from the $1.3\times10^6$ molecule cm$^{-3}$ OH radical concentrations computed for Byrd (Frey et al., 2005). Because the mixing ratios of HO$_x$ for WAIS Divide are estimates, we calculated the steady-state NO$_2$ concentrations with reported HO$_x$ levels from Halley ($2.3\times10^7$ molecule cm$^{-3}$) (Bloss et al., 2007) and South Pole ($8.3\times10^7$ molecule cm$^{-3}$) (Eisele and Davis, 2008), resulting in NO$_2$ concentrations of 5.1 and 6.1 pptv, respectively. These results show that this steady state relationship depends essentially on NO and $O_3$ mixing ratios.

Note that halogen radicals were not considered in these calculations, and no records have yet been reported above WAIS. Coastal sea ice is the main source of active halogen (Saiz-Lopez and von Glasow, 2012, and references therein) and measurements from Antarctic coastal sites showed a strong spatial variability in concentrations of halogens and their potential impact on boundary layer photochemistry. Summertime measurements of IO and BrO ranged from 0.7 to 5.5 pptv at Halley (Saiz-Lopez et al., 2007), while mixing ratios at Dumont D’Urville were observed between 0.04 and 0.15 pptv for IO and below or equal to 2 pptv for BrO (Grilli et al., 2013). The back-trajectory analysis described in Sect. 4.2.1 shows that marine intrusions over WAIS are frequent (58% of all back trajectories) and have therefore the potential to export halogen compounds over the ice sheet. However, from measurements by the satellite spectrometer SCIAMACHY (Schönhardt et al., 2012), monthly mean IO and BrO vertical column amounts averaged over 2004 to 2009 revealed insignificant concentrations of both species over WAIS Divide between December and January. Thus, it is reasonable to assume that little or insignificant halogen chemistry is occurring over WAIS Divide.

The NO: NO$_2$ ratio of 3.8 is significantly higher than the ratios of 1.3–2 and 1.5 observed respectively at Halley (Bauguitte et al., 2012, and references therein) and Dome C (Frey et al., 2013). These studies showed evidence of large discrepancies between observed and steady-state estimated ratios that are attributed to halogen chemistry for the coastal station and to the presence of HO$_2$ + RO$_2$ on the plateau. Therefore, the estimated atmospheric NO$_2$ from the steady-state assumption has to be considered as a lower limit of true value.

### 4.1.4 Potential NO$_x$ lifetime

The NO$_x$ lifetime was calculated to investigate the factors controlling its atmospheric concentration. Assuming that halogen chemistry is not significant at WAIS Divide, Reactions (R7)–(R8) are the main sink of NO$_2$ during solar day:

$$\text{NO}_2 + \text{OH} + \text{M} \rightarrow \text{HNO}_3 + \text{M} \quad (R7)$$

$$\text{NO}_2 + \text{HO}_2 + \text{M} \rightarrow \text{HNO}_4 + \text{M} \quad (R8)$$

Based on these reactions, the lifetime of NO$_2$ ($\tau_{\text{NO}_2}$) was estimated with $[\text{OH}]=1.3\times10^6$ molecule cm$^{-3}$ and $[\text{HO}_2]=4.9\times10^7$ molecule cm$^{-3}$ (Sect. 4.1.3). The lifetime of NO$_x$ ($\tau_{\text{NO}_x}$) was then deduced through Eq. (2) (Seinfeld and Pandis, 1998):

$$\tau_{\text{NO}_x} = \tau_{\text{NO}_2} \times \left(1 + \frac{[\text{NO}]}{[\text{NO}_2]} \right).$$

The resulting lifetime of NO$_x$ at WAIS Divide of 15 h represents an upper bound, given that the NO: NO$_2$ ratio of 3.8 is an upper limit. This value is less than the 24 h estimated by Jones et al. (2000) from a snow block at Neumayer, but it is longer than the 6.4 h estimated at Halley (Bauguitte et al., 2012) and 8 h at South Pole (Davis et al., 2004). Bauguitte et al. (2012) showed that NO$_2$ lifetime at Halley is mainly controlled by halogen oxidation processes, while low temperatures at South Pole prevent thermal decomposition of pernitric acid and therefore enhance NO$_2$ removal through its oxidation with HO$_2$ (Slusher et al., 2002). Thus, our potential NO$_x$ lifetime seems plausible since we expect to observe none of these conditions at WAIS Divide. NO$_x$ lifetimes computed with reported HO$_x$ concentrations from South Pole and Halley (Sect. 4.1.3) are respectively 9 and 33 h and represent the potential lower and upper bounds for the NO$_x$ lifetime in the inner WAIS.

Considering a photochemical lifetime of other NO$_x$ precursors, such as HNO$_3$ and HNO$_2$ (HNO$_3$ compared to dry deposition), of less than a day above the East Antarctic Plateau (Davis et al., 2008; Slusher et al., 2002) and an air mass transport between WAIS Divide and the Plateau longer than 33 h (Sect. 4.2), outflows from this region are not expected to be a major NO$_x$ source for the boundary layer at WAIS.

### 4.1.5 Calculation of upper-limit NO$_x$ emission from NO$_3^-$ and NO$_2^-$ photolysis

The simplified reaction scheme (R9) to (R12) summarizes the currently known NO$_3^-$ photochemistry in near-surface snow:

$$\text{NO}_3^- + h\nu \rightarrow \text{NO}_2 + \text{O}^- \quad (R9)$$
\[ \text{NO}_3^- + h\nu \rightarrow \text{NO}_2^- + \text{O}(^3\text{P}) \]  
\[ \text{(R10)} \]

\[ \text{NO}_2^- + h\nu \rightarrow \text{NO} + \text{O}^- \]  
\[ \text{(R11)} \]

\[ \text{NO}_2^- + \text{OH} \rightarrow \text{NO}_2 + \text{OH}^- \]  
\[ \text{(R12)} \]

While \( \text{NO}_3^- \) photolysis is the major source of \( \text{NO}_2 \) (Reaction R9) (Grannas et al., 2007), recent studies also pointed at the potential contribution of the photolysis of \( \text{NO}_3^- \) to produce nitric oxide (Reaction R11) (France et al., 2012, and references therein). Therefore, calculations of the \( \text{NO}_x \) emission flux in snow were based on Reactions (R9) and (R11) for the respective wavelengths 280–360 nm and 280–400 nm (Chu and Anastasio, 2003, 2007, and references therein). The first-order rate constants for the photolysis of \( \text{NO}_3^- \) at the snowpack surface \( j_{\text{NO}_3^-} \) were calculated as defined in Seinfeld and Pandis (1998):

\[ j_{\text{NO}_3^-} = \frac{\lambda I}{\lambda I} \int \sigma_{\text{NO}_3^-}(\lambda, T) \phi_{\text{NO}_3^-}(\lambda, T) I(\lambda, \theta, z_0) \text{d}\lambda. \]  
\[ \text{(3)} \]

Based on the results of Chu and Anastasio (2003), the spectral UV absorptivity \( \sigma_{\text{NO}_3^-}(\lambda, T) \) was derived between 280 to 360 nm from the measured molar absorption coefficients of aqueous \( \text{NO}_3^- \) at 278 K and a quantum yield \( \phi_{\text{NO}_3^-}(\lambda, T) \) of \( 2.79 \times 10^{-3} \) was estimated for \( T = 259 \text{K} \), the average temperature over the three sampling days. The first-order rate constants for the photolysis of \( \text{NO}_2^- \) at the snow surface \( j_{\text{NO}_2^-} \) were calculated with the same approach. \( \sigma_{\text{NO}_2^-}(\lambda, T) \) was calculated between 280 to 400 nm from reported molar absorptivities for aqueous \( \text{NO}_2^- \) at 274 K and a range of \( \phi_{\text{NO}_2^-}(\lambda, T) \) of \( 3.4 \times 10^{-2} - 0.9 \times 10^{-2} \) for \( \lambda \) between 280 and 400 nm was also determined for \( T = 259 \text{K} \) (Chu and Anastasio, 2007). The actinic fluxes \( I(\lambda, \theta, z_0) \) were computed with the NCAR/ACD radiative transfer model TUV version 5.0 under clear-sky conditions for a 23° solar elevation angle (SEA = 90° − \( \theta \), with \( \theta \) the solar zenith angle) averaged over the three sampling days.

The transmission of light in snow that drives the photochemistry in the top snowpack is controlled by optical processes: scattering and absorption (Dominé et al., 2008; Grannas et al., 2007, and references therein). These processes are summarized in the parameter \( e \)-folding depth (EFD), which in turn depends on snow physical properties – grain size, density, liquid water content – and the concentration of light-absorbing impurities such as black carbon. Studies at Dome C (France et al., 2011) and Barrow, Alaska (Reay et al., 2012), showed that absorption in snow layers, and therefore EFD, is inversely proportional to the content of black carbon in snow. Reported concentrations of black carbon in snow at WAIS Divide of 0.08 ± 0.4 ng g\(^{-1}\) (average ±1σ) are in the range of those measured at South Pole, 0.2–0.3 ng g\(^{-1}\) (Bisiaux et al., 2012, and references therein). Values of \( j_{\text{NO}_3^-} \) and \( j_{\text{NO}_2^-} \) were then scaled as a function of depth (\( z \)) using an EFD of 30 cm (Table 3), based on the similar value calculated for South Pole by Zatko et al. (2013):

\[ j_{\text{NO}_3^-} = j_{\text{NO}_3^-}(30 \text{ cm}) \exp \left( -\frac{z}{\text{EFD}} \right). \]  
\[ \text{(4)} \]

The depth-integrated emission flux of \( \text{NO}_x \) was calculated for 18 and 28 December 2008 and 4 January 2009 with

\[ F_{\text{NO}_2} = \frac{\int_{z=0}^{30 \text{ cm}} [\text{NO}_3^-](z) j_{\text{NO}_3^-}(z) \text{d}z}{30 \text{ cm}} \]  
\[ \text{(5)} \]

\[ F_{\text{NO}} = \frac{\int_{z=0}^{30 \text{ cm}} [\text{NO}_2^-](z) j_{\text{NO}_2^-}(z) \text{d}z}{30 \text{ cm}} \]  
\[ \text{(6)} \]

with respectively \( [\text{NO}_3^-] \) and \( [\text{NO}_2^-] \) in molecule cm\(^{-3}\) measured at each depth \( z \) (Fig. 6).

In order to convert flux \( F_{\text{NO}_x} \) (molecule cm\(^{-2}\) s\(^{-1}\)) into a volumetric production rate \( P_{\text{NO}_x} \) (ppv h\(^{-1}\)), \( F_{\text{NO}_x} \) was multiplied by the height of the boundary layer estimated from prior balloon soundings above the West Antarctic Ice Sheet in summer, with an average height of 250 m (range 13–354 m) between morning and late afternoon (Frey et al., 2005). Note that atmospheric boundary layer height as low as 13 m may be infrequent at WAIS Divide. Therefore, estimation of the potential \( \text{NO}_x \) production from snow for an atmospheric boundary layer height of 13 m will be an upper bound.

Recent studies (Chu and Anastasio, 2007; Boxe and Saiz-Lopez, 2008, and references therein) suggest that only \( \text{NO}_3^- \) and \( \text{NO}_2^- \) in quasi-liquid layers are available for photolysis as photochemistry of these species does not occur in the bulk ice. Because the distribution of \( \text{NO}_3^- \) and \( \text{NO}_2^- \) between the quasi-liquid layers and the lattice ice is unknown, these calculations assumed that all \( \text{NO}_3^- \) and \( \text{NO}_2^- \) can be photolyzed. It is also assumed that the totality of the \( \text{NO}_x \) produced escapes from the snowpack and is released into the overlying atmosphere without considering any chemical loss. These fluxes and production rates are therefore an upper limit of the potential \( \text{NO}_x \) source from the snow.

### 4.1.6 Potential \( \text{NO}_x \) production from top snowpack

The resulting potential daily \( \text{NO}_2 \) emission fluxes from \( \text{NO}_3^- \) photolysis in the surface snowpack were \( 8.6 \times 10^6 \) molecule cm\(^{-2}\) s\(^{-1}\) (Table 3). This value is in agreement with previous reported \( \text{NO}_2 \) emission fluxes on the Antarctic continent (Table 4). For \( \text{NO}_x \), the estimated value is \( 33.9 \times 10^8 \) molecule cm\(^{-2}\) s\(^{-1}\). Maximum daily \( \text{NO}_x \) emission fluxes from snow are therefore \( F_{\text{NO}_x} \) of \( 42.5 \times 10^8 \) molecule cm\(^{-2}\) s\(^{-1}\), with 20% and 80% from...
Table 3. Potential depth integrated NOx emission fluxes and equivalent NOx production rates from NO3 and NOO3 photolysis for an EFD of 30 cm, a solar elevation angle of 23° and three different atmospheric boundary layer heights (ABL) from previous measurements (Frey et al., 2005).

<table>
<thead>
<tr>
<th>Compound</th>
<th>Potential emission fluxes $(10^8$ molecule cm$^{-2}$ s$^{-1}$)</th>
<th>Potential production rates (pptv h$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NO2</td>
<td>8.6</td>
<td>114</td>
</tr>
<tr>
<td>NO</td>
<td>33.9</td>
<td>452</td>
</tr>
</tbody>
</table>

NO$^-$ and NOO$^-$ photolysis, respectively. This contribution is similar to that found by France et al. (2012) at Barrow.

The resulting local NOX production is 30 pptv h$^{-1}$ for a mean atmospheric boundary layer height of 250 m (Table 3). Average production rates of 21 and 566 pptv h$^{-1}$ were calculated for boundary layer depths of 354 and 13 m, respectively. Comparison between the potential NOX production rate and the mixing ratios of NOX estimated from the extended Leighton mechanism with measured NO (Sect. 4.1.3) indicates that the snow source contributes significantly to the NOX budget at WAIS Divide. However, values of $P_{NOx}$ suggest that the estimated production rates are an upper limit since the observed levels of NO (19 pptv) and the estimated NO2 mixing ratios (5 pptv) are inconsistent with an NOX production rate of 566 pptv h$^{-1}$ for atmospheric boundary layer height of 13 m.

These calculated NOX emissions are higher than reported values from other Antarctic sites, where modeled and measured fluxes reached between $2.4 \times 10^8$ and $22 \times 10^8$ molecule cm$^{-2}$ s$^{-1}$ (Table 4). In a recent study, Zatko et al. (2013) suggested that local contamination from nearby stations increases the content of impurities in snow, therefore reducing the EFD of the actinic flux in the top snowpack, resulting in a possible underestimation of the NOX emissions modeled or measured nearby stations by a factor of 1.4–2.4. Thus, high NOX emissions could be expected in our study since the sampling site was 5 km upwind of the main WAIS camp emissions (Sect. 2). However, the results from Zatko et al. (2013) can be discussed since most of the measurements or estimations of NOX nearby Antarctic stations were from sites located in clean air sector, upwind of the stations (Table 4). With said precautions, local contamination from nearby stations should not reach these sites and impact on the measured or calculated snowpack emissions.

The above estimates do have uncertainties: a) daily surface-snow samples revealed that the local-scale spatial variability of NO$^-$ in the near-surface snowpack is significant, and b) our calculations assume that all of the NO$^-$ and NOO$^-$ in snow is available for photolysis followed by emission into the atmosphere of all the photo-produced NO and NO2. Concerning b), Anastasio and Chu (2009) suggested that 30% of the NO2 produced from NOO$^-$ photolysis may be converted to NO$^-$ within the snow matrix before being released into the atmosphere. Also, the decrease of 5% per day of NO$^-$ in surface snow (Sect. 3.2) indicates that only a small amount of NOO$^-$ may actually undergo photolysis to produce NO. Nevertheless, these results show the key role of the boundary-layer depth in determining the contribution of snowpack emissions to the overlying photochemistry. Our estimates imply that both the variability of boundary layer height with an observed range in summer of 13 to 354 m and local NOX emissions from snow impact the atmospheric NOX mixing ratios.

### 4.1.7 Local production of O3

The oxidation of methane is the main in situ chemical source of tropospheric O3. Thus, Reaction (R1) leads to production of O3 while Reaction (R13) leads to its destruction (Seinfeld and Pandis, 1998):

$$
\text{HO}_2 + \text{O}_3 \rightarrow \text{OH} + 2\text{O}_2.
$$

(R13)

The ratio between these two rates indicates whether WAIS Divide is an O3 production or destruction site:

$$
\frac{\text{R13}}{\text{R1}} = \frac{k_{\text{R13}} [\text{O}_3]}{k_{\text{R1}} [\text{NO}]},
$$

(7)

This ratio was calculated with $k_{\text{R1}} = 9.49 \times 10^{-12}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ and $k_{\text{R13}} = 1.62 \times 10^{-15}$ cm$^3$ molecule$^{-1}$ s$^{-1}$, derived from Sander et al. (2006), and an average temperature of 255.6 K measured during the campaign. Note that this estimation from the photo stationary state relation accounts for the NOX emitted from the snowpack. In the case of Summit, Thomas et al. (2012) showed that NOX emitted from the snowpack can increase the boundary layer O3 by an additional 2–3 ppbv. Frey et al. (2013) show that under calm conditions typical for Dome C, East Antarctica, NO and NO2 emitted by surface snow reach steady-state at 1 m. Thus, it is expected that our measured mixing ratios of O3 at 1 m above surface snow already reflect the contribution from the NOX emissions. Also, it is assumed that halogen chemistry does not occur at WAIS Divide, otherwise the reaction of BrO with O3 and NOX in the snowpack would reduce the local O3 production (Thomas et al., 2012).

The ratio (R13)/(R1) equals 0.216 and confirms that WAIS Divide is a surface ozone production site (Sect. 4.1.1).
Table 4. Potential emission flux of NO$_x$ from NO$_3^−$ and NO$_2^−$ photolysis in snow at WAIS Divide compared to other Antarctic sites.

<table>
<thead>
<tr>
<th>Site &amp; reference</th>
<th>$F_{NO_x} \pm 1\sigma$ (10$^8$ molecule cm$^{-2}$ s$^{-1}$)</th>
<th>Period</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>WAIS Divide</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>This study</td>
<td>$F_{NO_x} = 42.5^a$</td>
<td>12/08–01/09</td>
<td>Daily average, 23° SEA</td>
</tr>
<tr>
<td>Neumayer Halley V:</td>
<td>$3(+0.3/−0.9)^b$</td>
<td>05–07/02/99</td>
<td>Noon maxima</td>
</tr>
<tr>
<td>Jones et al. (2001)</td>
<td>2.42$^a$</td>
<td>18/01/05</td>
<td>Daily average</td>
</tr>
<tr>
<td>Bauguitte et al. (2012)</td>
<td>7.3$^b$</td>
<td>02/02/05</td>
<td>Noon maxima (07:05–15:05 UTC) average</td>
</tr>
<tr>
<td>South Pole</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oncley et al. (2004)</td>
<td>3.2–4.2$^e$</td>
<td>11–12/03</td>
<td>Noon maxima, remote</td>
</tr>
<tr>
<td>Wang et al. (2008)</td>
<td>3.3–9.8$^f$</td>
<td>January</td>
<td>Noon maxima, near station</td>
</tr>
<tr>
<td>Zatko et al. (2013)</td>
<td>7.7–22$^f$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dome C</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Frey et al. (2013)</td>
<td>6.9 ± 7.2$^b$</td>
<td>12/09–01/10</td>
<td>Noon maxima, near station</td>
</tr>
<tr>
<td>Zatko et al. (2013)</td>
<td>3.2–12$^f$</td>
<td>January</td>
<td>Noon maxima, remote</td>
</tr>
<tr>
<td></td>
<td></td>
<td>4.4–17$^f$</td>
<td></td>
</tr>
<tr>
<td>WAIS Divide</td>
<td>$F_{NO_2} = 8.6^a$</td>
<td>12/08–01/09</td>
<td>Daily average, 23° SEA</td>
</tr>
<tr>
<td></td>
<td>$F_{NO} = 33.9^a$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>South Pole</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oncley et al. (2004)</td>
<td>$F_{NO} = 2.6 ± 0.3^d$</td>
<td>26–30/11/00</td>
<td>Overall average</td>
</tr>
<tr>
<td>Dome C</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>France et al. (2011)</td>
<td>$F_{NO_2} = 2.4–3.8^g$</td>
<td>12/09–01/10</td>
<td>Values for 22° SEA</td>
</tr>
</tbody>
</table>

$^a$ Depth-integrated $F_{NO_x}$ with spectral irradiance from the TUV model.

$^b$ From measured gradients of NO$_x$ concentrations and turbulent diffusivity.

$^c$ From 1-D model of NO$_x$ concentrations based on production from NO$_3^−$ photolysis and chemical loss.

$^d$ Based on observed NO gradients and assuming photochemical steady-state.

$^e$ From 1-D chemistry-diffusion model based on trace gas measurements.

$^f$ Based on depth-dependent actinic flux profiles derived from snowpack radiative transfer model and snow-impurity measurements.

$^g$ Depth-integrated $F_{NO_x}$ with measured spectral irradiance.

From Eq. (7), a threshold value of 2 pptv for NO is found to trigger O$_3$ production at WAIS Divide, whereas this threshold is reached at 5 pptv in the remote continental troposphere of mid-latitude regions (Seinfeld and Pandis, 1998) and of high latitude sites, such as Dome C (derived from Chen et al., 2007). An atmospheric HO$_2$ concentration of 4.9×10$^7$ molecule cm$^{-3}$ leads to a potential O$_3$ production of 0.8 ppbv day$^{-1}$. This is about 5% of the observed O$_3$ mixing ratios and respectively 20% and 50% of the rates calculated above the East Antarctic Plateau. O$_3$ production rates are ∼4 ppbv day$^{-1}$ at South Pole (Chen et al., 2004, and references therein) and ∼1.5 ppbv day$^{-1}$ at Dome C (Legrand et al., 2009). Our result is consistent with these previous studies since WAIS Divide is a lower-altitude site with a deeper atmospheric boundary layer that is influenced by a diel cycle of UV irradiance.

It is apparent that local O$_3$ production is too small to account for the observed increases between 24–25 and 27–29 December of 0.3 ppbv h$^{-1}$. We therefore consider the impact of air mass transport, as detailed below.

4.2 Impacts of air mass transport

Analysis of wind direction and O$_3$ mixing ratios indicate that air masses from the ENE-SWS sector have typically the highest O$_3$ concentrations (Sect. 2, Fig. 4). O$_3$ can be considered a long-lived chemical species compared to NO$_x$ (Sect. 4.1.5) with a lifetime of about 22 days in the polar regions (Grannas et al., 2007). It is therefore expected that local O$_3$ is affected by transport, so the origin and transport of air at WAIS Divide were further investigated using 4-day back trajectories.

A total 168 trajectories were computed with the NOAA Hysplit/HYbrid Single-Particle Lagrangian Integrated Trajectory; see http://ready.arl.noaa.gov/hysplit4.html model (R. R. Draxler and G. D. Rolph, 2003) every 4 h using global meteorological data from the NCEP Global Data Assimilation System (GDAS) with a 3 h temporal resolution, a
1° × 1° (latitude × longitude) spatial resolution, an endpoint at the latitude and longitude of WAIS Divide and 10 m above ground level from 10 December 2008 to 8 January 2009. A comparison of the resulting 168 trajectories with another set generated from the NCEP-NCAR archived data shows similar results (Fig. 9), while Sinclair et al. (2013) showed that back trajectories from NCEP-NCAR and ECMWF Interim Re-Analysis (ERA-Interim) were comparable during austral summer. Therefore, the following discussion is based on trajectories produced from the GDAS archives. The back trajectories were combined to create daily maps (Fig. 10) and we distinguish three regions of air mass origin based on elevation, slope and previous O$_3$ and temperature measurements (Fig. 9).

We refer to the East Antarctic Plateau as the East Antarctic region above 2500 m a.m.s.l. with a slope of less than 1° since elevated O$_3$ has not been observed below 2500 m a.m.s.l. during the Antarctic summer (Frey et al., 2005). Surface O$_3$ is produced over the East Antarctic Plateau and possibly exported through air outflow (Sect. 4.2.2). The inner WAIS refers to the West Antarctic area above 1750 m a.m.s.l. with a slope inferior to 1°. An average temperature of about −25 °C was reported in the inner WAIS compared to −15 °C in the lower-elevated regions (Frey et al., 2005). Finally, based on a slope higher than 1° and on the previous observations, the areas below 2500 m a.m.s.l. in East Antarctica and 1750 m a.m.s.l. in West Antarctica are referred to as Antarctic coastal slopes.

Figure 9 shows that over the campaign, 58% of the air masses originated from the Antarctic coastal slopes with 2% possibly flowing from the distant King Haakon VII coast. An estimated 17% was transported from the inner WAIS and 25% of the air was advected from the East Antarctic Plateau. About 71% of the air reaching WAIS Divide flowed from East Antarctica and 29% from West Antarctica.

4.2.1 Air mass origins related to low O$_3$ levels

Except for outflows from the East Antarctic Plateau, all air mass origins are associated with low O$_3$, averaging 13 ± 3 ppbv (Fig. 11). This represents therefore 75% of the air mass reaching WAIS Divide, which includes air from the coastal slopes (Fig. 10a, b) and the inner WAIS (Fig. 10e). In coastal Antarctica, halogen-catalyzed chemistry has been found to prevent O$_3$ production in summer and even leads to the well-known episodic O$_3$ depletion events during spring (Jones et al., 2008, and references therein). The little vertical-column amounts of IO and BrO observed at WAIS Divide from satellite in December and January 2004–2009 (Schönhardt et al., 2012) suggest that contribution of halogens from intrusions of coastal air (58% of all back trajectories) to the WAIS Divide photochemistry is not significant. Thus, reduction of O$_3$ levels at WAIS Divide by halogen chemistry may not occur, as assumed in the previous sections.

![Fig. 9. Map of the different origins of air reaching WAIS Divide identified by the 4-day back-trajectory analysis and topography: East Antarctic Plateau (yellow), inner WAIS (red) and the Antarctic coastal slopes (green). Comparison between the results from analyses using the GDAS and NCAR meteorological archived data is shown.](image)

Air masses from the interior of WAIS (17% of all back trajectories, Fig. 10e) are also associated with low O$_3$ mixing ratios. Influence from local halogen chemistry or halogens exported from the Antarctic coasts is not consistent with observations of Schönhardt et al. (2012).

It is therefore suggested that the O$_3$ levels of the air masses from the coastal slopes and the interior of WAIS are low, so these airflows can reduce the O$_3$ levels of WAIS Divide when reaching the site. From these results and the wind-speed record (Fig. 11), it is also expected that O$_3$ production over WAIS is less than or equal to our estimated O$_3$ production rate for WAIS Divide.

Airflows from long-distance sources such as King Haakon VII sea coast (2% of all back trajectories, Fig. 10f) appear infrequently, with no obvious impact on local O$_3$. The O$_3$ signature of these air masses likely disappeared during their transport over more than 2500 km.

4.2.2 Air mass origins associated with elevated O$_3$ levels

Approximately 25% of all back trajectories point to airflows from the East Antarctic Plateau (Fig. 10c, d and Fig. 9), which are mainly related to the highest O$_3$ measured at WAIS Divide, with an average ±1σ of 19 ± 4 ppbv (Fig. 11).

This is consistent with the O$_3$ production above the East Antarctic Plateau (Helmig et al., 2008a, and references therein) that enhances the O$_3$ levels of WAIS through air transport, as previously reported by Legrand et al. (2009), who only observed high O$_3$ mixing ratios at Dumont...
D’Urville for air masses originating from the Antarctic Plateau. Such an oxidizing environment is confirmed by the O$_3$ observations at South Pole (S. J. Oltmans, http://ds.data.jma.go.jp/gmd/wdegg) over the same period that shows an average concentration of 31 ppbv, 17 ppbv more than the average observed at WAIS Divide (Fig. 11).

Further back-trajectory analysis reveals two conditions in which high O$_3$ concentrations are observed above WAIS Divide. First, elevated O$_3$ mixing ratios were only observed for lower atmosphere air coming from the East Antarctic Plateau (below 1500 m above ground level). The trajectories of these near-surface airflows are in good agreement with the katabatic streamlines described by Parish and Bromwich (2007). This suggests that air exported off the East Antarctic Plateau via katabatic outflows raises significantly the O$_3$ mixing ratios at WAIS Divide. The ozone depletion recorded between 25 and 27 December 2008 (Fig. 10d and Fig. 11) is attributed to an air mass coming from altitude as high as $\sim$3000 m above ground level.

Secondly, high O$_3$ was only observed at WAIS Divide when the transport time from the East Antarctic Plateau was less than 3 days (Fig. 10c and d). To investigate this condition, the total O$_3$ loss during transport from the Antarctic Plateau to WAIS Divide was estimated. The back trajectories indicate that O$_3$ maxima during 24–28 December at WAIS Divide are due to rich O$_3$ air originating from the South Pole region, as local records show similar but higher peaks about 2 days before (Fig. 11). From this delay, a loss rate of 0.25 ($\pm$0.04) ppbv h$^{-1}$ was calculated. This represents an O$_3$ loss rate of 5.5 ($\pm$0.9) ppbv day$^{-1}$ when averaged O$_3$ mixing ratios were 30 and 14 ppbv, respectively, at South Pole and WAIS Divide over the campaign. If these loss rates (from dilution, net destruction, etc.) are typical, then it follows that O$_3$ export off the Plateau has to be rapid (less than 3 days) in order to have an impact on WAIS Divide’s levels.

Observations of elevated O$_3$ levels are frequently related to vertical transport from the free troposphere/lower stratosphere at Summit, Greenland (Helmig et al., 2007b) and

![Figure 10](image_url)
Fig. 11. The 1 h averages of observed (a) wind speed at WAIS Divide, (b) O₃ at WAIS Divide, and (c) O₃ at South Pole (data available at: http://ds.data.jma.go.jp/gmd/wdcgg) for austral summer 2008–2009. Air mass origins are reported with the identical color coding used in Fig. 9: East Antarctic Plateau (yellow), inner WAIS (red) and Antarctic coastal slopes (green). Events of air transport less than 3 days between East Antarctic Plateau and WAIS Divide (τ_{air} < 3 days) and of high-altitude air origin (<2 km above ground level) are also reported.

other Arctic sites (Helmig et al., 2007a, and references therein). These stratospheric-intrusion events are promoted by boundary-layer instabilities, which are caused by significant diel radiation cycles and by the size of the Greenland ice sheet (Helmig et al., 2007b). All of the vertical O₃ profiles over the East Antarctic Plateau from tethered-balloon measurements (Helmig et al., 2008a; Johnson et al., 2008; Oltmans et al., 2008) and an airborne study (Slusher et al., 2010) present higher O₃ mixing ratios in the atmospheric boundary layer than in the free troposphere, indicating that no contribution of stratospheric O₃ occurs over the Antarctic Plateau. Enhancement of the O₃ levels at WAIS Divide from vertical transport is neither supported by the back-trajectory analysis nor by the vertical O₃ profiles measured at South Pole during this period (http://www.esrl.noaa.gov/gmd/dv/data/index.php), confirming the previous observations from the East Antarctic Plateau.

However, not all elevated O₃ mixing ratios at this site could be explained with the air origin. This is the case for the 5 h rise of O₃ mixing ratios that reached 24 ppbv on 12 December and occurred during the only snowfall observed during the campaign (Fig. 5). With low wind speeds (Fig. 11) and airflow from the inner WAIS observed during this event, it is possible that more O₃ accumulated in the boundary layer from local production. However, given the O₃ production rate we estimated previously (Sect. 4.1.7), it seems unlikely that only the local production of O₃ could cause this specific increase.

5 Conclusions

Being from a geographically intermediate site between the Antarctic coast and the East Antarctic Plateau, our results show that concentrations of atmospheric NO, O₃, H₂O₂ and skin-layer NO₃⁻ at WAIS Divide during summer 2008–2009 are similar to coastal levels.

Comparison of local potential NOₓ production from daily average depth-integrated emission fluxes of NOₓ and the NOₓ mixing ratios estimated from a steady-state assumption shows that NO₃⁻ and NO₂ photolysis in the local snowpack are a significant source of NOₓ at WAIS Divide. The upper limit of 15 h NOₓ lifetime confirms that short-lived NOₓ species are more sensitive to local production above WAIS than to outflows from the East Antarctic Plateau. Lower NO₃⁻ preservation in snow at WAIS Divide than at Summit is coherent with the potential NOₓ emission flux and suggests stronger lateral export of emitted NOₓ than local recycling.

O₃ production at WAIS Divide can be triggered with NO mixing ratios higher than 2 ppbv, but the small O₃ production rate cannot explain the observed O₃ variability, which suggests a significant air mass transport contribution. Air flows from the East Antarctic Plateau are only associated with high O₃ mixing ratios if the air transport is near surface by gravity driven winds and less than 3 days from the East Antarctic Plateau. With levels of O₃ over the East Antarctic Plateau twice those observed at WAIS Divide, outflows from the interior of the continent therefore have the potential to enhance the mixing ratios of long-lived atmospheric chemical species such as O₃ over WAIS. During this 28-day campaign, outflows from the Antarctic Plateau were observed over 7.5 days, raising the summer O₃ average by about 20 % (2 ppbv). The possibility of halogen chemistry above inland WAIS that would impact the atmospheric photochemistry is not consistent with satellite observations and cannot therefore explain the low O₃ levels that are associated with air mass origins from the interior of WAIS.

Results from the back-trajectory analysis over December 2008 and January 2009 at WAIS Divide (Fig. 9) compare well with those of Markle et al. (2012) calculated over thirty years in the Ross Sea region. For the same months, the authors found that 45 % of the air mass trajectories have an oceanic/West Antarctic origin and 55 % a continental/East Antarctic source, while we observed respective values of 29 % and 71 % at WAIS Divide. These results confirm the minor contribution of oceanic/West Antarctic inflows to the WAIS atmosphere during austral summer shown by Nicolas and Bromwich (2011), and the predominance of continental/East Antarctic air masses that may significantly impact the oxidative capacity of the atmosphere above the interior of WAIS.
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