1	HO_2NO_2 and HNO_3 in the coastal Antarctic winter night: A "lab-in-					
2	the-field" experiment					
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14 Abstract

- 15 Observations of peroxynitric acid (HO₂NO₂) and nitric acid (HNO₃) were made during a 4 month
- 16 period of Antarctic winter darkness at the coastal Antarctic research station, Halley. Mixing ratios of
- 17 HNO₃ ranged from instrumental detection limits to ~8 parts per trillion by volume (pptv), and of
- HO₂NO₂ from detection limits to \sim 5 pptv; the average ratio of HNO₃:HO₂NO₂ was 2.0(±0.6):1, with
- 19 HNO₃ always present at greater mixing ratios than HO₂NO₂ during the winter darkness. An extremely
- 20 strong association existed for the entire measurement period between mixing ratios of the
- respective trace gases and temperature: for HO₂NO₂, $R^2 = 0.72$, and for HNO₃, $R^2 = 0.70$. We focus
- 22 on three cases with considerable variation in temperature, where wind speeds were low and
- constant, such that, with the lack of photochemistry, changes in mixing ratio were likely to be driven
- by physical mechanisms alone. We derived enthalpies of adsorption (ΔH_{ads}) for these three cases.
- The average ΔH_{ads} for HNO₃ was -42±2 kJ.mol⁻¹ and for HO₂NO₂ was -56±1kJ.mol⁻¹; these values are extremely close to those derived in laboratory studies. This exercise demonstrates i) that adsorption
- to/desorption from the snow pack should be taken into account when addressing budgets of
- 28 boundary layer HO_2NO_2 and HNO_3 at any snow-covered site, and ii) that Antarctic winter can be used
- as a natural "laboratory in the field" for testing data on physical exchange mechanisms.
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33

35 1 Introduction

- 36 Peroxy nitric acid (HO₂NO₂, also written as HNO₄) and nitric acid (HNO₃) are acidic gases that are of
- 37 increasing interest to polar tropospheric chemistry. Their primary relevance is that they act as
- 38 reservoir species for HOx and NOx, which are now recognised to drive the surprisingly vigorous
- 39 oxidation chemistry that has been observed during Antarctic summer (e.g. Davis et al., 2001; Chen et
- 40 al., 2001). The spatial and temporal distribution of HO_2NO_2 and HNO_3 across the polar regions thus
- 41 becomes important for understanding the overall atmospheric chemical system, and models require
- 42 details of their sources, and any physical exchange process by which they move from one
- 43 environmental compartment to another. Currently, many of these details are unknown.
- 44 The gas-phase chemistry of HO₂NO₂ and HNO₃ is relatively straightforward. Peroxy nitric acid is a
- 45 somewhat unstable molecule that forms and dissociates through its temperature-dependent46 equilibrium reaction:

47
$$HO_2 + NO_2 + M \leftrightarrow HO_2NO_2 + M$$
 R1, -1

48 which renders an increased stability for HO₂NO₂ at lower temperatures.

There are a number of photodissociation pathways which drive HO₂NO₂ and HNO₃ chemistry under
 sunlit conditions. The most important are thought to be:

51	$HO_2NO_2 + hv \rightarrow$	$HO_2 + NO_2$	R2a

52 \rightarrow OH + NO₃ R2b

53 Peroxynitric acid can also be lost through reaction with OH:

54
$$HO_2NO_2 + OH \rightarrow H_2O + NO_2 + O_2$$
 R3

55 Gas-phase production of nitric acid proceeds via:

56 $OH + NO_2 + M \rightarrow HNO_3 + M$ R4

57 The major loss processes are reaction with OH and photolysis:

58
$$HNO_3 + OH \rightarrow H_2O + NO_3$$
 R5

59
$$HNO_3 + hv \rightarrow OH + NO_2$$
 R6

60 Both HO₂NO₂ (Ulrich et al., 2012) and HNO₃ (Bartels-Rausch et al., 2002; Hudson et al., 2002;

61 Ullerstam et al., 2005) have been shown in laboratory experiments to adsorb to ice surfaces. This

62 conclusion is supported by field observations which have confirmed uptake of both HNO₃ and

- HO_2NO_2 to snow surfaces (Huey et al., 2004; Slusher et al., 2002), and of HNO_3 to cirrus clouds
- 64 (Weinheimer et al., 1998; Popp et al., 2004; Ziereis et al., 2004). In general, therefore, in snow
- 65 covered areas, or indeed regions of the atmosphere with lofted snow/ice, such as cirrus clouds or
- blowing/precipitating snow, physical adsorption of HNO_3 and HO_2NO_2 from the air to the snow/ice is
- 67 likely to occur. The details of this uptake will differ somewhat between the two molecules because

- 68 the partitioning coefficient of HNO_3 is greater than that for HO_2NO_2 . HNO_3 is thus more sticky than
- HO_2NO_2 , and a higher fraction of HNO_3 can be expected on the ice surface compared with HO_2NO_2 .
- High resolution observations of HNO_3 and HO_2NO_2 in the polar regions are scarce. Critically, both
- HO₂NO₂ and HNO₃ have been measured together during a number of Antarctic studies at high
- temporal resolution. These studies have included both ground-based experiments at South Pole
- 73 (Slusher et al., 2002; Huey et al., 2004) and airborne measurements across the wider Antarctic
- 74 Plateau (Slusher et al., 2010).
- 75 The ground-based studies have revealed considerable inter-annual variability in summertime HNO₃
- and HO₂NO₂ mixing ratios, but always of the order 10s pptv at the South Pole. For example, the
- $\label{eq:median} 77 \qquad \text{median observed } HNO_3 \text{ between 16 and 31 December in 2000 was 18.2 pptv and for } HO_2NO_2 \text{ was}$
- 23.5 pptv (Davis et al., 2004); over the equivalent time period in 2003, the median HNO_3 was 84
- pptv, and for HO_2NO_2 was 39 pptv (Eisele et al., 2008). Considerably greater mixing ratios have also
- 80 been observed; for example, the median mixing ratio of HNO_3 between 15 and 30th November 2003
- was 194 pptv, and of HO₂NO₂ was 63 pptv (Eisele et al., 2008). While the specific mixing ratios will be
 strongly influenced by boundary layer height, overall, these high mixing ratios are fuelled by in situ
- $rac{1}{2}$ production from elevated levels of NO_x and HO_x within the South Pole boundary layer, in turn driven
- 84 by photochemical release of trace gases from the surrounding snowpack (Davis et al., 2001, 2008).
- 85 The airborne measurements assessed the three-dimensional distribution of HO₂NO₂ and HNO₃
- 86 across the Antarctic Plateau during the ANTCI 2005 campaign (Slusher et al., 2010). They revealed
- 87 significant vertical gradients in both species, with higher concentrations at the ground, consistent
- 88 with a source associated with emissions from the snowpack. The measurements also showed a
- 89 widespread distribution of both HNO₃ and HO₂NO₂ across the Plateau region.
- 90 To date there have been no measurements of high temporal resolution HO_2NO_2 and HNO_3 in coastal
- 91 Antarctica, and no measurements at all from Antarctica outside the summer season. We report here
- 92 observations of HO_2NO_2 and HNO_3 made using a chemical ionisation mass spectrometer (CIMS) at
- Halley research station in coastal Antarctica (75° 35' S, 26° 39' W) from 24 May to 18 September
- 94 2007. The data allow us to assess whether HNO_3 and HO_2NO_2 are present in significant
- 95 concentrations at other Antarctic locations and seasons than the Antarctic Plateau in summer. They
- 96 also provide an opportunity to test laboratory-derived physical exchange parameters under semi-
- 97 constrained, but genuine real-world conditions. At Halley, the sun remains below the horizon from
- 98 April 30th to August 13th, such that this new data set includes many weeks of winter darkness. Under
- these conditions of 24 hour per day darkness, atmospheric photochemistry stalls, and trace gas
- 100 concentrations are controlled entirely by either transport or physical air-snow exchange.
- 101
- 102 2 Experimental

103 2.1 CIMS instrumentation

- 104 The CIMS instrument used in this study has been described in detail elsewhere (Buys et al., 2013). It
- 105 was installed in the Clean Air Sector Laboratory (CASLab), which is located roughly 1 km from the

- 106 main Halley station, and in a sector that rarely receives air from the base (Jones et al 2008). The
- 107 CIMS inlet extended ~20 cm above the roof of CASLab, at a height roughly 5 m above the
- 108 surrounding snowpack. The inlet system was designed to minimise residence time and surface losses
- 109 (Neuman et al., 1999).
- 110 The instrument employed the SF_6^- method to detect both HNO_3 and HO_2NO_2 , using the NO_4^- (HF)
- 111 cluster at m/z 98 to detect HO_2NO_2 , and $NO_3^-(HF)$ at m/z 82 to detect HNO_3 (as per Slusher et al.,
- 112 2001, 2002). Calibration was achieved using the SO₂ method as described by Slusher et al. (2001)
- and Kim et al. (2007). Background measurements, or zeros, were obtained every 10 minutes. These
 were achieved by passing sampled air for 3 minutes through a customized filter filled with activated
- 115 coarse charcoal and nylon glass wool coated in NaHCO₃. This scrubbing method has previously been
- shown to be efficient at removing both HO_2NO_2 and HNO_3 from sampled air (Slusher et al., 2001).
- 117 The instrument detection limit derived from background data averaged over 10-minutes was 0.7
- 118 pptv for HNO_3 and 0.4 pptv for HO_2NO_2 . Total estimated uncertainty in the CIMS observations is
- 119 ±40%.
- 120 While the SF₆⁻ method has been used successfully in previous field campaigns (e.g. Slusher et al.,
- 121 2002; Slusher et al., 2010), it has been demonstrated in laboratory studies (Slusher et al., 2001) that
- 122 SF_6^- reacts with H_2O in the sample air flow. This introduces an interferent into the technique, the
- 123 non-linearity of which is evident in the unfiltered data (not shown). However, with their instrument
- reaction time of ~25 ms, Slusher et al. (2001) also concluded that this interferent was significant only
- at dewpoints greater than -25°C, and that at lower dewpoints, the interferent was negligible. During
- the period of measurements at Halley, the CIMS instrument also operated with a reaction time of
 ~25 ms, such that the interferent would be equivalent to that of Slusher et al. (2001). At Halley,
- dewpoint temperatures varied from -12° C to -52° C (mean -31° C), but were below -25° C for 81% of
- 129 the time. To remove the potential for H_2O interference in our data, all measurements made at
- 130 dewpoints above $-25^{\circ}C$ are filtered out from the dataset.
- 131

132 **2.2 Boundary layer meteorology**

- 133 Measurements of near-surface boundary layer meteorology were made on a 32-m profiling mast
- located ~25 m from the CASLab. Bulk sensors were located at 1, 2, 4, 8, 16, and 32 m above ground
- level, recording at 1Hz and averaged to 10 minute means; temperatures and humidity were
- 136 measured with platinum resistance thermometers (0.1 K resolution) and solid state humidity probes
- 137 (2% resolution), respectively, using an aspirated HMP35D from Vaisala Corp. 10 minute vector
- average wind speed and direction were measured with R.M. Young propeller vanes at 0.1 ms⁻¹ and
- 139 2° resolution .
- 140 Three 3-axis Metek USA-1 ultrasonic anemometer/thermometers were deployed at the 4, 16, and
- 32-m levels, sampling at 20 Hz. The data were tilt corrected and the relevant co-variances calculatedover 1 minute means.
- 143

144 **3 Results and discussion**

145 **3.1 Overall data series**

- 146 The time series of filtered HO₂NO₂ and HNO₃ data, averaged to hourly means, is shown in Figure 1,
- 147 together with hourly means of ambient temperature, dew point temperature, and solar zenith
- 148 angle. The solar zenith angle shows that the sun was below the horizon for the majority of this time
- 149 period, and observations of NO and NO₂ (made using a Sonoma Tech. Dual channel
- 150 chemiluminescence analyser (Bauguitte et al., 2012; Cotter et al., 2003), with detection limits of 2
- 151 pptv for NO and 6 pptv for NO₂, not shown) were consistently below the instrumental detection
- 152 limits.
- Regardless of this apparent lack of photochemical activity, there is considerable variability in the
 HNO₃ and HO₂NO₂ observations. Mixing ratios of HNO₃ ranged from instrumental detection limits to
 parts per trillion by volume (pptv) and of HO₂NO₂ varied from detection limits to ~5 pptv. These
- values are considerably lower than those observed at South Pole in summer, where photochemical
- production is fuelled by emissions of NOx from the snowpack, and where mixing ratios of HNO_3 and
- HO_2NO_2 were generally in the 10s of pptv, and sometimes over 100 pptv, as discussed earlier.
- 159 It is also noticeable in the Halley data that the pattern of variability in both HO_2NO_2 and HNO_3 was
- 160 very similar, with each time series tracking the other closely. Indeed the correlation between the
- 161 two chemical species was high, as shown in Fig. 2, where the correlation coefficient, R², for the
- hourly averages was 0.70. This finding is consistent with the data of Slusher et al., (2002), which
- showed that, although the range of mixing ratios at South Pole in summer were considerably higher
- than observed at Halley during the winter (<5 to 54 pptv for HO_2NO_2 , and <5 pptv to 68 pptv for
- HNO₃), the variability observed in both species during the measurement period was, again, highlycoupled.
- 167 The timeseries presented in Fig. 1 also clearly shows the very strong association between mixing
- 168 ratios of HNO₃ and HO₂NO₂ and ambient (and dewpoint) temperature. This association is further
- demonstrated in Fig. 3, which shows both HNO_3 and HO_2NO_2 plotted against ambient temperature,
- again for the entire period of measurements. Calculated correlation coefficients with temperature
- are high, with $R^2 = 0.70$ and 0.72 for HNO₃ and HO₂NO₂ respectively.
- 172 Given that mixing ratios of HNO₃ and HO₂NO₂ are so strongly associated with ambient temperature,
- 173 Table 1 gives the mean and standard deviation of HNO₃ and HO₂NO₂ measured between 24 May and
- 174 18 September 2007, calculated within specific temperature ranges. The statistics are derived using
- 175 only the filtered data that were above 3-sigma detection limits. At South Pole during the summer,
- ambient temperature ranged from -31.5°C to -23.6°C, with a mean of -27.7°C; within this
- temperature range, HNO_3 mixing ratios ranged from <5 to 54 pptv (mean 25 pptv) and HO_2NO_2
- 178 ranged from <5 to 68 pptv (mean 22pptv). For the equivalent temperature range at Halley (also with
- a mean of -27.7°C), mean and maximum mixing ratios for HNO₃ were 4.4 pptv and 9 pptv
- $180 \qquad \text{respectively, and for } HO_2NO_2 \text{ were } 2.5 \text{ pptv and } 5 \text{ pptv respectively, clearly significantly below those}$
- 181 observed at South Pole, for the reasons outlined above.
- 182 Differences between South Pole summer and Halley winter are also evident in the ratio of
- 183 HNO₃:HO₂NO₂, arising through differences in the species' lifetimes. Throughout the Halley
- measurement period, the average ratio of HNO_3 : HO_2NO_2 was 2.0(±0.6):1, with HNO_3 always (apart

- from a few outliers) present at greater mixing ratios than HO_2NO_2 during the winter darkness (see 185
- Fig. 4). This finding is in contrast to observations from South Pole during sunlit summer time, when 186
- mixing ratios of HNO₃ and HO₂NO₂ were roughly equal for much of the measurement period (Slusher 187
- et al., 2002). Figure 1 of Slusher et al. (2002) shows that HO₂NO₂ was present at higher mixing ratios 188
- 189 than HNO₃ for roughly 2 out of the 7 days of measurements; during the roughly 4 months of
- measurements at Halley, the only occasion when the mixing ratio of HO₂NO₂ exceeded that of HNO₃ 190
- was on the 6th and 7th September, a period when temperatures were particularly low but there was a 191
- 192 limited amount of sunlight.
- 193

194 3.2 Short-term variability in HNO₃ and HO₂NO₂ and link to ambient temperature

- 195 The short-term variability in the HNO₃ and HO₂NO₂ is shown more clearly in Figure 5. The three
- 196 examples show periods when ambient air temperatures varied rapidly and considerably, but where
- 197 they remained below the -25° C dewpoint threshold such that no chemical data filtering was
- 198 required. These 10-minute averages show that even very small-scale features of temperature
- change are reflected in the chemical measurements. For example, at midnight on 5th June, the short-199
- lived peak in temperature is reflected also in HNO₃ and HO₂NO₂; the temperature peak around 11am 200
- 201 on 21^{st} June is apparent with similar, small, peaks in HNO₃ and HO₂NO₂; and the short-lived temperature peak around noon on 15th July is also evident in short-lived increases in HNO₃ and 202
- 203 HO_2NO_2 mixing ratios. While large-scale variability in HNO_3 and HO_2NO_2 could be linked to air mass
- 204 origin, such fine-scale variability can only be explained by a local, fast-acting, source/sink
- 205 mechanism. The association between variability in HNO₃ and HO₂NO₂ and changes in ambient
- 206 temperature strongly suggest a temperature-dependent mechanism. Given our understanding of the
- 207 interaction between acidic gases and ice gained through laboratory studies (e.g. Huthwelker et al.,
- 208 2006), one possible mechanism is temperature-dependent adsorption/desorption at the snow surface.
- 209

210

211 3.3 Evidence for HO₂NO₂ and HNO₃ air/snow exchange

- 212 To probe in more detail the response of HNO_3 and HO_2NO_2 to changes in temperature, we examined
- 213 periods in the data where ambient temperatures changed, but where wind speeds were relatively
- 214 low and invariable. By adopting this approach, we minimise any influence that air flow through the
- 215 snow (e.g. via ventilation/wind pumping) may have on air/snow exchange processes. We derive a
- 216 mixing diffusivity to determine the timescale for vertical mixing (via turbulent diffusion) between the
- 217 snow surface and the CIMS inlet height, in order to confirm that the CIMS HNO_3 and HO_2NO_2
- 218 observations can be used to analyse processes occurring at the ground-level air/snow interface. The
- 219 mixing diffusivity is roughly equal to $k.z.u_*$, where k = von Karman's constant (=0.4), z = CIMS inlet
- 220 height (=5 m) and u_* , the friction velocity, is derived from the sonic anemometer data (Stull 1988).
- The e-folding time scale, t_{sc} , is given by z^2 /diffusivity, that is $t_{sc} = z/(k.u_*)$. t_{sc} will vary during each case 221
- study, but cannot be negative: this range is presented below derived from log means and standard 222 deviations. Figure 6a shows observations made on 30th May 2007, with a clear gradual increase in 223
- 224 both HNO₃ and HO₂NO₂ as ambient temperatures rose from \sim -44°C to \sim -30°C. On this day, data from

- the boundary layer mast (not shown) show that between the surface and 8 m height, there was little
- or no temperature gradient; to first order, therefore, 8 m temperatures can be used as a surrogate
- for those at the ground. Wind speeds were between 0 and 2 ms⁻¹ from the surface to 4 m, and
- remained at around 2 m.s⁻¹ at 8 m height. For 95% of the time, t_{sc} was between 100 s and 300 s.
- Figure 6b) shows data for the period from 9am to midnight on 21st June, discussed briefly in section
- 3.2 above. Again, a gradual increase in mixing ratios of HNO_3 and HO_2NO_2 is evident (upper panel), as
- ambient temperatures rose gradually from \sim -38°C to \sim -26°C. Data from the boundary layer met mast
- show that, during this period, there was no vertical gradient in temperature between the surface and 8 m height; wind speeds from the surface to 8 m were below 2 ms⁻¹. Data from the sonic
- anemometers show that vertical mixing was again very weak, with mixing time scales between 90 s
 and 600 s.
- Figure 6c) shows observations from midnight to 9am on 18th July, another quiescent period, with
- wind speeds in the lowest 8 m below 1 ms⁻¹, and with no temperature gradient below 32 m. Gradual
- 238 increases in both HNO₃ and HO₂NO₂ proceed as ambient temperatures rise from -38°C to -30°C. t_{sc}
- varied between 30 s and 100 s during the event.
- 240 Correlation coefficients between mixing ratios of HNO₃ (and HO₂NO₂) and temperature, are
- extremely high for the time periods presented in Figure 6: R² for the correlation between HNO₃ and
- temperature is 0.72 (30th May), 0.90 (21st June) and 0.72 (18th July); for the correlation between
- HO_2NO_2 and temperature, R² was 0.88 (30th May), 0.94 (21st June), and 0.92 (18th July). The values of
- R^2 show that between 72% and 90% of the variability in HNO₃ can be explained by variability in
- temperature; and between 88% and 94% of the variability in HO₂NO₂ can be explained by the
- 246 variability in temperature.
- 247

248 **3.4 Deriving enthalpy of adsorption from the Halley field data**

- 249 The enthalpies of adsorption between HNO₃/ice and HO₂NO₂/ice have been derived in laboratory
- 250 experiments carried out under environmentally-relevant conditions. Ulrich et al. (2012) studied
- 251 uptake of HO₂NO₂ at low concentrations and temperatures between 230 K and 253 K while Bartels-
- Rausch et al. (2002) and Ullerstam et al. (2005) studied the adsorption enthalpy of HNO₃. Field
- studies carried out during the 24-hour per day darkness of Antarctic winter provide optimum
- 254 conditions for validating such laboratory-derived physical air/snow exchange parameters under
- 255 "real-world" conditions.
- At equilibrium, the partitioning of HO_2NO_2 or HNO_3 molecules between the gas phase (C_g) and the snow/ice surface (C_s) can be expressed as:

$$K_{part} = \frac{C_s}{C_g}$$

As the partition constant will obey the van't Hoff equation, a new equation can be written:

$$\frac{dlnK_{part}}{d\frac{1}{T}} = -\frac{\Delta H_{ads}}{R}$$

- 259 where T is the temperature (K), ΔH is the enthalpy of adsorption (J/mol), R is the gas constant (8.314 $J.K^{-1}.mol^{-1}$) 260
- 261 Given our definition of K_{part} above, the equation can then be re-formulated as:

$$\frac{dln\frac{C_s}{C_g}}{d\frac{1}{T}} = -\frac{\Delta H_{ads}}{R}$$

262 At 240 K, roughly the temperatures of our observations, Ulrich et al. (2012), in their Figure 4, show

 $C_s/C_g \approx 20$ cm for HO₂NO₂ and 8000 cm for HNO₃. In the firn, the ratio of the surface area of snow to 263 the volume of air is approximately 0.5 to 5 cm⁻¹ (based on a density of snow of 0.3 g cm⁻³ and a 264

specific surface area of 100 to 1,000 cm²g⁻¹ (Domine et al., 2008)). We can therefore calculate that

265 the ratio of the number of molecules of HNO₃ adsorbed to the snow surface to that in the gas phase, 266

is approximately 4×10^5 to 4×10^6 ; for HO₂NO₂, this ratio is 1000 to 10,000. As a result, for both 267

HNO₃ and HO₂NO₂, exchange between the air and snow will thus have little effect on C_s, which can 268

- 269 therefore be considered as a constant relative to C_g.
- 270 If we also assume that, over several hours, and under low and constant wind conditions, the
- 271 concentration at our inlet tracks the concentration in the firn, then:

$$\frac{dlnC_g}{d\frac{1}{T}} = \frac{\Delta H_{ads}}{R}$$

- 272 It is then possible to derive ΔH_{ads} from the slope of InC_g vs 1/T multiplied by R.
- 273 For the Halley data, Figure 7 shows plots of both ln(HNO₃) vs 1/T and ln(HO₂NO₂) vs 1/T for the time 274 periods discussed in section 3.3 above. As a reminder, these periods are characterised by 24-hour 275 per day darkness, low wind speeds, and limited vertical mixing from turbulent diffusion, so are as

276 close to laboratory conditions as could be found in our dataset. They were also chosen as they

277 spanned a reasonably large temperature range, which would improve the constraint on the linear fit.

- 278 The values of ΔH_{ads} derived from these fits are given in Table 2. The average ΔH_{ads} for HNO₃ is -42±2 279 kJ.mol⁻¹ which can be compared with laboratory-derived values of Bartels-Rausch et al. (2002) and
- Ullerstam et al. (2005). Bartels-Rausch et al. (2002) derived ΔH_{ads} of -44 kJ.mol⁻¹ (with random error 280
- 2.3 kJ.mol⁻¹, systematic error 13 kJ.mol⁻¹); Ullerstam et al. (2005), working at lower concentrations of 281
- HNO₃, relevant to the natural atmosphere, derived ΔH_{ads} of -30.6 ± 6.0 kJ.mol⁻¹. For HO₂NO₂, the 282
- average ΔH_{ads} derived from our field data is -56±1 kJ.mol⁻¹ which can be compared with the 283
- laboratory-derived value (Ulrich et al., 2012) of -59 kJ.mol⁻¹. For both HO₂NO₂ and HNO₃, the 284
- agreement between laboratory and field-derived enthalpies of adsorption is remarkably good. 285

286

4. Summary and conclusions 287

- 288 We present the first high time resolution observations of HNO_3 and HO_2NO_2 in coastal Antarctica,
- 289 and the first from Antarctica during the dark winter period. Mixing ratios of HNO₃ ranged from
- 290 instrumental detection limits to \sim 8 parts per trillion by volume (pptv) and of HO₂NO₂ varied from
- 291 detection limits to ~5 pptv. These values are on average lower than those observed at South Pole in
- 292 summer, where mixing ratios of HNO₃ and HO₂NO₂ were generally in the 10s of pptv, and sometimes
- 293 over 100 pptv.

294 The Antarctic, during winter, is an ideal natural laboratory for studying physical air/snow exchange

- 295 processes. The environmental system is considerably simplified compared with other times of the
- 296 year because of the lack of photochemical activity which must otherwise be taken into account
- 297 when interpreting data.
- 298 In our study, we considered whether adsorption/desorption of HNO₃ and HO₂NO₂ to snow/ice
- 299 surfaces could be invoked to explain our observations; we did not consider formation of solid
- 300 solutions from solid ice, or take-up to liquid NaCl aerosols. In a follow-up study that further analysed
- 301 our field data, Bartels-Rausch (2014) considered both the case of Langmuir adsorption to the ice
- 302 surface, and solubility in ice forming a solid solution. He found that equilibrium air/snow partitioning
- 303 was able to describe our field data well, both in terms of absolute mixing ratios and trend with
- 304 temperature. He also found that the reservoir of adsorbed HNO_3 and HO_2NO_2 in the upper snow
- 305 pack was sufficient to fuel the observed emissions. In contrast, while calculations based on
- 306 reversible solid-solution/air partitioning were able to describe mixing ratios of HNO₃, they were not 307 able to reproduce the observed trend with temperature. Further, the reservoir of HNO₃ in the outer
- 308 part of the snow crystals was too small to explain observed increases in mixing ratio.
- 309 The measurements of HNO₃ and HO₂NO₂ from Halley are consistent with laboratory experiments
- 310 showing a temperature-dependence in the partitioning of both HNO_3 and HO_2NO_2 to ice. They
- 311 further support the conclusion that HO₂NO₂/ice interactions are stronger than those between HNO₃
- 312 and ice, as shown by the higher enthalpy of adsorption of HO_2NO_2 compared with HNO_3 (Ulrich et
- 313 al., 2012). On short-timescales, therefore, HNO_3 and HO_2NO_2 that is adsorbed to snow/ice can be re-
- 314 released as temperatures rise. The snowpack can thus act as a source of HNO_3 and HO_2NO_2 to the
- 315 overlying atmosphere at all times of the year given sufficient reservoir in the snowpack and changing
- 316 temperatures. Similarly, HNO₃ and HO₂NO₂ adsorbed to cirrus clouds would be desorbed should 317 temperatures rise. Such a reversible, temperature-dependent partitioning also provides a
- 318
- mechanism for re-distributing HNO₃ and HO₂NO₂ on a local or regional scale across Antarctica. Snow 319 can be transported considerable distances by storm systems, and adsorbed HO₂NO₂ and HNO₃ can
- 320 be desorbed as a function of changing temperature experienced along the transport pathway.
- 321 Indeed, transport of snow from inland Antarctica is likely to contribute to the HO₂NO₂ and HNO₃
- 322 reservoir in the coastal snowpack. The other likely source for HO_2NO_2 to the winter snowpack is the
- 323 general shift in equilibria, as temperatures fall from summer to winter, from gas-phase HO_2 and NO_2
- 324 towards gaseous HO₂NO₂, and then, by temperature-dependent partitioning, towards snowpack-
- 325 adsorbed HO₂NO₂.
- 326 While clearly a controlling mechanism during polar night, the importance of air/snow partitioning
- 327 relative to photochemistry will vary according to time of year and location. However, adsorption
- 328 to/desorption from the snow pack should be taken into account when addressing budgets of

- boundary layer HO_2NO_2 and HNO_3 at any snow-covered site, as all are likely to experience varying ambient temperature which would drive such air/snow exchange.
- 331

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- Table 1. Mean and standard deviation of nitric acid and peroxy nitric acid according to ambient air
 temperature range. The statistics were derived using only data above the 3-sigma detection limit.
- Table 2. The enthalpies of adsorption to ice for HO₂NO₂ and HNO₃ as derived from three periods of
 the Halley measurements. The number of observations used for each derivation is also given.
- Fig. 1. Time series of HNO₃, HO₂NO₂, ambient temperature, dewpoint, and solar zenith angle (SZA)
 (hourly averages) for the entire measurement period discussed in this paper, 24 May 2007 to 18
- 433 September 2007.
- 434 Fig. 2. Nitric acid vs peroxy nitric acid; 1-houly averages of measurements made from 24 May 2007
 435 to 18 September 2007.
- Fig. 3. Nitric acid and peroxy nitric acid vs ambient temperature; 10-minute averages of
 measurements made from 24 May 2007 to 18 September 2007.
- Fig. 4. Ratio of HNO₃ : HO₂NO₂ (hourly average data) for the May to September measurement
 period. Also shown, for reference, are ambient temperature and solar zenith angle.
- Fig. 5. Three examples of short-term variability in HNO₃, HO₂NO₂ and ambient air temperature (10minute data), from a) early June; b) mid June; c) mid July. All three periods are during the winter 24hour per day darkness.
- Fig. 6. Detail of changes in HNO₃, HO₂NO₂ and temperature on a) 30th May, b) 21st June, and c) 18th
 July. These three periods in the measurement series were characterised by low and invariant wind
 speeds and 24-hours per day darkness.
- 446 Fig. 7. Plots of ln(HNO₃) and ln(HO₂NO₂) vs 1/T for the time periods shown in Fig. 6, i.e. a) 30th May,
 447 b) 21st June, and c) 18th July.

Ambient air	Nitric acid		Peroxy nitric acid			
temperature (°C)		mean	std.		mean	std.
	no. points	(pptv)	deviation	no. points	(pptv)	deviation
-45 to -47.4	24	0.96	0.12	0		
-40 to -45.9	579	1.36	0.42	472	0.58	0.14
-35 to -39.9	1575	2.13	0.74	1615	0.99	0.38
-30 to -34.9	3638	3.09	0.92	3732	1.6	0.44
-25 to -29.9	4821	4.32	1.18	4942	2.29	0.62
-21.2 to -24.9	1954	5.67	1.19	2013	3.23	0.62

Table 1. Mean and standard deviation of nitric acid and peroxy nitric acid according to ambient air

450 temperature range. The statistics were derived using only data above the 3-sigma detection limit.

	HO ₂	NO ₂	HNO ₃		
	ΔH_{ads}	no. observations	ΔH_{ads}	no. observations	
30 th May	-58 ± 2 kJ.mol ⁻¹	144	-36 ± 2 kJ.mol ⁻¹	144	
21 st June	-52 ± 2 kJ.mol ⁻¹	90	-50 ± 2 kJ.mol ⁻¹	82	
18 th July	-58 ± 2 kJ.mol ⁻¹	55	-41 ± 4 kJ.mol ⁻¹	55	

453 Table 2. The enthalpies of adsorption to ice for HO_2NO_2 and HNO_3 as derived from three periods of

454 the Halley measurements. The number of observations used for each derivation is also given.



Fig. 1. Time series of HNO₃, HO₂NO₂, ambient temperature, dewpoint, and solar zenith angle (SZA)

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Fig. 7. Plots of $ln(HNO_3)$ and $ln(HO_2NO_2)$ vs 1/T for the time periods shown in Fig. 6, i.e. a) 30th May, 502 b) 21st June, and c) 18th July.