Nitrous acid formation in a snow-free wintertime polluted rural area

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Abstract. Nitrous acid (HONO) photolysis is an important source of hydroxyl radicals (OH) in the lower atmosphere, in particular in winter when other OH sources are less efficient. The nighttime formation of HONO and its photolysis in the early morning have long been recognized as an important contributor to the OH budget in polluted environments. Over the past few decades it has become clear that the formation of HONO during the day is an even larger contributor to the OH budget and additionally provides a pathway to recycle NOx. Despite the recognition of this unidentified HONO daytime source, the precise chemical mechanism remains elusive. A number of mechanisms have been proposed, including gas-phase, aerosol, and ground surface processes, to explain the elevated levels of daytime HONO. To identify the likely HONO formation mechanisms in a wintertime polluted rural environment we present LP-DOAS observations of HONO, NO₂, and O₃ on three absorption paths that cover altitude intervals from 2 to 31, 45, and 68 m above ground level (a.g.l.) during the UBWOS 2012 experiment in the Uintah Basin, Utah, USA. Daytime HONO mixing ratios in the 2–31 m height interval were, on average, 78 ppt, which is lower than HONO levels measured in most polluted urban environments with similar NO₂ mixing ratios of 1–2 ppb. HONO surface fluxes at 19 m a.g.l., calculated using the HONO gradients from the LP-DOAS and measured eddy diffusivity coefficient, show clear upward fluxes. The hourly average vertical HONO flux during sunny days followed solar irradiance, with a maximum of \((4.9 \pm 0.2) \times 10^{10} \text{ molec. cm}^{-2} \text{ s}^{-1}\) at noontime. A photo-stationary state analysis of the HONO budget shows that the surface flux closes the HONO budget, accounting for 63 ± 32 % of the unidentified HONO daytime source throughout the day and 90 ± 30 % near noontime. This is also supported by 1-D chemistry and transport model calculations that include the measured surface flux, thus clearly identifying chemistry at the ground as the missing daytime HONO source in this environment. Comparison between HONO surface flux, solar radiation, NO₂ and HNO₃ mixing ratios, and results from 1-D model runs suggest that, under high NOₓ conditions, HONO formation mechanisms related to solar radiation and NO₂ mixing ratios, such as photo-enhanced conversion of NO₂ on the ground, are most likely the source of daytime HONO. Under moderate to low NO₂ conditions, photolysis of HNO₃ on the ground seems to be the main source of HONO.

1 Introduction

HONO is an important precursor of OH in the atmosphere. Observations in urban and rural areas show that HONO photolysis can contribute up to 55 % to the daytime radical budget, both from the photolysis of HONO accumulated throughout the night and from the photolysis of HONO continuously formed throughout the day (e.g., Alicke et al., 2002, 2003; Kleffmann et al., 2005; Kleffmann, 2007; Ren et al., 2003; Volkamer et al., 2010; Young et al., 2012). The
The importance of HONO as an OH precursor varies depending on the environment, but it generally appears to be most prominent during winter, when other OH sources are less efficient (e.g., Ren et al., 2003; Elshorbany et al., 2009). HONO chemistry can also play a significant role in the recycling of NO, as recently demonstrated by Ye et al. (2016b).

Despite the clear significance of atmospheric HONO, there is still considerable debate on its temporal and spatial distribution and its formation pathways. Observations typically show formation and accumulation of HONO in the nocturnal boundary layer, followed by a rapid decrease due to photolysis and enhanced vertical mixing in the morning (e.g., Wong et al., 2011). The nighttime formation of HONO is generally assumed to occur via the heterogeneous disproportionation of NO in the presence of surface-adsorbed water, with the following stoichiometry:

\[ 2\text{NO}_2 + \text{H}_2\text{O}_{\text{surface}} \rightarrow \text{HONO} + \text{HNO}_3. \quad (R1) \]

Nocturnal HONO formation is partly counteracted by HONO uptake on the ground, with reported surface uptake coefficients in the range of \(2 \times 10^{-5}\) to \(2 \times 10^{-4}\) (VandenBoer et al., 2013; Donaldson et al., 2014; Wong et al., 2012). Uptake on aerosol is also likely, but little information on this process is available.

During the day, the loss of HONO is dominated by its photolysis into OH and NO at wavelengths below 400 nm (Cox and Derwent, 1976):

\[ \text{HONO} + h\nu (\lambda < 400\text{nm}) \rightarrow \text{OH} + \text{NO}. \quad (R2) \]

This photolysis is, in part, balanced by the gas-phase reaction of OH and NO:

\[ \text{OH} + \text{NO} \rightarrow \text{HONO}. \quad (R3) \]

For many years, Reactions (R2), (R3), and the less important reaction of HONO + OH were assumed to be the only significant daytime HONO reactions. HONO concentrations were thus believed to be in a pseudo-steady state (PSS) determined by its photolysis rate \( (J_{\text{HONO}}) \), [OH], and [NO2]:

\[ [\text{HONO}]_{\text{SS}} = \frac{k_{\text{HONO}}[\text{NO}][\text{OH}]}{k_{\text{HONO}} + k_{\text{HONO} + \text{OH}}[\text{OH}]}. \quad (1) \]

Over the past decade, measurements have shown that daytime HONO mixing ratios tend to be 10 to 100 times higher than this PSS (e.g., Kleffmann et al., 2005; Kleffmann, 2007; Wong et al., 2012; Zhou et al., 2011). However, the precise chemical mechanism(s) responsible for the additional HONO in the atmosphere has yet to be clearly identified. A number of gas-phase, aerosol, and ground surface processes have been proposed to explain daytime HONO mixing ratios above its PSS.

Suggested daytime gas-phase formation pathways of HONO (Bejan et al., 2006; Li et al., 2008, 2014) have been found to be unimportant in the atmosphere (e.g., Sörgel et al., 2011; Wong et al., 2012; Ye et al., 2015). Daytime HONO formation pathways on aerosol and ground surfaces include the photo-enhanced heterogeneous conversion of NO2 to HONO on TiO2 (Bedjanian and El Zein, 2012; Langridge et al., 2009; Ndour et al., 2008, 2009), soot (Ammann et al., 1998; Aubin and Abbatt, 2007; Gerecke et al., 1998; Khalizov et al., 2010; Meng et al., 2010), humic acid (Bartels-Rausch et al., 2010; Stemmler et al., 2006, 2007), and organic films (Brigante et al., 2008; George et al., 2005; Gutzwiller et al., 2002). While we know little about the detailed mechanism, Stemmler et al. (2006) proposed that the formation of HONO from NO2 to HONO conversion on surface-adsorbed humic acid is first order in NO2 at low NO2 mixing ratios and linearly dependent on irradiance:

\[ P_{\text{HONO}} \propto \text{surface area} \times [\text{NO}_2] \times \text{irradiance} \quad (2) \]

Photolysis of surface-adsorbed HNO3 / nitrate and aerosol HNO3 / nitrate is another proposed heterogeneous daytime source of HONO (Zhou et al., 2002, 2003, 2007; He et al., 2006; Beine et al., 2002; Dibb et al., 2002). The proposed mechanism is similar to that of aqueous nitrate photolysis (Mack and Bolton, 1999):

\[ \text{HNO}_3(\text{ads}) + h\nu \rightarrow [\text{HNO}_3]^{*}(\text{ads}), \quad (3) \]
\[ [\text{HNO}_3]^{*}(\text{ads}) \rightarrow \text{HNO}_2(\text{ads}) + \text{O}_3(\text{ads}), \quad (4) \]
\[ [\text{HNO}_3]^{*}(\text{ads}) \rightarrow \text{NO}_2(\text{ads}) + \text{OH}. \quad (5) \]

The linear relationship between HONO production rate and the product of particulate nitrates and its photolysis was recently confirmed by Ye et al. (2016b) in the marine boundary layer, where this mechanism plays an important role in recycling NOx. While there is still some debate about the absorption cross section of surface-adsorbed HNO3 / nitrate, Ye et al. (2016a) report that the photolysis of HNO3 / nitrate is 1 to 3 orders of magnitude faster than that of gas-phase HNO3, confirming that this process is a significant source of HONO in many environments.

Su et al. (2011) proposed that soil nitrate, through a reversible acid–base reaction of nitrite produced by biological nitrification and denitrification processes, can serve as a strong source of atmospheric HONO. Their chamber experiment shows that agricultural soils with low pH are particularly efficient sources of HONO. VandenBoer et al. (2015) proposed that HONO deposited on the surface during the night could act as a HONO source the following day, through the displacement of HONO from surface nitrate by atmospheric strong acids, such as HCl and HNO3. Another proposed HONO source is the uptake of peroxynitric acid (HO2NO2) on the ground or aerosol surfaces that results in the formation of surface nitrite, which then reacts with hydrogen ions to produce HONO (Zhu et al., 1993). Lastly, HONO can also be directly emitted by combustion sources, such as vehicles, power plants, and biomass burning (Kirchstetter et al., 1996; Neuman et al., 2016).
The multitude of proposed HONO formation mechanisms motivates additional field observations to shed light on the significance of the various processes and to elucidate the question of the nature of the reactive surface, i.e., ground or aerosol. In addition, advances in our understanding of HONO formation requires observations that allow the analysis of the HONO budget, the HONO formation rate, and the identification and quantification of specific sources. Observations are also somewhat skewed to urban environments and summertime periods despite the fact that the significance of HONO is likely higher in winter.

Here we address some of these questions by presenting observations of HONO and HONO precursor concentrations, as well as vertical fluxes, at a polluted wintertime rural location in the Uintah Basin (UB), Utah, USA. The environment is substantially different than that of most other observations in that temperatures were below freezing during most of the experiment, but little to no snow was present. The overarching goal of our observations was to determine whether HONO surface fluxes are able to close the daytime HONO budget, i.e., if in this specific environment the missing HONO source is the ground, and to investigate whether any of the HONO formation mechanisms proposed in literature dominate the HONO formation.

2 Experimental

The 2012 Uintah Basin Wintertime Ozone Study (UBWOS) took place between January and early March 2012 in the Uintah Basin, Utah, USA, to study elevated wintertime O$_3$ formation (e.g., Edwards et al., 2014; NOAA, 2012). The UB is located in the northeast corner of Utah and is part of the Colorado Plateau. The floor of the UB is about 1570 m above sea level (Lyman and Shorthill, 2013; Stoeckenius, 2015). The basin is mostly rural, with a population of about 50,000, mainly living in three towns: Duchesne, Roosevelt, and Vernal. The economy of the UB is primarily based on energy production from petroleum resources, with approximately 10,000 active oil and gas wells. The Bonanza 500 MW coal-fired power plant is located in the UB, about 20 km southeast of the UBWOS main site.

The measurement main site, also known as Horse Pool site (40°8'37.339", 109°28'1.849"W), was located in the northern part of the UB, about 35 km south of Vernal. A variety of measurements were performed at the site, including observations of basic meteorological and radiometric parameters, as well as measurements of different trace gas concentrations. Here we will focus on the observations of concentrations and concentration gradients and fluxes made by UCLA’s Long-Path Differential Optical Absorption Spectroscopy (LP-DOAS) instrument (Stutz and Platt, 1997; Wong et al., 2011, 2012; Tsai et al., 2014).

2.1 LP-DOAS instrument

The LP-DOAS is an active DOAS system which consists of a Newtonian telescope that sends and receives a collimated light beam emitted by a high-pressure Xe-arc lamp. The light beam is folded once by an array of quartz corner cube retro-reflectors mounted at distances from 0.5 to 10 km. The light returned by these retro-reflectors arrays is fed into a spectrometer via a 350 µm quartz fiber that is part of a mode mixer (Stutz and Platt, 1997). Atmospheric absorption spectra were recorded at a resolution of about 0.5 nm by a Czerny–Turner type spectrometer with a focal length of 0.5 m (Acton Spectra Pro 500) in the wavelength range of 300–380 nm. A Princeton Instrument Pixis 256 CCD camera was used for the detection of the spectra. The multichannel scanning technique (MCST) was employed to overcome pixel-to-pixel sensitivity variations (Brauers et al., 1995; Platt and Stutz, 2008). Details on the instrument and the MCST can be found in Wong et al. (2012), Stutz et al. (2002), and Alicke et al. (2003).

The LP-DOAS system measured mixing ratios of NO$_2$, O$_3$, HONO, HCHO, and SO$_2$ at the Horse Pool site from 22 January to 28 February 2012. Here we will focus on the NO$_2$, O$_3$, and HONO data. The LP-DOAS telescope was located 2 m above the ground, and three retro-reflectors arrays were installed at the base, midway, and top of a bluff located about 2.5 km northwest of the main site (Fig. 1), at heights of 31, 45, and 68 m above the height of the LP-DOAS telescope. Figure 1 shows the location of the retro-reflectors relative to the location of the instrument and the three light paths (lower, middle, and upper). The retro-reflectors arrays were installed horizontally close to each other (within ~64 m). Surface topography, determined from online maps, is very similar under each light path (Fig. 1a). The light paths cross frozen natural soil and dry wintertime vegetation.
The LP-DOAS sequentially aimed at the three reflector arrays with a repetition time of 20–45 min, depending on visibility. The observations were continuous except for periods with heavy fog and times at which instrument maintenance was performed. A measurement on one retro-reflector consisted of a sequence of an atmospheric absorption spectrum, a background spectrum measurement with the lamp blocked and, at low NO₂ mixing ratios, an additional Hg reference spectrum. This sequence was repeated nine times, each with a wavelength offset of 0.8 nm. Each LP-DOAS absorption spectra underwent correction for electronic offset and atmospheric background light. The MCST procedure (Alicke et al., 2002; Platt and Stutz, 2008) was used to construct the final absorption spectrum from one sequence of nine different wavelengths. The MCST composite spectrum then underwent a low-pass filter using a 20-fold triangular smoothing (Platt and Stutz, 2008).

Trace gas reference spectra were calculated from literature absorption cross sections (Table 1) after convolution with the instrument function, which was determined by measuring the 334 nm mercury atomic emission line, and a MCST simulation with the same wavelength shift parameters as the atmospheric spectrum (Platt and Stutz, 2008). A linear combination of trace gas reference spectra and a polynomial function and a spectrum of the Xe-arc lamp was fitted to the atmospheric spectrum using a combination of a linear and non-linear least-squares fit (Stutz and Platt, 1996; Platt and Stutz, 2008). A small (< 1 pixel) joint spectral shift was allowed for the trace gases in the fit. The fit was performed in wavelength intervals specific for each trace gas (see Table 1) to optimize the detection limits and stability of the fit. Table 1 lists the references included in each fit window and the average detection limit for each spectrum. We will use these 1σ errors throughout the study. The total uncertainty of NO was determined to be 4 %, and, at low NO mixing ratios, an additional Hg reference spectrum is 144 ± 22 ppt.

Figure 2. Result of a DOAS analysis of HONO and NO₂ of a spectrum measured on 2 February 2012, 02:43 mountain standard time (MST). (a) Atmospheric spectrum with broadband features measured on the upper light path. (b) NO₂ fit result (red line) superimposed on the atmospheric spectrum without broadband features (blue line). (c) Fit result for HONO (red line) superimposed on sum of fitted HONO and unexplained residual structures in the fit (blue line); residual spectrum (black) shifted 7.5 × 10⁻⁴ on the y axis for better comparison with fitted HONO. The retrieved mixing ratio of HONO for this spectrum is 144 ± 22 ppt.

2.2 Ancillary measurements

Extensive chemical and meteorological measurements were made at the Horse Pool site. This section describes the specific measurements that were used in this study for the data interpretation and modeling efforts. A summary of all the measurements made during UBWOS 2012 can be found in Lyman and Shorthill (2013).

2.2.1 Meteorological and micrometeorological observations

Temperature, relative humidity, wind direction, and wind speed were measured at the top of a sampling tower. Temperature and relative humidity were measured with a Vaisala model HMP45AC and pressure was measured with a Vaisala model PTB101B at 13.5 m a.g.l. Wind direction and wind speed were measured with a vane anemometer (RM Young, model 05103) at 19 m a.g.l. and a three-dimensional sonic anemometer (RM Young model 81000) at 13.5 m a.g.l. Data are reported as 1-minute averages and the uncertainty is ±3 %. Turbulent heat flux, Monin–Obukhov length, and friction velocity were calculated using micrometeorological data collected by a second sonic anemometer (Applied Technologies, Inc) installed on a separate tower at 16 m a.g.l. Persson et al. (2002) estimated a random error of ±3 W m⁻² for turbulent heat flux and ±0.05 m s⁻¹ for friction velocity.

2.2.2 NOₓ and HNO₃ measurements

NOₓ measurements were made with a two-channel O₃–NO chemiluminescence detector (Williams et al., 2009). Only the NO observations from the NOₓ instruments were used in this study. The total uncertainty of NO was determined to be 4 %, with a detection limit of 6 ppt (1-minute average).
Table 1. Wavelength windows, cross sections, and fitted trace gases used for DOAS retrieval during UBWOS 2012. The quoted detection limits are campaign averages.

| Trace gases | Wavelength range (nm) | Cross section reference and uncertainty | Fit spectral references | Average detection limits |
|-------------|-----------------------|----------------------------------------|-------------------------|-------------------------|
| HCHO        | 324–346              | Meller and Moortgat (2000), 280 K, 5 % | NO2, HONO, HCHO, O3, O4 | 0.3 ppb                 |
| HONO        | 338–344 and 351–371  | Stutz et al. (2000), 298 K, 5 %        | NO2, HONO, HCHO, O3, O4 | 0.03 ppb                |
| NO2         | 338–344 and 351–371  | Vandaele et al. (1998), 294 K, 2–5 %;  | NO2, HONO, HCHO, O3, O4 | 0.07 ppb                |
| O3          | 324–346              | Voigt et al. (2002), 260 K, 4–6 %      | NO2, HONO, HCHO, O3, O4 | 1.4 ppb                 |
| SO2         | 305–310 and 313–321  | Vandaele et al. (1994), 296 K, 2.4 %   | NO2, HCHO, O3, and SO2  | 0.05 ppb                |

O3 literature cross section: Hermans et al. (1999), 298 K.

Nitric acid (HNO3) was measured by negative-ion proton-transfer chemical ionization mass spectrometry (NI-PT-CIMS), with acetate as the reagent ion. This method is described in detail by Roberts et al. (2010) and Veres et al. (2008). The NI-PT-CIMS was placed on the tower at 10 m a.g.l. The uncertainties for both HNO3 was ±20 %.

2.2.3 Photolysis rates

Photolysis rates of NO2 and O3 (O1D channel) were measured by commercial filter radiometers (Metcon, Inc.) mounted at 10 m a.g.l. The rates were measured at a frequency of 1 Hz from 15 January until 22 February, when strong wind gusts toppled the tower that held the instrument. Photolysis rates of HONO were calculated based on filter radiometer data (Stark et al., 2007). Total uncertainties for 1-minute average data are ±15 % for JNO2, ±25 % for JO1D, and ±14 % for JHONO.

2.2.4 Carbon monoxide (CO) and volatile organic compound (VOC) measurements

CO was measured with a commercial vacuum resonance fluorescence instrument; the accuracy and precision of the CO measurements were ±4 % and ±0.5 ppbv, respectively. A broad set of VOCs were measured in situ by a gas chromatograph–mass spectrometer instrument (GC-MS). The inlet of the GC-MS was set up at approximately 20 m a.g.l. A total of 65 VOCs were measured including alkanes, alkenes, cycloalkanes, aromatics, oxygenated VOCs, and nitrogen- and halogen-containing species. Uncertainties were about 15–25 % for hydrocarbons (HCs) and 20–25 % for oxygenates compounds. A detailed description of the instrument is given in the work of Gilman et al. (2010) and Goldan et al. (2004).

2.2.5 Aerosol number size distribution

Aerosol number size distribution was measured with a Scanning Mobility Particle Sizer (SMPS, TSI 3080 coupled to a TSI 3010 CN counter) and an Aerodynamic Particle Sizer (APS, TSI 3321). Aerosol particles were sampled at 12 m a.g.l. The SMPS counted particles ranging from 20 to 500 nm geometric diameter; the APS collected particles into 34 size bins with aerodynamic diameters between 0.7 and 10.37 nm. The number size distributions were collected every 5 min.

2.3 Chemical models

To aid in the interpretation of the UBWOS 2012 observations two chemistry and transport models were used. A one-dimensional chemical and transport model (RCAT8.2, run for this study) modeled vertical HONO concentration profiles and fluxes for direct comparison with the observations, while the OH output from the Weather Research and Forecasting (WRF) model coupled with chemistry was used in the HONO budget calculations. The following sections will briefly describe both models.

2.3.1 RCAT

RCAT8.2 is a highly resolved one-dimensional chemistry and transport model that has been used to investigate the impact of nocturnal processes and daytime chemistry of HONO and NO3 (e.g., Wong and Stutz, 2010; Wong et al., 2013; Tsai et al., 2014). It is based on the gas-phase Regional Atmospheric Chemistry Mechanism (RACM), which contains 77 species, 237 reactions, and aggregates atmospheric VOCs into 23 classes: four alkanes, four alkenes, three biogenic, three aromatics, and nine carbonyls (Stockwell et al., 1997). RCAT8.2 also includes biogenic and anthropogenic emissions and the Mainz Isoprene Mechanism (MIM) (Geiger et al., 2003). Aerosol uptake rates in RCAT8.2 are calculated.
using the mass-transfer equation by Fuks and Sutugin (1971). The uptake rates were calculated based on the vertical profile of the aerosol surface to air volume ratio and a particle diameter of 150 nm. A unique feature of RCAT is that heterogeneous chemistry on the ground is modeled through direct molecular interactions. The vertical grid subdivides the lowest 2000 m of the atmosphere into 23 boxes with a logarithmic spacing below 1 m to account for the inefficient vertical transport near the ground (grid points: $1 \times 10^{-4}$, $1 \times 10^{-3}$, $1 \times 10^{-2}$, 0.1, 1, 3, 6, 10, 20, 33, 50, 78, 90, 110, 121, 150, 175, 255, 300, 556, 750, 1000, and 2000 m). Vertical transport was constrained by observed meteorology, with a predominately neutrally stable atmosphere during the day. The canopy height was set to 3 m; the displacement height was set to 4.27 m and the surface roughness length was 0.03 m, to account for vegetation and small-scale topography. The chemistry portion of the model was initialized at sunset with NOAA in situ measurements of HCs (measured by GC-MS), CO (measured by CO instrument), CH$_4$ (measured by the Picarro CO$_2$/CH$_4$ instrument), PAN (measured by PAN GC), and UCLA LP-DOAS measurements of NO$_2$, O$_3$, HCHO, HONO, and SO$_2$. The initial concentrations of the trace gases changed depending on the modeled day, with sunny day averages at sunset of 2320 ppb for CH$_4$, 92.7 ppb for alkanes, 41 ppb for aromatics, 1.6 ppb for carbonyls, 0.45 ppb for PAN, and negligible biogenic VOC. Hourly NO emissions were assumed to be constant up to the planetary boundary level (PBL) height and decreased exponentially, to one-third of its surface concentration, above the PBL. The hourly aerosol surface area density diurnal values were calculated using SMPS and APS measurements. Soil NO emissions, α-pinene, and isoprene emissions were assumed to be $1 \times 10^{16}$ molec. cm$^{-2}$ s$^{-1}$, $1 \times 10^6$, and $1 \times 10^5$ molec. cm$^{-3}$ s$^{-1}$, respectively. The aerosol and ground uptake coefficients adopted in RCAT8.2 were $2 \times 10^{-3}$ for NO$_3$, $2 \times 10^{-5}$ for NO$_2$, $10^{-4}$ for HONO, 0.1 for H$_2$O, and $5 \times 10^{-5}$ for O$_3$.

2.3.2 WRF model coupled with Chemistry (WRF-Chem)

A recent version of the WRF-Chem model (version 3.5.1 http://ruc.noaa.gov/wrf/WG11/) was used to simulate O$_3$ levels during UBWOS 2012 (Ahmadov et al., 2015). Gas-phase chemistry was based on RACM (Stockwell et al., 1997), and the photolysis rates were simulated using the Tropospheric Ultraviolet and Visible (TUV) photolysis scheme (Madronich, 1987). The initial meteorological and boundary conditions were based on the North American Mesoscale analysis fields (www.emc.ncep.noaa.gov). The OH mixing ratios and PBL height model outputs were used in this study. A detailed description of the model results for UBWOS 2012 can be found in Ahmadov et al. (2015).

2.4 Analysis of the vertical distribution of HONO

LP-DOAS instruments have been used to retrieve deposition velocities, vertical gradients, and fluxes of SO$_2$ (Platt and Perner, 1979) and reactive nitrogen species (Stutz et al., 2004; Wang et al., 2006; Wong et al., 2012, 2013; Tsai et al., 2014). As illustrated in Fig. 1, the LP-DOAS observes trace gas concentrations averaged over the different light paths, i.e., averaged from the ground to the reflector height of each light path. It is thus necessary to convert these vertically averaged trace gas observations to vertical concentrations gradients and profiles. This is typically achieved using a geometric approach, where the ground is considered flat (Stutz et al., 2004; Wang et al., 2006; Wong et al., 2012, 2013; Tsai et al., 2014). While the setup of the LP-DOAS system is similar to that of previous studies, the terrain under the light paths during UBWOS exhibited significant topographical variations compared to the height interval covered by the light paths (Fig. 1). This setup thus makes a geometric approach challenging, and a better method to retrieve HONO vertical gradients was therefore developed.

2.4.1 Retrieval of HONO vertical gradients

Our retrieval approach is based on the assumption that the vertical HONO concentration profile ($C(z)$) is a function of altitude above the surface ($z$) for each point along the light path. From published field observations (VandenBoer et al., 2013; Zhang et al., 2009) as well as model results (Wong et al., 2013) we know that HONO concentrations typically decay from the surface upwards in a nonlinear way. This decay is also supported by the negative gradients in the path-averaged HONO concentrations we observed during UBWOS 2012 (see below). While a direct functional dependence of [HONO] with altitude cannot be established, an exponential decay with altitude seems to appropriately describe the profile. This is particularly true for the model results of Wong et al. (2013). We will show later that an exponential decay shape also agrees well with 1-D modeling studies for UBWOS. We therefore parameterized the HONO concentration profile using the following exponential function:

$$C(z) = C_0 + C_1 \cdot \exp \left( -\frac{z}{C_2} \right),$$  

where $C_0$, $C_1$, and $C_2$ represent the HONO background concentrations, HONO surface concentrations, and a scale height, respectively. To simulate the path-averaged concentration observed by the LP-DOAS, the light path of each reflector with length, $L_i$, was subdivided in 50 m intervals. For each interval, $j$, the height of the light path above ground, $z_j$, was determined based on the topography shown in Fig. 1 and the light-path geometry. Considering the slanted path with an elevation angle, $\alpha_i$, the path-averaged concentration is then calculated:
\[ M(\text{HONO})_i = \frac{1}{L_i} \sum_{j=1}^{L_i/50\text{m}} \frac{50\text{m}}{\cos(\alpha_i)} C(z_j). \quad (7) \]

To determine the factors \( C_0, C_1, \) and \( C_2 \) from the observed path-averaged concentrations \( M(\text{HONO})_i \) on each light path, a nonlinear least-squares fit was performed, minimizing \( \chi^2 \):

\[ \chi^2 = \sum_i (M(\text{HONO})_i - [\text{HONO}])_i)^2, \quad (8) \]

where \( i \) is the light-path index (lower, middle, and upper light path). The trust-region-reflective algorithm, which is based on the interior-reflective Newton method described in Coleman and Li (1996), was used in this optimization (MATLAB, lscurvefit). A lower allowable bound of 0 for both \( C_0 \) (HONO background) and \( C_1 \) (HONO surface concentrations) were specified in the fit. The decay term \( C_2 \) was left unbound (varying from \(-\infty\) to \(+\infty\)). The fit thus allows for decreasing, constant, or increasing altitude profiles, as long as the concentrations remain positive. Figure 3 shows the results of such a retrieval for 7 February 2012, at 11:00, 12:00, and 13:00 MST (mountain standard time, which is the local time zone in Utah and will be used throughout this paper). The comparison with results from RCAT 1-D model for these specific times confirms that Eq. (6) describes the expected HONO concentration profile well. The slight differences between the retrieved and model profiles at 11:00 and 12:00 MST can be explained by the uncertainty in the vertical mixing parameters used in the model; i.e., stronger vertical mixing in the model leads to modeled profiles which are less steep than the one retrieved from the observations. The modeled and retrieved profiles at 13:00 MST, however, show an excellent agreement, implying that the combination of surface source and vertical mixing in the model match those during the observations well.

### 2.4.2 Retrieval of HONO surface fluxes

The gradient method (Businger, 1986) was used to calculate HONO surface fluxes, i.e., the HONO flux is the product of the vertical concentration gradient, \( \frac{dC}{dz} \), and the eddy diffusivity coefficient \( (K_e) \):

\[ F_c(z) = -K_e(z) \frac{dC}{dz}, \quad (9) \]

where the eddy diffusivity \( K_e(z) \) at height \( z \) can be calculated using the von Kármán constant \( (k = 0.37) \) (Telford and Businger, 1986). Monin–Obukhov length \( (L) \), friction velocity \( (u^* \) ), displacement height \( (D) \), and the dimensionless stability correction factor \( (\Theta(\frac{z}{L})) \), which is a function of stabil-

\[ K_e = \frac{k \cdot u^* \cdot (z - D)}{\Theta(\frac{z}{L})}, \quad (10) \]

\[ \Theta(\frac{z}{L}) = \left(1 - \frac{16 \cdot (z - D)}{L}\right)^{-\frac{1}{2}} \text{ for } \frac{z}{L} < 0 \text{ (unstable)}, \quad (11) \]

\[ \Theta(\frac{z}{L}) = \left(1 + \frac{5 \cdot (z - D)}{L}\right) \text{ for } \frac{z}{L} > 0 \text{ (stable)}. \quad (12) \]

\( u^* \) and \( L \) were determined from the sonic anemometer observations described in Sect. 2.2.1. The displacement height for UBWOS was determined using the zero-plane displacement height equation by Stanhill (1969):

\[ \log(D) = 0.9793 \times \log(h) - 0.1536, \quad (13) \]

where \( h \) is the vegetation height. The displacement height for UBWOS, \( D = (0.39 \pm 0.17 \text{ m}), \) was calculated based on the average of 11 popular vegetation species in the southern UB (mainly shrubs and grass; Butler and England, 1979). \( K_e(z) \) was calculated for \( z = 19 \text{ m} \), the altitude of the micrometeorological measurements. The uncertainty of \( K_e(z) \) was determined by propagating the random error of friction velocity and Monin–Obukhov length from the observations and the uncertainty in \( D \).
To determine the HONO gradient in Eq. (9) the retrieved HONO concentration profile was averaged in two vertical height intervals. The lower interval extended from 2 m (height of the instrument a.g.l.) to 16 m a.g.l. (height of the anemometer a.g.l.), and the upper interval extended from 16 m to 40 m a.g.l. (maximum height difference between the upper light path and the ground). The difference between the average HONO mixing ratios in each box was divided by the average heights of the intervals ($z_{\text{upper}} = 28$ m, $z_{\text{lower}} = 9$ m):

$$\frac{\text{dHONO}}{dz} = \frac{\text{HONO}_{\text{upper}} - \text{HONO}_{\text{lower}}}{z_{\text{upper}} - z_{\text{lower}}}. \quad (14)$$

HONO gradients and fluxes were calculated at 19 m a.g.l. The uncertainty of the HONO gradient was determined by propagating the uncertainty associated with the average concentration of HONO in the lower and upper box. We assumed that the uncertainties in the boxes are equal to the HONO error of the DOAS analysis on the lower, middle, and upper light paths, weighted based on its contribution to the box height.

3 Results

3.1 Meteorology during UBWOS 2012

UBWOS 2012 was dominated by weak winds, with average wind speeds ranging between 2.5 m s$^{-1}$ during the night to 4 m s$^{-1}$ in the evening (Fig. 4b). Temperatures ranged between 259 and 286 K, with diurnally average temperatures varying between 270 K in the early morning and 277 K in the mid afternoon (Fig. 4a). Temperature measurements across the UB showed that daytime temperature inversions conducive to ozone formation were rare. Snow cover was almost absent during UBWOS 2012. Snow fell on 19 and 29 February but thawed quickly due to the high temperatures. For most of the field study the ground consisted of frozen natural soil and dry vegetation.

3.2 NO$_2$, O$_3$, and HONO light-path average mixing ratios

Daytime and nighttime NO$_2$ mixing ratios were, on average, ~2 ppb and ranged from $0.10 \pm 0.07$ to $28.8 \pm 0.08$ ppb on the lowest light path (Fig. 4c). While these high levels of NO$_2$ are comparable to NO$_2$ levels found in some polluted urban areas, their sources are different. In general, transportation is the main source of NO$_2$ pollution in urban areas but, in the UB, NO$_2$ was mainly emitted from oil and gas operations (57–61 %) and to a lesser extent by emissions from the Bonanza power plant, although the power plant emissions often remain aloft and are not observed at the ground in the basin (Lyman and Shorthill, 2013).

O$_3$ mixing ratios ranged from 2.4 ± 1.7 to 60.4 ± 3.3 ppb, with daytime ozone maxima consistently below the US National Air Quality Standard (75 ppb) (Fig. 4d). O$_3$ levels measured in the UB in 2012 were lower than those observed in the subsequent two winters. The absence of snow likely led to more convective mixing, which prevented pollutants from building up near the surface and reacting with sunlight to form O$_3$ (Lyman and Shorthill, 2013; Edwards et al., 2014).

Furthermore, without snow cover, photochemical formation of O$_3$ was less efficient, as the actinic flux was 1.6 to 2 times smaller (Lyman and Shorthill, 2013; Edwards et al., 2014). HONO levels were also low compared to levels measured in polluted urban areas. The nighttime and daytime path-averaged HONO mixing ratios were 56 and 78 ppt, respectively, and ranged from below the detection limits to 274 ± 33 ppt (Fig. 4e). The diurnally averaged mixing ratios of daytime HONO in the 2–70 m vertical height interval (or LP-DOAS upper height interval) ranged from 35 to 81 ppt, with an average of 48 ppt. These HONO levels are comparable to HONO observed in rural areas (e.g., Zhou et al., 2011) but lower than those observed in polluted urban environments (e.g., Su et al., 2008; VandenBoer et al., 2013; Wong et al., 2012).

3.3 Diurnal profile during sunny days

To simplify the analysis of the behavior of HONO, its precursors, and the various environmental parameters impacting its chemistry, we will concentrate only on sunny days (27 January, 4 February, 6 February, 7 February, and 17 February) from here on, when the meteorological conditions were fairly constant and the relatively smooth actinic flux simplified the interpretation of the data. The restriction to sunny days also has the advantage that we can use the PSS for budget calculations due to the short photolytic lifetime of HONO (about 13 min near noontime and less than 21 min between 09:00 and 15:00 MST). Figure 5 shows the diurnal profile of $J_{\text{HONO}}$ (Fig. 5a), NO$_2$ (Fig. 5b), O$_3$ (Fig. 5c), and HONO (Fig. 5d) on each light path (lower light path in black, middle light path in blue, upper light path in red, and the 1σ mixing ratio variation represented by the grey shadow) averaged over all sunny days.

The average diurnal profile of O$_3$ follows the expected behavior, with low mixing ratios of ~30 ppb at night and increasing O$_3$ levels after sunrise, mostly due to the mixing of the PBL (Edwards et al., 2013). As the day progresses, O$_3$ levels reach a maximum of ~46 ppb in the afternoon due to its photochemical formation. During the day, O$_3$ did not exhibit vertical gradients within our measurement uncertainty due to well mixed daytime atmospheric conditions. However, O$_3$ showed clear positive gradients at night. Together with the decreasing O$_3$ at night, this indicates that surface deposition likely played an important role (Sillman, 1999).

The sunny day diurnal average of NO$_2$ is higher at night, with maximum mixing ratios in the morning of ~4 ppb and lower values later during the day of ~1 ppb, when NO$_2$ undergoes photolysis and the boundary layer mixes. In contrast
Figure 4. Overview of meteorological parameters (a, b) and mixing ratios of (c) NO$_2$, (d) O$_3$, and (e) HONO observed on the lower light path throughout the study. Error bars in the trace gas panels (a–e) denote the statistical error of each observation derived by the DOAS analysis. Please note that the NO$_2$ and O$_3$ error bars are smaller than the data markers.

HONO shows a diurnal profile similar to that reported from urban locations. It accumulates during the night with the largest hourly average HONO mixing ratio of 0.10 ppb at 06:00 MST. A distinctive decrease after sunrise due to its efficient photolysis and daytime mixing is observed, leading to minimum HONO levels of 0.03 ppb around 15:00 MST. Daytime HONO mixing ratios are smaller than those observed in polluted urban areas. For example, in Los Angeles the UCLA LP-DOAS measured on average about 100 ppt of daytime HONO on its lowest light path (33–78 m a.g.l.). The hourly average diurnal profile of HONO generally shows larger HONO mixing ratios in the lower than in the upper light path, both during the day and night (Fig. 5d). This negative HONO gradient indicates an upward flux and the presence of a surface source of HONO. The observations of daytime negative HONO gradients (Fig. 3) are in agreement with HONO observations made in Boulder, Colorado (VandenBoer et al., 2013), where daytime mixing ratios were 100 ± 80 pptv at the surface and 35 pptv aloft, and measurements in Houston, Texas (Wong et al., 2012), that also show larger HONO mixing ratios near the surface.

3.4 HONO vertical fluxes

Both the daytime vertical profiles and negative gradients of HONO (Fig. 5d) point toward the existence of HONO surface fluxes. We therefore applied the methods outlined in Sect. 2.4 to calculate vertical HONO fluxes on sunny days. Figure 6 shows the components of such a calculations for 6 February 2012, along with NO$_2$ mixing ratios measured on the lower light path (Fig. 6a). HONO mixing ratios in the lower (black markers) and upper (red markers) boxes calculated from retrieved HONO vertical profiles (Fig. 6b) and the gradient calculated using Eq. (14) clearly show the negative gradient of HONO. By combining this gradient with the observed eddy diffusivity coefficients (Fig. 6d) in Eq. (9), the vertical HONO flux was determined. The relatively constant eddy diffusivity coefficients between 10:00 and 15:00 MST, along with the temporal trend of the HONO gradients, results in HONO surface fluxes that follow the solar irradiance temporal trend, with a maximum of (4.9 ± 0.2) × 10$^{10}$ molec. cm$^{-2}$ s$^{-1}$ near noontime. It should be noted that the temporal variation of the flux seems to be controlled by that of the HONO gradient. The source of the higher-frequency variation, i.e., at 13:30 MST, is unclear,
but one can suspect that advection could have played a role. Whether the apparent anticorrelation of the HONO gradient and flux with NO$_2$ after 11:00 MST is meaningful will be discussed further below. It should also be noted that the turbulent mixing timescale at 16 m altitude is around 160 s and is thus shorter than the photolytic lifetime of HONO. We thus report the flux without specifically considering HONO photolysis.

To allow a more thorough interpretation of our HONO flux calculations we averaged the hourly flux data over all sunny days of the experiment. The hourly average sunny day HONO flux (Fig. 7) approximately follows solar irradiance, with a maximum of $(1.7 \pm 0.3) \times 10^{10}$ molec. cm$^{-2}$ s$^{-1}$ at noontime. This value is comparable to the average HONO flux (about $1.15 \times 10^{10}$ molec. cm$^{-2}$ s$^{-1}$) measured above a northern Michigan forest canopy (Zhang et al., 2009; Zhou et al., 2011) and lower than average noontime HONO flux ($3.01 \times 10^{10}$ molec. cm$^{-2}$ s$^{-1}$) measured over an agricultural field in Bakersfield (Ren et al., 2011). The upward HONO fluxes observed during sunny days implies that daytime HONO formation occurs at the ground and is driven by solar irradiance, as we will discuss further below.
3.5 PSS daytime HONO budget

Similar to previous field observations, daytime HONO levels during UBWOS exceeded those of its photochemical steady-state levels (Eq. 1). The difference between the calculated \([\text{HONO}]_{\text{ss}}\) (Fig. 8, red line) and HONO measured in the lowest light path (Fig. 8, black line) points towards the presence of an unknown HONO source during the day. To investigate this missing source and to determine whether the upwards HONO flux from the surface explains the missing source, we performed a HONO budget analysis using a PSS approach similar to that used in Wong et al. (2012). The short lifetime of HONO justifies the use of the PSS approach (Eq. 15). It is assumed that HONO reaches a PSS between the known (\(P_{\text{unknown}}\)) sources of HONO and loss rates of HONO (Eqs. 17 and 18). Direct emissions of HONO are not considered in these studies because our measurements are not significantly influenced by traffic.

\[
\frac{d[\text{HONO}]}{dt} = P_{\text{unknown}} + \text{HONO}_{\text{formation}} - \text{HONO}_{\text{photolysis}} - \text{HONO}_{\text{oxygenation}} = 0 \tag{15}
\]

\(\text{HONO}_{\text{formation}} = k_{\text{NO} + \text{OH}}[\text{NO}][\text{OH}]\)

With \(k_{\text{NO} + \text{OH}} = (1.8 \pm 0.4) \times 10^{-13} \text{ molec.}^{-1} \text{ cm}^3 \text{ s}^{-1}\)

(275 K, 843 hPa) \tag{16}

\(\text{HONO}_{\text{photolysis}} = J_{\text{HONO}}[\text{HONO}]\)

\(\text{HONO}_{\text{oxygenation}} = k_{\text{HONO} + \text{OH}}[\text{HONO}][\text{OH}]\)

With \(k_{\text{HONO} + \text{OH}} = (4.3 \pm 0.8) \times 10^{-12} \text{ molec.}^{-1} \text{ cm}^3 \text{ s}^{-1}\)

(275 K) \tag{18}

The HONO formation and loss rates were calculated using in situ NO measurements, \(J_{\text{HONO}}\), LP-DOAS lower light-path HONO mixing ratios, and OH concentrations modeled by WRF-Chem. Modeled OH mixing ratios were used due to the lack of OH in situ measurements. An uncertainty of 20% was assigned to the WRF-Chem OH concentrations, which was determined by comparing the difference between OH calculated by RCAT8.2 and WRF-Chem during sunny days. The HONO rate constants and their uncertainties were determined using data from Sander et al. (2011) and the uncertainty in temperature.

Following Eq. (15), the HONO formation rate needed to balance the budget, \(P_{\text{unknown}}\), was calculated using

\[
P_{\text{unknown}} = J_{\text{HONO}}[\text{HONO}] + k_{\text{HONO} + \text{OH}}[\text{HONO}][\text{OH}]
- k_{\text{NO} + \text{OH}}[\text{NO}][\text{OH}] \tag{19}
\]

The uncertainties of the formation and loss rates of HONO and \(P_{\text{unknown}}\) were calculated by propagating the uncertainties of their rate constants and measurement errors. Figure 8 shows the sunny day hourly average of HONO formation (red line), HONO photolysis (green line), HONO oxidation (blue line), and \(P_{\text{unknown}}\) (black line). Surface fluxes of HONO were converted into column HONO formation rates (Fig. 8, magenta line) by dividing them by the height \((H)\) at which the influence of HONO surface fluxes on the total HONO column becomes negligible. This height was determined from two RCAT8.2 model runs for 27 January, one with and one without HONO surface fluxes. For each hour, an \(H\) was set at the height where the difference between the HONO modeled with and without HONO surface fluxes was less than 1 ppt. The average \(H\) between 09:00 and 15:00 (273 ± 113 m) was used for the calculation of the HONO surface flux rate. The uncertainty of \(H\) is the 1σ deviation of \(H\) from 09:00 to 15:00. The uncertainty of HONO flux rate was calculated by propagating the uncertainty of HONO flux and \(H\), which are the main uncertainties in this calculation.
The comparison of the various terms in Eq. (15) are shown in Fig. 8, where the grey bars indicate morning and evening periods when the HONO lifetime is too long to allow a PSS interpretation. The dominant HONO loss process is, as expected, photolysis. The HONO gas-phase formation from OH + NO does not fully balance this loss, thus clearly showing the need for an additional HONO source to close the budget. The comparison of this missing HONO source, P\text{unknown}, with the rate from surface emissions shows that HONO formation on the surface can explain this missing source, within the uncertainties of the calculation, between 11:00 and 13:00 and is somewhat lower at 10:00, when the photolysis of HONO formed at night may still contribute a small amount of HONO. The fact that surface formation cannot fully explain P\text{unknown} at 14:00 is less clear but may be a consequence of an increase of P\text{unknown} due to a decrease of P\text{NO}_2+\text{OH} caused by lower NO observations and lower modeled OH in the late afternoon.

3.6 Model daytime HONO budget

We used our 1-D chemistry and transport model, RCAT8.2, to verify that HONO surface fluxes can account for the HONO missing daytime source during UBWOS 2012. Initially, the model was run with its basic setup. NO\textsubscript{x} emissions were chosen to reproduce the observed NO\textsubscript{2} mixing ratios. The simulations (Fig. 9a, b) generally reproduce NO\textsubscript{2} mixing ratios well, except during advection events. For example, on 4 February between 10:00 and 12:00 the model has difficulties reproducing the fast concentration changes associated with the passing of a power plant plume. Similarly, the drop in NO\textsubscript{2} in the morning of 27 January is likely due to a change in air mass, which is not captured well by our 1-D model. HONO emissions are calculated using a HONO / NO\textsubscript{x} = 0.008 emission ratio (Kurtenbach et al., 2001), gas-phase formation of HONO from reaction of NO with OH, dark conversion of NO\textsubscript{2} on aerosol and ground surfaces (with a yield of 50%; Wong et al., 2011), and HONO...
loss reactions which included HONO photolysis, reaction with OH, and uptake on aerosol and ground surfaces (with reactive uptake coefficients of $10^{-3}$ and $2 \times 10^{-5}$, respectively; Wong et al., 2011). It should be noted that the surface HONO formation from the dark conversion of NO$_2$ was negligible for the daytime runs. The comparison of HONO mixing ratios from the base model run and observations for 2 days of the experiment, 4 February and 27 January (Fig. 9, blue line), confirms that gas-phase chemistry alone cannot explain the observed daytime HONO levels near noontime, when the photolytic source of HONO is largest. To test whether the calculated HONO flux can explain this difference, a HONO surface flux term was implemented in the 1-D model as a ground surface release. This flux was determined from the calculated fluxes for the 2 days, scaled by a factor to account for the difference (caused by photolytic loss of HONO) between HONO fluxes at the surface and at the measurement height of 19 m a.g.l. To determine the appropriate factor, we compared the measured and modeled HONO flux at 19 m a.g.l. in initial test runs and then applied the factor for the final run. Figure 9e and f show that the modeled HONO flux at 19 m agrees well with the observations after this scaling step.

As expected, the model run with the HONO flux setup (Fig. 9, green line) produces more HONO than the basic model run (Fig. 9, blue line). Within the uncertainties of the observations, the model with additional HONO flux reproduces the observed HONO mixing ratios well on 27 January. The model also reproduces HONO well after noon on 4 February but overestimates HONO between 09:00 and 11:00 MST, when a plume of NO$_2$ was encountered. We will discuss this event in more detail below. The disagreement during the early morning of 4 February is likely due to challenges of modeling the early morning transition from a stable to a neutral boundary layer during a time of fast changing actinic fluxes. Despite these shortcomings, the comparison of modeled and observed HONO mixing ratios confirms the findings of the PSS analysis that the observed vertical HONO fluxes explain the missing daytime HONO source in this environment.

4 Discussion

The vertical HONO concentration profiles (Fig. 3) and the derived vertical fluxes (Fig. 6e) clearly show the presence of an important ground surface source of daytime HONO. The strong daytime variation of the sunny day HONO fluxes, with an average upward fluxes at solar noon of $1.5 \times 10^{10}$ molec. cm$^{-2}$ s$^{-1}$, implies a source that is driven by solar irradiance (Fig. 7). The precise mechanism of this HONO source, however, deserves further analysis.

Two previous studies of HONO fluxes measured with relaxed eddy accumulation systems reported analyses of daytime HONO surface formation. Zhou et al. (2011) measured HONO fluxes at 11 m above a forest canopy in Michigan during summer 2008. Their observed HONO flux correlated positively ($R^2 = 0.69$) with the product of photolysis rate constant of HNO$_3$ and leaf surface nitrate loading, suggesting that the photolysis of HNO$_3$ / nitrate on forest canopies is a significant daytime source of HONO to the lower atmosphere in low NO$_x$ rural environments. In contrast, Ren et al. (2011) measured the HONO flux 18 m a.g.l. above an agricultural field in Bakersfield, California, during the spring of 2010. Their daytime HONO flux showed a strong correlation ($R^2 = 0.99$) with the product of NO$_2$ concentrations and solar radiation instead. A strong correlation of the upwards HONO flux over agricultural fields with the product of NO$_2$ concentration with the NO$_2$ photolysis rate was recently also reported by Laufs et al. (2017).

We performed a similar analysis based on our observations. Because positive HONO fluxes point to formation on the ground (rather than in the gas phase or on aerosols), we used solar irradiance instead of photolysis rates for our analysis (see Wong et al., 2012, for further explanation of this approach). Only visible solar radiance was measured during UBWOS, which can be used to describe photo-enhanced NO$_2$ conversion; but because HNO$_3$ / nitrate species absorb in the UV range and UV solar irradiance was not measured in this study, its value was estimated by scaling visible solar irradiance with the measured $J_{\text{HONO}}$ and $J_{\text{NO}_2}$ (UVSolarRad = SolarRad x ($J_{\text{HONO}}$ / $J_{\text{NO}_2}$)). We used gas-phase NO$_2$ and HNO$_3$ mixing ratios as proxies for surface-
adsorbed \( \text{NO}_2 \) and \( \text{HNO}_3 \)/nitrate; i.e., we assumed that gas-phase and surface-adsorbed concentrations were in an equilibrium.

Comparison of our calculated HONO flux, with the product of \( \text{NO}_2 \) mixing ratios and visible solar irradiance, and the product of gas-phase \( \text{HNO}_3 \) mixing ratios and UV solar irradiance (Fig. 10) shows that they have a similar variation pattern and generally follow the same diurnal variation trend as solar irradiance. Daytime HONO flux strongly correlates with both \( \text{[NO}_2\text{]} \times \text{ solar radiation} \) \( (R^2 = 0.58) \) (Fig. 11a) and \( \text{[HNO}_3\text{]} \times \text{ UV solar radiation} \) \( (R^2 = 0.66) \) (Fig. 11b). The correlation coefficient is slightly larger for the \( \text{[HNO}_3\text{]} \times \text{ UV solar radiation} \) vs. \( \text{[NO}_2\text{]} \times \text{ solar radiation} \) correlation between the two species, suggesting that the source of the plume emitted both of these trace gases. Further analysis shows that the plume had not aged significantly, as illustrated by the missing peak of \( \text{HNO}_3 \). In addition, aircraft measurements (made by NOAA’s Global Monitoring Division and not shown here) show CO levels consistent with levels observed in coal-fired power plants. Hence we concluded that the plume originated at the Bonanza power plant (Lyman and Shorthill, 2013).

We observed a peak of HONO mixing ratios and HONO flux (Fig. 9c, e) that coincided with the plume of \( \text{NO}_2 \) around 10:00 (Fig. 9a). The correlation of HONO mixing ratios, HONO flux, as well as \( \text{NO}_2 \) mixing ratios, and the absence of a gas-phase \( \text{HNO}_3 \) peak at this time, suggests that the source of the HONO peak was from the ground and that it involved \( \text{NO}_2 \) molecules.

To further confirm this interpretation we used the RCAT model to simulate this specific event. The model was constrained by meteorological and radiative observations and initialized with the trace gas observations as described in Sect. 2.3.1 (Fig. 9a). The model also included HONO formation from photolytic conversion of \( \text{NO}_2 \) on the ground, which was parameterized by a solar-irradiance-dependent reactive uptake coefficient. The solar irradiance was described by a cubic dependence on \( J_{\text{NO}_2} \) (Wong et al., 2013), and the nighttime reactive uptake coefficient was adjusted to a somewhat smaller value \( (1.4 \times 10^{-5}) \) than the one used in Wong et al. (2013), as the value used in their study led to an overprediction of HONO. A lower coefficient can likely be explained by the very different environments between our study, i.e., low soil temperature (in this study) compared to high temperatures in the urban environment of Houston (Wong et al., 2013).

Taking into account the hourly results of our model, the model simulations driven by the observed fluxes as well as the photo-enhanced \( \text{NO}_2 \) conversion capture the HONO mixing ratio peak between 10:00 and 11:00 MST. The agreement between the prescribed and model flux also supports the conclusion that photo-enhanced \( \text{NO}_2 \) conversion on the surface explains HONO formation in this plume.

We then applied the same model to the observations on 27 January 2012, a day characterized by average wind speeds and \( \text{NO}_2 \) mixing ratios around 2 ppb (Fig. 9). We again adapted the emissions to simulate the general \( \text{NO}_2 \) mixing ratios. The model overpredicts \( \text{NO}_2 \) in the morning but captures the observations well after 11:00 MST. Despite the high \( \text{NO}_2 \), the base run (blue) and photo-enhanced run (red) underestimate measured HONO mixing ratios (black), suggesting that on days of moderate \( \text{NO}_2 \) levels, photo-enhanced conversion of \( \text{NO}_2 \) does not seem to play an important role in HONO production. The strong correlation between HONO flux and \( \text{[HNO}_3\text{]} \times \text{ UV solar radiation} \) suggests that photolysis of surface \( \text{HNO}_3 \)/nitrate might explain the observed HONO flux. However, without further information, such as concentration of surface-adsorbed \( \text{HNO}_3 \)/nitrate, it is not possible to unequivocally determine whether this mechanism was the source for the vertical HONO flux on 27 January.

An alternative HONO formation mechanism is the daytime displacement of HONO by strong acids. VandenBoer et al. (2015) showed evidence of uptake of HONO on soil surface at night and subsequent HONO displacement from soil by deposition of strong acids, such as \( \text{HNO}_3 \) and \( \text{HCl} \), during the day. They argue that HONO formed through the acid displacement mechanism is proportional to the product of solar radiation and \( \text{NO}_2 \) concentration, as production of \( \text{HNO}_3 \) is proportional to the concentration of \( \text{NO}_2 \) and \( \text{OH} \), the latter of which is proportional to solar radiation. For this mechanism to contribute significantly to the next day daytime HONO surface flux, it requires a substantial deposition of HONO during the previous night. To determine whether this mechanism played an important role in UBWOS 2012, we calculated the total number of HONO molecules deposited during the night of 27 January using the nighttime model results and compared it to the total number of HONO molecules released by the daytime HONO formation.
surface flux. The integrated surface HONO concentration from 19:00 of the previous night to 07:00 MST was about $1.84 \times 10^{14}$ molec. cm$^{-2}$. This number is lower than the integrated daytime HONO flux, calculated from the net HONO flux from 07:00 to 17:00 MST, of $2.84 \times 10^{14}$ molec. cm$^{-2}$. If every HONO molecule deposited at night would be re-released, this mechanism would be able to explain up to 65% of the vertical flux. However, in this case one would expect that, as the surface reservoir is depleted throughout the day, the flux would also decrease, which we did not observe. Nevertheless, we cannot rule out that the re-release of surface deposited HONO contributes to the overall vertical HONO flux during the day.

Another potential HONO formation mechanism is its formation from soil nitrite (Su et al., 2011). Su et al. (2011) showed that gas-phase HONO released by soil depended strongly on the concentration of nitrite in soil and its pH. They proposed that biological nitrification and denitrification processes are the main sources of nitrite ions in the soil, and showed that fertilized soils with low pH appeared to be particularly strong sources of HONO. We believe that emission of HONO from soil nitrite played a minor role in UBWOS 2012, as the soil was alkaline and cold during this field study. Measurements by NOAA measured a pH of 8.06 for a soil sample collected near the Horse Pool site in 2014 (Jonathan Raff, personal communication, 18 August 2015) and a report by Utah State University Extension stated that the pH of a large portion of soils in Utah ranged between 7.8 and 8.2 (https://extension.usu.edu/files/publications/publication/AG-SO-07.pdf). Like most biological reactions, nitrification and denitrification processes are strongly influenced by soil temperatures, with these processes ceasing at soil temperatures below 5°C (Western Plant Health Association, 2002). The hourly average soil temperature at noon in 2014 was about $-4°C$ (measured at 5 cm below the surface); therefore we can assume that soil temperatures in 2012, which were not directly measured, were not as cold as in 2014 but, nevertheless, cold enough to render nitrite production from biological reactions negligible.

Comparison of the missing source of HONO and HONO surface flux rate (Fig. 8) shows that HONO surface flux rate (magenta line) accounts for a significant portion of the $P_{\text{unknown}}$ (63 ± 32%) throughout the day (from 09:00 to 15:00). Furthermore, near noontime (from 11:00 to 13:00), surface flux rate accounts for an average of 90 ± 30% of $P_{\text{unknown}}$, suggesting that photolytic surface sources of HONO, and not HONO formation pathways in the gas phase or on aerosols, were the dominant sources of the unknown HONO in 2012.

Under high NO$_x$ events, our HONO flux analysis favors HONO formation mechanisms that are related to solar radiation and NO$_2$ concentrations, such as photo-enhanced conversion of NO$_2$ and nighttime uptake of HONO followed by daytime acid displacement. Under moderate or low NO$_2$ level conditions, photolysis of HNO$_3$ / nitrate seems to be the main HONO source. The $P_{\text{unknown}}$ peak at 08:00–09:00 MST is probably the result of rapid photolysis of HONO that accumulated in the nocturnal boundary layer and ineffective HONO production (through NO + OH) in the morning. Nitrophenols were not measured during UBWOS 2012, but they were measured in 2014 in UB during the winter (Yuan et al., 2016). Yuan et al. (2016) determined that the main source of nitrophenols in UB in 2014 was the oxidation of aromatics and that primary emissions of nitrophenols were negligible. Even though aromatics such as benzene were quite large in 2012 (median benzene in UB 2012 was 0.72 ppbv), photochemistry in 2012 was much less active compared to 2014, “suggesting that secondary formation of nitrophenols would be a relatively small source in 2012 and that the subsequent photolysis of nitrophenols would not be an important source of HONO” (Yuan et al., 2016).

5 Conclusions

HONO vertical concentration profiles and fluxes were measured during the winter of 2012 in the UB, Utah, using UCLA’s LP-DOAS instrument. The UB is highly impacted by oil and gas activities and thus exhibits NO$_2$ mixing ratios similar to those of polluted urban and suburban areas. Snow was almost completely absent in 2012, thus allowing the study of HONO chemistry over cold natural soil.

HONO mixing ratios near the surface ranged from below detection limits to 270 ± 20 ppt, with a campaign average of 74 ppt. These levels are lower than HONO measured in polluted urban areas but comparable to levels measured in rural environments. The sunny day hourly average HONO fluxes generally followed the diurnal variation of solar irradiance, with a noontime maximum of $(1.7 \pm 0.3) \times 10^{10}$ molec. cm$^{-2}$ s$^{-1}$. The upward fluxes supported the idea that HONO is formed by a photolytically driven surface source. A PSS and model HONO budget calculation clearly show that this surface source closes the daytime HONO budget; i.e., it is responsible for the commonly identified missing daytime HONO source.

The determination of the precise chemical mechanisms responsible for the surface flux is challenging due to the dominating correlation with solar irradiance of most of the mechanisms. However, our results seem to indicate that both photolysis of surface HNO$_3$/nitrate and photo-enhanced NO$_2$ conversion are likely active. Acid displacement of HONO deposited during the night, as proposed by VandenBoer et al. (2014), can contribute to the observed HONO flux, but a calculation of the total deposited HONO throughout the night with the integrated daytime flux reveals that this mechanism by itself cannot explain the observed daytime HONO formation.

The low temperatures and alkaline soil make the bacterial production of nitrite and its release (Su et al., 2011) likely unimportant.
Our results illustrate the significance of heterogeneous chemistry on the ground in converting adsorbed reactive nitrogen species into HONO, thus impacting the OH radical and nitrogen budget in the surface layer. Our analysis points to the difficulty of clearly identifying the precise chemical reaction mechanism forming HONO due to their strong correlation with solar irradiance and the cross correlation of the involved nitrogen species. More focused experiments that include a better characterization of the chemical composition of the surface are needed to further investigate HONO chemistry.

Data availability. Data available are at https://esrl.noaa.gov/csd/groups/csd7/measurements/2012ubwos/Ground/DataDownload/ (NOAA, 2012).

Competing interests. The authors declare that they have no conflict of interest.

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