Strong wintertime ozone events in the Upper Green River basin, Wyoming


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Abstract. During recent years, elevated ozone (O₃) values have been observed repeatedly in the Upper Green River basin (UGRB), Wyoming, during wintertime. This paper presents an analysis of high ozone days in late winter 2011 (1 h average up to 166 ppbv – parts per billion by volume). Intensive operational periods (IOPs) of ambient monitoring were performed, which included comprehensive surface and boundary layer measurements. On IOP days, maximum O₃ values are restricted to a very shallow surface layer. Low wind speeds in combination with low mixing layer heights (∼50 m above ground level around noontime) are essential for accumulation of pollutants within the UGRB. Air masses contain substantial amounts of reactive nitrogen (NOₓ) and non-methane hydrocarbons (NMHC) emitted from fossil fuel exploration activities in the Pinedale Anticline. On IOP days particularly in the morning hours, reactive nitrogen (up to 69 %), aromatics and alkanes (∼10–15 %; mostly ethane and propane) are major contributors to the hydroxyl (OH) reactivity. Measurements at the Boulder monitoring site during these time periods under SW wind flow conditions show the lowest NMHC/NOₓ ratios (∼50), reflecting a relatively low reactive NMHC mixture, and a change from a NOₓ-limited regime towards a NMHC-limited regime as indicated by photochemical indicators, e.g., O₃/NOₓ, O₃/NO₂, and O₃/HNO₃ and the EOR (extent of reaction). OH production on IOP days is mainly due to nitrous acid (HONO). On a 24 h basis and as determined for a measurement height of 1.80 m above the surface HONO photolysis on IOP days can contribute ∼83% to OH production on average, followed by alkene ozonolysis (∼9%). Photolysis by ozone and HCHO photolysis contribute about 4% each to hydroxyl formation. High HONO levels (maximum hourly median on IOP days: 1096 pptv – parts per trillion by volume) are favored by a combination of shallow boundary layer conditions and enhanced photolysis rates due to the high albedo of the snow surface. HONO is most likely formed through (i) abundant nitric acid (HNO₃) produced in atmospheric oxidation of NOₓ, deposited onto the snow surface and undergoing photo-enhanced heterogeneous conversion to HONO (estimated HONO production: 10.2 ± 40 ppbv h⁻¹) and (ii) combustion-related emission of HONO (estimated HONO production: ∼0.1 ± 30 ppbv h⁻¹). HONO production is confined to the lowermost 10 m of the boundary layer. HONO, serves as the most important precursor for OH, strongly enhanced due to the high albedo of the snow cover (HONO photolysis rate 10.7 ± 30 ppbv h⁻¹). OH radicals will oxidize NMHCs, mostly aromatics (toluene, xylenes) and alkanes (ethane, propane), eventually leading to an increase in ozone.
1 Introduction

The Upper Green River basin (UGRB) has one of the largest natural gas reserves of the United States (US). In 2009, the proven gas reserves for the Jonah field (11.1 billion m$^{-3}$) ranked seventh and for the Pinedale Anticline field (13.8 billion m$^{-3}$) ranked third among the top 100 natural gas fields in the US (EIA, 2009). Oil and gas extraction including drill rigs, production equipment and compressor stations are operating continuously and represent the only significant emission source in the UGRB with overall emissions of 9.9 t day$^{-1}$ of reactive nitrogen (NO$_x$) and 41.7 t day$^{-1}$ of volatile organic compounds (VOC) (WDEQ, 2011).

The UGRB is a high plateau located about 2000 m a.s.l. (above sea level). It is surrounded by mountain ranges, which reach heights of up to 3500 m a.s.l. to the west (Wyoming Peak) and 4200 m a.s.l. to the northeast (Gannett Peak). Winters are usually cold and frequently associated with snow cover. During recent years, elevated hourly ozone values above 150 ppbv (parts per billion by volume) have been observed in the UGRB during wintertime (Schnell et al., 2009; Carter and Seinfeld, 2012). As of July 2012, the U.S. Environmental Protection Agency (EPA) declared the UGRB as a non-attainment area for the 2008 ground-level 8 h ozone standard, which is 75 ppbv. Recent publications have focused on some observational findings in the UGRB in the year 2008 (Schnell et al., 2009) or sensitivity analyses using a box model approach together with VOC (volatile organic compound) incremental reactivities for selected ozone episodes in 2008 and 2011 (Carter and Seinfeld, 2012). Still, there are major uncertainties in our understanding of the occurrence of high ozone levels in the UGRB under wintertime conditions. Apart from specific meteorological conditions for the UGRB (i.e., low mixing layer heights, light winds, extensive snow cover, at times recirculation of air masses), these include processes in the nitrogen oxide (NO$_x$) and VOC cycles, such as the role of nitric acid (HNO$_3$) and the radical precursors such as formaldehyde (HCHO) and nitrous acid (HONO). In particular, high daytime HONO levels were found (Rappenglück, 2010 and 2011). Although HCHO levels were moderate, the sources and role of HCHO in the UGRB is not fully understood, particularly with regard to the overall relatively low alkene reactivity as shown by Carter and Seinfeld (2012). A better quantification of these hydroxyl (OH) sources is needed to improve the description of ozone chemistry in the UGRB, which is required to develop efficient strategies to reduce pollution in that area.

In this paper we analyze high ozone days in late winter 2011 (1 h average up to 166 ppbv) observed in the area of the Boulder station and describe the meteorological and chemical processes leading to these extreme events using the comprehensive surface and boundary layer measurements collected during the Upper Green Winter Ozone Study (UGWOS) 2011 (MSI, 2011).

2 Methods

Surface air quality data used in this paper were collected continuously at the Boulder site and Boulder South Road site from January to March 2011 (for Boulder South Road site data see also Field et al., 2011). Boundary layer measurements including radiosonde and ozonesonde launches were performed at the Boulder site during intensive operational periods (IOPs). Data from the tethered balloon were obtained at the “Tethered Balloon” site. Supplement S1 lists the details of the instrumentation and Fig. 1 shows the location of these sites in the UGRB including the locations of oil and gas wells.

Surface measurements included routine measurements for ozone ($O_3$), reactive nitrogen compounds (NO/NO$_2$/NO$_x$), total non-methane hydrocarbon (NMHC), methane (CH$_4$) and trace level measurements for nitrogen monoxide (NO), nitrogen dioxide (NO$_2$), and total reactive nitrogen (NO$_x$). Additional measurements included HNO$_3$, HONO, HCHO, carbon monoxide (CO) and online speciated NMHC. If not otherwise indicated the term NMHC denotes total, i.e., non-speciated NMHC measured at the Boulder or Tethered Balloon site.

At all sites, basic meteorological measurements were made. Additional details beyond the information provided in Supplement S1 can be found in MSI, 2011. Here we briefly describe the methodology for the HONO measurements. For the UGWOS 2011 study a commercially available LOPAP (Long Path Absorption Photometry) instrument was used (QUA Elektronik & Analytik GmbH, Wuppertal, Germany). The LOPAP is described thoroughly in Heland et al. (2001) and Kleffmann et al. (2002). It is a wet-chemical in situ instrument, which consists of an external sampling unit where ambient gaseous HONO is directly sampled in a stripping coil using a mixture of sulfanilamide in hydrochloric acid. No sampling lines are used thus minimizing sampling artifacts on surfaces. The stripping reagent is transferred through an insulated transfer line (length: 3 m; outer diameter 5 cm; kept at 20°C) to the instrument where it is converted to an azo dye by the reaction with N-naphthylethylendiamine-dihydrochloride. The absorption, more precisely the logarithm of the ratio between the spectral intensity at 650 and 550 nm, is measured in long-path absorption tubes made of Teflon AF2400 using a two-channel minispectrometer (Ocean Optics S2000). In the external sampling unit two stripping coils are used in series. In the first channel HONO as well as possible interferences are determined, while in the second channel only the interferences are quantified. The difference of these two channels yields the HONO signal.

For routine zeroing ultra-high purity nitrogen (UHP N2) was applied every 8 h for 30 min directly to the inlet of the external sampling unit by a 1/32″ PFA (perfluoroalkoxy) tubing which was partially inserted into the tip of the stripping coil. A linear (or polynomial) fit was calculated for the stable
Figure 1. The monitoring sites Boulder, Boulder South Road, Tethered Balloon, Tall Tower and other WDEQ (Wyoming Department of Environmental Quality) monitoring sites relative to the locations of oil- and gas-well locations operating during the period January–March 2011.

reading and used as the zero baseline. Calibrations were always preceded by zero-measurements. For calibration itself the stripping solution was replaced by 0.01 mg NO$_2$ per liter of stripping solution, while UHP N2 was flowing into the inlet of the external sampling unit. Final calibration values were calculated using the calibration standard concentration and the measured gas- and liquid-flow rates.

The sampling time is 30 s. The response time, i.e., the time it takes for the signal to go from 100 to 10 % of the initial value or from 0 to 90 % of the final value, changes every time a new set of peristaltic pump tubing is installed, yet is
Table for the weather and ozone levels during the study period

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Figure 2. Time series for ozone and temperature based on hourly data at the Boulder site for January–March 2011.

Figure 3a. Mean diurnal variation of selected trace gases for IOP and non-IOP days.

3 Results and discussion

3.1 General observations

The Upper Green River basin had continuous snow cover throughout the winter months January–March 2011. Figure 2 shows that ambient temperature was below freezing most of the time, ranging from −28.0 °C in early February to +5.1°C in mid-March. Also displayed in the same figure are ambient ozone observations. A slight increase in background ozone from 45 ppbv in January to about 50 ppbv in March is discernible. Apart from this longer-term variation, it can clearly be seen that at times significant day-to-day variations in the ozone mixing ratios may occur. While decreases of ozone are reflecting NO titration effects, strong increases of ozone well above the ozone background level are likely due to a combination of dynamic and photochemical processes. During the time frames 28 February–2 March and 9–12 March 2011, the highest hourly ozone readings were observed with up to 166 ppbv at the Boulder surface site on 2 March. IOPs were performed during these two periods (IOP#1: 28/2–2/3; IOP#2: 9/3–12/3), which included additional information about the vertical distribution of meteorological parameters as well as some selected trace gases. The following discussions will focus on the time period 28 February–16 March 2011. This time period includes the two IOPs and has the most complete data availability with regard to continuous as well as discrete measurements. The non-IOP days during this time frame will be used as reference days. Figure 3a and b display mean diurnal variations of O₃, NO, NOₓ, NMHC, CH₄, NO₂, HNO₃, HONO, and HCHO split into IOP and non-IOP days. Most species show 2–3 times higher mixing ratios on IOP days compared with non-IOP days throughout the day. Ozone shows this enhancement during the afternoon and early evening only. Primarily emitted
species like NO show maximum values during the time frame 09:00–12:00 MST (Mountain Standard Time). Species that can also be formed secondarily (e.g., NO\textsubscript{y}, HNO\textsubscript{3}, HONO, and HCHO) exhibit enhanced levels during sunlit daytime hours from about 07:00 to 18:00 MST, in particular on IOP days. Ozone mixing ratios start to increase by 09:00 MST, reach maximum levels around 15:00 MST and remain at higher levels until early evening. Quite surprisingly, HONO mixing ratios are high during the daytime on non-IOP days and even higher on IOP days (maximum median around 275 and 186 %, respectively). HONO and HCHO are also enhanced (98 and 64 %, respectively). Overall, on IOP days trace gases relevant to O\textsubscript{3} formation are all significantly enhanced throughout the day compared with non-IOP days.

The only major anthropogenic emissions in the remotely located UGRB are associated with oil and gas extraction. As a benchmark for this study we compare some results to a highly polluted urban area. Luke et al. (2010) report results of reactive nitrogen compounds for Houston, Texas, a city exposed to complex emissions including emissions from large petrochemical sources (e.g., Parrish et al., 2009; Lefer and Rappenglück, 2010; Olaguer et al., 2014). An investigation of the same daytime periods shows that, while NO\textsubscript{y} is significantly higher in the urban air of Houston (about 2–4 times higher when compared to IOP days at Boulder), the picture is different for NO\textsubscript{3}. Although NO\textsubscript{3} is higher in Houston in the morning and during the night (275 and 186 %, respectively), it is lower in Houston than in Boulder during daytime on IOP days. These differences are largely due to HONO and HNO\textsubscript{3}. While HONO is slightly lower in the morning and during the night, it is about three times higher during the daytime on IOP days. HNO\textsubscript{3} however is higher throughout the day on IOP days in Boulder compared to the Houston case. On non-IOP days at Boulder, HONO and HNO\textsubscript{3} levels are mostly lower than in Houston. The HNO\textsubscript{3} fraction of NO\textsubscript{y} is around 18 % during daytime on non-IOP days, while it is around 22 % during daytime on IOP days (Luke et al., 2010, report an overall daytime HNO\textsubscript{3} fraction of 15.7 % of NO\textsubscript{y} in Houston). For HONO the corresponding numbers are around 2 (non-IOP days) and 4.5 % (IOP days), while Luke et al. report an overall daytime HONO fraction of 1.7 % of NO\textsubscript{y}.

Luke et al. (2010) also report temporally highly resolved data for particulate nitrate (p-NO\textsubscript{3}\textsuperscript{−}). Median values were 0.234 (05:00–09:00 CST – Central Standard Time), 0.091 (11:00–17:00 CST), and 0.135 ppbv for nighttime (21:00–05:00 CST). At Boulder, 24 h samples were collected. Although not exactly comparable, it may provide us with some estimate. On IOP days, the median value for p-NO\textsubscript{3}\textsuperscript{−} was equivalent to 0.58 ppbv (maximum equivalent to 1.54 ppbv), while on non-IOP days the corresponding values were equivalent to 0.47 and 0.74 ppbv, respectively, which indicates that p-NO\textsubscript{3}\textsuperscript{−} may be higher at the Boulder site than in Houston.

Supplement S3 shows a comparison of NO\textsubscript{y} measurements versus individual NO\textsubscript{y} compounds for the Boulder site. Particulate data was only collected on a 24 h basis. Therefore this data set only comprises a small number of
observations. The deviation from the 1:1 line is within the accuracy of the NO\textsubscript{x} and the combined individual NO\textsubscript{x} measurements (see Supplement S1) for ranges above 20 ppbv. For NO\textsubscript{x} values below 20 ppbv, NO\textsubscript{x} tends to be larger than the sum of the individually measured NO\textsubscript{x} components. According to Supplement S3, still some fraction of NO\textsubscript{x} compounds may be missing, even when particulate NO\textsubscript{3} was included. Potential candidates for this NO\textsubscript{x} deficit include the nitrate radical (NO\textsubscript{3}), dinitrogen pentoxide (N\textsubscript{2}O\textsubscript{5}), nitryl chloride (ClNO\textsubscript{2}), peroxy acetyl nitrate (PAN) and alkyl nitrates. While NO\textsubscript{3}, N\textsubscript{2}O\textsubscript{5}, and ClNO\textsubscript{2} would be present at nighttime (e.g., McLaren et al., 2004; Edwards et al., 2013), PAN and alkyl nitrates are produced photochemically and tend to reach maximum values during daytime (e.g., Hayden et al., 2003; Sommariva et al., 2008, and references therein). No measurements of these species were performed at the Boulder site, however some assumptions can be made. Maximum NO\textsubscript{3} and N\textsubscript{2}O\textsubscript{5} levels typically range from about 50 and 300 pptv, respectively, in polluted continental air masses (McLaren et al., 2004) to about 150 and 500 pptv, respectively, in urban areas (Stutz et al., 2010b). NO\textsubscript{3} and N\textsubscript{2}O\textsubscript{5} can constitute 7–30 % of NO\textsubscript{y} (McLaren et al., 2004, and references therein). NO\textsubscript{3} is formed through the reaction of NO\textsubscript{2} with O\textsubscript{3}. It is likely that this reaction may be efficient at the Boulder site due to the observed appreciable ambient levels of NO\textsubscript{2} and O\textsubscript{3} (see Supplement S2 and Fig. 2). Loss mechanisms for NO\textsubscript{3} include reactions with alkynes and biogenic hydrocarbons, and reaction with NO\textsubscript{2} to form N\textsubscript{2}O\textsubscript{5}. As alkynes and biogenic hydrocarbons are found at low levels in the UGRB (Field et al., 2012a, 2012b), while abundant NO\textsubscript{2} is present, the latter reaction to form N\textsubscript{2}O\textsubscript{5} is likely the dominant removal process for NO\textsubscript{3}. In addition, low ambient temperatures would favor this process. The homogeneous and heterogeneous hydrolysis of N\textsubscript{2}O\textsubscript{5} may ultimately lead to appreciable amounts of gaseous HNO\textsubscript{3} and particulate nitrate according to McLaren et al., 2004. While no direct measurements of ClNO\textsubscript{2} were performed at the Boulder site, particulate chloride (Cl\textsuperscript{−}) data show low median levels of 0.015 µg m\textsuperscript{−3} (based on 6 24 h filter samples). This may indicate that Cl\textsuperscript{−} is not abundantly available to form ClNO\textsubscript{2} in appreciable amounts, and likely found at less than 600–800 pptv in the Uintah Basin, Utah (Edwards et al., 2013). Earlier studies in the UGRB also included PAN measurements (Environ, 2010). The results showed up to 1 ppbv PAN during daytime and up to 400 pptv as an estimated 24 h average, while corresponding ozone values were 85 and 51 ppbv, respectively, i.e., PAN was ~1 % of the ozone mixing ratios, which lies in the range observed in areas (e.g., Rappenglück et al., 2003). Most important precursor candidates for PAN in the UGRB will be ethane, toluene and xylenes. As high PAN values would also coincide with high NO\textsubscript{y} values, we assume that PAN would only account for a negligible amount of around 1 % of the NO\textsubscript{y} budget based on the results shown in Supplement S3. Alkyl nitrates may account for about 10 % of NO\textsubscript{y} (Sommariva et al., 2008, and references therein). As alkyl nitrates have an atmospheric lifetime of more than a week they may accumulate under favorable meteorological conditions and contribute to the NO\textsubscript{y} budget not only during the day, but also at night. We predominantly found lower values for NO\textsubscript{y} at night (see Fig. 3b), i.e., at a time when the missing NO\textsubscript{y} components NO\textsubscript{3}, N\textsubscript{2}O\textsubscript{5}, ClNO\textsubscript{2}, and also alkyl nitrates would contribute most to the NO\textsubscript{y} budget. This may explain the deficit found in the NO\textsubscript{y} budget at NO\textsubscript{y}-mixing ratios below 20 ppbv.

Corresponding analyses for NO\textsubscript{x} oxidation products NO\textsubscript{y} (NO\textsubscript{2} = NO\textsubscript{x} – NO\textsubscript{y}) versus the sum of individual NO\textsubscript{x} compounds NO\textsubscript{z}\textsubscript{i} (i.e., HNO\textsubscript{3}, HONO and p-NO\textsubscript{3}) based on a 24 h database yielded NO\textsubscript{y} [ppbv] = Σ NO\textsubscript{z}\textsubscript{i} [ppbv] · 1.09–0.38 ppbv (r\textsuperscript{2} = 0.87) for IOP days and NO\textsubscript{y} [ppbv] = Σ NO\textsubscript{z}\textsubscript{i} [ppbv] · 1.12 + 0.28 ppbv (r\textsuperscript{2} = 0.82) for non-IOP days, which is of similar magnitude as found in Houston (Luke et al., 2010, for instance. Luke et al., 2010 point out that higher nighttime NO\textsubscript{y}/NO\textsubscript{z}\textsubscript{i} ratios and the magnitude of NO\textsubscript{y}/NO\textsubscript{z}\textsubscript{i} differences may point to the presence of nighttime ClNO\textsubscript{2}. Based on our limited data we cannot make similar statements, but cannot rule out this possibility. Our measurements did not include PAN. Using our previous assumptions of about 400 pptv as an estimated 24 h average value, PAN could contribute up to 5 % of the NO\textsubscript{y} budget. The atmospheric lifetime of PAN (τ\textsubscript{pan}) is primarily determined by the following reactions:

\[ \text{CH}_3\text{C(O)OONO}_2 \rightarrow \text{CH}_3\text{C(O)OO} + \text{NO}_2, \]
\[ \text{CH}_3\text{C(O)OO} + \text{NO}_2 \rightarrow \text{CH}_3\text{C(O)OONO}_2, \]
\[ \text{CH}_3\text{C(O)OO} + \text{NO} \rightarrow \text{CH}_3\text{C(O)O} + \text{NO}_2. \]

The atmospheric lifetime of PAN thus critically depends on the NO\textsubscript{y}/NO ratio and also the ambient temperature, since the reaction constant k\textsubscript{1} is proportional to \exp(−1/T). Based on reactions (1–3), τ\textsubscript{pan} can be calculated according to Ridley et al. (1990):

\[ \tau_{\text{pan}} = \frac{1}{k_1} \left( 1 + \frac{k_2 [\text{NO}_2]}{k_3 [\text{NO}]} \right) \]

For the Boulder site, the shortest τ\textsubscript{pan} for IOP#1 was 10.6 days and for IOP#2 2.6 days and reached almost 1 day on the last day of the field study (March 16) when temperature reached + 5.1 °C. It is thus unlikely that PAN may serve as a NO\textsubscript{2} source and contribute to ozone formation under the environmental conditions found during the study. However, it can serve as a reservoir for NO\textsubscript{x} and radicals and remove NO\textsubscript{x} and radicals out of the UGRB.

### 3.2 Role of dynamic processes

#### 3.2.1 Dependence on wind direction

Supplement S4a and b display daytime and nighttime wind roses of O\textsubscript{3}, primarily emitted pollutants NO\textsubscript{x}, NMHC, CH\textsubscript{4}, and trace gases that may have both primary and secondary
sources, i.e., NO\textsubscript{y}, HNO\textsubscript{3}, HONO, and HCHO. Supplement S4a and b include both IOP and non-IOP days in order to obtain larger and more representative data sets. Regardless of IOP and non-IOP days, nighttime data will most likely be useful to point to potential emission sources as photochemical processes are at a minimum.

The daytime ozone wind rose clearly shows enhanced mixing ratios for SSW–WSW and SE–ESE directions (median 64–73 ppbv). During nighttime, maximum median ozone of about 60 ppbv occurs under NNE–ENE wind directions. These wind directions only have small amounts of NO\textsubscript{2} (median about 1.5 ppbv; not shown) and are likely more aged air masses. Primarily emitted NO\textsubscript{x}, NMHC, and CH\textsubscript{4} display pronounced enhancements during the daytime in the similar directions as for O\textsubscript{3}. During nighttime there are significant peaks under SW–W wind directions. While NO\textsubscript{x} and NMHC are distinctly enhanced (NO\textsubscript{x} ∼ 10 ppbv, NMHC ∼ 900 pptv), still CH\textsubscript{4} with about 4 ppm (parts per million) is a factor of 2 above its background levels. This hints to sources that emit nitrogen oxides, NMHC and CH\textsubscript{4} or different sources located in the same area, which may overlap. This is likely consistent with locations of compressors and drill rigs operating during the period January–March 2011 (which emit primarily NO\textsubscript{x}) relative to wellhead production equipment (which emits primarily CH\textsubscript{4} and NMHC).

NO\textsubscript{y}, HNO\textsubscript{3}, HONO, and HCHO largely follow the same directional pattern as NO\textsubscript{x} and hydrocarbons (Supplements S4a–b). However, contrary to the primarily emitted pollutants such as NO\textsubscript{x}, NMHC, and CH\textsubscript{4}, species that may be formed secondarily (i.e., HNO\textsubscript{3}, HONO, and HCHO) generally show higher values during the daytime. It is known that HCHO may be emitted primarily from incomplete combustion in either mobile or stationary sources (Zweidinger et al., 1988; Altschuller, 1993; Chen et al., 2004; Dasgupta et al., 2005; Rappenglück et al., 2013). Also, HONO can be emitted primarily from various combustion processes (Kirchstetter et al., 1996; Kurtenbach et al., 2001; Rappenglück et al., 2013) with traffic emissions being an important contribution to ambient HONO (Sarwar et al., 2008).

NO\textsubscript{y} behaves somewhat similarly to the primary pollutants, most likely due to the large fraction of NO\textsubscript{x} in NO\textsubscript{y} (see Supplement S2). Interestingly, HONO clearly shows an anisotropic dependence on wind direction, both during night- and daytime, which indicates a point source rather than an area source like the surface. During daytime, HONO mixing ratios are highest from WSW to SSW (medians of 477 and 545 pptv, respectively), while at night the maximum mixing ratios occur under a WSW wind direction (median 347 pptv). The nighttime HONO peak for WSW flow is in accordance with those for other trace gases and hints at a point source, the high daytime HONO levels (see also Supplement S2) are at odds with the conventional understanding of HONO diurnal variability, suggesting that additional processes other than point source emissions (e.g., Czader et al., 2012, and references therein) may be important.

Figure 4. HYSPLIT backward trajectories calculated for 50 m a.g.l. and for the following arrival times at Boulder: 07:00 MST (red), 12:00 MST (blue), 14:00 MST (light green), and 17:00 MST (dark green).

3.2.2 Backward trajectories

For further analysis, we calculated HYSPLIT (HYbrid Single-Particle Lagrangian Integrated Trajectory) backward trajectories (Draxler and Hess, 2013; Rolph, 2013) based on 12 km resolution meteorological data provided by NAM (North American Mesoscale model). Uncertainties stated by Draxler and Hess are in the range of 15–30 % of the travel distance. As an additional test, we calculated back and forward trajectories within the UGRB for the times of interest and found only negligible difference among those. Although, there might be some limitations associated with the accuracy of backward trajectories, they at least may provide information about major regimes. The results of trajectory calculations are shown in Fig. 4 and they indicate three major regimes during the 28 February–3 March 2011 period. Non-IOP days mostly showed consistent air mass flows out of the NW wind sector (example shown: 3 March 2011). Also, they
are associated with higher wind speeds and enhanced atmospheric dispersion. This coincides with the generally low to modest levels of pollutants (Supplement S4a, b). The days with the highest ozone levels during each IOP may show recirculation processes during the day (as for 3 March 2011); however, they are usually characterized by the fact that at least once during the day, trajectories eventually pass through the area southwest of Boulder before arriving at Boulder. According to Fig. 1, air masses would then pass the oil- and gas-well locations of the Pinedale Anticline and also a large number of compressor facilities (Fig. 5). According to the trajectories on 12 March, air masses would have passed this sector at the most 1–2 h before arrival at Boulder and would have stayed in that region for about 1 h. As seen in Supplement S4a and b, air masses coming from the SW quadrant carry maximum amounts of both primary and secondary pollutants. As shown later, they are also associated with the highest VOC reactivity and a change of NO$_x$ sensitivity towards a VOC sensitivity regime.

### 3.2.3 Boundary layer height

Tethersonde data taken on IOP days indicate very shallow boundary layer heights. Based on the data in S5 and using approaches by Stull (1988), average morning boundary layer heights (09:00–11:00 MST) would be estimated to be about 35 m a.g.l., around 50 m a.g.l. around noon (11:00–13:00 MST), reach a daily maximum around 80 m a.g.l. in the early afternoon (13:00–15:00 MST), and decrease to around 45 m a.g.l. in the late afternoon (15:00–17:00 MST). This is largely in agreement with SODAR (sonic detection and ranging) data (not shown). Figure 6 shows vertical profiles of ozone and wind direction on IOP days. Generally, IOP days were characterized by W–NW flows above 600 m a.g.l. and appreciable temporal and spatial changes in wind direction below this level. These lower level changes are also reflected in the backward trajectories previously shown in Fig. 4. Below 600 m a.g.l., wind speed was generally less than 5 m s$^{-1}$. Above 600 m a.g.l., wind speed rapidly increased above 10 m s$^{-1}$ (not shown). The strong changes in atmospheric flow around 600 m a.g.l. certainly helps to maintain inversions and stable boundary layer conditions beneath this layer throughout IOP days. Since surrounding mountain ranges of the UGRB reach heights of up to 3500 m a.s.l. to the west (Wyoming Peak) and 4200 m a.s.l. to the northeast (Gannett Peak), the boundary layer air masses are easily trapped within the basin and recirculation processes as shown for 2 March in Fig. 4 are likely. This is in accordance with earlier wintertime meteorological studies in similar areas of the Rocky Mountains (Yu and Pielke, 1986).

Figure 6 shows that ozone exhibits a distinct behavior associated with the atmospheric dynamic pattern: above 600 m a.g.l., ozone mixing levels are quite stable around 60 ppbv regardless of the day and time of day; it represents background ozone levels in the well-mixed lower troposphere. Below 600 m a.g.l., ozone mixing ratios clearly show diurnal changes with lower than background values in the early morning and strongly enhanced values in the afternoon. The maximum deviations are clearly restricted to a very shallow surface layer below 70 m a.g.l. There is no evidence of ozone carry over from previous days. Ozone deposition over dry surfaces is about 0.4 cm s$^{-1}$ (Hauglustaine et al., 1994); however, it is about 0.07 cm s$^{-1}$ over snow surfaces (Hauglustaine et al., 1994); i.e., ozone removal through turbulent diffusion is significantly reduced. Recent studies over polar snow indicate even lower deposition velocities in the range of 0.01–0.07 cm s$^{-1}$ (Helming et al., 2009). As deposition may play a minor role during snow covered periods in the UGRB, the ozone deviation from the background ozone level is instead determined by removal (e.g., nighttime titration) and daytime in situ formation processes.

As an example, Fig. 7 shows selected data obtained by the tethersonde system for the IOP day, 2 March 2011. The results for $\Delta T$ (Delta-T in the plot) indicate well-defined inversion and stable conditions in levels with $\Delta T > 0 ^\circ C$. Again, it can be seen that highest ozone levels are restricted to the surface levels. Precursors such as NO$_2$ and NMHC may exhibit a layered distribution. The highest levels may occur close to the mixing layer height as defined by $\Delta T$. This hints to pollution plumes that may have originated from either elevated sources and/or sources, which show some plume rise due to higher than ambient temperatures. Potential ozone O$_x$, as expressed by the sum of O$_3$ and NO$_2$ provides information about the potential presence of O$_3$ formation processes. Using the tethersonde data and segregating it into time frames like in Supplement S5, we obtain S6. This figure shows two main features: (i) formation of O$_3$ from early morning to the afternoon throughout the surface layer, and (ii) apart from the late afternoon, a tendency towards higher O$_3$ values in higher layers of the surface layers. The highest O$_3$ mixing ratios at the surface occur only in the late afternoon.

In an attempt to investigate other trace gases for which no vertically resolved data is available, we segregated surface data into 5 m bins of hourly SODAR mixing layer height (MLH) data and split it into day- and nighttime observations. Results are shown in Supplement S7. For ozone, daytime data is clearly higher than nighttime data regardless of the MLH. However, during daytime, maximum ozone levels are observed under lowest MLH. Speciated NMHC data obtained at the Boulder South Road site generally show enhanced values under MLH < 40 m a.g.l. and decreasing concentrations with increasing MLHs regardless of the time of day. However, while higher nighttime concentrations of speciated NMHC classes are found at lower MLHs, during daytime, the lowest MLHs are associated with lower concentrations of NMHC classes. Assuming NMHC sources that do not change emission strength during the day, this points to photochemical degradation of NMHCs in the lowermost surface layers. The change in the median values is about $-20\%$ for alkanes, $-45\%$ for alkenes, and almost $-50\%$ for
aromatics. CO does not change significantly with MLH and time of day. NO\textsubscript{x} shows significantly higher values during nighttime and, contrary to NMHC classes, there is no change at the lowermost MLHs when comparing daytime with nighttime observations. However, nighttime data clearly shows decreasing NO\textsubscript{x} levels with increasing MLHs. Both HCHO and HONO show higher values during the daytime than nighttime regardless of MLHs. In accordance with other primary pollutants like NMHC and NO\textsubscript{x}, HCHO and HONO exhibit a trend towards lower mixing ratios with increasing MLHs during nighttime.

In general, results from IOP days, such as 2 and 12 March, indicate that slightly variable wind directions (including recirculation) and low wind speeds in combination with low boundary layer heights are essential for accumulation of both primary and secondary pollutants. The occurrence of low boundary layer heights in presence of snow cover in mountainous regions is in accordance with previous model simulations (Bader and McKee, 1985). These studies also showed that these conditions can hold a stable layer until very late in the day.
3.3 Role of emissions and chemistry

3.3.1 Potential combustion emissions for radical precursors HCHO and HONO

Primary sources for HCHO and HONO are important sources for the hydroxyl radical (e.g., Ren et al., 2013; Czader et al., 2013). While HCHO may have a variety of primary sources, its secondary formation during daytime usually exceeds primary HCHO emissions significantly (e.g., Rappenglück et al., 2010, and references therein). This is also reflected in the Boulder site data when comparing daytime HCHO data on IOP vs. non-IOP days (see Supplement S2 and Fig. 3b). However, secondarily produced HCHO does not provide a net OH source as the formation of OH already consumed OH.

Unfortunately, there were no CO measurements taken at the Boulder site, but linear regression analysis of CO vs. NO_x calculated for the Boulder South Road site indicate CO [ppbv] = 6.14 × NO_x [ppbv] + 140 [ppbv] \( (r^2 = 0.55) \) over all wind directions for the entire time period 28 February–3 March 2011. The slope is very close to values obtained for urban traffic rush hour, which is 6.01 ± 0.15 ppbv CO/1 ppbv NO_x (Rappenglück et al., 2013). While the correlation coefficient is significantly weaker than that found in the Rappenglück et al. (2013) study \( (r^2 = 0.91) \), which may be due to higher degree of mixing of different air masses, it indicates the presence of combustion sources; under remote unpolluted conditions CO and NO_x would be almost unrelated (see e.g., Villena et al., 2011).

As previously mentioned, compressors and drill rigs were operating during the period January–March 2011 in the area located upwind of the Boulder site under SW flow conditions. These units emit primarily NO_x relative to wellhead production equipment, which primarily emits CH_4 and NMHC. As indicated in Supplement S4a and b, enhanced levels of primary pollutants occur under SSW–W flows during the daytime and under SW–W flows during the night. In an attempt to identify potential emission sources for HCHO and HONO, we performed correlation analyses with various other trace gases measured at the Boulder site focusing on nighttime data, to exclude daytime photochemical processes and SW–W wind directions, which showed peak values for all species in accordance with Supplement S4a and b.

NO_y/NO_x ratios were around 0.35 under these wind flow conditions and thus, significantly less than 0.6, indicating freshly emitted pollutants.

Supplement S8 clearly shows the close relationship between HONO and NO_2, NO_x, and HNO_3 as observed at the Boulder site. The good, although somewhat weaker, correlation of HONO with total NMHC at the Boulder site is most
Figure 7. Results from tethersonde measurements of ozone, NMHC, NO\textsubscript{2}, and $\Delta T$ (Delta-T) on 2 March 2011. $\Delta T$ is defined as the temperature at a given level minus the temperature at the surface. Mixing layer heights as determined by SODAR data are indicated as “x” in the $\Delta T$ plot.

likely due to the fact that emission sources that emit NO\textsubscript{x} may be located close to NMHC sources. HCHO shows overall weaker correlations with NO\textsubscript{x}, total NMHC, and CH\textsubscript{4} compared with HONO. HCHO has a longer atmospheric lifetime than HONO ($\sim 6–7$ h for HCHO vs. $\sim 5–7$ min for HONO at local noontime conditions during the UGRB study) and thus some appreciable background, which may also include remnants of the previous day’s formation of HCHO. This background may get mixed into freshly emitted plumes and cause higher data scatter. In any case, according to Supplement S8, HCHO shows stronger correlation with total NMHC and CH\textsubscript{4} than with NO\textsubscript{x}, while for HONO it is the opposite behavior. This hints to different source categories for HONO and HCHO.

Based on the CO/NO\textsubscript{x} observations at the Boulder South Road site, we assume that NO\textsubscript{x} at the Boulder site would also be primarily related to combustion sources. In a recent traffic-related emission study, measurements of HONO versus NO\textsubscript{x} yielded a slope of $15.86 \pm 0.82$ pptv HONO/1 ppbv NO\textsubscript{x} ($r^2 = 0.75$) and a slope of $25.00 \pm 1.06$ pptv HCHO/1 ppbv NO\textsubscript{x} ($r^2 = 0.80$) for HCHO versus NO\textsubscript{x} (Rappenglück et al., 2013). The slopes of HONO versus NO\textsubscript{x} for the Boulder nighttime data under wind direction 180–270\textdegree~and the Houston traffic-related measurements are almost identical. In the traffic emissions study, Rappenglück et al. argued that the high HONO/NO\textsubscript{x} emission ratios were likely due to heavy duty vehicles. In the case of the Upper Green River basin, stationary diesel-powered compressors may act similarly. This may likely be favored in addition by altitude effects, as it has been demonstrated for diesel vehicles up to an altitude of 2400 m a.s.l. (e.g., Bishop et al., 2001; Wang et al., 2013) In particular it seems that NO\textsubscript{2} emissions increase at the expense of NO emissions (Yin et al., 2013). For HCHO versus NO\textsubscript{x}, the slopes are significantly lower than found in traffic related combustion emissions. The correlation coefficient is also weaker. It is likely that different emission sources overlap. Since HCHO shows a closer correlation with total NMHC at the Boulder site and even closer with CH\textsubscript{4} as shown in Supplement S8, it is likely that wellhead equipment, in addition to co-located compressors, may contribute to HCHO emissions. Interestingly, contrary to HCHO,
HONO displays a stronger correlation with total NMHC than with CH₄. This may further support the ideas that HONO emissions are more related to combustion (compressors) and that HCHO has some overlapping emissions originating from drill-rig operations.

### 3.3.2 NMHC versus NOₓ limitation of ozone production

Observation-based methods like photochemical indicators such as O₃/NOₓ, O₃/NO₂, O₃/HNO₃ and the EOR (extent of reaction) can be used to evaluate whether ozone production at a given location may be VOC or NOₓ limited (Sillman, 2002; Sillman and He, 2002). The EOR is defined as

\[
\text{EOR} = \left(1 - \frac{\text{NO}_x}{1.3 \cdot \text{NO}_y}\right)^{0.67}. \tag{5}
\]

Sillman and He (2002) point out that the photochemical indicators also depend on the chemical environment, in particular for O₃/NOₓ (which may not be applicable for highly polluted scenarios). They classified relatively clean environments (O₃ < 80 ppbv), moderately polluted (10–200 ppbv O₃) and highly polluted environments (O₃ > 200 ppbv). According to Sillman and He, photochemical indicators would tend to increase with decreasing ozone values. It is not certain, if these photochemical indicators can be applied to the environment found in the UGRB without modifications, as the radical pool is largely dominated by the photolysis of HONO, at least in the lowermost layer of the Planetary Boundary Layer (PBL) (see discussion in Sect. 3.3.4). While there might be some limitations to this concept, we followed this approach in an attempt to explore whether an ensemble of photochemical indicators would at least point towards common tendencies with regard to VOC and NOₓ limitation. Supplement S9 lists ranges of indicator values that are representative of different regimes. We calculated the photochemical indicators for the Boulder site. For this analysis, we focus on the IOP days, as those were the days with maximum ozone levels. As shown in Fig. 3, a ozone mixing ratios are usually below 80 ppbv before 11:00 MST and afterwards above 80 ppbv. It is common to all photochemical indicators that they display significant changes throughout the day depending on the wind direction (Fig. 8). While there are no consistent patterns throughout the day among all photochemical indicators, some general tendencies for the most critical time periods with regard to regulatory purposes can be deduced. In most cases during morning hours (07:00–11:00 MST; O₃ < 80 ppbv) and/or under SW flow conditions, all indicators reach minimum values. Supplement S10 shows that O₃/NOₓ, O₃/NO₂ and the EOR would indicate VOC limitation; the values of O₃/HNO₃ would fall into the transitional regime according to Supplement S9, if not in the VOC limited regime assuming that photochemical indicators would be higher in the case, where O₃ < 80 ppbv. From 13:00 to 15:00 MST, i.e., the time period with O₃ > 80 ppbv, all indicators either point towards a NOₓ limited or a transitional regime. Any change from a VOC-limited regime towards a NOₓ-limited regime or vice versa implies passing the transitional regime, where ozone production can be most efficient and can reach maximum values as shown by Mao et al. (2010). The periods with the lowest values for photochemical indicators generally coincide with minimum values of the NMHC/NOₓ ratio as observed at the Boulder site (Fig. 9). This minimum value is about 50 and represents a relatively high value compared to urban areas (Mao et al., 2010). This could be due to relatively low NMHC reactivity, either caused by low temperatures or a NMHC mixture that is mostly composed of slow-reacting hydrocarbons (e.g., alkanes such as ethane and propane), or a combination of both. Carter and Seinfeld (2012) found that regimes sensitive to VOC or NOₓ may vary from year to year and depend on the specific location in the UGRB. For the Boulder site in 2011 they determined a VOC sensitive regime, which largely agrees with our analysis for the morning hours on IOP days. Carter and Seinfeld also state that under such conditions additional OH generated by an increased level of HONO would favor O₃ formation, while NOₓ saturation would also have the potential to cause additional HONO formation. Carter and Seinfeld (2012) modeled the Boulder 2011 case based on the HONO data presented in this paper and found much better simulations of O₃ compared to the baseline scenario, when HONO was added, which is very encouraging. In addition to the Carter and Seinfeld analysis our paper also considers diurnal variations of the NMHC/NOₓ ratio, including any fluctuations in the NMHC mix, which is also subject to wind-directional changes, at least for the Boulder site. As mentioned above, our data shows that there is likely a transition phase after the morning NOₓ saturation conditions towards a VOC saturation regime, which is not described by Carter and Seinfeld and would require diversified speciated regulatory mitigation strategies.

### 3.3.3 OH reactivity

In order to describe the importance of individual or classes of trace gases with regard to photochemical processes, it is important to consider both the reactivity and the amount of these trace gases in ambient air. One way to do this is to calculate the propene equivalent \( J \) as proposed by Chameides et al. (1992):

\[
J = C_J \cdot \frac{k_{OH}(J)}{k_{OH}(C\_3H\_6)} \tag{6}
\]

with \( C_J \) being the mixing ratio of hydrocarbon compound \( J \) in ppbC or any other compound \( J \) in ppbv, the reaction rate of this compound with OH (\( k_{OH} \)) and normalized to the reaction rate of propene with OH (\( k_{OH}(C\_3H\_6) \)). This approach expresses any individual or classes of trace gases in terms of propene units. However, it is an approximation as reactions other than with OH, i.e., with O₃, NO₃, and Cl, are neglected.
Figure 8. Diurnal and wind directional variation of photochemical indicators $\text{O}_3/\text{NO}_y$, $\text{O}_3/\text{NO}_z$, $\text{O}_3/\text{HNO}_3$ and EOR on IOP days. Units shown in brackets for the wind directional variation plots refer to the radial direction of the corresponding trace gas plot.
In our approach, we merge data from the Boulder South Road site (speciated NMHC, CH$_4$, CO, NO$_x$) with data collected at the Boulder site (HCHO, HONO) to have a more comprehensive data set. During the winters of 2011 and 2012 19 canister surveys of speciated NMHC were performed at sites throughout the UGRB. The Boulder and Boulder South Road sites are located three miles from each other. The Boulder site is closer to the spine of the Pinedale Anticline development and concentrations at this site are generally slightly higher than at Boulder South Road. While the $r^2$ for the comparison of data from these two sites for each of four surveys during February and March 2011 were each greater than 0.97, there is some variation of slope between the two sites with values ranging from 1.2 to 1.7. The two Boulder sites share the characteristic of measuring air that is influenced by a wide range of oil- and gas-emission sources (Field et al., 2012a, b) with similar contributions between NMHC classes. When considering speciated NMHC alkanes are most correlated and aromatic are least correlated. Although there might be slight differences in the atmospheric chemistry settings of both sites, we believe that this will not have major impacts on the general findings. Figure 10 displays the diurnal variation of the fractions of the propene equivalent on IOP days. Within the speciated NMHC obtained at the Boulder South Road site, the most important contributions to the overall OH reactivity stem from aromatics (50–60 %), alkanes (30–40 %) and the remaining $\sim$ 10 % alkenes. As expected, the contribution from alkynes and isoprene is negligible due to their low reactivity (alkynes) and low mixing ratios (isoprene). Although CH$_4$ accounts for the largest fraction of all hydrocarbons (ranging from 90 % around noon to 95 % in the morning hours on a ppb basis), its propene-equivalent fraction is quite modest with 1–2 % due its low reactivity. This pattern does not change much throughout the day and is not much different on non-IOP days (not shown). It should be noted that in terms of absolute propene equivalents, OH reactivity on IOP days reaches up to 48 ppbC before noon compared to 31 ppbC around the same time on non-IOP days. While alkanes contribute to the propene equivalent about twice as much
on IOP days than on non-IOP days, the contribution by aromatics increases by about 30%, whereas the contribution by alkenes stays about the same. When other compounds are included (CO, NOx, HCHO, HONO), it becomes apparent that NOx plays an important role both in terms of contribution to reactivity as well as to the diurnal change of contributions. On IOP days aromatics contribute about 35–40%, alkanes 20–30%, NOx 20–40%, and alkenes about 5–10% to the overall propene equivalent. The remaining ∼5% are due to CO, HONO, HCHO, alkenes, and CH4. The contribution from NOx reaches its maximum from 05:00 to 09:00 MST. The picture is different on non-IOP days, where NOx contributions are significantly lower throughout the day. More specifically, the contributions to the propene-equivalent on non-IOP days would be aromatics 40–60%, alkanes, 20–40%, NOx 10–35%, alkenes 5–10%, and ∼5% for the remaining trace gases CO, HONO, HCHO, alkenes, and CH4.

Figure 11 displays OH reactivity based on the sum of the reaction-rate coefficients multiplied by the mixing ratios of reactants with OH (Mao et al., 2010):

$$k_{OH} = \sum k_{OH-VOC}\left[\text{VOC}_i\right] + k_{OH-\text{CO}}[\text{CO}] + k_{OH-\text{NOx}}[\text{NO}] + k_{OH-\text{NOx}}[\text{NO}_2] + k_{OH-\text{HONO}}[\text{HONO}] + \ldots$$  \hspace{1cm} (7)

Figure 11 clearly shows that on IOP days OH reactivity is higher than on non-IOP days by a factor of 2. In the daytime during IOP days, the magnitude of OH reactivity is comparable to that found in direct OH measurements for urban air in Houston, Texas (Mao et al., 2010), or in Mexico City (Sheehy et al., 2010). Contrary to the urban Houston case, where NOx may contribute up to 50% to OH reactivity during the rush hour, for the Boulder case NOx may contribute up to 69% throughout the day. This may be due to NOx emission sources in the Boulder area whose emission strength does not change much throughout the day (e.g., compressors). Another distinction is that, at Boulder, alkenes may contribute around 5% to the overall OH reactivity while for the Houston case, it can be up to 35%. The contribution from alkanes is in the 10–15% range at Boulder, while in Houston it is about 5–10%. The contribution of aromatics is about the same for both cases (10–15%).

In accordance with Supplement S4 a and b, also the absolute propene equivalent displays similar distribution with the highest values under SW wind directions. The reactivity mix does not change that much with wind direction (i.e., mostly aromatics, alkanes, and NOx), mostly from the SW. Figure 12 shows that while alkanes contribute 80–85%, aromatics around 10–15%, and alkenes less than ∼5% of NMHCs on a mass basis, aromatics contribute 50% and alkanes around 45% to the propene equivalent as observed at the Boulder South Road site. Still, contributions by alkenes are low. Contributions from CH4 and isoprene are negligible.

The major finding is that on IOP days in the morning hours in particular, NOx (and to some extent HONO as it is closely associated with NOx), aromatics and alkanes (the latter ones largely dominated by ethane and propane) are major contributors to the OH reactivity and propene equivalent at Boulder. The highest reactivities are found in air masses arriving at the Boulder site under SW flow conditions. This time period and wind flow condition largely coincides with the lowest NMHC/NOx ratios observed at the Boulder site and a switch from NOx-limited to transitional, if not VOC-limited regime, with the largest ozone production efficiency as described previously in Sect. 3.4. Under VOC-limited conditions, it is likely that highly reactive aromatics, such as toluene and xylenes, may be most efficiently competing with other NMHCs in reactions with OH. NOx reactions with OH in turn will cause substantial formation of HNO3.

3.3.4 Role of HONO

As discussed, major removal processes for OH at the Boulder site involve reactions with NOx, aromatics, and alkanes. Sources for OH are photolysis processes of O3, aldehydes (foremost HCHO), H2O2, CINO2, and HONO. Alkene
ozonolysis may also contribute to OH formation. Recent studies suggest that major OH formation occurs through HONO photolysis in the morning, HCHO photolysis in late morning, and O$_3$ photolysis throughout the day, whereas photolysis of H$_2$O and alkene ozonolysis are of minor importance during the day (Elshorbany et al., 2009; Ren et al., 2013; Czader et al., 2013). According to an analysis by Elshorbany et al. (2009), HONO photolysis can contribute to about 52% of radicals on a 24 h average, followed by alkene ozonolysis (about 20%), HCHO photolysis (about 15%) and ozone photolysis (about 4%) in an urban area. However, in unpolluted polar regions HONO is suggested to be a major radical source (98%) and ozone photolysis would account for the remaining 2% (Villena et al., 2011). Mao et al. (2010) report that contributions to the radical pool from HONO photolysis may be highest in high NO$_x$ environments.

The primary production of hydroxyl radicals from the photolysis of O$_3$, P(OH)$_{prim,O3}$, the photolysis of HCHO, P(HO$_2$)$_{prim,HCHO}$, the photolysis of HONO, P(OH)$_{prim,HONO}$, and from alkene ozonolysis, P(OH)$_{prim,alkenes}$ can be calculated as follows (see also Elshorbany et al., 2009; Sörgel et al., 2011):

$$
P(OH)_{prim,O3} = \frac{2 \cdot J(O(1D)) [O_3] \cdot k_{O(1D)+HCHO}}{k_{O(1D)+HCHO} [HCHO] + k_{D}}$$

where photolysis rates for photolysis of O$_3$ ($J(O(1D))$), HCHO ($J(HCHO)$), and HONO ($J(HONO)$), temperature-dependent reaction rates (Atkinson et al., 2004, 2006) and $\Phi_{OH}$, the OH yield of gas-phase reactions of ozone with alkenes, are used (Rickard et al., 1999).

Figure 13 displays results of the calculation of OH-productions due to the above processes for IOP and non-IOP days. Photolysis of ClNO$_2$ is not included. As mentioned earlier we presume that ClNO$_2$ may not be present in appreciable amounts based on the low levels of particulate chloride. In Fig. 13, median data for ozone, HCHO, HONO, and alkenes were taken. The results clearly indicate that OH production on IOP days is mainly due to HONO at the measurement height of 1.80 m above the surface. Between 09:30 and 15:30 MST, HONO photolysis contributes between 83 and 94% of the entire OH production, i.e., slightly less than observed in polar areas (Villena et al., 2011), whereas the contributions by other processes range between 2 and 7% each. On a 24 h basis HONO photolysis on IOP days can contribute ~83% to OH production on average, followed by alkene ozonolysis (~9%). Photolysis by ozone and HCHO photolysis contributes about 4% each to hydroxyl formation. On non-IOP days the picture is different and closer to the results by Elshorbany et al. (2009) for an urban area. The contributions to hydroxyl formation on non-IOP days were as follows: HONO photolysis ~54%, alkene ozonolysis ~28%, ozone photolysis ~13%, and HCHO photolysis ~5%. It is important to reiterate that the values for OH production as shown in Fig. 13 are calculated for the measurement height of 1.80 m above the surface and may not necessarily be regarded as an integrated quantity throughout the PBL. It is also worth noting that the calculated OH-production due to HCHO photolysis represents an upper limit as we did not distinguish between primary and secondary HCHO in these calculations. These results are different from the simulation
analyses for a site in the Uintah Basin, Utah (Edwards et al., 2013). Potential reasons may include that Edwards et al. assumed a uniform increase of O3 precursors by a factor of 2 under cold pool events, that the site may have been exposed to a different ambient chemical matrix (e.g., HCHO is significantly higher, while HONO is significantly lower at the Horse Pool site, Utah, compared to the Boulder site, Wyoming) and that they assumed a primary fraction of 50 % of the observed daily integrated HCHO mixing ratio.

In environments with significant amount of NOx the dominant terminal loss mechanism for OH is the reaction with NO3:

$$\text{NO}_2 + \text{OH} + \text{M} \rightarrow \text{HNO}_3 + \text{M} \quad (\text{M}=\text{N}_2, \text{O}_2), \quad (12)$$

which translates into

$$L(\text{OH}) = k_{\text{NO}_2+\text{OH}}[\text{NO}_2]. \quad (13)$$

Thus a photostationary state OH concentration can be calculated as follows:

$$[\text{OH}]_{\text{PSS}} = P(\text{OH})_{\text{prim-NO}_2} + P(\text{OH})_{\text{prim-HONO}} + P(\text{OH})_{\text{prim-alkenes}} + P(\text{OH})_{\text{prim-HCHO}}$$

$$\frac{L(\text{OH})}{P(\text{OH})_{\text{prim-NO}_2}}. \quad (14)$$

Figure 14 shows that OH concentration on IOP days can reach very high values during the time period 11:00–16:00 MST on IOP days. These are in the ranges found in New York City (Ren et al., 2003), Houston (Mao et al., 2010), Mexico City (Sheehy et al., 2010) and in the Pearl River delta (Lu et al., 2012), and at times exceed those. On non-IOP days maximum OH mixing ratios are significantly lower, and are confined to noontime. The OH chain length ranges between 10 and 13 between 12:00–15:00 MST on IOP days, which is of similar magnitude as found in Houston and Mexico City, and significantly higher than in New York City (Mao et al., 2010). Supplement S11 clearly shows that the OH net production occurs around noontime and is in the order of $4 \times 10^7$ molecules cm$^{-3}$ s$^{-1}$. In urban areas the peak OH net production occurs during morning rush hours (Mao et al., 2010) at a similar or higher magnitude than in our study, and decreases significantly afterwards. In NOx-high environments, such as New York City the OH net production is maintained around noontime, albeit lower than during rush-hours and slightly lower than in our study.

Based on the calculated OH concentrations it is possible to determine the photostationary state concentration of HONO using the following equation:

$$[\text{HONO}]_{\text{PSS}} = \frac{k_{\text{OH+NO}}[\text{OH}] \cdot [\text{NO}]}{L(\text{OH}) + k_{\text{OH+HONO}}[\text{OH}].} \quad (15)$$

[\text{HONO}]_{\text{PSS}} was $\approx$ 26 % of the observed ambient HONO mixing ratios on IOP days and 44 % on non-IOP days between 11:30 and 15:30 MST, when P(\text{OH})_{\text{prim-HONO}} was at a maximum, which is significantly less than observed in the urban area of Santiago de Chile, for instance ($\approx$ 66 %; Elshorbany et al., 2009). These low fractions of [HONO]_{PSS} indicate that a strong daytime source for HONO must be present. High daytime HONO/NOx ratios are also usually considered an indicator for strong additional photochemical HONO sources (Elshorbany et al., 2009). Figure 15 shows that the HONO/NOx ratio is about the same value for IOP and non-IOP days throughout the nighttime. In particular, in the morning hours from 05:00 to 09:00 MST, when the maximum levels of primary pollutants are present (Fig. 3a), the HONO/NOx ratio tends to be at a minimum and is between 2 and 5 % in both cases, which is similar to other locations. However, it increases significantly from 12:00 to 14:00 MST on non-IOP days (median values $\approx$ 10 %) and from 11:00 to 16:00 on IOP days (median values up to 30 %). According to Villena et al. (2011) the magnitude of the HONO/NOx
ratio is similar to the magnitude of the ratio of the corresponding atmospheric lifetimes. Based on the calculated $J_{\text{HONO}}$ photolysis rates an atmospheric lifetime for HONO of about 5 min between 11:30 and 15:30 MST on IOP days can be deduced. Using the calculated OH concentrations and temperature-dependant reaction rates for reactions with OH and $O_3$ a corresponding atmospheric lifetime of $\sim$ 19 min for NO$_2$ can be retrieved, which translates into a ratio of both lifetimes in the order of $\sim 26\%$, which is close to the observed average median HONO/NO$_x$ ratios for that time period, which is $\sim 21\%$. On non-IOP days, the atmospheric lifetime of NO$_x$ is much longer (almost 1 h), the ratio of the HONO and NO$_x$ lifetimes is $\sim 12\%$, the observed HONO/NO$_x$ ratio is $\sim 9\%$. This indicates that more HONO is being formed in photochemical processes, most likely through NO$_2$ conversion, on IOP days during these time periods. At the same time, NO$_x$ levels decrease (Figure 3a). As meteorological conditions (PBL height; SW wind flow) do not vary much during this time of the day on IOP days, and NO$_x$ deposition velocity onto snow surfaces is low (Hauglustaine et al., 1994), rapid oxidation of NO$_x$ leads to a decrease in NO$_x$ mixing ratios. Reactions of NO$_2$ with OH and $O_3$ will lead to the formation of HNO$_3$ accelerated through the enhanced availability of radicals due to the strong HONO photolysis source. This decrease in ambient NO$_x$ levels indicate that these removal processes exceed other sources for NO$_x$, e.g., through photolysis of HONO.

For the Boulder site, it is the photolysis of HONO that makes a significant difference to hydroxyl formation, as it is up to a magnitude higher on IOP days than on non-IOP days around noontime and the overall 24 h production of OH due to HONO photolysis is 6 times higher on IOP days versus non-IOP days, at least at the sampling height of 1.80 m above the surface. While also the contribution by HCHO photolysis to hydroxyl formation is enhanced by 3–5 times, the contribution by ozone photolysis and alkene ozonolysis remains at the same level.

The high OH-production rate of about 10.7 ppbv h$^{-1}$ by HONO-photolysis must be balanced by HONO sources of the same magnitude. At the Boulder site no speciated photolysis-rate measurements were available. As a surrogate, we plotted measured incoming ultraviolet (UV) radiation in Fig. 15. On IOP days, incoming UV radiation was on the
order of 10% higher than on non-IOP days. We calculated photolysis rates using the TUV model (for specific details see figure caption of Fig. 13). Median nocturnal levels on IOP days were 3.14 × 10^{-3} s^{-1} for $J_{\text{NO}_2}$, 1.78 × 10^{-2} s^{-1} for $J_{\text{O}_3}$, 2.87 × 10^{-3} s^{-1} for $J_{\text{NO}_3}$, 3.64 × 10^{-3} s^{-1} for $J_{\text{HONO}}$, and 4.74 × 10^{-5} s^{-1} for $J_{\text{HCHO}}$. Likewise, photolysis rates were reduced by about 10% on non-IOP days. Those high photolysis rates are primarily due to the high snow albedo, already relatively high solar zenith angles in March, high altitude (2160 m) and relatively low latitude (42.7° N) of the UGRB. Some of these factors differ significantly from conditions found in polar regions. Generally, data in Fig. 15 shows that HONO increases as incoming UV radiation increases. Also during the same time periods O$_3$ increases as well. It appears likely that HONO is being produced through photo-enhanced formation processes, which are most efficient when snow cover is present.

It has become evident that ambient HONO concentrations are higher than can be accounted for by direct emissions and that heterogeneous processes on surfaces may lead to enhanced ambient HONO levels. In particular, these processes tend to occur on surfaces with adsorbed water in the dark (Finlayson-Pitts et al., 2003; Jenkin et al., 1988; Kleffmann et al., 1998) based on the following reaction (Goodman et al., 1999; Kleffmann et al., 1998):

$$2\text{NO}_2(\text{ads}) + \text{H}_2\text{O}(\text{ads}) \rightarrow \text{HONO}(\text{ads}) + \text{HNO}_3(\text{ads}).$$

Recently, an additional nighttime formation of HONO that was not related to NO$_2$ was reported on aqueous surfaces in the marine boundary layer by Wojtal et al. (2011), assuming a surface nano-layer saturated with NO$_2$ precursors and no irreversible loss of HONO from that layer. Wojtal et al. (2011) observed that HONO/NO$_2$ ratios would increase during the night from 3 to 30% and even higher on some occasions, with HONO levels about 1 ppbv at night. While we suspect that the snow surface could have been saturated with NO$_2$ precursors due to appreciable amounts of these compounds in ambient air (see Fig. 3a, b), we did not observe increasing HONO/NO$_3$ ratios throughout the night as shown in Fig. 15. Also, HONO mixing ratios were 3–10 times lower compared to the Wojtal et al., 2011.

As pointed out by Stutz et al. (2004), relative humidity may be favorable for HONO formation, at least in urban environments. Supplement S12 shows that on IOP and non-IOP days, HONO levels are higher during the day than at night. During nighttime, photochemical processes are at a minimum. A trend towards higher HONO levels with increasing relative humidity can be discerned, with a maximum around 80%. A dependency of the HONO/NO$_2$ ratio on relative humidity as described by Stutz et al. (2004) could not be identified in our data set, likely due to the fact that the snow cover itself provided a constant amount of water, which would be in line with observations by Wojtal et al. (2011) for aqueous surfaces.

Over the last decade a variety of photo-enhanced HONO formation mechanisms have been discussed to explain observations of elevated daytime HONO levels (Kleffmann, 2007). These include (i) the photolysis of surface-adsorbed nitrate or nitric acid (Zhou et al., 2001; Zhou et al., 2002; Ramazan et al., 2006); (ii) the photolysis of ortho-nitrophenols (Bejani et al., 2006); (iii) light-induced NO$_2$ reduction on surface-adsorbed humic acid films (Stemmeler et al., 2007); (iv) gas-phase reaction of electronically excited NO$_2$ due to photolysis, with water (Li et al., 2008); and (v) the conversion of HNO$_3$ to HONO on primary organic aerosol (Ziemba et al., 2010).

As mentioned in Sect. 3.1, observations at the Boulder site were characterized by remarkably high ambient levels of HONO, HNO$_3$, and p-NO$_3^-$, which on IOP days were higher than in urban air measurements, e.g., in Houston, Texas. HONO showed close correlation with HNO$_3$ not only during nighttimes (Supplement S8) but, contrary to NO$_2$ and NO$_x$, also during daytime regardless of IOP or non-IOP days with correlation coefficients ranging from 0.73 to 0.80. Contrary to Ziemba et al. (2010), who found that HONO was anti-correlated with HNO$_3$ during morning traffic rush hours, our observations do not show any anticorrelation during any time period. We thus assume that conversion of HNO$_3$ to HONO on primary organic aerosol will be negligible in the UGRB. HNO$_3$ on IOP days formed a higher fraction of NO$_3$ than in Houston (22 vs. 16%). Its fraction of NO$_3$ ranged between 30 and 40%, with maximum values found under SW wind directions. Again, corresponding values in Houston were only about 20% (Luke et al., 2010). p-NO$_3^-$ at the Boulder site was closely related to EC, elemental carbon, ($r^2 = 0.85$) and OC, organic carbon ($r^2 = 0.73$), but only showed poor correlation with sulfur ($r^2 = 0.09$) and sulfate ($r^2 = 0.03$). This fingerprint is likely associated with fossil fuel combustion in industrial processes (Jacobson, 2012). Björkman et al. (2013) studied dry deposition velocities for HNO$_3$ and p-NO$_3^-$ onto snow surfaces in the arctic. They found that the dry deposition velocity for p-NO$_3^-$ critically depends on the particle size; still, for particles of around 7 µm size, the dry deposition velocity is significantly lower than for HNO$_3$. Overall, the dry deposition of p-NO$_3^-$ only accounts for 1–7% of the total nitrate dry deposition (Björkman et al., 2013). According to Reaction (8) HNO$_3$(ads) can already be formed through NO$_2$(ads). Dry deposition velocity of HNO$_3$ onto snow surfaces is about 50 times greater than for NO$_2$ (Hauglustaine et al., 1994). Using median data for NO$_2$ and HNO$_3$ mixing ratios as shown in Supplement S2 dry deposition flux of HNO$_3$ onto the snow surface may be about 8–65 times greater than for NO$_2$, with maximum values during daytime, as HNO$_3$ mixing ratios surpass those of NO$_2$. While Reaction (8) would likely contribute to HONO formation throughout the day as shown by Wojtal et al. (2011) over aqueous surfaces, we assume that deposition of HONO would be of
When adsorbed on a surface, the following reactions may occur (Mack and Bolton, 1999; Zhou et al., 2002; Zhou et al., 2003):

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

Reaction (17) requires sunlit daytime conditions. NO$_2$(ads) formed in Reaction (19) may then further react according to Reaction (16) and subsequently form HONO. According to Zhou et al. (2003), the production yield of HONO from surface HNO$_3$ photolysis is almost independent of relative humidity between 20 and 80%, although some moisture is necessary. Thus, snow cover would provide a favorable substrate. However, the negligible dependence on relative humidity may explain, at least partly, why we did not find a clear correlation of HONO with relative humidity.

Supplement Fig. S13 shows that generally the extra HONO source, which balances the OH production rate due to HONO photolysis (dOH/dt-HONO; see Fig. 13) correlates better with $J_{\text{NO}_2}$ ($R^2_{\text{IOP}} = 0.86$; $R^2_{\text{non-IOP}} = 0.85$) and $J_{\text{HNO}_3}$ ($R^2_{\text{IOP}} = 0.82$; $R^2_{\text{non-IOP}} = 0.88$) than with $J_{\text{NO}_2}$ ($R^2_{\text{IOP}} = 0.71$; $R^2_{\text{non-IOP}} = 0.85$). According to Villena et al. (2010) and Elshorbany et al. (2012) a better correlation of the daytime HONO extra source with $J_{\text{NO}_2}$ would point to photolysis of HNO$_3$ as a potential HONO source, whereas a better correlation with $J_{\text{NO}_2}$ would point to photosensitized conversion of NO$_2$ on humic acid surfaces.

Following the approach by Villena et al. (2010) we determined the correlation of the HONO extra source with photolysis rates for each day. In our case we used photolysis rates based on the TUV calculation. In Supplement S14 the slopes $m$ of each daily correlation of the HONO extra source versus $J_{\text{NO}_2}$ ($m_{\text{IOP}}$) are plotted against median daytime NO$_2$ mixing ratios for the same time periods. For these plots any days with daily median NO mixing ratios > 500 pptv were excluded to avoid potential bias due to freshly polluted air masses. The results are segregated for IOP and non-IOP days. A similar approach is shown for $m_{\text{HNO}_3}$ against median daytime HNO$_3$ mixing ratios. The following observations can be deduced: (i) in both cases IOP days show higher $m_{\text{NO}_2,\text{HNO}_3}$ values, (ii) the slopes of $m_{\text{NO}_2,\text{HNO}_3}$ are steeper on IOP days than on non-IOP days, and (iii) correlation coefficients $R^2$ of $m_{\text{NO}_2,\text{HNO}_3}$ versus NO$_2$ and HNO$_3$, respectively, are higher on IOP than non-IOP days. While this indicates photo-enhanced HONO formation on IOP days, the stronger correlation coefficients in the case of $m_{\text{HNO}_3}$ versus HNO$_3$ mixing ratios ($R^2_{\text{IOP}} = 0.97; R^2_{\text{non-IOP}} = 0.89$) compared with the case of $m_{\text{NO}_2}$ versus NO$_2$ mixing ratios ($R^2_{\text{IOP}} = 0.84; R^2_{\text{non-IOP}} = 0.39$) would indicate that photolysis of HNO$_3$ may play an important role for HONO formation. According to Villena et al. (2010) the correlation of $m_{\text{HNO}_3}$ against the product of [NO$_2$] × 1/v (v being the horizontal wind speed) used as an indication for turbulent vertical mixing, was even stronger, suggesting a ground surface source. In our study, correlations of $m_{\text{HNO}_3}$ versus HNO$_3$ mixing ratios ($R^2_{\text{IOP}} = 0.98; R^2_{\text{non-IOP}} = 0.97$) as well as $m_{\text{NO}_2}$ versus NO$_2$ mixing ratios ($R^2_{\text{IOP}} = 0.95; R^2_{\text{non-IOP}} = 0.85$) also became stronger when wind speed was included.

Additional estimates on PBL properties can be deduced based on meteorological measurements obtained at the Tall Tower site located about 10 km southeast of the Boulder site (2149 m; 42.4241° N, −109.5609° W; see Fig. 1). For describing atmospheric stability the bulk Richardson number $R_B$ can be applied (Stull, 1988):

\[ R_B = \frac{g \Delta \Theta_v \Delta z}{\Theta_v \left( (\Delta U)^2 + (\Delta V)^2 \right)}. \]

where $\Delta \Theta_v$ is the virtual potential temperature averaged over a given time interval, $\Delta z$ the difference between two altitudes, $\Delta \Theta_v$ the difference of the virtual potential temperature over $\Delta z$, averaged over a given time interval, $\Delta U$ and $\Delta V$ the differences of the wind speed in $u$ and $v$ directions, respectively, over $\Delta z$ averaged over a given time interval, and $g$ the gravitational constant. The bulk Richardson number relates the thermal buoyancy to mechanical shear. For our calculations we used the meteorological data obtained at the 3 and 25 m levels. In the absence of relative humidity measurements at the 25 m level we calculated $R_B$ using potential temperature instead of virtual potential temperature. Figure 14 indicates that on IOP days most of the time the PBL is stable and behaves like a laminar stratified flow. It is only during the time frame 11:00–16:00 MST that the PBL becomes turbulent. During this time frame also the mixing layer height tends to increase, albeit slightly, until the maximum atmospheric instability is reached. More importantly, this time frame also coincides with the time period of maximum OH mixing ratios as shown in Fig. 14. This is the time frame when vertical exchange processes from the surface may occur. During other times of the day these processes are largely suppressed.

The distribution of trace gases within the atmosphere depends on their atmospheric lifetime and the atmospheric turbulent diffusion. The turbulent diffusion time $\Delta t$ is determined as

\[ \Delta t = \frac{(\Delta z)^2}{2K_z}. \]

Likewise, a diffusion distance can be defined:

\[ \Delta z = \sqrt{2K_z \cdot \Delta t}, \]

where $\Delta z$ is the vertical distance traveled by an eddy and $K_z$ is the turbulent diffusion coefficient or eddy diffusivity. $K_z$
relates vertical fluxes of matter with properties of the PBL (e.g., Stull, 1988, and references therein):

\[ F = -K_z \frac{\partial C}{\partial z} = -\frac{ku_z}{\Phi_M} \frac{\partial C}{\partial z}, \]

where \( F \) is the vertical flux, \( C \) the mixing ratio of a trace gas, \( k = 0.4 \) the von Kármán constant, \( u_z \) the friction velocity, and \( \Phi_M \) an empirically derived stability function. \( \Phi_M \) can be described as a function of the bulk Richardson number for specific cases (Arya, 1998):

\[ \Phi_M = I - R_B \quad 0.25 < R_B < 1, \quad (23) \]

\[ \Phi_M = (1 - 15 \cdot R_B)^{\frac{1}{2}} \quad R_B < 0.25, \quad (24) \]

In our study \( K_z \) can only be calculated for cases \( R_B < 1 \), i.e., for the time frame 11:00–16:00 MST (see Fig. 14). For this time frame and for a height of 3 m the median \( K_z \) was 348 cm\(^2\) s\(^{-1}\), ranging from 253 to 840 cm\(^2\) s\(^{-1}\), which is about 5–10 less than observed above the snowpack at Summit, Greenland (Honrath et al., 2002), but of the same magnitude found in Alert, Canada (Zhou et al., 2001). Given the maximum mixing layer height of about 58 m (Fig. 14) this translates into a turbulent diffusion time of more than 13 h likely causing the development of vertical gradients of trace gases whose atmospheric lifetime is shorter than 13 h. Given an atmospheric lifetime of 5–7 min for HONO around noon-time, when turbulent mixing is present, HONO mixing ratios may have decreased due to photolysis by 15 % by the time it reaches the 1.80 m sampling level and by 63 %, when it reaches about 4.40 m, which is significantly lower than found in Alert, which was 11 m (Zhou et al., 2001). At Boulder HONO will almost be completely removed (by 99.6 %) when it reaches 10 m above the surface level. This means that OH formation due to HONO is likely confined to the lowermost 10 % of the PBL, while other OH formation processes due to photolysis of HCHO, \( \text{O}_3 \) and alkene ozonolysis may be active throughout the PBL.

We calculated \( K_z \) for the surface layer. \( K_z \) usually tends to increase with height within the boundary layer (e.g Stull, 1988, and references therein). Using meteorological data obtained at the 25 and 50 m levels at the Tall Tower site, we obtained a median value for \( K_z \) of 2.6 \times 10^3 \text{ cm}^2 \text{ s}^{-1} \pm 35 \%, which is 7.5 times larger than those obtained in the surface layer. This enhanced vertical transport process may partly contribute to the occurrence of an ozone peak between 50 and 70 m above the ground level as seen in the ozone profiles (Fig. 6).

Zhou et al. (2003) calculate HONO formation rates based on surface HNO\(_3\) photolysis. We adopted this approach to explore whether this process may balance the median HONO photolysis rate of about 10.7 ppbv h\(^{-1}\) at the Boulder site around noon-time on IOP days. According to Zhou et al. (2003) the upward flux \( F_{up} \) of NO\(_x\) and HONO can be described as follows:

\[ F_{up} = \alpha J \cdot v \cdot C \cdot t, \]

where \( \alpha \) represents the fraction of the average of the individual median values for the diurnal variation of the UV radiation (Fig. 15) versus the noontime maximum value of the UV radiation. This value is about 25 % and is used to determine the fraction of deposited HNO\(_3\) exposed to the noontime photolysis rate \( J_{HNO3} \) (s\(^{-1}\)). The parameter \( v \) is the average HNO\(_3\) dry deposition velocity, \( C \) is the HONO concentration (mol m\(^{-3}\)) and \( t(s) \) is the accumulation time since the last precipitation. Zhou et al. (2003) applied the following values for their calculation: \( J_{HNO3} = 2.5 \times 10^{-5} \text{ s}^{-1} \), \( v = 2 \times 10^{-2} \text{ m s}^{-1} \), \( C = 700 \text{ pptv} = 3.1 \times 10^{-8} \text{ mol m}^{-3} \), and \( t = 24 \text{ h} = 86400 \text{ s} \). Zhou et al. (2003) calculated a HONO production rate of \( \sim 150 \text{ ppbv h}^{-1} \) for a 100 m boundary layer height. For the estimate at the Boulder site, we consider a different HONO concentration and also assume a different photolysis rate \( J_{HNO3} \) due to higher albedo. The median HONO mixing ratio on IOP days at Boulder was 2.1 ppbv (which equals 9.37 \times 10^{-8} \text{ mol m}^{-3} \). According to the TUV model \( J_{HNO3} \) is \( \sim 2.5 \) times higher over surfaces with 75 % albedo (e.g., over snow) compared with standard surfaces. For the Boulder site the effective active layers for HONO are the lowermost 10 m of the PBL. This would result in a HONO emission flux of 10.2 \pm 40 \% ppbv h\(^{-1}\), with the largest uncertainties associated with the estimates for \( J_{HNO3} \) and the accuracy of the HNO\(_3\) measurements. The magnitude of the HONO emission flux is very close to the observed loss of HONO due to its photolysis. Apart from surface emissions, combustion related HONO emissions may contribute to the HONO flux. As outlined earlier we found a robust HONO/NO\(_x\) emission ratio of 15 ppbv HONO/1 ppbv NO\(_x\) during nighttime under SW flow conditions. In the quadrant southwest of Boulder multiple facilities related to oil-drilling activities are distributed over an area of \( \sim 300 \text{ km}^2 \). According to the WDEQ inventory (WDEQ, 2011) the overall NO\(_x\) emission from these facilities is \( \sim 1180 \text{ kg h}^{-1} \) with about 91 % originating from drill rigs and completion emissions. The overall NO\(_x\) flux in this area would be \( \sim 4000 \mu \text{g m}^{-2} \text{ h}^{-1} \). Assuming a boundary layer height of 50 m a.g.l. this would result in an increase of \( \sim 39 \text{ ppbv h}^{-1} \) for NO\(_x\) and an increase of \( \sim 0.6 \text{ ppbv h}^{-1} \) for HONO using the emission ratio of 15 ppbv HONO/1 ppbv NO\(_x\). We estimate the uncertainty of the combustion-related HONO emissions on the order of 30 %, with major uncertainties associated with the height of the boundary layer. The combustion-related HONO source largely contributes to nighttime HONO abundance. During daytime, radiative processes prevail between 07:00 and 17:00 MST and decrease this contribution. Given an average wind speed of 1.4 m s\(^{-1}\) between 11:00 and 16:00 MST on IOP days and a distance of about 3–5 km between the Boulder site and most local sources we estimate that photolysis will have removed 85 % (for the closest sources) and up to 90 % (for the most distant sources) of ambient HONO by the time air masses would reach the Boulder site, based on a HONO lifetime of \( \sim 5–7 \) min during the same time period. Thus, during the transport
HONO photolysis may likely have contributed to radical formation. At the Boulder site an estimated HONO production of $\sim 0.1 \text{ ppbv h}^{-1}$ would still be due to combustion-related HONO emissions. Together with the surface HONO flux due to photolysis of HNO$_3$ adsorbed on the snow surface this would account for a total HONO production rate of $10.3 \pm 0.4 \text{ ppbv h}^{-1}$ at around noontime on IOP days, which is in very good agreement with the photolysis rate of HONO of $10.7 \pm 30 \text{ ppbv h}^{-1}$. We therefore conclude that, ultimately, NO$_x$ emitted into the extremely shallow boundary layer during the wintertime season in the UGRB is causing high HONO levels. This occurs from two processes: (i) HNO$_3$ produced during atmospheric oxidation of NO$_x$ can be deposited onto the snow surfaces, which facilitate subsequent photo-enhanced heterogeneous conversion to HONO and (ii) through combustion related emission of HONO. HONO, in turn, will serve as the most important precursor for OH, strongly enhanced due to the high albedo of the snow cover. The first process is the most dominant process.

4 Conclusions

During the periods 28 February–2 March and 9–12 March 2011, designated IOPs, high hourly ozone readings of up to 166 ppbv were observed at the Boulder surface site, located in the Upper Green River basin, Wyoming. These values were well above the estimated surface background ozone of $\sim 40 \text{ ppbv}$ and lower tropospheric ozone background of $\sim 60 \text{ ppbv}$ in March. During the IOPs, comprehensive surface measurements were performed and observations of vertical distributions of meteorological parameters and selected trace gases were carried out. The observed high ozone levels likely result from a combination of multiple factors.

Maximum ozone values are restricted to a very shallow surface layer. There is no evidence of ozone carryover from previous days. Ozone levels may remain stable as ozone deposition velocity onto snow surfaces is low. On IOP days, slightly variable wind directions (including recirculation) and low wind speeds in combination with low mixing layer heights ($\sim 50 \text{ m a.g.l.}$ around noontime) are essential for accumulation of both primary and secondary pollutants within the UGRB. IOP days are usually characterized by the fact that, at least once during the day, trajectories pass through the area southwest of Boulder before arriving at the Boulder site. Recirculation processes within the basin may also occur. Air masses would then pass the oil- and gas-well locations of the Pinedale Anticline and also a large number of compressor facilities at the most 1–2 h before arrival at Boulder and would have stayed in that region for about 1 h. Compressors and drill rigs primarily emit NO$_x$, relative to wellhead production equipment, which primarily emits CH$_4$ and NMHC. While compressors and drill rigs would also emit HONO and to some extent HCHO, wellhead production equipment would only contribute to emissions of HCHO. This is largely supported in an analysis of nighttime ratios of HONO and HCHO versus NO$_x$, NMHC, and CH$_4$ in air masses coming from this area. In general, these air masses contain maximum amounts of both primary and secondary pollutants. They are also associated with the highest VOC reactivity and a change of NO$_x$ sensitivity towards a VOC-sensitivity regime.

On IOP days, particularly in the morning hours, NO$_x$ (up to 69 %) followed by aromatics and alkanes ($\sim 10\text{–}15 \%$; mostly ethane and propane) are the major contributors to the OH reactivity and propene equivalent at Boulder. The highest OH reactivities (up to $22 \text{ s}^{-1}$) are found in air masses arriving at the Boulder site under SW flow conditions. This time frame (and also wind direction) largely coincides with the lowest NMHC/NO$_x$ ratios at the Boulder site and a change from a NO$_x$-limited regime towards a VOC-limited regime, which implies reaching or passing the transitional regime where ozone production can be most efficient and can reach maximum values. This is supported by photochemical indicators such as O$_3$/NO$_x$, O$_3$/NO$_x$, O$_3$/HONO and the EOR. The NMHC/NO$_x$ at the Boulder site during these periods is $\sim 50$ and represents a relatively high value compared to urban areas. This could be due to relatively low NMHC reactivity, either caused by low temperatures or a NMHC mixture that is mostly composed of slow-reacting hydrocarbons (e.g., alkanes), or a combination of both. Under VOC-limited conditions, it is likely that highly reactive aromatics, such as toluene and xylene, may be most efficiently competing with other NMHC in reactions with OH. OH reactions with OH in turn will cause substantial formation of HNO$_3$.

OH production on IOP days is mainly due to HONO. Until noon HONO photolysis contributes between 83 and 94 % of the entire OH-production, whereas the contributions by other processes range between 2 and 7 % each. On a 24 h basis and as determined for a measurement height of 1.80 m above the surface, HONO photolysis on IOP days can contribute $\sim 83 \%$ to OH production, on average, followed by alkene ozonolysis ($\sim 9 \%$). Photolysis by ozone and HCHO photolysis contribute about 4 % each to hydroxyl formation. The contributions to hydroxyl formation on non-IOP days were as follows: HONO photolysis $\sim 54 \%$, alkene ozonolysis $\sim 28 \%$, ozone photolysis $\sim 13 \%$, and HCHO photolysis $\sim 5 \%$.

We conclude that, ultimately, NO$_x$ emitted into the extremely shallow boundary layer during the winter season in the UGRB is causing high HONO levels (maximum hourly median on IOP days: 1096 pptv) through (i) HNO$_3$ produced in atmospheric oxidation of NO$_x$, deposited onto the snow surface and undergoing photo-enhanced heterogeneous conversion to HONO (estimated HONO production: $10.2 \pm 40 \text{ ppbv h}^{-1}$) and (ii) combustion-related emission of HONO at the Boulder site (estimated HONO production: $\sim 0.1 \pm 30 \text{ ppbv h}^{-1}$). HONO production is likely confined to the lowermost 10 m of the boundary layer. HONO, in turn, serves as the most important precursor for OH, strongly
enhanced due to the high albedo of the snow cover (HONO photolysis rate: 10.7 ± 30 % ppbv h\(^{-1}\)). OH radicals oxidize NMHCs, mostly aromatics (toluene, xylenes) and alkanes (ethane, propane), eventually leading to an increase in ozone. The data does not suggest that relative humidity favors the presence of high levels of HONO. This may be due to the assumptions that the surface-HNO\(_2\) photolysis is almost independent of relative humidity (Zhou et. al., 2003) or to the limitation of our data, which always showed relative humidity well above 50 % during the nighttime.

In our analysis we only considered radical production at the sampling height of 1.80 m, not as an integrated quantity throughout the PBL. Our study suggests that future work would need to include vertically resolved HONO measurements throughout the boundary layer to describe in detail the vertical distribution of the atmospheric oxidation potential.

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