On the contribution of nocturnal heterogeneous reactive nitrogen chemistry to particulate matter formation during wintertime pollution events in Northern Utah

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Abstract. Mountain basins in Northern Utah, including Salt Lake Valley (SLV), suffer from wintertime air pollution events associated with stagnant atmospheric conditions. During these events, fine particulate matter concentrations (PM_{2.5}) can exceed national ambient air quality standards. Previous studies in SLV have found PM_{2.5} is primarily composed of ammonium nitrate (NH₄NO₃), formed from the condensation of gas-phase ammonia (NH₃) and nitric acid (HNO₃). Additional studies in several western basins, including SLV, have suggested that production of HNO₃ from nocturnal heterogeneous N₂O₅ uptake is the dominant source of NH₄NO₃ during winter. The rate of this process, however, remains poorly quantified, in part due to limited vertical measurements above the surface, where this chemistry is most active. The 2017 Utah Winter Fine Particulate Study (UWFPS) provided the first aircraft measurements of detailed chemical composition during SLV wintertime pollution events. Coupled with ground-based observations, analysis of day and nighttime research flights confirm that PM_{2.5} during wintertime pollution events is principally composed of NH₄NO₃, limited by HNO₃. Here, observations and box-model analyses assess the contribution of N₂O₅ uptake to nitrate aerosol during pollution events using the NO₃⁻ production rate, N₂O₅ heterogeneous uptake coefficient (γ(N₂O₅)), and production yield of ClNO₂ (ϕ(ClNO₂)), which had medians of 1.6 μg m⁻³ hr⁻¹, 0.076, and 0.220, respectively. While fit values of γ(N₂O₅) may be biased high by a potential under-measurement in aerosol surface area, other fit quantities are unaffected. Lastly, additional model simulations suggest nocturnal N₂O₅ uptake produces 3.9 μg m⁻³ of nitrate per day, when considering the possible effects of dilution. This nocturnal production is sufficient to account for 86% of the daily observed surface-level build-up of aerosol nitrate, though accurate quantification is dependent on modeled dilution and mixing processes.
Introduction

Over 80% of Utah’s population lives in counties that experience periods of elevated fine particulate matter (PM\textsubscript{2.5} < 2.5 μm in diameter) during the winter season (U.S. Census Bureau, 2018; Whiteman et al., 2014). In these counties, the highest levels have been limited to three northern valleys along the Wasatch Mountains, shown in Figure 1 (north to south: Cache Valley (Logan Non-attainment area (NAA)), Salt Lake Valley (Salt Lake NAA), and Utah Valley (Provo NAA)). These valleys were designated by the U.S. EPA as “Moderate” non-attainment areas (NAA) in December 2009, with the Salt Lake and Provo areas reclassified from moderate to “Serious” in May 2017 (Utah Department of Environmental Quality). Elevated PM\textsubscript{2.5} concentrations in these regions impact public health and are associated with increases in emergency room visits for asthma (Beard et al., 2012). Short-term exposure has also been shown to increase the chance of triggering acute ischemic heart disease events by 4.5-6% per 10 μg m\textsuperscript{-3} of PM\textsubscript{2.5} in sensitive populations living in the Wasatch region (Pope et al., 2006; Pope et al., 2015).

Elevated wintertime PM\textsubscript{2.5} concentrations in these valleys typically correspond to multi-day events of high atmospheric stability (e.g. Whitman et al., 2014; Silcox et al., 2012; Gillies et al., 2010; Wang et al., 2012; Green et al., 2015; Silva et al., 2007; Baasandorj et al., 2017), associated with large, synoptic-scale high-pressure systems that transit from west to east, simultaneously impacting multiple basins across the intermountain western U.S. (e.g. Reeves and Stensrud, 2009). Warm temperatures aloft cause boundary layer stratification that reduces mixing and traps cold air and emissions near the surface, illustrated in Figure 2 and discussed further below. These events, termed persistent cold air pools (PCAPs), typically mix-out after 1-5 days but have been observed to persist for as long as 18 days (Whiteman et al., 2014). Similar meteorological patterns have been linked to wintertime PM\textsubscript{2.5} accumulation in basins across the western U.S. (e.g. Chen et al., 2014) before plateauing after ~6 days into an event (Baasandorj et al., 2017). Average 24-hour concentrations reported during PCAP events between 2001 and 2016 have been as large as 40-80 μg m\textsuperscript{-3} in Salt Lake (Baasandorj et al., 2017; Silcox et al., 2012) and Utah Valleys (Malek et al., 2006), and up to 132.5 μg m\textsuperscript{-3} in Logan, Utah (Cache Valley) (Malek et al., 2006).

Previous ground-based studies have identified ammonium nitrate (NH\textsubscript{4}NO\textsubscript{3}) as the main component of PM\textsubscript{2.5} (70 - 80% by mass) during PCAP events in all three Northern Utah Valleys (Silva et al., 2007; Hansen et al., 2010; Kuprov et al., 2014; Kelly et al., 2013; Long et al., 2003; Long et al., 2005a; Long et al., 2005b; Baasandorj et al., 2017). Ammonium nitrate formation is thermodynamically favorable under cold wintertime conditions from the equilibrium between gas-phase ammonia (NH\textsubscript{3}) and nitric acid (HNO\textsubscript{3}), shown in Reaction (R1) in Figure 2 (e.g. Kuprov et al., 2014; Nowak et al., 2012; Mozurkewich, 1993). PM\textsubscript{2.5} mitigation strategies that are based on control of these gas-phase species are expected to be more effective if the limiting reagent and its sources can be identified. Both model- and observationally-informed ground-based analyses have suggested that NH\textsubscript{4}NO\textsubscript{3} formation in Cache and Salt Lake Valleys is limited by the production of HNO\textsubscript{3} (Kuprov et al., 2014; Mangelson et al., 1997; Martin, 2006; Utah Division of Air Quality, 2014b, a, c; Franchin et al., 2018), though uncertainties remain in how this limitation may be impacted by temporal and spatial variations.

While NH\textsubscript{3} is directly emitted from agricultural sources, industrial processes, waste disposal, and automobile emissions (Behera et al., 2013; Livingston et al., 2009), HNO\textsubscript{3} forms chemically in the atmosphere from the oxidation of NO\textsubscript{x} (=NO + NO\textsubscript{2}), which in turn arises mainly from combustion emissions. There are two mechanisms by which this formation occurs, illustrated by reactions (R2 - R6) in Figure 2. The first is through daytime NO\textsubscript{2} oxidation by the hydroxyl radical (OH) (Figure 2, R2) and the second is through the nocturnal heterogeneous uptake of dinitrogen pentoxide (N\textsubscript{2}O\textsubscript{5}) (R6), which itself is a product of nocturnal
NO$_x$ oxidation (R3 – R5). The former is relatively more important during the summer (Brown et al., 2004) whereas the latter, the focus of this study, may be relatively more important in winter (e.g. Wagner et al., 2013) due to reduced OH concentrations, colder temperatures that favor N$_2$O$_5$ in its equilibrium with NO$_3$ (R5), and longer nights that allow more time for nocturnal reactions to occur. The nocturnal heterogeneous production of HNO$_3$ is also expected to be largest in the residual layer (RL), due to the near surface accumulation of NO$_x$ which titrates O$_3$ (R3) and reacts with NO$_3$ (R7), the precursor to N$_2$O$_5$ (e.g. Brown and Stutz, 2012).

The role of this nocturnal reactive nitrogen chemistry in the formation of PM$_{2.5}$ has been considered in previous wintertime studies, though nocturnal, vertically-resolved measurements have been limited. Previous studies using ground and tower-based observations, as well as mid-morning aircraft vertical profiles have identified heterogeneous chemistry and subsequent morning transport from aloft as a major source of surface-level NH$_4$NO$_3$ in California’s San Joaquin Valley (e.g. Brown et al., 2006; Prabhakar et al., 2017; Pusede et al., 2016; Watson and Chow, 2002). Nocturnal heterogeneous chemistry has also been considered as a source for PM$_{2.5}$ in northern Utah (Baasandorj et al., 2017; Kuprov et al., 2014), though vertically resolved measurements have been limited to ground-based observations at different elevations along the Wasatch Mountains (Baasandorj et al., 2017). In an analysis of ground-based HNO$_3$ and PM$_{2.5}$ observations in SLV, Kuprov et al. (2014) suggested that daytime HNO$_3$ formation was dominant over the contribution from nocturnal heterogeneous chemistry. Baasandorj et al. (2017), however, noted that ground-based measurements in this region may not capture the extent of heterogeneous chemistry aloft in the RL, which is expected to be distinct from the surface composition (e.g. Brown et al., 2007; Brown and Stutz, 2012; Stutz et al., 2004).

Therefore, vertical gradients in NO$_x$ and oxidants could promote efficient HNO$_3$ and NH$_4$NO$_3$ formation aloft, which could contribute to enhanced surface-level PM$_{2.5}$ concentrations the following day. Regardless of altitude, the absolute contribution will depend on 1) the rate of NO$_x$ and N$_2$O$_5$ production, 2) the efficiency of N$_2$O$_5$ uptake onto aerosol ($\gamma$(N$_2$O$_5$)) and 3) the heterogeneous production yield of HNO$_3$ relative to ClONO$_2$ ($\phi$(ClONO$_2$)) (Osthoff et al., 2008; Behnke et al., 1997). Net accumulation surface-level NH$_4$NO$_3$, however, also depends on mixing and dilution associated with growth of the convective boundary layer and mixing of the RL down to the surface during the following day. Quantification of these processes is a key component in designing effective mitigation strategies for Utah’s wintertime air pollution and requires vertically resolved observations of chemical composition at night.

In this study, we present results from the Utah Winter Fine Particulate Study (UWFPS), which consisted of aircraft and ground-based observations throughout Cache, Salt Lake, and Utah Valleys during January and February 2017. This analysis focuses on data from 16 aircraft flights (5 at night) during two pollution events between 16 January and 1 February 2017. These flights were carried out in SLV, the most populated of the three Utah Non-Attainment Areas. An overview of PM$_{2.5}$ during winter 2016-2017 is presented in the first section. Ambient mixing ratios of total (gas and particle-phase) oxidized and reduced nitrogen are shown in the second section to assess the limiting reagent to NH$_4$NO$_3$ aerosol formation, as well as its spatial and temporal trends. The final section presents upper-limit NH$_4$NO$_3$ production rate estimates from aircraft observations and results from a chemical box model that is fit to observations to calculate $\gamma$(N$_2$O$_5$), $\phi$(ClONO$_2$), and an estimated contribution of nocturnal heterogeneous chemistry to NH$_4$NO$_3$ formation in SLV. The contribution of nocturnal production relative to photochemically-driven NO$_2$ oxidation will have consequences for the development of effective mitigation strategies as day and nighttime production processes may have different sensitivities to NO$_x$ emissions and VOC radical sources (Pusede et al., 2016; Womack et al., 2019), such that net sensitivities will be determined by the dominant formation mechanism.
2 Methods

2.1 UWFPS Campaign Overview and Instrumentation

The UWFPS campaign included both aircraft and ground-based measurements throughout Salt Lake, Cache, and Utah Valleys during January and February 2017 (Figure 1). A total of 23 research flights were conducted during both day and night with the NOAA Twin Otter (TO) aircraft, equipped with aerosol and gas-phase instrumentation (summarized in Table 1) to probe the regional sources and formation mechanisms of PM$_{2.5}$. While flights were conducted over three valleys, the focus of this analysis will be on the more densely populated SLV, with relevant flight tracks highlighted in the right panel of Figure 1.

Briefly, the TO payload included gas-phase measurements of NO$_x$, NO$_2$, NO$_3$, and O$_3$ (1 Hz sample frequency) from a NOAA Cavity Ring Down Spectrometer (NOxCaRD) (Wild et al., 2014), NH$_3$ (1 Hz sample frequency) measurements from an Aerodyne mid infrared absorption instrument (QC-TIDLAS) from the University of Toronto (Ellis et al., 2010), and N$_2$O$_5$, HNO$_3$, and ClNO$_2$ (1 Hz sample frequency) measured with an iodide Time-of-Flight Chemical Ionization Mass Spectrometer (ITOF-CIMS) from the University of Washington (Lee et al., 2014; Lee et al., 2018). Accuracies for NO$_x$, NO$_2$ and O$_3$ were 5% and 12% for NO$_x$, with stated detection limits of 60 pptv (2σ) (Wagner et al., 2011; Wild et al., 2014) in the boundary layer. Gas-phase NH$_3$ was measured with a detection limit of 450 pptv (1s 3σ), as described in further detail by Moravek et al. (2019). Accuracy and detection limits for N$_2$O$_5$, CINO$_2$, and HNO$_3$ were similar to those reported from the same instrument deployed during the Wintertime INvestigation of Transport, Emissions, and Reactivity (WINTER) campaign (≤ 0.6 pptv (1σ)).

Non-refractory sub-micron aerosol composition (sampled every ~ 10 s) was measured with the NOAA Aerosol Mass Spectrometer (AMS) (Bahreini et al., 2009; Middlebrook et al., 2012) and aerosol size (sample every ~ 3 s) with a commercial Ultra-High Sensitivity Aerosol Spectrometer (UHSAS) (Brock et al., 2011). Average detection limits for AMS aerosol composition were 0.04, 0.09, 0.33, 0.03, and 0.07 μg sm$^{-3}$ (sm$^{-3}$ refers to m$^3$ under standard conditions (1 atm and 273.15 K)) for particulate nitrate, ammonium, organics, sulfate, and chloride, respectively. Uncertainties were ~20% for all species (Franchin et al., 2018). Ambient temperature and pressure (1 Hz sample frequency) were measured with a commercial (Avantech) meteorological probe. The accuracy of the commercial UHSAS instrument was also expected to be similar to that used during WINTER (dry surface area density: ~34%).

Additional ground-based measurements used in this analysis include hourly PM$_{2.5}$, NO$_x$, O$_3$, and temperature from the Utah Department of Air Quality (UDAQ) instrumentation at the Hawthorne (HW) monitoring site (Figure 1). Total PM$_{2.5}$ mass was measured with a Thermo Scientific 1405-DF Dichotomous Ambient Air Monitor, NO$_2$ with a Teledyne API T200U Chemiluminescence detector, and O$_3$ with a Teledyne API T400 UV absorption spectrometer, all in accordance with EPA guidelines (Environmental Protection Agency, 2018). Select volatile organic compounds (VOCs) were also measured at the University of Utah (UU) ground site by a Proton-Transfer Reaction Mass Spectrometer. Further information about the UWFPS campaign and aircraft and ground-based instrumentation can be found in additional publications (Franchin et al., 2018; UWFPS Science Team, 2018; Womack et al., 2019; Moravek et al., 2019).
2.2 Box Model

2.2.1 Description

A zero-dimension chemical box model has been developed to simulate the nocturnal chemical evolution of an air parcel from sunset until the time of aircraft measurement. Extensive model details have been previously discussed in McDuffie et al. (2018b). Briefly, the model forward-integrates the chemical mechanism (13 reactions, Table S1) starting 1.3 hours prior to sunset (see below), iteratively adjusting the initial concentrations of O₃ and NO₂, until the model-predicted concentrations of both are within 0.5% of the aircraft observations. Holding these initial concentrations constant, the model next adjusts the total heterogeneous loss rate constant of N₂O₅ (k_{N₂O₅}) and production rate constant of ClNO₂ (k_{ClNO₂}) until the model output simultaneously reproduces ambient nighttime observations of N₂O₅ and ClNO₂ to within 1%. The N₂O₅ uptake coefficients (γ(N₂O₅)) and ClNO₂ production yields (ψ(ClNO₂)) are then calculated following Eqs. (1) and (2), where c is the mean molecular speed and SA is the ambient wet PM₄ surface area density (described below). The model repeats this process every 10 seconds for all flights conducted at night, as determined by time and aircraft GPS altitude.

\[ γ(N₂O₅) = \frac{4 \times k_{N₂O₅}}{c \times SA} \]  
\[ ϕ(ClNO₂) = \frac{k_{ClNO₂}}{k_{N₂O₅}} \]  

Holding the derived \( k_{N₂O₅} \) and \( k_{ClNO₂} \) values constant, the model can further simulate the total nitrate produced overnight by forward-integrating the model until the time of sunrise, as shown for a representative SLV point in Figure 3. Here, total nitrate (gas + particulate-phase) is represented as HNO₃ only, as this model does not include aerosol thermodynamics that partition nitrate between the gas and particle phases. Modeled gas-phase HNO₃ is assumed to partition to the particle phase with 100% efficiency, following observations presented in Franchin et al. (2018) that show > 90% of total nitrate is in the particle phase during wintertime pollution events in SLV. As modeled nitrate is initialized with a concentration of 0 μg m⁻³, concentrations predicted at sunrise, therefore represent the total amount of nitrate produced from nocturnal chemistry over the course of a single night (i.e. nocturnal nitrate production rate). These base case values assume no overnight loss from dilution and constant values of \( γ(N₂O₅) \) and \( ϕ(ClNO₂) \), as discussed further in Section 3.3.3.

2.2.2 Model Simplifications and Uncertainties

For the UWFPS campaign, the box model was run in a similar manner to that described previously in McDuffie et al. (2018b), for nocturnal aircraft observations collected in the RL over the eastern U.S. coast during the 2015 WINTER campaign. Due to more limited instrumentation during UWFPS than WINTER, a larger number of box model assumptions and simplifications were required, which are summarized below.

First, the wet SA density for the base case simulations was calculated by applying a hygroscopic growth curve as a function of RH (Figure S2) to the dry PM₄ SA measured by the UHSAS (details in Section S1.3). The growth curve was derived with the Extended-AIM Aerosol Thermodynamic Model (Wexler and Clegg, 2002), assuming pure NH₄NO₃ particles.
Alternatively, estimating the growth factor from AMS organic measurements and estimates of aerosol organic density and hygroscopicity constant ($k_{org}$) (described in S1.3, (Jimenez et al., 2009; Mei et al., 2013; Cerully et al., 2015; e.g. Kuwata et al., 2012; Brock et al., 2016; Shingler et al., 2016)), resulted in only a ~3% change in the total wet SA for night flights during UWFPS (Figure S2a). For the 1031 measurement periods with simultaneous values of $\gamma(N_2O_5)$ and $\phi(ClNO_2)$, the median dry aerosol SA was 151 $\mu$m$^2$ cm$^{-3}$, which increased to 353 $\mu$m$^2$ cm$^{-3}$ when accounting for hygroscopic growth (Figure S2b). Additional uncertainties associated with aerosol SA are discussed below in Section 3.3.2 and only impact the $\gamma(N_2O_5)$ values reported here (not the nitrate production rates) as $k_{N_2O_5}$ and $k_{ClNO_2}$ are independent of the aerosol SA.

Second, loss of the nitrate radical (NO$_3$) from its reaction with volatile organic compounds (VOCs) was assumed to occur with a single first-order rate constant ($k_{NO_3}$), calculated for each flight from a combination of historical ground-based VOC measurements and select VOC measurements from a PTR-MS at the UU site (see Supplemental Section S1.2 for details; Atkinson and Arey (2003)). At night, NO$_3$ serves as one of the primary tropospheric oxidants for VOCs and can react with RO$_2$ and HO$_2$ radicals to contribute to nocturnal NO$_3$ recycling (Vaughan et al., 2006). In this analysis, NO$_3$-VOC reactions were lumped and treated as a net NO$_3$ sink with values of the first order loss rate constant, $k_{NO_3}$, ranging from 1.5×10$^{-3}$ - 9.5×10$^{-3}$ s$^{-1}$ (NO$_3$ lifetime ~100 – 1000 s). These rate constants are slightly larger than average values measured during the WINTER campaign (1.3×10$^{-4}$ to 4.6×10$^{-4}$ s$^{-1}$) (McDuffie et al., 2018b) and within the range previously reported (3×10$^{-3}$ to 1×10$^{-2}$ s$^{-1}$) during winter 2012 at a ground site in Colorado (Wagner et al., 2013). Additional NO$_3$-regeneration from reactions of NO$_3$ with HO$_2$ and RO$_2$ radicals were not included in this analysis due to a lack of radical measurements. An under-prediction in $k_{NO_3}$ from these uncertainties would cause both an over-prediction in the loss rate constant of N$_2$O$_5$, as well as the subsequent production of nitrate. While uncertainties in $k_{NO_3}$ can lead to large model uncertainties during summertime conditions (e.g. Phillips et al., 2016), NO$_3$-VOC reactivity is largely reduced during the winter season as a result of lower biogenic emissions and colder temperatures that favor N$_2$O$_5$ in its equilibrium with NO$_3$. Sensitivity studies discussed below showed 0.2% changes in the median model-predicted nocturnal nitrate production rate associated with ± 50% changes in $k_{NO_3}$ (Table S4). The possibility of varying VOC reactivity with time was also investigated (Section S1.4.5), but resulted in a minimal (<0.1%) impact on nitrate production results presented below. The potential for other rate constants to vary with time may additionally lead to increased variability in the results presented in Section 3.3.

Third are uncertainties in assumptions regarding the start time and duration of each simulation. All simulations were initialized at 1.3 hours prior to sunset, assuming no initial concentrations of N$_2$O$_5$ or ClNO$_2$. The pre-sunset time of 1.3 hours was derived for the WINTER campaign, based on the time when predicted daytime N$_2$O$_5$ concentrations (described in Section S1.4.4 and Brown et al. (2005)) diverged from ambient observations when approaching sunset. This value was not recalculated for UWFPS simulations as daytime N$_2$O$_5$ calculations require measurements of j(NO$_3$) photolysis rates, which were not measured during UWFPS. The median nocturnal nitrate production rate, however, changed by <0.3% when this pre-sunset time was varied between 0 and 2 hours. Photolysis rates during this time were also calculated from those measured during the WINTER campaign (Section S1.4.3; Schter and Müller (1999)). While WINTER photolysis rates may have been larger than those during Utah PCAP events, the median modeled nocturnal nitrate production rate showed a small sensitivity (<2.8%) to ± 40% changes in these values (Section S1.4.3). Additional uncertainties in air age (i.e. simulation start time and duration), however, may still serve to over-predict N$_2$O$_5$ loss rates and nocturnal nitrate based on previous sensitivity studies (McDuffie et al., 2018b). A combination of these assumptions will lead to a greater uncertainty in model results near sunset, as discussed in Section 3.3.2.
Fourth, air parcel mixing and deposition of gas-phase nitric acid were not included in base case simulations. Additional simulations, described in Section S1.4.2, included deposition using a first order nitric acid loss constant of 2.6×10^{-6} s^{-1}, calculated from a boundary layer height of 800 m, deposition velocity of 2.7 cm s^{-1} (Zhang et al., 2012), and gas/particle nitrate fraction of 8% from Franchin et al. (2018). The median nocturnal nitrate production rate increased by < 8% when this depositional loss of HNO_3 was included. In contrast, modeled nitrate production was most sensitive (-42.2% reduction) to the addition of a 1st order loss processes, meant to simulate air parcel dilution and O_3 entrainment from vertical mixing between the RL and free troposphere (Table S4). Based on a previous analysis by Womack et al. (2019), the dilution rate constant here was estimated to be 1.3×10^{-5} s^{-1} in the RL, with a possible range of 1.2-2.5×10^{-5} s^{-1} (described in Section S1.4.1). Results from the simulations that include dilution are discussed further in the final section.

Finally, the absolute uncertainty associated with each individual nocturnal nitrate production rate was calculated from the quadrature addition of the uncertainties associated with sensitivity tests described above and in Section S1.4, as well as uncertainties in the NO_2, O_3, N_2O_5, and ClNO_3 measurements used as model fit parameters (< 6% for all tests). Production rates derived from model fits to observations, as well as the absolute uncertainties associated with all 17 sensitivity tests are shown as a time series in Figure S3, with dilution contributing 92% of the total uncertainty (light blue in Figure S3) on average. Both the base case results (black dots) and those from simulations including the effects of air parcel dilution are discussed in Section 3.3.3.

3 Results and Discussions

3.1 PM_{2.5} in Salt Lake Valley – Winter 2017

To provide an overview of wintertime pollution events in SLV, Figure 4 shows a time series of total PM_{2.5} mass (1-hour and 24-hour averages) measured at the UDAQ Hawthorne (HW) site (Figure 1) from 1 December 2016 to 22 February 2017. Additional time series of ground-based PM_{2.5} measurements for all three Utah NAAs are provided in Franchin et al. (2018). The SLV data in Figure 4 show four pollution events that exceeded the NAAQS during the 2016-2017 winter. Calculated from 24-hour measurements, the four largest pollution events during December 2016 and January 2017 had daily PM_{2.5} build-up rates that ranged from 3.7 – 15.6 µg m^{-3} day^{-1} (see Figure 4), encompassing the daily rates reported previously in the same valley (Whiteman et al., 2014; Silcox et al., 2012; Baasandorj et al., 2017). The last two major pollution events (10 - 22 January (Event #3) and 25 January - 5 February (Event #4)) overlapped with the flights during UWFPS, shown by the gray shading in Figure 4. Average non-refractory (NR) PM_{1} aerosol mass fractions measured during these periods by the TO AMS showed that the aerosol was primarily composed of NH_4NO_3 (Figure 4 pie charts). The sum of NO_3^- and NH_4^+ contributed to 76.6% and 74.0% of the total PM_{1} mass measured during the last two pollution episodes (74% average (Franchin et al., 2018)), which agree with previous ground-based observations (e.g. Baasandorj et al., 2017) of past seasons. Nitrate alone contributed to an average 57% and 58% of the total aerosol mass during pollution episodes #3 and #4, respectively. During the relatively clean period sampled between 8 and 12 February 2017, the combined NH_4^+ + NO_3^- fraction decreased to an average of 57%, with a larger relative contribution from aerosol organics. The remaining analyses here will focus on aircraft flights during the two late January pollution events (#3 and #4) to evaluate the contribution of nocturnal RL heterogeneous nitrogen chemistry to observed surface-level nitrate during pollution events.
3.2 Limiting and Excess Reagents for NH₄NO₃ Aerosol

As NH₄NO₃ was the principle component of PM₂.₅ during pollution events in SLV (Figure 4), the contribution from heterogeneous reactive nitrogen processes is dependent on whether NH₄NO₃ formation is limited by the availability of gas-phase NH₃ or HNO₃. Under ambient conditions, gas-phase NH₃ and HNO₃ are assumed to be in a thermodynamic equilibrium with their particulate equivalents (NOₓ(p) and NH₄⁺(p)). The limiting reagent can therefore be inferred from the ratio of total oxidized (HNO₃(g) + NOₓ(p)) to total reduced nitrogen (NHₓ = NH₃(g) + NH₄⁺(p)), shown in Eq (3). This ratio does not account for other aerosol components such as (NH₄)₂SO₄, NH₄HSO₄, and NH₄Cl, but should generally represent the NH₄NO₃ aerosol system when particulate concentrations of sulfate and inorganic chloride are low, as was observed during UWFPS 2017 (Figure 4 and Franchin et al. (2018)). A nitrogen ratio greater than 1 indicates that oxidized nitrogen is in excess and NH₄NO₃ particle formation is limited by the presence of NH₃. Conversely, a ratio smaller than 1 indicates that formation is limited by the presence of HNO₃, which itself is limited by the oxidation rate of NOₓ. In a HNO₃-limited system, NH₄NO₃ formation will be sensitive to changes in HNO₃ concentrations resulting from both day and nighttime NOₓ oxidation processes. Daytime NOₓ oxidation rates during winter will depend on specific conditions but are generally slower, such that nighttime oxidation may play a dominant role (e.g. Wood et al., 2005; Kenagy et al., 2018).

\[ N_{Ratio} = \frac{HNO_3(g) + NO_x(p)}{NH_3(g) + NH_4^+(p)} \]  

A time series of nitrogen ratios in SLV between 17 January and 1 February is shown in Figure 5a, calculated from 10s averaged (AMS frequency) measurements of gas and particle-phase compounds aboard the TO aircraft. Figure 5a shows that NH₄NO₃ particle formation in SLV during pollution episodes was largely limited by HNO₃ (median ratio 0.77), but highly variable (range of 0.1 - 1.9) and time dependent, with the frequency of NH₃-limited conditions increasing throughout both late January pollution events. The color scale in Figure 5a and the vertical profiles of average and 10-90th percentile nitrogen ratios in Figure 5b further show that the lowest nitrogen ratios corresponded to the lowest altitudes. These results of HNO₃-limitation near the ground are consistent with all previous ground-based observations that show exclusive HNO₃-limitation in SLV (Kelly et al., 2013; Utah Division of Air Quality, 2014b). The increased frequency of NH₃-limited points throughout both pollution episodes (Figure 5a), however, is opposite the trend predicted by Baasandorj et al. (2017), who suggested that observed surface-level oxidant depletion should lead to more HNO₃-limited conditions over time. Events of NH₃-limitation (excess HNO₃) during 2017, however, only occurred at the highest altitudes (panel b) and their increasing frequency with time (panel a) is consistent with these events reflecting negative NH₃ gradients away from the surface and/or the production of HNO₃ aloft from nocturnal N₂O₅ chemistry. The rate of HNO₃ production from nocturnal heterogeneous chemistry is expected to be maximized at higher altitudes, removed from NO emissions and O₃ titration at the surface (Figure 2). Results here are also consistent with aerosol thermodynamic modeling studies by Franchin et al. (2018) who found that simulations of total PM₁ mass during UWFPS flights over SLV were proportionally sensitive to 50% reductions in total nitrate. Additional simulations, however, also resulted in near 50% PM₁ reductions with 50% reductions in total ammonium (NH₃+NH₄⁺) (Franchin et al., 2018), indicating that 50% ammonium reductions may be enough to shift SLV from the HNO₃ to NH₃-limited regime, consistent with nitrogen ratios in Figure 5 approaching and exceeding 1.
3.3 Nitrate Production via Heterogeneous Reactive Nitrogen Chemistry

The absolute amount of nitrate chemically produced from heterogeneous nitrogen chemistry will depend on the production rate of the nitrate radical and gas-phase N₂O₅ (Section 3.3.1), N₂O₅ aerosol uptake efficiency (Section 3.3.2), and yields of ClNO₂ and HNO₃ (Section 3.3.2), which are quantified below. The final section (Section 3.3.3) presents forward-integrated box model simulations to further quantify the nocturnal nitrate production rate and estimate the contribution of this chemistry to NH₄NO₃ formation during January 2017 in SLV.

3.3.1 Maximum Instantaneous Nitrate Production Rates

An upper limit estimate of the instantaneous rate of aerosol nitrate production from heterogeneous N₂O₅ chemistry, \( P_{N_2O_5^-} \), can be calculated as two times the gas-phase N₂O₅ production rate, \( P_{N_2O_5} \). These instantaneous production rates are calculated from the rate limiting reaction between NO₂ and O₃, which forms the nitrate radical (Eqs. (4) – (6)). In Eq. (4), \( P_{N_2O_5} \) is calculated in units of molec. cm⁻³ s⁻¹ but is typically reported in units of ppbv hr⁻¹ after conversion using the ambient air concentration (ND) and conversion factors for seconds to hours (3600) and mixing ratio to ppbv (1 × 10⁹). The reaction kinetics in Eq. (5) between NO₂ and O₃ are from the 2008 IUPAC recommendation (IUPAC, 2008). In Eq. (6), \( P_{NO_3^-} \) is then calculated as two times \( P_{N_2O_5} \) after it has been converted to units of \( \mu g \) m⁻³ hr⁻¹, as detailed in Supplemental Section S2. This calculation assumes: 1) N₂O₅ is produced quantitatively from NO₃ (i.e. no competing reaction of NO₃ + VOC), 2) N₂O₅ is produced at the rate of NO₂ production (valid under cold conditions that shift the NO₂-N₂O₅ equilibrium to favor of N₂O₅), 3) N₂O₅ is efficiently taken up onto aerosol, and 4) aqueous-phase reactions form two molecules of HNO₃ for every molecule of N₂O₅ (i.e. \( \phi(CINO_2) = 0 \)).

\[
P_{N_2O_5} \text{[ppbv hr}^{-1}] = \frac{k_d[O_2][NO_2]}{[N_D] \text{[molec. cm}^{-3}]} \times 3600 \text{[s hr}^{-1}] \times 1 \times 10^9 \text{[ppbv]}
\]

\[
k_4 \text{[cm}^3\text{molec}^{-1} \text{s}^{-1}] = 1.4 \times 10^{-13}e^{(-2470/T)}
\]

\[
P_{NO_3^-} \text{[} \mu g \text{m}^{-3} \text{hr}^{-1}] = 2 \times \left( P_{N_2O_5} \text{[} \mu g \text{m}^{-3} \text{hr}^{-1}] \right)
\]

The value of \( P_{NO_3^-} \) is expected to vary with altitude due to boundary layer dynamics and surface NO₃ emissions that can deplete O₃ at night near the surface, as described previously in Baasandorj et al. (2017). The time series in Figure 6a illustrates that the O₃ measured at HW was frequently absent at night during the 3rd and 4th pollution events in January 2017. As surface-level O₃ was titrated overnight, ground-site data cannot provide direct information about \( P_{NO_3^-} \) aloft in the RL. In the absence of vertical observations during pollution events in 2016, a previous analysis by Baasandorj et al. (2017) used late afternoon measurements at the HW ground site to predict N₂O₅ production rates (\( P_{N_2O_5} \)) in the RL that varied from 0 up to ~ 2 ppbv hr⁻¹ (~ 0 – 5 \( \mu g \) m⁻³ hr⁻¹), but with values frequently < 1 ppbv hr⁻¹. These values correspond to instantaneous nitrate production rates (\( P_{NO_3^-} \)) of ~ 0 – 10 \( \mu g \) m⁻³ hr⁻¹, with typical values closer to 5 \( \mu g \) m⁻³ hr⁻¹. Late afternoon estimates from the same site during 2017 (dashed lines in Figure 6, from Eq. (6)), suggest smaller \( P_{NO_3^-} \) rates in 2017 than in 2016, with values between 1 and 5 \( \mu g \) m⁻³ hr⁻¹ during UWFPS pollution events (Figure 6a).

The bottom panels of Figure 6b show the binned, vertical profiles of median, 25th, and 75th percentile instantaneous \( P_{NO_3^-} \) values, along with aircraft observations of O₃, NO₂, and PM₁ for all UWFPS night flights (red shaded regions in Figure 6a). The vertical profiles show a relatively uniform distribution of \( P_{NO_3^-} \) with altitude through the lowest 600 m. The dashed black lines
also show that the number of points in each altitude bin was weighted toward the 100-500 m altitude range. The median instantaneous $P_{NO_2^{-}}$ value in this polluted layer (0-600 m AGL) was 1.6 µg m$^{-3}$ hr$^{-1}$ ($N = 21666$). This value is at the low range of estimates of 1.6 - 5 µg m$^{-3}$ hr$^{-1}$ that are predicted from late afternoon ground-based observations on each flight day (dashed line in the middle panel of Figure 6a), following the method of Baasandorj et al. (2017).

Vertical profiles in Figure 6b do not show evidence for a reduction in $P_{NO_2^{-}}$ or $O_3$ near the surface, as is expected for $O_3$ titration near the ground level (shown in panel a). The distribution in panel b, however, is affected by the location of the missed approaches / landings in the SLV (Salt Lake International and South Valley Regional airfields), which are further from the urban center of Salt Lake City than the HW ground site (see Figure 1). Vertical profiles to the surface over urban Salt Lake City were not possible due to a lack of airfields for missed approaches. Instead, the SLV flights often executed box patterns over the eastern Salt Lake basin at several altitudes. Figure 7 shows the vertical distribution of $P_{NO_2^{-}}$ values from these boxes on January 28 - 29 between 21:20 – 00:30 local time, compared to $P_{NO_2^{-}}$ measured at surface level during the same interval. At 300 and 500 m AGL, the median (and interquartile range) $P_{NO_2^{-}}$ was 2.2 (2.1 to 2.4) and 1.9 (1.8 to 2.1) µg m$^{-3}$ hr$^{-1}$, while at 650 m, slightly above the most concentrated pollution layer, it was 0.5 (0.3 to 0.7) µg m$^{-3}$ hr$^{-1}$. The median value at the HW ground site, directly below the aircraft, was 0.02 (0.01 to 0.2) µg m$^{-3}$. These plots demonstrate that $P_{NO_2^{-}}$ is typically low or zero at night near the surface within the urban area of Salt Lake City, but large within the RL. Away from the urban area, the vertical distributions of $P_{NO_2^{-}}$ are also likely more uniform (Figure 6b) due to the lack of $O_3$ titration within the nocturnal boundary layer. In the final section below, nightly integration of these instantaneous $P_{NO_2^{-}}$ values are compared to box model predictions of total nitrate.

### 3.3.2 Modeled Uptake Coefficients and Production Yields

Both the aerosol uptake efficiency of $N_2O_5$ ($\gamma(N_2O_5)$) and the production yield of CINO$_2$ ($\phi(CINO_2)$) are highly variable, dependent on aerosol composition, and can impact the absolute amount of nitrate formed from nocturnal heterogeneous nitrogen chemistry. The nighttime formation of nitrate, however, is only limited by these processes when $N_2O_5$ uptake is inefficient and is instead limited by the oxidation rate of NO$_2$ ($R_1$) (discussed above) at sufficiently large values of $\gamma(N_2O_5)$.

As described in Section 2.2, an iterative box model was fit to observations of NO$_2$, $O_3$, $N_2O_5$, and CINO$_2$ to quantify $\gamma(N_2O_5)$ and $\phi(CINO_2)$ during pollution events. For SLV alone ($N = 1030$), the distribution in Figure 8 shows that $\gamma(N_2O_5)$ values ranged four orders of magnitude from $1 \times 10^{-1}$ to $>1$. Values approaching or exceeding 1 are unphysical and suggest artifacts in the $\gamma(N_2O_5)$ determinations for UWFPS (see below), at least for the largest values. Values of $\phi(CINO_2)$ encompassed the entire possible range of 0 to 1 (Figure 8). The medians for this subset were 0.076 and 0.220 for $\gamma(N_2O_5)$ and $\phi(CINO_2)$, respectively. For all UWFPS flights between 16 January and 1 February 2017, the median $\gamma(N_2O_5)$ and $\phi(CINO_2)$ values in the RL ($N = 2195$) were 0.049 and 0.256, respectively, derived from box-model fits to observations. These values are compared to multiple derivation methods further below.

Compared to previous studies, the median $\gamma(N_2O_5)$ over SLV was twice as large as the mode derived with a similar model using data from the Nitrogen, Aerosol Composition, and Halogens on a Tall Tower (NACHTT) campaign near Denver, Colorado in winter 2011 (Wagner et al., 2013). Similarly, the median was over 5 times larger than the median calculated using the same model from the 2015 WINTER campaign (McDuffie et al., 2018b). The largest values during UWFPS exceeded those from both WINTER and NACHTT studies, while the smallest values were also larger than either of the respective minimums. The two most common suppression mechanisms that lead to reductions in $\gamma(N_2O_5)$ are associated with the presence of organic material and nitrate
in the aerosol phase. Insoluble aerosol organics have been shown to suppress N\textsubscript{2}O\textsubscript{5} uptake in previous laboratory studies (e.g. Griffiths et al., 2009; Thornton et al., 2003; McNeill et al., 2006; Thornton and Abbatt, 2005; Cosman et al., 2008; Badger et al., 2006; Folkers et al., 2003) and large organic mass fractions have been associated with \(\gamma(\text{N}_2\text{O}_5)\) reductions in past field studies (Bertram et al., 2009; McDuffie et al., 2018b). The average dry mass fraction of aerosol organics (i.e. organic mass / total dry aerosol mass) during SLV pollution events was less than half of that observed during the WINTER campaign (~18% vs 40%) and 40% lower than the average during NACHTT (27%, (Wagner et al., 2013)). Aerosol nitrate can also suppress uptake as soluble nitrate facilitates the reformation of gas-phase N\textsubscript{2}O\textsubscript{5} (Bertram and Thornton, 2009; Griffiths et al., 2009), and nitrate mass fractions have been negatively correlated with \(\gamma(\text{N}_2\text{O}_5)\) in previous field-studies (Wagner et al., 2013; Morgan et al., 2015; Riedel et al., 2012; Bertram et al., 2009; McDuffie et al., 2018b). The presence of sufficient aerosol water, however, can offset this nitrate suppression by promoting N\textsubscript{2}O\textsubscript{5} aqueous solvation and reaction (e.g. Bertram and Thornton, 2009; Griffiths et al., 2009; Mentel et al., 1999; Wahner et al., 1998), resulting in increases in \(\gamma(\text{N}_2\text{O}_5)\) with the ratio of water to nitrate (McDuffie et al., 2018b). The average dry mass fraction of aerosol nitrate was much larger during UWFPS (60%) than during NACHTT (30%, Wagner et al. (2013)) or WINTER (15%, McDuffie et al. (2018b)). High humidity conditions during UWFPS (77% average RH during pollution events) resulted in average aerosol water mass fractions (i.e. water mass / aerosol dry mass + water mass) near 70%, as calculated with an aerosol thermodynamic model, described in Franchin et al. (2018). This higher RH likely contributed to efficient N\textsubscript{2}O\textsubscript{5} uptake during UWFPS despite the presence of aerosol nitrate. In fact, the largest 25% of UWFPS \(\gamma(\text{N}_2\text{O}_3)\) values exceed the largest value (0.175) that has been reported from recent field studies (Figure 4 in McDuffie et al. (2018b)).

The median \(\phi(\text{ClNO}_2)\) value of 0.220 during SLV pollution events was 4 times larger than during the NACHTT campaign (Riedel et al., 2013; Wagner et al., 2013), but within a factor of 2 larger than the median derived during WINTER over the U.S. east coast (McDuffie et al., 2018a). The SLV median was also similar to medians reported from previous ground-based studies across North America (Mielke et al., 2016; Mielke et al., 2011; Mielke et al., 2013; Wagner et al., 2012; Thornton et al., 2010). Heterogeneous CINO\textsubscript{2} production requires aerosol chloride (R6) (e.g. Behnke et al., 1997) and though a consistent geographic pattern in \(\phi(\text{ClNO}_2)\) has not emerged from past studies (Figure 2 in McDuffie et al. (2018a)), heterogeneous chemistry in the vicinity of the Great Salt Lake appears to produce CINO\textsubscript{2} with the same efficiency as comparable measurements near North American ocean coastlines. CINO\textsubscript{2} production yields, however, remain smaller than those predicted based on measured aerosol composition, as discussed below.

While large \(\gamma(\text{N}_2\text{O}_3)\) and moderate \(\phi(\text{ClNO}_2)\) values indicate efficient nitrate production from heterogeneous chemistry during UWFPS, these values may be upper and lower limits, respectively. As discussed in Section 2.1, limited observations of VOC and photolysis rates, as well as uncertainties in air age, and dilution may cause the \(k_{\text{N}_2\text{O}_3}\) and \(k_{\text{ClNO}_2}\) values (and subsequent \(\gamma(\text{N}_2\text{O}_3)\) and \(\phi(\text{ClNO}_2)\)) to be over- and under-predicted, respectively. This is more likely near sunset where the model has an increased sensitivity to assumptions in simulation start time (McDuffie et al., 2018b). Uncertainties in gas-phase measurements may also contribute to uncertainties in the model predictions, though the level of uncertainty associated with these parameters is small (Table S4). As summarized in Table S4, the box model is not highly sensitive to most sources of uncertainty, and the model-derived values of \(k_{\text{N}_2\text{O}_3}\) are consistent with those derived from observations (discussed below).

Independent of the model fits of \(k_{\text{N}_2\text{O}_3}\) and \(k_{\text{ClNO}_2}\), unphysically large \(\gamma(\text{N}_2\text{O}_3)\) values (> 0.1 in Figure 8) may alternatively be an artifact arising from under-measurement of ambient aerosol SA. Low aerosol SA would bias high the \(\gamma(\text{N}_2\text{O}_3)\) calculation in Eq. (1) without influencing the model derivations of \(k_{\text{N}_2\text{O}_3}\) and \(k_{\text{ClNO}_2}\). In this study, wet aerosol SA was calculated as described.
above by applying a relative humidity-dependent growth factor curve to the measured dry PM$_1$ SA. Despite large concentrations of total dry SA (Figure S2), an under-prediction in the wet SA could arise from uncertainties in the hygroscopic growth curve or additional unmeasured SA from large particles (≥ 1 μm). Both factors would be exacerbated by the high humidity conditions encountered during UWFPS since large, hydrated particles would not be sampled efficiently by the aerosol inlet and hygroscopic growth curves are highly uncertain above ~95% RH (corresponding to 6.7% of the SLV data). A third possible cause of under-measured SA is the presence of fog under high humidity conditions. Fog is well known to promote rapid heterogeneous processes (Lelieveld and Crutzen, 1990), and is associated with surface areas that can be orders of magnitude larger than accumulation mode aerosol. For example, fog has been demonstrated to lead to rapid N$_2$O$_5$ loss at a ground site in Hong Kong, during November - December 2013 (Brown et al., 2016). It is therefore possible that unmeasured SA under high humidity conditions could bias the calculated γ(N$_2$O$_5$) values high relative to values reported in previous literature. Any bias caused by aerosol SA, however, would not impact the model-derived $k_{N_2O_5}$ and $k_{ClNO_2}$ values that are used in the final analysis below.

To further evaluate the UWFPS γ(N$_2$O$_5$) and ϕ(ClNO$_2$) values, box model determinations are compared to two other derivation methods in Figures 8 and S5. The first method calculates γ(N$_2$O$_5$) from NO$_2$, O$_3$, and N$_2$O$_5$ observations, based on the steady state approximation (γ(N$_2$O$_5$)$_{ss}$), described by Brown et al. (2003) and defined in Supplemental Section S4.1. This method shows excellent agreement with box model results (Figure 8 and S5). The steady state method has been shown in previous analyses to over-predict γ(N$_2$O$_5$) values under cold, high NO$_2$ conditions, but only if the first order rate constants for NO$_2$ and N$_2$O$_5$ loss ($k_{NO_2}$ and $k_{N_2O_5}$) are modest (Brown et al., 2003). Both the steady state and box model γ(N$_2$O$_5$) values are consistent with a rapid first order loss constant of N$_2$O$_5$ (median $k_{N_2O_5}$ = 1.1×10$^{-3}$ s$^{-1}$), suggesting the steady state approach is valid for SLV conditions. The corresponding median lifetime (1/$k_{N_2O_5}$) of 14 minutes is, for example, much shorter than the lifetimes of 2-18 hours calculated from a previous steady state analysis of aircraft measurements over Texas in fall 2006 (Brown et al., 2009). Nevertheless, the color scale in Figure S5 shows that the largest γ(N$_2$O$_5$) values (≥ 0.1) were exclusively derived for air sampled within 3 hours of sunset (4.3 hr simulation time), where previous analysis has shown the steady state approximation to be least reliable. As Figure S5 shows large γ(N$_2$O$_5$) determinations from both the box model and the steady state analysis during this time, there may be a common bias between the methods if these values are indeed too large.

The second method calculates both γ(N$_2$O$_5$) and ϕ(ClNO$_2$) using laboratory-based parameterizations by Bertram and Thornton (2009) (BT09), based on aerosol volume-to-surface area ratio, N$_2$O$_5$ solubility (Fried et al., 1994), aerosol molarities of water, nitrate, and chloride (calculated as described in Section S4.2), and laboratory-derived reaction rate constant ratios. Further details of each parameterization are provided in Supplemental Section S4.2. These parameterizations have had mixed success in reproducing previous field-derived values (e.g. Bertram et al., 2009; Riedel et al., 2012; McDuffie et al., 2018b; McDuffie et al., 2018a), but are commonly used to predict N$_2$O$_5$ uptake and ClNO$_2$ production on internally-mixed inorganic aerosol when N$_2$O$_5$ chemistry is included in global models (e.g. Sarwar et al., 2014; Shah et al., 2018; Wang et al., 2018).

Results in Figure S5 show that the median γ(N$_2$O$_5$) value predicted by the BT09 parameterization is within a factor of 2 of the box model median, but that this parameterization does not reproduce the observed variability (Figures 8). For ϕ(ClNO$_2$), the BT09 parameterization largely over-predicts model-derived values with a median of 0.66 relative to the model median of 0.22 (Figure S5). This over-prediction is consistent with all previous studies to compare parameterized and field-derived ϕ(ClNO$_2$) results (Wagner et al., 2013; Wang et al., 2017b; Ryder et al., 2015; Thornton et al., 2010; Riedel et al., 2013; Wang et al., 2017a; Tham et al., 2018; McDuffie et al., 2018a). These results also suggest that the parameterization would need to be reduced by 68%
for agreement with the box model median, similar to the 74-85% reduction required for agreement of this parameterization with the WINTER campaign median (McDuffie et al., 2018a). The possible presence of additional, refractory-phase chloride (i.e. NaCl, CaCl\textsubscript{2}, and KCl) in the accumulation mode would increase the predicted \(\gamma(N_2O_3)\) and improve agreement with the box model, but would further degrade the agreement of \(\phi(CINO_2)\).

Lastly, the empirically-based \(\gamma(N_2O_3)\) parameterization from McDuffie et al. (2018b) was applied to UWFPS data, though only an estimated range for the campaign median is presented here due to uncertainties in the aerosol O:C ratio and aerosol organic density, both required for this calculation (discussed in Section S4.2). This parameterization models N\textsubscript{2}O\textsubscript{3} uptake onto an aqueous inorganic particle with a resistive organic coating, with a thickness determined by the volume ratio of inorganic to total aerosol components (McDuffie et al., 2018b; Riemer et al., 2009; Anttila et al., 2006). By estimating a range of O:C ratios using the improved-ambient O:C ratio method from Canagaratna et al. (2015) and AMS organic m/z 44 fraction (Figure 6, Franchin et al. (2018)), assuming an organic density of 1.3 g/cm\textsuperscript{3} (e.g. Kawata et al., 2012) to estimate the organic-associated volume, and applying additional constants described in Section S4.2, this parameterization estimated a median \(\gamma(N_2O_3)\) between 60 and 85% lower than the box model. Though there are large uncertainties in the required parameters, these results suggest that during pollution events: 1) aerosol organics are not surface active, 2) aerosol organics are not resistive toward N\textsubscript{2}O\textsubscript{3}, or 3) box model \(\gamma(N_2O_3)\) values are over-predicted due to missing SA (e.g. fog, Section 3.3.2) or other simplifying assumptions (e.g. dilution) discussed above.

Despite disagreement between the box model and parameterizations, the \(\gamma(N_2O_3)\) values predicted by all three methods are large enough, in combination with the large measured aerosol SA, to fall within the range where models of nighttime chemistry are insensitive to variation in uptake efficiency (e.g. Macintyre and Evans, 2010; Riemer et al., 2003). Under these conditions, the NO\textsubscript{2} gas-phase oxidation rate (i.e. \(P_{NO_2}^-\)) becomes the limiting factor to HNO\textsubscript{3} formation relative to N\textsubscript{2}O\textsubscript{3} uptake. As further evidence, the median lifetime of NO\textsubscript{2} with respect to O\textsubscript{3} (\(\tau_{NO_2}^- = 1/(k_3[O_3])\)) was 9 hours during pollution events in SLV, while the equivalent lifetime of N\textsubscript{2}O\textsubscript{3} (\(\tau_{N_2O_3} = 1/k_{N_2O_3}\)) was 14 minutes. Further to this point, explicit box modeling of day and nighttime chemical processes during UWFPS by Womack et al. (2019) showed that the production of O\textsubscript{4,\textsubscript{max}} (\(= NO_2 + O_3 + 2*NO_2 + 1.5*(HNO_3 + particulate nitrate) + CINO_2 + 3*N_2O_3 + others\)) was insensitive (<1.5%) to order-of-magnitude changes in \(\gamma(N_2O_3)\). Short lifetimes of N\textsubscript{2}O\textsubscript{3} relative to NO\textsubscript{2}, as well as nitrate insensitivity to \(\gamma(N_2O_3)\), both indicate that nocturnal heterogeneous chemistry contributes to NH\textsubscript{4}NO\textsubscript{3} formation, but that absolute production is limited by gas-phase kinetics rather than aerosol composition and \(\gamma(N_2O_3)\). This insensitivity to \(\gamma(N_2O_3)\) provides confidence in the ability of the box model to predict the magnitude of nocturnal nitrate production in SLV, regardless of uncertainties in \(\gamma(N_2O_3)\).

### 3.3.3 Modeled Nocturnal Nitrate Production Rates and Contribution of Heterogeneous Chemistry to Total NH\textsubscript{4}NO\textsubscript{3} Aerosol Accumulation Rates

As described in Section 2.2 and shown in Figure 3, the box model simulates the amount of total nitrate (HNO\textsubscript{3} + NO\textsubscript{3}) produced from heterogeneous chemistry over the course of a single night. This amount of nitrate, in units of \(\mu g\ m^{-3}\) night\textsuperscript{-1}, is in addition to any nitrate present at sunset from the previous day (e.g. Figure 3). Figure 9 shows the distribution of nightly nitrate production predicted by base case simulations (\(N = 1033\)), with a median of 9.9 \(\mu g\ m^{-3}\) nitrate night\textsuperscript{-1}.

Comparisons between the base case results and integrated \(P_{NO_2}^-\) from Section 3.3.1 also suggest that nocturnal nitrate production is limited by the rate of NO\textsubscript{2} oxidation rather than the efficiency of N\textsubscript{2}O\textsubscript{3} aerosol uptake. Based on the calculations in
Section 3.1, upper-limit \( P_{\text{NO}_3^-} \) values, integrated over an average 14 hour night and reduced to account for a \( \phi(\text{CINO}_2) \) value of 0.2, ranged from < 0.5 to > 40 \( \mu g \) m\(^{-3}\) night\(^{-1}\), with a median of 20.2 \( \mu g \) m\(^{-3}\) night\(^{-1}\) (\( N = 21666 \)). To more directly compare with box model results, the subset of points with simultaneous \( \gamma(\text{N}_2\text{O}_3) \) determinations had a median of 10.6 \( \mu g \) m\(^{-3}\) night\(^{-1}\), which is slightly larger, but agrees well with the box-model predicted median of 9.9 \( \mu g \) m\(^{-3}\) night\(^{-1}\). As described in Section 3.3.1, the \( P_{\text{NO}_3^-} \) calculation assumes efficient \( \text{N}_2\text{O}_5 \) uptake and only considers nitrate production to be limited by gas-phase kinetics. Observed agreement between the integrated \( P_{\text{NO}_3^-} \) values and box model-predicted production rates, therefore suggests that nitrate production may be largely limited by gas-phase oxidation rather than multi-phase processes.

Uncertainties associated with base case production rates are discussed in Section 2.2.2 and shown as a time series in Figure S3. Air parcel dilution associated with vertical mixing was the largest source of uncertainty (Table S4, Figure S3). This process was not included in base case simulations, though mixing / dilution has been observed and predicted in an analysis of WINTER nighttime flights (Kenagy et al., 2018; McDuffie et al., 2018b). Estimating the impact of dilution by including a single first order dilution rate constant \( k_{\text{dilution}} \) of 1.3\( \times \)10\(^{-5} \) s\(^{-1}\) reduced the median nocturnal nitrate production rate by 42% to 5.7 \( \mu g \) m\(^{-3}\) night\(^{-1}\), shown in comparison to base case simulations in Figure 9. This \( k_{\text{dilution}} \) rate constant was derived, as described in Womack et al. (2019), by fitting a box model to best reproduce the build-up of \( O_{\text{iso}} \), observed between 28 January and 1 February at the UU ground site in SLV. Following Womack et al. (2019), this entrainment rate constant of 8\( \times \)10\(^{-6} \) s\(^{-1}\) was then scaled up by 40% to represent the reduced volume of the nocturnal RL relative to the mixed daytime boundary layer, for which this rate constant was derived. The resulting \( k_{\text{dilution}} \) of 1.3\( \times \)10\(^{-5} \) s\(^{-1}\) is ~60% lower than \( k_{\text{dilution}} \) from the WINTER campaign, derived from observations of \( \text{NO}_x \) (\( \equiv \text{NO} + \text{NO}_2 + \text{NO}_3 + 2\text{N}_2\text{O}_5 + \text{CINO}_2 + \text{RONO}_2 \ldots \) ) overnight in a single RL air parcel over the eastern U.S. coast (McDuffie et al., 2018b). As processes relevant to RL dilution were not directly measured during UWFPS, there are uncertainties associated with this \( k_{\text{dilution}} \) estimation. For instance, based on the modeled surface albedo in Womack et al. (2019), \( k_{\text{dilution}} \) could have reproduced observed \( O_{\text{iso}} \), mixing ratios with scaled values ranging between 1.2\( \times \)10\(^{3} \) and 2.5\( \times \)10\(^{-5} \) s\(^{-1}\) (Figure S10, Womack et al., 2019). This particular range of loss rate constants predicts median nitrate production rates in SLV between 3.6 and 6.1 \( \mu g \) m\(^{-3}\) night\(^{-1}\).

Modeled nitrate production rates are further compared in Figure 10 to the average daily accumulation of surface-level nitrate aerosol during pollution event #4 at the HW ground site. This ground-based accumulation rate (red diamond in Figure 10a) was taken as the slope of the 24-hour average PM\(_{2.5}\) observations at HW (scaled by 0.58; average \( \text{NO}_3^- \) fraction from Figure 4) during the first six days of event #4, before it began to degrade on 1 February 2017 (Figure 10b). Only data from event #4 data are here assessed as this was the only PCAP sampled with the aircraft on multiple nights. Figure 10a shows this average, 24-hour surface accumulation rate of 4.6 \( \mu g \) m\(^{-3}\) day\(^{-1}\) (red diamond) compared to the 10\(^{th}\) – 90\(^{th}\) percentile distributions, medians, and averages of the nocturnal production rates predicted by base case box model simulations (gray) and simulations including the effects of 24-hour dilution (blue), described below.

Comparing modeled RL chemical nitrate production to the observed ground-based accumulation rate can provide an estimate for the \( \text{N}_2\text{O}_5 \) uptake contribution to total particulate nitrate production in SLV. Direct comparison is difficult, however, as the 24-hour ground-based accumulation rate includes contributions from night- and daytime chemical production, but also depends on dilution and mixing processes. For example, the amount of nocturnally produced nitrate observed at the surface will depend on mixing of nitrate aerosol to the surface from the RL during morning boundary layer expansion (Figure 2). As a result, base case box model predictions (no dilution consideration) in Figure 10a (gray) had a median of 8.6 \( \mu g \) m\(^{-3}\) night\(^{-1}\), nearly twice
as large as the observed, 24-hour average ground-based accumulation rate. Therefore, to more directly compare box model predictions and ground-based observations, Figure 10a also shows the results from simulations that included loss from both nocturnal and daytime dilution. At night, $k_{\text{dilution}}$ values of $1.2 \times 10^{-5}$ s$^{-1}$ (L), $1.3 \times 10^{-5}$ s$^{-1}$ (M), and $2.5 \times 10^{-5}$ s$^{-1}$ (H) (blue) were applied to all modeled species as described above. Modeled nighttime nitrate (e.g. Figure 9) was then further diluted for ~10 hours (24 – 14 night) at 60% of the nocturnal dilution rate ($k_{\text{dilution}} = 0.6 = 8 \times 10^{-6}$ s$^{-1}$) to reflect the increased volume of the daytime boundary layer, following Womack et al. (2019). For a single 24-hour period, this resulted in a net median of 3.9 µg m$^{-3}$ nitrate produced from nocturnal heterogeneous N$_2$O$_5$ uptake, with a range of medians between 1.9 and 4.2 µg m$^{-3}$ day$^{-1}$ when considering the extended range of dilution rate constants from Womack et al. (2019).

Comparison of modeled rates to the observed, daily surface build-up of 4.6 µg m$^{-3}$ day$^{-1}$, suggests that on average, nitrate produced from heterogeneous chemistry can generally account for the nitrate accumulation observed at the surface. This result is qualitatively consistent with an observational analysis by Pusde et al. (2016), who determined that nocturnal heterogeneous chemistry was the main source of regional aerosol nitrate during wintertime pollution events in the San Joaquin Valley. A box model analysis of this same event by Womack et al. (2019), however, also showed that photochemical nitrate production is also occurring during these events with roughly equal contributions between photochemical and nocturnal nitrate production pathways. Therefore, while results in Figure 10a (including dilution) predict a median nocturnal fractional contribution of 86% (ranging between 42 and 91%), confirmation and further quantification of this result will require additional, vertically resolved measurements of aerosol composition, gas-phase precursors, and physical parameters, as well as more sophisticated modeling of these multi-day pollution accumulation events with 3D-chemical transport models.

4 Summary and Conclusions

Aerosol and gas-phase measurements collected during the 2017 UWFPS campaign showed multiple pollution events that exceeded PM$_{2.5}$ standards in SLV, the most populated region in the state of Utah. During these events, aerosol particles were largely composed of NH$_4$NO$_3$, which forms from the reaction between gas-phase NH$_3$ and HNO$_3$. While NH$_3$ is emitted from surface sources, HNO$_3$ is chemically formed from the oxidation of NO$_x$ emissions. This oxidation can occur through daytime reactions with the photochemical OH radical, or nocturnal heterogeneous reactions involving NO$_3$ and N$_2$O$_5$. The contribution of nocturnal chemistry to PM$_{2.5}$ formation in SLV is dependent on whether NH$_4$NO$_3$ formation is NH$_3$- or HNO$_3$-limited, as well as the NO$_3$ production rate, N$_2$O$_5$ uptake efficiency, CINO$_2$ and HNO$_3$ production yields, and loss processes such as air parcel dilution.

Vertically resolved measurements of gas and particulate phase oxidized and reduced nitrogen in SLV showed that NH$_4$NO$_3$ formation during pollution events was nearly always HNO$_3$ limited, but that oxidized and reduced nitrogen approached equivalence as pollution events progressed. This reagent balance analysis is consistent with aerosol thermodynamic modeling presented in Franchin et al. (2018), which predicted that all three major valleys in Wasatch region were sensitive to nitrate reductions, and that SLV was also sensitive to NH$_3$ reductions. Both observation and modeling-based analyses agreed that NH$_4$NO$_3$ formation in the RL was largely HNO$_3$-limited during pollution events, providing the possibility of a large contribution from nocturnal heterogeneous chemistry to HNO$_3$ and PM$_{2.5}$ mass.

Analysis of vertically-resolved, calculated nitrate production rates (an upper-limit estimate due to heterogeneous HNO$_3$ formation, $P_{\text{NO}_3^-}$) and results from an observationally-constrained chemical box model, suggest that nocturnal chemistry is an efficient mechanism for PM$_{2.5}$ production in SLV during pollution events. Nitrate production rates had a median of 1.6 µg m$^{-3}$ hr$^{-1}$.
\[ \gamma(N_2O_3) \text{ and } \phi(ClNO_2) \] had medians of 0.076 and 0.220, respectively, during pollution events. Values of \( \gamma(N_2O_3) \) were larger than previous field-based determinations (e.g. McDuffie et al., 2018b) and those predicted from the Bertram and Thornton (2009) parameterization, but were in agreement with values derived using the \( N_2O_3 \) steady state approach. The median \( \phi(ClNO_2) \) value was larger than that derived from aircraft observations over the eastern US coast, but were simultaneously overpredicted by 68\% by the Bertram and Thornton (2009) parameterization, which uses measurements of aerosol chloride and water estimations.

While the box model has uncertainties associated with limited available measurements and model assumptions, the large measured aerosol SA, efficient \( N_2O_3 \) uptake coefficients, and moderate \( ClNO_2 \) yields resulted in nightly modeled nitrate production rates that were largely insensitive to specific values of derived parameters. Agreement between base case modeled nightly nitrate production (9.9 \( \mu g \) m\(^{-3}\) night\(^{-1}\)) and that calculated from \( P_{N_2O_3^-} \) values (10.6 \( \mu g \) m\(^{-3}\) night\(^{-1}\)) alternatively suggests that nitrate production is more sensitive to gas-phase \( NO_2 \) oxidation rates than \( \gamma(N_2O_3) \), providing confidence in the model’s predictions of nocturnal nitrate. Of the parameters tested, the model was most sensitive to loss through air parcel dilution, with a 42\% reduction to 5.2 \( \mu g \) m\(^{-3}\) nitrate night\(^{-1}\) when including a nocturnal \( k_{dilution} \) rate constant of 1.3\times10\(^5\) s\(^{-1}\). When considering the possible effects of 24-hour dilution, model simulations predicted a reduced median of 3.9 \( \mu g \) m\(^{-3}\) nitrate day\(^{-1}\), corresponding to 86\% (median) of the net aerosol nitrate accumulation that was observed at a SLV ground site. Due to model uncertainties and sensitivities to dilution, further quantification of this result will require additional vertically-resolved measurements and photochemical / 3D modeling analyses. These results however, highlight the importance of nocturnal chemistry in the formation of \( PM_{2.5} \) in SLV and can provide constraints for regulatory models of \( PM_{2.5} \), used to assess control strategies in this populated non-attainment area.

**Author Contributions**

During the UWFPS campaign, EEM, CCW, DLF, and WPD were responsible for the CRD gas-phase measurements, AF and AM for the AMS particle measurements, LG, BHL, and JAT for the I-TOF-CIMS measurements, and AM and JM for the \( NH_3 \) instrument. MB and SSB organized the UWFPS campaign with technical support from WPD. EEM developed the box model code and preformed the analyses with support from CCW and SSB. EEM prepared the manuscript with contributions from co-authors.

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Figure 1. (Left) Elevation Map of Utah’s Wasatch region (Utah State in insert), with the Great Salt Lake (north) and Utah Lake (south) shown in blue and county borders in black. U.S. EPA designated non-attainment areas (NAA) for PM$_2.5$ are shown by red boundaries. From north to south these NAAs include the Logan NAA: “Moderate” status, Salt Lake City NAA: “Serious” status, and Provo NAA: “Serious” status. UWFPS TO flight tracks are shown in pink. Purple markers indicate the locations of major cities, including Logan in Cache Valley, Ogden and Salt Lake City in SLV, and Provo in Utah Valley. The location of missed approaches conducted with the aircraft are shown by dark pink circles. The Hawthorne (HW) measurement site in SLV is labeled. (Right) Expanded view of SLV, with analyzed flight tracks highlighted in green.
Figure 2. Illustration of the day-night dynamics and chemical cycles of reactive nitrogen oxides, O₃, and NH₄NO₃ during PCAP conditions in SLV. The development of the nocturnal boundary layer and morning growth and mix-out are illustrated by the dashed lines. Figure is not to scale. (R6) represents the reaction: $N_2O_5 + \gamma(N_2O_5) \rightarrow 2 \cdot (1 - \varphi) \cdot HNO_3 + \varphi \cdot ClNO_2$.

Figure 3. Example simulation of total nitrate production from sunset to sunrise for an air parcel sampled over SLV on 28 January 2017. Model derived $\gamma(N_2O_5)$ and $\phi(ClNO_2)$ values were 0.05 and 0.21, respectively. Modeled nocturnal nitrate (blue) is the total nitrate produced by heterogeneous chemistry in the box model, with the nocturnal production rate ($\mu g \cdot m^{-3} \cdot night^{-1}$) represented by the blue diamond. Pre-existing nitrate (yellow) represents the nitrate present at sunset and is calculated as the difference between total measured nitrate from the aircraft (red diamond) and the model-predicted nitrate at the time of aircraft measurement (vertical black line). Assuming pre-existing nitrate is constant overnight (i.e. no deposition or dilution) and constant values of $\gamma(N_2O_5)$ and $\phi(ClNO_2)$, the fractional contribution of nitrate production from a single night to the total observed is calculated as the ratio of modeled nitrate (blue diamond) to total nitrate (gold diamond) at sunrise.
Figure 4. Time series of total PM$_{2.5}$ mass (µg m$^{-3}$) (1-hr and 24-hr averages) for the 2016-2017 winter, measured at the Hawthorne (HW) UDAQ site in SLV. Dashed black lines are daily PM$_{2.5}$ accumulation rates (rates given in Figure). The 24-hour EPA national ambient air quality standard for PM$_{2.5}$ (35 µg m$^{-3}$) is shown by the dashed gray line. Gray shading indicates days when the TO aircraft was flying during UWFPS. Average aerosol mass fractions measured by the AMS aboard the TO are given in pie charts for polluted and clean conditions. Aerosol components are colored by nitrate (blue), ammonium (gold), sulfate (red), non-refractory chloride (pink), and organics (green).

Figure 5. (a, top) Time series of ratio of total oxidized (HNO$_3$ + NO$_3^-$) to reduced (NH$_3$ + NH$_4^+$) nitrogen between 16 January and 1 February 2017 (10s averages), calculated from TO observations over SLV. Individual nitrogen ratios are colored by aircraft altitude (mAGL). Yellow and gray shading indicate times of day and night, respectively. (a, bottom) PM$_{2.5}$ mass (24-hour average) measured at the HW ground-site (bottom). (b) Vertical profile of oxidized to reduced nitrogen ratios from panel (a). Diamonds represent the average values in each altitude bin and gray shading shows the 10th-90th (light gray) and 25th-75th (dark gray) percentiles. The number of points in each bin is shown by the gray dashed line. The vertical black line illustrates a nitrogen ratio of 1.
Figure 6. (a) Time series of NO$_2$, O$_3$ (top), P$_{\text{NO}_3^-}$ (middle, see text for definition), and PM$_{2.5}$ (bottom) measured at the HW ground site during 16 January – 6 February 2017. O$_3$ data during the middle January pollution event were corrected to account for a 4.5 ppbv offset in the HW measurements, as shown in Figure S4. Aircraft flight times are shown by red shading. Dashed blue line shows the calculated P$_{\text{NO}_3^-}$ rates that would occur during the day if this mechanism were operative. Solid blue line assumes nitrate production from this mechanism during the day is zero. Late afternoon P$_{\text{NO}_3^-}$ at the surface (dashed line), is roughly equivalent to the P$_{\text{NO}_3^-}$ expected in the RL at night. (b) Vertical profiles of O$_3$, NO$_2$, P$_{\text{NO}_3^-}$, and PM$_1$ measured from the aircraft on all night flights over SLV. In each panel, light shaded regions show the 10$^{th}$-90$^{th}$ percentile ranges, dark shaded regions are the 25$^{th}$-75$^{th}$ percentile ranges, and the solid lines are the 50$^{th}$ percentile. Dashed black lines show the number of points at each altitude.
Figure 7. Vertical Profiles of NO$_2$, O$_3$, P$_{NO_3^-}$, and PM$_{1}$ measured from the TO aircraft during 5 box patterns, flown over the SLV urban core between 21:20 and 00:30 MST on 28 and 29 January. Percentiles and number of points at each altitude are shown as in Figure 6. Square markers and error bars represent the median and 25th-75th percentile range of NO$_2$, O$_3$, P$_{NO_3^-}$, and PM$_{2.5}$ measured concurrently at the HW ground site.
Figure 8. (Top) Histograms of $\gamma(N_2O_5)$ determinations from SLV during pollution events, calculated with the box model (green), steady state approximation (pink), and parameterization from Bertram and Thornton (2009), (Bottom) Histograms of $\phi(ClNO_2)$ determinations from SLV during pollution events calculated with the box model (gold) and parameterization from Bertram and Thornton (2009) (gray).

Figure 9. Histograms of nocturnal nitrate production rates ($\mu g \text{ m}^{-3} \text{ night}^{-1}$) predicted by base case simulations and simulations incorporating a first-order dilution loss process with rate constant $k_{\text{dilution}} = 1.3 \times 10^{-5} \text{ s}^{-1}$.
Figure 10. (a) For pollution event #4, comparison of model-predicted nocturnal nitrate production ($\mu$g m$^{-3}$ day$^{-1}$) for base case simulations (gray), simulations with 24-hours of dilution (blue), and the average daily nitrate build-up observed at HW (red). Dilution cases are for simulations that incorporate nocturnal dilution rate constants of $1.2\times10^{-5}$ (L), $1.3\times10^{-5}$ (M), and $2.5\times10^{-5}$ (H) s$^{-1}$, scaled by 60% during the day. Box and whisker plots show the 10th–90th percentile distributions of each set. The red diamond shows the ground-based build-up rate, calculated from 24-hr averaged data at HW in panel b. (b) Observed concentrations and average daily build-up rate of nitrate aerosol mass (total mass $\times$ 0.58) at HW during event #4.
Table 1. Aircraft measurements used in this analysis

<table>
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<tr>
<th>Compound</th>
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<th>Accuracy</th>
<th>Meas. Frequency</th>
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<td>1s</td>
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<tr>
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<td>1s</td>
<td>Aircraft</td>
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<sup>a</sup>NOAA, Cavity Ring down Spectrometer (CRDS, NOxCaRD)
<sup>b</sup>Hawthorne
<sup>c</sup>University of Washington I Time of Flight Chemical Ionization Mass Spectrometer
<sup>d</sup>University of Toronto, Quantum Cascade Tunable Infrared Laser Differential Absorption Spectrometer
<sup>e</sup>NOAA, Aerosol Mass Spectrometer
<sup>f</sup>Droplet Measurement Techniques, Ultra-High Sensitivity Aerosol Spectrometer
<sup>g</sup>Estimated according to the performance of a different UHSAS in the WINTER campaign