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### **Observations of gas- and aerosol-phase organic nitrates at BEACHON-RoMBAS 2011**

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Abstract. At the Rocky Mountain Biogenic Aerosol Study (BEACHON-RoMBAS) field campaign in the Colorado front range, July-August 2011, measurements of gas- and aerosol-phase organic nitrates enabled a study of the role of  $NO_x$  ( $NO_x = NO + NO_2$ ) in oxidation of forest-emitted volatile organic compounds (VOCs) and subsequent aerosol formation. Substantial formation of peroxy- and alkylnitrates is observed every morning, with an apparent 2.9% yield of alkyl nitrates from daytime RO<sub>2</sub> + NO reactions. Aerosol-phase organic nitrates, however, peak in concentration during the night, with concentrations up to 140 ppt as measured by both optical spectroscopic and mass spectrometric instruments. The diurnal cycle in aerosol fraction of organic nitrates shows an equilibrium-like response to the diurnal temperature cycle, suggesting some reversible absorptive partitioning, but the full dynamic range cannot be reproduced by thermodynamic repartitioning alone. Nighttime aerosol organic nitrate is observed to be positively correlated with  $[NO_2] \times [O_3]$  but not with  $[O_3]$ . These observations support the role of nighttime NO<sub>3</sub>-initiated oxidation of monoterpenes as a significant source of nighttime aerosol. Nighttime production of organic nitrates is comparable in magnitude to daytime photochemical production at this site, which we postulate to be representative of the Colorado front range forests.

### 1 Introduction

Nitrogen oxides affect both daytime and nighttime volatile organic compound (VOC) oxidation mechanisms, controlling tropospheric ozone formation and the branching ratios for formation of VOC oxidation products, such as organic peroxides (ROOR and ROOH), alcohols and ketones (ROH and R=O), and multifunctional organic nitrates (RONO<sub>2</sub>). These products can have reduced volatility making them good aerosol precursors. Figure 1 shows the general mechanisms of NO<sub>x</sub> control of VOC oxidation. First, NO<sub>x</sub> enhances nighttime or low-light oxidation by providing a source  $(NO_2 + O_3)$  of elevated NO<sub>3</sub> (blue), adding an additional oxidation pathway. Second, during the day, directly emitted and photolytically produced NO can be elevated (red) and can control the fate of RO2 radicals, preferentially enhancing  $RO + NO_2$  and  $RONO_2$  reaction channels over non-nitrate channels. With sufficiently high NO<sub>3</sub>, another nighttime NO<sub>x</sub> effect is possible: NO<sub>3</sub> may react with



Fig. 1. Overview of potential  $NO_x$  effects on the oxidation of a volatile organic compound.

 $RO_2$ , effectively playing the role of daytime NO, in competition with  $RO_2 + RO_2$  reactions. These reactions remain very poorly understood (Sommariva et al., 2012).

 $NO_3$  is the least understood of the atmospheric oxidants; its concentration is much more variable in space and time and is therefore poorly characterized. Further, since it exists at night in a stable rather than well-mixed boundary layer, surface measurements are insufficient to predict the vertical distribution of  $NO_3$ .

Due to its high reactivity, the atmospheric lifetime of  $NO_3$ in forests is likely to be limited by its reaction with unsaturated hydrocarbons, especially biogenic VOCs (Winer et al., 1984; Golz et al., 2001; Fuentes et al., 2007). In urban forests,  $NO_3$  is responsible for a substantial fraction of total VOC degradation (e.g., 28 % in Pabstthum near Berlin, Germany; Geyer et al., 2001).

A better understanding of the effects of  $NO_x$  on secondary organic aerosol (SOA) production is a key goal of this study. A growing consensus is emerging that NO<sub>3</sub>-initiated oxidation of biogenic volatile organic compounds (BVOCs) is an important source of secondary organic aerosol (Fry et al., 2009, 2011; Pye et al., 2010; Rollins et al., 2012), suggesting that organic nitrates may be a significant fraction of aerosol composition. These nitrates may also contribute to the remaining underprediction of SOA sources: models based on laboratory-measured SOA yields from known precursor VOCs underestimate aerosol loading by from a factor of 2 (Farina et al., 2010) to up to an order of magnitude in polluted regions (de Gouw et al., 2005; Heald et al., 2005; Volkamer et al., 2006) and cannot explain the evolution of SOA (Jimenez et al., 2009). Recent explicit chemical modeling of SOA formation in Mexico City indicates SOA loading increases up to several days downwind of urban areas (Lee-Taylor et al., 2011). All of this suggests the existence of additional mechanisms of SOA formation not yet quantified in the laboratory, yields that are higher under ambient conditions than as measured in the laboratory, or the presence of significant additional VOC mass that is the precursor for aerosol formation (Donahue et al., 2006; Robinson et al., 2007). More recent modeling results can close the gap with the measurements in polluted regions, but it is unclear if this is for the right reasons (Dzepina et al., 2009; Hodzic et al., 2010). A substantial *enhancement* of biogenic SOA formation due to anthropogenic pollution has been suggested as a controlling factor for SOA formation (de Gouw et al., 2005; Weber et al., 2007; Spracklen et al., 2011; Hoyle et al., 2011). Because  $NO_x$  can either suppress or enhance aerosol formation, field measurements that probe  $NO_x/SOA$  interactions in the real atmosphere are key to evaluating the magnitude of this effect.

Here, we address the question of the fate of  $NO_x$  in a pine forest, with particular focus on its influence on biogenic SOA formation. We measured selected  $NO_y$  species ( $NO_2$ , peroxynitrates ( $\Sigma PNs$ ), alkyl nitrates ( $\Sigma ANs$ ), and their gas/aerosol partitioning) over a 6-week summertime campaign in a Colorado front range forest, and interpret their daily cycles with the aid of auxiliary data on  $NO_3$ ,  $N_2O_5$ , meteorology,  $O_3$ , BVOCs, radicals, and regional transport modeling. We demonstrate and quantify distinct daytime and nighttime mechanisms of organic nitrate formation, showing that even at this forested site 40 km from the nearest urban area (peak [ $NO_2$ ]  $\approx 2$  ppb),  $NO_x$  chemistry has a substantial impact on BVOC fate and SOA formation.

### 2 Experimental design

#### 2.1 BEACHON-RoMBAS field site description

The Bio-hydro-atmosphere interactions of Energy, Aerosols, Carbon, H<sub>2</sub>O, Organics, and Nitrogen (BEACHON) project is a long-term, multi-institution, collaborative effort, spearheaded by the National Center for Atmospheric Research (NCAR). In July and August 2011, the Rocky Mountain Biogenic Aerosol Study (BEACHON-RoMBAS, hereinafter RoMBAS) field campaign took place, with research groups from 25 institutions participating in measurements with focus particularly on aerosol emissions and formation. The RoMBAS campaign was located in the United States Forest Service (USFS) Manitou Forest Observatory (MFO) in Pike National Forest, Colorado (39.10° N, 105.10° W). The site is at 2370 m elevation, 40 km northwest of Colorado Springs and 70 km southwest of Denver, CO (Fig. 2). Vegetation at the site is almost entirely open-canopy ponderosa pine, but nearby areas include Douglas fir, aspen, oak, spruce, willow, and grass which may have some impact at the site. Previous studies at the site have observed dominant VOC emissions to be monoterpenes (34% of total VOC mixing ratio) and 2-methyl-3-butene-2-ol (MBO, 50%) during the day, with monoterpene concentrations dominant at night ( $\approx 66\%$  of VOC), comprised of a roughly equal mix of  $\alpha$ -pinene,  $\beta$ pinene, and  $\Delta$ -3-carene (Kim et al., 2010). Nighttime winds at the surface were dominated by local drainage flows from the south.

During July and August 2011 the circulation and precipitation patterns over the MFO site were strongly influenced by the North American Monsoon (Hodzic et al., 2013). During



**Fig. 2.** Map of Manitou Forest Observatory site. Top panels show topography and point sources for  $NO_x$ . Regional  $NO_x$  point sources are shown as black/grey dots with size proportional to emissions rate. The Four Corners power plant (lower left of larger map) is not shown on this scale; its emissions of 136 tons day<sup>-1</sup> dwarf all other power plants in this region. The site is 40 km northwest of Colorado Springs and 70 km southwest of Denver, the two largest urban areas (outlined in blue) in the domain. For discussion of transport to the site, note the South Platte River Valley (rivers shown in aqua) southwest from the Denver urban area towards the field site (red/green diamond). Bottom panels show monoterpene emission potential, calculated using the BEIS model.

25–28 July and 2–5 August, the presence of a strong anticyclone over the four-corners region in the southwestern US was associated with the strong southerly mid-level flow, and brought significant amount of moisture from the southeast and southwest into this region. The largest rain event occurred on the evening of 4 August and was characterized by heavy hail and rain whose 5 min intensity values exceed  $120 \text{ mm}\text{h}^{-1}$ . During the rest of the campaign, the site was experiencing more zonal synoptic flow with mid-level winds mainly from the west and very little large-scale advection of moisture into the region.

The near-surface circulation was strongly influenced by the daytime thermally induced mountain flow that was established a few hours after sunrise on days when the synoptic forcing was weak. The upslope transport of the front range (plains) air and pollution was observed on all nights in NO<sub>2</sub> and some days in CO and SO<sub>2</sub> trace gas measurements (see discussion in Sect. 3.1.2 and Fig. 4).

The site was specifically chosen to have large biogenic emissions and limited anthropogenic influence, and one of the objectives of RoMBAS was studying the ambient submicron aerosol, presumed to be dominantly biogenic. Most chemical instruments were arranged in and on top of trailers with inlets oriented into the dominant wind, towards the south, or on a 30 m tower at the center of the site ("Chemistry tower").

### 2.2 Instrumentation

Nitrogen dioxide, the sum of total peroxy nitrates ( $\Sigma PNs$ ), and the sum of total alkyl nitrates ( $\Sigma ANs$ ), as well as aerosolphase-only organic nitrates, were measured by a 3-channel thermal dissociation laser-induced fluorescence (TD-LIF) instrument similar to that described in Day et al. (2002), with selective measurement of aerosol phase enabled by a modification described in Rollins et al. (2010). Briefly, the instrument thermally dissociates the different classes of NO<sub>v</sub> into NO<sub>2</sub> at ambient pressure in three inlet channels, samples this flow through a critical orifice (dropping the pressure to  $\approx 2$  Torr) into a 38-pass White cell, and measures the NO<sub>2</sub> mixing ratio by LIF at 408 nm. One of these channels can be optionally operated as an "aerosol-only" channel by passing sampled air first through a 10 cm-long, 2.2 cm outer diameter cylindrical charcoal denuder (Mast Carbon Ltd, UK, NovaCarb F honeycomb monolith, 44 % of cross-sectional area open). This denuder removes gases more efficiently than the original design; using calculations described and verified in Rollins et al. (2010), we find that at 10L per minute total flow, < 0.001 % of NO<sub>2</sub> gets through this honeycomb charcoal denuder. Contributions of each class of NO<sub>v</sub> are determined by calculating the difference in NO<sub>2</sub> concentration detected between the channels. Because peroxynitrates dissociate between 100 and 150 °C and alkyl nitrates between 250 and 300 °C, these classes of NO<sub>v</sub> can be separately measured; e.g., the difference in NO<sub>2</sub> signal between ambient air sampled through a 180 °C oven ( $\Sigma PNs + NO_2$ ) and ambient temperature (NO<sub>2</sub>) is attributed to  $\Sigma PNs$  alone. The instrument is calibrated every 3h using an NO2 gas standard (4.74 ppm  $\pm 2\%$  NO<sub>2</sub> in N<sub>2</sub>, NIST characterized, March 2011, Scott-Marrin) diluted with zero air to 5 known concentrations in the 1–25 ppb range, resulting in a measurement uncertainty of 5 % for NO2. The accuracy of this measurement and effectiveness of the separation has been demonstrated by comparison to summed individually measured speciated peroxynitrates (Wooldridge et al., 2010) and alkyl nitrates (Perring et al., 2009; Beaver et al., 2012).

The TD-LIF instrument was run in two modes during the campaign: (1) "3-gas mode", in which the inlet ovens were held at ambient temperature, 180 °C, and 320 °C, with the denuder bypassed, to obtain 1 min averaged time series of separated NO<sub>2</sub>,  $\Sigma$ PNs, and  $\Sigma$ ANs; and (2) "gas/aerosol mode", in which both heated channels are held at 320 °C, with air sampled into one of the heated channels passed through the denuder, to obtain 1 min averaged time series of total  $\Sigma$ PNs+ $\Sigma$ ANs and aerosol-phase-only  $\Sigma$ PNs+ $\Sigma$ ANs (ambient temperature NO<sub>2</sub> channel subtracted from each). It is assumed that the aerosol-phase organic nitrates will consist predominantly of alkyl nitrates, due to the short lifetime of PANs. To the best of our knowledge, PANs have not been measured in the aerosol alkyl nitrates (FTIR) with  $\Sigma$ PNs+ $\Sigma$ ANs

(TD-LIF) finds that aerosol-phase  $\Sigma PNs + \Sigma ANs$  are dominated by alkyl nitrates (Rollins et al., 2012).

NO<sub>3</sub> and N<sub>2</sub>O<sub>5</sub> were measured in situ by cavity ringdown spectroscopy (CRDS) using the 662 nm absorption band of NO<sub>3</sub>, detecting N<sub>2</sub>O<sub>5</sub> via its thermal conversion to NO<sub>3</sub> (Brown et al., 2001, 2003b, 2007; Wagner et al., 2011). The instrument used a diode laser, tunable over a narrow range near the 662 nm absorption of NO<sub>3</sub>, to pump two separate optical cavities, each constructed of two high-reflectivity mirrors (R = 99.999% or better). The laser is modulated as a square wave at 500 Hz, and the single exponential ringdown transients following the falling edge of the square wave are co-added and fit at 4 Hz to measure total optical extinction,  $\alpha$ , according to Eq. (1):

$$\alpha = [\text{NO}_3]\sigma_{\text{NO}_3} = \frac{R_{\text{L}}}{c} \left(\frac{1}{\tau} - \frac{1}{\tau_0}\right). \tag{1}$$

Here  $\tau$  and  $\tau_0$  are exponential intensity decay time constants in the presence and absence of the absorber, respectively; c is the speed of light;  $R_{\rm L}$  is a geometric design factor to account for the fraction of the cavity length occupied by purge volumes that keep the mirrors clean (typically  $R_{\rm L} = 1.15$ ; and [NO<sub>3</sub>] and  $\sigma$ (NO<sub>3</sub>) are the NO<sub>3</sub> number density (molecules  $cm^{-3}$ ) and absorption cross section (cm<sup>2</sup> molecule<sup>-1</sup>), respectively. Values of  $\tau_0$  vary from 250 to 450 µs, equivalent to 75–135 km in effective path length, to provide a limit of detection of 1 pptv  $(2\sigma)$  for NO<sub>3</sub> and N<sub>2</sub>O<sub>5</sub>. The  $\tau_0$  is measured by addition of NO to chemically destroy NO<sub>3</sub> via the rapid reaction NO + NO<sub>3</sub>  $\rightarrow$  2NO<sub>2</sub>. The combination of 662 nm extinction and NO titration is a specific technique for ambient air NO3 measurement. The concentration of the sum of NO3 and N2O5 is measured simultaneously by thermal decomposition of N<sub>2</sub>O<sub>5</sub> to NO<sub>3</sub> in a second optical cavity that samples through a heated section of tubing (120 °C) and that is maintained at 75 °C. To achieve high sensitivity, both channels sample through a Teflon membrane filter that removes aerosol; filters are changed hourly using an automated device to maintain their cleanliness and transmission of NO<sub>3</sub> and N<sub>2</sub>O<sub>5</sub>. The instrument has an overall accuracy of -9/+12% for NO<sub>3</sub> and -8/+11% for N<sub>2</sub>O<sub>5</sub>, calibrated by separately measuring the NO<sub>2</sub> produced from the same chemical titration reaction used to zero the instrument (Fuchs et al., 2008).

Ambient NO<sub>2</sub> was also measured continuously by a chemiluminescent NO<sub>x</sub> box (Thermo 17i), which agreed within 8% with the TD-LIF ambient temperature (NO<sub>2</sub>) channel for all periods when both instruments were sampling ambient air,  $\approx$  14 days over the course of the campaign (the TD-LIF was used for other measurements for substantial periods). Because the NO<sub>x</sub> box monitored ambient air continuously, producing a 32-day record, while agreeing well with the more selective TD-LIF measurement of NO<sub>2</sub> during periods of overlap, the longer NO<sub>x</sub> box time series will be used in e.g. computing diurnal averages. Inlets for this instrument as well as CO and SO<sub>2</sub> monitors were located at  $\approx$  3m. Ozone was measured by UV absorption spectroscopy (2B Technologies, Model 205) and sulfur dioxide by pulsed fluorescence (Thermo Environmental Model 43C-TLE) at 6 different heights along the Chemistry tower (z = 1.6, 5.0, 8.5, 12.0, 17.7 and 25.1 m). These inlets were sampled sequentially every 5 min, yielding a complete profile every 30 min; for this work, only concentration data at the lowest height of 1.6 m were used. Carbon monoxide was measured at 3.5 m by a Thermo Environmental Gas Filter Correlation CO analyzer (Model 48 equipped with a heated Pt catalytic converter). Wind speed/direction, temperature and relative humidity were measured at 4 heights (z = 1.8, 7.0, 14.1 and 27.8 m; Vaisala, Model WXT520). Net radiation (visible + IR, incoming - outgoing) was measured at 27.8 m (Radiation and Energy Balance Systems, Inc., Model Q\*7.0 net radiometer). Turbulence measurements were made at z = 25.1 m,  $\approx 9 \text{ m}$  above the canopy height, by a 3dimensional sonic anemometer (Campbell Scientific, CSAT-3). From these flux measurements, the Obukhov length (L)can be determined (Stull, 1998), which is a surface layer scaling parameter that gives the ratio of buoyant to mechanically (shear) generated turbulence, which can be used as a measure of atmospheric stability. When  $L > \pm 130 \text{ m} (|z/L| < 0.1)$ , shear forces dominate and the atmosphere has nearly neutral thermal stability. At values less than  $\pm 100$  m, convective forces become increasingly important and denote an atmo-

VOCs were measured using two proton-transfer-reaction time-of-flight mass spectrometers (PTR-ToF-MS, Ionicon Analytik GmbH, Austria, Jordan et al., 2009; and University of Innsbruck, Graus et al., 2010). Ambient air was sampled at a flow rate of  $\approx 9$  SLPM through a 40 m-long Teflon (PFA) line (1/4'' OD) mounted at 25.3 m on the Chemistry tower. Both instruments were sampling off the same line with a sampling period of about 10s (NCAR) and 0.1s (UIBK). The merged dataset was averaged to 6 min. The drift tube was operated at 2.3 mbar (both instruments) and a drift voltage of 580 V (UIBK) and 550 V (NCAR) and a drift tube temperature of 60 °C (both instruments). Calibration was performed by dynamically diluting the VOC standards to ppbv levels using purified ambient air. Details about the data evaluation can be found in Müller et al. (2010), Kaser et al. (2013), and Cappellin et al. (2011).

sphere that is either thermally stable (|z/L| > 0) or unstable

(|z/L| < 0).

Submicron particle size distributions (20–800 nm) were measured with a Brechtel Manufacturing, Inc. scanning electrical mobility spectrometer (SEMS, Model 2002). From these size distributions, total particle surface area per volume can be calculated, and assuming a density of  $1.3 \text{ g cm}^{-3}$ , aerosol mass loading can be determined.

An Aerodyne high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS; hereafter AMS; DeCarlo et al., 2006), was operated in close vicinity ( $\approx 10 \text{ m NW}$ ) to the TD-LIF instrument during RoMBAS. Ambient aerosols in the AMS are analyzed by impaction/vaporization of the aerosol on a porous tungsten vaporizer at 600 °C and subsequent ionization of the gas plume by electron impact; ions are mass analyzed by a high-resolution time-of-flight mass spectrometer. The AMS quantifies non-refractory submicron aerosol, which includes organic species and most inorganic salts and organic species, but not black carbon. Refractory nitrates such as Ca(NO<sub>3</sub>)<sub>2</sub> and NaNO<sub>3</sub> can be detected by the AMS; however, they are typically associated with reacted mineral dust and marine aerosol, respectively, both of which only have small fractions in the submicron regime that were likely negligible during RoMBAS.

The AMS sampled ambient aerosol from an inlet at the same height inside the forest canopy as the TD-LIF instrument; a PM<sub>2.5</sub> cyclone was used to prevent large dust particles from entering the instrument, and sample air was drawn from the inlet into the temperature-controlled trailer through a 3/8-inch o.d. copper tube at 10 lpm. Ambient data used in this work was acquired every 10 min for 2.5 min-long intervals using the lower mass-spectral resolution, highersensitivity mode of the AMS ("V-mode"). The AMS was calibrated every four days with ammonium nitrate aerosol of known mass, and calculated submicron aerosol volumes derived from these calibrations agreed well with volumes from a collocated SMPS. The overall accuracy of the AMS for ground measurements has been estimated to be about 30 % for all AMS species (Middlebrook et al., 2012; Hayes et al., 2012), with better accuracy for ratios due to error cancellation. The precision is significantly better: about  $4 \text{ ng m}^{-3}$  for aerosol nitrate and  $30 \text{ ng m}^{-3}$  for OA for the time resolution used here. The low nitrate and high organic aerosol concentrations require careful analysis of the HR spectra, taking into account the contributions of otherwise unimportant isotopic air interferences such as  $C^{18}O^+$  and  $C^{18}OO^+$  to the signals at m/z 30 and 46, respectively.

#### 2.3 Regional modeling

The 3-D Weather Research and Forecasting model coupled with chemistry (WRF-Chem; Grell et al., 2005) was applied to study the transport and chemistry of biogenic and anthropogenic pollutants at the MFO site from 25 July to 26 August 2011. The chemistry was simulated using the SAPRC99 gas-phase chemical mechanism (Carter, 2000) and the MO-SAIC aerosol module with 4 size bins (Zaveri et al., 2008). Organic aerosols were treated as described in Hodzic et al. (2012), and several anthropogenic CO tracers were added to monitor the transport of anthropogenic pollution to the site.

The simulations were performed using two nested domains: a coarse-scale 36 km horizontal resolution grid covering the western and central US and a fine-scale 4 km grid covering Colorado. The anthropogenic emissions come from the 2005 US EPA National Emission Inventory (NEI), while the biogenic emissions are calculated online using the MEGAN model (Guenther et al., 2006). The initial and boundary conditions for the meteorological variables are taken every 6 h from the NCEP North American Regional Reanalysis, and gas-phase and aerosol variables were obtained from the MOZART4 global chemistry-transport model. The model performance in simulating the meteorology, boundary layer height, and the concentrations of the main gas-phase pollutants during RoMBAS has been evaluated in detail in Hodzic et al. (2013).

# 2.4 Predicting gas/aerosol partitioning of organic nitrates

Direct measurements of gas/aerosol partitioning of these organic nitrates can provide insights into relative volatilities of daytime vs. nighttime RONO<sub>2</sub>, and hence the role of NO<sub>x</sub> in SOA formation. Interpreting these observations in terms of chemical mechanisms, however, requires some way to connect chemical structures to a predicted partitioning. We choose to model gas/aerosol partitioning in terms of an absorptive partitioning formalism (assuming no solid phase), following Pankow (1994) and Capouet and Müller (2006). The partitioning coefficient,  $K_p$ , is defined as

$$K_{\rm p} = \frac{F/{\rm TSP}}{A} = \frac{760 \cdot R \cdot T \cdot f_{\rm om}}{{\rm MW}_{\rm om} \cdot 10^6 \cdot \zeta \cdot p_{\rm vap}}.$$
(2)

*F* and *A* are the total aerosol-phase and gaseous concentrations of the compound of interest, and TSP is the concentration of total suspended particulate matter. In the second expression showing the equilibrium constant in terms of thermodynamic properties, *R* is the universal gas constant  $(8.314 \text{ J} \text{ mol}^{-1} \text{ K}^{-1} = 8.206 \times 10^{-5} \text{ atm m}^3 \text{ K}^{-1} \text{ mol}^{-1})$ , *T* is temperature (K),  $f_{\text{om}}$  is the weight fraction of organic matter in the total aerosol (assumed = 1), MW<sub>om</sub> is the average molecular weight of the absorbing organic material (g mol<sup>-1</sup>),  $\zeta$  is the activity coefficient of the compound of interest in the condensed phase (assumed = 1), and  $p_{\text{vap}}$  is the subcooled vapor pressure of the compound of interest (Torr); 760 (Torr atm<sup>-1</sup>) and  $10^6 (\mu \text{ g} \text{ g}^{-1})$  are conversion factors to give  $K_p$  in units of m<sup>3</sup>  $\mu \text{ g}^{-1}$ .

Vapor pressures calculated in this paper are also reported as saturation mass concentrations ( $C^*$ ) for ease of comparison with volatility basis set parameterizations (Donahue et al., 2006). These saturation densities can be calculated as

$$C^* = \frac{1}{K_{\rm p}} \tag{3}$$

using the same variables shown above to determine  $K_p$ , yielding  $C^*$  in units of  $\mu g m^{-3}$ .

As pointed out by Barley and McFiggans (2010), a significant uncertainty in this method can arise from the variation in vapor pressure values predicted by different group contribution estimation methods. We use the SIMPOL.1 method here (Pankow and Asher, 2008), with primary focus on the temperature and TSP dependence over a daily cycle for a given molecular structure, rendering the exact predicted  $p_{vap}$  values less important than relative changes.

#### 3 Results and discussion

### 3.1 Reasons for elevated NO<sub>x</sub> and organic nitrate concentrations at RoMBAS

### 3.1.1 Observed diurnal cycles in oxidants, BVOCs, meteorology, and photochemical age markers

Figure 3 shows averaged daily cycles in the concentrations of oxidant and terpenoid gases that form the background and reactants for the observed NO<sub>y</sub> chemistry, as well as relevant meteorological variables and an indicator of airmass age with respect to anthropogenic pollution. Ozone (measured at h = 1.6 m) shows a typical photochemistry- and depositiondriven daily cycle, peaking at an average concentration of 60 ppb in the late afternoon and decreasing to 30 ppb during the night. NO<sub>2</sub> shows the opposite diurnal pattern, peaking at an average concentration of 2 ppb at night and sinking below 0.5 ppb during the day. The diurnal NO<sub>2</sub> pattern is explored below in Sect. 3.1.2.

BVOC concentrations measured by PTR-ToF-MS are dominated at night by the sum of all monoterpenes (measured as m/z 137.134 (C<sub>10</sub>H<sup>+</sup><sub>17</sub>) and m/z 81.071 (C<sub>6</sub>H<sup>+</sup><sub>9</sub>), peak diurnal average concentration of about 0.6 ppb) and during the day by the sum of MBO and isoprene (measured as m/z87.081 (C<sub>5</sub>H<sub>11</sub>O<sup>+</sup>) and m/z 69.070 (C<sub>5</sub>H<sub>0</sub><sup>+</sup>), peak average concentration of about 1.8 in the early morning,  $\approx 1.5 \text{ ppb}$ throughout the day). This suggests a very large daytime photo-induced emission of MBO, since this morning peak occurs simultaneously with an increase in convective vertical mixing which effectively dilutes all species. Furthermore, this morning peak (9:00-10:30 LT) coincides with an observed downward flux of  $CO_2$  above the canopy (not shown), suggesting a maximum in photosynthetic activity. This transition is shown by the Obukhov stability parameter (z/L) in Fig. 3) changing from a positive to negative sign. At this point, the temperature structure switches from a stable nighttime inversion to a daytime positive lapse rate with higher temperatures at the surface. Local wind direction is consistently from the south, with greater variability during the day and early evening.

We also use PTR-ToF-MS measurements of two aromatic hydrocarbons, xylene and benzene, to establish the diurnal changes in average photochemical age of the pollution contained in the airmass. Because xylene and benzene are typically co-emitted by fossil fuel and combustion sources which are common in urban areas, their ratio can provide information about photochemical airmass age. Xylene is measured at m/z 107.086 (C<sub>8</sub>H<sup>+</sup><sub>11</sub>), as a mixture of *ortho-*, *para-*, and *meta-*isomers, ethylbenzene, and negligible contributions from the pinonaldehyde fragment. Benzene is measured at m/z 79.055 (C<sub>6</sub>H<sup>+</sup><sub>7</sub>). Both were measured with concentrations in the 10–100 ppt range over the course of the campaign. The time rate of change in the ratio of xylene : benzene provides information on the photochemical





**Fig. 3.** Hourly binned average diurnal trends in NO<sub>2</sub> (blue), O<sub>3</sub> (orange), monoterpenes (purple), methyl butenol and isoprene (pink), wind direction (grey), radiation (yellow, arbitrary units), Obukhov length (green), xylene-to-benzene ratio (aqua), and average day and night windroses at MFO, 21 July–24 August 2011. Error bars are  $1\sigma$  based on data averaged to a one-hour timebase; all points are averaged for the same hour intervals but offset slightly on some plots to make error bars easier to read. Concentrations of NO<sub>2</sub> and monoterpenes peak at night when they are concentrated in a shallow boundary layer, while O<sub>3</sub> and MBO peak during the day. Winds are dominantly from the south, with more variability during the day; every morning at sunrise the Obukhov length shows a transition from a stable nighttime inversion to a convectively mixed boundary layer, and decreasing xylene : benzene ratio shows aging.

age of the airmass; since the OH rate constants of these species are different, the xylene : benzene decreases with aging. We note that this ratio provides no information on the age of the biogenic gases or aerosol in the airmass. This analysis is described below in Sect. 3.1.3.

#### 3.1.2 Information from tracers: modeled and measured

Results from the WRF-Chem simulation from the early part of the campaign are shown in Fig. 4. The anthropogenic front range CO tracers suggest the frequent occurrence of transport of polluted air from both Denver and Colorado Springs to MFO. This transport occurs in the early afternoon within a well-developed boundary layer. Peaks in observed [SO<sub>2</sub>] occur most often during the day, and often coincide with peaks in anthropogenic tracers from either Denver or Colorado Springs, indicating the arrival of pollution plumes at the site. The sharper spikes in predicted surface NO<sub>2</sub> concentrations are also coincident with the presence of anthropogenic tracers at MFO; e.g., the NO<sub>2</sub> spike that was measured and predicted on 27 July is coincident with the modeled arrival of a concentrated Denver plume.

The regular, broader nighttime peaks in NO<sub>2</sub> are also predicted by the model (see e.g. 6-8 August), though the accuracy of the concentration prediction varies. Rather than individual concentrated plumes as for SO<sub>2</sub>, this NO<sub>2</sub> appears to arise from a more regional background, consistent with the importance of a few point sources (coal-fired power plants) for SO<sub>2</sub> vs. the much larger contribution of area sources (vehicles) for NO<sub>x</sub>. Also note that SO<sub>2</sub> peaks are often not coincident with NO2 or Denver/Colorado Springs anthropogenic tracer peaks, suggesting distinct sources. Examination of regional topography (Fig. 2) reveals that the Denver plume could be transported up the Platte River canyon during daytime upslope flow that is characteristic of the Rocky Mountain foothills, and dispersed into the valley where the MFO site is located in late afternoon and evening, potentially leading to these higher nighttime concentrations of urban pollutants. Diurnal variation in measured CO concentration (not shown) almost always tracks NO2, showing the same broad nighttime peaks, consistent with this idea of a dispersed urban source of NO<sub>2</sub> to MFO.

An alternative explanation for the broad nighttime peaks in NO<sub>2</sub> concentration could be natural emissions of nitrogen oxides from local soils, emitted as NO but immediately titrated to NO<sub>2</sub> by O<sub>3</sub>. This appears to be contradicted, however, by the correlation of NO<sub>2</sub> with CO, as well as the observed sharp early evening increases in NO<sub>2</sub> concentration, followed by either flat or decreasing concentrations overnight. If the cause were NO emissions into a shallow nocturnal boundary layer, the concentration increase should continue throughout the night, as observed by, e.g., Aneja et al. (1996). In that study in the southeastern US, the authors used a dynamic flux chamber to quantify the nitrogen flux from soils, which they found to be  $\approx 0.1$  ppb NO<sub>2</sub> h<sup>-1</sup>.



**Fig. 4.** Time series of NO<sub>2</sub> (black, Thermo 17i) and SO<sub>2</sub> (blue) surface concentrations as observed at the MFO site during the early stages of RoMBAS from 26 July to 14 August 2011. The WRF-Chem-predicted NO<sub>2</sub> is also shown (red, ppbv), as well as the levels of advected anthropogenic tracers (arbitrary units) from Denver (green) and Colorado Springs (yellow). Grey shadings indicate nighttime periods (8 p.m.–8 a.m.).

Rates of increase observed at RoMBAS were typically of order 1 ppb NO<sub>2</sub> h<sup>-1</sup>, with large variability in timing and rate of increase. The terrain and transport at RoMBAS is complex, with downslope drainage flows advecting past the site at night. This makes it far more likely that observed sharp increases in NO<sub>2</sub> arrive with this advected plume, rather than resulting from local soil NO<sub>x</sub> production: a buildup of soil NO<sub>x</sub> would be diluted out by this drainage flow and cause *decreasing* NO<sub>x</sub>.

### 3.1.3 Observed ambient PNs and ANs

While NO<sub>2</sub> peaks during the night, the diurnal pattern in peroxynitrates ( $\Sigma$ PNs) and alkyl nitrates ( $\Sigma$ ANs) is different (top panel of Fig. 5). Both  $\Sigma$ PNs and  $\Sigma$ ANs peak during the day, with increases of  $\approx 200 \,\mathrm{ppt}$  in both cases, relative to nighttime concentrations. This increase in concentration occurs coincident with the increased vertical mixing accompanying the switch from nighttime temperature inversion to daytime positive lapse rate, as described in Sect. 3.1.1. A recent comparison (Wooldridge et al., 2010) of ambient  $\Sigma$ PN measurements shows typical mean concentrations below 500 ppt (except in urban areas where  $\Sigma$ PNs are up to a few ppb), with peak concentration during the day. At another pine forested site downwind of an urban area (BEARPEX, 75 km northeast of Sacramento), mean  $\Sigma$ PN concentration was  $\approx$  400 ppt, while  $\Sigma$ ANs were found to increase from  $\approx 300$  to  $\approx 600$  ppt in the morning (Beaver et al., 2012).

There is no substantial change in local wind direction coincident with the increase in [ $\Sigma$ PNs] and [ $\Sigma$ ANs], though the local winds are overall weaker and more mixed in direc-



**Fig. 5.** Hourly binned average diurnal trends in (top): NO<sub>2</sub> (blue),  $\Sigma$ PNs (green), and  $\Sigma$ ANs (red) from times when the TD-LIF was running in "3-gas" mode (13–17 and 19–20 August); and (bottom): total  $\Sigma$ PNs+ $\Sigma$ ANs (red), aerosol-phase-only  $\Sigma$ PNs+ $\Sigma$ ANs (brown), and fraction of organic nitrates in the aerosol phase (black) from times when the TD-LIF was running in "gas/aerosol" mode (28 July–1 August, 20–22 August). Error bars are 1 $\sigma$  based on data averaged to a one-hour timebase; all points are averaged for the same hour intervals but offset slightly on some plots to make error bars easier to read. The NO<sub>2</sub> trace looks slightly different than Fig. 3 because this data comes only from the subset of campaign days when the TD-LIF was running in "3-gas" ambient mode.

tion during the day than at night. This rules out the possibility that sampling air from a different source region is responsible for the increase. Because the increase in concentration *is* coincident with increasing vertical mixing, a second possible explanation is that higher-NO<sub>y</sub> air from above mixes down in the morning. The observations would require that the residual layer above the nocturnal boundary layer consist of essentially NO<sub>2</sub>-free air with higher concentrations of  $\Sigma$ PNs and  $\Sigma$ ANs, more "aged" forms of NO<sub>y</sub>.

The available data do not support a solely vertical mixing explanation for the increase in [ $\Sigma$ PNs] and [ $\Sigma$ ANs]; the ratio of xylene to benzene is observed to decrease relatively slowly at daybreak in the mean (Fig. 3). Roberts et al. (1984) showed that fresh emissions have a characteristic ratio of aromatic hydrocarbons, which change with age. In the case of the xylene-to-benzene ratio, fresh urban emissions have xylene : benzene of 1.3 (Harley et al., 1992) to 2 (Fraser et al., 1998; Schauer et al., 2002). Peak observed xylene : benzene at MFO occurs at night, with an average ratio of  $\approx 1$ . This ratio decreases at daybreak, consistent with photochemical aging, as the rate constant of xylene with OH ( $1.5 \times 10^{-11}$  molecules cm<sup>-3</sup> s<sup>-1</sup>; Roberts et al., 1984) is an order of magnitude faster than benzene with OH ( $1.2 \times 10^{-12}$  molecules cm<sup>-3</sup> s<sup>-1</sup>; Atkinson et al., 2004). At the observed morning MFO [OH] of  $2.5 \times 10^6$  molecules cm<sup>-3</sup> (Kim et al., 2013), these rate constants lead to the prediction that photochemical aging from xylene : benzene of 1 to 0.5 would take  $\approx 5$ -6 h. In fact, we observe this decrease in the ratio over 5 h in the diurnally averaged (mean) data (Fig. 3). Because this increase in average photochemical age corresponds to the actual clock age and tracks with the increase in [ $\Sigma$ PNs] and [ $\Sigma$ ANs], actual aging is more likely to explain this timing than the downward mixing of more aged air aloft in the morning. We note, however, that there are some days on which the xylene : benzene ratio decreases more rapidly, which would be consistent with aged aloft air contributing on occasion.

The final possible explanation for the morning increase in [ $\Sigma$ PNs] and [ $\Sigma$ ANs] is that a significant photochemical source exists that initiates at daybreak, converting local VOCs that have built up overnight to these more highly oxidized NO<sub>y</sub>. This is corroborated by an observed increase in [OH] from 1 to  $3 \times 10^6$  between 7:00–9:00 LT (Kim et al., 2013). This photochemical processing is intertwined with tropospheric ozone production, and that chemistry can be used to determine a branching ratio of alkyl nitrate formation in the oxidation of the ambient mixture of VOCs.

# 3.1.4 Using morning $\Sigma AN$ rise to determine RONO<sub>2</sub> yield and impact on O<sub>3</sub> production

Tropospheric ozone formation is governed by the coupled  $RO_x$  and  $NO_x$  catalytic cycles. Without radical sinks, typically two  $O_3$  molecules are produced for each time around the  $RO_x$  radical cycle ( $RO_2 \rightarrow RO \rightarrow HO_2 \rightarrow OH \rightarrow RO_2$ ), because this drives the  $NO \rightarrow NO_2 \rightarrow NO$  cycle twice (Seinfeld and Pandis, 1998; Sect. 5.10). The radical sink (R2) occurs as a minor secondary channel in the  $RO_2 + NO$  reaction, the yield of which (A = R2/(R1 + R2)) depends on the structure of the initial VOC.

$$RO_2 + NO \rightarrow RO + NO_2$$
 (R1)

$$RO_2 + NO \rightarrow RONO_2$$
 (R2)

Values of A range from near zero for small hydrocarbons like methane to, e.g., 18 % for  $\alpha$ -pinene (Atkinson and Arey, 2003). Because of the coupling of RO<sub>x</sub> and NO<sub>x</sub> cycles leading to ozone formation, A can be determined empirically by comparing ozone production to alkyl nitrate production when losses (including mixing) of O<sub>3</sub> and RONO<sub>2</sub> can be neglected (Rosen et al., 2004; Perring et al., 2013):

$$\frac{\Delta O_3}{\Delta \Sigma A N_8} \approx \frac{2(1-A)}{A} \approx \frac{2}{A}.$$
(4)

A is the fractional yield of RONO<sub>2</sub> from RO<sub>2</sub> + NO reactions. This analysis would yield a tight, linear correlation if the airmass contained a single reactive hydrocarbon and all radical losses were due to RO<sub>2</sub> + NO. A caveat is that this yield estimate will incorporate any  $\Sigma$ AN losses that occur on



Fig. 6. Correlation plot of measured ambient  $O_3$  vs.  $\Sigma$ ANs, for the  $\approx$  6 days during the campaign when the TD-LIF was running in "3-gas" ambient mode, with standard linear regression including 95% confidence limits, slope =  $70 \pm 38$ . Only data from 12 noon-6 p.m. LT are used (see text).

the six-hour timescale of the correlation plot. This may introduce variation in measured yields from different hydrocarbon mixtures and in different climates: especially if monoterpene nitrate products are multifunctional, and in a humid environment, losses could be significant and depress the apparent yield.

Figure 6 shows ambient O<sub>3</sub> vs.  $\Sigma$ ANs for the  $\approx$  6-day period during which ambient  $\Sigma$ ANs were separately measured (3-gas mode). Much of the variability is assumed to be due to the changing ambient mix of VOCs, but additional contributions to the variability may arise from mixing, especially during the morning breakup of the nighttime thermal inversion layer; hence, in order to omit periods where mixing or deposition of O<sub>3</sub> compete with photochemical production, we include only data from noon to 18:00. The fitted slope of  $70 \pm 38$  (= 2/A) implies an effective alkyl nitrate yield of A = 2.9 + 3.4 / -1.0%, similar to a previously observed effective alkyl nitrate yield of 2.4% in an isoprene-dominated region (Horowitz et al., 2007). The major daytime VOCs at this site are MBO + isoprene (assumed to be dominantly MBO; Kim et al., 2010), with ten times lower concentration of monoterpenes. The OH rate constant with MBO is comparable to that with monoterpenes, but MBO is present at an order of magnitude larger concentration, so RO<sub>2</sub> reactivity is assumed to be dominated by that from MBO. Measured organic nitrate yields from these precursors are available but not well constrained. Alvarado et al. (1999) report a nitrate vield of 5.2 % from MBO; Chan et al. (2009) report  $10 \pm 4$  % yield. The ambient yield of 2.9% is lower than both MBO lab studies, suggesting either other VOCs contributing to RO<sub>2</sub> chemistry or additional radical losses at the RoMBAS site.

This relatively small ambient alkyl nitrate yield implies that most of the  $NO_x$  at the MFO site remains available for ozone production. In the absence of other chain-terminating reactions, the RONO<sub>2</sub> yield of 2.9% would imply a RO<sub>x</sub> cycle length of 34; however, other reactions (e.g.,  $OH+NO_2$ ,  $HO_2 + HO_2$ ) are likely to contribute and reduce this cycle chain length.

### **3.2** NO<sub>3</sub> radical fate, nighttime organic nitrate and SOA production

#### 3.2.1 Organic nitrate gas/aerosol partitioning

Examining the NO<sub>y</sub> data from gas/aerosol mode, we find another interesting diurnal pattern. While total (gas + aerosolphase)  $\Sigma PNs + \Sigma ANs$  peaks during the day, aerosol-phase-only  $\Sigma PNs + \Sigma ANs$  peaks during the night (bottom panel of Fig. 5). This can be seen most clearly in the diurnal cycle of the fraction of  $\Sigma PNs + \Sigma ANs$  in the aerosol phase (black trace, bottom panel of Fig. 5). Up to 30 % of the organic nitrate is in the aerosol phase at night, but less than 10 % during the day. Some of this may be due to repartitioning of semivolatile organic nitrates to the gas phase as temperature increases; however, we show in Sect. 3.2.3 that partitioning thermodynamics alone cannot account for the diurnal pattern observed in the bottom panel of Fig. 5.

### **3.2.2** Observed and calculated ambient NO<sub>3</sub> and lifetime

Early in the field campaign, the cavity ringdown  $NO_3/N_2O_5$  instrument was deployed to measure incanopy ambient concentrations for several nights (it was otherwise deployed to monitor oxidant exposure on a  $NO_3$  potential aerosol mass flowtube reactor, manuscript in preparation). On one of those nights, between 23 and 24 July 2011,  $NO_3$  and  $N_2O_5$  displayed a sharp peak just after midnight (shown in the bottom panel of Fig. 7), coincident with a sharp peak in  $NO_2$  and during a period when  $O_3$  was anomalously elevated (55 ppb) until past midnight. In contrast, for most of the remainder of the ambient measurement period, no  $NO_3$  was detectable due to high levels of biogenic VOCs that provided a rapid  $NO_3$  sink.

We can use the peak measured  $[NO_3]=5ppt$ ,  $[NO_2]=6.5ppb$ , and  $[O_3]=55ppb$  to determine the lifetime of NO<sub>3</sub> due to reactive losses, following the steady-state assumption derived in Eq. (8) of Brown et al. (2003a):

$$\tau_{\rm SS}(\rm NO_3) = \frac{[\rm NO_3]}{k_1[\rm O_3][\rm NO_2]} \approx k_x^{-1},$$
(5)

where  $k_1$  is the rate constant for  $O_3 + NO_2$  and  $k_x$  is the rate constant for reactive loss of NO<sub>3</sub>. The left-hand equality of this equation is the definition of the NO<sub>3</sub> steady-state life-time, and the right-hand approximation holds when  $k_x$  is the only significant loss term, i.e., when sinks of N<sub>2</sub>O<sub>5</sub> are negligible.

In this field campaign, the sinks of  $N_2O_5$  indeed appear to be smaller than reactive loss of  $NO_3$ . There are both homogeneous and heterogeneous reactions of  $N_2O_5$  with



Fig. 7. Calculated lifetime based on BVOC reactive losses (top, purple), production rate from  $NO_2 + O_3$  (middle, red) and predicted steady-state concentration (lower, blue) of  $NO_3$  throughout the RoMBAS campaign, nights only. Lower panel also shows measured  $NO_3$  (aqua) and  $N_2O_5$  (red) from two nights early in the campaign.

water, which have been examined extensively (e.g., Brown et al., 2009; Thornton et al., 2003; Wahner et al., 1998; Tuazon et al., 1983; Hjorth et al., 1987). However, the homogeneous  $N_2O_5 + H_2O$  rate constant is negligible for this dataset, especially since at 283 K absolute humidity is low (Brown et al., 2007; Crowley et al., 2011). We calculate relative rates of N2O5 heterogeneous uptake compared to NO<sub>3</sub> reaction with BVOCs (dominated at night by monoterpenes), using typical nighttime monoterpene (MT), MBO, and sesquiterpene (SQT) concentrations (measured by PTR-ToF-MS) of 0.6 ppb, 0.4 ppb, and 0.01 ppb, respectively, and the N<sub>2</sub>O<sub>5</sub>/NO<sub>3</sub> ratio of 4.1. This ratio is calculated based on the 283 K equilibrium constant ( $K_{eq} =$  $[N_2O_5]/[NO_2][NO_3])$  of  $2.05 \times 10^{-10}$  cm<sup>3</sup> molec<sup>-1</sup> (Sander et al., 2006) and 1 ppb NO2. The rate constant used for NO<sub>3</sub> with monoterpenes is  $6.1 \times 10^{-12} \text{ cm}^3 \text{ molec}^{-1} \text{ s}^{-1}$ , which is an average of the rate constants with  $\alpha$ -pinene,  $\beta$ -pinene, and  $\Delta$ -3-carene (Calvert et al., 2000), since they were present in roughly equal concentrations at RoMBAS (based on gas chromatology/mass spectrometry canister samples, not shown). For MBO, the 283 K rate constant of  $1.1 \times 10^{-14}$  cm<sup>3</sup> molec<sup>-1</sup> s<sup>-1</sup> is used; for sesquiterpenes, we use the rate constant for  $\beta$ -caryophyllene of  $1.9 \times 10^{-11}$  cm<sup>3</sup> molec<sup>-1</sup> s<sup>-1</sup>. At the reported average concentrations, this gives an average NO<sub>3</sub> lifetime ( $\tau_{NO_3,BVOC}$ ) of 14 s, dominated by MT losses ( $k_i$ [i] for MT : MBO : SQT is 1 : 0.001 : 0.05).

$$\tau_{\text{NO}_3,\text{BVOC}} = \frac{1}{k_{\text{NO}_3+\text{BVOC}}[\text{BVOC}]}$$
$$= \frac{1}{k_{\text{NO}_3,\text{MT}}[\text{MT}] + k_{\text{NO}_3,\text{MBO}}[\text{MBO}] + k_{\text{NO}_3,\text{SQT}}[\text{SQT}]} \quad (6)$$

The heterogeneous uptake of N<sub>2</sub>O<sub>5</sub> onto aqueous particles depends on the uptake coefficient ( $\gamma$ ), molecular speed of N<sub>2</sub>O<sub>5</sub> (v), and the aerosol surface area per volume (SA), parameterized as a first-order rate constant:

$$k_{\rm het} = \frac{\gamma v}{4} SA. \tag{7}$$

Measured uptake coefficients on liquid water range from 0.02 to 0.04 around 283 K (Sander et al., 2006); we use an upper limit value  $\gamma = 0.04$ . At a typical observed (SEMS) nighttime aerosol surface area per air volume of  $240 \,\mu\text{m}^2 \,\text{cm}^{-3}$ , N<sub>2</sub>O<sub>5</sub> molecular speed of  $2.3 \times 10^4 \,\text{cm} \,\text{s}^{-1}$ , this gives an average N<sub>2</sub>O<sub>5</sub> lifetime ( $\tau_{N_2O_5,\text{het}}$ ) of 440 s.

$$\tau_{\rm N_2O_5,het} = \frac{1}{k_{\rm het}} \tag{8}$$

Applying the N<sub>2</sub>O<sub>5</sub>/NO<sub>3</sub> ratio of 4.1 and comparing to the  $\tau_{NO_3,BVOC}$  to  $\tau_{N_2O_5,het}$ , this estimate suggests that the rate of NO<sub>3</sub> with BVOC is 15 times faster than uptake of N<sub>2</sub>O<sub>5</sub> to aqueous particles.

We apply the NO<sub>3</sub> steady-state approximation (Eq. 5) to the case of the high-NO<sub>3</sub> night (23–24 July), to compare the calculated lifetime of NO<sub>3</sub> from steady state to that calculated from reactive loss via  $k_x$ . For the average ambient nighttime temperature of  $\approx 283$  K, the rate constant of O<sub>3</sub> with NO<sub>2</sub> is  $k_1 = 2.3 \times 10^{-17}$  cm<sup>3</sup> molec<sup>-1</sup> s<sup>-1</sup> (Atkinson et al., 2004). At the concentrations listed above, this gives  $k_x^{-1} = \tau_{SS}(NO_3) = 30$  s. Calculating the NO<sub>3</sub> lifetime instead from the dominant MT loss ( $k_x = k_{NO_3,MT}$ ), at the observed average MT concentrations that night (7p–5a local) of 0.28 ppb, we find  $\tau_{NO_3,MT} = 29$  s, in excellent agreement with the steady-state prediction. Hence, N<sub>2</sub>O<sub>5</sub> + NO<sub>3</sub> losses are in fact dominated by reactive loss of NO<sub>3</sub> to biogenic VOCs, with a lifetime in the canopy on the order of seconds, making this a very efficient sink of both species.

Other than the period described above, no  $NO_3$  was observed on four additional nights the CRDS instrument was used to monitor ambient  $NO_3$  and  $N_2O_5$ , so it was subsequently deployed to make other measurements. In order to estimate ambient concentrations of  $NO_3$  at RoMBAS throughout the campaign, we can use Eq. (6) to determine an estimate of the NO<sub>3</sub> lifetime, using the known mix, concentration time series, and NO<sub>3</sub> rate constants of monoterpenes, sesquiterpenes, and MBO as described above. This calculation shows that maximum NO<sub>3</sub> lifetimes throughout the campaign are consistently < 200 s during the night. The top panel of Fig. 7 shows the result of this calculation,  $\tau_{NO_3}$ , for the entire campaign. Daytime results have been omitted in this plot, since NO<sub>3</sub> lifetime will be limited by photolysis and reaction with NO during the day.

We also calculate a full time series of the production rate of NO<sub>3</sub> based on measured O<sub>3</sub>, NO<sub>2</sub>, and temperature (P(NO<sub>3</sub>)= $k_1$ [O<sub>3</sub>][NO<sub>2</sub>]), shown in the middle panel of Fig. 7. We predict production rates of up to 0.15–0.2 ppt s<sup>-1</sup> (0.5–0.7 ppb h<sup>-1</sup>), with peaks generally in the early evening when NO<sub>2</sub> rise and O<sub>3</sub> decline intersect (see Fig. 3). Combining these production and loss terms, we derive the time series shown in the bottom panel of Fig. 7, the predicted nighttime steady-state nitrate concentration:

$$NO_{3,SS} = \frac{P(NO_3)}{(\tau_{NO_3,BVOC})^{-1}}.$$
(9)

Because NO<sub>3</sub> will also be reactively lost to later generation oxidation products, this represents a conservative upper limit on the steady-state concentration of NO<sub>3</sub>. During the night of 24 July when a sustained peak NO<sub>3</sub> of 5 ppt was measured, our steady-state calculation accurately predicts the timing of the peak, but overestimates its magnitude by almost a factor of 4 (predicted NO<sub>3</sub> concentration was 18 ppt). For most of the campaign, the steady-state NO<sub>3</sub> predicted in this way is significantly lower, near the detection limit of the CRDS instrument, so it may not be surprising that we did not detect it. Golz et al. (2001) also sought to measure NO<sub>3</sub> using DOAS in a European forest and were unable to detect it.

# **3.2.3** Evidence for NO<sub>3</sub>-initiated organic nitrate SOA production at night: predicted organic nitrate structures and modeled thermodynamics

When NO<sub>3</sub> oxidation of monoterpenes is modeled in the Leeds Master Chemical Mechanism (MCMv3.2; http://mcm. leeds.ac.uk/MCM; Jenkin et al., 1997; Saunders et al., 2003), the major organic nitrates produced by daytime (largely OHinitiated) and nighttime (largely NO<sub>3</sub>-initiated) oxidation are distinct (Fig. 8). We carried out two 12h model runs with the chemical conditions observed at RoMBAS (temperature,  $[O_3]$ ,  $[NO_2]$ , [monoterpenes]) for daytime photochemistry and dark nighttime chemistry, and tracked the > 900 products, identifying the top six organic nitrates in terms of peak concentration during that time period. Daytime simulations were run at 25 °C and 30 % RH, nighttime simulations at 10 °C and 80 % RH. For consistency, both were initiated with the same gas-phase concentrations of reactive precursors: 35 ppb O<sub>3</sub>, 4 ppb NO<sub>2</sub>, 0.5 ppb  $\alpha$ -pinene, 0.4 ppb  $\beta$ pinene, and 0.1 ppb limonene ( $\Delta$ -3-carene is not yet in the



**Fig. 8.** Major organic nitrate oxidation products predicted by the MCM for daytime and nighttime simulations, initialized with observed monoterpene and oxidant concentrations at MFO. Structures are shown in order of peak concentration, with highest on the left. Outlined structures show the more volatile daytime nitrates (orange, predicted  $p_{\rm vap} > 10^{-3}$  Torr;  $C^* > 10^4 \,\mu {\rm g m}^3$ ) and less volatile nighttime nitrates (blue, predicted  $p_{\rm vap} \approx 10^{-5}$  Torr;  $C^* \approx 100 \,\mu {\rm g m}^3$ ).

MCM, and MBO was omitted because it is not expected to produce condensable products).

In both day- and nighttime cases, C10 hydroxynitrates are among the top products (Fig. 8). The remaining top organic nitrates show some differences, however: during the day, two fragmentation products are predicted (outlined in orange), which SIMPOLv.1 predicts to have higher volatility. The structure on the left, a C9 keto-nitrate, has a predicted vapor pressure of  $5.7 \times 10^{-3}$  Torr at daytime average temperature of 298 K, which is equivalent to a  $C^*$  of 6.2  $\times$  $10^4 \,\mu g \,\mathrm{m}^3$ ; the structure on the right, a multifunctional peroxynitrate produced glycolaldehyde, which has a predicted vapor pressure of  $1.5 \times 10^{-3}$  Torr ( $C^* = 1.2 \times 10^4 \,\mu g \,m^3$ ). At night, in contrast, two hydroperoxynitrates are observed (outlined in blue), which are predicted to have lower volatility  $(1.0 \times 10^{-5}$  Torr at nighttime average temperature of 283 K,  $C^* = 130 \,\mu \text{g}\,\text{m}^3$ ). Hence, accounting for day/night temperature differences observed during RoMBAS, nighttime NO3produced organic nitrates have substantially lower volatility, consistent with the observation that a larger fraction of organic nitrates is in the aerosol phase at night.

Although daytime- and nighttime-produced organic nitrates are predicted to include distinct structures, some of the observed diurnal cycle in aerosol-phase fraction of nitrates (bottom panel of Fig. 5) may be explained by thermodynamics: as temperature cools at night, total organic nitrate repartitions to favor the condensed phase. We can test this by modeling the fraction in the aerosol phase for a series of organic nitrate structures deemed likely as monoterpene oxidation products. We employ the absorptive partitioning formalism introduced in Eq. (2) to predict background aerosoldependent (using AMS-measured organic aerosol) partitioning of hypothesized structures, using SIMPOLv.1 to predict



**Fig. 9.** Observed (black, same as in Fig. 5) and modeled fraction of organic nitrate in the aerosol phase over the diurnal cycle in temperature (blue) and background organic aerosol mass (measured by AMS, brown), based on calculated vapor pressures for a series of possible nitrate structures (indicated in legend). None of these structures can capture the observed dynamic range in aerosol fraction. Bottom panel shows aerosol fraction for the same series of molecules, demonstrating that the observed aerosol fraction does not fully scale with temperature as simple repartitioning of the same nitrates would predict. Data are normalized to peak fraction to facilitate comparison of the lineshapes.

the temperature-dependent vapor pressures of hypothesized structures. We calculate the average diurnal cycle in several multifunctional nitrates in this way (Fig. 9), using average diurnal cycles in temperature and ambient organic aerosol mass loading as inputs. These model predictions (colors) are shown with comparison to the observed aerosol nitrate fraction (black).

First, we note that neither daytime ( $C_9$  keto-nitrate or PN) nor nighttime ( $C_{10}$  hydroperoxy-nitrate) MCM-predicted products would partition appreciably to the aerosol phase

in this simple model: the predicted volatilities are simply too high. As discussed above, Barley and McFiggans (2010) have pointed out that vapor pressure estimation methods differ widely; in a test of sensitivity of calculated OA formation to vapor pressure prediction, Barley and McFiggans observed that predicted multifunctional compounds were 100– 1000 times more volatile than needed to contribute significantly to OA.

We find a similar result here. In order to bracket the observed aerosol fraction, one or two additional functional groups must be added ( $C_{10}$  keto-hydroxy-nitrate,  $2.0 \times 10^{-6}$  Torr,  $C^* = 27 \mu \text{gm}^3$  at 283 K; or  $C_{10}$  dihydroxy-nitrate,  $8.0 \times 10^{-8}$  Torr,  $C^* = 1.0 \mu \text{gm}^3$ ). Alternatively, similarly low volatilities can be realized if the precursor BVOC is a sesquiterpene ( $C_{15}$ ) with only two functional groups (e.g.,  $C_{10}$  hydroperoxy-nitrate,  $4.6 \times 10^{-8}$  Torr,  $C^* = 0.7 \mu \text{gm}^3$  at 283 K). Sesquiterpenes were measured at RoMBAS at much lower concentrations than monoterpenes (nighttime peak < 0.01 ppb, PTR-ToF-MS), but may have a disproportionate impact on aerosol formation because less functionalization is required to achieve lower volatility.

Furthermore, we observe that while the fraction in the aerosol phase falls in the range of these selected trifunctional molecular structures, none of these individual species captures the full dynamic range of aerosol fraction observed. As shown in the bottom panel of Fig. 9, the aerosol fraction is not simply a function of temperature: at the lowest temperatures the observed aerosol fraction is higher than the linear extrapolation that individual structures would predict. This again suggests that at least a subset of the organic nitrates contributing to aerosol during the day and night are in fact of different molecular structures with differing aggregate volatility. After condensation, these organic nitrates may also continue to evolve via condensed-phase reactions, resulting in a different aerosol composition than dictated by the initial condensing species. No such continuing transformations are treated in this simple thermodynamic model.

This apparent combination of thermodynamic repartitioning and different production mechanisms of organic nitrates contrasts with a recent study (Perraud et al., 2012), in which observed SOA formation from  $\alpha$ -pinene could only be explained by irreversible, kinetically limited uptake of organic nitrates onto aerosol. This difference may be explained by the BVOC precursor mix, if the mechanism is unique to  $\alpha$ pinene, or by the elevated oxidant (1.5 ppm O<sub>3</sub>) and aerosol mass (30–2000 µg m<sup>3</sup>) in those chamber experiments causing the SOA formation mechanism to be different than under ambient conditions at RoMBAS (30 ppb O<sub>3</sub> and 2 µg m<sup>3</sup> PM<sub>1</sub> at night).

### **3.2.4** Further evidence for NO<sub>3</sub>-initiated organic nitrate SOA production at night: oxidant contributions

We find further observational evidence that nighttime chemistry is dominated by NO<sub>3</sub>-initiated oxidation rather than



**Fig. 10.** Rates of reaction of NO<sub>3</sub> and O<sub>3</sub> with monoterpenes averaged for entire campaign, nighttime data only, using measured [O<sub>3</sub>] and [monoterpenes], and using the production rate of NO<sub>3</sub> ( $k_1$ [NO<sub>2</sub>][O<sub>3</sub>]) to stand in for the rate of NO<sub>3</sub>+MT, since it is assumed that every NO<sub>3</sub> produced reacts with MT. Right axis: the fraction of total MT reactivity due to reactions with NO<sub>3</sub> (black) suggests a greater role for NO<sub>3</sub> than O<sub>3</sub> as the initial oxidant in nighttime oxidation.

ozonolysis when we evaluate their relative rates for all of the campaign nighttime data (Fig. 10). We approximate the rate of NO<sub>3</sub> + MT as the rate of NO<sub>3</sub> production ( $P_{NO_3} = k_1[NO_2][O_3]$ ), because we assume that at night nearly every NO<sub>3</sub> produced is consumed by reaction with monoterpenes. We calculate the ozonolysis rate ( $R_{O_3+MT}$ ) as  $k_O[O_3][MT]$ , where  $k_O = 5.7 \times 10^{-17} \text{ cm}^3 \text{ molec}^{-1} \text{ s}^{-1}$ , the average of rates of ozonolysis of  $\alpha$ -pinene,  $\beta$ -pinene and  $\Delta$ -carene (Calvert et al., 2000), the three dominant monoterpenes observed at MFO. We calculate the fraction of monoterpenes oxidized by NO<sub>3</sub> as  $P_{NO_3}/(P_{NO_3} + R_{O_3+MT})$ and find that throughout the campaign,  $60\% \pm 10\%$  of the nighttime monoterpene reactivity goes via NO<sub>3</sub>.

### 3.3 Aerosol organic nitrate concentrations derived from AMS measurements

The nighttime peak in aerosol organic nitrate concentration is further corroborated by AMS measurements of this species (Fig. 11). Farmer et al. (2010) reported that due to fast thermal dissociation of organic nitrates prior to ionization, the AMS will quantify the nitrate moiety almost exclusively as NO<sup>+</sup> and NO<sub>2</sub><sup>+</sup> ions. Several groups have reported that the NO<sub>2</sub><sup>+</sup>/NO<sup>+</sup> ratios observed in AMS spectra for organic nitrates are typically 2–3 times lower than for NH<sub>4</sub>NO<sub>3</sub> (Fry et al., 2009; Bruns et al., 2010; Farmer et al., 2010; Liu et al., 2012). Assuming these ratios are stable, the measured ratio can be used to separately quantify ammonium and organic nitrates as (Farmer et al., 2010)

$$\text{RONO}_{2,\text{frac}} = \frac{(R_{\text{ambient}} - R_{\text{NH}_4\text{NO}_3})(1 + R_{\text{RONO}_2})}{(R_{\text{RONO}_2} - R_{\text{NH}_4\text{NO}_3})(1 + R_{\text{ambient}})}.$$
 (10)



**Fig. 11. (a)** Concentration of AMS organic and inorganic nitrate, calculated based on the  $NO_2^+/NO^+$  field ratios and  $R_{RONO_2} = 0.13$ , compared to aerosol organic nitrate measurements with the TD-LIF system, for the six days the ambient measurements of both instruments overlapped during RoMBAS. (b) Scatter plot of calculated AMS organic and inorganic nitrate with TD-LIF aerosol organic nitrate. (c) Total organic aerosol mass as measured by the AMS, and  $NO_2^+/NO^+$  ( $R_{ambient}$ ) used to calculate organic nitrate fraction in (a). Also shown are the  $NO_2^+/NO^+$  ratio determined in calibration with inorganic nitrate as well as the estimated  $R_{RONO_2}$  ratio. A 3-point smoothed version of the ratio was used when calculating RONO\_2. (d) Cation vs. anion balance, evaluated by plotting measured AMS ammonium vs. predicted ammonium for two cases: assuming all AMS nitrate is inorganic and assuming it is all organic.

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According to Eq. (10),  $R_{\text{ambient}}$  (the ratio of NO<sub>2</sub><sup>+</sup>/NO<sup>+</sup> in ambient data) can be used to calculate the fraction of organic nitrate, RONO<sub>2,frac</sub>, if both  $R_{\text{NH}_4\text{NO}_3}$  (for the ammonium nitrate calibrant) and  $R_{\text{RONO}_2}$  (measured or assumed average NO<sub>2</sub><sup>+</sup>/NO<sup>+</sup> for organic nitrates) are known. It should be noted that this expression only applies if NH<sub>4</sub>NO<sub>3</sub> is the solely important inorganic nitrate in the submicron mode, since other nitrate salts have different fragmentation ratios. However, as noted above, it is unlikely that reacted dust or sea salt contributed significantly to the submicron aerosol during RoMBAS.

Both  $R_{\rm NH_4NO_3}$  and  $R_{\rm RONO_2}$  depend on the particular AMS instrument and AMS tuning; hence they need to be determined at the same instrument settings as  $R_{\rm ambient}$ . While  $R_{\rm NH_4NO_3}$  can be determined from the routine in-field calibrations, estimating  $R_{\rm RONO_2}$  is more difficult as there is some variability in the ratios found for pure organic nitrates (Farmer et al., 2010), and standards for the specific organic nitrates important at a field location are typically not available at the time of the field study. One possible way to circumvent this issue is to use the observation that  $R_{\rm NH_4NO_3}$ and  $R_{\rm RONO_2}$  typically co-vary proportionally between instruments. Based on that observation, we derive field values for  $R_{\text{RONO}_2}$  based on previous laboratory measurements of the "ratio of ratios"  $\chi$ :

$$\chi = \frac{R_{\rm NH_4NO_3}}{R_{\rm RONO_2}}.$$
(11)

Based on the calibrations reported by Farmer et al. (2010), we calculate a value for  $\chi$  of 2.25 ± 0.35. Using the (very constant)  $R_{\rm NH_4NO_3}$  of 0.295±0.005 measured throughout the in-field IE calibrations, this would result in  $R_{\rm RONO_2} = 0.13 \pm 0.02$ .

Figure 11c shows  $R_{\text{ambient}}$  for ambient data for the six days where the AMS and the TD-LIF ambient aerosol datasets overlap.  $R_{\text{ambient}}$  is quite constant within the noise at  $\approx 0.13$ , with the exception of a brief spike around midnight of 20 August. This implies that the fraction of total nitrate that is organic was essentially constant, with no diurnal cycle, except for a few brief inorganic nitrate episodes such as on 20 August. Given the very different night and day chemistry, it seems unlikely that this fraction would stay constant, unless it happens to correspond to 100 % organic nitrate, which is consistent with the  $R_{\text{RONO}_2}$  value derived above.

Another piece of evidence that supports the dominance of organic nitrates in the submicron nitrate at this location is the anion vs. cation balance. A linear regression of measured AMS ammonium vs. predicted, assuming that all measured nitrate is inorganic, leads to a negative intercept and a slope of less than 1 (Fig. 11d; slope = 0.92). The deviation of the slope from 1 is substantially larger than the 2% observed for NH<sub>4</sub>NO<sub>3</sub> calibrations. Such a deviation is normally associated with partially acidic sulfate, but in those cases the deviation typically varies substantially in time (e.g., Zhang et al., 2007), and that would also be expected here given the large variability of  $SO_2$  (Fig. 4). The slope can also be due to mistakenly assuming RONO<sub>2</sub> to be inorganic nitrate (Farmer et al., 2010). If we instead exclude all of the aerosol nitrate from the neutralization calculation, the slope becomes very close to 1 (0.99) and the negative intercept disappears. As can be inferred from the tightness of the scatter plot, for most of the campaign this ratio was very constant. While this could reflect an aerosol of fairly constant free acidity, that would be highly unusual for such a long time period. Hence, the most likely explanation is that the anions were fully neutralized and thus most of the nitrate was organic.

Calculated AMS organic nitrate concentrations are compared with the TD-LIF data in Fig. 11a. The overall agreement between the two organic nitrate measurements, both in amplitude and time dependence, is very good. Figure 11b compares AMS organic and inorganic nitrates with the TD-LIF aerosol nitrate. As expected, there is little correlation of TD-LIF nitrate with AMS inorganic nitrate  $(r^2 = 0.1)$ , while the correlation with AMS organic nitrate correlation is fair  $(r^2 = 0.53)$ , despite separate inlets, the low concentrations involved, and the difference in analytical methods. Both datasets were averaged to a longer timebase (i.e., longer than the 10 min original database). The slope for the resampled data, 1.16, seems to indicate a slightly lower AMS response, but is well within the stated accuracies for the instruments (30% for the AMS, 25% for the TD-LIF). The dominance of aerosol organic nitrate over the inorganic form at this location contrasts with polluted urban locations such as Riverside, CA, where the aerosol nitrate is dominated by the inorganic form (Farmer et al., 2010).

The concentrations of aerosol organic nitrate concentrations shown in Fig. 11a for both instruments only quantify the NO<sub>3</sub> functional group and do not reflect the actual mass of organic molecules containing nitrate functional groups that are present in the aerosol (since both techniques detect the nitrate functional group after it has dissociated from the organic parent structure). Estimating a total mass for the organic molecules containing nitrate groups is difficult, since the average molecular mass of the organic species per NO<sub>3</sub> group is needed for this calculation. Assuming that the  $C_{10}$ keto-hydroxy-nitrate as shown in Fig. 9 is a representative organic nitrate molecule, the molecular weight of the molecule minus the nitrate group is  $168 \text{ g mol}^{-1}$ , 2.7 times larger than the mass of the nitrate group. For the periods considered here, that implies a concentration of  $\approx 0.2-0.5 \,\mu g \, m^{-3}$  of organic species with nitrate groups, representing 6-20% of the total organic aerosol mass at a given time. As first reported by Farmer et al. (2010), this also has implications for the O/C and N/C ratios that the AMS reports, since the default analysis algorithm used for field studies assumes that only CHO, CHN, and CHON ions are organic and all of the NO<sup>+</sup><sub>x</sub> ions are assigned to the "nitrate" species. For RoMBAS, including nitrate from RONO<sub>2</sub> in the calculation of organic N/C would increase by 0.013 on average (up to 0.025 at night and about 0.009 during the day), while typical AMS N/C values which do not include organic nitrate are  $\approx$  0.01–0.02 (e.g., Hayes et al., 2012; Docherty et al., 2011). Likewise, O/C ratios would be enhanced 0.03 on average, with spikes of up to 0.07 at night, resulting in about a 5 % increase for the campaign average.

# 3.4 Estimated alkyl nitrate production rate, daytime vs. nighttime

Finally, we wish to put the nighttime production of alkyl nitrates into context with the total nitrate production. We use both our empirically determined net daytime yield of RONO<sub>2</sub> from RO<sub>2</sub> + NO reactions A = 2.9% (Sect. 3.1.4) as well as higher literature values from chamber studies, to determine two estimates of daytime production rates of organic nitrates from OH-initiated reactions followed by RO<sub>2</sub>+NO, for comparison to nighttime NO<sub>3</sub>-initiated chemistry.

We calculate the daytime production rate of alkyl nitrates as

 $P_{\text{day}} = (k_{\text{OH}_1}[\text{OH}][\text{MT}]\alpha_{\text{MT}} + k_{\text{OH}_2}[\text{OH}][\text{MBO}]\alpha_{\text{MBO}})F_{\text{RO}_2 + \text{NO}}$ (12)

where  $k_{\text{OH}_1} = 7.4 \times 10^{-11} \text{ cm}^3 \text{ molec}^{-1} \text{ s}^{-1}$  (Calvert et al., 2000) is the average of the rate constants for OH reaction with  $\beta$ -pinene,  $\alpha$ -pinene, and  $\Delta$ -carene at this temperature (these three dominant monoterpenes are present in approximately equal concentrations at RoMBAS), and  $k_{\text{OH}_2} = 6.5 \times 10^{-11} \text{ cm}^3 \text{ molec}^{-1} \text{ s}^{-1}$  (293 K, Rudich et al., 1995) is the rate constant of OH with MBO. The values for  $\alpha_{\text{MT}}$  and  $\alpha_{\text{MBO}}$  are either chosen to both be equal to *A*, the empirically determined branching ratio of alkyl nitrate production from RO<sub>2</sub> + NO reactions, or to be equal to values from literature chamber studies (23 % for monoterpenes, Perring et al., 2013; and 7 % for MBO, Alvarado et al., 1999; Chan et al., 2009). The former production rate is lower by about a factor of 5.  $F_{\text{RO}_2+\text{NO}}$  is the fraction of RO<sub>2</sub> radicals that react with NO rather than other RO<sub>2</sub> or HO<sub>2</sub>.

The calculation of  $F_{\text{RO2+NO}}$  is based on diurnal RO<sub>2</sub> and HO<sub>2</sub> data with calculated NO steady-state concentrations:

$$[NO]_{ss} = \frac{[NO_2]j_{NO_2}}{k_{O_3+NO} [O_3] + k_{R/HO_2+NO} [RO_2 + HO_2]},$$
 (13)

where  $j_{\text{NO}_2}$  is the photolysis rate of NO<sub>2</sub>,  $k_{\text{O}_3+\text{NO}} = 1.4 \times 10^{-12} \exp(-1310/T) \text{ cm}^3 \text{ molec}^{-1} \text{ s}^{-1}$  and  $k_{\text{R/HO}_2+\text{NO}} = 7.9 \times 10^{-12} \text{ cm}^3 \text{ molec}^{-1} \text{ s}^{-1}$ , which is the average of rate constants of HO<sub>2</sub> + NO and CH<sub>3</sub>O<sub>2</sub> + NO (Sander et al., 2006), using instantaneous measured temperature, [O<sub>3</sub>],

[RO<sub>2</sub>], [NO<sub>2</sub>] and [HO<sub>2</sub>]. Typical daytime [HO<sub>2</sub>] peaked midday at 100 ppt; [RO<sub>2</sub>] was 45–55 ppt throughout the day, and [NO] peaked at 80 ppt at 8:30 and 14:30 and was > 40 ppt throughout the day. To determine the relative rates of RO<sub>2</sub>+NO reactions, we use rate constants for CH<sub>3</sub>O<sub>2</sub> (Atkinson et al., 2004). We use the CH<sub>3</sub>O<sub>2</sub> rate constants because those reactions have been well characterized; however, since they are for a smaller RO<sub>2</sub> than the ambient mix, this likely introduces some error.

Nighttime production rate of alkyl nitrates is calculated as

$$P_{\text{night}} = [\text{NO}_3](k_{\text{MT}}[\text{MT}]Y_{\text{MT}} + k_{\text{MBO}}[\text{MBO}]Y_{\text{MBO}}), \quad (14)$$

where  $k_i$  and  $Y_i$  are NO<sub>3</sub> rate constants and organic nitrate yields, respectively, from monoterpenes and MBO. Again, we use  $6.1 \times 10^{-12}$  cm<sup>3</sup> molec<sup>-1</sup> s<sup>-1</sup> for the NO<sub>3</sub> + monoterpene rate, the average of rate constants with  $\alpha$ -pinene,  $\beta$ -pinene, and  $\Delta$ -3-carene (Calvert et al., 2000). The rate constant for NO<sub>3</sub> with MBO is  $1.1 \times 10^{-14}$  cm<sup>3</sup> molec<sup>-1</sup> s<sup>-1</sup> (Rudich et al., 1996) at 283 K. Yields of organic nitrates from MT and MBO are taken to be 45 % (Fry et al., 2009; measured nitrate yield for  $\beta$ -pinene) and 13 % (Fantechi et al., 1998), respectively. The few measurements of organic nitrate yields from [NO<sub>3</sub>] + BVOC that have been made are reviewed in Atkinson and Arey (2003), and span the range of 14 % for  $\alpha$ -pinene to 66 % for  $\Delta$ -3carene, making this  $\beta$ -pinene figure a reasonable average behavior for the monoterpenes present at RoMBAS.

Rollins et al. (2012) also recently assessed aerosol organic nitrate yield from NO<sub>3</sub>-initiated SOA formation, in Bakersfield, California, an urban site in the agricultural San Joaquin Valley with much higher NO<sub>2</sub>, O<sub>3</sub>, and biogenic VOC concentrations, and particle loading. At that site, BVOC concentrations are so high that RONO<sub>2</sub> production is limited by NO<sub>3</sub> production via NO<sub>2</sub> + O<sub>3</sub>, which proceeds at typical rates of  $1-2 \text{ ppb} \text{h}^{-1}$  (Rollins et al., 2012). Significantly lower organic nitrate production is observed at RoMBAS (0.05 ppbh<sup>-1</sup>), due to much lower pollutant concentrations, but at both sites the resulting contribution of RONO<sub>2</sub> to organic aerosol loading is substantial.

A comparison of daytime (OH-initiated) and nighttime (NO<sub>3</sub>-initiated) rates of alkyl nitrate production is shown in Fig. 12, showing a diurnal cycle that indicates the rates are comparable in magnitude, with the nighttime production rate falling between the two estimates of daytime organonitrate production. The difference in these two daytime production rate estimates suggests that MBO and monoterpenes are not the only sinks of OH at this site: since the observed overall A is lower than the yield for MBO, it is likely that at this site, substantial OH loss goes to RO2 which have a low or 0% yield of organic nitrate, bringing the overall yield down. For example, the OH oxidation of first-generation products of MBO oxidation may be important. Note also that this analysis omits contributions from ozone, which may contribute to alkyl nitrate formation both during the day and night, but which we cannot constrain.



**Fig. 12.** Calculated rates of production of organic nitrates for daytime OH-initiated (orange/yellow) and nighttime NO<sub>3</sub>-initiated (blue) mechanisms. The fraction of RO<sub>2</sub> radicals that react with NO ( $F_{\rm RO_2+NO}$ , black) ranges from  $\approx 40\%$ –90% during the day. The daytime production of organic nitrates is calculated using the observed *A* of 2.9% (orange) and, alternately, using literature chamber organic nitrate yields of 23% for monoterpenes and 7% for MBO (yellow).

These production rates are both likely underestimates since they consider only first-generation oxidation chemistry initiated by OH or NO<sub>3</sub>, and omit other VOC precursors that may contribute to additional organic nitrate production, e.g., sesquiterpenes. We note further that the empirically derived A, the daytime organic nitrate branching ratio, implicitly includes any fast chemical losses of daytime RONO<sub>2</sub>, including potentially rapid hydrolysis in the aerosol phase (Liu et al., 2012), while the nighttime source calculation is the gross formation rate, with any nighttime RONO<sub>2</sub> losses unaccounted for. Based on structural differences - daytime organic nitrates are derived from shorter-chain hydrocarbons and are thus likely to have more oxidized functional groups per carbon and hence be more hydroscopic - we anticipate wet deposition losses to be faster for daytime than nighttime RONO<sub>2</sub>. Thus, we believe it is more important that these losses are incorporated for the daytime RONO<sub>2</sub> production rate, and we expect the error in net nighttime production rate due to omission of chemical losses to be relatively small.

Because this organic nitrate production is occurring in very different volumes during the day (afternoon boundary layer height of 800–1500 m, radiosonde measurements) vs. night (shallow nocturnal boundary layer height of order 50 to 90 m), the total mass of organic nitrate produced during the day is likely to be larger if the NO<sub>x</sub> or BVOC level aloft during the night is significantly smaller than that at the surface. Nevertheless, the lower-volatility nighttime organic nitrates produced in the shallower boundary layer lead to a peak in observed aerosol organic nitrate at night. This is consistent with the hypothesis that nighttime NO<sub>3</sub>-initiated chemistry

is a dominant source of aerosol organic nitrate surface concentrations in the Colorado front range.

#### 4 Conclusions

Substantial morning organic nitrate production is observed at a pine forest site with NO<sub>2</sub> mixing ratios in the 0.5-3 ppb range in the Colorado front range, attributed to BVOC reactions with oxidants present in the urban outflow of Denver and Colorado Springs. The fraction of this organic nitrate in the aerosol phase is highest during nighttime, suggesting that nighttime NO3-initiated chemistry produces more condensable organic nitrates, in addition to thermal repartitioning of all organic nitrates. Comparison of TD-LIF and AMS measurements of organic nitrates shows that aerosol nitrate signal at this site is dominated by organic nitrate, and the aerosol organic nitrate concentrations measured by these two distinct instruments agree very well. Despite the distance from urban centers, NO<sub>3</sub>-initiated oxidation is observed to be an important reaction at this site, which is assumed to be representative of forested areas within the Colorado front range foothills. NO<sub>3</sub> radical is responsible for the majority of nighttime monoterpene losses, and contributes substantially to organic aerosol loading.

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