Controlled nitric oxide production via O(1D) + N2O reactions for use in oxidation flow reactor studies

Andrew Lambe1,2, Paola Massoli1, Xuan Zhang1,a, Manjula Canagaratna1, John Nowak1,b, Conner Daube1, Chao Yan3, Wei Nie4,3, Timothy Onasch1,2, John Jayne1, Charles Kolb1, Paul Davidovits2, Douglas Worsnop1,3, and William Brune5

1Aerodyne Research, Inc., Billerica, Massachusetts, USA
2Chemistry Department, Boston College, Chestnut Hill, Massachusetts, USA
3Physics Department, University of Helsinki, Helsinki, Finland
4Joint International Research Laboratory of Atmospheric and Earth System Sciences, School of Atmospheric Sciences, Nanjing University, Nanjing, China
5Department of Meteorology and Atmospheric Sciences, The Pennsylvania State University, University Park, Pennsylvania, USA

aCurrent address: Atmospheric Chemistry Observations & Modeling Laboratory, National Center for Atmospheric Research, Boulder, Colorado, USA
bCurrent address: Chemistry and Dynamics Branch, NASA Langley Research Center, Hampton, Virginia, USA

Correspondence to: Andrew Lambe (lambe@aerodyne.com) and William Brune (whb2@psu.edu)

Received: 30 November 2016 – Discussion started: 6 January 2017
Revised: 11 May 2017 – Accepted: 23 May 2017 – Published: 22 June 2017

Abstract. Oxidation flow reactors that use low-pressure mercury lamps to produce hydroxyl (OH) radicals are an emerging technique for studying the oxidative aging of organic aerosols. Here, ozone (O3) is photolyzed at 254 nm to produce O(1D) radicals, which react with water vapor to produce OH. However, the need to use parts-per-million levels of O3 hinders the ability of oxidation flow reactors to simulate NOx-dependent secondary organic aerosol (SOA) formation pathways. Simple addition of nitric oxide (NO) results in fast conversion of NOx (NO + NO2) to nitric acid (HNO3), making it impossible to sustain NOx at levels that are sufficient to compete with hydroperoxy (HO2) radicals as a sink for organic peroxy (RO2) radicals. We developed a new method that is well suited to the characterization of NOx-dependent SOA formation pathways in oxidation flow reactors. NO and NO2 are produced via the reaction O(1D) + N2O → 2NO, followed by the reaction NO + O3 → NO2 + O2. Laboratory measurements coupled with photochemical model simulations suggest that O(1D) + N2O reactions can be used to systematically vary the relative branching ratio of RO2 + NO reactions relative to RO2 + HO2 and/or RO2 + RO2 reactions over a range of conditions relevant to atmospheric SOA formation. We demonstrate proof of concept using high-resolution time-of-flight chemical ionization mass spectrometer (HR-ToF-CIMS) measurements with nitrate (NO3) reagent ion to detect gas-phase oxidation products of isoprene and α-pinene previously observed in NOx-influenced environments and in laboratory chamber experiments.

1 Introduction

Recent atmospheric observations supported by experimental and theoretical studies show that highly oxidized molecules (HOMs), together with sulfuric acid, are involved in the initial nucleation steps leading to new particle formation (NPF) (Donahue et al., 2013; Riccobono et al., 2014; Kurten et al., 2016). HOMs form rapidly in the gas phase via auto-oxidation processes (Crounse et al., 2013; Rissanen et al., 2014), and many tend to condense irreversibly (Ehn et al., 2014). Following NPF, semivolatile organic compounds (SVOCs) with higher vapor pressures condense on newly formed aerosols at rates influenced by their volatil-
ity (Donahue et al., 2012), ultimately driving nanoparticle growth towards formation of cloud condensation nuclei (CCN) (Pierce et al., 2012; Riipinen et al., 2012). NPF events may produce as much as 50% of global CCN (Merikanto et al., 2009; Yu and Luo, 2009). However, mechanisms that govern the formation of specific HOMs and condensation of SVOCs in various source regions are largely unknown.

The extent to which NPF and growth are influenced by natural and anthropogenic emissions, separately and together, is still unknown. In some locations, biogenic secondary organic aerosol (SOA) formation is enhanced by anthropogenic carbaceous aerosol particles, SO₄, and/or NOₓ (Carlton et al., 2010; Shilling et al., 2013; Xu et al., 2015). At the moment, one can only speculate about some of the possible synergistic or antagonistic chemical mechanisms regulating these processes. For example, anthropogenic emissions can enhance biogenic SOA formation by providing seed particles for condensable biogenic vapors. On the other hand, isoprene can slow down the formation of SOA from other volatile organics, possibly by depleting the local concentration of OH without itself producing significant SOA yields (Pugh et al., 2011). Globally the source strength of anthropogenic SOA is poorly constrained, with an uncertainty of at least a factor of 2 or 3 (Spracklen et al., 2011). Large uncertainties in pre-industrial aerosol emissions and processes further confound our understanding of the direct and indirect effects of anthropogenic aerosol emissions (Carslaw et al., 2014) and the impact of aerosols on climate (Andreae and Gelencsér, 2006).

To date, environmental chamber experiments have generated most of the laboratory SOA yield data used in atmospheric models, especially in simulations of polluted atmospheric conditions with elevated NOₓ concentrations. However, NOₓ-dependent chamber studies are complicated by the need to use multiple OH radical precursors such as hydrogen peroxide (H₂O₂) and nitrous acid (HONO) or methyl nitrite (CH₃ONO) to span the relevant range of NOₓ levels (typically, H₂O₂ for low-NOₓ conditions and HONO or CH₃ONO for high-NOₓ conditions) (Ng et al., 2007). Additionally, chambers have relatively low throughput and are limited to residence times of several hours due to chamber deflation and/or loss of particles and oxidized vapors to the chamber walls (Zhang et al., 2014). This restricts environmental chambers to simulating atmospheric aerosol particle lifetimes and SOA yields only up to 1 or 2 days, therefore limiting the study of the formation of highly oxygenated SOA characteristic of aged atmospheric organic aerosol particulate matter (PM) (Ng et al., 2010) unless very low VOC precursor concentrations are used (Shilling et al., 2009; Pfafffenberger et al., 2013).

Oxidation flow reactors have recently been developed to study SOA formation and evolution over timescales ranging from hours to multiple days of equivalent atmospheric OH exposure. In these reactors, O₃ is photolyzed at 254 nm to produce O(¹D) radicals, which react with water vapor to produce OH radicals. OH concentrations are typically 10⁸ cm⁻³ or greater. Under these conditions, atmospheric photochemical aging timescales up to ~10 days can be simulated at flow tube residence times of a few minutes or less. Recent experimental studies suggest that flow-reactor-generated SOA particles have compositions similar to SOA generated in smog chambers (Bruns et al., 2015; Lambe et al., 2015) and in the atmosphere (Tkacik et al., 2014; Ortega et al., 2016; Palm et al., 2016). Modeling studies suggest that flow reactors can simulate tropospheric oxidation reactions with minimal experimental artifacts (Li et al., 2015; Peng et al., 2015, 2016). A limitation of flow reactors is the need to use parts-per-million levels of O₃, hindering the possibility to efficiently simulate NOₓ-dependent SOA formation pathways. Simple addition of NO to flow reactors, while possible (Liu et al., 2015), cannot sustain NOₓ mixing ratios at levels that are sufficient to compete with hydroperoxy (HO₂) radicals as a sink for organic peroxy (RO₂) radicals due to fast conversion of NOₓ to nitric acid (HNO₃) via the reactions NO + O₃ → NO₂ + O₂ and NO₂ + OH → HNO₃. Here, we present a new method well suited to the characterization of NOₓ-dependent SOA formation pathways in oxidation flow reactors. By utilizing O(¹D) radicals that are generated from O₃ photolysis, we add N₂O to generate NO via the reaction O(¹D) + N₂O → 2NO with no additional method modifications. We validate the concept using high-resolution time-of-flight chemical ionization mass spectrometer measurements (HR-ToF-CIMS) to detect gas-phase oxidation products of isoprene and α-pinene that have been observed in NOₓ-influenced environments and laboratory chamber experiments.

2 Experimental

Experiments were conducted using an Aerodyne Potential Aerosol Mass (PAM) oxidation flow reactor, which is a horizontal 13.3 L aluminum cylindrical chamber (46 cm long × 22 cm ID) operated in continuous flow mode (Kang et al., 2007; Lambe et al., 2011a). The average residence time was 80 s. The relative humidity (RH) in the reactor was controlled in the range of 3–35% at 22 °C, corresponding to H₂O mixing ratios of approximately 0.07–1%. The irradiance in the reactor was measured using a photodiode (TOCON-C6, sglux GmbH). The gas-phase SOA precursors used in these studies include two biogenic compounds (isoprene, α-pinene) that were prepared in compressed gas cylinders and introduced to the reactor at controlled rates using a mass-flow controller. Mixing ratios of the gas-phase precursors entering the reactor were 36 ppb for isoprene (diluted from 1000 ppm in N₂, Matheson) and 15 ppb for α-pinene (diluted from 150 ppm in N₂, Matheson). These mixing ratios are a factor of 3 to 10 lower than mixing ratios that are typically required to induce homogenous nucleation of condensable oxidation products in related oxidation flow reactor.
studies (Lambe et al., 2011b). Minimizing precursor mixing ratios also decreases the rate of RO2 self-reactions relative to RO2 + HO2 and RO2 + NO reactions. This is a goal for most laboratory experiments that is not specific to the method proposed here. However, this goal takes on added importance when RO2 can be formed via OH, O3, and/or NO3 oxidation using this method as discussed in Sect. 2.1.

2.1 OH radical and NO3 generation

OH radicals were produced in the reactor via the reaction O(1D) + H2O → 2OH, with O(1D) radicals produced from the reaction O3 + hν → O3 + O(1D). O3 (∼ 1–5 ppm) was generated outside the flow reactor by O2 irradiation at 185 nm using a mercury fluorescent lamp (GPH212T5VH, Light Sources, Inc.). O(1D) was produced by photolysis of O3 at 254 nm inside the reactor using two or four mercury fluorescent lamps (GPH436T5L, Light Sources, Inc.). A fluorescent dimming ballast was used to regulate current applied to the lamps. To vary [OH] inside the reactor, I254 was varied by changing the dimming voltage applied to the ballast between 1.6 and 10 VDC. At these conditions, I254 ranged within approximately (0.042–2.1) × 1015 ph cm−2 s−1. The highest I254 value was calculated from the internal surface area of the reactor and the lamp output at maximum intensity (e.g., 10 VDC) specified by the manufacturer. Lower I254 values were calculated from the measured irradiance at lower dimming voltage relative to the measured irradiance and manufacturer-specified lamp output at 10 VDC.

NO and NO2 were produced via the reaction O(1D) + N2O → 2NO, followed by the reaction NO + O3 → NO2 + O2. N2O (99.5 %) was introduced from a compressed gas cylinder at flow rates ranging from 0 to 648 cm3 min−1, corresponding to mixing ratios of 0 to 5.6 % at the carrier gas flow rates that were used. Using N2O as the NO3 precursor has the following advantages over the simple addition of NO to the carrier gas. First, due to continuous production of O(1D) from O3 photolysis inside the reactor (along with minor consumption of N2O), the spatial distribution of NO and NO2 is more homogenous. Second, attainable steady-state mixing ratios of NO from O(1D) + N2O reactions (parts-per-billion levels) are orders of magnitude higher than simple NO injection (sub-parts-per-trillion levels) as inferred from photochemical model simulations described below in Sect. 2.3. Third, photolysis of N2O at 185 nm (if used) provides an additional source of O(1D) from the reaction N2O + hν → N2 + O(1D). We assume background [NO] < 0.05 ppb in the reactor based on separate [NO] measurements and calculate additional NO formed from O(1D) + N2O reactions using the model described in Sect. 2.3. Gradients in [O(1D)] due to its reaction with H2O and N2O may alter spatial distributions of O3, HOx, and NO3 in the reactor. To first order, gradients in [O(1D)] decrease both [HO2] and [NO] to a similar extent, keeping the relative rates of RO2 + HO2 and RO2 + NO3 termination pathways the same.

In most cases, oxidation of VOCs by O3 is slower than oxidation by OH radical, even with parts-per-million levels of O3 present (Peng et al., 2016). NO3 radicals, which are produced as a byproduct of NO2 + O3 or HNO3 + OH reactions, can potentially convolute interpretation of results if the relative oxidation rates of isoprene/α-pinene by OH and NO3 are comparable. For results presented in Sects. 3.3 and 3.4, calculated OH, O3, and NO3 exposures combined with published OH, O3, and NO3 rate constants (Atkinson, 1986, 1991; Grosjean and Grosjean, 1996) suggest that the relative contribution of NO3 to isoprene and α-pinene oxidation ranges from approximately 0 to 11 and 0 to 67 % as a function of [N2O]. Thus, reaction rates of α-pinene with OH, O3, and NO3 may be comparable under a subset of experimental conditions. Potential implications are discussed in more detail in Sects. 3.3.4 and 3.4.4.

2.2 NO3 and chemical ionization mass spectrometer (CIMS) measurements

In one set of experiments, [NO] and [NO3] were measured downstream of the reactor with a Thermo Scientific Model 42i chemiluminescent analyzer and an Aerodyne Cavity-Attenuated Phase Shift (CAPS) NO2 analyzer, which measures NO2 absorption at λ = 450 nm (Kebabian et al., 2008). During these experiments, the following operating conditions were used: I254 = 4 × 1015 ph cm2 s−1, [O3] = 1 ppm, [H2O] = 0.07 and 1 %, and [N2O] = 0 to 3 %. These conditions assess a subset of the attainable operating conditions for comparison with outputs of the photochemical model described in Sect. 2.3. The measured NO2 mixing ratio was decreased by 10 ppb due to absorption by 1 ppm O3 at 450 nm in the absence of NO2. The measured NO mixing ratio was scaled by a factor of 3.2 for depletion downstream of the reactor due to 1.2 s reaction time with 1 ppm O3 in the sample line, assuming kNO = 1.8 × 10−14 cm3 mole−1 s−1 and pseudo-first-order conditions (Atkinson et al., 2004). Additional NO depletion inside the chemiluminescent analyzer (∼ 47 % at 1 ppm O3) was accounted for in a separate experiment where known mixing ratios of NO (50 ppb) and O3 (0 to 6.9 ppm) were added at the inlet of the instrument (Fig. S1 in the Supplement). Because the combined NO depletion in the sample line and the chemiluminescent analyzer is significantly higher at higher [O3] (e.g., ∼ 90 % at [O3] = 2 ppm and ∼ 99.6 % at [O3] = 5 ppm), accurate experimental characterization of [NO] is more difficult above [O3] ∼ 1 ppm.

In another set of experiments, mass spectra of isoprene and α-pinene gas-phase oxidation products were obtained with an Aerodyne high-resolution time-of-flight mass spectrometer (Bertram et al., 2011) coupled to an atmospheric pressure interface with a nitrate ion chemical ionization source (NO3−-HR-ToF-CIMS, hereafter abbreviated as
“NO$_3^-$-CIMS”) (Eisele and Tanner, 1993; Ehn et al., 2012). Nitrate (NO$_3^-$) and its higher-order clusters (e.g., HNO$_3$NO$_3$) generated from X-ray ionization of HNO$_3$ were used as the reagent due to the selectivity to highly oxidized organic compounds, including species that contribute to SOA formation (Ehn et al., 2014; Krechmer et al., 2015). Isoprene and α-pinene oxidation products were detected as adducts with NO$_3^-$ or HNO$_3$NO$_3$. CIMS data were analyzed using the Tofware software package (ToFwerk AG, Aerodyne Research, Inc.) implemented in IGOR Pro 6 (WaveMetrics, Inc.). The output of the PAM oxidation flow reactor was sampled at 10.5 L min$^{-1}$ through a 2 ft length of 0.75 in. OD stainless-steel tubing inserted directly into the rear feedthrough plate of the reactor.

Ambient NO$_3^-$-CIMS measurements were conducted during the Southern Oxidant and Aerosol Study (SOAS) at the forest site in Centreville, AL, USA (1 June–15 July 2013). At this site, emissions were dominated by local biogenic volatile organic compounds (BVOCs) with occasional influence from nearby anthropogenic sources (Hansen et al., 2003). The mixing of biogenic and anthropogenic emissions at the forest site promotes the formation of organic nitrates via oxidation of BVOCs in the presence of NO$_3$ (Lee et al., 2016).

2.3 Photochemical modeling

We used a photochemical model (Li et al., 2015; Peng et al., 2015) implemented in MATLAB (Mathworks) to calculate concentrations of radical/oxidant species produced in the reactor. Model input parameters included pressure, temperature, [H$_2$O], [O$_3$], [N$_2$O], $l_{254}$, mean residence time, and the input mixing ratios of isoprene and α-pinene. Differential equations used to describe the radical/oxidant chemistry were integrated at 5 ms time steps. The following reactions and associated kinetic rate constants (Sander et al., 2000, 2006) were implemented to describe NO$_3$ chemistry in the reactor.

\[
\begin{align*}
N_2O + hv &\rightarrow N_2 + O(1D) \quad (R1) \\
N_2O + O(1D) &\rightarrow 2NO \quad (R2) \\
N_2O + O(1D) &\rightarrow N_2 + O_2 \quad (R3) \\
NO + OH + M &\rightarrow HONO + M \quad (R4) \\
NO + HO_2 &\rightarrow OH + NO_2 \quad (R5) \\
NO + O_3 &\rightarrow NO_2 + O_2 \quad (R6) \\
NO_2 + O &\rightarrow NO + O_2 \quad (R7) \\
NO_2 + O_3 &\rightarrow NO_3 + O_2 \quad (R8) \\
NO_3 + O &\rightarrow NO_2 + O_2 \quad (R9) \\
HONO + OH &\rightarrow H_2O + NO_2 \quad (R10) \\
\end{align*}
\]

\[
\begin{align*}
NO_2 + HO_2 + M &\rightarrow HNO_4 + M \quad (R14) \\
N_2O_3 + H_2O &\rightarrow 2HNO_3 \quad (R15) \\
HNO_3 + OH &\rightarrow H_2O + NO_3 \quad (R16) \\
NO_3 + NO_3 &\rightarrow 2NO_2 + O_2 \quad (R17) \\
\end{align*}
\]

The model also includes simplified RO$_2$ chemistry, which is incorporated using the reactions listed below (IUPAC, 2013). The addition of these reactions constrains the effects of added isoprene or α-pinene (species “X” below) on steady-state [OH], [HO$_2$], and [NO]. Second-generation organic radical products of initial organic radical reactions (“RPHO”, “RPO$_2$”, “RPO”) are not reacted further in the model.

\[
\begin{align*}
OH + X &\rightarrow RO_2 + H_2O \quad (R18) \\
RO_2 + NO &\rightarrow RO + NO_2 \quad (R19) \\
RO_2 + HO_2 &\rightarrow ROOH + O_2 \quad (R20) \\
ROOH + OH &\rightarrow RO_2 + H_2O \quad (R21) \\
ROOH + OH &\rightarrow RPHO + OH + H_2O \quad (R22) \\
RO$_2$ + OH &\rightarrow RPO$_2$ + H_2O \quad (R23) \\
RO$_2$ + RO$_2$ &\rightarrow ROOR \quad (R24) \\
RO + O$_2$ &\rightarrow RPO + HO$_2$ \quad (R25) \\
RO$_2$ + NO + M &\rightarrow RNO$_2$NO + M \quad (R26) \\
RO + NO + M &\rightarrow RNO + M \quad (R27) \\
RO + NO$_2$ + M &\rightarrow RNO$_2$O + M \quad (R28) \\
\end{align*}
\]

Calculated steady-state OH exposures (product of mean OH concentration and residence time) ranged from $6.3 \times 10^9$ to $1.4 \times 10^{12}$ molec cm$^{-3}$ s or approximately 1 h to 11 days of equivalent atmospheric exposure at [OH] = 1.5 × 10$^9$ cm$^{-3}$ (Mao et al., 2009). These steady-state OH exposure values incorporate OH suppression due to reaction with isoprene and α-pinene and assume zero HO$_2$ regeneration. Steady-state [NO] and [HO$_2$] ranged from 0 to 6.3 and 0.02 to 2.3 ppb, respectively, depending on [N$_2$O], [H$_2$O], [O$_3$], and $l_{254}$. We assumed ±25% uncertainty in the calculated OH exposure and ±60% uncertainty in other model outputs (Peng et al., 2015). For ratios of model outputs with independent ±60% uncertainties (e.g., NO: HO$_2$), propagated uncertainties of ±85% were assumed. Addition of N$_2$O at the highest mixing ratios that were used suppressed [OH] because N$_2$O competes with H$_2$O as a sink for O(1D). Potential consequences of OH suppression are discussed where applicable in Sects. 3.3 and 3.4.

3 Results and discussion

3.1 Comparison of measured and modeled [NO] and [NO$_2$] values following O(1D) + N$_2$O and NO + O$_3$ reactions

Figure 1 compares modeled and measured NO mixing ratios obtained following 80 s residence time in the reac-
tor at the operating conditions described in Sect. 2.2. The corresponding integrated OH exposures are approximately 2.6 × 10^{11} and 2.4 × 10^{12} molec cm^{-3} s, respectively, in the absence of added N_{2}O. Symbols are colored by [N_{2}O], which ranged from 0 to 3 %. Measured [NO] ranged from 0 to 10.4 ppb and increased with increasing [N_{2}O], as expected, at both [H_{2}O] = 0.07 and 1 %. The mean ratio of modeled to measured [NO] was 0.94 ± 0.19 at [H_{2}O] = 0.07 % and 3.85 ± 2.33 at [H_{2}O] = 1 %.

NO_{2}, which is formed by the NO + O_{3} reaction, is more straightforward to measure under these conditions because NO_{2} reacts approximately 500 times slower than NO with O_{3}. Thus, a comparison of modeled and measured [NO_{2}] provides additional method evaluation with less uncertainty than [NO] measurements. Figure 2 compares corresponding modeled and measured NO_{2} mixing ratios obtained during the same experiments described in Fig. 1. As expected, [NO_{2}] increased with increasing [N_{2}O] because of faster NO + O_{3} reaction rate from increasing [NO]. At [H_{2}O] = 0.07 %, measured [NO_{2}] ranged from 0 to 291 ppb, whereas at [H_{2}O] = 1 %, measured [NO_{2}] ranged from 0 to 59 ppb. [NO_{2}] was lower in the latter case because additional OH was formed from O^{(1)D} + H_{2}O reactions (Sect. 2.1), which increased the rate of the OH + NO_{2} reaction. The mean ratio of modeled to measured [NO_{2}] was 0.72 ± 0.39 at [H_{2}O] = 0.07 % and 1.05 ± 0.50 at [H_{2}O] = 1 %. These results, combined with results shown in Fig. 1, suggest that an uncharacterized H_{2}O- or HNO_{3}-related artifact negatively biased the measured [NO] values at [H_{2}O] = 1 % and that the photochemical model described in Sect. 2.3 may be used to evaluate a wider range of reactor operating conditions.

The model also constrains mixing ratios of radical species such as HO_{2} that are difficult to measure directly or require additional measurement techniques (Mauldin et al., 1999; Sanchez et al., 2016).

### 3.2 Optimal reactor operating conditions for O^{(1)D} + N_{2}O reactions

To investigate optimal operating conditions for NO_{x} generation, we implemented the model described in Sect. 2.3 over operating conditions I_{254} = 3.2 \times 10^{13} to 6.4 \times 10^{15} ph cm^{-2} s^{-1}, [O_{3}] = 0.5 to 50 ppm, and [H_{2}O] = 0.07 to 2.3 % at 22°C, respectively, as a function of [N_{2}O] = 0 to 5 %. These values span the nominal range of operating conditions that can be achieved with the PAM reactor. To facilitate independent evaluation of the effects of [O_{3}] and I_{254} on [NO], we restricted our analysis to conditions that use only 254 nm photolysis. Using both 185 and 254 nm photolysis provides additional sources of O^{(1)D} and OH from N_{2}O and H_{2}O photolysis at 185 nm, respectively, at the expense of independent control of [O_{3}] and I_{254}.

Figure 3 shows the modeled steady-state [NO] in the reactor as a function of [N_{2}O] = 0 to 5 %, assuming a mean residence time of 80 s, [H_{2}O] = 1 %, and [O_{3}] = 5 ppm. In addition, Figs. S1–S3 in the Supplement show modeled NO : HO_{2} and OH : NO_{3} ratios as a function of input [N_{2}O], with I_{254}, [O_{3}], and [H_{2}O] each varied individually, while other input conditions are fixed. The following observations that are obtained from Figs. 3 and S1–S3 were used to identify the optimal operating conditions:

1. At fixed [O_{3}], [H_{2}O], and [O^{(1)D}]), [N_{2}O] and [NO] increase with increasing I_{254} (Figs. 3 and S1).
2. At fixed I_{254}, [H_{2}O], and [N_{2}O], increasing O_{3} increases the production and loss rates of NO from O^{(1)D} + N_{2}O and NO + O_{3} reactions, respectively. The relative importance of NO + OH, NO + O_{3}, and NO + NO_{3} reactions, which depend on [N_{2}O] and [O_{3}], further influence [NO]:
   - At [N_{2}O] ~ 1 %, increasing [O_{3}] from 0.5 to 5 ppm increases [NO] because the reaction rate of NO + OH decreases relative to NO + O_{3} (Fig. S2a).
   - At [N_{2}O] > 1 %, increasing [O_{3}] from 5 to 50 ppm decreases [NO] because the reaction rate of NO + NO_{3} increases relative to NO + O_{3} (Fig. S2a).
3. At fixed I_{254}, [H_{2}O], and [N_{2}O], increasing [O_{3}] decreases [NO] : [HO_{2}] and [OH] : [NO_{3}] by increasing NO_{2} and NO_{3} formation from NO + O_{3} and NO_{2} + O_{3} reactions.
4. At fixed I_{254}, [O_{3}], and [N_{2}O], increasing [H_{2}O] increases [OH] : [NO_{3}] by increasing OH production from H_{2}O + O^{(1)D} reactions (Fig. S3).
The relative importance of these operating conditions is situationally dependent on the relative OH, O₃, and NO₃ rate constants of the target species and photochemical age. To demonstrate proof of principle, we present NO₃⁻-CIMS spectra of isoprene and α-pinene oxidation products in the following sections.

3.3 NO₃⁻-CIMS spectra of isoprene oxidation products

Figure 4 shows NO₃⁻-CIMS mass spectra of products generated from the oxidation of isoprene (C₅H₈) that cluster with NO₃⁻ ions to form NO₂⁻-species adducts. Ion signals are plotted as a function of mass-to-charge ratio (m/Q). NO₃⁻ adduct formation is a relatively low-energy process that does not result in fragmentation of the analyte (Eisele and Tanner, 1993; Kurtén et al., 2011). Thus, the measured ion signals are directly related to the chemical formulas of individual species that are generated in the reactor. Ion signals corresponding to isoprene oxidation products shown in Fig. 4 were colored based on classification in ion groups containing two–five carbon atoms with zero (C₃H₄,6,8O₄–7 and C₅H₆,8,10,12O₃–8), one (C₂–3H₃,5NO₅ and C₃H₇,9,11NO₆–11), and two (C₅H₁₀NO₈–10) nitrogen atoms, where we assumed that nitrogen atoms were associated with nitrate functional groups and not heterocyclic compounds. We also assumed that nitrate functional groups are formed from RO₂ + NO or RO₂ + NO₂ reactions (Sect. 2.1). To examine changes in relative contributions of C₄H₄,6,8O₄–7, C₅H₆,8,10,12O₃–8, C₆H₇,9,11NO₆–11, and C₅H₁₀NO₈–10 ions as a function of added NO₃⁻, we made two simplifying assumptions: (1) the NO₃⁻-CIMS had the same sensitivity to all species that were detected, and (2) HNO₃ generated in the reactor did not alter the relative selectivity of the CIMS to different classes of oxidation products, as has been observed in some cases (Hyttinen et al., 2015).

To generate spectra shown in Fig. 4, the reactor was operated at I₂₅₄ = 4.2 × 10¹³ and 2.1 × 10¹⁵ phosph cm⁻² s⁻¹, [H₂O] = 1 %, [N₂O] = 0 and 3 %. As shown in Figs. S4 and S5, corresponding OH exposures ranged within (5.6–6.3) × 10⁹ (Fig. 4a and c; calculated > 43 % of isoprene reacted) and within (0.43–1.4) × 10¹² molec cm⁻³ s⁻¹ (Fig. 4b and d; calculated ~ 100 % of isoprene reacted), respectively. At low OH exposure, the OH suppression at “high NO₃⁻” relative to “low NO₃⁻” was comparatively minor (11 %), whereas at high OH exposure, the OH suppression at high NO₃⁻ relative to low NO₃⁻ was larger (69 %). At the high-NO₂, OH exposure of 4.3 × 10¹¹ molec cm⁻³ s⁻¹, isoprene can react with OH up to 43 times in the reactor. This presumably exceeds the number of OH reactions (followed by RO₂ + NO reactions) that are necessary to fragment or condense oxidation products to the point where they are no longer detected with NO₃⁻-CIMS. Thus, it is unlikely that OH suppression at high OH and high NO₃⁻ significantly affected the NO₃⁻-CIMS spectra shown in Fig. 4. To aid interpretation of results shown in Fig. 4, Fig. 5 summarizes several known isoprene + OH reaction pathways that are terminated by reactions of RO₂ with HO₂, NO, or NO₂. As will be discussed in the following sections, these pathways yield multigenerational oxidation products with chemical formulas corresponding to the major ions that are plotted in Fig. 4.
Figure 4. NO$_3^-$-CIMS mass spectra of isoprene oxidation products generated in the PAM reactor at [H$_2$O] = 1% and [O$_3$] = 5 ppm, and mean residence time = 80 s: (a) $I_{254} = 4.2 \times 10^{13}$ ph cm$^{-2}$ s$^{-1}$, [NO$_2$] = 0%; (b) $I_{254} = 2.1 \times 10^{15}$ ph cm$^{-2}$ s$^{-1}$, [NO$_2$] = 0%; (c) $I_{254} = 4.2 \times 10^{13}$ ph cm$^{-2}$ s$^{-1}$, [NO$_2$] = 3.2%; and (d) $I_{254} = 2.1 \times 10^{15}$ ph cm$^{-2}$ s$^{-1}$, [NO$_2$] = 2.9%. NO$_3^-$-CIMS mass spectra of the same compounds detected at the SOAS ground site in Centreville, Alabama, USA, during “low-NO” and “high-NO” conditions (see text for additional details; C$_{5}$H$_{6}$O$_{5-7}$ ions removed from SOAS spectra due to larger contributions from α-pinen + OH oxidation products; Fig. 6). “C$_x$” or “O$_y$” indicates number of carbon or atoms in labeled ions (not including oxygen atoms in nitrate functional groups).

### 3.3.1 NO$_3^-$-CIMS spectral features observed at low-NO$_x$ conditions

C$_{4-5}$H$_{4-12}$O$_{3-8}$ ions comprised 93 and 97% of the signals at low and high OH exposure (Fig. 4a and c, respectively). The C$_{5}$H$_{7-11}$NO$_{6-11}$ signals that were observed here may be due to background NO$_x$ in the reactor (Sect. 2.1). The signal at $m/Q = 230$, C$_{5}$H$_{12}$O$_{6}$ (NO$_3^-$ omitted for brevity here and elsewhere), was the largest signal detected at both low and high OH exposures at low-NO$_x$ conditions. Figure 5 suggests this species is a second-generation oxidation product generated from two reactions with OH and two RO$_2$ + HO$_2$ termination reactions (Krechmer et al., 2015; St. Clair et al., 2016) and is typically associated with isoprene SOA formation and growth under low-NO$_x$ conditions (Liu et al., 2016). Signals in Fig. 4b and d are approximately 10 times higher than in Fig. 4a and c because additional OH exposure produces higher yields of multigeneration oxidation products that are detected with NO$_3^-$-CIMS.

Previously identified multigeneration isoprene oxidation products such as C$_{5}$H$_{10}$O$_5$, C$_{5}$H$_{12}$O$_5$, and C$_{5}$H$_{10}$O$_8$ (Suratt et al., 2006; Krechmer et al., 2015; St. Clair et al., 2016) were also detected at significant intensity under low-NO$_x$ conditions. These species are formed after two reactions with OH, one RO$_2$ + HO$_2$ termination reaction, and one RO$_2$ + RO$_2$ termination reaction (Fig. 5). When the OH exposure was increased from $6.3 \times 10^{9}$ to $1.4 \times 10^{12}$ molec cm$^{-3}$ s$^{-1}$, the signal at C$_{5}$H$_{12}$O$_6$ increased by a factor of 10, and the signal at $m/Q = 246$, C$_{5}$H$_{12}$O$_7$, increased by a factor of 5. At high OH exposure, C$_{5}$H$_{12}$O$_7$ was the second-largest peak in the spectrum. These highly oxygenated isoprene oxidation products are likely also important in SOA formation processes. C$_{5}$H$_{10}$O$_5$ is a proposed tri-hydroperoxy carbonyl product formed after one reaction with OH, two hydrogen shifts, and one RO$_2$ + HO$_2$ termination reaction as shown in Fig. 5 (Peeters et al., 2014).

We hypothesize that C$_{5}$H$_{10}$O$_7$, C$_{5}$H$_{12}$O$_7$, and C$_{5}$H$_{10}$O$_8$ are more prominent in our spectra than in other studies because NO$_3^-$ is more selective to highly oxidized species than other reagent ions (Suratt et al., 2006; Liu et al., 2016).

### 3.3.2 NO$_3^-$-CIMS spectral features observed at high-NO$_x$ conditions

Following addition of N$_2$O at $\sim 3%$ mixing ratio, the NO$_3^-$-CIMS spectra changed significantly at low and high OH exposures (Fig. 4b and d). The signals of C$_{4-5}$H$_{4-12}$O$_{1-8}$-NO$_x$ oxidation products decreased, although the C$_{4}$H$_{4-6}$O$_{4-7}$ : C$_{5}$H$_{6-10}$O$_{3-8}$ ratio increased, presumably due to decomposition of alkoxy (RO) radicals generated from C$_5$ RO$_2$ + NO reactions into C$_4$ products. The C$_{2-3}$H$_{3-5}$NO$_5$ (peroxyacetyl nitrate (PAN) and peroxypropionyl nitrate (PPN)), C$_{5}$H$_{7-9}$NO$_{6-11}$, and C$_{5}$H$_{10}$N$_2$O$_{8-10}$ signals increased. At low OH exposure, C$_{2-3}$H$_{3-5}$NO$_5$, C$_{5}$H$_{7-9}$NO$_{6-11}$, and C$_{5}$H$_{10}$N$_2$O$_{8-10}$ signals constituted 2, 38, and 7% of the NO$_3^-$-CIMS signals, respectively (Fig. 4c). The largest signal in this spectrum was $m/Q = 259$, C$_{5}$H$_{11}$NO$_7$. Fig. 5 suggests this compound is a second-generation oxidation product that is formed after two reactions with OH, one RO$_2$ + NO termination reaction, and one RO$_2$ + HO$_2$ termination reaction (Xiong et al., 2015). The signal observed at $m/Q = 288$, C$_{5}$H$_{10}$N$_2$O$_8$, is a second-
Figure 5. Simplified reaction scheme summarizing known isoprene + OH reaction pathways yielding multigeneration oxidation products. Four peroxy radical (RO₂) isomers are generated following initial OH addition to isoprene: 1,2-RO₂, 4,3-RO₂, 1,4-RO₂, and 4,1-RO₂. The 1,2-RO₂ and 4,3-RO₂ isomers follow the same reaction pathways, yielding chemical formulas with green text that were detected with NO₃⁻-CIMS. The 4,1-RO₂ isomer yields C₅H₁₀O₇, also detected with NO₃⁻-CIMS. Chemical formulas with red text may be generated through the methacrolein (MACR) channel but were not detected with NO₃⁻-CIMS.

generation oxidation product that is formed after two reactions with OH and two RO₂ + NO termination reactions (Fig. 5) (Xiong et al., 2015). Other ion signals associated with dinitrate species included \( m/Q = 304, C_4H_{10}N_2O_6 \), and \( m/Q = 320, C_5H_{10}N_2O_{10} \). Related signals were detected at \( m/Q = 351 \) and 367 (not shown), which we assume represent \((HNO₃NO₃⁻)C_5H_{10}N_2O_6\) and \((HNO₃NO₃⁻)C_5H_{10}N_2O_9\) because we are not aware of other feasible \((NO₃⁻)C_5\) adducts at these mass-to-charge ratios.

At high OH exposure, the same \( C_4H_{7,9,11}NO_6−11 \) and \( C_5H_{10}N_2O_{8−10} \) species observed at low OH exposure were detected, albeit at higher concentrations and at higher dinitrate: nitrate. This is presumably due to higher \( NO:HO₂ \) achieved at higher \( I_{254} \) and fixed \([N₂O] \) (Figs. 3, S2, S5–S6). \( C_2−3H_{3,5}NO_5, C_5H_{7,9,11}NO_6−11, \) and \( C_5H_{10}N_2O_{8−10} \) signals made up 0.3, 33, and 56 %, respectively, of the NO₃⁻-CIMS spectrum shown in Fig. 4d, where \( C_5H_{10}N_2O_5 \) was the largest signal that is detected.

To demonstrate our ability to mimic atmospheric NO₃⁻-dependent photochemistry, Fig. 4e and f show \( C_4H_{4,6,8}O_{4−7}, C_5H_{6,8,10,12}O_{3−8}, C_2−3H_{3,5}NO_5, C_5H_{7,9,11}NO_6−11, \) and \( C_5H_{10}N_2O_{8−10} \) ion signals detected in NO₃⁻-CIMS spectra at the SOAS ground site in Centre ville, Alabama, USA. The spectra shown were obtained on 25 June 2013 (07:30–11:00) and 4–5 July 2013 (12:00–00:00) which represented periods with sustained high and low NO mixing ratios of 0.53 ± 0.17 and 0.024 ± 0.025 ppb, respectively, measured at the site. Figure 4a, c, and e indicate that adding \( N₂O \) to the reactor increases the similarity between the composition of isoprene oxidation products generated at lower photochemical age in the reactor (Fig. 4a and c) and under low-NO ambient conditions (Fig. 4e). Likewise, Fig. 4b, d, and f indicate that adding \( N₂O \) to the reactor increases the similarity between the composition of isoprene oxidation products generated at higher photochemical age in the reactor (Fig. 4b and d) and at high-NO ambient conditions (Fig. 4f). \((HNO₃NO₃⁻)C_5H_{10}N_2O_{8−9}\) adducts were also observed in Fig. 4f (not shown).

3.3.3 Influence of acylperoxy nitrates from RO₂ + NO₂ reactions

Acylperoxy nitrates (APNs) may be generated from reactions of aldehydic, biogenic VOC oxidation products with OH followed by RO₂ + NO₂ termination reactions (e.g., LaFranchi et al., 2009). PAN \((C_2H₃NO₃)\) and PPN \((C_3H₅NO₃)\) are minor components (<2 %) of the spectra shown in Fig. 4e–f. A comparison of Fig. 4e and e suggests that yields of PAN and PPN are not enhanced in the reactor compared to atmospheric conditions. Additional APNs may be generated following the OH oxidation of methacrolein, a first-
generation isoprene oxidation product. Methacryloyl peroxynitrate (MPAN, C$_4$H$_5$NO$_3$) is a second-generation oxidation product formed after one methacrolein + OH reaction and one RO$_2$ + NO$_2$ termination reaction (Orlando et al. 1999). C$_4$-hydroxynitrate-PAN (C$_4$H$_8$N$_2$O$_9$) is a third-generation oxidation product formed through the methacrolein channel after three reactions with OH, two RO$_2$ + NO termination reactions, and one RO$_2$ + NO$_2$ termination reaction (Surrat et al., 2010). Neither C$_4$H$_7$NO$_3$ nor C$_4$H$_6$N$_2$O$_9$ was detected in the laboratory and ambient NO$_3^-$-CIMS spectra shown in Fig. 4c–f. Either these compounds were oxidized or thermally decomposed prior to detection, or their signals were below detection limit. C$_4$H$_7$NO$_3$, which is formed after one methacrolein + OH reaction and one RO$_2$ + NO termination reaction (Surrat et al., 2010), was detected (Fig. 5). Taken together, these observations suggest that yields of APNs were not significantly enhanced in the reactor compared to atmospheric conditions.

3.3.4 Influence of isoprene + NO$_3^-$ reactions

Based on the calculated isoprene + OH and isoprene + NO$_3^-$ reaction rates (Figs. S5–S6), we assume that isoprene + NO$_3^-$ reactions had a minor influence on the NO$_3^-$-CIMS spectra shown in Fig. 4c and d. This assumption is further supported by the similarity between laboratory and ambient NO$_3^-$-CIMS spectra, the latter of which was obtained during the daytime and thus with minimal NO$_3^-$ exposure (07:30–11:00 for the high-NO spectra shown in Fig. 4f). Specific operating conditions different than those used in this study could increase the relative influence of isoprene + NO$_3^-$ reactions. In this hypothetical situation, enhanced yields of C$_5$H$_7$NO$_3$, C$_5$H$_8$N$_2$O$_6$, and C$_5$H$_{10}$N$_2$O$_8$ might occur following two reactions with NO$_3^-$ (Rollins et al., 2009). In addition, C$_5$H$_{10}$N$_2$O$_9$ may be generated from one isoprene + NO$_3^-$ reaction followed by one RO$_2$ + HO$_2$ termination reaction (Schwantes et al., 2015). All four of these ions are detected in the spectra shown in Fig. 4, although C$_5$H$_8$N$_2$O$_8$ (not shown in Fig. 4) is present at 0.5 % of the intensity of C$_5$H$_{10}$N$_2$O$_8$. If C$_5$H$_8$N$_2$O$_8$:C$_5$H$_{10}$N$_2$O$_8$ is significantly different under NO$_3^-$-dominated conditions, this ratio could distinguish the relative rates of isoprene + OH and isoprene + NO$_3^-$ reactions. Otherwise, it is not clear that the expected product distributions are significantly different whether isoprene is oxidized by OH or NO$_3^-$ in the presence of NO$_3^-$.

3.4 NO$_3^-$-CIMS spectra of α-pinene oxidation products

Figure 6 shows NO$_3^-$-CIMS mass spectra of products generated from the oxidation of α-pinene (C$_{10}$H$_{16}$). Ion signals corresponding to α-pinene oxidation products were colored based on classification in C$_{4}$H$_{8}$O$_{5}$–7, C$_{6}$–O$_{4}$–12, and C$_{10}$H$_{14}$–16,18,19,20–14 ion groups containing zero nitrogen atoms; C$_{3}$–H$_{3}$NO$_{3}$, C$_{3}$H$_{7}$NO$_{6}$–11, C$_{6}$–19,20HO$_{3}$–18 ion groups containing one nitrogen atom; and a C$_{10}$H$_{16}$,18,20NO$_{6}$–13 ion group containing two nitrogen atoms. As was the case with isoprene oxidation products, we assumed nitrogen atoms present in α-pinene oxidation products were associated with nitrate functional groups formed from RO$_2$ + NO or RO$_2$ + NO$_2$ reactions. Additionally, we again assumed that the NO$_3^-$-CIMS had the same sensitivity to all species that were detected and that HNO$_3$ generated in the reactor did not

www.atmos-meas-tech.net/10/2283/2017/

alter the relative selectivity of the CIMS to different classes of oxidation products. Because the oxidation pathways leading to α-pinene-derived HOMs are significantly more complex than those leading to isoprene-derived HOMs, the analogous figure to Fig. 5 for α-pinene-derived HOMs is beyond the scope of this paper.

To generate the spectra shown in Fig. 6, the reactor was operated at $I_{254} = 1.8 \times 10^{15}$ ph cm$^{-2}$ s$^{-1}$, [H$_2$O] = 0.07 %, and [N$_2$O] = 0 and 3.2 %. In this experiment, lower [H$_2$O] was used to minimize [OH] and facilitate closer comparison with spectra from previous NO$_3^-$-CIMS studies of α-pinene + O$_3$ oxidation products generated at low-NO$_x$ conditions (Ehn et al., 2012, 2014). As shown in Fig. S7, corresponding OH and O$_3$ exposures ranged within (0.13–1.1) $\times$ 10$^{11}$ molec cm$^{-3}$ s and within (7.2–9.3) $\times$ 10$^{16}$ molec cm$^{-3}$ s for the low- and high-NO$_x$ conditions, respectively. To first order, at OH and O$_3$ exposures of 1.3 $\times$ 10$^{10}$ and 7.2 $\times$ 10$^{15}$ molec cm$^{-3}$ s that are attained at [N$_2$O] = 3.2 %, α-pinene should react once with each oxidant in the gas phase. Thus, at the highest [N$_2$O] used, yields of second-generation (or later) α-pinene + OH oxidation products detected with the NO$_3^-$-CIMS were minimized relative to α-pinene + O$_3$ first-generation oxidation products, as desired (Jokinen et al., 2015). However, a potential consequence of using O($^1$D) + N$_2$O reactions to study the NO$_x$ dependence of chemical systems similar to those examined by Ehn et al. (2012, 2014) is that RO$_2$ may be produced from α-pinene + NO$_3$ reactions in addition to α-pinene + O$_3$ or α-pinene + OH reactions (Sect. 2.1 and Fig. S7).

3.4.1 NO$_3^-$-CIMS mass spectral features observed at low-NO$_x$ conditions

C$_{5}$H$_{6}$,O$_5$–7, C$_{6}$–9H$_8$,10,12,14O$_6$–12, C$_{10}$H$_{14}$,16,18O$_5$–14, and C$_{19}$–20H$_{28}$,30,32O$_6$–18 ion groups comprised 5, 36, 46, and 4 % of the signal detected at low-NO$_x$ conditions (Fig. 6a), respectively, assuming equal CIMS sensitivity and transmission to all detected species. The C$_{10}$ monomers and C$_{19}$–20 dimers compounds that were observed are often associated with atmospheric new particle formation events (Ehn et al., 2014). The prominent C$_{10}$H$_{14}$,16O$_7$–9 signals detected at $m/Q = 308, 310, 324, 326, 340$, and 342 in our measurements were dominant signals in previous laboratory and field experiments influenced by the ozonolysis of α-pinene emissions (Ehn et al., 2010, 2012; Jokinen et al., 2015). Other signals that were observed correspond to C$_{5}$–9 species that were generated following carbon–carbon bond cleavage of the C$_{10}$ backbone (Ehn et al., 2012). The remaining 10 % of the signal was classified into C$_{2}$–3H$_3$,S$_2$NO$_5$, C$_{5}$H$_2$NO$_6$–11, C$_{6}$–9H$_9$,11,13,15NO$_5$–10, and C$_{10}$H$_{15}$,17NO$_4$–14 ion groups and may be due to background NO$_x$ in the reactor (Sect. 2.1).

3.4.2 NO$_3^-$-CIMS mass spectral features observed at high-NO$_x$ conditions

As was the case with NO$_3^-$-CIMS spectra of isoprene oxidation products, the addition of N$_2$O to the reactor significantly changed the mass spectrum of α-pinene oxidation products (Fig. 6b). At [N$_2$O] = 3.2 %, organic nitrates and dinitrates comprised 65 % of the total ion signal. We observed reduction in C$_{6}$–9H$_8$,10,12,14O$_6$–12, C$_{10}$H$_{14}$,16,18O$_5$–14, and C$_{19}$–20H$_{28}$,30,32O$_6$–18 signals, along with increases in C$_5$H$_6$,O$_5$–7, C$_{5}$H$_2$O$_6$–11, C$_{6}$–9H$_9$,11,13,15NO$_5$–10, C$_{10}$H$_{15}$,17NO$_4$–14, and C$_{10}$H$_{16}$,18N$_2$O$_6$–13 signals. The C$_{10}$ dinitrates may originate from two α-pinene + OH reactions followed by two RO$_2$ + NO reactions, but they may also include contributions from one α-pinene + NO$_3$ reaction followed by one RO$_2$ + NO reaction. The largest signal in Fig. 6b was observed at $m/Q = 240, C_6H_8O_7$. The largest organic nitrates signals in this spectrum were at $m/Q = 329, C_6H_13NO_8$, followed by C$_{10}$H$_{15}$NO$_3$ ($m/Q = 355$), C$_{10}$H$_{16}$N$_2$O$_6$ ($m/Q = 354$), and C$_{10}$H$_{15}$NO$_8$ ($m/Q = 339$).

Figure 6c shows C$_5$H$_6$O$_5$–7, C$_{6}$–9H$_8$,10,12,14O$_6$–12, C$_{10}$H$_{14}$,16,18O$_5$–14, C$_{19}$–20H$_{28}$,30,32O$_6$–18, C$_2$H$_3$NO$_5$–11, C$_{5}$H$_2$NO$_6$–11, C$_{6}$–9H$_9$,11,13,15NO$_5$–10, C$_{10}$H$_{15}$,17NO$_4$–14, and C$_{10}$H$_{16}$,18N$_2$O$_6$–13 signals detected with NO$_3^-$-CIMS at the Centreville site during the SOAS campaign. The spectra shown here were obtained during the sampling period shown in Fig. 4f and, given the large number of compounds, may include contributions from HOM precursors other than α-pinene. A comparison of Fig. 6a–c indicates that adding N$_2$O to the reactor increases the similarity between the composition of α-pinene oxidation products generated in the reactor and under high-NO ambient conditions, especially in regards to the enhanced C$_5$H$_6$O$_5$–7, C$_{6}$–9H$_8$,11,13,15NO$_5$–10, C$_{10}$H$_{15}$,17NO$_4$–14, and C$_{10}$H$_{16}$,18N$_2$O$_6$–13 signals.

3.4.3 Detection of acylperoxy nitrates from RO$_2$ + NO$_2$ reactions

Figure 6b and c indicate that PAN ($m/Q = 183, C_2H_3NO_5$) and PPN ($m/Q = 197, C_3H_4NO_5$) are formed at lower yields (< 0.4 %) than were observed with isoprene (Fig. 4c and d), suggesting that PAN and PPN formation from reaction of α-pinene-derived RO$_2$ with NO$_2$ is not enhanced in the reactor compared to atmospheric conditions. C$_{6}$H$_3$NO$_5$ and C$_{10}$H$_{15}$NO$_6$–8 are APNs generated following OH oxidation of pinonaldehyde, a major first-generation oxidation product of α-pinene, with termination by RO$_2$ + NO$_2$ reaction (Eddingsas et al., 2012). All four compounds are detected in the reactor and ambient NO$_3^-$-CIMS spectra shown in Fig. 6b and c, with C$_{10}$H$_{15}$NO$_6$–8 signals among the largest in the spectra. If these signals represent APNs, they appear to be important in both laboratory and atmospheric conditions.
3.4.4 Influence of α-pinene + NO3 reactions

Our calculations suggest that α-pinene + NO3 reactions compete with α-pinene + OH reactions at the experimental conditions used to generate the NO3−-CIMS spectrum shown in Fig. 6b (Fig. S7). If this were the case, enhanced yields of C10H12NO6 would be anticipated from α-pinene + NO3 reaction to generate pinonaldehyde, followed by pinonaldehyde + NO3 reaction and RO2 + NO3 termination (Perraud et al., 2010; Nah et al., 2016). Other minor α-pinene + NO3 products detected with CIMS include C10H13NO5, C9H13NO6, C10H16NO2O2, and C10H15NO9 (Nah et al., 2016). We hypothesize that, if α-pinene + NO3 reactions influence the spectrum shown in Fig. 6b, C10H15NO6: C10H13NO5 should be higher in Fig. 6b than in Fig. 6c. Instead, the C10H15NO6: C10H13NO5 ratio was 0.12 in the reactor and 0.28 at the Centreville site during a daytime period (07:30–11:00) with presumably negligible NO3 influence.

Dinitrates (C10H16,18N2O6–13) shown in Fig. 6b may originate from two α-pinene + OH reactions followed by two RO2 + NO terminations, or one α-pinene + NO3 reaction followed by one RO2 + NO termination. Given comparable calculated OH and NO3 reaction rates under these conditions (Fig. S7e), we hypothesize that the majority of dinitrate signals should originate from α-pinene + NO3 reactions if their yields are not oxidant-dependent. If this were the case, C10H16,18N2O6–13 : C10H15,17NO4–14 should be larger in Fig. 6b than in c. However, C10H16,18N2O6–13 : C10H15,17NO4–14 was 0.23 in the reactor and 0.61 at the Centreville site. Thus, while the calculated α-pinene + NO3 oxidation rate is significant (Fig. S7e), it is not clear that α-pinene + NO3 oxidation products significantly affect the spectrum shown in Fig. 6b. This may be due to significantly lower organic nitrate yields from α-pinene + NO3 than from α-pinene + OH reactions in the presence of NO (Fry et al., 2014; Rindelaub et al., 2015).

3.5 Transition from RO2+HO2- to RO2+NO-dominant regimes observed in isoprene and α-pinene oxidation products

Figures 7 and 8 show normalized signals of the representative groups of isoprene and α-pinene oxidation products as a function of increasing NO : HO2, which may be influenced by NO + HO2, NO + RO2, and HO2 + RO2 reactions in the reactor. For each group of compounds, signals obtained at a specific NO : HO2 were normalized to the maximum observed signal. NO : HO2 is correlated with the relative branching ratios of RO2 + HO2 and RO2 + NO reactions that govern the distribution of oxidation products observed in Figs. 4 and 6. As is evident from Figs. 7 and 8, different ion families were characterized by different trends as a function of NO : HO2. The normalized signals of C5 (isoprene), C6–10...
(α-pinene), and C_{19–20} (α-pinene) species decreased monotonically with increasing NO : HO_2. In Fig. 8, the abundance of C_{19–20} dimers decreased significantly faster than the C_{6–10} species. Because dimers are products of RO_2 + RO_2 self-reactions, their yield is quadratic with respect to [RO_2] and therefore was more affected by competing RO_2 + NO reactions than species formed from RO_2 + HO_2 reactions.

The normalized signals of C_5 (isoprene) and C_{10} (α-pinene) organic nitrates reached their maximum values at NO : HO_2 ≈ 0.5–2 prior to decreasing. Maximum signals of C_{6–9} organic nitrates (α-pinene) were obtained at NO : HO_2 = 2.2, and maximum signals of C_5 (isoprene) and C_{10} (α-pinene) dinitrates were obtained at NO : HO_2 = 2.1 and 2.2. The formation of dinitrates was favored when RO_2 + NO ≫ RO_2 + HO_2, as expected, and regardless of whether RO_2 was formed from oxidation of α-pinene by OH, O_3, or NO_3. We hypothesize that NO : HO_2 ≫ 1 favored RO_2 + NO → RO + NO_2 fragmentation reactions that led to formation of smaller, more volatile C_5H_{6–8}O_{5–7} and C_5H_2NO_6–11 α-pinene oxidation products (Atkinson, 2007; Chacon-Madrid and Donahue, 2011), whose signals continuously increased with increasing NO : HO_2 along with other products not detected with NO_3−CIMS. This pathway apparently competed with RO_2 + NO → RO_2NO reactions that led to formation of C_5 isoprene dinitrates, C_6–C_{10} α-pinene nitrates, and C_{10} α-pinene dinitrates.

Isoprene oxidation products such as C_5H_9NO_3 and C_5H_11NO_3 contain one peroxide and one nitrate functional group, and C_5H_3NO_5 contains two peroxides and one nitrate functional group. The formation of these species, as well as C_{6–10} α-pinene-derived organic nitrates, was favored at NO : HO_2 ≈ 0.5–2, where the relative rates of RO_2 + NO and RO_2 + HO_2 reactions were similar. This correlation suggests that the C_{6–10} α-pinene organic nitrates detected with NO_3−CIMS contained a combination of peroxide and nitrate functional groups, whereas C_5 (isoprene) and C_{10} (α-pinene) dinitrates contained fewer functional groups that were specifically formed from RO_2 + HO_2 reactions.

4 Atmospheric Implications

The use of O(1D) + N_2O reactions facilitates systematic control of NO : HO_2 over the range of “RO_2 + HO_2-dominant” to “RO_2 + NO-dominant” conditions. Further, this is accomplished with the use of a single OH radical precursor (O_3) that has previously hindered characterization of NO_3-dependent chemistry in oxidation flow reactors. Our results suggest that this method can be used to identify molecular tracers for processes influenced by RO_2 + NO and/or RO_2 + NO_2 reactions (Figs. 4 and 6). This method will be used in future work to investigate the influence of NO_3 on physicochemical properties of secondary organic aerosols such as hygroscopicity and refractive indices over an atmospherically relevant range of NO : HO_2. Care should be taken to use experimental conditions that minimize the relative contributions of unwanted NO_3-initiated oxidation chemistry – for example, [O_3] ≫ 5 ppm and [H_2O] ≪ 1 % (Figs. S2 and S3) – particularly when using species such as α-pinene that are highly reactive to NO_3. While potential formation of dinitrates from α-pinene + NO_3 reactions at high-NO conditions was not the primary goal of this experiment, we note that this chemical fingerprint has been observed in ambient measurements (Yan et al., 2016) and thus represents a potential application of O(1D) + N_2O reactions in future work. Additionally, studies that require multiple days of equivalent atmospheric OH oxidation at NO : HO_2 ≫ 1 should consider implementing 185 nm photolysis of H_2O and N_2O to provide additional sources of O(1D) and OH that may decrease OH suppression due to competing O(1D) + H_2O and O(1D) + N_2O reactions.

Data availability. Data presented in this manuscript are available upon request to the corresponding authors.

The Supplement related to this article is available online at https://doi.org/10.5194/amt-10-2283-2017-supplement.

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

Acknowledgements. This research was supported by the Atmospheric Chemistry Program of the US National Science Foundation under grants AGS-1536939, AGS-1537446, and AGS-1537099, and by the US Office of Science (BER), Department of Energy (Atmospheric Systems Research), under grants DE-SC0006980 and DE-SC0011935. Andrew Lambe thanks Gabriel Isaacman-VanWertz and Jesse Kroll (Massachusetts Institute of Technology) for helpful discussion, Zhe Peng and Jose Jimenez (University of Colorado – Boulder) for preliminary input on method operation, Karsten Baumann and Eric Edgerton (Atmospheric Research and Analysis, Inc.) for the use of ambient NO measurements during the SOAS campaign, Thomas Hanisco for handling editor responsibilities, and the anonymous reviewers for reviewing this manuscript.

Edited by: Thomas F. Hanisco
Reviewed by: two anonymous referees

References


