Technical note: Gas-phase nitrate radical generation via irradiation of aerated ceric ammonium nitrate mixtures

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Abstract.

We present a novel photolytic source of gas-phase NO_3 suitable for use in atmospheric chemistry studies that has several advantages over traditional sources that utilize $NO_2 + O_3$ reactions and/or thermal dissociation of dinitrogen pentoxide (N_2O_5). The method generates NO_3 via irradiation of aerated aqueous solutions of ceric ammonium nitrate ($(NH_4)_2Ce(NO_3)_6$,

- 5 "CAN") and nitric acid (HNO₃) or sodium nitrate (NaNO₃). We present experimental and model characterization of the NO₃ formation potential of irradiated CAN/HNO₃ and CAN/NaNO₃ mixtures containing [CAN] = 10^{-3} to 1.0 M, [HNO₃] = 1.0 to 6.0 M, [NaNO₃] = 1.0 to 4.8 M, photon fluxes (*I*) ranging from 6.9×10^{14} to 1.0×10^{16} photons cm⁻² s⁻¹, and irradiation wavelengths ranging from 254 to 421 nm. NO₃ mixing ratios ranging from parts per billion to parts per million by volume were achieved using this method. At the CAN solubility limit, maximum [NO₃] was achieved using [HNO₃] ≈ 3.0 to 6.0 M
- 10 and UVA radiation (λ_{max} = 369 nm) in CAN/HNO₃ mixtures or [NaNO₃] ≥ 1.0 M and UVC radiation (λ_{max} = 254 nm) in CAN/NaNO₃ mixtures. Other reactive nitrogen (NO₂, N₂O₄, N₂O₅, N₂O₆, HNO₂, HNO₃, HNO₄) and reactive oxygen (HO₂, H₂O₂) species obtained from the irradiation of ceric nitrate mixtures were measured using a NO_x analyzer and an io-dide adduct high-resolution time-of-flight chemical ionization mass spectrometer (HR-ToF-CIMS). To assess the applicability of the method for studies of NO₃-initiated oxidative aging processes, we generated and measured the chemical composition of
- 15 oxygenated volatile organic compounds and secondary organic aerosols from the β -pinene + NO₃ reaction using a Filter Inlet for Gases and Aerosols (FIGAERO) coupled to the HR-ToF-CIMS.

1 Introduction

The importance of NO₃ as a nighttime atmospheric oxidant is well established (Wayne et al., 1991; Brown and Stutz, 2012; Ng et al., 2017; Wang et al., 2023). NO₃ is generated via the reaction NO₂ + O₃ \rightarrow NO₃ + O₂, followed by achievement of

- 20 temperature-dependent equilibrium between NO₃, NO₂, and dinitrogen pentoxide (N₂O₅). N₂O₅ also hydrolyzes efficiently to HNO₃ on aqueous surfaces (Brown et al., 2004). Thus, any investigation of the influence of NO₃ chemistry in a specific source region necessarily must account for the local temperature, humidity, and particle surface area along with other factors. Despite these complications, for decades, laboratory studies investigating gas-phase NO₃ chemistry have utilized the same $NO_2 + O_3$ reactions and/or N₂O₅ thermal decomposition to produce NO₃ as occurs in the atmosphere, and accommodated the
- inherent limitations associated with N₂O₅; namely, that it must be stored under cold and dry conditions until use. Few viable alternative methods for the generation of gas-phase NO₃ have been identified. Reactions between fluorine atoms and nitric acid (F + HNO₃ \rightarrow HF + NO₃), or chlorine atoms and chlorine nitrate (Cl + ClNO₃ \rightarrow Cl₂ + NO₃) require handling and/or synthesizing hazardous halogen-containing compounds (Burrows et al., 1985; Bedjanian, 2019). F and Cl can also compete with NO₃ for the oxidation of target analytes, as can O₃ if its reaction with NO₂ is used as the NO₃ source.
- In the 1960s and 1970s, following earlier research into the properties of ceric solutions (Meyer and Jacoby, 1901; Wylie, 1951; Hinsvark and Stone, 1956; Blaustein and Gryder, 1957), Thomas Martin and coworkers discovered that irradiating solutions containing ceric ammonium nitrate (CAN, $(NH_4)_2Ce(NO_3)_6$) generates aqueous NO₃ (Henshall, 1963; Martin et al., 1963, 1964; Glass and Martin, 1970; Martin and Glass, 1970; Martin and Stevens, 1978). In $\geq 6M$ nitric acid (HNO₃), CAN is thought to dissociate primarily into NH⁺₄ cations and hexanitratocerate (Ce(NO₃)²⁻₆) anions (Henshall, 1963). The
- 35 $Ce(NO_3)_6^{2-}$ is subsequently reduced to $Ce(NO_3)_5^{2-}$ upon irradiation by ultraviolet light, and NO₃ is generated as a primary photolysis product. A similar process occurs in other solvents, although the ensuing ceric composition in solution is complex and influenced by several factors. For example, in glacial acetic acid (CH₃COOH), CAN dissociates into primarily Ce(NO₃)₄ (Henshall, 1963). Additionally, ceric ions containing complexed hydroxyl (OH) or H₂O, CH₃COOH, or acetonitrile (CH₃CN) molecules are formed in aqueous, acetic acid, or CH₃CN media, respectively (Henshall, 1963; Glebov et al., 2021). Higher
- 40 solution acidity and/or CAN concentration appears to promote the formation of $Ce(NO_3)_6^{2-}$ (Wylie, 1951) and ceric nitrate dimers (Blaustein and Gryder, 1957; Demars et al., 2015). The following generalized mechanism was proposed by Glass and Martin (1970) to describe ceric nitrate photochemistry:

$$Ce^{(IV)} + h\nu \rightarrow Ce^{(III)} + NO_3$$
 (R1)

$$Ce^{(III)} + NO_3 \rightarrow Ce^{(IV)}$$
 (R2)

$$NO_3 + NO_3 \rightarrow N_2O_6$$
 (R3)

$$N_2O_6 + 2Ce^{(IV)} \rightarrow 2NO_2 + O_2 + 2Ce^{(III)}$$
(R4)

$$NO_3 + NO_2 + H_2O \rightarrow 2HNO_3$$
 (R5)

where $Ce^{(IV)}$ represents ceric nitrates as diverse as $Ce(NO_3)_4$, $Ce(NO_3)_6^{2-}$, $(NO_3)_5CeOCe(NO_3)_5^{4-}$, and $(H_2O)_3(NO_3)_3CeOCe(NO_3)_3(H_2O)_3$ that are potentially formed in solution (Henshall, 1963; Blaustein and Gryder, 1957;

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50 Demars et al., 2015). Similarly, Ce^(III) represents cerous nitrates such as Ce(NO₃)₃ and Ce(NO₃)²⁻. The rate of Reaction R2 is [HNO₃]-dependent (Martin and Glass, 1970), and the dinitrogen hexaoxide (N₂O₆) intermediate was proposed on the basis of supporting observations without direct measurements (Glass and Martin, 1970).

CAN is used routinely as an oxidizing agent in organic synthesis due to its widespread availability and low cost, high oxidative potential, and low toxicity (Nair and Deepthi, 2007). However, its usage in atmospheric chemistry to date is limited

- to studies of NO₃-initiated oxidative aging processes in solution, e.g. Alexander (2004). Given the potential simplicity of irradiating $Ce^{(IV)}$ mixtures relative to synthesizing and storing N₂O₅ under cold and dry conditions or reacting NO₂ + O₃ under carefully controlled conditions, $Ce^{(IV)}$ irradiation could in principle enable more widespread studies of NO₃ oxidation chemistry, which is understudied compared to OH chemistry (Ng et al., 2017). Here, for the first time, we investigated the use of $Ce^{(IV)}$ irradiation as a source of gas-phase NO₃. First, we designed a photoreactor that generates gas-phase NO₃ from
- 60 irradiated CAN/HNO₃ and CAN/NaNO₃ mixtures. Second, we characterized NO₃ concentrations achieved over a range of reactor operating conditions and mixture composition. Third, we characterized gas-phase reactive nitrogen and reactive oxygen species generated following $Ce^{(IV)}$ irradiation. Fourth, we demonstrated application of the method to generate and characterize OVOCs and SOA from the β -pinene + NO₃ reaction.

2 Methods

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65 2.1 Photoreactor design and operation

Figure 1 shows a schematic of the experimental setup used in this study. A zero air carrier gas flow of 0.5 L min⁻¹ was bubbled through a gas dispersion line consisting of 6.35 mm OD x 4.8 mm ID FEP tubing into approximately 10 mL of aqueous CAN/HNO₃ or CAN/NaNO₃ mixtures placed at the bottom of a 12.7 mm OD x 11.1 cm ID FEP tube. The FEP tube was surrounded by low-pressure mercury fluorescent lamps installed vertically in a custom enclosure. These lamps had a 35.6 cm illuminated length. At these operating conditions, the calculated gas transit time in the illuminated portion of the reactor was approximately 3 s. After exiting the photoreactor, the carrier gas flow was passed through a filter holder (Savillex, 401-21-47-10-21-2) containing a 47 mm PTFE membrane filter (Pall Gelman, R2PJ047) to transmit NO₃ (Wagner et al., 2011) while removing stray droplets from the sample flow. At the end of each experiment, the lamps were turned off, the gas dispersion line was removed from the top of the reactor, and FEP tubing and filter holder were flushed with distilled H₂O to remove residual Ce^(III) precipitate. Initial studies were conducted using a Cavity Attenuated Phase Shift (CAPS) NO₂ monitor operating at $\lambda = 405$ nm (Kebabian et al., 2005) and a second retrofitted CAPS monitor operating at $\lambda = 630$ nm which established that NO₂ and NO₃ were produced from irradiated Ce^(IV). Subsequent studies described in the next section used a 2B Technologies Model 405 analyzer to measure NO and NO₂ (Birks et al., 2018).

Depending on the specific experiment, lamps with peak emission output centered at $\lambda = 254$, 313, 369, or 421 nm, respectively (GPH436TL/4P, Light Sources, Inc.; F436T5/NBUVB/4P-313, F436T5/BLC/4P-369, F436T5/SDI/4P-421, LCD Lighting, Inc.) were used. Emission spectra from the manufacturer are shown in Figure S1. A fluorescent dimming ballast (IZT-2S28-D, Advance Transformer Co.) was used to regulate current applied to the lamps. To quantify the photon flux I_{i} in

the photoreactor for studies that used $\lambda = 254$, 313, or 369 nm radiation, we measured the rate of externally added O₃ ($\lambda = 254$ nm) or NO₂ photolysis ($\lambda = 313$ or 369 nm) as a function of lamp voltage under dry conditions (RH < 5%). The photon flux

85 was not quantified in studies that used $\lambda = 421$ nm radiation. NO₂ photolysis measurements were conducted in the absence

of oxygen to avoid O₃ formation. *I*-values Photon flux values were then calculated using methods described in Lambe et al. (2019); maximum $I_{254} = 1.0 \times 10^{16}$ photons cm⁻² s⁻¹, $I_{313} = 6.0 \times 10^{15}$ photons cm⁻² s⁻¹, and $I_{369} = 7.0 \times 10^{15}$ photons cm⁻² s⁻¹ were obtained.

2.2 Characterization studies

- In one set of experiments, the 0.5 L min⁻¹ photoreactor effluent was mixed with a 6.5 L min⁻¹ zero air carrier gas and injected 90 into a dark Potential Aerosol Mass oxidation flow reactor (OFR; Aerodyne Research, Inc.), which is a horizontal 13 L Tefloncoated aluminum cylindrical chamber operated in continuous flow mode. Approximately 6.5 L min⁻¹ of sample flow was pulled from the reactor, resulting in a calculated mean residence time in the OFR ($\tau_{\rm OFR}$) of approximately 120 s. To constrain NO₃ mixing ratios, a mixture of 10 VOC tracers with NO₃ reaction rate coefficients (k_{NO_3}) ranging from 3.01×10^{-19} to 2.69×10^{-11} cm³ molecules⁻¹ s⁻¹ at T = 298 K (Table S1) was injected through a 10.2 cm length of 0.0152 cm ID Teflon 95 tubing at a liquid flow rate of 0.94 μ L hr⁻¹ using a syringe pump. The tracer mixture was then evaporated into a 1 L min⁻¹ zero air carrier gas prior to injection into the OFR. The total external NO_3 reactivity (NO_3R_{ext}), which is the summed product of each tracer mixing ratio and its $k_{\rm NO_3}$, was approximately 5 s⁻¹. VOCs with proton affinities greater than that of H₂O were chosen to enable their measurement with a Tofwerk/Aerodyne Vocus proton transfer-reaction time-of-flight mass spectrometer (hereafter referred to as "Vocus PTR") operated using H_3O^+ reagent ion chemistry (Krechmer et al., 2018) and ~ 8000 (Th/Th) 100 resolving power. NO_3 mixing ratios were calculated from the measured decrease in VOC mixing ratios using the Vocus PTR. Here, we assumed that the total concentration of reacted VOCs was equal to the concentration of NO₃ injected into the OFR; because . Because NO_3 may additionally react with organic peroxy radicals (RO_2) generated from VOC + NO_3 reactions as well as OVOCs, these calculated NO₃ concentrations represent lower limits. Modeling calculations suggest that the fractional
- 105 consumption of NO_3 by RO_2 ranged from <0.01 to 0.17 over the range of conditions that were studied (Fig. S2). A subset of OVOCs generated from VOC + NO_3 reactions that had proton affinities greater than that of H_2O were also detected with the Vocus PTR.

In a separate set of experiments, the photoreactor effluent was diluted into 4 Lmin^{-1} zero air carrier gas and sampled with an Aerodyne iodide-adduct high-resolution time-of-flight chemical ionization mass spectrometer (HR-ToF-CIMS; hereafter

- 110 referred to as "CIMS"; Bertram et al. (2011)) and the NO_x analyzer. The CIMS was operated at a ~ 4000 (Th/Th) resolving power. Iodide-adduct reagent ion chemistry was used due to its high sensitivity and selectivity towards nitrogen oxides and multifunctional organic nitrates (Lee et al., 2014). To demonstrate application of the method to study NO₃-initiated oxidative aging processes, the chemical composition of β -pinene + NO₃ gas-and condensed-phase oxidation products was measured with a Filter Inlet for Gases and Aerosols (FIGAERO) coupled to the CIMS (Lopez-Hilfiker et al., 2013). Gas sampling and
- 115 simultaneous particle collection was performed for 1 min intervals, followed by thermal desorption of the particle sample from a PTFE filter membrane (15 min ramp from room temperature to 200°C, 10 min holding time, 8 min cooldown to room temperature).

2.3 Photochemical model

To supplement our measurements, and to characterize aqueous phase concentrations of species produced in the photoreactor

- 120 that were not measured, we developed a photochemical box model that was implemented in the KinSim chemical kinetic solver (Peng and Jimenez, 2019). The KinSim mechanism shown in Table S2 contains 79 reactions to model concentrations of Ce^(IV), Ce^(III), NO, NO₂, NO₃, N₂O₃, N₂O₄, N₂O₅, HNO₂, HNO₃, HNO₄, H, O, OH, HO₂, and H₂O₂. We assumed that HNO₃ that was present in solution prior to irradiation completely dissociated into H⁺ and NO₃⁻. When possible, we used condensedphase rate coefficients in the mechanism. For reactions that we assumed occurred but did not have published condensed-phase
- rate coefficients (e.g. $NO_3 + OH \rightarrow NO_2 + HO_2$) we used published gas-phase rate coefficients instead with no modifications aside from unit conversion. Gas-phase wall loss rates of NO_x , NO_y , and HO_x species were not explicitly considered in the mechanism. UV/Vis extinction cross sections (σ_{ext}) of CAN/HNO₃ and CAN/NaNO₃ mixtures were separately obtained between $\lambda = 200$ and 600 nm using an Agilent Cary 5000 UV/Vis/NIR spectrophotometer. Because of the high absorptivity and concentrations of the mixtures, samples were prepared in a 0.01 mm short-path-length cuvette (20/C-Q-0.01, Starna) to
- 130 minimize saturation of the photodetector relative to a cuvette with a standard 10 mm path length. Even with the cuvette that was used, CAN dilution was necessary in some cases in order to obtain σ_{ext} without photodetector saturation at shorter wavelengths. Spectra were obtained as a function of [CAN] (0.047 to 0.526 M), [HNO₃] (0 to 6.0 M), and [NaNO₃] (0 to 4.0 M) to cover the approximate range of mixture compositions that were characterized in Section 2.2. The σ_{ext} -values of the mixtured were then calculated using the Beer-Lambert law and applied in the KinSim mechanism. Model outputs were obtained over a total 135 experimental time of 14400 s at 1 s intervals.

3 Results and Discussion

The maximum NO₃ quantum yield (φ_{NO₃}) of UVA-irradiated CAN/HNO₃ mixtures is obtained at 6.0 M HNO₃ (Martin and Stevens, 1978); thus, this mixture composition served as the basis from which additional characterization studies were conducted. We found that 0.5 M CAN was the approximate solubility limit in 6.0 M HNO₃ at 25°C. Because 1.1 M CAN
is the solubility limit in H₂O and CAN is almost nearily in HNO₃ (Martin and Glass, 1970), 0.7 M CAN is the estimated solubility limit in 6.0 M HNO₃ in the absence of changes in ceric nitrate composition in solution. Thus, the reduction in CAN solubility (0.7 M → 0.5 M) observed in our studies was presumably associated with significant conversion of CAN to dimeric ceric nitrates in 6.0 M HNO₃ (Blaustein and Gryder, 1957; Demars et al., 2015).

3.1 NO₃ characterization studies

Figure 2a shows time series of thiophene (C₄H₄S), 2,3-dihydrobenzofuran (C₈H₈O), cis-3-hexenyl acetate (C₈H₁₄O₂), isoprene (C₅H₈), dimethyl sulfide (C₂H₆S), 2,5-dimethylthiophene (C₆H₈S), α -pinene (C₁₀H₁₆), and guaiacol (C₇H₈O₂) concentrations following injection into the OFR and exposure to NO₃ generated in the photoreactor from irradiation of a mixture of 0.5 M CAN and 6.0 M HNO₃ at $I_{369} = 7 \times 10^{15}$ photons cm⁻² s⁻¹. Here, concentrations of each VOC were first normalized

to the acetonitrile concentration to correct for changes in the syringe pump output over time and then normalized to the VOC

- 150 concentration prior to NO₃ exposure. Aside from C_6H_8S , whose relative decay was less pronounced than expected (Table S1), and butanal (C_4H_8O , not shown), whose signal decreased by approximately 30% and did not recover for reasons that are unclear, the oxidative loss of each tracer increased with increasing k_{NO_3} . Maximum tracer consumption was observed at the beginning of the experiment due to maximum NO₃ production from $Ce^{(IV)}$ irradiation. As the experiment progressed and $Ce^{(IV)}$ was reduced to $Ce^{(III)}$, the NO₃ concentration and corresponding VOC oxidative loss decreased. Compared to the
- 155 other VOCs, the initial increase in $C_{10}H_{16}$ and $C_7H_8O_2$ concentrations over the first 2 hours was delayed because of their higher k_{NO_3} values that resulted in >95% consumption and lower sensitivity to changes in [NO₃] in the initial stage of the experiment. To confirm that VOC degradation shown in Fig. 2a was due to reaction with NO₃, Figure S3 shows the relative NO₃ rate coefficients obtained from the decay of C_4H_4S , C_8H_8O , and $C_8H_{14}O_2$ measured with the Vocus PTR. We measured relative rate coefficients of 3.59 between C_8H_8O and C_4H_4S and 6.92 between $C_8H_{14}O_2$ and C_4H_4S , which are in agree-
- ment with relative rate coefficient values of 3.44±1.20 and 7.68±2.84 calculated from their absolute NO₃ rate coefficients (Atkinson, 1991; D'Anna et al., 2001). Time series of ions corresponding to nitrothiophene (C₄H₃NO₂S), C₅H₇NO₄₋₆ and C₁₀H₁₅NO_{5,6} organic nitrates, and nitroguaiacol (C₇H₇NO₄), which are known NO₃ oxidation products of C₄H₄S, C₅H₈, C₁₀H₁₆, and C₇H₈O₂ (Atkinson et al., 1990; Jenkin et al., 2003; Saunders et al., 2003; Cabañas et al., 2005), along with C₈H_{5,7}NO₄₋₆ and C₈H₁₃NO₅₋₆ ions that may be associated with NO₃ oxidation products of C₈H₈O and C₈H₁₄O₂, respectively, were anticorrelated with those of their respective VOC precursors (Figure S4). Tracer decay experiments similar to the
- one shown in Figure S3 were used to obtain results that are discussed in more detail in Sections 3.2, 3.3, and 3.4.

3.2 Effect of irradiation wavelength

- Figure 3a shows normalized [NO₃] values obtained following irradiation of mixtures containing CAN and 6.0 M HNO₃ or 4.8 M NaNO₃ as a function of irradiation wavelength. In CAN/HNO₃ mixtures, [NO₃] was a factor of 2.4-3.5 higher
 following irradiation at λ = 369 compared to the other wavelengths. On the other hand, [NO₃] decreased with increasing irradiation wavelength following irradiation of CAN/NaNO₃ mixtures; at λ = 254 nm, [NO₃] was a factor of 3.2-42 times higher than at the other irradiation wavelengths that were used. These differences in [NO₃] were larger than the differences in calibrated *I*-values photon flux values at the maximum ouptut output of each lamp type (±40%; Sect. 2.1). Different Ce^(IV) in CAN/HNO₃ and CAN/NaNO₃ mixtures may have influenced these trends, as suggested by their UV/Vis spectra (Fig. 3b).
- 175 The σ_{ext} curves of CAN/HNO₃ mixtures were generally larger, broader, and red-shifted relative to those of CAN/NaNO₃ mixtures, with the extent of red-shifting increasing with increasing [HNO₃], possibly due to higher yields of Ce(NO₃)₆²⁻ and/or ceric nitrate dimers (Blaustein and Gryder, 1957; Henshall, 1963; Demars et al., 2015). For $\lambda > 250$ nm, CAN/HNO₃ mixtures had $\sigma_{\text{ext,max}}$ values between $\lambda = 306$ 311 nm, whereas CAN/NaNO₃ solutions had $\sigma_{\text{ext,max}}$ values at $\lambda = 296$ nm. However, if [NO₃] was simply proportional to σ_{ext} , irradiation of CAN/HNO₃ mixtures at $\lambda = 313$ nm should have produced
- 180 the highest [NO₃]; this was not the case. Instead, model calculations suggest that higher [NO₂] obtained from significantly faster photolysis of HNO₃ at $\lambda = 254$ and 313 nm relative to $\lambda > 350$ nm suppressed NO₃ downstream of the photoreactor when shorter irradiation wavelengths were used (Sander et al. (2011), Table S2). At a photon flux of 10¹⁶ photons cm⁻² s⁻¹,

model-calculated [NO₃] values were within \pm 13% of each other for irradiation wavelengths ranging from λ = 254 to 369 nm. However, higher [NO₂] values obtained following Ce^(IV) irradiation at λ = 254 and 313 nm suppressed NO₃ by >96%

185 relative to the $\lambda = 369$ nm case during 120 s of simulated NO₂ + NO₃ reactions in the OFR. Thus, although the measured NO₃ suppression at these other irradiation wavelengths was less substantial than the model output, the measurement and model trends, along with achievement of maximum [NO₃] following $\lambda = 254$ nm irradiation of CAN/NaNO₃ mixtures that had lower [HNO₃], qualitatively support this explanation for the wavelength-dependent NO₃ yields observed in CAN/HNO₃ mixtures.

3.3 Effect of mixture composition

- 190 To characterize the influence of individual reagents on NO₃ formation, tracer decay experiments similar to the measurements shown in Figure 2 were repeated as a function of [CAN], [HNO₃], and [NaNO₃]. Figure 4a shows [NO₃] obtained from irradiated 6.0 M HNO₃ solutions containing 0.001 to 0.5 M CAN ($I_{369} = 7 \times 10^{15}$ photons cm⁻² s⁻¹), and irradiated 1.0 M NaNO₃ solutions containing 0.5 to 1.0 M CAN ($I_{254} = 1 \times 10^{16}$ photons cm⁻² s⁻¹). Results were normalized to [NO₃] achieved with solutions containing 0.5 M CAN and 6.0 M HNO₃. Control experiments conducted with irradiated 6.0 M HNO₃
- 195 or 1.0 M NaNO₃ solutions at $I_{254} = 1 \times 10^{16}$ photons cm⁻² s⁻¹ in the absence of CAN suggest that a fraction of the NO₃ obtained in CAN mixtures was generated via the reactions HNO₃ + $h\nu \rightarrow$ OH + NO₂ and HNO₃ + OH \rightarrow NO₃ + H₂O. The remaining NO₃ was clearly obtained from CAN irradiation because [NO₃] increased with increasing [CAN], as expected from Reaction R1. Overall, [NO₃] increased by approximately a factor of 3 as [CAN] was increased from 0.001 to 0.5 M in 6.0 M HNO₃.
- Figure 4b shows [NO₃] obtained in irradiated solutions containing 0.5 M CAN as a function of [HNO₃] ranging from 1.0 to 6.0 M or [NaNO₃] ranging from 1.0 to 4.8 M at the same I₃₆₉ and I₂₅₄ values used to obtain results shown in Fig. 4a. Irradiated CAN solutions containing 3.0 M and 6.0 M HNO₃ generated the same [NO₃] concentrations within measurement uncertainties, presumably because the NO₃ quantum yield (φ_{NO₃}) ranged from 0.92-1.00 over this range of acidity (Martin and Stevens, 1978; Wine et al., 1988). [NO₃] decreased by a factor of 2 as [HNO₃] was decreased from 3.0 M to 1.0 M, consistent with a reduction in φ_{NO₃} from 0.92 to 0.46 (Martin and Stevens, 1978). On the other hand, in irradiated CAN/NaNO₃ mixtures with uncharacterized φ_{NO₃}, [NO₃] was constant within measurement uncertainties between 1.0 and 4.8 M NaNO₃.

Other mixture components that were tested or considered included substitution of CH_3CN in place of H_2O and HNO_3 , ammonium nitrate (NH_4NO_3) instead of $NaNO_3$, ceric potassium nitrate ($K_2Ce(NO_3)_6$) instead of CAN, and addition of sodium persulfate ($Na_2S_2O_8$) to generate additional NO_3 via $S_2O_8^{2-} + h\nu \rightarrow 2SO_4^-$ followed by $SO_4^- + NO_3^- \rightarrow NO_3 +$

SO₄²⁻ (Gaillard de Sémainville et al., 2007). CAN/CH₃CN mixtures are commonly used in organic synthesis applications, perhaps even more so than CAN/HNO₃ (Baciocchi et al., 1988; Choidini et al., 1993; Alexander, 2004). In limited testing, CAN/CH₃CN appeared to generate significantly less NO₃ than CAN/HNO₃ or CAN/NaNO₃, possibly due to lower φ_{NO₃} of irradiated Ce^(IV)-CH₃CN complexes (Glebov et al., 2021) and/or suppression of NO₃ due to its reaction with CH₃CN in solution. K₂Ce(NO₃)₆ is less widely available and less water-soluble than CAN and so was not considered further. Irradiation of CAN/NH₄NO₃ and CAN/NaNO₃ mixtures generated similar [NO₃], but we prefer NaNO₃ due to its lower volatilty.

Finally, ternary mixtures containing 0.5 M CAN + 2.0 M NaNO₃ + 0.5 M Na₂S₂O₈ irradiated at λ = 254 nm generated negligible additional NO₃ compared to binary CAN/NaNO₃ mixtures.

3.4 Effect of photon flux

Figure 5 shows normalized [NO₃] values obtained from UVA-light-irradiated irradiated mixtures of 0.5 M CAN & 6.0 M

HNO₃ and UVC-light-irradiated mixtures of (λ = 369 nm) and 0.5 M CAN & 1.0 M NaNO₃ (λ = 254 nm) as a function of photon flux ranging from 6.9×10¹⁴ to 7.5×10¹⁵ photons cm⁻² s⁻¹. Results for both CAN/HNO₃ and CAN/NaNO₃ mixtures were normalized to [NO₃] achieved with 0.5 M CAN, 6.0 M HNO₃ and I₃₆₉ = 6.8×10¹⁵ photons cm⁻² s⁻¹. Symbols are colored by the NO₃ lifetime (τ_{NO₃}), defined here as the time it took for [NO₃] to experience one e-fold decay relative to the maximum [NO₃] that was measured. Figure 5 shows that [NO₃] increased with increasing photon flux, consistent with the fact that it is a primary photolysis product, along with a concurrent decrease in τ_{NO₃} due to faster reduction of Ce^(IV) to Ce^(III). For the CAN/HNO₃ system, [NO₃] increased by a factor of 1.5 as *I*₃₆₉ was increased from 6.9×10¹⁴ to 6.8×10¹⁵ photons cm⁻² s⁻¹, in agreement with the model-calculated increase in [NO₃] within measurement uncertainty. τ_{NO₃} decreased from 9 to 5 hr. For the CAN/NaNO₃ system, [NO₃] increased by a factor of 1.9 as *I*₂₅₄ was increased from 1.0×10¹⁵ to 7.5×10¹⁵

photons cm⁻² s⁻¹, and τ_{NO_3} decreased from 10 to 3 hr.

- To examine concentrations of NO₃ and a subset of additional gas-phase photolysis products obtained over a wider range of conditions, Figure 6 plots model-calculated [NO₃], NO₂:NO₃, HO₂:NO₃, and N₂O₅:NO₃ values as a function of photon flux ranging from 1×10^{14} to 1×10^{17} photons cm⁻² s⁻¹ following $\lambda = 254$, 313, 369 and 421 nm irradiation of a mixture of 0.5 M CAN and 6.0 M HNO₃. Figure 6a also plots the measured [NO₃] obtained from irradiation of a mixture of 0.5 M CAN and 6.0 M HNO₃ at $I_{369} = 7 \times 10^{15}$ photons cm⁻² s⁻¹ (Fig. 2) after correcting for dilution between the photoreactor and
- 235 the OFR (Sect. 2.2) and application of a NO₃ wall loss rate coefficient of 0.2 s⁻¹ within the photoreactor (Dubé et al., 2006) . At this photon flux value, the model-calculated [NO₃] = 1.4 ppmv agrees with [NO₃] = 1.7 ± 0.6 ppmv obtained from measurements. When considering only the primary photochemical process (Reactions R1-R5), maximum [NO₃] values within ± 10% of each other were achieved at photon fluxes ranging from 5×10^{15} (λ = 313 nm) to 4×10^{16} photons cm⁻² s⁻¹ (λ = 421 nm). [NO₃] values decreased at higher *I*-values photon flux values due to conversion of NO₃ to NO₂ via photolysis. As
- shown in Fig. 6b, significant additional NO₂ production was obtained via HNO₃ photolysis at shorter irradiation wavelengths above $I \approx 10^{15}$ photons cm⁻² s⁻¹, resulting in NO₂:NO₃ > 10 (λ = 254 nm) and 1 (λ = 313 nm). Given additional reaction time downstream of the photoreactor, high NO₂ may suppress NO₃ (Sect. 3.2) and increase N₂O₅:NO₃ beyond the range of values shown in Fig. 6c. We also calculated OH:NO₃ and HO₂:NO₃ following irradiation of CAN/HNO₃ mixtures over the range of conditions shown in Figure 6. Aqueous OH:NO₃ \approx 0.1 and did not change significantly as a function of photon flux
- or irradiation wavelength, and aqueous HO₂:NO₃ values ranged from 0.05 ($\lambda = 254$ nm) to 0.25 ($\lambda \ge 369$ nm). While OH influenced aqueous-phase chemistry inside the photoreactor via formation of reactive oxygen species (Sect. 3.5), OH probably did not influence downstream gas-phase chemistry due to significant wall losses inside the photoreactor: assuming a lower-limit OH wall loss rate coefficient of 5 s⁻¹ (Schwab et al., 1989), the estimated OH penetration efficiency through the reactor was

less than 10^{-6} . Similarly, in studies involving the generation of via VOC + reactions, is unlikely to significantly influence fate

250 because + reactions are several times slower than those of + reactions (Orlando and Tyndall, 2012).

3.5 Characterization of reactive nitrogen and reactive oxygen photolysis products

Figure S5-7 shows time series of ,,,,,,, and obtained with the CIMS reactive nitrogen and reactive oxygen species detected following irradiation of a the same mixture of 0.5 M CAN and 1.0 M NaNO₃ (I₂₅₄ ≈ 10¹⁶ photons cm⁻² s⁻¹). Signals of , and decreased following irradiation of the CAN/mixture, whereas , , , , and signals increased. One potential source of is +
reactions in the CIMS ion-molecule reactor (IMR); if this reaction was the sole source of here, we estimate an upper limit mixing ratio of approximately 15 ppbv present in the IMR (Dörich et al., 2021). is generated following the reaction of and/or with (Abida and Osthoff, 2011), and is generated from the reaction of and/or with multiple nitrogen oxides, including , , , and (Huey et al., 1995; Veres et al., 2015; Dörich et al., 2021), all of which are generated following irradiation as discussed later

in this section. Figure 7 shows time series of reactive nitrogen and reactive oxygen species detected following irradiation of

- 260 the same mixture of 0.5 M CAN and 1.0 M, shown here because the signal-to-noise in CIMS measurements of irradiated CAN/NaNO₃ mixtures was generally better than in measurements of irradiated CAN/HNO₃ mixtures due to reagent ion depletion by HNO₃. A time series of [NO₃] obtained separately from VOC tracer decay measurements under similar irradiation conditions is also shown. The NO₂ mixing ratio reached a maximum value and NO₃ mixing ratios reached maximum values of 26 and 58 ppbv shortly after the lights were turned on (Fig. 7a), suggesting an initial NO₂:NO₃ \approx 0.37 0.45 (Fig. 4)that was
- 265 similar to modeled := 0.33 obtained from irradiated CAN/(Fig. 6)... Multiple reactions may generate NO₂, including Reaction R3, HNO₃ and/or NO₃ photolysis, and other reactions listed in Table S2. While NO₂ and/or HNO₂ photolysis generated NO, its concentration was negligible in these experiments.

Figure 7b shows time series of $IN_2O_5^-$ and $IN_2O_6^-$ signals measured with the CIMS. Figure S5 additionally shows a time series of , and Figures S6, S7, and S8 shows high-resolution CIMS spectra at m/Q = 235, 251, and 267. $IN_2O_5^-$ was formed

- 270 from $NO_2 + NO_3 \rightarrow N_2O_5$ reactions in the photoreactor and $N_2O_5 + I^- \rightarrow IN_2O_5^-$ reactions in the CIMS IMR. As expected, $IN_2O_5^-$ followed a similar profile as NO_2 and NO_3 . Given : $\approx 10^{-3}$ coupled with similar and temporal profiles (Fig. S5), we hypothesize that + reactions in the IMR were the primary source of . $IN_2O_6^-$ was either generated from $NO_3 + NO_3 \rightarrow N_2O_6$ reactions in the photoreactor (Glass and Martin, 1970) followed by $N_2O_6 + I^- \rightarrow IN_2O_6^-$ reactions in the IMR, or from the following series of reactions in the IMR: $HNO_3 + IO^- \rightarrow NO_3^- + HOI$, $HOI + NO_3^- \rightarrow INO_3 + OH^-$, and $INO_3 + NO_3^-$
- 275 \rightarrow IN₂O₆⁻ (Ganske et al., 2019). If is related to , its signal increased faster than because is a primary photolysis product, then decreased faster than because the production rate decreased quadratically as a function of decreasing , whereas the production rate remained constant following processes that converted to . Additionally, because the aqueous phase +reaction rate is approximately 2000 times slower than that of + (Martin and Stevens, 1978; Katsumura et al., 1991), even a small amount of would favor the formation of compared to . To further explore the plausibility of N₂O₆ formation in this system, we conducted
- a theoretical investigation of the gas-phase $NO_3 + NO_3 \rightarrow N_2O_6$ reaction at T = 298 K and p = 1 atm. Quantum chemical calculations were performed using the Q-Chem 5.2 software package (Epifanovsky et al., 2021), and molecular geometries were obtained using the B3LYP density functional (Becke, 1993) and the 6-31G* basis set (Hariharan and Pople, 1973). All

stationary points were refined by single point calculations applying the B3LYP density functional and the cc-pVTZ basis set (Dunning, 1989) as well as CCSD(T) (Jeziorski and Monkhorst, 1981) and the cc-pVTZ basis set. For $+ \rightarrow$, the calculated

- 285 enthalpy of reaction (ΔH_{rxn}) was -35.8 kcal mol⁻¹ using the CCSD(T) method, and -21.9 kcal mol⁻¹ using the B3LYP method. By comparison, we calculated ΔH_{rxn} values of -26.5 (CCSD(T)) and -18.1 (B3LYP) kcal mol⁻¹ for the + \rightarrow reaction; the corresponding energy change (ΔE_{rxn}) values agreed within 5% of previously obtained experimental and computational ΔE_{rxn} values for this reaction (Jitariu and Hirst, 2000; Glendening and Halpern, 2007). Thus, regardless of the quantum chemical method that was used, + \rightarrow appears to be an exothermic reaction and found that this reaction is exothermic, even more so than
- NO₃ + NO₃ → N₂O₅. While the reverse reaction → 2 is possible (although endothermic, as is → +) our analysis suggests that the thermodynamically favored reaction pathway is → +, which had ΔH_{rxn} values ranging from -7.02 (CCSD(T)) to -6.15 (B3LYP) keal mol⁻¹. By contrast, the reaction → 2 + had ΔH_{rxn} = 5.28 (CCSD(T)) 5.58 (B3LYP) keal mol⁻¹; however, because → 2 is fast (Poskrebyshev et al., 2001; Atkinson et al., 2004), the overall reaction → 2 + is the favored removal pathway in the gas phase, and in solution may occur in addition to or instead of Reaction R4. Additional details
 regarding this analysis are provided in Sect. S1.
- Figure 7c shows time series of IHNO₂⁻, HNO₂NO₃⁻, IHNO₄⁻, and HNO₄NO₃⁻. These ions are associated with nitrous acid (HNO₂) and peroxynitric acid (HNO₄) respectively (Veres et al., 2015). Because rapid formation of HNO₂₋₄NO₃⁻ ions was observed following Ce^(IV) irradiation, and because IO_x⁻ signals were relatively low (Sect. S2.1), we hypothesize that I⁻ + NO₃ eharge transfer and/or I⁻ + HNO₃ reactions were the main source of NO₃⁻ (Lee et al., 2014) (Lee et al., 2014; Dörich et al., 2021)
 and that subsequent competitive NO₃⁻ + HNO₂₋₄ and I⁻ + HNO₂₋₄ reactions in the IMR generated both IHNO₂₋₄ and HNO₂₋₄NO₃⁻. HNO₄ was generated following the reactions HNO₃ + hv → OH + NO₂, OH + NO₃ → HO₂ + NO₂, and HO₂ + NO₂ → HNO₄. This hypothesis is supported by the similarity between NO₂ and IHNO₄⁻ time series coupled with the relatively constant concentrations of HO₂ generated via OH + OH → H₂O₂ and OH + H₂O₂ → HO₂ + H₂O reactions. H₂O₂, detected as IH₂O₂⁻, also behaved similarly as IHO₂⁻ (Figure 7d). HNO₂ had a different temporal profile than the other nitro-gen oxides: IHNO₂⁻ increased throughout the experiment, and HNO₂NO₃⁻ increased and then decreased. We hypothesize that NO₂+NO₂→N₂O₄ and N₂O₄+H₂O→HNO₂+HNO₃ reactions were the main source of HNO₂ (Sect. S2.2), was not detected
- with the CIMS following irradiation of aqueous, presumably because its hydrolysis rate was too fast (Park and Lee, 1988). In an attempt to decrease the hydrolysis rate, separate experiments were conducted in which the effluent of 40 g of irradiated solid CAN was sampled with the CIMS. At the sample sizes that were used, the solid CAN contained enough solvated and/or
- 310 that its irradiation provided sufficient production of nitrogen oxides for CIMS detection. As shown in Figures S9, S10 and S11, CIMS , , , and signals were significantly higher following irradiation at $\lambda = 254$ nm than at the other wavelengths, and Fig. S12 confirms that was the dominant ion signal at m/Q = 219. Taken together, these observations support our hypothesis that was generated following fast hydrolysis in aqueous solution.
- To compare measurements of reactive nitrogen and reactive oxygen species obtained from irradiated CAN/and CAN/mixtures, 315 Figure S13 shows time series of the same ions plotted in Figure 7 following irradiation of a solution containing 0.5 M CAN and 3.0 M HNO₃ ($I_{369} \approx 7 \times 10^{15}$ photons cm⁻² s⁻¹). Here, 3.0 M HNO₃ was used because 6.0 M HNO₃ depleted the CIMS reagent ion too much (IHNO₃⁻:I⁻ \approx 15) to achieve signal-to-noise that was sufficient for comparison to CAN/NaNO₃ mix-

tures (IHNO₃⁻:I⁻ \approx 3). The same gas-phase nitrogen oxides and reactive oxygen species were observed in this reaction system as with the irradiated CAN/NaNO₃ mixture. The relative yields of each compound plotted in Figures 7 and S13 were within a

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factor of 3 of each other, although signals of nitrogen oxides and reactive oxygen species obtained from irradiated CAN/HNO₃ mixtures decreased at a slower rate than the same compounds obtained from irradiated CAN/NaNO₃ mixtures. These trends may be due to different Ce^(IV) composition (Fig. 3 and Sect. 3.2) and/or enhanced rate of Ce^(III) + NO₃ \rightarrow Ce^(IV) reactions in HNO₃ relative to NaNO₃ (Reaction R2).

3.6 OVOC/SOA generation from β -pinene + NO₃

- To demonstrate proof of principle for NO₃-initiated oxidative aging studies, we generated NO₃ via irradiation of a mixture of 0.5 M CAN and 3.0 M HNO₃ (I₃₆₉ = 7×10¹⁵ photons cm⁻² s⁻¹), reacted it with β-pinene in a dark OFR, and obtained FIGAERO-CIMS spectra of gas- and condensed-phase β-pinene + NO₃ oxidation products (Sect. 2.2). Figure 8a shows a spectrum of gas-phase β-pinene/NO₃ oxidation products detected between m/Q = 320 and 420, where the majority of the signal was observed; signals shown are unmodified (M+I)⁻ formulas. The largest ion detected was at m/Q = 356 (IC₁₀H₁₅NO₅⁻), which represents a major first-generation dicarbonyl nitrate oxidation product with a relative abundance of 0.31 and a calculated saturation vapor pressure of 2×10⁻⁷ atm (C* = 1900 µg m⁻³; Claflin (2018)). Other ions corresponding to first-generation hydroxycarbonyl nitrate (IC₁₀H₁₇NO₆⁻, C* = 4.7 µg m⁻³), and hydroxycarbonyl nitrate acid (IC₁₀H₁₇NO₇⁻, C* = 0.29 µg m⁻³) products were detected in addition to IC₉H₁₃NO₅⁻ and a suite of additional previously characterized C₈ and C₉ organic nitrates (Nah
- et al., 2016; Takeuchi and Ng, 2019; Shen et al., 2021). The $IC_{10}H_{16}N_2O_7^-$ dinitrate was obtained following reaction of the β -nitrooxyperoxy radical with NO or hydroxy dinitrate, which was also previously observed in FIGAERO-CIMS spectra of α -pinene/NO₃ (Nah et al., 2016; Bates et al., 2022)SOA (Nah et al., 2016), was generated via an unknown reaction pathway. Because model-calculated NO:NO₃ was on the order of 10⁻⁵ under these conditions, its formation from the RO₂ + NO₃ reaction seems more likely (Orlando and Tyndall, 2012). Overall, the high molar yield and vapor pressure of $C_{10}H_{15}NO_5$ (Claffin,
- 2018) are consistent with it having the highest relative abundance in the gas phase (Fig. 8a), whereas the other $C_{10} \beta$ -pinene oxidation products were semivolatile under our experimental conditions.

Figure 9a shows a spectrum of condensed-phase β -pinene/NO₃ oxidation products obtained with the FIGAERO-CIMS; signals were averaged over the entire thermal desorption cycle and are plotted on logarithmic scale and represent unmodified (M+I)⁻ formulas. To aid interpretation of the major features of the spectrum, bands of ion signals corresponding to

- 345 $IC_{10}H_{15}NO_x^-$, $IC_{20}H_{32}N_2O_x^-$, and $IC_{30}H_{47}N_3O_x^-$ oxidation products were highlighted and colored by the number of oxygen atoms in their chemical formulas. Here, the largest ion detected was at m/Q = 372 ($IC_{10}H_{15}NO_6^-$), which is the condensedphase component of the same tricarbonyl nitrate detected in the gas-phase (Fig. 8a). $IC_{10}H_{15}NO_5^-$ and $IC_{10}H_{15}NO_{7-9}^-$ signals were also detected. The second largest ion signal was measured at m/Q = 571 ($IC_{20}H_{32}N_2O_9^-$), an acetal dimer obtained from the condensed-phase reaction of two $C_{10}H_{17}NO_5$ monomers followed by H_2O elimination (Claffin and Ziemann,
- 2018). Similar accretion reactions between other C_{10} organic nitrates yielded $IC_{20}H_{32}N_2O_8^-$ and $IC_{20}H_{32}N_2O_{10-13}^-$ signals. Likewise, reactions between C_{10} monomers and C_{20} dimers generated C_{30} trimers detected between m/Q = 768 864

 $(IC_{30}H_{47}N_3O_{12-18}^-)$. The largest trimer-related ion, $IC_{30}H_{47}N_3O_{12}^-$, was generated from $C_{10}H_{17}NO_4 + C_{20}H_{32}NO_9 - H_2O$ or $C_{10}H_{17}NO_5 + C_{20}H_{32}NO_8 - H_2O$ reactions (Claffin and Ziemann, 2018). A fourth cluster of ion signals at m/Q > 984was also observed. Unambiguous assignment of chemical formulae to these signals was challenging due to the limited range of the CIMS m/z calibration and lack of available information about $C_{>30}$ β -pinene/NO₃ oxidation products. However, it seems plausible that these signals are associated with tetramers.

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To compare our results with those obtained using a conventional NO_3 generation method (room temperature N_2O_5 thermal decomposition) in an environmental chamber study, Figures 8b and 9b show reference gas- and condensed-phase FIGAERO-I⁻-CIMS spectra of OVOCs and SOA generated from NO_3 oxidation of β -pinene in the Georgia Tech environmental chamber

- 360 (Takeuchi and Ng, 2019). The spectra obtained here and by Takeuchi and Ng (2019) exhibit an overall high degree of similarity, with linear correlation coefficients of 0.87 and 0.96 between the respective gas- and condensed-phase spectra. Clusters of $IC_{10}H_{15}NO_x^-$, $IC_{20}H_{32}N_2O_x^-$, and $IC_{30}H_{47}N_3O_x^-$ ion signals were present in both Figs. 9a and 9b. The main differences between the gas-phase spectra shown in Figs. 8a and 9a were the different abundances of $IC_{10}H_{17}NO_4^-$, a first-generation hydroxynitrate product (Claffin and Ziemann, 2018), and $IC_{10}H_{16}N_2O_7^-$. Because $C_{10}H_{17}NO_4$ is formed from RO_2+RO_2
- 365 reactions (DeVault et al., 2022) and is sufficiently volatile ($C^* = 750 \ \mu g \ m^{-3}$) to partition into the gas phase (Claflin, 2018), differences in gas-phase $C_{10}H_{17}NO_4$ and $C_{10}H_{16}N_2O_7$ yields were likely related to differences in the relative rates-importance of $RO_2 + RO_2$ and versus $RO_2 + NO_3$ reaction pathways in the study by Takeuchi and Ng (2019) compared to this work.

To further investigate the fate of RO₂ generated from VOC + NO₃ reactions as a function of CAN irradiation conditions, we calculated the fractional oxidative loss of generic alkyl and acyl RO₂ species due to reaction with HO₂, NO₃ and NO₂ $(F_{RO_2+HO_3}, F_{RO_2+NO_3}, F_{RO_2+NO_3})$ using Equations 1-3:

$$F_{\rm RO_2+HO_2} = \frac{k_{\rm RO_2+HO_2}[\rm HO_2]}{k_{\rm RO_2+HO_2}[\rm HO_2] + k_{\rm RO_2+NO_3}[\rm NO_3] + k_{\rm RO_2+NO_2}[\rm NO_2]}$$
(1)

$$F_{\rm RO_2+NO_3} = \frac{k_{\rm RO_2+NO_3}[\rm NO_3]}{k_{\rm RO_2+HO_2}[\rm HO_2] + k_{\rm RO_2+NO_3}[\rm NO_3] + k_{\rm RO_2+NO_2}[\rm NO_2]}$$
(2)

$$F_{\rm RO_2+NO_2} = \frac{k_{\rm RO_2+NO_2}[\rm NO_2]}{k_{\rm RO_2+HO_2}[\rm HO_2] + k_{\rm RO_2+NO_3}[\rm NO_3] + k_{\rm RO_2+NO_2}[\rm NO_2]}$$
(3)

Here, $k_{RO_2+HO_2}$, $k_{RO_2+NO_3}$, and $k_{RO_2+NO_2}$ are reaction rate coefficients for the corresponding $RO_2 + HO_2$, $RO_2 + NO_3$ and **375** $RO_2 + NO_2$ forward reactions whose values are summarized in Table S3. Several simplifying assumptions were made. First, we assumed that $RO_2 + NO$ reactions were negligible. Second, we did not consider RO_2 isomerization/autooxidation and $RO_2 + RO_2$ reactions that are influenced by external factors. Third, we set $F_{RO_2+NO_2} = 0$ for alkyl-RO₂-generated RO_2NO_2 , which thermally decompose on timescales of seconds or less (Orlando and Tyndall, 2012). Fourth, we assumed that vapor wall losses of acyl-RO₂-generated RO_2NO_2 were a minor RO_2 sink because the OFR residence time ($\tau_{OFR} \approx 120$ s, Sect. 2.2) was

380 significantly shorter than their estimated wall loss timescale (($\tau_{wall} \approx 400 \text{ s}$, Palm et al. (2016)). Figure A shows calculated

 $F_{\rm RO_2+HO_2}$, $F_{\rm RO_2+NO_a}$, and $F_{RO_2+NO_2}$ values for alkyl-RO₂ and acyl-RO₂ as a function of photon flux over the range of NO₃ generation conditions presented in Fig. 6. For alkyl-RO₂, $F_{\rm RO_2+HO_2}$ decreased and $F_{\rm RO_2+NO_3}$ increased with increasing photon flux and decreasing irradiation wavelength. On the other hand, for acyl-RO₂, $F_{\rm RO_2+NO_2}$ increased while $F_{\rm RO_2+HO_2}$ and $F_{\rm RO_2+NO_3}$ decreased over the same irradiation conditions. Overall, at the optimal NO₃ generation conditions (e.g. $\lambda = 369$ nm and $I_{369} \approx 10^{16}$ photons cm⁻² s⁻¹), our calculations suggest that $F_{\rm RO_2+HO_2} \approx F_{\rm RO_2+NO_3}$ (Fig. Ac) and that $F_{\rm RO_2+HO_2}$

 $\approx F_{\rm RO_2\pm NO_3} \approx F_{\rm RO_2\pm NO_2}$ was significant for acyl-RO₂ (Fig. Ag).

4 Conclusions

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 $Ce^{(IV)}$ irradiation complements $NO_2 + O_3$ reactions and N_2O_5 thermal dissociation as a customizable photolytic NO_3 source. Important method parameters were [CAN], [HNO₃] or [NaNO₃], UV intensity, and irradiation wavelength. By contrast, important parameters for NO₂+O₃ and N₂O₅-based methods are [O₃], [NO₂], temperature, and humidity. Because Ce^(IV) irra-390 diation already generates NO_3 in aqueous solution, its performance is not hindered by humidity to the same extent (if at all) as N_2O_5 -based methods, where hydrolysis of N_2O_5 to HNO₃ decreases the efficacy of the source. Additionally, the NO₃ + H₂O reaction rate in solution or on surfaces is slow relative to other NO_3 loss pathways. Another advantage of $Ce^{(IV)}$ irradiation is that it does not involve the use of O_3 as a reagent, therefore eliminating the possibility of competing O_3 and NO_3 oxidation 395 of compounds that are reactive towards both oxidants if NO_2+O_3 reactions and/or online N_2O_5 synthesis are used as the NO_3 source (Lambe et al., 2020). To identify optimal operating conditions for maximizing $[NO_3]$, we characterized concentrations of NO₃ at [CAN] = 10^{-3} to 1 M, [HNO₃] = 1.0 to 6.0 M, [NaNO₃] = 1.0 to 4.8 M, photon flux = 6.9×10^{14} to 1.0×10^{16} photons cm⁻² s⁻¹, and irradiation wavelengths of $\lambda = 254, 313, 369$, or 421 nm. With CAN/HNO₃ mixtures, maximum [NO₃] was achieved with [CAN] ≈ 0.5 M, [HNO₃] ≈ 3.0 to 6.0 M, and $I_{369} = 8 \times 10^{15}$ photons cm⁻² s⁻¹ (4.3 mW cm⁻²). With CAN/NaNO₃ mixtures, maximum [NO₃] was achieved with [CAN] ≈ 1.0 M, [NaNO₃] ≥ 1.0 M, and $I_{254} \approx 1 \times 10^{16}$ photons 400 $cm^{-2} s^{-1}$ (7.8 mW cm^{-2}). Thus, for applications such as environmental chamber or OFR studies of NO₃-initiated oxidative aging processes, where significant NO3 production over relatively short time periods is beneficial, irradiation of concentrated $Ce^{(IV)}$ solutions at high photon flux is advantageous. Other applications that require sustained NO₃ production at lower concentrations and/or over longer time periods may benefit from using lower [Ce^(IV)] and photon flux. Overall, because Ce^(IV) irradiation generates NO₃ at room temperature using widely-available, low-cost reagents and light sources (including high 405

- power light-emitting diodes in addition to, or instead of, UV fluorescent lamps) it is easier to apply than other NO_3 generation techniques - especially in field studies - and it may therefore enable more widespread studies of NO_3 oxidation chemistry. Adapting a photoreactor to operate with continuous injection of fresh $Ce^{(IV)}$ or alternative photolytic NO_3 precursors (e.g. Hering et al. (2015)) rather than in batch mode as was done here may further enhance its performance and will be investigated
- 410 in future work.

Code and data availability. Data and KinSim mechanisms presented in this manuscript are available upon request. The KinSim kinetic solver is freely available at http://tinyurl.com/kinsim-release.

Author contributions. AL, BB, and PL conceived and planned the experiments. AL, BB, MA, and PL carried out the experiments. AL conceived, planned, and carried out the KinSim model simulations. NO and PZ conceived, planned, and carried out the quantum chemical

415 calculations. AL, BB, MT, NO, PZ, MC, DW, and PL contributed to the interpretation of the results. AL took the lead in writing the manuscript. All authors provided feedback on the manuscript.

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Figure 1. Overview of experiments conducted in this study. Aqueous mixtures of ceric ammonium nitrate (CAN) and nitric acid (HNO₃) or sodium nitrate (NaNO₃) were irradiated in a photoreactor to generate nitrate radicals (NO₃) in solution. Air was bubbled through the solution to evaporate NO₃ and other volatile photolysis products into the gas phase. The photoreactor effluent was then (i) injected into a dark oxidation flow reactor (OFR) along with a VOC mixture to characterize [NO₃] via tracer decay measurements using a Vocus proton transfer-reaction time-of-flight mass spectrometer (PTR-ToF) (ii) sampled with an iodide adduct high-resolution time-of-flight chemical ionization mass spectrometer (HR-ToF-CIMS) (iii) injected into a dark OFR to characterize β -pinene/NO₃ oxidation products with a Filter Inlet for Gases and Aerosols (FIGAERO) coupled to the HR-ToF-CIMS. Supporting measurements were obtained using a NO_x analyzer.



Figure 2. Example results from an experiment in which a mixture of 0.5 M CAN and 6.0 M HNO₃ was irradiated to generate NO₃ (λ_{max} = 369 nm, I₃₆₉ = 7×10¹⁵ photons cm⁻² s⁻¹) that was injected into the OFR along with a reactive VOC tracer mixture. (**a**) Time series of the fractional consumption of VOC tracers measured with the Vocus following irradiation: thiophene (C₄H₄S), 2,3-dihydrobenzofuran (C₈H₈O), cis-3-hexenyl-1-acetate (C₈H₁₄O₂), isoprene (C₅H₈), dimethyl sulfide (C₂H₆S), 2,5-dimethylthiophene (C₆H₈S), α -pinene (C₁₀H₁₆), guaiacol (C₇H₈O₂). Signals of each tracer were normalized to their initial concentrations prior to NO₃ exposure and to acetonitrile concentrations to account for changes in the syringe pump output. (**b**) Time series of [NO₃] calculated from (**a**) and Tab. S1.



Figure 3. (a) [NO₃] values obtained from irradiated CAN & 6.0 M HNO₃ and CAN & 4.8 M NaNO₃ mixtures as a function of irradiation wavelength. Results were normalized to [NO₃] achieved with irradiation of CAN/HNO₃ mixtures at λ = 369 nm or CAN/NaNO₃ mixtures at λ = 254 nm. Error bars represent ±1 σ uncertainty in binned [NO₃] values. (b) Extinction cross sections (σ_{ext}) of CAN/HNO₃ and CAN/NaNO₃ mixtures (for details see Sect. 2.3). Additional figure notes: ¹: The black dot corresponds to data from Wine et al. (1988).



Figure 4. [NO₃] obtained from (**a**) irradiated 6.0 M HNO₃ solutions containing 0.001 to 0.5 M CAN ($I_{369} = 7 \times 10^{15}$ photons cm⁻² s⁻¹), and irradiated 1.0 M NaNO₃ solutions containing 0.5 to 1.0 M CAN ($I_{254} = 1 \times 10^{16}$ photons cm⁻² s⁻¹). (**b**) irradiated 0.5 M CAN solutions containing 1.0 to 6.0 M [HNO₃] or 1.0 to 4.8 M [NaNO₃] at the same I_{369} and I_{254} values used to obtain results shown in (**a**). Results were normalized to [NO₃] achieved with mixtures of 0.5 M CAN and 6.0 M HNO₃. Error bars represent estimated $\pm 35\%$ uncertainty in [NO₃] values obtained from CAN/HNO₃ mixtures, $\pm 15\%$ uncertainty in [NO₃] values obtained from CAN/NaNO₃ mixtures, and $\pm 10\%$ uncertainty in [CAN], [HNO₃], and [NaNO₃] values.



Figure 5. Normalized [NO₃] values obtained from irradiated mixtures of 0.5 M CAN and 6.0 M HNO₃ ($\lambda = 369$ nm) or 0.5 M CAN and 1.0 M NaNO₃ ($\lambda = 254$ nm) as a function of photon flux ranging from 6.9×10^{14} to 7.5×10^{15} photons cm⁻² s⁻¹. Results were normalized to [NO₃] achieved with 0.5 M CAN, 6.0 M HNO₃ and $I_{369} = 6.8 \times 10^{15}$ photons cm⁻² s⁻¹. Symbols are colored by the time it took for [NO₃] to experience one e-fold decay relative to the maximum [NO₃] that was measured (τ_{NO_3}). Error bars represent estimated ±35% uncertainty in [NO₃] values obtained from CAN/HNO₃ mixtures, ±15% uncertainty in [NO₃] values obtained from CAN/NaNO₃ mixtures, and ±30% uncertainty in photon flux values.



Figure 6. Model-calculated (**a**) [NO₃], (**b**) NO₂:NO₃, (**c**) HO₂:NO₃, and (**d**) N₂O₅:NO₃ values in solution as a function of photon flux ranging from 1×10^{14} to 1×10^{17} photons cm⁻² s⁻¹ following $\lambda = 254$, 313, 369 and 421 nm irradiation of a mixture containing 0.5 M CAN and 6.0 M HNO₃. [NO₃] obtained from measurements shown in Fig. 2 is plotted in (**a**). For details see Sect. 2.3 and Tab. S2.



Figure 7. Time series of **a** (a) NO₂ and NO₃, (b) N₂O₅ and N₂O₆, (c) HNO₂ and HNO₄, and (d) HO₂ and H₂O₂ detected following irradiation of a mixture containing 0.5 M CAN and 1.0 M NaNO₃. N₂O₅, N₂O₆, HO₂ and H₂O₂ were detected as I⁻ adducts, and HNO₂ and HNO₄ were detected as both I⁻ and NO₃⁻ adducts with HR-ToF-CIMS. CIMS signals detected as iodide adducts were normalized to the I⁻ signal prior to the start of the experiment, and CIMS signals detected as nitrate adducts were normalized to the maximum NO₃⁻ obtained during the experiment (see Fig. S5).



Figure 8. HR-ToF-CIMS spectra of gas-phase β -pinene/NO₃ oxidation products obtained following β -pinene reaction with NO₃ generated via (a) irradiation of a mixture of 0.5 M CAN and 3.0 M HNO₃ and subsequent injection into the OFR (b) thermal decomposition of N₂O₅ injected into the Georgia Tech environmental chamber. Signals shown are unmodified (M+I)⁻ formulas.



Figure 9. FIGAERO-HR-ToF-CIMS spectra of condensed-phase β -pinene/NO₃ oxidation products obtained following β -pinene reaction with NO₃ generated via (a) irradiation of a mixture of 0.5 M CAN and 3.0 M HNO₃ and subsequent injection into an OFR (b) thermal decomposition of N₂O₅ injected into the Georgia Tech environmental chamber. Signals shown are unmodified (M+I)⁻ formulas. Bands of ion signals corresponding to C₁₀H₁₅NO_x, C₂₀H₃₂N₂O_x, and C₃₀H₄₇N₃O_x oxidation products are highlighted and colored by the number of oxygen atoms in their chemical formulas.



Figure 10. Fractional oxidative loss of alkyl and acyl organic peroxy radicals (RO₂) due to reaction with HO₂, NO₃ and NO₂ ($F_{RO_2+HO_2}$. $F_{RO_2+NO_3}$ and $F_{RO_2+NO_2}$) generated following $\lambda = (a)$, (e) 254 (b), (f) 313 (c), (g) 369 (d), (h) 421 nm irradiation of a mixture containing 0.5 M CAN and 6.0 M HNO₃ as a function of photon flux ranging from 1×10^{14} to 1×10^{17} photons cm⁻² s⁻¹.