Real time in situ detection of organic nitrates in atmospheric aerosols

Andrew W. Rollins,† Jared D. Smith,‡ Kevin R. Wilson,‡ and Ronald C. Cohen*,†,¶

Department of Chemistry, University of California, Berkeley, Chemical Sciences Division, Lawrence Berkeley National Laboratory, and Department of Earth and Planetary Science, University of California, Berkeley

E-mail: rccohen@berkeley.edu

Abstract

A novel instrument is described that quantifies total particle phase organic nitrates in real time with a detection limit of 0.11 µg m\(^{-3}\) min\(^{-1}\), 45 ppt min\(^{-1}\) (-ONO\(_2\)). Aerosol nitrates are separated from gas phase nitrates with a short residence time activated carbon denuder. Detection of organic molecules containing -ONO\(_2\) subunits is accomplished using thermal dissociation coupled to laser induced fluorescence detection of NO\(_2\). This instrument is capable of high time resolution (seconds) measurements of particle phase organic nitrates, without interference from inorganic nitrate. Here we use it to quantify organic nitrates in secondary organic aerosol generated from high-NO\(_x\) photooxidation of limonene, \(\alpha\)-pinene, \(\Delta\)-3-carene, and tridecane. In these experiments the organic nitrate moiety is observed to be 6-15% of the total SOA mass.

*To whom correspondence should be addressed
†Department of Chemistry, University of California, Berkeley
‡Chemical Sciences Division, Lawrence Berkeley National Laboratory
¶Department of Earth and Planetary Science, University of California, Berkeley
Introduction

Organic nitrates (RONO$_2$) are formed in the atmosphere as a minor product of reactions between organic peroxy radicals and nitric oxide (NO), as well as a major product from reactions of alkenes with the nitrate radical (NO$_3$). As a result, RONO$_2$ molecules are chemically produced during the day and the night and are a significant fraction of the total oxidized nitrogen (NO$_y$) anywhere that non-methane organic compounds are available (e.g. (I)). Specific individual gas phase RONO$_2$ molecules have been observed using gas chromatography (e.g. (2–6)) and chemical ionization mass spectrometry (7–9), and the total of gas and aerosol RONO$_2$ (total Alkyl Nitrates, ΣAN) has been observed by Thermal Dissociation Laser Induced Fluorescence (TD-LIF) (I, 10–12). Recently there has been growing interest in understanding the partitioning of RONO$_2$ to atmospheric particles (13–18). RONO$_2$ molecules with 2 - 4 functional groups and a carbon backbone of $\geq$ C$_4$ have low enough vapor pressures to condense under ambient conditions and mechanistic studies show that RONO$_2$ can be among the most important molecules to SOA formation (e.g. (13, 19)).

Despite the theoretical and laboratory evidence that RONO$_2$ should be important to the composition of atmospheric organic aerosols, there remains little evidence that RONO$_2$ are widespread components of ambient SOA. The most compelling evidence derives from direct FTIR analysis of particles impacted on ZnSe plates. This technique has been used to identify organic nitrates as ubiquitous components of submicron particles sampled in the Los Angeles basin (20, 21) and in Houston (22). FTIR analysis provides quantitative measurements of positively identified organic functionalities, with time resolution limited to the rate at which filters are changed (typically hours - days). Interferences due to evaporative loss from the filters or heterogeneous reactions on the filters during sampling are known issues and difficult to quantify. Individual large multifunctional nitrates have been observed in ambient aerosol samples collected on filters using electrospray ionization mass spectrometry (23, 24). However the most widely used methods for aerosol chemical characterization including Aerosol Mass Spectrometry (AMS) have had limited success. The difficulty of distinguishing inorganic and organic nitrates using AMS has been recently discussed in the literature (16–18). Nitrates have not yet been reported by Thermal Desorption Aerosol Gas-
Chromatography (TAG) (25) or other methods of detailed realtime aerosol characterization.

In this paper a new technique for the detection of aerosol organic nitrate is described. The new method quantifies particulate organic nitrate with high time resolution, low background signal, and no interference from inorganic nitrate ion ($\text{NO}_3^-$). Here we apply the method to measure the organic nitrate content of SOA produced in the laboratory from high NO$_x$ photooxidation of one alkane and three alkenes. In these experiments, we show that the fraction of the total organic mass which is organic nitrate is consistent with the expected organic nitrate yields from high NO$_x$ photooxidation.

**Instrument Design and Evaluation**

The selective measurement of particle phase nitrates is achieved by removing gas phase NO$_2$ and nitrates (both RO$_2$NO$_2$ and RONO$_2$) from the sample stream with an activated carbon denuder. The aerosol flow is then heated so that particles are vaporized to their gas phase precursors and detected using TD-LIF by which RONO$_2$ is thermally converted to RO + NO$_2$ and NO$_2$ is detected by LIF (10). A schematic of the instrument inlet system used in this study is shown in Figure 1.

---

Figure 1: Schematic of particle organic nitrate inlet system. Gas and particles are sampled through $\approx$10 cm of 0.25" o.d. stainless tube. The flow enters the denuder where particles (green dots) pass while gas phase NO$_y$ (brown dots) is removed by uptake on the denuder walls. Organic nitrate particles are converted to NO$_2$ in a heated quartz tube. A critical orifice reduces the pressure to $\approx$2 torr for measurement by LIF.
Thermal Dissociation Laser Induced Fluorescence

The core measurement used in TD-LIF is the laser induced fluorescence detection of NO$_2$ (26). Details of the specific LIF instrument used for this study are described in Wooldridge et al. (27). Briefly, gas is sampled through a critical orifice into a 38 pass White cell held at $\approx 2$ torr. NO$_2$ is excited with a 30 mW diode laser centered at 408 nm (Power Technologies Inc.). Red shifted NO$_2$ fluorescence passes through a 650 nm long pass dielectric filter and a red glass filter which block scatter and is detected with a photomultiplier tube (Hamamatsu H7421-50). Alkyl nitrates and nitric acid (HNO$_3$) are quantified by sampling through a heated inlet which thermally dissociates these molecules to NO$_2$ and a corresponding radical. The method has been compared to independent approaches for detection of NO$_2$ (28, 29), total peroxy nitrates (ΣPN) (27) and ΣAN (9). In this work we used an inlet which consisted of a 4 mm i.d. quartz tube wrapped in nichrome wire providing a heated length of 20 cm. Using a flow rate of 125 sccm the residence time in the heated region was 1.2 s. The setpoint used to thermally dissociate all organic nitrates, while rejecting HNO$_3$ was determined by scanning the inlet temperature while sampling a mixture of ethyl nitrate and HNO$_3$ (Figure 2).

TD-LIF has previously been used to quantify various classes of gas + particle NO$_y$ by taking the difference in LIF signals measured with two inlets at two different temperatures (10). For the application described in this work, gas phase species including NO$_2$ are separated from particles before entering the heated portion of the inlet, and thus no subtraction is required.

Gas/Particle Separation

Gas phase NO$_y$ is removed using a single channel cylindrical denuder, 25 cm in length, and 1.0 cm i.d., which provides 9.3 s residence time at a flow rate of 125 sccm. We tested the effectiveness of the denuder for removing NO$_2$ and organic nitrates by using an injector inside the denuder in a concentric flow tube configuration. We measured the penetration of NO$_2$ and n-propyl nitrate (NPN) as a function of length of denuder exposed and gas flow rate. The data were compared to the predicted penetration as calculated by the expression obtained by Ingham et al.(30) (equations
Figure 2: Inlet temperature scan on a mixture of gas phase n-propyl nitrate and nitric acid.

23 and 24 in that paper). The experiments showed that uptake of NO\textsubscript{2} and NPN by the denuder walls could be accurately predicted assuming removal is limited only by gas phase diffusion. In Figure 3 we show observations and model predictions of gas penetration from experiments using NO\textsubscript{2} and NPN. The diffusivity of NO\textsubscript{2} in air is 0.154 cm\textsuperscript{2} s\textsuperscript{-1} (31) and that of NPN was estimated using a hard sphere model and the method of Chapman and Cowling (32) to be 0.070 cm\textsuperscript{2} s\textsuperscript{-1}. The largest deviations between the model and data occur at the higher flow rates and short denuder lengths and are likely due to the entrance length required to establish laminar flow, which was not accounted for here but is estimated to be less than 1 cm at 125 sccm. At the lowest flow rates of 255-260 sccm used in the experiments the model and measurements are in excellent agreement. We use the diffusion limited model to calculate the removal efficiency as a function of diffusivity at 125 sccm. The bottom panel of Figure 3 shows that at this flow rate even large gas phase molecules are predicted to be removed efficiently.

Once particles leave the denuder they enter the heated quartz tube at 325°C where they evaporate. The approximate time required to completely evaporate the particles was calculated using a model which integrates the Hertz-Knudsen equation (Eq. (1)) assuming evaporation and condensation onto spherical particles.

\[
J = \frac{\gamma p}{\sqrt{2\pi mk_BT}}
\]  

(1)
Figure 3: Top: Measured penetration of gas phase NO$_2$ and NPN in the denuder. Bottom: Calculated removal of gas phase nitrates at a flow rate of 125 sccm as a function of diffusivity. NO$_y$ compounds are noted by their measured (NO$_2$, HNO$_3$) and approximated (others) diffusivities in air.

Here, $J$ is the molecular flux across the particle surface (molecules cm$^{-2}$ s$^{-1}$), $\gamma$ is the evaporation coefficient, $p$ is the vapor pressure of the compound (saturation vapor pressure for evaporation ($p_{sat}$) and actual partial pressure in the vapor phase for condensation), $m$ is the molecular mass (g), $k_B$ is the Boltzmann constant, and $T$ is the temperature. Details of the particle evaporation calculations are included in the Supporting Information. We calculate that for $\gamma=1$, particles with $C^* > 10^{-5}$ $\mu$g m$^{-3}$ evaporate in < 10 ms. Complete evaporation of $\alpha$-pinene SOA in the 1.2 s inlet was verified in one experiment by using an additional heating section with a 2 minute residence time at 150$^\circ$C which is calculated to evaporate particles with log$_{10}C^*$ greater than approximately -7.5. In this experiment no differences in signal were observed between having this additional section held at room temperature or at 150$^\circ$C, indicating that particles are completely evaporated in the shorter 325$^\circ$C inlet.
Particle Transmission Efficiency

Because semivolatile particles exist in equilibrium with their gas phase precursors, removal of the gas in the denuder might lead to evaporative loss of particles in the system (33). To estimate the importance of this process we used the same model as described here for calculating the evaporation of particles in the heated part of the inlet, this time setting the condensation rate of vapors onto the particles to zero. Figure S2 shows the calculated fraction of particle mass remaining as a function of $C^*$ assuming 9.3 s at 300 K for 4 initial particle sizes. For this calculation we set the evaporation coefficient to 1, which is the upper limit of evaporative loss. For $C^* < 1$ all particle mass is calculated to be transmitted with >97% efficiency. At $C^* = 10$, 25% of the mass of the smallest particles is lost, but for particles larger than 100 nm, more than 88% of mass is transmitted. Significant particle loss may occur for more volatile particles of $C^* > 100$. However, these more volatile particles are not expected to be an appreciable fraction of the mass in ambient aerosols (34, 35), and while particles smaller than 100 nm may be important by number, in a mass weighted distribution they are also of little importance.

Loss of particles due to diffusion or thermophoresis in the heated part of the inlet are expected to be non-negligible. To determine an upper limit to the loss of particles due to these combined effects we measured the transmission of non-volatile NaCl particles through the denuder and thermal-dissociation heater sections of the instrument. Details and data from this experiment are provided in the Supporting Information. Significant losses were observed for the smallest particles, while particles with diameters greater than 100 nm were transmitted with >85% efficiency. The transmission efficiency observed with NaCl is expected to represent a lower limit to the transmission and detection efficiency for semivolatile particles, as semivolatile particles are expected to evaporate and dissociate to NO$_2$ rapidly rather than deposit on the walls of the tube in regions of strong thermal gradients.

The effects of thermophoretic and evaporative loss were combined to evaluate the total penetration of particle mass through the inlet. The results show that the evaporative losses are negligible compared to deposition of particles in the inlet for $C^* < 1$. At $C^* = 10$, the difference between
deposition only and deposition combined with evaporation is largest for particles with diameters between 100 and 200 nm, with a difference of about 10% of the mass for a 100 nm particle and about 5% for a 200 nm particle. Generally, the depositional losses we observed dominate the total mass loss in all conditions, though as discussed the detection efficiency of semivolatile particles is likely higher than NaCl particles.

**Detection Limit**

The lower limit of detection of the instrument is determined by the ratio of the NO$_2$ LIF signal relative to the to noise produced from scatter and single photon counting statistics. Using $c$ as the NO$_2$ mixing ratio (ppb), $A$ as the calibration constant (counts s$^{-1}$ ppb$^{-1}$) and $B$ as the background count rate (counts s$^{-1}$) the signal to noise ratio for some integration time $t$ (s) is given by

$$\frac{\text{Signal}}{\text{Noise}} = \frac{Act}{\sqrt{Bt + Act}}$$

We define the detection limit as the minimum NO$_2$ concentration required for a signal:noise ratio of more than 2. The instrument used in this study had a typical background signal of 675 counts s$^{-1}$ and a typical calibration constant of 150 counts s$^{-1}$ ppb$^{-1}$. These values result in a detection limit of 110 ppt for 10 seconds of averaging, or 45 ppt for one minute. Higher sensitivity NO$_2$ instruments are available and have been employed in other studies (26, 27, 29). At standard conditions (1013 hPa, 300 K) the detection limit for this instrument converts to 0.11 $\mu$g -ONO$_2$ m$^{-3}$ min$^{-1}$. It is worth noting that since the instrument is sensitive to the mixing ratio of NO$_2$ produced by the evaporation and thermal dissociation of particles as opposed to the volumetric density, the sensitivity in $\mu$g m$^{-3}$ improves at increased altitude. For example, at 300 hPa the detection limit would be 0.033 $\mu$g -ONO$_2$ m$^{-3}$ min$^{-1}$. The total inlet residence time is calculated to be $\approx$ 10 sec and experiments demonstrated that the instrumental response time to ambient changes was changes was in good agreement with this value. No extended residual signal was observed upon sudden changes such as overflowing the inlet with zero air. Figure 4 shows sample observations with 10
SOA Experiments

A series of secondary organic aerosol experiments was performed to measure the contribution of organic nitrates to SOA formed under high-NOₓ conditions. Experiments were carried out in a quartz flow tube reactor with an inner diameter of 65 mm. The tube was surrounded by 4, T10 Hg vapor lamps, 130 cm long (UHP, LLC) providing \( \approx 12 \) mW cm\(^{-2}\) of 254 nm light. OH was generated through the photolysis of H\(_2\)O\(_2\) which was produced by flowing 100 sccm N\(_2\) through a mixture of urea hydrogen peroxide (Sigma) and sand which was warmed slightly (30-40\(^\circ\)C). Continuous streams of limonene, \( \alpha \)-pinene, \( \Delta \)-3-carene, or tridecane vapors were produced by flowing 1000 sccm N\(_2\) over the output of a syringe pump loaded with the respective liquids. The concentrations of the organic compounds added to the flow tube were controlled by varying the pumping rate of the syringe pump, and/or the fraction of the 1000 sccm N\(_2\)/organic mixture which was exhausted rather than sent to the flow tube. 10 ppm NO was added at 100 sccm along with N\(_2\) and O\(_2\) to balance the total flow in the reactor at 2000 sccm and achieve an O\(_2\) concentration of 20\% and an NO concentration of 500 ppb. Though the concentrations of the organics were typically <100 ppb, the 500 ppb NO concentrations were used to be assured the experiments were high-NO\(_x\). Average OH concentrations were \( \approx 10^9 \) molecules cm\(^{-3}\) as was verified by measuring the loss rate of hexane. The steady state HO\(_2\) produced by H\(_2\)O\(_2\) photolysis converts a large fraction of the NO to NO\(_2\). The Master Chemical Mechanism (MCM) (36) was used with modified photolysis rates for 254 nm to calculate that >90\% of the organic peroxy radicals reacted with NO in the experiments. The MCM simulations were run for \( \alpha \)-pinene and n-dodecane (C\(_{12}\)H\(_{26}\)). Detailed reaction mechanisms are not currently available in the MCM for tridecane, limonene or \( \Delta \)-3-carene. However the tridecane and dodecane mechanisms are expected to be almost identical, and \( \alpha \)-pinene, limonene and \( \Delta \)-3-carene have very similar photochemistry (37, 38).

A scanning mobility particle sizer (SMPS) system (TSI) was used to measure particle size...
Figure 4: Sample time series from Δ-3-carene experiment. The time series begins sampling air from the irradiated H₂O₂ / air mixture. NO is added just after 9:45, and then at 9:50 more NO is added along with carene. The different aerosol concentrations were achieved by varying the syringe pump flow rate used to deliver the liquid organic material to the air flow stream, and by flowing various amounts of the organic vapor / air mixture into the flow tube vs. to the exhaust.
distributions. Total particle mass was calculated from the SMPS data assuming a particle density of 1.2 g cm\(^{-3}\) which is similar to that reported in a number of studies for high-NO\(_x\) SOA of terpenes (39–44). Particle number distributions were typically centered at 100 - 150 nm. Figure 4 shows a sample time series of NO\(_2\), particle -ONO\(_2\) and total aerosol mass from the Δ-3-carene experiment. In this experiment, the organics, H\(_2\)O\(_2\) and air were initially flowing into the flow tube reactor. Just after 9:45 the NO flow was also added and due to a slight NO\(_2\) impurity 12 ppb NO\(_2\) was observed. The bottom panel of Figure 4 shows a magnified view of the first 15 minutes of this experiment confirming that NO\(_2\) is not an interference for the particle -ONO\(_2\) measurement. At 9:51 the lights were turned on, generating OH and HO\(_2\), which produced SOA and NO\(_2\). After 10:00 the concentration of aerosol produced was varied by adding additional the amount of organic precursor to the flow tube at 10:15, 10:25 and 10:37.

Figure 5 shows regression plots of the measured particle -ONO\(_2\) vs. total particle mass from the four different organic experiments. In all four experiments the particle -ONO\(_2\) and total particle mass were well correlated (\(R^2 > 0.94\)). The slopes of the linear fits (y-intercepts were held at zero) are the fraction of aerosol mass which is the -ONO\(_2\) functional group. The observed -ONO\(_2\) content was 6.2% for α-pinene, 8.5% for Δ-3-carene, 12% for limonene and 15% for tridecane. The assumed SOA density used to calculate the total mass from the SMPS data has a direct impact on these calculations. Based on the range of reported values for α-pinene SOA (39–44), 1.2 ± 0.2 g cm\(^{-3}\) is the maximum expected range of particle density produced here, resulting in a ±17% accuracy in the total calculated OA mass and fractional -ONO\(_2\) content.

To estimate the fraction of SOA molecules which are organic nitrates, we make some simple assumptions about the presumed oxidation products of the reactions. The monoterpenes are expected to react with OH and then O\(_2\) to produce hydroxy alkyl peroxy radicals. Under high NO\(_x\) conditions the dominant pathway is for the peroxy radicals to react with NO to form either a hydroxy-nitrate with a molecular weight of 215, or a hydroxy-alkoxy radical which reacts with O\(_2\) to form a hydroxy-carbonyl compound (MW = 168) (38). Analogous assumptions for the tridecane + OH reaction which is initiated by hydrogen abstraction results in an alkyl nitrate (MW = 245)
Figure 5: Regressions of particle NO$_3$ and total aerosol mass measurements from the high-NOx photooxidation experiments.
and a hydroxy-carbonyl (MW = 215) (45). If we assume that the observed -ONO$_2$ content of the aerosol is due to a linear combination of these two types of molecules, we can calculate the fraction of SOA molecules which are hydroxy-nitrates. Under these assumptions we calculate 18% of the molecules are organic nitrates for $\alpha$-pinene SOA, 25% for $\Delta$-3-carene, 36% for limonene and 56% for tridecane. No doubt the product distribution which contributed to the observed SOA was more complicated than described here, but our analysis relies only on the approximate molecular weights. If for example we assume that all SOA (nitrate and non-nitrate) from the $\alpha$-pinene reaction had a molecular weight of 150, we calculate 15% instead of the 18% for condensed organic nitrates. These values are in agreement with previously reported yields of total organic nitrates from high-NO$_x$ photooxidation of e.g. $\alpha$-pinene (18 ± 9%) (46). The correspondence between the previously observed and calculated total alkyl nitrate yields, and our calculated contribution of alkyl nitrates to SOA (i.e. 18% total RONO$_2$ (46) and 18% of SOA is RONO$_2$), suggests that the organic nitrate products of high-NO$_x$ photooxidation are roughly as condensable as the non-nitrate products. To our knowledge, the organic nitrate yields from limonene and $\Delta$-3-carene have not been reported. Total alkyl nitrate yield for the first generation products of tridecane are expected to be $\approx$ 35% (10). Lim et al. (13) have studied the SOA formed from high-NO$_x$ photooxidation of a series of n-alkanes and concluded that all observed SOA was organic nitrate.

In summary, we have described and evaluated a new technique which quantifies the total organic nitrate component of aerosols with high time resolution. The technique is sensitive, accurate and specific to organic nitrates. Near unit efficiency was observed for the detection of particles larger than 100 nm. The measurement was used to show that organic nitrates can be a significant contributor to high-NO$_x$ SOA, with the nitrate group itself being an important contribution to the total aerosol mass. These experiments suggest that in general, the organic nitrate fraction of SOA molecules produced in photooxidation, may be similar to the total yield of RONO$_2$ from the RO$_2$ + NO reaction, at least for molecules that are C$_{10}$ or larger.
Acknowledgement

The Berkeley authors were supported by NSF ATM-0639847. Part of this work utilized equipment at the Chemical Dynamics Beamline, which is supported by the Director, Office of Energy Research, Office of Basic Energy Sciences, and Chemical Sciences Division of the U.S. Department of Energy under contracts No. DE-AC02-05CH11231

Supporting Information Available

Additional details of the evaporation model as well as experimental methods and data associated with measuring inlet transmission is included in the Supporting Information. This material is available free of charge via the Internet at http://pubs.acs.org/.

References


(13) Lim, Y. B.; Ziemann, P. J. Products and Mechanism of Secondary Organic Aerosol Formation


LEGAL DISCLAIMER

This document was prepared as an account of work sponsored by the United States Government. While this document is believed to contain correct information, neither the United States Government nor any agency thereof, nor The Regents of the University of California, nor any of their employees, makes any warranty, express or implied, or assumes any legal responsibility for the accuracy, completeness, or usefulness of any information, apparatus, product, or process disclosed, or represents that its use would not infringe privately owned rights. Reference herein to any specific commercial product, process, or service by its trade name, trademark, manufacturer, or otherwise, does not necessarily constitute or imply its endorsement, recommendation, or favoring by the United States Government or any agency thereof, or The Regents of the University of California. The views and opinions of authors expressed herein do not necessarily state or reflect those of the United States Government or any agency thereof or The Regents of the University of California.