



Evidence for a Kinetically Controlled Burying Mechanism for Growth of High Viscosity Secondary Organic Aerosol

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Abstract

Secondary organic aerosol (SOA) particles are ubiquitous in air and understanding the mechanism by which they grow is critical for predicting their effects on visibility and climate. The uptake of three organic nitrates into semi-solid SOA particles formed by α -pinene ozonolysis either with or without an OH scavenger was investigated. Four types of experiments are presented here. In Series A, uptake of the selected organic nitrates (2-ethylhexyl nitrate (2EHN); β-hydroxypropyl nitrate (HPN); β-hydroxyhexyl nitrate (HHN)) into impacted SOA particles was interrogated by attenuated total reflectance (ATR)-FTIR. In this case, equilibrium was reached and partition coefficients ($K_{SOA} = [-ONO_2]_{SOA}/[-ONO_2]_{air}$) were measured to be $K^{2EHN} = (3.2-11) \times 10^4$, $K^{HPN} = (4.4-5.4) \times 10^5$, and $K^{HHN} = (4.9-9.0) \times 10^6$. In Series B, SOA particles were exposed on-the-fly to gas phase organic nitrates for comparison to Series A, and uptake of organic nitrates was quantified by HR-ToF-AMS analysis, which yielded similar results. In Series C (AMS) and D (ATR-FTIR), each organic nitrate was incorporated into the SOA as the particles formed and grew. The incorporation of the RONO₂ was much larger in Series C, D (*during growth*), exceeding equilibrium values determined in Series A, B (*after* growth). This suggests that enhanced uptake of organic nitrates during SOA formation and growth is due to a kinetically controlled "burying" mechanism, rather than equilibrium partitioning. This has important implications for understanding SOA formation and growth under conditions where the particles are semi-solid, which is central to accurately predicting properties for such SOA.

Environmental Significance

Secondary organic aerosol (SOA) particles are known to have deleterious effects on human health and to impact climate and visibility. Understanding the processes by which gases are

incorporated into these particles to grow them to diameters large enough to have these impacts is therefore critical for predicting and addressing their effects. In this work, we examine how gas phase organic nitrates interact with SOA particles either during the particle growth process in a flow reactor or after particle growth has occurred. It is shown that co-condensation of organics that grow the particles enhances particulate organic nitrate content relative to that expected from equilibrium partitioning, and is best described by a kinetically controlled 'burying' mechanism.

Introduction

 Secondary organic aerosol (SOA) is a major contributor to airborne particles, which are known to impact human health,¹⁻⁷ visibility⁸⁻¹¹ and climate.^{7-9, 12, 13} Yet the molecular processes that lead to SOA particle formation from the oxidation of volatile organic compounds and the subsequent growth of these particles is not well understood.¹³⁻¹⁷ Specifically, there is not yet a comprehensive understanding of the mechanism by which gases are taken up into the particles to grow them.¹⁸

Many models have been developed to explain the mechanisms by which particles grow and the various physical and chemical processes involved.¹⁹⁻³⁴ In some cases, particles may have relatively high viscosity and a semi-solid or glassy physical state, and hence the growth process is subject to diffusion limitations.^{25, 26, 35-47} This has important implications for particle growth as well as for chemical reactions and interactions occurring both between the gas and condensed phases, and within the condensed phase.^{25, 26}

Organic nitrates are known to be formed by NO₃ radical oxidation of volatile organic compounds (VOCs) and OH radical oxidation of VOCs in the presence of NO_x.⁴⁸⁻⁵⁴ Additionally, organic nitrates (including alkyl nitrates and multifunctional hydroxy nitrates) have been measured in air.⁵⁵⁻⁶⁶ Recently, initial uptake and bulk partition coefficients for three organic nitrates into several thin film substrates of selected organic compounds and into particles from the ozonolysis of α -pinene were measured.⁶⁷ The organic nitrates included the alkyl nitrate (2-ethylhexyl nitrate, 2EHN) and two isomeric β -hydroxy nitrates (β -hydroxyhexyl nitrate, HHN, and β -hydroxypropyl nitrate, HPN), shown in Figure 1. It was found that the trend in uptake coefficients did not uniformly track the trend in partition coefficients, suggesting that

 interactions controlling initial uptake (e.g., surface of the particle) were different from those determining the equilibrium partitioning into the substrates (bulk properties).

The goal of this study was to elucidate the mechanism of incorporation of those three organic nitrates into high viscosity semi-solid SOA particles formed in the α -pinene ozonolysis. This was accomplished by measuring the organic nitrate content of SOA in particles *during their formation and growth*, and by comparison, the uptake of the organic nitrates into "fully grown" SOA. Zelenyuk *et al.*⁶⁸ previously demonstrated that when polycyclic aromatic hydrocarbons (PAHs) were incorporated *during SOA formation*, they remain trapped inside the particles due to the highly viscous semi-solid nature of the SOA and remained shielded from oxidation and evaporation; however, when the PAHs interacted with the SOA already formed, they remained on the surface and were subject to greater evaporation rates. In the current study, we show that a higher partition coefficient is obtained when the organic nitrate tracer is present *during formation and growth* than when equilibrium partitioning into fully grown SOA is measured. This suggests that a kinetically controlled "burying" mechanism is responsible for uptake into highly viscous, semi-solid particles.

Experimental

Figure 2 shows a schematic of the four experimental configurations used in these studies. In Series A, nitrate-free SOA particles formed in the α -pinene ozonolysis were generated in a large volume, slow flow stainless steel aerosol flow reactor⁶⁹ and impacted on an attenuated total reflectance (ATR) crystal to generate a thin film of SOA particles, over which gas phase organic nitrate was flowed and uptake of the organic nitrate was measured until equilibrium was reached. This method was limited to higher concentrations of organic nitrate (approaching saturation vapor pressure) in order to detect -ONO₂ peaks by ATR-FTIR. A benefit of this method over in

situ production of organic nitrates is that it avoids oxidation of the organic nitrate compounds from gas phase OH that is generated during ozonolysis in the reactor.

In Series B, SOA particles formed in the α -pinene ozonolysis also from the large flow reactor were first passed through a monolith carbon denuder (NovaCarbTM; MAST Carbon, Ltd.) to remove gas phase species and then flowed into a smaller glass flow tube where they were exposed to gaseous 2EHN. This limited series was carried out to ensure that the observations were in agreement with Series A results on impacted particles.

In Series C and D, the organic nitrates were introduced into the stainless steel flow reactor and were incorporated into SOA particles from α -pinene ozonolysis as they formed and grew in the reactor. Smaller concentrations of the organic nitrates than those in Series A were used here due to the much larger volumes of air and hence higher dilution factors that are associated with the large flow reactor. In Series C, the organic nitrate in the particles was quantified *on-the-fly* by high resolution time-of-flight aerosol mass spectrometry (HR-ToF-AMS). In Series D the same particles as in Series C were simultaneously impacted on an ATR crystal and quantification of each organic nitrate (RONO₂) was carried out by FTIR.

Aerosol Generation and Particle Size Distributions

SOA from the ozonolysis of α -pinene (AP) was generated in the stainless steel flow reactor with a total flow rate of 34 L min⁻¹, and all reactants were introduced in the initial mixing section of the reactor. Gas-phase AP (250 ppb) was generated by injection of the pure liquid (Sigma Aldrich, >99%) from an automated syringe pump (New Era Pump System Inc., Model NE-1000) into a 10 L min⁻¹ flow of clean, dry air. Ozone was generated by flowing 0.4 L min⁻¹ O₂ gas (Praxair, 99.993%) through a Pen-Ray® mercury lamp (UVP, LLC), and was

subsequently diluted with 9.6 L min⁻¹ of air before being added to the system. An additional 14 L min⁻¹ of air was introduced to create a total flow rate of 34 L min⁻¹, and the resulting reactor concentrations were 250-300 ppb O₃ measured using an ozone monitor (Teledyne Photometric O₃ Analyzer 400E; Advanced Pollution Instrumentation, Inc. Photometric O₃ Analyzer 400). The reaction of AP with O₃ produces OH radicals which, unless scavenged, will react with the organic nitrates.⁷⁰ Thus in some experiments, cyclohexane (CH, Fisher Scientific, 99.9%), used as an OH scavenger was evaporated into the flow of air to give a concentration of 2.5 × 10¹⁵ molecules cm⁻³ (100 ppm).

Three organic nitrates (Fig. 1) were used as "tracer" compounds because of their ability to be detected via both FTIR and mass spectrometry. An alkyl nitrate, 2-ethylhexyl nitrate (2EHN, Sigma Aldrich, 97%) was used as purchased. Two multifunctional organic nitrates, β hydroxyhexyl nitrate and β-hydroxypropyl nitrate (HHN and HPN, respectively, 82-93% purity in the liquid phase) were synthesized using the method of Cavdar and Saracoglu⁷¹ as described previously.⁶⁷ In brief, epoxypropane (Sigma Aldrich, \geq 99%) or epoxyhexane (Sigma Aldrich, 97%) were reacted with Bi(NO₃)₃ • 5 H₂O (Sigma Aldrich, 98%) in dichloromethane (Macron, \geq 99.5%) for 16-24 hours under N₂ gas (Praxair, 99.999%) at room temperature, after which the solvent was evaporated off. After synthesis, each organic nitrate was stored under an inert atmosphere of N₂ in a freezer (T \sim -20 °C). The purity of the resulting liquid organic nitrate was quantified by NMR, with the major impurities identified as the corresponding di-alcohols. Compared to the impurities in the liquid phase, analysis of the gas-phase directly flowing out of a glass trap by direct analysis in real time mass spectrometry (DART-MS) did not detect impurities.⁶⁷ The isomeric ratios of the hydroxy-terminated to nitrate-terminated isomer of the pure liquid estimated by NMR were $\sim 2:1$ for HPN and $\sim 4:3$ for HHN.

For series A experiments, each individual organic nitrate was introduced by flowing 0.06 L min⁻¹ air through the glass trap to yield high gas-phase concentrations approaching the saturation vapor pressure before being introduced into the ATR-FTIR cell. The concentration of each gas-phase organic nitrate was controlled by keeping the trap at room temperature using a water bath, and the concentration of organic nitrate exiting the trap was measured by GC-MS as described below. For the limited series B experiments, a reservoir of liquid 2EHN was placed inside the mini flow tube in order to expose particles *on-the-fly* to the saturation vapor pressure of 2EHN in a total flow of air of 0.4 - 1.5 L min⁻¹. In a separate experiment for Series B, a glass trap containing the pure 2EHN liquid was used in place of the reservoir to flow 0.04-0.15 L min⁻¹ into the mini flow tube to provide more dilute 2EHN concentrations. For series C and D experiments, each organic nitrate was introduced by flowing 1 L min⁻¹ air through the glass trap containing the pure liquid into a stream of air totaling 10 L min⁻¹ simultaneously with AP, either with or without CH.

Experiments were performed under ambient temperature (295-298 K) and pressure, and dry conditions (RH < 5%), without seed particles. The particles were sampled along the length of the flow system at 7 min and 31 min reaction time. Particle size distributions were monitored using a scanning mobility particle sizer (SMPS, TSI), equipped with a model 3080 classifier, a 3081 long differential mobility analyzer, and a 3776 butanol-based CPC.

Gas Phase Measurements

Gas phase concentrations of AP and CH were measured using EI GC-MS (Agilent 7890A GC system with a 5975C MS detector) in a dual total ion/single ion monitoring (SIM) system (total ion monitoring was used for CH while m/z 93 was followed in SIM mode for AP), with the

 particles and ozone filtered out using a quartz filter (TissuQuartz®; 37 mm; PallFlex) and a KI (Fisher Chemical, 100.0%) ozone scrubber, respectively.

The gas phase concentrations of all three organic nitrates were measured by flowing air at 1 L min⁻¹ through the trap containing each nitrate into a 1 mL sampling loop on the GC-MS and comparing to a calibration using the synthesized liquid standards. The concentrations measured in this manner are shown in Table 1. Independently, the vapor pressures of all three organic nitrates were estimated using two group contribution methods⁷²⁻⁷⁴ and are also listed in Table 1. The estimated vapor pressures are in reasonable agreement with the measured values, indicating that these group contribution methods are good predictors of vapor pressure for these species. Hereafter, the concentration measured by GC-MS of each organic nitrate exiting the trap is used in all calculations, factoring in any additional dilution factors. For example, after the dilution factor of 34 in the flow reactor, the final gas phase concentrations are 1.4×10^{14} (2EHN), 1.2×10^{14} 10^{14} (HPN) and 5.0×10^{12} (HHN) molecules cm⁻³. Concentrations of 2EHN measured by GC-MS directly from the flow reactor were in good agreement with the calculated values. However, this comparison was not possible for HPN and HHN due to greater losses in the sampling line. Although some losses of the organic nitrate are expected to occur to the walls of the flow reactor, for some experiments the walls were conditioned with a flow of the organic nitrate overnight. There was no statistical difference in the partition coefficients for these experiments compared to experiments where the walls were not conditioned.

Incorporation into Impacted Particles

Previously, partition coefficients (K) and uptake coefficients (γ) were measured using ATR-FTIR for uptake of these organic nitrates into impacted particles from AP ozonolysis in the absence of an OH scavenger.⁶⁷ For comparison, the partition coefficients for these organic

nitrates into impacted particles from AP ozonolysis in the presence of CH as an OH scavenger have been measured in the same manner here (Series A). In brief, the polydisperse particles were collected onto a Ge ATR crystal using a custom-designed impactor with a 50% cut-off diameter of 240 nm.⁴¹ This resulted in >60% of the total mass concentration of the particles in the flow reactor being collected by the impactor under the reaction conditions used here. The particles were sampled at a total flow of 30 L min⁻¹ for 10-30 min at the end of the reactor (~20-60 µg total impacted mass), corresponding to a reaction time of 31 min. The amount impacted onto the crystal was varied to ensure the film thickness was smaller than the depth of penetration (d_n) of the infrared evanescent wave, and thus ensuring that the entire film was probed by the IR beam. Using the wavelength of the IR beam and the refractive indices of the Ge crystal and air, the d_p was calculated to be 0.35 µm at 1730 cm⁻¹, 0.37 µm at 1630 cm⁻¹, and 0.47 µm at 1280 cm⁻¹ ^{1.75} The path length (l) of the IR beam through the impacted particles can be estimated using d_p and factoring in the 10 bounces of the beam within the ATR crystal, giving total path lengths of 3.5 µm at 1730 cm⁻¹, 3.7 µm at 1630 cm⁻¹, and 4.7 µm at 1280 cm⁻¹.⁷⁵ After impaction, the Ge crystal was placed in an ATR cell inside an FTIR spectrometer (Nicolet 6700), and the spectrum of the impacted SOA particles was acquired at 4 co-added scans and a resolution of 8 cm⁻¹. The selected organic nitrate was introduced by flowing clean, dry air over the pure liquid and subsequently over the impacted particles at a constant flow of 0.060 ± 0.005 L min⁻¹. The 1280 cm⁻¹ peak was used for analysis of the organic nitrates since there was some overlap of the carbonyl peaks with the 1630 cm⁻¹ peak of the $-ONO_2$ group. The partition coefficients were calculated once the organic nitrate signal had reached steady-state (~1000 seconds) based on the intensity of the infrared peaks for the -ONO₂ symmetric stretch (1280 cm⁻¹)^{48, 76} and the carbonyl stretch (1733 cm⁻¹),⁷⁶ as follows:

$$\mathbf{K} = \frac{[-\mathsf{ONO}_2]_{\mathsf{SOA}}}{[-\mathsf{ONO}_2]_{\mathsf{air}}} \tag{1}$$

In equation (1), $[-ONO_2]_{SOA}$ is the concentration of organic nitrate in the impacted particles in units of moles per liter of SOA and $[-ONO_2]_{air}$ is the concentration of organic nitrate in air in units of moles per liter of air. Note that for Series D when the organic nitrate is present in the flow reactor, $[-ONO_2]_{SOA}$ can include the parent organic nitrate or a product from oxidation by OH. The $[-ONO_2]_{air}$ is the gas phase concentration measured from the trap by GC-MS. The $[-ONO_2]_{SOA}$ was calculated using equations (2), (3) and (4):

$$\frac{\frac{A_{nit}}{l_{nit} \times \sigma_{nit}}}{\frac{A_{C=0}}{l_{C=0} \times \sigma_{C=0}} + \frac{A_{nit}}{l_{nit} \times \sigma_{nit}}} = \frac{n_{nit}}{n_{C=0} + n_{nit}}$$
(2)

$$n_{C=0} \times \frac{N_{sub}}{N_{C=0}} = n_{sub}$$
(3)

$$\frac{n_{nit}}{n_{sub} \times \frac{M_{sub}}{\rho_{sub}} + n_{nit} \times \frac{M_{nit}}{\rho_{nit}}} = [-ONO_2]_{SOA}$$
(4)

In equation (2), A_{nit} and $A_{C=O}$ are the absorbances for the -ONO₂ and the C=O, respectively. The absorption cross-section σ is in units of cm² mole⁻¹ (base 10), *l* is the pathlength of the IR beam through the film in cm at the selected wavenumbers, and n_{nit} and $n_{c=o}$ concentrations are in moles –ONO₂ cm⁻³ and moles C=O cm⁻³ of film, respectively. The $n_{c=o}$ is converted into n_{sub} (moles substrate cm⁻³) in eq. (3) using the number of C=O groups (N_{C=O}) per substrate molecule. This includes any carbonyl, acid, anhydride, or ester functional groups that may be present in α -pinene SOA products. The value of N_{C=O}/N_{sub} is taken to be 2 for this SOA based on the literature.^{77, 78} In equation (4), the number of moles of substrate and nitrate in one cubic centimeter are converted to volume (in units of liters) using the molecular weights (M = 175 g mole⁻¹ for 2EHN; 121 g mole⁻¹ for HPN; 163 g mole⁻¹ for HHN, assuming M = 200 g mole⁻¹ for SOA),⁷⁷⁻⁷⁹ and the densities ($\rho = 9.6 \times 10^2$ g L⁻¹ for 2EHN; 1.2×10^3 g L⁻¹ for HPN; 1.1 × 10³ g L⁻¹ for HHN, and using $\rho = 1.2 \times 10^3$ g L⁻¹ for SOA).^{67, 80} The IR cross sections for all three organic nitrates and the proxy used for SOA, were previously reported.⁶⁷

AMS Measurements

 An HR-ToF-AMS (Aerodyne)⁸¹⁻⁸³ was used to characterize the particles formed in the absence or presence of the gas-phase organic nitrates and in the absence or presence of the OH scavenger. Particles were sampled at a flow rate of ~0.082 L min⁻¹ into the AMS and focused with an aerodynamic lens, vaporized at 600°C, and ionized via electron impact (EI, 70 eV) ionization. The data presented were acquired in V-mode without HEPA-filter dilution. Measurements were taken with a particle filter for each experiment to adjust the isotope ratio of ¹⁵N¹⁴N that interferes with the CHO⁺ fragment using the "Improved-Ambient" method of Canagaratna *et al.*⁸⁴ Data were analyzed using Igor Pro v. 6.3 and 6.37 (Wavemetrics, Inc.) with SQUIRREL (v. 1.57I and 1.62A) and PIKA (v. 1.16I and 1.22A). Elemental analysis was carried out using the default calibration factors for O:C and H:C.

Previous studies have shown that organic nitrates fragment in EI ionization to yield NO⁺ and NO₂⁺ as major fragments,⁸⁵⁻⁸⁹ with small CHNO⁺ or CHN⁺ fragments.^{86, 88} The ratio of NO⁺/NO₂⁺ can be used to differentiate organic nitrates in the particles from inorganic nitrates or nitric acid.^{86, 88, 90} Details on this measurement and the measured ratios (Table S1) are found in the electronic supporting information. The ratios indicate that the organic nitrate functional group remains unreacted once taken up into the particles. This is also supported by FTIR data showing the lack of detectable peaks due to inorganic NO₃⁻ in the infrared spectra (Figure S1) by comparison to NaNO₃ (Fisher Scientific, 99.9%).

To quantify the amount of organic nitrate in the particles and to compare to the FTIR data, the AMS mass concentrations of NO⁺ and NO₂⁺ were expressed as moles -ONO₂ per liter of SOA. Thus, the mass loading (μ g m⁻³) of NO⁺ and NO₂⁺ were converted using eq. (5) into moles m⁻³ air of organic nitrate using the molecular weights of NO⁺ and NO₂⁺ (30 and 46 g mole⁻¹, respectively), assuming that each organic nitrate has only one nitrate group which will give either an NO⁺ or an NO₂⁺ fragment. The mass concentration of SOA (μ g m⁻³ air), represented by HROrg, is converted to volume concentration of SOA (L m⁻³ air) using its density ($\rho = 1.2 \times 10^3$ g L⁻¹)⁸⁰ as shown in eq. (5):

$$\frac{\left|NO^{+} \times \frac{1}{MW_{NO^{+}}}\right| + \left|NO_{2}^{+} \times \frac{1}{MW_{NO_{2}^{+}}}\right|}{HROrg} \times \rho_{SOA} \times \frac{RIE_{Org}}{RIE_{Nit}} = \frac{moles - ONO_{2}}{Volume SOA}$$
(5)

The default value for the relative ionization efficiency (RIE) of organics (1.4) was used for SOA, while an RIE of 1.0 was used for all organic nitrates, assuming their respective ionization efficiency is similar to that of inorganic nitrate as assumed by other researchers.⁵¹ To calculate the partition coefficient, K, the moles RONO₂ per liter of SOA from eq. (5) was divided by the gas phase concentration in the flow reactor in moles -ONO₂ L⁻¹ air (eq. 1). These concentrations were determined by measuring the concentration exiting the trap and factoring in the dilution into the flow reactor.

To examine changes in composition as a function of particle diameter, the high-resolution particle time-of-flight (HR-PToF) feature was used, which allows size-dependent composition analysis of specific fragments. The particle size distribution was separated into 12 evenly spaced bins between 157 and 822 nm D_{va} , and the high-resolution mass spectrum collected up to m/z120. The data for diameters < 157 nm and diameters >822 nm D_{va} were omitted due to the very small mass loading which caused large contributions from noise, and due to decreased lens transmission efficiency in these diameter ranges.⁹¹

For Series D (incorporation *during growth*), the polydisperse particles were impacted at a total flow of 30 L min⁻¹ for 10-30 min at the end of the reactor. The partition coefficient was then quantified using the -ONO₂ and carbonyl stretches as described above in equations 1-4.

Results and Discussion

I. IN THE PRESENCE OF AN OH SCAVENGER, CYCLOHEXANE.

a. Incorporation of organic nitrate after SOA formation and growth. The uptake of the organic nitrates into SOA particles impacted on an ATR crystal was studied by ATR-FTIR as described above (Series A). Figure 3a shows typical ATR-FTIR spectra after equilibrium was achieved (after ~1000 seconds of exposure to the organic nitrate), and Figure 3b shows typical time profiles for the number of -ONO₂ per cm² crystal surface area during uptake and subsequent desorption. The high concentrations of organic nitrates used in Series A are expected to induce a plasticizing effect as previously reported, i.e. lowering the viscosity of the impacted particles and increasing diffusion rates into the particles which allows equilibrium with the gas-phase organic nitrate to be established on a faster timescale.⁶⁷ Partition coefficients (K_A) were calculated to be $(3.2 \pm 1.5) \times 10^4$, $(4.4 \pm 2.0) \times 10^5$ and $(4.9 \pm 0.8) \times 10^6$ for 2EHN, HPN, and HHN, respectively $(\pm 1\sigma)$. The increasing trend from 2EHN to HPN to HHN is not surprising given the more polar nature of the hydroxy nitrates compared to 2EHN. It is noteworthy that the partition coefficient is larger for HPN than for 2EHN by about an order of magnitude, despite their similar vapor pressures (Table 1),⁷²⁻⁷⁴ indicating that vapor pressure alone is not sufficient for predicting SOA growth and composition. This is not surprising in that vapor pressure is a measure of the attractive forces between the same molecules in the liquid, while interactions in the particles are between the organic nitrate and the SOA functional groups.

To establish that the uptake into impacted particles is similar to particles suspended in air, experiments were carried out in which the SOA particles were denuded (to remove the α -pinene gas phase oxidation products), diverted to a mini glass flow tube and subsequently exposed to gas phase 2EHN (Series B). In this experiment, the measured 2EHN concentration (~3 × 10¹⁵ molecules cm⁻³) was similar in magnitude to that used for the impacted particles in Series A. The partition coefficient for Series B was determined using AMS for times of exposure to 2EHN of approximately one to eight minutes. These exposure times encompass the timeframe for 2EHN to reach equilibrium in Series A experiments on impacted particles, which was approximately 2-3 min (Fig. 3b). The average partition coefficient (K_B^{2EHN}) is in excellent agreement with that for Series A (Table 2), confirming that suspended particles come to the same equilibrium as the thin film of impacted particles upon exposure to high concentrations of 2EHN. It also demonstrates that the AMS and the FTIR measurements are in excellent agreement with each other.

The concentration of 2EHN was reduced by removing the reservoir of 2EHN in Fig. 2b and instead introducing the 2EHN into the mini glass flow tube using a trap and flowing clean air over the pure liquid. This diluted the 2EHN by about a factor of 10, giving a concentration of $\sim 3 \times 10^{14}$ molecules cm⁻³, comparable to the concentration used in Series C which will be discussed below. Under these conditions, the organic nitrate signal in the particles became undetectable as would be expected if diffusion into the particles is slow. Indeed, earlier experiments have shown that SOA from the ozonolysis of AP under dry conditions is a high viscosity semi-solid.^{35, 37, 41, 92-96} As discussed previously, exposure to high concentrations of the organic nitrates can have a plasticizing effect.⁶⁷ The undetectable nitrate signal suggests that the concentration of 2EHN was not sufficiently high to produce a plasticizing effect, leaving the SOA particles as a high

viscosity material. Diffusion coefficients for particles from AP ozonolysis formed under dry conditions range from 10⁻¹⁴-10⁻¹⁷ cm² s⁻¹.^{40, 42, 92, 97} Using the Stokes-Einstein relation^{26, 37} and assuming a molecular radius of 1 nm, this results in viscosities ranging from 10⁵-10⁸ Pa s, consistent with measured viscosities for SOA from AP ozonolysis.^{37, 98-102} The resulting characteristic timescale for diffusion²⁶ through a semi-solid 200 nm particle is at least half an hour, much longer than the maximum residence time for Series B of ~8 minutes.

b. Incorporation of organic nitrate during SOA formation and growth. Partition coefficients (K_C) were calculated using the AMS data (eq. 5) for Series C where the organic nitrates were present in the flow reactor while particles were forming and growing. Table 2 summarizes these partition coefficients at 31 min reaction time, which shows the same increasing trend from 2EHN to HPN to HHN seen for the incorporation after growth (Series A).

The results show that the partition coefficients in Series C are much larger than those in Series A, which might seem surprising since they imply a larger than equilibrium concentration in the particles. As described above, the difference observed is not associated with any bias in the two analytical techniques (AMS and ATR-FTIR) that were used. Additionally, Series C used much lower gas phase concentrations than for Series A, so the explanation cannot be a significant plasticizing effect. In support of this, Figure S2 shows typical impaction patterns for SOA formed either with or without an organic nitrate or OH scavenger present. Also shown in Fig. S2 for comparison are the impaction patterns for deliquesced NaSO₄ particles, dry carboxylate-modified latex particles, and SOA particles formed at 87% relative humidity which is known to decrease viscosity.^{41, 99, 100, 103} Upon impaction, particles hit and may stick to the substrate directly below the orifice plate to form spots, form midlines due to multi-orifice interactions, or they may bounce and either be re-entrained into the gas stream or subsequently

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be re-captured on the substrate to form a cloud or halo.¹⁰⁴ The impaction patterns in Fig. S2 are distinctly different from both the deliquesced NaSO₄ and the SOA formed at high relative humidity, indicating the particles in the current study are highly viscous. The patterns do not change across the experimental conditions, suggesting there was no significant change in the viscosity upon addition of the organic nitrates or cyclohexane. One might therefore expect that the uptake into high viscosity semi-solid particles would be hindered; however, our results (Series C) show the opposite, which highlights that the incorporation of organic nitrates is driven by a different phenomenon than diffusion.

When the 2EHN concentration used in Series B was lowered to that used in Series C, the organic nitrate signal became undetectable, consistent with a higher viscosity limiting uptake into and diffusion through the pre-formed and denuded particles.³⁸ K_C values from Series C taken at 7 min and 31 min are not significantly different, and thus the higher partition coefficients in Series C are not resulting from the longer time spent in the flow reactor. The presence of the gas-phase ozonolysis products in the large flow reactor in Series C must therefore play a central role in the incorporation of the organic nitrates *during* particle formation and growth that enhances organic nitrate uptake beyond the expected equilibrium established in Series A/B.

II. IN THE ABSENCE OF AN OH SCAVENGER.

a. Incorporation of organic nitrate after SOA formation and growth. The incorporation of the three organic nitrates into pre-formed SOA particles impacted on an ATR crystal was previously studied for SOA formed without an OH scavenger.⁶⁷ In order to make a direct comparison with the present Series A data (pre-formed SOA particles formed with an OH scavenger), those partition coefficients,⁶⁷ which were based solely on the vapor pressures, have been adjusted here

to reflect our measured concentrations. Both sets of partition coefficients are reported in Table 3. In these measurements, OH reaction with the organic nitrates does not occur since the particles are pre-formed and impacted, and thus the uptake is that of the unoxidized parent organic nitrate (2EHN, HPN or HHN).

 For HPN, the partition coefficient on impacted particles formed without CH is not statistically different from that measured on impacted particles formed in the presence of CH, and is in excellent agreement with the air-octanol partition coefficient of $(4.2 \pm 0.3) \times 10^5$ reported by Treves *et al.*¹⁰⁵ This suggests that the magnitude of the HPN partitioning into the particles is unaffected by any composition changes in the SOA resulting from changes in the chemistry in the presence of the OH scavenger. It is interesting that the partition coefficients into SOA and octanol are similar and both larger than into water,^{106, 107} suggesting that even with the hydrogen bonding possibility to the alcohol group of HPN, dispersion forces between HPN and SOA are important as well.

For 2EHN and HHN, the partition coefficients decrease when the SOA is pre-formed in the presence of the OH scavenger by factors of ~3 and ~2, respectively. This reduction in partitioning indicates decreased solubility of 2EHN or HHN into the bulk of the film, which could reflect differences in the SOA composition when CH is added to scavenge the OH. Table S2 shows the O:C and H:C ratios and the oxidation state of carbon (OS_c) determined by AMS.¹⁰⁸ Only very small changes in the bulk elemental composition were exhibited, consistent with previous reports for AP ozonolysis SOA formed with or without an OH scavenger.¹⁰⁹ However, this may simply reflect that functional group changes important in determining solubility are not detected in these bulk measurements. The percent change of a few major fragments by AMS when OH scavenger is present (Figure S3) shows that there are some changes in the SOA bulk

composition. Why this results in changes in the partition coefficients for 2EHN and HHN, and not for HPN, is not clear, but illustrates the need for a detailed molecular level understanding of the composition of SOA.^{110, 111}

b. Incorporation of organic nitrate during SOA formation and growth. Figure 4a shows the ATR-FTIR spectra for impacted particles from AP ozonolysis alone (without organic nitrate or OH scavenger) and for particles formed in the presence of 2EHN, HPN or HHN (Series D, no CH). Figure 4b shows the ATR-FTIR spectra for particles formed in the presence of an OH scavenger (Series D, with CH). Comparison of the SOA itself shows some change in the -CH region (3000-2800 cm⁻¹) in the presence of CH. Although CH is very volatile and unlikely to partition into the particle phase, some of its OH oxidation products may be taken up and contribute to the particle growth. Figure S4 shows the ATR-FTIR spectra for cyclohexanone and cyclohexanol which are among the expected products from the CH + OH reaction, ¹¹²⁻¹¹⁴ showing that the CH + OH products may be contributing to the changes seen in the particles formed in the presence of CH. While the organic nitrate peaks at 1630 cm⁻¹ and 1280 cm⁻¹ are seen in the spectrum when 2EHN is present during particle formation, there is much less organic nitrate after the addition of the OH scavenger. Similar results were obtained for SOA formed in the presence of HHN. For HPN, the organic nitrate signal was below the limit of detection by FTIR either with or without the OH scavenger.

When the gas-phase organic nitrate is present in the flow reactor in the absence of an OH scavenger (Series C and D, no CH), the organic nitrates can react in the gas phase with the OH radical generated in the ozonolysis. The calculated OH rate constants for 2EHN and HHN are similar, and are a factor of approximately three larger than that for HPN (Table S3).¹¹⁵ This is due to the long alkyl chains in 2EHN and HHN which provide a number of potential sites for

hydrogen abstraction by OH. Many of the products formed from OH oxidation of 2EHN and HHN are expected to be more functionalized and have lower volatility than the parent organic nitrate. Scheme 1 shows some of the likely routes for oxidation of those organic nitrates. Thus, the combination of higher rate constants and lower volatility OH oxidation products are such that the organic nitrate-OH oxidation products for 2EHN and HHN can contribute significantly to the nitrate content of the SOA in the absence of an OH scavenger. As a result, suppressing OH by the addition of cyclohexane significantly lowers the formation of the OH oxidation products of the organic nitrates.

The effect is more dramatic for 2EHN because its concentration in the experiments is about 28 times greater than that of HHN so that 2EHN itself is a more efficient OH scavenger. While HPN can also react with OH, oxidation at the tertiary carbon of the molecule is most likely and will lead to smaller, more volatile species which may not partition significantly into the particles (Scheme 1).

The trends seen in the organic nitrate signal measured by FTIR (Series D) in the presence or absence of the OH scavenger are also supported by the AMS data (Series C). Figure 5 shows the number of moles RONO₂ taken up per liter of SOA calculated from the AMS data (eq. 5) at 31 min reaction time (there was no statistical difference between 7 min and 31 min reaction time). Consistent with the FTIR data, the signal for 2EHN increases significantly (by a factor of ~14) when the SOA is formed in the absence of the OH scavenger. This factor is larger than that of Series A, indicating that it is the oxidation products of 2EHN that are incorporated into the SOA, and is not simply due to a difference in the SOA bulk composition. A similar but smaller trend is seen for HHN (a factor of ~2.5), whereas the amount of HPN in the particles is unaffected by the presence of the OH scavenger. However, it should be noted that the detection

 methods used here (FTIR and AMS) focus on functional group analysis and are not direct measurements of the parent organic nitrate.

The much smaller effect seen for HHN is due to the smaller gas phase concentrations that were able to be added to the flow system, which results in HHN not competing very effectively with the α -pinene for the OH radical generated in the ozonolysis. Thus, the initial first order rates of loss of OH, estimated using k[X]₀, where X = AP or RONO₂, are 3.3 × 10² s⁻¹ for AP and 8.8 × 10² s⁻¹ for 2EHN under the conditions shown in Fig. 5 (the rate for AP decreases with time due to reaction with O₃). As a result, 2EHN competes with AP for OH, forming oxidized alkyl nitrate products that are incorporated into the SOA. On the other hand, k[X]₀ for HHN is only 27 s⁻¹ so that relatively small amounts are oxidized by OH whose removal is now mainly via reaction with AP, and incorporation of the unoxidized parent HHN contributes relatively more than its oxidation products to SOA growth. This is consistent with the magnitude of change in the incorporation of HHN into SOA formed with versus without CH being similar for Series A (uptake into preformed SOA) and Series C/D (incorporation during SOA formation). While the first order loss rate for HPN by OH is $1.9 \times 10^2 \text{ s}^{-1}$, its oxidation products are sufficiently small that they will not be efficiently incorporated into the SOA.

The contribution of OH radical chemistry is also manifested in the decrease in the particle number concentration in the presence of the OH scavenger, and in the presence of the organic nitrates which can also scavenge OH (Fig. 6). Note that the total particle number concentration is smaller in the presence of CH (Fig. 6a). This is consistent with previously reported work on the impact of the OH chemistry on SOA formation.^{116, 117} For example, Berndt *et al.*¹¹⁸ showed that the OH radical plays an important role in the formation of highly oxidized multifunctional organic compounds (HOMs), and thus scavenging the OH suppresses the formation of these

HOMs, lowering SOA yields. Figure 6b, 6d, and 6f show the particle size distributions when organic nitrates are present during SOA formation without CH. A decrease in SOA is seen in the presence of 2EHN in a manner that is qualitatively similar to that due to addition of CH alone (Fig. 6a), while there is little impact for HPN, and no impact at all for HHN. This is consistent with the relative rate constants for OH reaction with the organic nitrates versus CH (Table S3) and the initial concentrations of these compounds. As described above, 2EHN competes with AP for OH, but HHN does not. The rate of loss of OH by CH is 1.8×10^4 s⁻¹, which overwhelms the reaction of OH with AP or the organic nitrates. Based on kinetic modeling using Kintecus^{®119} and a simplified reaction scheme (Table S3), less than 0.1% of the organic nitrates reacts with OH after 31 min reaction time when 100 ppm CH is present. Additional experiments were done for 2EHN using 500 ppm CH, and there was no statistical difference in the amount of 2EHN incorporated, indicating that 100 ppm CH is enough to adequately scavenge the OH. At 31 min reaction time without CH present approximately 1.3% of the 2EHN has reacted with OH, compared to ~0.8% of the HPN and ~4.3% of the HHN. This is consistent with the trend in the impacts on SOA, given the much lower initial gas phase concentration of HHN compared to the other organic nitrates.

In the absence of an OH scavenger, partition coefficients for 2EHN and HHN cannot be reliably quantified due to contributions from the $RONO_2 + OH$ oxidation products whose identity and gas phase concentrations are not known. Instead, the concentrations of $RONO_2$ in SOA (moles of $RONO_2$ per liter of SOA) were measured using both AMS (Fig. 5) and FTIR as described in the experimental (Series C and D). The values for 2EHN and HHN are summarized in Table 4, showing the two methods are in agreement within 50%. Only the AMS value (Series C) is reported for HPN, as it was below the detection limit by FTIR.

 In short, OH oxidation of these organic nitrates forms more oxidized organic nitrates that, in the case of 2EHN and HHN, partition to a greater extent into the SOA The rate constant for OH with HPN is smaller than for 2EHN and HHN, and in addition, its oxidation is expected to lead to smaller, higher volatility products, which will not be readily taken up into SOA (Scheme 1).

Physical Mechanism for Particle Growth

As described previously, the growth of the SOA particles by ozonolysis products and the incorporation of the organic nitrates into the particles in the stainless steel flow reactor may be best described by a kinetically limited "burying" mechanism. Figure 7 shows a simplified schematic that describes this mechanism, where RONO₂ represents the organic nitrate, and P1 is a proxy low volatility organic from the ozonolysis of α -pinene. When the organic nitrate is present *during particle growth* in the flow reactor (Series C/D), condensing P1 molecules can bury the organic nitrate and hinder re-evaporation into the gas phase, resulting in larger partition coefficients than those measured after particle growth at equilibrium (Series A/B).

Figure 8 shows the HR-PToF data for both the total organic and the ratio of (NO⁺ + NO_2^+) to total organics as a function of particle size for SOA formed in the presence of 2EHN at 7 min reaction time, either with or without CH. The ratio (NO⁺ + NO_2^+)/HROrg is a measure of the relative concentrations of organic nitrate in the SOA. This ratio is approximately constant across the range of particle sizes, confirming that relative rates of incorporation of the organic nitrates and the organics that grow the particles do not vary significantly as the particles grow across this diameter range. The corresponding data for SOA formed in the presence of HPN and HHN are found in Figure S5 and S6, respectively, and show similar results. For comparison, Fig. S7 shows HR-PToF analysis for two major fragments, $C_2H_3O^+$ (a marker for carbonyl

groups) and CO_2^+ (a marker for carboxylic acid groups),^{120, 121} and the ratio of these two fragments. In contrast to the uniform organic nitrate composition over all particle diameters, smaller diameter SOA particles are composed of more acid groups on average than the larger particles, either in the absence or presence of OH. The combination of data suggests that while condensation of low volatility organics onto particles leads to non-uniform particle composition, organic nitrates can become physically buried during particle growth.

Conclusions

Uptake of organic nitrate tracers into highly viscous, semi-solid SOA particles during their formation in the ozonolysis of AP offers new insights into the molecular interactions between gases and particles that ultimately lead to particle growth. The role of the OH radical resulting from the ozonolysis reaction is important not only for the bulk composition of the particles and overall SOA number concentration, but also plays an important role in the gasphase chemistry of the organic nitrates. In the case of the smallest organic nitrate, HPN, the partitioning was unaffected both by the reaction of HPN with OH and by any changes in the particle composition by scavenging OH. However, the two long-chain organic nitrates were affected by both of these factors, with the effect being most pronounced for the 8-carbon 2EHN.

The amount of organic nitrate taken up into growing particles relative to the gas phase concentration was found to be larger than expected based on the equilibrium partition coefficients into pre-existing impacted particles. This may be attributed to the evolution of particles during growth, such that continued uptake of organics leads to 'burying' of the organic nitrate, hindering re-evaporation into the gas phase. This is consistent with the HR-PToF analysis which shows that the organic nitrate was evenly distributed across all particle diameters. This could play a role in cases where mechanisms in addition to thermodynamic partitioning

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 have been implicated.^{34, 60, 68, 122} The results of these studies highlight the importance of a molecular level understanding of the interactions of gases with particle surfaces and their bulk as the foundation for accurately predicting their impacts on air quality and climate.

Conflicts of Interest

There are no conflicts to declare.

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Table 1: Gas phase concentrations measured at the exit of the trap containing the pure liquids^a and estimated vapor pressures using two group contribution methods⁷²⁻⁷⁴ for 2EHN, HPN and HHN.

Organic Nitrates	Gas Phase Concentration at Trap Exit (10 ¹⁵ molecules cm ⁻³ and Pa) ^b	Vapor Pressure using Moller ^{73, 74} (Pa)	Vapor Pressure ^c using SIMPOL.1 ⁷² (Pa)
2EHN	4.7 ± 0.2	14	18
	$(19 \pm 1.0 \text{ Pa})$		
HPN	3.9 ± 0.2	12 ^d	16
	$(16 \pm 1.0 \text{ Pa})$	<u>35</u> ^e	
		Average: $f 24 \pm 16$	
HHN	0.17 ± 0.05	0.35 ^d	0.85
	$(0.70 \pm 0.21 \text{ Pa})$	<u>0.65</u> ^e	
		Average: f 0.50 ± 0.21	

^aMeasured concentrations are from the average of triplicate measurements.

^bError bars are $\pm 1\sigma$ from the average of triplicate measurements

^cSIMPOL.1 does not distinguish between isomers

^dHydroxy-terminated isomer

^eNitrate-terminated isomer

^fError bars are $\pm 1\sigma$

Table 2: Comparison of partition coefficients into SOA formed in the presence of an OH

scavenger when the organic nitrates are incorporated after growth (Series A, K_A, and Series B,

K_B), or *during growth* (Series C, K_C).

Organic Nitrate	K _A (Series A) ^{a,b}	K _B (Series B) ^{c,d}	K _C (Series C) ^{b,c,e}	Ratio K _C /K _A
2EHN	$(3.2 \pm 1.5) \times 10^4$	$(2.9 \pm 0.7) \times 10^4$	$(4.7 \pm 1.0) \times 10^5$	15 ± 7.6
HPN	$(4.4 \pm 2.0) \times 10^5$	n/a ^f	$(1.7 \pm 0.2) \times 10^{6}$	3.9 ± 1.8
HHN	$(4.9\pm0.8)\times10^6$	n/a ^f	$(1.6\pm0.3)\times10^7$	3.3 ± 0.81

^aUsing ATR-FTIR.

^bError bars are $\pm 1\sigma$ from the average of three experiments.

^cUsing AMS.

^dError bars are $\pm 1\sigma$ from the average of ~1-8 minutes exposure time

^eValues are taken at 31 minutes reaction time in the stainless steel flow reactor ^fSeries B was only done for 2EHN.

Table 3: Partition coefficients (K_A) calculated for the uptake of the organic nitrates into impacted particles formed either with or without CH (Series A).

Organic Nitrate	Partition Coefficient (K _A) ^a for SOA with CH	Partition Coefficient (K _A) ^{a,b} for SOA without CH ⁶⁷
2EHN	$(3.2 \pm 1.5) \times 10^4$	$(1.1 \pm 0.1) \times 10^5$
HPN	$(4.4 \pm 2.0) \times 10^5$	$(5.4 \pm 2.0) \times 10^5$
HHN	$(4.9\pm0.8)\times10^6$	$(9.0 \pm 1.0) \times 10^{6}$

^a Error bars are $\pm 1\sigma$ from the average of three experiments.

^bThese values have been adjusted from previous concentrations using estimated vapor pressures to the newly measured gas phase concentrations in this study.

Table 4: Organic nitrate content (moles L⁻¹ SOA) for particles formed in the presence of 2EHN, HHN, or HPN, in the absence of an OH scavenger (Series C and D, no CH).

	RONO ₂ content (Moles L ⁻¹ SOA) ^a Series C (AMS)	RONO ₂ content (Moles L ⁻¹ SOA) ^{a,b} Series D (FTIR)
2EHN	1.6 ± 0.3	1.6 ± 0.2
HPN	0.29 ± 0.03	n/a ^c
HHN	0.34 ± 0.05	0.50 ± 0.05

^aError bars are $\pm 2\sigma$ from the average of three experiments.

^bFTIR quantification used the absorption cross section of the parent organic nitrate.⁶⁷

^cHPN was below the detection limit for FTIR.



Scheme 1: Simplified reaction scheme showing some pathways for OH oxidation of 2EHN, HPN and HHN. The vapor pressures for each compound were estimated using SIMPOL.1.⁷² Hydroxyl radical attack on a secondary carbon is shown for 2EHN and HHN due to the higher relative contribution of the sum of the secondary carbons to the total OH rate constant compared to that of the one tertiary carbon.

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Figure 1: Structures of the gas phase organic nitrates used in this study. The synthesis of the hydroxynitrates resulted in the presence of the two isomers.



Figure 2: Schematic of the four experiment types. In series A, SOA particles are formed in the absence of organic nitrates (RONO₂) in the stainless steel flow reactor and impacted on an ATR crystal to generate a thin film of particles, over which RONO₂ at near saturation vapor pressures $(4.7 \times 10^{15} \text{ (2EHN)}, 3.9 \times 10^{15} \text{ (HPN)} \text{ and } 1.7 \times 10^{14} \text{ (HHN)} \text{ molecules cm}^{-3})$ were flowed and the uptake of RONO₂ was measured by ATR-FTIR. In series B, SOA particles were generated in the stainless steel flow reactor and then passed through a charcoal denuder before subsequently being flowed into a 1-m long glass flow tube and exposed to 2EHN through either a reservoir with a pure liquid or a trap flowing air over the pure liquid, and analyzed by HR-ToF-AMS. In series C, RONO₂ at lower levels $(1.4 \times 10^{14} (2\text{EHN}), 1.2 \times 10^{14} (\text{HPN}))$ and 5.0×10^{12} (HHN) molecules cm⁻³) were incorporated into SOA particles as they form and grow in the stainless steel flow reactor. These particles were quantified by HR-ToF-AMS. In series D, the same particles as in Series C were impacted on an ATR crystal and partition coefficients were determined from the quantification of RONO₂ by ATR-FTIR. For Series A, the gas-phase organic nitrate concentration was measured using GC-MS from the trap. For Series B, the gasphase 2EHN concentration was measured directly from the mini flow tube. For Series C/D, the gas-phase organic nitrate concentration is calculated from the measured concentration exiting the trap and accounting for dilution (a factor of 34).





Figure 3: a) ATR-FTIR spectra of the impacted particles formed in the presence of an OH scavenger, after exposure to 2EHN (4.7×10^{15} molecule cm⁻³, red), HPN (3.9×10^{15} molecule cm⁻³, blue), or HHN (1.7×10^{14} molecule cm⁻³, green) after equilibrium was reached for all organic nitrates (~1000 seconds) (Series A). Also shown is the spectrum of SOA alone (black). The inset shows an expanded view of the 1280 cm⁻¹ peak characteristic of the -ONO₂ stretch. The region between 2500 – 2000 cm⁻¹ is not shown due to variations in the CO₂ in the sampling compartment. b) Concentrations of the {-ONO₂} functional group in molecules per cm² after exposure of impacted particles (with total impacted mass of ~30 µg) to the gaseous organic nitrates, and subsequent desorption by exposure to clean, dry air. The dashed black line indicates the experimentally-determined limit of detection for the nitrates. Solid lines are fits to guide the eye, and error bars are ± 2 σ determined from the uncertainty in the measured absorption cross sections of 2EHN, HPN and HHN.



Figure 4: ATR-FTIR spectra for a) impacted particles alone and formed in the presence of 2EHN (1.4×10^{14} molecules cm⁻³), HPN (1.2×10^{14} molecules cm⁻³), or HHN (5.0×10^{12} molecules cm⁻³) in the stainless steel flow reactor (Series D) without an OH scavenger, and b) impacted particles alone and formed in the presence of 2EHN, HPN or HHN at the same concentrations in the stainless steel flow reactor in the presence of CH as an OH scavenger (Series D). The region between 2500 - 2000 cm⁻¹ is not shown due to variations in the CO₂ in the sampling compartment.



Figure 5: Quantification of moles of organic nitrate per liter of SOA (moles RONO₂ L⁻¹ SOA) for SOA formed in the presence of 2EHN (1.4×10^{14} molecules cm⁻³), HPN (1.2×10^{14} molecules cm⁻³), or HHN (5.0×10^{12} molecules cm⁻³), with or without OH scavenger at 31 minutes reaction time (Series C). Error bars are $\pm 2\sigma$ from the average of three experiments.



Figure 6: Particle number distributions (# cm⁻³) at 31 minutes reaction time for a) SOA alone and formed in the presence of CH as an OH scavenger, b) SOA alone and formed in the presence of 2EHN (1.4×10^{14} molecules cm⁻³), c) SOA formed in the presence of 2EHN either with or without CH, d) SOA alone and formed in the presence of HPN (1.2×10^{14} molecules cm⁻³), e) SOA formed in the presence of HPN either with or without CH, f) SOA alone and formed in the presence of HHN (5.0×10^{12} molecules cm⁻³), and g) SOA formed in the presence of HHN either with or without CH. Error bars are $\pm 1\sigma$ from the average of three scans, and solid lines are best fit distributions to guide the eye.



SOA growth and incorporation of organic nitrates

Figure 7: Schematic of "burying" mechanism for incorporation of organic nitrate tracers as SOA particles are forming. RONO₂ represents the organic nitrate, and P1 is a proxy for low volatility organics from the ozonolysis of α -pinene. When the organic nitrate is present during particle growth by P1, the P1 can bury the organic nitrate and hinder re-evaporation into the gas phase.



Figure 8: The HR-PToF mass distribution of total HROrg (green) and the HR-PToF mass ratio of $(NO^+ + NO_2^+)$ to HROrg (red) for a) SOA formed in the presence of 2EHN $(1.4 \times 10^{14} \text{ molecules cm}^{-3})$ at 7 min reaction time, and b) SOA formed in the presence of 2EHN and CH $(2.5 \times 10^{15} \text{ molecules cm}^{-3})$ at 7 min reaction time. Note the NO⁺ and NO₂⁺ signals in the presence of CH have high uncertainty due to weak signal.



TOC

The incorporation of organic nitrates into viscous secondary organic aerosol during particle formation is enhanced relative to expected equilibrium partitioning, and is best described by a kinetically controlled "burying" mechanism.