Nonequilibrium atmospheric secondary organic aerosol formation and growth

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Airborne particles play critical roles in air quality, health effects, visibility, and climate. Secondary organic aerosols (SOA) formed from oxidation of organic gases such as α-pinene account for a significant portion of total airborne particle mass. Current atmospheric models typically incorporate the assumption that SOA mass is a liquid into which semivolatile organic compounds undergo instantaneous equilibrium partitioning to grow the particles into the size range important for light scattering and cloud condensation nuclei activity. We report studies of particles from the oxidation of α-pinene by ozone and NO3 radicals at room temperature. SOA is primarily formed from low-volatility ozonolysis products, with a small contribution from higher volatility organic nitrates from the NO3 reaction. Contrary to expectations, the particulate nitrate concentration is not consistent with equilibrium partitioning between the gas phase and a liquid particle. Rather the fraction of organic nitrates in the particles is only explained by irreversible, kinetically determined uptake of the nitrates on existing particles, with an uptake coefficient that is 1.6% of that for the ozonolysis products. If the nonequilibrium particle formation and growth observed in this atmospherically important system is a general phenomenon in the atmosphere, aerosol models may need to be reformulated. The reformulation of aerosol models could impact the predicted evolution of SOA in the atmosphere both outdoors and indoors, its role in heterogeneous chemistry, its projected impacts on air quality, visibility, and climate, and hence the development of reliable control strategies.

Results and Discussion

Airborne particles are well-known to negatively affect human health (1) and to contribute to "haze" associated with urban and regional pollution, leading to a reduction in visibility (2). On a global scale, airborne particles scatter solar radiation and can act as cloud condensation (CCN) and ice nuclei (IN), influencing the radiative balance of the atmosphere (3, 4). Currently these effects represent the largest uncertainty in calculations of climate change (5). A major component of atmospheric particles is secondary organic aerosol (SOA) formed via the oxidation of gaseous anthropogenic and biogenic precursor compounds. The SOA material is formed from low-volatility oxidation products (3, 4). However, the processes and species leading to SOA formation and growth are not fully understood, which precludes reliable quantitative predictions of their impacts on climate, visibility, and human health.

Regional and global chemical models have generally underpredicted SOA concentrations compared to those from field measurements (6–9). Inclusion of a number of additional factors such as new SOA precursors, condensed phase chemistry, updated gas-phase chemistry and SOA yields, new primary semivolatile and intermediate volatility species, and improved emissions inventories of both gases and primary organic aerosols have lessened the magnitude of the disagreement (10–18). However, there is still significant uncertainty in predicting ambient SOA levels, and model-measurement discrepancies of a factor of two or more remain common.

One possible source of this uncertainty is that current models typically assume instantaneous equilibrium partitioning of semivolatile organic compounds (SVOCs) between existing liquid airborne particles and the gas phase using the theory of absorptive, activity coefficient-corrected, gas/liquid partitioning described in detail by Pankow (19, 20). Equilibrium partitioning is justified for particles with viscosities in the range of 0.01–100 Pa·s and diffusion coefficients ranging from 10−5 to 10−9 cm²·s⁻¹ (21). A volatility basis set approach has recently been developed for representing the partitioning of SVOCs, which also depends on saturation mass concentration of liquid particles (22).

In this paper, we present laboratory studies of particles formed in the simultaneous oxidation of α-pinene by ozone and NO3 radicals using an aerosol flow system (23). Quantification of the organic nitrate contributions to the SOA provides unique insight into the mechanisms by which particles form and grow, which has important implications for model formulations of SOA, both outdoors and indoors, and the associated impacts predicted based on the model outputs.


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(\(K_{NO2} = 6.2 \times 10^{-12} \text{ cm}^3\text{ molecule}^{-1}\text{ s}^{-1}\); ref. 24) so that the reaction is >90% complete in terms of \(\alpha\)-pinene loss by port 1 (Table S1). This loss is confirmed by experimental measurements of \(\alpha\)-pinene by GC-MS. The validity of the kinetics model is also verified by using the model-predicted concentrations of NO\(_3\) and N\(_2\)O\(_5\) (Fig. S1) to predict how much organic nitrate should be formed and by comparing these concentrations to those measured using FTIR (Fig. S2).

The size distributions of SOA formed by the oxidation of \(\alpha\)-pinene are shown in Fig. 1 (and Fig. S3) when NO\(_2\) is varied from 6.3 to 0 ppm at constant O\(_3\). At the highest NO\(_2\) concentration, few particles are formed, which is expected because multifunctional organic nitrates, formed in the NO\(_3\) reaction with \(\alpha\)-pinene, have sufficiently high vapor pressures that they do not readily nucleate to form new particles. As the NO\(_2\) have vapor pressures between 2 \(\times\) \(10^{-6}\) and \(1 \times 10^{-7}\) atm at 295 K (26, 31). However, ozonolysis products typically have lower vapor pressures (e.g., \(26, 31\)). However, ozonolysis products typically have lower vapor pressures (e.g., \(ca. 1 \times 10^{-7}\) and \(6 \times 10^{-10}\) atm for pinonic and pinic acids, respectively; ref. 31) and more readily undergo homogeneous nucleation to form new particles. As the NO\(_2\) concentration decreases and the contribution of O\(_3\) to the \(\alpha\)-pinene loss increases, the particle number and mass concentration due to ozonolysis increase (Fig. 1 and Fig. S4) as expected.

Organic nitrate products from the NO\(_3\) chemistry comprise a fraction of the SOA. For example, Fig. 2 shows FTIR spectra from particles collected at the end of the flow tube as a function of initial NO\(_2\) concentration. In addition to the aliphatic –C–H stretches in the 2,800–3,000 cm\(^{-1}\) region, three infrared bands corresponding to the –ONO\(_2\) asymmetric stretch (1,630 cm\(^{-1}\)), the –ONO\(_2\) symmetric stretch (1,280 cm\(^{-1}\)), and RO–NO\(_2\) stretch (860 cm\(^{-1}\)) characteristic of organic nitrates are seen (27, 29, 32–34). As described in the SI Text, these spectra show that the relative number of –ONO\(_2\) groups to –C–H groups, \(n_{\text{–ONO}_2}/n_{\text{–CH}}\), increases with the NO\(_2\) concentration (Table S2).

Similarly, mass spectra acquired using real-time high-resolution time-of-flight mass spectrometry (HR-TOF-AMS) and single particle mass spectrometry (SPLAT-II) show that the SOA contains organic nitrates (Fig. S5). The spectra when NO\(_2\) is present (Fig. S5 A–C and E–G) are very similar to those for the ozonolysis reaction (Fig. S5 D and H), except that peaks at \(m/z\) 30 and 46 also appear, primarily from NO\(_3^+\) and NO\(_2NO_2^+\) (SPLAT-II spectra show primarily \(m/z\) 30). The NO\(_3^+\)/NO\(_2NO_2^+\) ratio from the high-resolution AMS mass spectra ranges from 4.9 to 7.6, consistent with a contribution from organic nitrate products (35–38) from the NO\(_2\) reaction with \(\alpha\)-pinene (26, 27, 35).

The similarity of the spectra in the presence and absence of NO\(_2\) suggests that the overall bulk composition of the aerosol (excluding the organic nitrate component) remains relatively constant. For example, the O:C ratios calculated from the AMS data (Fig. S6) average 0.34 ± 0.10 over the range of experimental conditions. This average value is similar to previous measurements of SOA from \(\alpha\)-pinene ozonolysis (39–41) and with pinonic and pinic acid (with O:C ratios of 0.3 and 0.4, respectively), which are known to be major products of the ozone reaction (42, 43). Although there may be a slight trend in the O:C ratio with the NO\(_2\) concentration, it is not significant within uncertainty of the measurements.

Extracts of particles collected on quartz-fiber filters are analyzed by liquid chromatography with UV detection (LC-UV). Chromatograms (Fig. S7) show a group of overlapping peaks with UV spectra attributable to organic nitrates (44) in all NO\(_2\) experiments. The mass concentrations of total organic nitrates in the particles derived from the LC-UV data (\(F_p\), {\(\mu\) g per m\(^3\) of air) (see Fig. S5) are shown in Table 1 as a function of the initial NO\(_2\) concentration. Using the size distributions and the densities measured by SPLAT-II, the total mass concentration of particles and the mass fraction of organic nitrates (\(f_{\text{organic}}\)) are also calculated. Organic nitrates comprise between 0.1–8.2% of the total SOA mass and follow the trend in NO\(_2\). This trend is also shown in the HR-TOF-AMS and FTIR analysis (Table S2).

In short, measurement of the SOA composition by three different techniques shows that the organic nitrate contribution to the SOA decreases as the NO\(_2\) concentration (and thus the available NO\(_3\) radicals to react with \(\alpha\)-pinene) decreases. Simultaneously, the amount of SOA formed increases due to the increasing contribution of the O\(_3\) reaction. Fig. 3 summarizes qualitatively the chemistry occurring in this system: Although ozonolysis is primarily
responsible for SOA formation, ozonolysis products (hereafter defined as “Prod1”) and organic nitrates (hereafter defined as “APONO2”) are both incorporated into the SOA, contributing to its growth. To probe the chemistry quantitatively, a simplified 96-step mechanism for the NO2 + O3 + α-pinene system is developed, and the rate equations are integrated using Acuchem (45). Two reaction channels are assumed for the NO3 reaction and two for the O3 reaction:

\[
\begin{align*}
\alpha\text{-pinene} + NO_3 & \rightarrow APONO_2 \\
& \quad \leftarrow other \text{ products} \\
& \quad k_1 = 1.0 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \\
\alpha\text{-pinene} + O_3 & \rightarrow OH + RO_2 \\
& \quad \leftarrow Prod1 \\
& \quad k_4 = 1.6 \times 10^{-17} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}. \\
\end{align*}
\]

The rate constants for these reactions are based on known rate constants for the overall reactions (24) and branching ratios for the formation of organic nitrates (APONO2) and OH radicals. Thus, the yield of organic nitrates in the NO3 reaction is taken to be 16.5% (46) and the OH yield in the ozone reaction is taken to be 82% (46). Prod1 is a marker for the O3 chemistry that represents, in part, low-volatility products that lead to particle formation and growth. The use of a first-generation product (Prod1) as a proxy for SOA is supported by the work of Ng et al. (47) who showed that the particles from α-pinene ozonolysis are due to first-generation low-volatility products. Fig. S2 compares the box model-predicted total (gas plus particle phase) concentrations of APONO2 as a function of the initial NO3 (gray dashed line) to the gas-phase organic nitrate concentrations measured using long-path FTIR (black circles). This comparison should be reasonable because most of the organic nitrates are in the gas phase (Table 1). As described in the SI Text, the model well represents the time evolution of the total concentration of organic nitrates as a function of the initial NO3 concentration.

Growth of SOA has often been described via adsorption of SVOCs, such as organic nitrates, onto the particle surface or absorption into the bulk. In most atmospheric models of SOA, the latter process (19, 20) is taken to be responsible for uptake of SVOCs into existing liquid particles. It is assumed that once a seed particle is formed (e.g., by homogeneous nucleation), SVOCs will partition into the bulk liquid phase of the particles, causing it to grow.

An equilibrium partitioning coefficient \( K_{p,i} \) for compound \( i \) is defined as the ratio of the concentrations of compound \( i \) in the gas and particle phases, assuming a reversible gas-particle partitioning based on Raoult’s law with activity corrections (19, 20):

\[
K_{p,i} = \frac{F_i/M}{A_i} = \frac{f_{om,RT}}{MW_{\text{om}}10^6 \xi_i P_{li}^{\text{sat}}}. 
\]

In Eq. 5, \( F_i \) and \( A_i \) are the mass concentrations (μg per m3 of air) of compound \( i \) in the aerosol and gas phase, respectively, and \( M \) (μg per m3 of air) is the mass concentration of particulate material, so \( K_{p,i} \) has units of m3 per μg. On the right-hand side of Eq. 5, \( f_{om} \) is the mass fraction of the presumably largely organic material portion of the particulate matter into which the partitioning is occurring (because of the absence of preexisting seed particles, \( f_{om} \) is unity for our experiments), \( R \) is the ideal gas constant (m3 atm K\(^{-1}\) mol\(^{-1}\)), \( T \) is the temperature (K), \( MW_{\text{om}} \) (g mol\(^{-1}\)) is the average molecular weight of the particulate matter phase into which the partitioning is occurring, \( \xi_i \) is the activity coefficient of compound \( i \), and \( P_{li}^{\text{sat}} \) is the vapor pressure of compound \( i \) (atm).

Values of \( F_i/M \) and \( A_i \) for the sum of all organic nitrates are calculated using \( F_i \) and \( M \) from Table 1, and the corrected gas-

![Fig. 3. Representation of SOA formation and growth based on the experimental observations. See text for details.](image-url)
phase concentrations \( (A_i; \text{see SI Text}) \) from the box model with \( A_i = [\text{APONO2}] - F_i \). Eq. 5 predicts that a plot of \( F_i/M \) versus \( A_i \) will be a straight line with slope \( K_p \), which is seen in Fig. 4 (black circles) not to be the case for the \( NO_2/O_3/\alpha\)-pinene system.

There are four possible sources of this unexpected variation that are associated with the physical description of \( K_p \) (right-hand side of Eq. 5): (i) The activity coefficients (\( \gamma_i \)) vary systematically as the \( NO_2 \) concentration changes; (ii) the gas-phase box model does not predict accurately \( A_i \) as a function of \( NO_2 \) concentration; (iii) the composition of the SOA into which the organic nitrates partition changes dramatically with \( NO_2 \) concentration; or (iv) the conditions for which Eq. 5 is developed do not apply in this system. As shown in Fig. S5 and discussed in the SI Text, a systematic variation in the bulk SOA composition (excluding the organic nitrates) and hence in \( \gamma_i \) is unlikely. With respect to (ii), experimental measurements of gas-phase organic nitrates show that the box model does predict accurately their trend with \( NO_2 \) (Fig. S2). The third possibility is ruled out by the data presented in Figs. S5 and S6, which show that the SOA composition does not change significantly, except for the contribution of organic nitrates (see discussion above). This process of elimination leaves the fourth possibility—i.e., that the equilibrium described by Eq. 5 does not apply in this system.

To test that organic nitrates do partition into particles that are known to be liquid and where equilibrium should be rapidly achieved, uptake into liquid poly(ethylene glycol) (PEG) particles (see SI Text) is studied. Organic nitrates are generated in a separate set of experiments by reacting \( \alpha\)-pinene with \( O_3 \), generated from the thermal decomposition of varying concentrations of \( N_2O_5 \). The mixture of products and remaining reactants are then exposed to PEG particles. After approximately 10 min, particles are collected and analyzed using the same technique as for the \( NO_2/O_3/\alpha\)-pinene experiments. Fig. 4 (gray triangles) shows that \( F_i/M \) depends linearly on \( A_i \), consistent with equilibrium partitioning into liquid particles as described by Eq. 5. This outcome is not surprising; for a liquid of the viscosity of the PEG (ca. 0.1 Pa·s), the diffusion coefficient is approximately \( 5 \times 10^{-6} \text{ cm}^2\text{s}^{-1} \) and the time to diffuse from the surface to the center of a 260-nm diameter particle (average geometric diameter of the PEG particle distributions) is only about 35 ms (21).

In short, although the organic nitrates partition in equilibrium fashion as expected into liquid PEG particles, the same is not the case for the incorporation of organic nitrates into the SOA formed in the ozonolysis of \( \alpha\)-pinene, at least on the timescale (ca. 10–60 min) of the experiments.

An alternative hypothesis to equilibrium partitioning is that the particles grow and incorporate organic nitrates via a nonequilibrium, kinetically limited mechanism, similar to what has been described historically as a “condensation” mechanism (3, 4). Gas molecules impinge on the surface of an existing particle, with some probability that they will be taken up. This probability, and the time they remain adsorbed on the surface, are determined by the nature of the attractive forces operating between the gas and the surface of the particle. In addition, reactions at the surface that convert the adsorbate to lower volatility compounds (for example, oligomer formation) may contribute (48–51). If the residence time on the surface is sufficiently long, the adsorbed species can become “buried” and hence incorporated into the bulk by semivolatile gaseous compounds that subsequently condense. Based on gas kinetic theory, the time between collisions of a gas molecule with a molecular weight of 220 g mol\(^{-1}\) with an adsorbed surface molecule is about 100 ms, assuming a 10 ppb gas-phase SVOC concentration and surface area per molecule of 1 nm\(^2\). If a molecule at the surface has a longer residence time than 100 ms, the probability of it becoming buried by an incoming product molecule becomes significant. An estimate of residence times on the surface can be obtained from the rate constant for desorption (52), \( k_{des} = A_{des} \exp(-\Delta H_{des}/RT) \) where \( k_{des} \sim \Delta H_{des} \) (or \( \Delta H_{vap} \), depending on the process).

Heats of sublimation for acids and diacids are typically about 150 kJ mol\(^{-1}\) (53, 54) and heats of vaporization and sublimation for simple alcohols are typically in the range of 80–120 kJ mol\(^{-1}\) (55). Using \( A \sim 10^{13} \text{s}^{-1} \), lifetimes on the surface will range from 10 to 10\(^3\) s, more than sufficient to be buried by incoming SVOCs.

To assess if this growth mechanism is consistent with the experimental data, the uptake of APONO2 and Prod1 are treated as irreversible and occurring with uptake probabilities of \( \gamma_{APONO2} \) and \( \gamma_{Prod1} \). A value of the relative uptake coefficients, \( \gamma_{APONO2}/\gamma_{Prod1} = 0.016 \), is found to provide an excellent fit to the experimentally measured mass fraction of organic nitrates in the SOA (\( f_{APONO2} \), Table 1) for all \( NO_2 \) concentrations (Fig. S8). These results show that a nonequilibrium kinetically determined mechanism determines the formation and growth of SOA in this atmospherically important system.

A kinetics mechanism implies that reevaporation of the organic nitrates back to the gas phase is negligible, at least on the timescale of these experiments. Given the earlier discussion of diffusion times under typical liquid viscosities (21), the lack of reevaporation shows that the SOA material must be very viscous; once the organic nitrate is buried, it does not readily diffuse back to the particle surface and into the gas phase. The characteristic times for diffusion in a 400-nm diameter particle for a liquid are 10 \( \mu \)s to 10 ms for diffusion coefficients in the range of \( 10^{-5} \) to \( 10^{-9} \text{ cm}^2\text{s}^{-1} \) (21). For the approximately 1 h residence time in our flow system, equilibrium should be reached if the diffusion coefficient was \( \geq 10^{-14} \text{ cm}^2\text{s}^{-1} \). The fact that our data show equilibrium is not reached is compatible with the SOA being solid or semisolid, where \( D < 10^{-14} \text{ cm}^2\text{s}^{-1} \) (21). Clearly, the phase of a particle is important as it impacts the interaction of gas-phase products with the seed particles, and thus their growth rate, optical properties, and CCN activity (56).

Many studies have examined the timescales for the various steps associated with the formation and growth of SOA (21, 57–60); the conclusion is that the time for gas-phase SVOCs to come to equilibrium with liquid particles under typical atmospheric conditions is sufficiently short that the assumption of instantaneous equilibrium partitioning in atmospheric models is justified. However, a variety of recent results from other laboratories indicate that SOA in both ambient air and laboratory systems from reactions such as \( O_3 \) with \( \alpha\)-pinene does not always behave like a liquid. Vaden et al. (61) show that SOA mixed with a hydrophobic liquid organic forms layered particles with SOA at the core coated with the hydrophobic organic, or a hydrophobic core coated with SOA, both of which were stable for many hours. Vaden et al. (62) also demonstrate that evaporation of ambient SOA particles from Sacramento, California as well as laboratory-generated SOA, is quite slow, and the size dependence of the eva-
poration is not consistent with that expected for liquid particles. Earlier studies of the desorption of organic compounds from particles collected in a highway tunnel showed that the diffusion coefficients were orders of magnitude smaller than expected for liquids (63). Virtanen et al. (64, 65) report that particles collected in Hyytiälä, Finland and those generated in the laboratory by ozonolysis of α-pinene, bounce off impactor plates as if they are solids. Pierce et al. (66) show that freshly nucleated particles from Hyytiälä, Finland and Egbert, ON, Canada have very low volatility. Cappa and Wilson (67) demonstrate that the composition of SOA from the ozonolysis of α-pinene does not change with heating as expected if the evaporation is determined by equilibrium partitioning between a liquid particle and the gas phase. Thermodesorber measurements from field studies (11, 68) and laboratory-generated SOA from ozonolysis of monoterpenes (69, 70) show that thermal evaporation of the SOA (specifically at low RH) is much smaller than predicted by models of liquid particles, indicating that a significant fraction of the SOA is essentially nonvolatile.

Laboratory studies of the interaction of water vapor with some species relevant to atmospheric particles have also suggested that atmospheric particles may form highly viscous amorphous material under atmospheric conditions (71–73). More recently, Koop et al. (74) have shown that pinic acid and pinonic acid, which are major products of the α-pinene ozonolysis (42, 43), have glass transition temperatures between 265 and 268 K and thus can form a “glass” at lower temperatures found under some conditions in the troposphere; even above the glass transition temperature, the viscosity may be quite high.

In short, the results from our studies suggest that the assumption of instantaneous equilibrium commonly applied to SOA formation and evolution in most atmospheric models may need to be revisited. A kinetically limited/condensation growth mechanism actually can provide a better fit to field data. For example, measurements of the evolution of the number concentrations and size distributions of aerosols in the Mexico City area are shown to be consistent with a condensation mechanism for particle growth (75). Similarly, the growth of ultrafine aerosols from Egbert, ON, Canada and Hyytiälä, Finland are best modeled if >50% of the ultrafine particle growth was because of condensation (76). Very recent modeling efforts (77–79) also suggest the importance of this growth mechanism. As discussed in detail in the SI Text, the SOA measured in Riverside, California is consistent with a kinetically limited growth mechanism with no reevaporation if the overall average uptake coefficient for the SVOCs that lead to particle growth is of the order of 0.5–0.6. The issue of thermodynamic equilibrium has also been discussed in detail with respect to modeling inorganic atmospheric aerosols. The first generation of aerosol models assumed thermodynamic equilibrium for the volatile compounds between the gas and aerosol phase (80–83). However, measurements by Tanner (84), Allen et al. (85), and Wexler and Seinfeld (86) showed that equilibrium is not always achieved, especially within the time step used by models. Consequently, gas-to-particle conversion for inorganic species is now generally represented in aerosol modules by a dynamic mass transfer between the gas and aerosol phases (e.g., Meng et al., ref. 87).

In short, the combination of experiments reported here suggests that uptake of SVOCs into ambient SOA is consistent with a kinetically limited/condensation growth mechanism. Adsorbed SVOCs become incorporated into the bulk by being buried by incoming gas molecules and do not reevaporate, at least on the timescale of the experiments. If this process proves to be a general phenomenon, then the current formation and growth of SOA is not appropriately represented in most atmospheric models that rely on instantaneous thermodynamic equilibrium of SVOCs into liquid particles. Thus, current treatments of SOA formation and growth in models for both indoor and outdoor environments and the predicted impacts based on these models may need to be revisited.

Materials and Methods
A brief description of the experimental procedure is provided below and details can be found in the SI Text. α-Pinene was reacted with mixtures of O2 and NO2 in an aerosol flow tube (23). The size distributions, densities, and composition of the particles formed were measured using a scanning mobility particle sizer, an aerodynamic particle sizer, two particle mass spectrometers (SPLAT-II-MS and HR-TOF-AMS), LC-UV, and FTIR. Gas-phase concentrations were measured using a chemiluminescence NO analyzer, a photometric O3 analyzer, GC-MS, and long-path FTIR. For comparison, experiments were performed in Teflon reaction chambers using liquid PEG seed particles and the use of N2O as the source of NO2 radicals.

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