Images reveal that atmospheric particles can undergo liquid–liquid phase separations


A large fraction of submicron atmospheric aerosol particles contains both organic material and inorganic salts. As the relative humidity cycles in the atmosphere and the water content of the particles correspondingly changes, these mixed particles can undergo a range of phase transitions, possibly including liquid–liquid phase separation. If liquid–liquid phase separation occurs, the gas-particle partitioning of atmospheric semivolatile organic compounds, the scattering and absorption of solar radiation, and the reactive uptake of gas species on atmospheric particles may be affected, with important implications for climate predictions. The actual occurrence of liquid–liquid phase separation within individual atmospheric particles has been considered uncertain, in large part because of the absence of observations for real-world samples. Here, using optical and fluorescence microscopy, we present images that show the coexistence of two noncrystalline phases for real-world samples collected on multiple days in Atlanta, GA as well as for laboratory-generated samples under simulated atmospheric conditions. These results reveal that atmospheric particles can undergo liquid–liquid phase separations. To explore the implications of these findings, we carried out simulations of the Atlanta urban environment and found that liquid–liquid phase separation can result in increased concentrations of gas-phase NO, and N2O5 due to decreased particle uptake of N2O5.

In the atmosphere, single particles containing both organic species and inorganic salts are abundant (1, 2). The number of different types of inorganic salts is relatively small, with ammonium sulfate considered to be one of the most important in particles less than 1 μm in diameter (3, 4). In contrast, the number of organic species in a single atmospheric particle is on the order of thousands, with only ~10% of these species identified at the molecular level (5). As the relative humidity (RH) cycles in the atmosphere through high and low values, the water content of the particles increases and decreases as a hygroscopic response. In response to variable water content, the mixed particles can undergo a range of phase transitions including crystallization (i.e., efflorescence), dissolution (i.e., deliquescent), and liquid–liquid phase separation (Fig. 1) (6–12).

Results of laboratory measurements or calculations for particles or solutions containing ammonium sulfate mixed with one or a few specific organic molecules suggest, after extrapolation to atmospheric conditions, that liquid–liquid phase separations can occur in atmospheric particles (6, 10, 13–16). Atmospheric particles, however, are far more complex than the simple proxies used in these studies. A few studies have inferred that liquid–liquid phase separation occurs in particles containing ammonium sulfate and secondary organic material (SOM) generated from the dark ozonolysis of α-pinene (17–20). In the present study, liquid–liquid phase separation is explored for real-world samples.

Results and Discussion

Up to 90% of the submicron particle-phase organic material in the atmosphere is SOM (5). In a first set of measurements focused on the laboratory samples, we investigated the phase behavior of particles consisting of ammonium sulfate mixed with SOM that had been produced for conditions designed to simulate natural and polluted atmospheres. For these experiments, α-pinene was ozonized, a process that is considered to be a major biogenic source of SOM in the atmosphere (5). As a surrogate of a polluted atmosphere, 1,2,4-trimethylbenzene was photooxidized (21). In both cases, the oxidation of the volatile organic compounds led to oxidized products having low vapor pressures, and these products condensed to form SOM. The SOM produced by these approaches contained tens to hundreds of oxygenated organic species (22) and, as such, can represent a good surrogate for oxygenated organic material widely prevalent in the atmosphere (5). The oxygen-to-carbon (O:C) elemental ratios of the SOM used in these experiments are listed in Table 1. These ratios fall within the range of values observed in the atmosphere (23).

The SOM was collected on filters, extracted with water, and combined with ammonium sulfate to prepare solutions having organic-to-sulfate (org: sulf) mass ratios of 1.4 and 3.0 for α-pinene and 1,2,4-trimethylbenzene, respectively. These org: sulf ratios were chosen to overlap with ratios often prevalent in the atmosphere (24, 25). The solutions were atomized onto Teflon slides, producing supermicron particles (10–30 μm in diameter). The Teflon slides were then placed in a flow cell having temperature and relative humidity control, and the microscopy experiments were carried out (13).

Shown in Fig. 2 are the optical and fluorescence images of the particles as a function of RH. Also included for comparison are images of pure ammonium sulfate particles. Included in SI Text are image movies of ammonium sulfate particles and ammonium sulfate–SOM particles recorded as the RH was decreased and increased (Movies S1, S2, S3, and S4).


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Particles of ammonium sulfate behaved as expected (Fig. 2 A and B). At 90%, 70%, and 50% RH the particles consisted of a single aqueous phase, and when the relative humidity was decreased from 90% to 50%, the particles decreased in size due to the loss of water. At 30% RH the particles were crystalline. In addition, the fluorescence signal was nearly negligible for the ammonium sulfate particles for all RH values. In comparison with these results for ammonium sulfate, images of particles consisting of SOM and ammonium sulfate (Fig. 2 C–D) indicated the presence of two distinct phases. In the optical images, the outer phase appeared as a dark perimeter ring. The phase separation was especially evident in the fluorescence images because of the contrast between the fluorescent outer phase and the nonfluorescent inner phase. The mixed particles of ammonium sulfate and SOM from the ozonolysis of α-pinene appeared to have two distinct phases, even at the upper end of the RH measurement at 90% RH (Fig. 2 C and D). The mixed particles of ammonium sulfate and SOM from the photooxidation of trimethylbenzene appeared to have two phases at RH values up to at least 70% (Fig. 2 E and F).

Several pieces of evidence indicate that the inner phase was inorganic rich and the outer phase was organic rich. First, the fluorescence signal, practically absent for pure ammonium sulfate particles, was confined mainly to the outer phase. Second, the inner phase crystallized at relative humidities between 35% and 40% RH (compare Movies S1, S2, S3, and S4). This range overlaps with the crystallization RH range of pure ammonium sulfate particles (~33–37%) (26, 27). Third, the outer phase did not crystallize for the full range of RH values studied in our experiments, suggesting that there was negligible ammonium sulfate present in the outer phase. On the basis of these several observations, we conclude that the inner phase was an inorganic-rich phase and that the outer phase was an organic-rich phase. This conclusion is in agreement with previous Raman studies of particles containing ammonium sulfate and oxidized organic species (13, 14, 16).

We can also conclude from our results that both phases are noncrystalline above 35–40% RH. The presence of two separate noncrystalline phases confirms that liquid–liquid phase separation occurred in the particles, as molecular mobility is required for two phases to separate.

A few previous studies have inferred from laboratory studies that liquid–liquid phase separation occurs in particles containing ammonium sulfate and SOM generated from the dark ozonolysis of α-pinene (17–19). In these previous studies submicron particles were investigated. The consistency between the current results, which use supermicron particles, and these previous studies suggests that the liquid–liquid phase separations studied here do not depend strongly on particle size.

In the next set of measurements, we investigated the phase behavior of atmospheric samples collected in Atlanta, GA during August 6 and 7, 2010. TABLE 1. Relevant information for samples

<table>
<thead>
<tr>
<th>Particle type</th>
<th>O:C elemental ratio</th>
<th>Org:sulf mass ratio</th>
<th>NH₄⁺ mass, mg</th>
<th>SO₄²⁻ mass, mg</th>
<th>NO₃⁻ mass, mg</th>
<th>Other inorganic ions mass*, mg</th>
<th>SRH, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>α-Pinene dark-ozonolysis SOM + (NH₄)₂SO₄</td>
<td>0.3</td>
<td>1.4</td>
<td>0.9</td>
<td>2.4</td>
<td>NA</td>
<td>NA</td>
<td>&gt;90</td>
</tr>
<tr>
<td>1,2,4-trimethylbenzene photooxidation</td>
<td>0.4</td>
<td>3.0</td>
<td>2.3</td>
<td>6.2</td>
<td>NA</td>
<td>NA</td>
<td>&gt;70</td>
</tr>
<tr>
<td>SOM + (NH₄)₂SO₄</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Atlanta organic sulfate July 28 and 29, 2010</td>
<td>0.5³</td>
<td>1.5</td>
<td>0.9</td>
<td>2.7</td>
<td>0.5</td>
<td>0.3</td>
<td>&gt;90</td>
</tr>
<tr>
<td>Atlanta organic sulfate August 6 and 7, 2010 Unknown</td>
<td>1.0</td>
<td>2.0</td>
<td>4.7</td>
<td>0.9</td>
<td>0.2</td>
<td>&gt;90</td>
<td></td>
</tr>
</tbody>
</table>

The table shows oxygen-to-carbon (O:C) elemental ratio of the organic material, organic-to-sulfate (org:sulf) mass ratio, mass of inorganic material, and the liquid–liquid separation relative humidities (SRH). NA, not applicable.

*Other inorganic ions: Na⁺, K⁺, Mg²⁺, Ca²⁺, and Cl⁻.
³The O:C is based on a filter collected on July 29 and 30, 2010. We assume that the O:C was similar on July 28 and 29, because meteorological conditions and particle chemistry, such as org:sulf, concentrations of water-soluble organic carbon, and concentrations of inorganic ions did not change considerably over the 2 d.

Fig. 1. Schematic of possible phase transitions of particles containing mixtures of oxygenated organic material and ammonium sulfate. Processes: (A) liquid–liquid phase separation, (B) liquid–liquid mixing, (C and D) inorganic efflorescence, and (E and F) inorganic deliquescence.
July and August 2010. The atmosphere in and around Atlanta is heavily influenced by both anthropogenic and biogenic emissions (28). Single-particle mass spectrometric measurements in Atlanta have shown that ambient particles typically occur as internal mixtures of organic molecules and sulfate (29, 30). For our study, atmospheric samples were collected on filters, and inorganic salts and water-soluble organic material were extracted with high-purity water. The resulting extract solutions were atomized onto Teflon slides, producing supermicron particles for the microscopy studies. The filter extracts were almost exclusively ammonium sulfate and organic material, with other species making up less than 10% of the total mass identified (see Materials and Methods and Table 1 for further details). Additional ammonium sulfate was not added to the Atlanta filter samples.

Shown in Fig. 3 are results from samples collected July 28 and 29 and August 6 and 7, 2010 in Atlanta. Included in the SI are movies for samples collected on August 6 and 7, 2010 in Atlanta, recorded as the RH was decreased and increased (Movies S5 and S6). The sequence of observations is qualitatively similar to that described for the laboratory-generated particles of mixed SOM and ammonium sulfate. Following a similar line of argument, we then conclude that the atmospheric samples from Atlanta also underwent liquid–liquid phase separation. A total of four samples collected on different days in Atlanta were analyzed, and in all cases results similar to those shown in Fig. 3 were observed.

As only the relatively water-soluble species were extracted from the filters, the experiments herein can be anticipated to be enriched in more-oxidized organic species in comparison to the atmospheric particles. However, less-oxidized species tend to favor liquid–liquid phase separation more strongly than do more-oxidized species (6, 13, 16). Therefore, our results showing that liquid–liquid phase separation occurs even for the more-oxidized fraction suggest that liquid–liquid phase separation will also occur for the whole composition, as present in ambient atmospheric particles.

### Atmospheric Implications

Phase separation can influence the relative humidity at which crystallization and dissolution of sulfate solids occur in atmospheric particles (13, 14, 18) and thus can influence the extinction of solar radiation (31). Phase separation in atmospheric particles can also change the partitioning of organic molecules between the gas and particle phases. Thermodynamic calculations suggest that separation into organic-rich and inorganic-rich phases can increase the mass of organic material partitioning into the condensed phase by as much as 50% (6, 7).

Phase separation can also influence the reactive uptake of N₂O₅ into atmospheric particles (32–34). N₂O₅ is known to react...
efficiently on aqueous inorganic particles. Laboratory experiments show a significant decrease in $N_2O_5$ uptake for aqueous ammonium bisulfate particles coated with SOM (32, 33, 35). In addition, Reimer et al. (34) present simulations for Europe showing that liquid–liquid phase separation can effectively suppress $N_2O_5$ uptake onto particles in the atmosphere, leading to changes in mixing ratios of $N_2O_5$, NO$_2$, particle phase nitrate, and volatile organic compounds. A decrease in particle nitrate concentrations by up to 90% was simulated. Possibly related, Brown et al. show an anticorrelation between $N_2O_5$ reactive uptake and the organic–sulfate mass ratio for the northeastern United States (36).

To extend the simulations introduced by Reimer et al. but for the Atlanta environment, we used a box model to investigate the implications of liquid–liquid phase separation. Two different simulations were carried out. In case a, for $N_2O_5$ reactivity a homogeneous particle was assumed, and the rate of reaction depended only on the ratio of sulfate and nitrate in the particle (34, 37). In case b, for $N_2O_5$ reactivity an organic-rich coating and inorganic-rich core were assumed, and the rate of reaction depended both on the ratio of sulfate and nitrate in the core and the solubility and diffusion of $N_2O_5$ in the organic coating (34). The method of describing $N_2O_5$ reactive uptake followed that of Reimer et al. However, the model formulations used to describe the organic and inorganic particle concentrations as well as to treat liquid–liquid phase separation were different (SI Text and Table S1).

Fig. 4 shows the simulation results. The effect of the organic coating is evident in the $N_2O_5$ concentrations at nighttime and early morning (Fig. 4B). The organic coating reduces the uptake of $N_2O_5$ by particles. There is a resultant increase in gas-phase $N_2O_5$ and NO$_3$ concentrations by up to 15%. There is also a moderate reduction of gas-phase ozone concentrations (1% of ~30–50 parts per billion) in the morning (Fig. 4C), as explained by ozone titration with NO$_3$ (NO$_3$ = NO + NO$_2$) under low-sunlight intensities. The box model studies, taken together with the prior study of Reimer et al. (34), illustrate that liquid–liquid phase separation can impact concentrations of important atmospheric species.

For case b above we assume, similar to previous work, that the organic-rich phase will form a complete coating on the inorganic-rich core. This assumption is consistent with recent laboratory experiments that have shown a significant decrease in $N_2O_5$ uptake for aqueous ammonium bisulfate particles coated with SOM (32, 33, 35). A caveat, however, is that recent experiments using supermicron levitated particles illustrate that a partially engulfed structure can occur for mixed particles composed of aqueous salt solutions and organic molecules of low O:C values (11, 38, 39). Atmospheric particles that form an engulfed structure rather than a core-shell structure might have a considerably higher $N_2O_5$ uptake. Additional studies focusing on the factors influencing morphology after liquid–liquid phase separation would be beneficial to help resolve this issue.

The work reported here shows that liquid–liquid phase separation can occur in the atmosphere when the O:C ratio of the organic material is roughly $\leq$0.5. Additional studies are needed to determine whether these phase transitions also occur in the atmosphere when the O:C ratio is $>$0.5.

Materials and Methods

Production of Secondary Organic Material. Dark ozonolysis of $\alpha$-pinene was performed in the Harvard Environmental Chamber to produce secondary organic material that then condensed on dry ammonium sulfate seed particles. Ammonium sulfate seed particles were introduced into the chamber by atomization of a solution of (NH$_4$)$_2$SO$_4$, which was then dried using a diffusion dryer. The $\alpha$-pinene and ozone were introduced to the chamber by a flow of purified air. The setup and experimental conditions of the dark ozonolysis experiments were similar to those used by Shilling et al. (40). Flow into the continuous-flow chamber was fixed at 20 standard liters per minute (slpm) during operation. The temperature and relative humidity inside the chamber were maintained at 25 °C and 40%, respectively. An Aerodyne high-resolution time-of-flight Aerosol Mass Spectrometer (HR-ToF-AMS) was used to determine the composition of the particles. The secondary organic material with ammonium sulfate seeds was collected at the outlet of the continuous-flow chamber onto
a quartz fiber filter. The collection time for sampling was 48 h at a flow rate of 9.0 1slpm.

Photooxidation of 1,2,4-trimethylbenzene was carried out in the Pacific Northwest National Laboratory (PNNL) Continuous-Flow Environmental Chamber to produce secondary organic material; 1,2,4-trimethylbenzene was injected into 24 slpm of pure airflow to produce concentrations of 4.8 ppmv (parts-per-million by volume) in the chamber before reaction. An additional 1 slpm flow of pure air was bubbled through a 50% (vol/vol) solution of hydrogen peroxide in water and added to the chamber. Photooxidation was initiated by 105 UV (Q-Laboratories UVA-340) lights surrounding the chamber, which generated OH radicals from the photolysis of hydrogen peroxide. The chamber temperature and relative humidity were maintained at 18.1 ± 0.2 °C and 6 ± 2%, respectively, over the course of the 4-d experimental run. NOx levels were below the 1-ppbv (parts-per-billion by volume) detection limit of a NOx chemiluminescence detector. Particle chemical composition was analyzed in real time, using an Aerodyne HR-ToF-AMS. The secondary organic material was collected at the outlet of the continuous-flow chamber onto a Teflon filter ( Pall; R2FL037). The collection time for sampling was 82 h at a flow rate of 4 slpm.

Collection of Samples from Atlanta, GA. Particles were collected on quartz fiber filters for ~24 h, using a high-volume sampler (Thermo Anderson; flow rate 1.13 × 103 1slpm). Collection occurred on the Georgia Institute of Technology Environmental Science and Technology rooftop laboratory located in central Atlanta, ~15 m above ground level. The concentration of organic carbon and inorganic salts on the filters was determined from 2.54-cm diameter filter punches, using a total organic carbon analyzer and ion chromatography. For the sample collected on July 28 and 29, 2010, the mass composition was the following: organic carbon = 3.98 mg; NH4+ = 0.859 mg; SO42− = 2.711 mg; NO3− = 0.471 mg; Na+ = 0.097 mg; K+ = 0.043 mg; Mg2+ = 0.019 mg; Ca2+ = 0.068 mg, and Cl− = 0.049 mg. The O:C elemental ratio of the organic component of the filters collected in Atlanta was determined by extracting the filters with water and then determining the O:C from the filter extracts, using HR-ToF-AMS (41, 42).

Production of Particles. Water-soluble species were extracted from the filter samples, using Millipore water (18.2 MΩ cm). For the samples collected in Atlanta, particles were generated directly from the filter extracts without the addition of ammonium sulfate. Filter extracts were passed through a Particle-on-Demand Generator (model MJ-ABL-01-120; MicroFab Technologies) to produce particles with sizes ranging from 10 to 30 μm in diameter. These particles were deposited on Teflon slides and then placed in a flow cell that had temperature and relative humidity control for the microscopy experiments (43).

To increase the sulfate-to-organic mass ratio to values prevalent in the atmosphere (24, 25), ammonium sulfate solution was added to the filter extracts from the environmental chambers. After addition of ammonium sulfate, the organic-to-sulfate mass ratio in the samples from α-pinene and 1,2,4-trimethylbenzene were 1.4 and 3.0, respectively. Once ammonium sulfate was added to these particles as described above.

Optical and Fluorescence Microscopy of Particles. Images of the particles were recorded with fluorescence microscopy (Zeiss LSM510; λexcitation = 543 nm, λemission = 650–710 nm, 293 ± 1 K) and optical light-reflectance microscopy (Zeiss AxioTech; 50x objective, 273 ± 1 K). At the beginning of an experiment, the RH in the flow cell was set to 90–100%, and the particles were allowed to equilibrate with the flow-cell RH for 15–20 min. The RH was then decreased at a constant rate (0.6%/min) while images were collected. Calibration of the absolute RH readings was performed using the
deliquescence relative humidity values for pure ammonium sulfate particles. Ammonium sulfate particles were studied for comparison with the experimental samples. First a mixture of ammonium sulfate and high-purity water was prepared. This solution was then added to a blank filter and processed the same way as the filter samples discussed above.

**Box Model Simulations.** A box model was used to simulate chemistry in the Atlanta atmosphere. The size of the modeling region was set to 36 × 36 km (1,296 km²) with a 600-m boundary layer height. Further details on the box model are given in SI Text and Table S1.

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Supporting Information

You et al. 10.1073/pnas.1206414109

SI Text

Box Model Simulations. A box model was used to simulate chemistry in the Atlanta atmosphere. The concentration of species \( i \) was described using the equation

\[
\frac{dc_i}{dt} = -\nabla \cdot (uc_i) + Q_{\text{chem},i} + Q_{\text{aerosol},i} + Q_{\text{depos},i} + E_i, \tag{S1}
\]

where \( c_i \) is the concentration of species \( i \), \( u \) is the air mass velocity, \( Q_{\text{chem},i} \) is the rate change of species \( i \) in the gas phase, \( Q_{\text{aerosol},i} \) is the rate change of species \( i \) in the aerosol phase, \( Q_{\text{depos},i} \) is the loss rate of \( i \) due to deposition, and \( E_i \) is the emission rate of \( i \).

The size of the modeling region was set to 36 × 36 km (1,296 km²) with a 600-m boundary layer height. It was assumed that the air mass entering Atlanta contained background levels of pollutant concentrations (1). Boundary conditions for biogenic species were obtained from Riemer et al. (2). These concentrations were incorporated in the model as Dirichlet boundary conditions (3). The dry deposition mechanism was adapted from the California Institute of Technology (CTT) model (4) and was incorporated in the modeling region as a bottom boundary condition. Rates of dry deposition were calculated assuming that the entire Atlanta area is a “mixed urban environment” with residential, commercial, and industrial structures. The top of the modeling region incorporated a no-flux boundary condition.

Meteorological conditions for Atlanta for the period from July 27 to July 31, 2010 were obtained from the National Oceanic and Atmospheric Administration (NOAA) National Weather Service. Information included temperature, relative humidity, wind speed, precipitation, and cloud coverage. The episode was characterized by clear skies and dry conditions, except for 0.1 inches of rain in the second day and variable cloud coverage in the last day. The average temperature during the episode was 28.4 °C, and the average humidity was 68.5%. To account for precipitation, a parameterization of wet deposition was implemented on the basis of Martin et al. (5). Cloud coverage was parameterized as a multiplication factor (HV) to total solar radiation. The parameterization is presented in Table S1.

Emission rates were calculated using the US Environmental Protection Agency National Emissions Inventory for 2002 (http://www.epa.gov/ttn/chief/eiinformation.html) and the Sparse Matrix Operational Emissions (SMOKE) model, to determine the specific emissions for the metropolitan area of Atlanta at the 36-km resolution. Size distribution of particle emissions was determined on the basis of measurements reported by Woo et al. (6).

The Caltech Atmospheric Chemistry Mechanism (CACM) (7) was used to describe the gas-phase chemistry. CACM is based on the work of Stockwell et al. (8), Jenkin et al. (9), and Carter (10) and includes \( O_3 \) chemistry and a state-of-the-art mechanism of the gas-phase precursors of secondary organic material (SOM). The full mechanism consists of 361 chemical reactions and 191 gas-phase species, which describe a comprehensive treatment of the oxidation of volatile organic carbon species.

Inorganic aerosol formation is calculated using the Simulating Composition of Atmospheric Particles at Equilibrium model (SCAPE2) (Meng et al. (11)), whereas organic aerosol formation is calculated using the Model to Predict the Multiphase Partitioning of Organics (MPMPO) (Griffin et al. (12) and Pun et al. (13)). MPMPO allows the simultaneous formation of SOM in a hydrophobic organic phase and a hydrophilic aqueous phase. In addition, MPMPO modifies SCAPE2 to account for the interaction between organic ions present in the aqueous phase and the inorganic aerosol components. The module consists of 37 size-resolved aerosol-phase species, in eight different-size bins ranging from 0.04 to 10 µm. It is assumed that all particulate material is internally mixed.

The MPMPO module treats explicitly the surrogate secondary organic material on the basis of their hydrophilic/hydrophobic character. The SOM surrogates in the model are described below.

Hydrophilic.

A1: anthropogenic, dissociative, low no. C (2)
A2: anthropogenic, dissociative, high no. C (8)
A3: anthropogenic, non dissociative
A4: biogenic, dissociative
A5: biogenic, non dissociative

Hydrophobic.

B1: anthropogenic, benzene based, low volatility
B2: anthropogenic, benzene based, higher volatility
B3: anthropogenic, naphthalene based
B4: anthropogenic, aliphatic
B5: biogenic, aliphatic.

The rate for \( \text{NO}_2 \) hydrolysis on particles to produce nitric acid followed the implementation described by Riemer et al. (14). The reaction rate was described by the equation

\[
\frac{d[\text{NO}_2]}{dt} = -\frac{1}{4} \gamma_{\text{het}} \text{NO}_2 \gamma_{\text{het}} \text{NO}_2 [\text{NO}_2], \tag{S2}
\]

where \( \gamma_{\text{het}} \) is the molecular speed of \( \text{NO}_2 \), \( S \) is the surface area of the aerosol, and \( \gamma_{\text{het}} \) is the uptake coefficient of \( \text{NO}_2 \) on particles.

Two different simulations were carried out. In case a, for \( \text{NO}_2 \) reactivity a homogeneous particle was assumed, and the uptake coefficient of \( \text{NO}_2 \) was described by the equation

\[
\gamma_{\text{het}} = 0.002 f + 0.02(1 - f), \tag{S3}
\]

where \( f \) = (nitrate mass/sulfate mass + nitrate mass). Eq. S3 is based on a simplified parameterization of the relative concentration of nitrate loading with respect to the total nitrate and sulfate loading (15). In case b, for \( \text{NO}_2 \) reactivity, an organic-rich coating and an inorganic-rich core were assumed, and the reactive uptake coefficient was described by Eq. S4,

\[
\frac{1}{\gamma_{\text{het}}} = \frac{1}{\gamma_{\text{core}}} + \frac{1}{\gamma_{\text{coat}}} \tag{S4}
\]

where \( \gamma_{\text{core}} \) and \( \gamma_{\text{coat}} \) were described by the equations

\[
\gamma_{\text{core}} = 0.002 f + 0.02(1 - f) \tag{S5}
\]

\[
\gamma_{\text{coat}} = \frac{4RTH_{\text{org}}D_{\text{org}}R_c}{c_{\text{NO}_2}R_p}, \tag{S6}
\]

where \( R \) is the ideal gas constant, \( T \) is temperature, \( H_{\text{org}} \) is Henry’s law coefficient for \( \text{NO}_2 \) on organic liquid, \( D_{\text{org}} \) is the diffusion coefficient of \( \text{NO}_2 \), \( R_c \) is the radius of the inorganic core, \( l \) is the thickness of the film, and \( R_p \) is radius of the particle. Eqs. S4–S6 assume that the rate of reaction depends on both the ratio of
sulfate and nitrate in the core and the solubility and diffusion of N$_2$O$_5$ in the organic coating. In case b the organic-rich coating was assumed to include all primary organic aerosol components and the hydrophobic portion of the secondary organic material, whereas the inorganic-rich core was assumed to include all other material in the particle phase.


### Table S1. Relation between cloud coverage and solar radiation used in the box model simulations

<table>
<thead>
<tr>
<th>Condition</th>
<th>HV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clear</td>
<td>0.9</td>
</tr>
<tr>
<td>Scattered clouds</td>
<td>0.8</td>
</tr>
<tr>
<td>Partly cloudy</td>
<td>0.5</td>
</tr>
<tr>
<td>Light rain</td>
<td>0.3</td>
</tr>
<tr>
<td>Mostly cloudy</td>
<td>0.3</td>
</tr>
<tr>
<td>Overcast</td>
<td>0.2</td>
</tr>
<tr>
<td>Rain</td>
<td>0.2</td>
</tr>
<tr>
<td>Haze</td>
<td>0.1</td>
</tr>
<tr>
<td>Heavy rain</td>
<td>0.1</td>
</tr>
<tr>
<td>Thunderstorm</td>
<td>0.1</td>
</tr>
</tbody>
</table>

Cloud coverage was parameterized as a multiplication factor (HV) to total solar radiation.

**Movie S1.** Optical image movie; a particle consisting of ammonium sulfate (diameter = 21.6 μm). Images were recorded as the relative humidity (RH) was decreased from 50% to 35% at a temperature of 270 ± 1 K. The ramp rate was −0.6% RH min$^{-1}$. For the particle shown, efflorescence occurred at 37.4 ± 4%. A rough indication of the RH during the movie is included.

You et al. www.pnas.org/cgi/content/short/1206414109
Movie S2. Optical image movie; a particle consisting of ammonium sulfate (diameter = 21.6 μm). Images were recorded as the RH was increased from 62% to 80% RH at a temperature of 270 ± 1 K. The ramp rate was ∼0.6% RH min⁻¹. For the particle shown, deliquescence occurred at 78.0 ± 4%. A rough indication of the RH during the movie is included.

Movie S3. Optical image movie; a particle (org: sulf = 1.4 and diameter = 28.0 μm) consisting of ammonium sulfate mixed with SOM generated by the ozonolysis of α-pinene. Images were recorded as the RH was decreased from 90% to 35% at a temperature of 270 ± 1 K. The ramp rate was ∼0.6% RH min⁻¹. For the particle shown, liquid–liquid phase separation occurred at an RH greater than 90 ± 4% and efflorescence occurred at 40.5 ± 4%. A rough indication of the RH during the movie is included.

Movie S4. Optical image movie; a particle (org: sulf = 1.4 and diameter = 28.0 μm) consisting of ammonium sulfate mixed with SOM generated by the ozonolysis of α-pinene. Images were recorded as the RH was increased from 60% to 80% RH at a temperature of 270 ± 1 K. The ramp rate was ∼0.6% RH min⁻¹. For the particle shown, complete deliquescence occurred at 77.0 ± 4%. A rough indication of the RH during the movie is included.
Movie S5. Optical image movie; a particle (org:sulf = 1.0 and diameter = 18.3 μm) generated from a filter sample collected August 6 and 7, 2010 in Atlanta. Images are shown as the RH was decreased from 68% to 35% at a temperature of 270 ± 1 K. The ramp rate was ∼0.6% RH min⁻¹. For the particle shown, liquid–liquid phase separation occurred at an RH greater than 90% ± 4%, whereas efflorescence occurred at an RH of 37.0 ± 4%. A rough indication of the RH during the movie is included.

Movie S5

Movie S6. Optical image movie; a particle (org:sulf = 1.0 and diameter = 18.3 μm) generated from a filter sample collected August 6 and 7, 2010 in Atlanta. Images are shown as the RH was increased from 50% to 80% at a temperature of 270 ± 1 K. The ramp rate was ∼0.6% RH min⁻¹. For the particle shown, deliquescence occurred at 71.6 ± 4%. A rough indication of the RH during the movie is included.

Movie S6