Rapid growth of organic aerosol nanoparticles over a wide tropospheric temperature range


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Nucleation and growth of aerosol particles from atmospheric vapors constitutes a major source of global cloud condensation nuclei (CCN). The fraction of newly formed particles that reaches CCN sizes is highly sensitive to particle growth rates, especially for particle sizes <10 nm, where coagulation losses to larger aerosol particles are greatest. Recent results suggest that some oxidation products from biogenic volatile organic compounds are major contributors to particle formation and initial growth. However, whether oxidized organic compounds can contribute to particle growth over the broad span of tropospheric temperatures remains an open question, and quantitative mass balance for particle growth over the broad span of tropospheric temperatures is compensated by the decreased volatility of all oxidation products from biogenic volatile organic compounds.

The global budget of cloud condensation nuclei (CCN) significantly influences the Earth’s radiative balance, as it affects the albedo and the lifetime of clouds. New particle formation by gas-to-particle conversion is the largest source of CCN (1). Especially the early steps of particle growth between 1 and 10 nm determine the survival chance of freshly formed particles and therefore their climatic relevance (2, 3). The major vapors driving particle growth are sulfuric acid and, maybe more importantly, low-volatility organics resulting from the oxidation of volatile organic compounds (VOCs) (4). Monoterpenes are an important class of atmospheric VOCs with copious emissions from vegetation (5). They are quickly oxidized in the atmosphere and, through a subsequent autoxidation process, rapidly contribute to particle growth over a wide range of tropospheric temperatures from molecular cluster sizes onward.

aerosols | nanoparticle growth | aerosol formation | CLOUD experiment | volatile organic compounds


The authors declare no conflict of interest.

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Aerosol particles can form and grow by gas-to-particle conversion and eventually act as seeds for cloud droplets, influencing global climate. Volatile organic compounds emitted from plants are oxidized in the atmosphere, and the resulting products drive particle growth. We measure particle growth by oxidized biogenic vapors with a well-controlled laboratory setup over a wide range of tropospheric temperatures. While higher temperatures lead to increased reaction rates and concentrations of highly oxidized molecules, lower temperatures allow additional, but less oxidized, species to condense. We measure rapid growth over the full temperature range of our study, indicating that organics play an important role in aerosol growth throughout the troposphere. Our finding will help to sharpen the predictions of global aerosol models.

In contrast to sulfuric acid plus ammonia or amines, where growth proceeds close to the kinetic limit (14), growth driven by organics is governed by the resulting volatilities of the wide variety of oxidation products. Therefore, temperature likely plays a decisive role, as the saturation concentration has a steep exponential temperature dependence as described by the Clausius–Clapeyron relation. Additionally, a recent study has shown that temperature crucially influences the chemical composition of the initially formed molecular clusters in α-pinene ozonolysis (15). Therefore, the contribution of biogenic organics to new particle formation might be strongly sensitive to temperature. This, in turn, may significantly influence the importance of new particle formation at high altitudes (16) and in outflow regions of deep-convective clouds—for example, over the Amazon Basin (17–19).

Here, we investigate in the Cosmics Leaving Outdoor Droplets (CLOUD) chamber (20) the effect of temperature on the production of oxygenated molecules and subsequent particle growth from dark α-pinene ozonolysis at three different temperatures (−25 °C, 5 °C, and 25 °C) for various precursor concentrations. The resulting volatility distributions are inferred by combining two types of chemical ionization (CI) high-resolution time of flight mass spectrometers (TOF-MS) (21, 22) using complementary ionization techniques to obtain a detailed representation of the gaseous oxidation products. Together with the precision measurement of particle growth rates (23) and analysis of the particle-phase composition (24), this allows identification of the underlying processes and their temperature dependence responsible for initial growth in biogenic ozonolysis systems (see Materials and Methods for details about the experimental setup and measurement procedures).

**Results**

**Observed Gas-Phase Composition and Volatility Distribution.** We measured gas-phase composition with a nitrate-CI atmospheric pressure interface (API)-TOF-MS (nitrate-CI) (21) and a proton transfer reaction (PTR)-TOF-MS (PTR3) (22) to obtain a more detailed overview of the neutral gas-phase species present during the α-pinene ozonolysis experiments. We obtained overlap for peaks observed in both instruments (SI Appendix, Fig. S3) and show a combined mass-defect plot of both instruments for three representative experiments at three different temperatures in SI Appendix, Fig. S4. The PTR3 introduces >200 previously undetected molecular ion signals, not only HOMs, which are usually specified by their high oxygen to carbon ratio (O:C > 0.7 for monomers), but mostly compounds toward lower oxidation states. For molecules with identified chemical composition, a volatility can be assigned according to the number of oxygen atoms \( n_0 \) and the number of carbon atoms \( n_C \) within the molecule (SI Appendix).

As volatilities of organic compounds observed in the atmosphere vary by >10 orders of magnitude and the combined mass spectra contain ∼500 different molecules, it is convenient to simplify considerations of gas-to-particle partitioning by grouping compounds together within a volatility basis set (VBS) (13, 25). Within this framework, the volatility bins are separated by one decade in \( C^* \) at 300 K, and for other temperatures, the binned distribution is shifted toward lower saturation mass concentrations. The saturation mass concentration of oxidized organics should follow the Clausius–Clapeyron relation at a constant evaporation enthalpy \( \Delta H_{\text{vap}} \), which in turn is linked to \( C^* \) at 300 K (13) (SI Appendix).

Fig. 1 shows the resulting binned volatility distribution of all observed organic gas-phase compounds for three representative experiments. We averaged observed gas-phase concentrations and, with increasing particle size, more and more low-volatility species can contribute to the growth process.

Fig. 1. Volatility distributions for representative experiments with similar α-pinene ozonolysis rate: 25 °C (A), 5 °C (B), and −25 °C (C). The green and blue bars show summed molecular ions observed in the nitrate-CI and PTR3, respectively. The highest and lowest bin are overflow bins. Volatility bins are defined at 300 K, shifted, and widened according to their corresponding temperature. The resulting saturation mass concentration is defined on the x axis, while log_{10} \( C^*_{\text{sat}} \) is specified by white numbers. Additionally, the bins in supersaturation with \( C^*/C^* > 1 \) are found left of the indicating arrow. ELVOC, extremely low-volatility organic compound; IVOC, intermediate-volatility organic compound; LVOC, low-volatility organic compound; SVOC, semi-volatile organic compound.
\( \text{C}^{\circ} \) over a period where comparable particle growth rates are measured with a differential mobility analyzer (DMA) train (23), and the \( \alpha \)-pinene ozonolysis rate is similar with \( k(T) \cdot [\text{ap}] \cdot [\text{O}_3] \approx 1.4 \cdot 2.0 \cdot 10^{9} \text{ cm}^{-3} \text{s}^{-1} \).

Due to the comparable growth rates of the three examples, the gas-particle partitioning is expected to be comparable, which is confirmed by the similarity of the observed total volatility distribution over the extremely low-volatility organic compound (ELVOC) and low-volatility organic compound (LVOC) ranges. Earlier work on growth of nucleated particles from \( \alpha \)-pinene oxidation at 5°C using only a nitrate-CI found that the measured HOMs could only explain a fraction of the growth and speculated that the nitrate detection efficiency was progressively lower for less polar (and, hence, more volatile) species (8). We confirmed the missing fraction and found that the PTR3 detected many new compounds, mainly less-oxygenated molecules with \( n_O \leq 7 \), not measured by the nitrate-CI, independent of temperature. At low temperature, fewer polar functional groups are required for a compound to have a low volatility, and thus at 5°C, and even more significantly at \(-25^\circ \text{C}\) (Fig. 1B and C, respectively), these species observed by the PTR3 contribute substantially in the LVOC and even ELVOC range.

**Particle-Growth Measurements.** We measured growth rates during the experiments with a DMA train over two different size intervals, 1.8–3.2 nm and 3.2–8 nm, by the appearance time method as it was done in comparable studies (8, 14). It gives robust apparent particle-growth rates for chamber experiments and is not affected by measurement uncertainties in absolute particle concentrations due to possible evaporation effects during the measurement procedure (SI Appendix). Fig. 2 shows the measured growth rates vs. several gas-phase variables. Fig. 2A and B shows the correlation with the estimated reaction rate of the \( \alpha \)-pinene ozonolysis during the growth rate measurement. Higher reaction rates, and hence higher product concentrations, lead to higher growth rates, following an exponential relation \( m(T, d_p) \cdot (k(T) \cdot [\text{ap}] \cdot [\text{O}_3])^g \) (SI Appendix). For a given \( \alpha \)-pinene ozonolysis reaction rate, we find lower growth rates at smaller sizes. The smaller size range also shows a more significant temperature dependency: The growth rates are higher at low temperatures at a given reaction rate. This indicates that the ozonolysis products at the three different temperatures have different properties influencing their ability to condense from molecular cluster sizes onward.

Fig. 2 C and D shows the measured growth rates vs. the total HOM signal observed in the nitrate-CI only, along with a kinetic curve showing the growth rate if all measured HOMs condensed irreversibly (26). The growth rates of the three different temperatures are clearly separated, but condensation at the kinetic limit for HOMs would give almost identical values. Thus, the total HOM concentration observed in the nitrate-CI cannot fully describe the observed growth at any temperature. At 25°C, several HOMs measured by the nitrate-CI are classified as semi-volatile organic compounds and might not be able to condense, and at \(-25^\circ \text{C}\), the nitrate-CI measures only a small fraction of the less-oxygenated \( \alpha \)-pinene oxidation products responsible for particle growth (Fig. 1).

Therefore, Fig. 2 E and F shows the growth rates vs. a sum, combining both mass spectrometers, over all VBS bins in supersaturation for a given particle size—that is, with \( S = K(D_p) \cdot C_{VBS,\text{tot}}^\circ \cdot C_{VBS,\text{tot}}^\circ > 1 \). A Kelvin term \( K(D_p) = 10^{(D_p)/D_p} \) accounts for the curvature of the particles, slowing growth of smaller particles. With this simple approach, it is possible to bring the growth measurements at these three different temperatures into reasonable agreement, aligning the data points roughly parallel to the kinetic line. This approach only accounts for bins in supersaturation, which should condense almost kinetically. Especially for the larger size interval, the measured growth rates were slightly higher than the supersaturated kinetic limits for all temperatures. However, some VBS bins below supersaturation will contribute as well by gas-particle partitioning, which was not considered in this simple approach.
Comparison with an Aerosol Growth Model. We modeled growth with the same framework as used in ref. 8. However, we modified the model to take real-time measured VBS distributions from both mass spectrometers as input, without any adjustments of unknown charging efficiencies (SI Appendix).

The most important remaining unknown in the condensation equations is the Kelvin term, $K(D) = 10^{4K_{10}/T}$, parametrized for simplicity by a decadal Kelvin diameter related to bulk liquid properties $D_{K10} = \log_{10}(s) \cdot (4\pi M) \cdot (RTp)^{-1}$. However, the observed size dependence and especially the growth measurements at diameters $< D_{K10}$ should provide a direct constraint on the curvature effect. For the three representative experiments, we found the best agreement with $D_{K10}(T) = (4.8 \pm 0.8) \cdot (300 \text{ K} \cdot T^{-1})$ nm. This corresponds to reasonable average properties (surface tension $\sigma = 0.03 \text{ N} \cdot \text{m}^{-1}$, molecular mass $M = 320 \text{ g} \cdot \text{mol}^{-1}$, and density $\rho = 1400 \text{ kg} \cdot \text{m}^{-3}$) for the many condensing species, ignoring other possible temperature dependencies in $\rho$, $\sigma$, and $M$. Fig. 3 shows the resulting predicted growth rates and their size dependence in comparison with the measurements. The agreement between modeled and measured growth rate at the smallest sizes is within the uncertainties of the measurements. Other values for $D_{K10}$—for example, $D_{K10}(300 \text{ K}) = 3.75 \text{ nm}$, used previously—lead to a significant overestimation of the observed growth rates at the smallest diameters for all temperatures. Another reason for the higher $D_{K10}$ could be an underestimation of the volatility of the most oxygenated compounds (27). Above 5 nm, the model agreed well with the observations at all temperatures. Considering the 1-decade uncertainty in saturation mass concentration (SI Appendix), we achieved reasonable mass balance for growth of freshly nucleated particles between 2 and 30 nm over a wide range of conditions.

Although there was no disagreement of the model with the measurements, within the uncertainties, there are several contributions that we have not considered. First, some condensable compounds might still be undetected by the two used ionization chemistries. Additionally, fragmentation of molecules within the instruments might disturb the volatility estimate. Second, the temperature dependence of organic volatilities is also subject to uncertainties (13). Third, we did not model any particle-phase reactions, such as oligomerization. Reactive uptake is thought to be more important at larger particle sizes (28), again in part because of the Kelvin effect (29).

Particle-Phase Composition Measurements. The predictions by the aerosol growth model were supported by the measurement of the particle-phase composition using a filter inlet for gas and aerosols (FIGAERO) (24) attached to a $O_{2}^{-}\text{-CI-API-TOF-MS}$ [FIGAERO-chemical ionization mass spectrometer (CIMS)] (SI Appendix). Fig. 4 shows the desorption profiles of three molecular ion signals corresponding to $C_{10}H_{16}O_{4}$, $C_{10}H_{16}O_{6}$, and $C_{10}H_{16}O_{8}$. We accumulated particles on the FIGAERO filter inlet at chamber temperature during the three experiments at $-25 \text{ °C}$, $5 \text{ °C}$, and $25 \text{ °C}$. The signal intensity was normalized to the accumulated mass on the filter inferred from the measured particle-size distributions and sample flow rate, which allowed for quantitative comparison of the three temperatures. We fit the desorption profiles with a bimodal distribution, where the first mode represented the monomer signal and the second mode was due to fragmentation products of less volatile dimers which therefore desorb at higher temperatures (SI Appendix). Desorption of the monomer clearly occurred earlier for the less-oxygenated products, which experimentally confirmed the volatility dependence on $n_{O}$ used for the volatility estimates.

It was evident that the less-oxygenated monomers only appeared at lower temperature in the particle phase. $C_{10}H_{16}O_{4}$ contributed significantly only at $-25 \text{ °C}$, and $C_{10}H_{16}O_{6}$ appeared already at $5 \text{ °C}$ in the particle phase. For $C_{10}H_{16}O_{8}$, a subsequent reduction in the normalized particle-phase intensity was observed for decreasing temperatures. Although the volatility of this molecule is low enough to contribute significantly even at $25 \text{ °C}$, its production in the gas phase was reduced at lower temperatures, and its contribution to the particle phase thus decreased with decreasing temperature.

These trends are supported by the comparison of gas- and particle-phase composition within the representative $C_{10}H_{16}O_{4-9}$ series, which contains the most dominant peaks of the particle-phase mass spectra. In Fig. 5, the fractional contributions of $C_{10}H_{16}O_{4-9}$ are shown for the gas and particle phase for the three investigated temperatures. Generally, the most abundant gas-phase product at all temperatures within this series is pinonic acid ($C_{10}H_{16}O_{4}$), having a similar yield to all HOM products together (7, 30). However, the fraction of the contribution of highly oxygenated products increased significantly at warmer temperatures (see also SI Appendix, Fig. S4), while at $-25 \text{ °C}$, less oxygenated products dominated the gas phase completely. In the particle phase, HOM products were mainly found at $25 \text{ °C}$, while at lower temperatures, the particle phase contained a large contribution of less-oxygenated molecules with even a strong contribution of the modestly oxygenated pinonic...
acids. This is not only because of the decrease of the availability of gas-phase HOM products, but mainly due to a strong drop in volatility of all compounds, resulting in condensation of less-oxygenated products. This is similar to the observations in ref. 15, where a significant decrease in O:C of the nucleating charged clusters was observed at colder temperatures.

**Conclusion**

Organics play a leading role in atmospheric new particle formation and growth and thus govern the global budget of CCN. VOC oxidation products in the atmosphere make up a substantial portion of condensing vapors causing growth of existing particles. Because oxidized organics span a wide range of volatilities, temperature is a crucial parameter. We use a combination of mass spectrometers, two using complementary ionization techniques for gas-phase measurements and one measuring particle-phase composition. Consideration of the volatility distribution of the measured gas-phase compounds, here with a volatility basis set, gives a sufficient constraint of the gas-phase products to comprehensively describe growth over a wide temperature range. The measured and modeled particle-phase compositions are self-consistent. The measurements are in good agreement with an aerosol growth model, and a direct estimate of the Kelvin distribution measurements in the crucial region of <10 nm provide important constraints for model predictions of the contribution of gas-to-particle conversion to the global budget of CCN.

**Materials and Methods**

The CERN, the European Organization for Nuclear Research, CLOUD chamber is a 26.1-m³ electropolished stainless-steel vessel, surrounded by a thermal housing capable of stabilizing temperature in a range from −65 °C to 100 °C with ±0.1 K precision (33). The chamber is equipped with a gas-control system achieving extremely high purities by mixing boil-off nitrogen and boil-off oxygen at the atmospheric ratio of 79:21. Highly pure trace gases can be precisely added at the parts per trillion (ppt) level. Before the experiments, the chamber was heated to 100 °C and rinsed with ultrapure water. This assured operation at contaminant levels of <5 × 10^16 cm⁻³ H₂SO₄ and total organics <150 ppt by volume (15, 20).

![Fig. 4](image)

**Fig. 4.** Mean thermal desorption profile of three compounds found in particle-phase composition measurements with a FIGAERO-CIMS and the corresponding SE (shaded areas). The signal intensity normalized (Norm.) by primary ion signal and collected particle mass vs. the desorption temperature is compared for three representative experiments with red indicating 25 °C, green indicating 5 °C, and dark blue indicating −25 °C. For all three temperatures, the mean of the median mass diameters during sampling was between 40 and 50 nm, which should be representative for sizes >0.1 μm. A shows the desorption profiles of C₁₀H₁₆O₄, B of C₁₀H₂₀O₄, and C of C₁₈H₂₄O₉. Fits for the monomer, dimer fragment, and background signal are indicated for a single temperature on each profile.

![Fig. 5](image)

**Fig. 5.** Overview of the competing processes and their temperature dependence and comparison between the relative (rel.) gas- and particle-phase contribution of the ozonolysis product group C₁₀H₁₆O₄−₉. Left shows the normalized relative contribution of the different oxygenated molecules within the gas phase, while Right shows the normalized relative contribution of the same compounds within the particle phase, inferred from monomer desorption fits.
Experiments were conducted as follows: At 38% relative humidity, with no SO₂ and no NOₓ present in the chamber, stable ozone concentrations of 30–40 parts per billion were established. Under dark conditions (i.e., without any additional OH radical production mechanism except from the ozonolysis itself), a high-voltage field cage inside the chamber was switched on to perform neutral experiments first. Injection of α-pinene initiated the ozonolysis reaction and the subsequent formation of particles. After steady-state α-pinene concentrations were reached and particle growth was measured up to at least 10 nm, the high-voltage field was switched off. Ions now present in the chamber led to a significant increase in nucleation rate (7). Therefore, two growth rate measurements could eventually be performed, as the size distribution will show two growing particle populations. The second measurement was independent of changing gas concentrations, as steady-state was reached during the neutral stage. As no significant effect on growth due to different ionization levels was found, all measurements are treated equally.

The key particle-size distribution and growth-rate measurements of this study were performed with a DMA train (23). The sheath flows of the six DMAs were conditioned to chamber temperature to avoid possible particle evaporation during particle sizing. A 50% systematic uncertainty was assumed on the apparent particle-growth rates inferred by the appearance time method when comparing the values to growth rates from pure condensation. This also covered uncertainties of possible evaporation during the measurement procedure. Details can be found in SI Appendix. Gas-phase composition was measured by two complementary mass spectrometers using different ionization techniques (21, 22). For molecular ion signals observed in both instruments, the stronger signal was used to account for a reduced charging efficiency by any of the two ionization chemistries. For more details, see SI Appendix. Particle-phase composition was measured with a FIGAERO-CIMS (24), which accumulated particles for 30 min on a filter inlet kept at chamber temperature. As all experiments were started with a particle-free chamber, all collected particles originated from new particle formation under similar experimental conditions. The accumulated particles were thermally desorbed by heating the filter, and their composition was analyzed by the connected mass spectrometer. Details can be found in SI Appendix.

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