Aging of secondary organic aerosol generated from the ozonolysis of \( \alpha \)-pinene: effects of ozone, light and temperature

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Abstract

A series of experiments was conducted in the CESAM simulation chamber to investigate the evolution of the physical and chemical properties of secondary organic aerosol (SOA) during different forcing. The present experiments represent a first attempt to comprehensively investigate the influence of oxidative processing, photochemistry, and diurnal temperature cycling upon SOA properties. SOA generated from the ozonolysis of α-pinene were exposed to (1) elevated ozone concentrations, (2) light (under controlled temperature conditions), or (3) light and heat (6°C light-induced temperature increase), and the resultant changes in SOA optical properties (i.e. absorption and scattering), hygroscopicity and chemical composition were measured using a suite of instrumentation interfaced to the CESAM chamber. The complex refractive index (CRI) was derived from integrated nephelometer measurements at 525 nm wavelength, using Mie scattering calculations and measured number size distributions. The particle size growth factor (GF) was measured with a hygroscopic tandem differential mobility analyzer (H-TDMA). An aerosol mass spectrometer (AMS) was used for the determination of the $f_{44}/f_{43}$ and O : C ratio of the particles bulk.

No change in SOA size or chemical composition was observed during O$_3$ and light exposure at constant temperature; in addition, GF and CRI of the SOA remained constant with forcing. By contrast, illumination of the SOA in the absence of temperature control led to an increase in the real part of the CRI from 1.35 ($\pm$0.03) to 1.49 ($\pm$0.03), an increase of the GF from 1.04 ($\pm$0.02) to 1.14 ($\pm$0.02) and an increase of the $f_{44}/f_{43}$ ratio from 1.73 ($\pm$0.03) to 2.23 ($\pm$0.03). These surprising results suggest that SOA properties may be governed more by local temperature fluctuations than by oxidative processing and photochemistry.
1 Introduction

Atmospheric aerosols influence climate directly by altering the absorption and scattering of solar and terrestrial radiation (Haywood and Ramaswamy, 1998) and indirectly by changing cloud properties (Lohmann and Feichter, 2005). One of the major uncertainties in estimating the aerosol radiative effect is associated with the contribution of secondary organic aerosols (SOA). SOA are formed by condensation of species formed during gas-phase oxidation of volatile organic compounds (VOCs) and are a major constituent of atmospheric organic aerosol (Kanakidou et al., 2005; Turpin and Huntzicker, 1995; Zhang et al., 2007). During their atmospheric lifetime, which ranges from 48–72 h (Wagstrom and Pandis, 2009), their physical and chemical properties do not stay constant but rather evolve in response to local atmospheric conditions. For example, studies have shown that SOA can grow by condensation of low-volatility oxidized species (Ellison et al., 1999) and by cloud processing (Cocker et al., 2001; Ervens and Volkamer, 2010; Volkamer et al., 2009). SOA can also be oxidized by gas-phase oxidants (Kalberer, 2004; Gao et al., 2004), undergo chemical reactions in the particle phase (Kalberer, 2004; Gao et al., 2004) or partially evaporate (Warren et al., 2009). These changes in aerosol properties have also been observed in the field: for example, more oxidized, less volatile and more hygroscopic SOA are typically observed in remote areas as a result of continuous aging in the atmosphere (Jimenez et al., 2009; Ng et al., 2011; Rudich et al., 2007). Because SOA contains a wide variety of organic compounds, which vary in terms of their size, structure, functionality, and oxidation state (Kroll and Seinfeld, 2008; Jimenez et al., 2009; de Gouw et al., 2005; Hallquist et al., 2009), the processes associated with SOA aging are very complex.

Experiments performed in simulation chambers have significantly improved our understanding of SOA aging processes (Donahue et al., 2012; Qi et al., 2012; Yasmeen et al., 2012). In order to provide modelers with accurate parameters for SOA aging in the atmosphere, these experiments must be atmospherically relevant (Kourtchev et al., 2014). While the O:C ratio of laboratory-generated SOA is similar to that of freshly generated SOA, their physical and chemical properties may differ significantly due to aging processes that occur in the atmosphere.
formed ambient SOA, it is generally lower than that of aged ambient SOA (Ng et al., 2010). In addition, while the representation of SOA in chemical transport models based on parameterization of chamber data showed a good agreement with nighttime SOA concentrations (in a rural site near Rotterdam, the Netherlands), an underprediction of SOA concentrations occurred during the day (Li et al., 2013).

Research has shown that the oxidative aging of SOA in the atmosphere has a major influence on its properties. For example, studies have shown that highly oxygenated organic particles are likely to have a higher hygroscopicity and CCN activity than freshly emitted particles, due to the increased polarity and solubility of their constituents (Massoli et al., 2010; Jimenez et al., 2009; Chang et al., 2010; Duplissy et al., 2011). Oxidative aging has also been shown to lead to changes in the real part of the complex refractive index (CRI) of SOA (Cappa et al., 2011; Lambe et al., 2013; Flores et al., 2014), and to an increase in its UV absorption via the formation of additional carbonyl compounds and oligomeric products (Sareen et al., 2013; Nozière and Esteve, 2005; Shapiro et al., 2009; Lambe et al., 2013). To date, laboratory studies have primarily focused on oxidative aging mediated by the heterogeneous reactive uptake of OH radical (Rudich et al., 2007; George and Abbatt, 2010; Smith et al., 2009). Oxidative aging by O₃, by contrast, has received less attention.

Light exposure has also been shown to influence SOA properties: by photodissociation of molecules such as carbonyls and organic peroxides, either in the gas or the particle phase, it has been shown to induce a decrease in SOA mass concentration (Kroll et al., 2006; Bateman et al., 2011). There are also indications of significant photolytic processing of carbonyl compounds in aerosols during long-range transport (Hawkins and Russell, 2010). In addition, laboratory studies have revealed that photochemical processes alter the chemical composition of SOA (Tritscher et al., 2011; Cappa et al., 2011; Qi et al., 2012; George and Abbatt, 2010; Donahue et al., 2012) and modify its hydrophilicity (Tritscher et al., 2011; George and Abbatt, 2010; George et al., 2009), optical properties (Cappa et al., 2011) and volatility (Tritscher et al., 2011). In these studies, H₂O₂ was used as a photolytic OH precursor. Since this OH source requires
the use of UV light, these experiments did not allow for the individual effects of heterogeneous reactions by OH and direct photolysis on SOA properties to be distinguished.

SOA properties can also be affected by local temperature variations via the evaporation of volatile products. A number of studies have used thermodenuder-based techniques to investigate the volatility of SOA and to determine the effect of temperature on SOA properties (Asa-Awuku et al., 2009; Huffman et al., 2009; Cappa and Wilson, 2011). Volatile Tandem Differential Mobility analyzers (V-TDMA) have been used to measure the shrinkage of monodisperse particles after heating (Cappa and Wilson, 2011; Salo et al., 2011). In these studies, particles were exposed to elevated temperatures (up to 300 °C) for a short time. It has been shown, however, that SOA exhibits a significantly slower response to changes in temperature than that predicted by models for liquid droplets (Cappa and Wilson, 2011). Recently, it has been suggested that SOA could be in an amorphous semi-solid or amorphous solid (glassy) state under dry conditions (Renbaum-Wolff et al., 2013; Saukko et al., 2012; Denjean et al., 2014b), which could limit the volatilization kinetics of the aerosol.

This study focuses on the SOA formed from the ozonolysis of α-pinene, which is an important source of SOA on both regional and global scales (Guenther et al., 1995; Hallquist et al., 2009). In our companion paper (Denjean et al., 2014b), we explored the evolution of the physical, chemical, optical and hygroscopic properties of α-pinene-O₃ SOA during the first hours after its formation. However, Wang et al. (2011) have shown that particle lifetime can vary from 10 h to 4 days in the CESAM simulation chamber. CESAM is a powerful tool for the study of SOA over longer timescales corresponding to their lifetime in the atmosphere (Yasmeen et al., 2012). In the present work, we investigate in the same chamber the effects of (i) additional ozone exposure, (ii) light exposure and (iii) temperature variation on the chemical composition, hygroscopicity, and optical properties of α-pinene-O₃ SOA, over timescales reaching 20 h.
2 Methods

2.1 CESAM atmospheric simulation chamber

The present experiments were performed in the CESAM (French acronym for Experimental Multiphasic Atmospheric Simulation Chamber) atmospheric simulation chamber, which has previously been described in detail by Wang et al. (2011). In brief, CESAM is a stainless-steel chamber with a volume of 4.2 m$^3$. Chamber illumination is accomplished using the borosilicate-filtered output of three high-pressure arc xenon lamps (4 kW, XPO 4000 W/HS, OSRAM), which provides a good reproduction of the solar energy distribution at the Earth’s surface over the 290–700 nm wavelength region (Fig. S1 in the Supplement). The inner walls of the chamber are polished in order to provide good reflection inside the chamber and thus enhance the radiation homogeneity. During our experiments, the NO$_2$ photolysis frequency $J_{\text{NO}_2}$ within the chamber was approximately $3 \times 10^{-3}$ s$^{-1}$, which corresponds to a solar zenith angle of $\sim 70^\circ$ (Carter et al., 2005). The simulation chamber was maintained at room temperature (±1 °C) using a refrigerating liquid (70 % water/30 % ethylene glycol), which was circulated in the double walls of the chamber. Temperature and relative humidity are monitored with a transmitter (HMP234, Vaisala) equipped with a thin-film capacitive humidity sensor (HUMICAP, Vaisala). During our experiments, the temperature accuracy is ±0.1 °C at 20°C and the RH accuracy is ±1.9 % (up to 90 % RH).

2.2 Experimental details

Prior to each experiment, the chamber was evacuated to a secondary vacuum (typical pressure $\sim 4 \times 10^{-4}$ mbar) and kept under vacuum overnight. The chamber was then filled to atmospheric pressure with a mixture of 200 mbar of Oxygen (Air Liquide, Alphagaz class 1, purity 99.9 %) and 800 mbar of Nitrogen produced from the evaporation of a pressurized liquid nitrogen tank (Messer, purity > 99.995 %, H$_2$O < 5 ppm). The background conditions were typically: particles concentration < 0.1 µg m$^{-3}$, ozone
mixing ratio < 5 ppb, gas phase organics < 5 ppb and relative humidity < 1 %. To avoid contamination, a slight overpressure of about 5 mbar with respect to the atmospheric pressure was maintained during each experiment by adding nitrogen (Messer, purity > 99.995 %, H₂O < 5 ppm).

All aging experiments were carried out under the same initial conditions with α-pinene and ozone reacting in the dark, with no seeds neither OH scavenger. O₃ was generated in an O₂ flow using a commercial dielectric ozone generator (MBT 802N, Messtechnik GmbH, Stahnsdorf, Germany) and introduced to the chamber through an injection port. Quantification of α-pinene was performed by evaporating precisely measured amounts of the terpene into a glass bulb held under vacuum. When the O₃ concentration within the chamber reached ∼ 250 ppb, α-pinene (Aldrich, 98 %) was flushed from the bulb into the chamber in a flow of oxygen to a concentration of ∼ 200 ppb within the chamber. In all experiments, SOA formed directly after α-pinene injection, and its precursors were essentially consumed after 4 h of reaction.

In our companion paper, we observed changes in the oxidative degree and optical properties during the formation of SOA, but these changes ceased after 9 h of reaction (Denjean et al., 2014b). In the present study, therefore, SOA was allowed to evolve for 14 h prior to simulate atmospheric processing, which was in turn conducted over a period of 6 h. Three different aging regimes were applied after 14 h of α-pinene ozonolysis: (1) 700 ppb of ozone was introduced into the simulation chamber, (2) the SOA was exposed to light for 6 h in the absence of temperature control, during which time the temperature increased by ∼ 6 °C, and (3) the SOA was exposed to light for 6 h under controlled temperature conditions (±1 °C). For comparison purposes, control experiments were performed: in these experiments, SOA was allowed to remain in the chamber for 20 h under dark conditions. The specific experimental conditions associated with each aging regime are shown in Table 1.

During these experiments, the concentrations of α-pinene and ozone were monitored using a Fourier-transform infrared spectrometer (FTIR) from Bruker GmbH (Ettlingen, Germany) coupled to a multi-reflection cell with an optical path of 192 m. Ozone was
also measured with a commercial instrument (Horiba APOA 370, Kyoto, Japan), with a detection limit of 0.2 ppb and a precision of 0.1 ppb.

2.3 Measurement of SOA properties

2.3.1 Size distribution

SOA number size distributions between 14 and 505 nm were monitored using a scanning mobility particle sizer (SMPS; DMA Model 3080, CPC Model 3010; TSI) operated at flow rates of 3/0.3 Lpm (sheath flow/aerosol sample flow). Instrument calibration was conducted using polystyrene latex spheres (PSL) (Duke Scientific). Since the PSL diameters measured during calibration were ~10% larger than the certified PSL diameters (for 100 nm PSL samples), a correction factor was applied to all measurements. Corrections for particle loss by diffusion in the SMPS tubing and the contribution of multiply charged particles were made using the SMPS software (Aerosol Instrument Manager, version 9, TSI). The number size distribution was used to obtain SOA mass concentrations, assuming homogeneous spherical particles and an effective density of 1.2 g m\(^{-3}\), as determined by Shilling et al. (2008), Saathoff et al. (2009) and Denjean et al. (2014b).

2.3.2 Chemical composition

SOA chemical composition was analyzed using a high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS, Aerodyne) (DeCarlo et al., 2006). Instrumental and data treatment details are given in our companion paper (Denjean et al., 2014b), and thus will be described only briefly here. The HR-ToF-AMS was used under standard conditions (vaporizer at 600 °C and electron ionization at 70 eV). The instrument was switched between two modalities: a single-reflectron configuration (V-mode), which offers higher sensitivity but lower resolving power (up to ~2100 at \(m/z\) 200), and a double-reflectron configuration (W-mode), which provides a higher resolving power (up to ~2100 at \(m/z\) 200).
(up to ~ 4300 at \( m/z \) 200) but a lower sensitivity (De Carlo et al., 2006). Default collection efficiencies (CE) and relative ionization efficiencies (RIE) were used for quantification of SOA composition. High-resolution analysis was performed using V-mode data, by integrating each \( C_xH_yO_z \) ion in the mass range 12–180 \( m/z \); W-mode data were used only to check for possible interferences. The method is based on the relative ionization efficiencies of molecules containing C, H and O atoms. The sum of the ion signal intensities from all fragments was used to estimate the \( O: C \) ratio of the SOA, and thus its degree of oxidation. Air interferences were removed by adjusting the fragmentation table (Aiken et al., 2007; Allan et al., 2004). Measurement uncertainties were estimated to be ±30 %, as determined by Aiken et al. (2007).

### 2.3.3 Optical properties

SOA optical properties were characterized by combining data obtained from an integrating nephelometer (Model M9003, Ecotech), a spectral aethalometer (Model AE31, Magee Scientific) and the SMPS described above. The integrating nephelometer measured the scattering coefficient (\( \sigma_{\text{scat}} \)) at 525 nm wavelength, as well as the temperature and relative humidity of the incoming airflow, and was calibrated prior to the experiments using filtered air and CO\(_2\). The nephelometer collected light only from particles at scattering angles between 10 and 170°. Measured scattering coefficients were corrected for this angular truncation using Mie calculations, which were performed using the measured SMPS size distribution. The aethalometer measured the SOA absorption coefficient at 7 wavelengths (370, 470, 520, 590, 660, 880 and 950 nm) by measuring the increase in attenuation of transmitted light through its quartz fiber filter as a function of particle exposure time. In order to avoid artifacts from the adsorption of ozone and VOCs on the filter, a charcoal denuder was installed upstream of the aethalometer (Weingartner et al., 2003). Measurements of light attenuation were corrected for aerosol scattering effects according to the method described by Collaud Coen et al. (2010) and using the parameters obtained by Denjean et al. (2014a, b) for \( \alpha \)-pinene-O\(_3\) SOA.
These measurements were ultimately used to calculate the real and imaginary parts of the CRI, which together describe the scattering and absorbing characteristics of SOA. The CRI retrieval procedure employed has been described and validated in Denjean et al. (2014a). In brief, the CRI of SOA was retrieved at 525 nm by comparing the measured scattering and absorbing coefficients (respectively $\sigma_{\text{scat}}$ and $\sigma_{\text{abs}}$) with those obtained from Mie scattering calculations (Bohren and Huffman, 1983) performed using the measured number size distribution. The absolute error associated with the real CRI was $\pm 0.03$ (Denjean et al., 2014b).

### 2.3.4 Hygroscopic properties

SOA hygroscopic properties were analyzed with a custom-built hygroscopic Hygroscopic Tandem Differential Mobility Analyzer (H-TDMA), which is described in detail in Denjean et al. (2014a). The first DMA was used to select particles with a mobility diameter of 200 nm, which were then humidified at a constant RH of 90±1 % (residence time $\sim 15$ s). The second DMA, which was coupled to a CPC, measured the humidified size distribution (with respect to the mobility diameter). Both DMAs were calibrated using monodisperse PSL particles (Duke Scientific) with size diameters of 70, 100, 200 and 300 and 500 nm. These size distributions were fitted to lognormal size distributions to obtain the dry geometric mean diameter, $D_{p,m}$ (dry), and the geometric mean diameter of the humidified aerosol, $D_{p,m}$ (90 % RH). These values were used to obtain the size growth factor (GF), which is defined as the ratio of $D_{p,m}$ (90 % RH) to $D_{p,m}$ (dry). The uncertainties in the calculated GF are associated with uncertainties in particle size distributions arising from DMA classification and calibration, as well as with uncertainties in the estimation of $D_{p,m}$ from size distributions, and are estimated to $\pm 0.02$. Prior to each experiment, the experimental set-up was validated by comparing measurements of the GF of ammonium sulfate particles to those predicted by Köhler theory (Denjean et al., 2014a).
2.4 Modeling SOA formation and aging

In order to assist in the interpretation of experimental results, the α-pinene ozonolysis experiments were simulated using a box model that included the Master Chemical Mechanism (MCM) (version 3.1) (Saunders et al., 2003) and the Generator for Explicit Chemistry and Kinetics of Organics in the Atmosphere (GECKO-A) (Aumont et al., 2005). The MCM v3.1 oxidation scheme for α-pinene contains 329 organic species and 973 reactions. The GECKO-A chemical scheme for α-pinene oxidation, which provides a more detailed description of the gaseous oxidation of organic species and takes into account minor reaction pathways not considered by the MCM, involves $5.7 \times 10^4$ organic species reacting according to $1.7 \times 10^6$ reactions (Valorso et al., 2011). Both models simulate gas/particle partitioning in terms of equilibria between the gas phase and an ideal liquid homogeneous condensed phase (Camredon et al., 2007).

Vapor pressures and boiling points for secondary organic species were estimated using the methods developed by Nannoolal et al. (2004, 2008), as they have been shown to provide the most reliable estimates for the purpose of SOA formation (Barley and McFiggans, 2010). Accretion, oxidation and photolysis in the condensed phase were not taken into account in this investigation in the models.

Time integration of the chemical schemes was solved using the TWOSTEP solver (Verwer et al., 1994, 1996). The gas/particle partitioning module was solved using the iterative method described in Pankow (2008). Simulations were initialized at a time point corresponding to α-pinene injection. This injection was implemented in the model as a constant flux over the injection time period that reproduced the observed concentration. Dilution within the chamber (arising from the periodical injection of N$_2$ to compensate for instrumental sampling flows) was described as a measurement-constrained first-order process and applied in the simulations to both gas and aerosols. The temperature used in the simulations was that observed in the CESAM chamber during the experiments. Ozone wall loss is significant in the CESAM chamber (Wang...
et al., 2011). The ozone loss rate employed in the simulations, therefore, was adjusted for each experiment in order to reproduce its measured decay.

3 Results

3.1 Changes in SOA size distribution during simulated atmospheric processing

Figure 1 shows the temporal evolution of SOA mass concentrations (normalized to the mass concentration at the beginning of each aging experiment and corrected for dilution within the chamber) during and after simulated atmospheric processing. As shown in this figure, the SOA mass concentrations in the control experiments decreased by ∼15% over the 6 h duration of the experiment. Since these measurements were performed after the total consumption of O$_3$, this mass decrease cannot be attributed to O$_3$-induced fragmentation reactions. It is thus probable that the SOA mass decrease observed in the control experiments arose via losses of particle and gaseous compounds to the chamber walls. Similar behavior was observed when SOA was exposed to ozone and to light under controlled temperature conditions, which suggests that these forcing did not lead to significant fragmentation or functionalization. By contrast, exposure of SOA to light and increasing temperature led to a 40% loss in total SOA mass concentration and, as shown in Fig. 2, to a change in the SOA number size distribution: as the temperature increased from 20 to 26°C, the geometric mean diameter of the normalized SOA number size distribution decreased from 286 to 249 nm, which also indicates that significant evaporation of SOA particles occurred.

3.2 Changes in SOA optical and hygroscopic properties during simulated atmospheric processing

Since the CRI is related to the aerosol chemical composition, density, molecular weight and polarization (Liu and Daum, 2008), it was thus expected to be influenced by atmospheric processing. The imaginary part of the SOA CRI over the 370–950 nm
wavelength range is shown in Fig. 3. Its value was almost zero at all the wavelengths studied, which indicates that SOA, even after simulated atmospheric processing, has a pure scattering effect in the visible to near-UV region. Interestingly, while one might have expected the formation of chromophores during the aging process, these results indicate that this phenomenon occurred only weakly and/or that the specific absorption of any potential aging products was not significant. The influence of atmospheric processing upon the real part of the SOA CRI is shown in Fig. 4. No discernible change in real CRI was observed when the SOA was exposed to O$_3$ or to light at constant temperature. By contrast, a significant increase in the real CRI was observed (1.35 ($\pm$0.03) to 1.49 ($\pm$0.03)) after photolytic forcing with increasing temperature.

The hygroscopic properties of SOA were studied by measuring the GF at a constant relative humidity (90±1%). As shown in Fig. 5, the GF remained constant ($\sim$ 1.04 ± 0.02) both during O$_3$ exposure and during photochemical aging at constant temperature. These results are consistent with our observations of constant mass concentration and CRI of SOA, and confirm that no significant processing of the SOA occurred under these conditions. In contrast, a significant increase in the GF, from 1.04 (±0.02) to 1.14 (±0.02), was observed with increasing temperature.

### 3.3 Changes in SOA chemical composition during simulated atmospheric processing

The results presented in the previous two sections suggest that the exposure of the aging of SOA to light and increasing temperature resulted in changes in its physical, optical and hygroscopic properties, which are likely to be linked to changes in its chemical composition. Indeed, as shown in Fig. 6a, the increase in the O : C ratio of bulk SOA during its exposure to light and increasing temperature (0.55 (±0.16) to 0.59 (±0.18)) was much larger than that observed for SOA under control conditions. The increase in the O : C ratio was, however, within the measurement uncertainties (±30 %) estimated from Aiken et al. (2007). This uncertainties may be overestimated compared to the experimental variability and even experimental reproducibility observed in this study.
In fact, we estimated the experimental uncertainties to be ±0.01 from the standard deviation of the experimental values before the forcing.

The fragments \( f_{44} \), defined as the ratio of the \( m/z \) 44 (a major fragment of organic acids and hydroperoxides) signal to the total organic aerosol signal, and \( f_{43} \), defined as the ratio of \( m/z \) 43 (associated with less-oxygenated groups, e.g. aldehydes and alcohols) signal to the total organic aerosol signal, \( (m/z \) 43 is associated to lower oxygenated groups, such as aldehydes and alcohols) have been widely used in laboratory and field studies as indicators of SOA functionality and degree of oxidation (Ng et al., 2010; Alfarra et al., 2013; Poulain et al., 2010; Pfaffenberger et al., 2013). As shown in Fig. 6b, in the experiment performed under illumination but in the absence of temperature control, the \( f_{44}/f_{43} \) ratio of SOA increased from 1.73 (±0.03) to 2.23 (±0.03) as the chamber temperature increased. This result implies a temperature-mediated increase in particle-phase oxidized species. By contrast, only a small increase in the \( f_{44}/f_{43} \) ratio (from 1.93 (±0.03) to 2.03 (±0.03)) was observed during the control experiment, which implies that the SOA composition in this experiment remained relatively constant.

### 3.4 Effect of phase partitioning on SOA properties

During the aging of SOA by photolysis with increasing temperature, we observed a decrease of the mass concentration (Fig. 1) which was associated to an increase of the \( f_{44}/f_{43} \) ratio of bulk particles (Fig. 6b). These evolutions may result from possible shifts of some semi-volatile organics to the gas phase as temperature increases. A second possible explanation would be the photochemical reactions that occur in the condensed phase combined to the evaporation of less oxidized compounds. In this case, photolysis could lead to a loss of semi-volatile and less oxidized compounds in the particle phase due to the fragmentation of condensed-phase species (Donahue et al., 2012; Henry and Donahue, 2012).

In order to investigate if the increase in temperature can explain the SOA mass decrease, the SOA formation and evolution were modelled with the detailed chemical schemes GECKO-A and MCM. In these models, the simulated SOA formation started...
at the initial step of α-pinene ozonolysis. The effects of temperature on phase partitioning were simulated with both models. The effects of photochemical reactions and heterogeneous reactions by O₃ were ignored in these simulations. As shown in Fig. 7 for the initial SOA formation period, the time profiles of α-pinene and ozone concentrations were well reproduced by both models within the measurement uncertainties. Although the temporal profile of the SOA formation was also well reproduced by the models, the simulated mass concentrations of SOA obtained using the GECKO-A and MCM models (302 and 337 µg m⁻³, respectively) were higher than those observed in the experiment (130 µg m⁻³). This overestimation of the simulated SOA mass concentration might be due to an underestimation of the vapor pressure with the Nannoolal method used in this study. In addition, some oxidation processes occurring in the aerosol phase and leading to fragmentation may not be implemented in the model.

Four experiments, each of which exhibited different temperature increases, were simulated using these models. SOA mass concentration loss was calculated as the difference between the mass concentration before and after each forcing. For the simulations, only the effects of the temperature on phase partitioning were simulated with both models. The effects of photochemical reaction on the gas and particle phase were omitted. A comparison between the observed and modelled loss of SOA mass concentration under these temperature conditions is presented in Fig. 8a. As shown in this figure, both the temporal profile and magnitude of SOA mass loss observed in these experiments were well-reproduced by the GECKO-A and MCM models. This finding suggests that the observed heating-induced loss in SOA mass concentration can be explained by the temperature-dependent gas–particle phase partitioning of the semivolatile components of SOA.

The measured and simulated O : C ratios for these experiments are shown in Fig. 8b. Before the forcing, both models predict an O : C ratio of 0.49, which is within the uncertainty of the experimental value (0.55 ± 0.16) obtained with the AMS. During the aging experiments, the O : C ratio simulated with the two models increased with increasing temperature: for example, experiment E200411, which showed the highest temperature
increase (+6.5°C: red in Fig. 8c), also showed the highest increase in simulated O:C ratio (from 0.49 to 0.51); by contrast, experiment E060512 exhibited constant values for both temperature and simulated O:C ratio (yellow in Fig. 8b and c). These results suggest that the changes in SOA physical and chemical properties observed in these experiments resulted from the heating-induced evaporation of semi-volatile and less oxidized SOA species, which in turn modified the optical and hygroscopic properties of the condensed phase.

4 Discussion

Our results have shown that exposure of SOA to increasing temperature enhances its hydrophilicity, degree of oxidation, and scattering properties. By contrast, the few previous studies investigating the changes in physical and chemical properties of α-pinene-O₃ SOA as a function of temperature have largely found little effect: Cappa and Wilson (2011) observed a reduction in the total mass concentration but no change in the mass spectra of α-pinene-O₃ SOA after heating at 170°C, and Kim and Paulson (2013) showed no significant change over temperatures ranging from 23 to 86°C in the real CRI of SOA. Using AMS data, Huffman et al. (2009) observed an increase in SOA oxygen content during SOA evaporation. This trend is in good agreement with the increase in the CO₂⁺ fragment (m/z 44) with SOA evaporation observed by (Kostenidou et al., 2009). Finally, Warren et al. (2009) showed that the GF of SOA decreased with decreasing temperature from 27 to 5°C. These disagreements with the present study may be attributable to two differences in experimental conditions. First, in the previous studies, SOA volatility was measured only several minutes after its formation. However, since the chemical composition of SOA has been observed to vary significantly during its formation (Shilling et al., 2008; Chhabra et al., 2010; Denjean et al., 2014b), it is likely that the volatility of the condensed species also varies during this time period. In the present study, by contrast, SOA volatility was measured only after 14 h of formation (i.e. at a point where the composition of the semivolatile component of SOA was likely
stable). Second, in most of the previous studies, temperature variation was performed using a thermodenuder. The timescale for aerosol evaporation within thermodenuders (∼16 s) is significantly lower than in simulation chambers (several hours), and, owing to mass transfer limitations, may not be long enough for SOA evaporation to occur (Lee et al., 2011).

Surprisingly, our results suggest that exposure of α-pinene SOA to light and O₃ does not significantly change its chemical, hygroscopic and optical properties. These observations suggest that, under our experimental conditions, neither significant photolysis nor ozonolysis of the particle-phase products occurred.

Several previous studies have reported a decline in α-pinene-O₃ SOA mass after irradiation under exposure to low OH concentrations (Donahue et al., 2012; Henry and Donahue, 2012). These authors used UV lights (∼360 nm) to initiate photochemistry; by contrast, a more realistic reproduction of the solar energy distribution at the Earth’s surface was used in our study (Wang et al., 2011). In a test of the effect of light source upon SOA aging, Donahue et al. (2012) observed a monotonic decrease in SOA mass during aging with 360 nm UV lights, but an increase in SOA mass during aging with sunlight or quasi-solar lamps. These observations can be explained by the fact that some oxygenated organics in the SOA condensed phase, such as carbonyls or peroxides, undergo photodissociation by photolysis at the specific wavelength 360 nm. Together, these results highlight the strong influence of the light source on the photodissociation of SOA, and underscore the utility of experiments performed under realistic illumination conditions.

In addition, recent literature has highlighted that α-pinene-O₃ SOA is not composed of a homogeneous chemical mixture. For example, when Maurin et al. (2014) exposed α-pinene-O₃ SOA to high concentrations of O₃, they observed a constant particle-phase concentration of verbenone, an unsaturated oxygenated product. Since this molecule would have been expected to be consumed by heterogeneous reaction with O₃, and since heterogeneous particle-phase reactions are predicted to occur mainly at the particle surface of the particles (Shiraiwa et al., 2013), these authors postulated
that reaction of O$_3$ with verbenone was kinetically limited by the diffusion of buried verbenone to the particle surface. This hypothesis is also supported by our observation that $\alpha$-pinene-O$_3$ SOA exhibits a core-shell structure, with less oxidized species at its surface (Denjean et al., 2014b). The effect of the forcing would depend not only on the chemical composition of the bulk particle, but also on the chemical surface and the phase of the particle.

5 Conclusions

We have demonstrated that the size, chemical composition, hygroscopicity and optical properties of $\alpha$-pinene-O$_3$ SOA change dramatically in response to relatively minor increases in temperature ($\sim 6^\circ$C). These changes, in turn, have implications for the role that SOA plays in climate. For example, the volatilization-induced increase in the real part of the SOA CRI is likely to enhance the direct radiative effect of SOA by increasing its ability to scatter radiation. The direct radiative effect of SOA can be parameterized using the mass extinction efficiency $k_{\text{ext}}$ at 525 nm, which is defined as the ratio of the measured $\sigma_{\text{scat}}$ at 525 nm to the SOA mass concentration. In our companion paper (Denjean et al., 2014b), we have estimated a value of $k_{\text{ext}} = 1.7 \pm 0.5$ m$^2$ g$^{-1}$ for unprocessed SOA. In the present experiments, we estimate a value of $k_{\text{ext}} = 2.8 \pm 0.5$ m$^2$ g$^{-1}$ for SOA exposed to light and temperature increase, which implies that this aging regime led to a $\sim 40\%$ enhancement in this parameter. We attribute this enhancement to both the decrease in SOA particle size and the increase in the scattering component of the CRI upon heating.

As shown in this work, exposure of SOA to increasing temperature also leads to an increase in SOA hygroscopicity: the SOA GF increased from 1.04 to 1.14 as a result of this forcing. In order to estimate the influence of these changes in hygroscopicity on the direct radiative effect of SOA, we made the simplified assumption that SOA exhibits a homogeneous mixing state with water. We used Mie scattering calculations for homogeneous spheres to determine $\sigma_{\text{scat}}$ at 90\% RH both from the measured GF...
and the CRI. The CRI calculations were based on volume-weighted CRI of values of α-pinene-O$_3$ SOA and water. We estimate that $k_{\text{ext}}(90\% \text{ RH}) = 2.83 \pm 0.70 \text{ m}^2 \text{ g}^{-1}$ after SOA volatilization, which is significantly higher than the value of $k_{\text{ext}}(90\% \text{ RH}) = 2.36 \pm 0.70 \text{ m}^2 \text{ g}^{-1}$ calculated in Denjean et al. (2014b) for unprocessed SOA. Since we have shown that volatilization of condensed-phase species takes place over a range of temperatures consistent with diurnal fluctuations, we suggest that these changes in $k_{\text{ext}}$ are representative of the diurnal evolution of SOA during its lifetime in the atmosphere.

Our results also suggest that α-pinene-O$_3$ SOA is quite insensitive to light- and ozone-induced aging under our experimental conditions, most likely as a result of kinetic limitations imposed by its physical state (semi-solid/glassy). Several recent studies have reported that SOA undergoes a transition from a more glassy state to a more liquid state with increasing RH, (Saukko et al., 2012; Renbaum-Wolff et al., 2013; Denjean et al., 2014b). On the basis of viscosity data and the Stokes–Einstein equation, Renbaum-Wolff et al. (2013) estimated that the change in α-pinene-O$_3$ SOA viscosity associated with its transition from a solid to a semi-solid state increases the bulk diffusion coefficient of particles by six orders of magnitude (from $< 10^{-17} \text{ cm}^2 \text{ s}^{-1}$ to $10^{-11} \text{ cm}^2 \text{ s}^{-1}$). Since an increase in bulk diffusion coefficient would be expected to be accompanied by an increase in particle reactivity, we suggest that future studies examine the effect of ozone and light exposure on SOA properties under humidified conditions (i.e. RH > 30\%).

The Supplement related to this article is available online at doi:10.5194/acpd-14-22437-2014-supplement.

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Table 1. Summary of experimental conditions.

<table>
<thead>
<tr>
<th>Type of forcing</th>
<th>Run</th>
<th>Cm_{t=0}^{a} (µg m^{-3})</th>
<th>[O_3]_{t=0}^{b} (ppb)</th>
<th>Lights on (min)</th>
<th>T_{t=0}/T_{t=6h}^{c} (°C)</th>
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<td>20.4/21.3</td>
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<td>0</td>
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<td>with increasing temperature</td>
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<td>19.8/20.1</td>
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</table>

^{a} Aerosol mass concentration estimated from the aerosol volume concentration corrected from dilution and by assuming a density of 1.2 g cm^{-3}.

^{b} Ozone concentrations determined using FTIR spectroscopy.

^{c} Temperature before forcing/temperature after forcing.
Figure 1. Temporal evolution of SOA mass concentrations (normalized to the mass concentration at the beginning of each forcing) during and after simulated atmospheric processing. In the “control experiment” (blue symbols), SOA was left to evolve in the chamber under dark conditions. In the “O₃ aging” experiments (green), SOA was exposed to an excess of ozone (∼ 700 ppb) under constant temperature conditions. In the “photochemical aging” experiments, SOA was exposed to light for 6 h, either under constant temperature conditions (yellow) or with light-induced heating (red). Here, the initial time $t = 0$ corresponds to the beginning of simulated aging, which was commenced after SOA was allowed to form and stabilize for 14 h.
Figure 2. Temporal evolution of SOA number size distribution (normalized to the total number concentration) (a) during photochemical aging in the absence of temperature control (Experiment E120312) and (b) during a control experiment (Experiment E160414).
Figure 3. Wavelength dependence of the imaginary part of the complex refractive index of SOA before and after simulated atmospheric processing. In the “control experiment” (blue symbols), SOA was left to evolve in the chamber under dark conditions. In the “O_3 aging” experiments (green), SOA was exposed to an excess of ozone (∼700 ppb) under constant temperature conditions. In the “photochemical aging” experiments, SOA was exposed to light for 6 h, either under constant temperature conditions (yellow) or with light-induced heating (red).
Figure 4. Measurements of the real part of the complex refractive index of SOA before and after simulated atmospheric processing. In the “control experiment” (blue symbols), SOA was left to evolve in the chamber under dark conditions. In the “O₃ aging” experiments (green), SOA was exposed to an excess of ozone (∼700 ppb) under constant temperature conditions. In the “photochemical aging” experiments, SOA was exposed to light for 6 h, either under constant temperature conditions (yellow) or with light-induced heating (red). Here, the initial time \( t = 0 \) corresponds to the beginning of simulated aging, which was commenced after SOA was allowed to form and stabilize for 14 h.
Figure 5. Temporal evolution of SOA hygroscopicity, here parameterized using the size growth factor (GF), during and after simulated atmospheric processing. In the “control experiment” (blue symbols), SOA was left to evolve in the chamber under dark conditions. In the “O$_3$ aging” experiments (green), SOA was exposed to an excess of ozone ($\sim$ 700 ppb) under constant temperature conditions. In the “photochemical aging” experiments, SOA was exposed to light for 6 h, either under constant temperature conditions (yellow) or with light-induced heating (red). Here, the initial time $t = 0$ corresponds to the beginning of simulated aging, which was commenced after SOA was allowed to form and stabilize for 14 h.
Figure 6. Evolution of the (a) O:C ratio and (b) $f_{44}/f_{43}$ ratio of SOA during the control experiment (blue circles) and during exposure to light with increasing temperature (red circles). The temperature profiles within the chamber during the two experiments are shown in (c). Here, the initial time $t = 0$ corresponds to the beginning of simulated aging, which was commenced after SOA was allowed to form and stabilize for 14 h.
Figure 7. Comparison of temporal profiles of measured α-pinene (red circles), ozone (blue circles) and SOA mass concentration (green circles) with those modeled using GECKO-A (dashed lines) and the Master Chemical Mechanism (MCM) (solid lines). The experimental data is taken from Experiment E160410.
Figure 8. (a) Loss in SOA mass concentration (calculated as the difference between the SOA mass concentrations before and after simulated atmospheric processing) and (b) O:C ratio of the bulk SOA for four experiments exhibiting (c) different temperature increases. Here, the experimental data (filled circles) is compared with results obtained using the MCM (solid lines) and the GECKO-A (dashed lines) models. The blue points refer to the control experiment (E160411), the yellow points refer to an experiment in which SOA was exposed to light under controlled temperature conditions (E060512), and the red and brown points refer to experiments in which SOA was exposed to light and increasing temperature (E200411; E120312). Here, the initial time $t = 0$ corresponds to the beginning of simulated aging, which was commenced after SOA was allowed to form and stabilize for 14 h.