

This discussion paper is/has been under review for the journal Atmospheric Chemistry and Physics (ACP). Please refer to the corresponding final paper in ACP if available.

## Real refractive indices and volatility of secondary organic aerosol generated from photooxidation and ozonolysis of limonene, $\alpha$ -pinene and toluene

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Received: 18 December 2012 - Accepted: 20 December 2012 - Published: 21 January 2013

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Published by Copernicus Publications on behalf of the European Geosciences Union.

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Thermodenuding particles can provide insights into aerosol composition, and may be a way to create particles in laboratory chambers that better mimic the atmosphere. The volatility of secondary organic aerosol (SOA) was investigated by evaporating organics from the particles using a thermodenuder (TD) at temperatures between  $\sim 60$  and 100 °C. Volatility was influenced by the parent hydrocarbon, oxidation chemistry and relative humidity (RH). For SOA generated from ozonolysis, limonene had lower volatility than  $\alpha$ -pinene, and OH scavengers had no influence on volatility. For photooxidation,  $\alpha$ -pinene SOA was slightly more volatile than limonene SOA and increasing RH also modestly increased volatility, while toluene SOA was unaffected by heating to 98°C. For both  $\alpha$ -pinene and limonene, the concentration of NO<sub>x</sub> and the HC/NO<sub>x</sub> ratio had no discernible effect on SOA volatility. Refractive indices for the original and denuded particles were retrieved from polar nephelometer measurements using parallel and perpendicular polarized 532 nm light. Retrievals were performed with a genetic algorithm method using Mie-Lorenz scattering theory and measured particle size distributions. Retrieved refractive indices for the SOA before thermodenuding varied between 1.35 and 1.61 depending on several factors, including parent hydrocarbon, oxidation chemistry, and SOA generation temperature. For high NO<sub>v</sub> SOA, as particles shrink, their refractive index returns to the value of the corresponding size particles before heating (limonene) or slightly higher ( $\alpha$ -pinene). For low NO<sub>x</sub> however, the resulting refractive index is 0.05 ± 0.02 lower than the corresponding size undenuded particles. Additionally, for  $\alpha$ -pinene SOA from ozonolysis with OH radical scavenger, resulting refractive indices were higher by about 0.03 after heating. Consistent with no change in size, refractive indices of toluene SOA were unaffected by heating. Finally, refractive index data available to date are reviewed and resulting in suggestions an  $m_r$  for biogenic SOA of 1.44 and 1.55 for anthropogenic SOA as the most representative values.

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### Introduction

Atmospheric aerosols play an important role in determining direct radiative transfer by scattering and absorbing solar radiation. Refractive indices are also necessary to determine the optical parameters relevant to radiative transfer such as single scatter albedo, asymmetry factor and specific absorption using Mie-Lorenz theory. Additionally, reliable phase function and polarization information is essential for the interpretation of satellite and aircraft measurements to infer aerosol optical depth, size and single scatter albedo (Mishchenko, 2007).

Organic compounds constitute 20-70 % of aerosol mass in the lower troposphere depending on space and time, of which roughly 70-90% is contributed by secondary organic aerosols (SOA) (Hallquist et al., 2009). Their contribution to the global radiation balance depends on their production and loss rates and size distribution, their interactions with other radiatively important atmospheric constituents, (via the "indirect and semi-direct" effects), and on their direct interactions with insolation and upwelling terrestrial infrared radiation. The radiative properties of these ubiquitous SOA particles are poorly understood (Kanakidou et al., 2005). This is due to the chemical and physical complexity of the organic aerosols as well as the limited number of measurements available.

Recently several methods to measure or estimate optical properties of aerosols (rather than bulk materials) have been developed, and some laboratory studies have begun to derive refractive indices for SOA (Kim et al., 2010, 2012). These studies showed that  $m_r$ s span a reasonably wide range, from 1.35 to 1.6, likely depending primarily on the chemical composition, which in turn is determined by precursors, oxidation chemistry, temperature, diameter and aerosol mass concentration. At this point it is not yet clear which values are most representative of ambient aerosols.

Many currently available optical measurements of SOA have been performed on particles generated in environmental chambers or flow tubes (Hallquist et al., 2009; Kim et al., 2012), often at unrealistically high aerosol mass concentrations. We note that

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smaller particles (e.g. < ~ 200 nm) do not scatter much light, which presents measurement challenges and in some cases requires high mass concentrations. Thus, while ambient species are believed to be comprised of primarily low volatility species, chamber generated SOA contain species with a wide range of volatilities, potentially reducing the atmospheric relevance of laboratory generated SOA. The thermodenuder (TD) is a tool that can be used to help unravel some aspects of volatility, and it has been applied in a number of recent laboratory and field studies (Cappa and Wilson, 2011; Huffman et al., 2009; Jonsson et al., 2007; Kostenidou et al., 2009; Lee et al., 2011). Chamber experiments produce aerosols with higher volatility material because they are commonly run at high precursor concentration. While it might seem that this problem could be solved by lowering the precursor concentration, this approach introduces a different problem, namely that low volatility species now have much more opportunity to deposit on the chamber walls (Matsunaga and Ziemann, 2010). By generating aerosols at higher concentration and thermodenuding them, we may obtain aerosols with the full range of volatilities, and then thermodenude them to obtain more atmospherically relevant lower volatility species (Matsunaga and Ziemann, 2010). In a typical TD, aerosols are heated at a fixed temperature for a specific period of time, evaporating the high volatility compounds from the particles. Following the heating section, the particles pass through an activated carbon denuder to remove volatilized gas phase organic species to minimize recondensation on the particles as they cool.

Several TD studies to date have provided insights into aerosol volatility. Jonsson et al. (2007) showed the impact of relative humidity (RH), OH scavenger and temperature of SOA formation on the volatility of limonene and  $\alpha$ -pinene SOA. Lee et al. (2011) showed that with sufficient residence time,  $\alpha$ -pinene,  $\beta$ -pinene and limonene SOA can be almost completely evaporated at ~90°C. They also found that the concentrations of the precursor, and NO<sub>x</sub> as well as RH each influenced SOA volatility. Several recent studies have related SOA volatility and chemical composition. By combining TD and aerosol mass spectrometer (AMS) measurements, Kostenidou et al. (2009) showed that the lower volatility components of  $\alpha$ - and  $\beta$ -pinene SOA that remained after heating

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up to 90 °C had similar chemical composition as ambient aerosol, but did not find this similarity for limonene SOA. Recently Cappa and Wilson (2011) characterized the SOA from the  $\alpha$ -pinene-ozone reaction without scavenger using vacuum ultraviolet photoionization MS as the particles were heated to 170 °C. The chemical composition of SOA was not changed by heating, although the aerosol mass concentration was reduced due to evaporation.

In this study we use a TD to evaporate the high volatility components from SOA generated in a smog chamber, and retrieve their refractive indices in an effort to obtain the optical properties of SOA similar to ambient aerosols. We investigate volatility and related  $m_r$ s for SOA generated from  $\alpha$ -pinene, limonene and toluene using several different oxidation chemistries (reaction with O<sub>3</sub> with and without scavenger, and photooxidation at different HC/NO<sub>x</sub> ratios).  $\alpha$ -Pinene and limonene are representative biogenic hydrocarbons, accounting for approximately 25 % and 16 % of global monoterpene emissions, respectively, and toluene is typically the most abundant aromatic compound emitted globally (Kanakidou et al., 2005). For dark ozone reactions of unsaturated hydrocarbons, conditions can be adjusted so that organics can react either solely with O<sub>3</sub> or with both O<sub>3</sub> OH (Paulson et al., 1997). The chemical composition of SOA generated photochemically varies depending on the branching of RO2 reactions with HO<sub>2</sub>/RO<sub>2</sub> vs. NO, which is in turn controlled by the HC/NO<sub>2</sub> ratio. Angular scattering, including polarization, was measured with a 2nd generation polar nephelometer (Kim et al., 2010, 2012). A genetic algorithm (GA) approach was used to retrieve refractive indices from the PN angular scattering data from the PN (Barkey et al., 2007).

### 2 Experimental

### 2.1 Secondary organic aerosol generation

Experiments were performed in a 24 m<sup>3</sup> Teflon chamber constructed on the roof of the Math Sciences Building at UCLA. The chamber is described in detail in Chung et



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al. (2008). Air is supplied to the chamber by two 33 gallon oil-free portable air compressors (Craftsman) after passing through a series of packed bed scrubbers filled with Purafil Triple Blend (Purafil Inc.), activated charcoal, and HEPA capsule filters (Gelman). The scrubbed air had  $< 5 \,\mathrm{particles\,cm^{-3}}$ , and NO<sub>x</sub> (Thermo Electron model 14B/E), O<sub>3</sub> (Dasibi 1001-RS), and VOC (GC, HP 5890-II) levels below the 1 ppb detection limits. Access to the inside of the chamber is provided by custom Teflon sampling ports. Teflon (gasses) and copper (particles) sampling lines extend 20 cm into the chamber and withdraw samples at 0.6 (NO<sub>x</sub>) and 2 (O<sub>3</sub> and organics) LPM. Between experiments, a vent is opened and the chamber is flushed with clean air for 10 h in full sun in preparation for the next experiment.

Tables 1 and 2 show initial conditions and summary statistics for 20 experiments. For photooxidation experiments, the chamber, covered with a black tarpaulin (supported on a frame above the chamber), was half filled with purified air. At this point, gas phase reagents such as NO (Scott Specialty Gasses) and propene (Aldrich, as a photochemical initiator for toluene) were added. Later, the SOA precursor hydrocarbon liquid was evaporated into the air stream filling the chamber. Limonene (Aldrich, 99.8%),  $\alpha$ -pinene (Fluka, 98.5%) and toluene (Aldrich, 99.8%) were used as received. The chamber contents were allowed to mix for about 40 min. Once the gas chromatograph returned two measurements within 2% of one another, the tarps were removed and photochemistry initiated. The overall oxidation time was 4-5 h for limonene and  $\alpha$ -pinene 5–6 h for toluene.

For ozone experiments, the hydrocarbon(s) (precursor with or without cyclohexane) for selected experiments were injected into the half-full chamber. After the chamber contents were well mixed, excess ozone was generated by flowing pure oxygen (0.5 L min<sup>-1</sup>) through a mercury lamp O<sub>3</sub> generator (Jelight, model 600) and introduced into the chamber. In order to minimize inhomogeneities, the chamber was mixed manually while ozone was injected.

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### Thermodenuder

The thermodenuder (TD) was constructed in-house using the design of Lee et al. (2011). The TD consisted of a variable temperature heated section (L = 55 cm, inner tube D = 3.5 cm, outer tube D = 6 cm) followed by an activated carbon denuder section (L = 45 cm, inner mesh D = 3.5 cm, outer tube D = 6 cm). Temperature was controlled with heating tape wrapped on outer tube of heating sections. The space in between inner tube/mesh and outer tube was filled with sand (heating section) and activated carbon (denuding section), respectively. The flow rate through the TD was 1 LPM, corresponding to a centerline residence time of 16 s. Measurements of volatility were made with the temperature of the heated section ranging from ambient up to 98 °C. The temperature profile was measured with a 60 cm long thermocouple (OMEGACLAD, KQXL-18E) positioned in the center of the heating tube. As shown on Fig. S1, the temperature reached its maximum at 27 cm from the entrance, and remained within ±3°C of this value to 45 cm, after which point it began to decrease somewhat. Temperatures reported here are the maximum temperatures, measured 35 cm from the entrance.

For terpenes, to minimize complications created by rapidly changing particle size distributions. TD measurements were made after the mass concentration reached its maximum. Toluene nucleated late and grew slowly and continuously throughout the 6 h experiments, so TD measurements were initiated once rapid particle growth ceased and the particle growth rate decreased below 2 nm min<sup>-1</sup>. Two or three different TD temperatures were explored between 60-110°C, as experimental conditions permitted. Within this temperature range, almost all of the more volatile species should be evaporated, in principle creating particles with similar chemical composition to ambient organic aerosol (e.g., Kostenidou et al., 2009; Lee et al., 2011). It took 6-9 min for the TD to stabilize at a new temperature (Fig. 1a). Once the temperature was stable, the PN and SMPS were switched to sample denuded and undenuded chamber particles every 3 min for about 30 min.

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mobility particle sizer (SMPS) and a microbalance. Figure S2 shows a measurement schematic; SOA are drawn either into a bypass line or through the thermodenuder and then to a 4-way value (Swagelok). The aerosol flow direction was controlled in such a way that the bypass line (original particles) is sampled by the polar nephelometer (PN, described below) while the SMPS (TSI model 3080) samples the thermodenuder line (denuded particles) or vice-versa. The valve was switched every three minutes so that the PN and SMPS sampled original or thermodenuded particles alternately each for three minutes of every six minutes. Since thermodenuded and chamber particles are sampled alternately by the SMPS or PN, before and after size distribution means and standard deviations for the same type of SOA were averaged to use in retrievals of the refractive index for each point (3 min of averaged data). As measurements were made later in the experiments, after rapid growth had stopped, the phase functions and size distributions exhibited only minimal changes from one cycle to the next. Additionally, any differences between the actual size distribution for the phase function is encompassed by the large (±30%) genetic algorithm size parameter search space.

Aerosols were characterized with polar and integrating nephelometers, a scanning

The upgraded polar nephelometer used in this study and its calibrations are described in detail in Kim et al. (2010, 2012). Every 16 s, the PN measures light intensities scattered at 21 discrete angles by a stream of aerosols intersecting the beam of a 532 nm 1 W laser. A 1/2 wave plate rotates the polarization plane of the incident light to be parallel or perpendicular to the measurement scattering plane. The 1 LPM aerosol sample flow, which intersects a 2 × 5 mm beam, is confined to the center of the scattering plane by a 10 LPM sheath flow. The SMPS measures particle size distributions between 19-948 nm every 3 min, and these are used to interpret the PN data as well as determine aerosol formation yields. A multiple charging correction was applied to all SMPS scans. The genetic algorithm (GA) approach used to retrieve refractive indices from the PN angular scattering data is described in detail in Barkey et al. (2007) and

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Kim et al. (2010). SOA are accepted to be reasonably spherical, as verified in Barkey et al. (2007). The absolute error associated with the  $m_r$  retrieval above is  $\pm 0.03$ , and the instrument has sufficient sensitivity to achieve reliable retrievals for about 20 µg m<sup>3</sup> of mass concentration with particles larger than about 100 nm. All of the retrieved  $m_r$ s had fitness values of at least 0.95.

The integrating nephelometer (Ecotech M9003) measures scattering at 700 nm as well as relative humidity and temperature with one minute time resolution. In order to check the SMPS mass, filter samples for gravimetric analysis were collected on prebaked (24 h at 550 °C) 25 mm quartz fiber filters (Pall Life Sciences) for 10-20 min at 20 LPM toward the end of the experiments. Filters were allowed to be equilibrate for 24 h in a temperature and humidity controlled room before weighting using a microbalance (±1 µq, Sartorius). As SOA is lost to the walls of the chamber during the experiment, first order loss coefficients vary between 0.0012 min<sup>-1</sup> and 0.0081 min<sup>-1</sup> (Kim et al., 2010, depending on particle size) were applied to correct wall loss of particles for yield calculations. For SOA volatility calculations, losses of SOA particle within the TD due to sedimentation, diffusion and thermophoresis were determined using NaCl (99%, Sigma Aldrich) aerosol, generated by nebulizing a 0.005 M NaCl solution (Collison spray nebulizer, BGI Inc.). Detailed on experiments and particle loss results are shown at the Supplement (Fig. S3).

### Results and discussion

Initial conditions and results for 20 experiments with three different precursors (limonene,  $\alpha$ -pinene and toluene) under five different conditions were performed: photooxidation at low NO<sub>x</sub> (initial HC/NO<sub>x</sub> = 32–33), intermediate NO<sub>x</sub> (HC/NO<sub>x</sub> = 13–14) and high NO<sub>x</sub> (HC/NO<sub>x</sub> = 6.5–6.9) and ozonolysis with and without OH scavenger are shown in Tables 1 and 2; Precursor concentrations ranged from 126-258 ppb for the terpenes and 2760-2770 ppb for toluene, which requires higher concentrations to produce sufficient SOA mass.

### Aerosol formation yields

For aerosol yields (aerosol mass/HC reacted, both in µg m<sup>3</sup>), SOA mass was calculated from SMPS size distributions assuming spherical particles with densities of 1.25, 1.2 and 1.24 g cm<sup>3</sup> for limonene,  $\alpha$ -pinene and toluene, respectively (Hallquist et al., 2009; Ng et al., 2006, 2007). Measured particle number concentrations were corrected for size-dependent wall loss (above). Gravimetric mass measurements averaged 18 ± 8 % higher than the SMPS mass measurements, almost identical to the difference observed in our recent limonene study (17 ± 7%, Kim et al., 2012). The gravimetric filter samples are expected to overestimate particle mass somewhat due to adsorption of gases during sampling (Chung et al., 2008). Our yield data follow the expected trends; yields appear to be higher at lower temperatures, higher initial HC/NO<sub>x</sub> ratios, and higher aerosol mass in the chamber (Hallquist et al., 2009; Kim et al., 2012). As expected, aerosol yields for limonene are significantly higher than  $\alpha$ -pinene (Tables 1 and 2). Also, because of higher initial hydrocarbon concentrations and resulting high aerosol mass concentrations, as expected, yields at the upper end of literature values (Griffin et al., 1999; Kim et al., 2012; Ng et al., 2006; Saathoff et al., 2009).

### Reaction profile of a photooxidation experiment

Figure 1 shows chamber and thermodenuded results for a typical photooxidation experiment (22 August). The experiment had initial concentrations of 152 ppb of limonene and 110 ppb of NO. SOA began to nucleate 84 min after the chamber was initially exposed to sunlight, and quickly grew to several hundred nm (Fig. 1b and c). The thermodenuder temperature was increased in three steps from 64-84 °C, beginning after the mass concentration had reached its maximum (Fig. 1a and b). Figure 1b shows NO, NO<sub>x</sub>, O<sub>3</sub>,  $\alpha$ -pinene and aerosol mass concentration, not corrected for wall and thermodenuder losses. Figure 1c shows the evolution of particle number, mean diameter, and integrated scattering ( $\beta_{sca}$ ). The particles continued to grow throughout, however particle number concentrations dropped due to coagulation and wall loss after

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### 3.3 Volatility measurements

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### 3.3.1 Volatility of photooxidation SOA

The SOA mass fraction remaining (MFR) corrected for particle losses in the TD (Fig. S3) as a function of TD temperature is shown in Fig. 2. Biogenic SOA was increasingly denuded as temperature was increased. In contrast, toluene exhibited virtually no evaporation over the range investigated. There are no investigations available in the literature to compare the volatility of toluene SOA, however, AMS measurements of chemical composition show that SOA from aromatic HC including toluene has consistently higher values of  $f_{44}$  compared to biogenic SOA, suggesting aromatic SOA may be less volatile (Chhabra et al., 2011).

12:40.  $\beta_{sca}$  is measured from particles from the chamber (undenuded) and continues

to increase as particle numbers drop until 14:18, at which point the decrease in numbers overtakes the increases in scattering due to growing particles (Fig. 1c). Reliable

refractive indices were first obtained when the particles had grown to about 344 nm at 13:07. For undenuded particles,  $m_r$  decreases very slowly from when retrievals are first obtained from about 1.50 to 1.48 at the end of the experiment. We consistently observe

that the refractive index changes only slowly after the initial growth phase has ended

and the aerosol mass concentration has reached its peak (Barkey et al., 2007; Kim et

al., 2010, 2012), suggesting that the chemical composition of the SOA changes only

slowly later in the experiments. Thermodenuded particles exhibit slightly higher  $m_r$ s, at

1.51–1.54 (Fig. 1b) and particle numbers and diameters are reduced to 74–78% and 73–82% of the original values, respectively, decreasing with increasing TD tempera-

 $\alpha$ -pinene SOA MFRs were 61–42% for the range 64–86°C; limonene SOA had MFRs were 65–31% as the temperature was increased from 58–113°C. For both  $\alpha$ -pinene and limonene SOA, there was no evidence of an impact of the HC/NO<sub>x</sub> ratio on volatility for temperatures between 60 and 85°C. A difference might be expected for low

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NO $_{\rm x}$  conditions, as a significant fraction of this SOA is believed to consist of hydroper-oxide groups from RO $_{\rm 2}$ + HO $_{\rm 2}$  that are less volatile than the species (organic nitrate and other products) produced via RO $_{\rm 2}$ + NO at lower HC/NO $_{\rm x}$  (Presto et al., 2005). Lee et al. (2011) reported that the HC/NO $_{\rm x}$  ratio did impact  $\alpha$ -pinene SOA volatility between HC/NO $_{\rm x}$  = 6 and NO $_{\rm x}$ -free over the TD temperature range 45–60°C, but did not observe a dependence above 60°C, consistent with our results. Lee et al. (2011) reported no impact of NO $_{\rm x}$  concentration on the volatility of limonene SOA between 25–100°C, and that higher temperatures were required to almost completely evaporate limonene than  $\alpha$ -pinene SOA, also in agreement with our results. The limonene SOA generated on 16 November was somewhat more volatile (by  $\sim$  13–15%) than other experiments. This experiment had significantly higher RH (by 10–19%), implying RH may have a modest effect on SOA volatility. Consistent with this result, Jonsson et al. (2007) and Lee et al. (2011) also reported moderate increases in volatility with

### 3.3.2 Volatility of SOA from ozonolysis

increasing RH.

 $\alpha$ –pinene SOA from ozonolysis had about 15% lower MFRs than limonene ozonolysis SOA, the reverse of the photooxidation volatility trend. For both terpenes, addition of a scavenger did not have a discernible effect (Fig. 2c). This is in agreement with Jonsson et al. (2007) who reported no differences at the moderate temperatures used here, although they did find a difference above 150 °C. SOA generated in the 10 November experiment was somewhat less volatile (by  $\sim$  10%) than SOA from other  $\alpha$ -pinene/ozone/no scavenger experiments (Table 2). This experiment had significantly lower RH (24–25% vs. 30–43%), adding further credence to the notion that increasing RH increases SOA volatility (above).

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Refractive indices for SOA cover a wide range, indicating significant, optically relevant variability in chemical composition. Aerosol chemical composition is expected to depend on the parent hydrocarbon, its oxidation chemistry, the particle mass concentration, size, the temperature, and under longer time scales, potentially in-particle reactions and heterogeneous aging (Kim et al., 2012).

### 3.4.1 Photochemically generated SOA

The photochemical experiments were performed over a range of relative humidities (15–24%) for which water uptake of larger particles is not expected to change rapidly (e.g., Mikhailov et al., 2009). The amount of water uptake may change considerably, however as the particles grow (VanReken et al., 2005; Varutbangkul et al., 2006). Fresher/smaller particles are expected to contain a higher fraction of low volatility highly oxidized, hygroscopic compounds, while older/larger particles may contain less hygroscopic, higher volatility condensing species and/or oligomers (Chhabra et al., 2011; Varutbangkul et al., 2006). Temperatures span a moderate range (33–39 °C) with the exception of two experiments with higher temperatures (26 August, 2 August), but this does not appear to have a pronounced effect on  $m_r$  and MFR (Figs. 2 and 4).

### **Toluene SOA**

Figure 3 shows  $m_r$ s of undenuded (filled symbols) and thermodenuded (open symbols) particles as a function of diameter for photochemically generated toluene SOA under different HC/NO<sub>x</sub> ratios. The numbers denote the TD temperature. Consistent with previous studies (Kim et al., 2010; Nakayama et al., 2010), the freshest toluene SOA have  $m_r$ s as low as 1.35. These grew to 1.61 as the experiments progressed, and then decreased somewhat toward the end of the experiments. The value of 1.61 is significantly higher than any  $m_r$ s observed for SOA generated any biogenic HC, but it is consistent

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with our earlier results (Kim et al., 2010, 2012). We note that there is no evidence of significant absorption at 532 nm by toluene SOA. GA retrievals (which can provide only a rough estimate of absorption) return imaginary components of order  $10^{-5}$ ; Nakayama et al. (2010) also reported a negligible absorption ( $m_i = 0.037$ ) at 532 nm.

Toluene SOA refractive indices were virtually unchanged by heating up to  $\sim 98\,^{\circ}$ C (Fig. 3), as might be expected given the particles exhibited no evaporation (above), and suggesting heating does not significantly change chemical composition.

### Limonene and $\alpha$ -pinene photochemical SOA

Figure 4 shows  $m_r$ s of photochemically generated limonene and  $\alpha$ -pinene SOA generated using different HC/NO<sub>x</sub> ratios. Figure 4S directly compares the (undenuded) m<sub>r</sub> data for limonene and  $\alpha$ -pinene with Kim et al. (2012); the two data sets are in good agreement, with the exception of the decrease in  $m_r$  with increasing size for limonene SOA, which has been previously observed for  $\alpha$ - and  $\beta$ -pinene and toluene (Kim et al., 2010, 2012), but not limonene. We note that HC/NO<sub>x</sub> ratio is a controlling factor for particle size (Kim et al., 2012); as HC/NO<sub>x</sub> ratios decrease (e.g., the middle and right panels in Fig. 4), the size ranges of the particles increase for both limonene and  $\alpha$ -pinene, with the exceptions of limonene on 16 November and  $\alpha$ -pinene on 21 September. The smaller size ranges of these experiments seem to be influenced by the lower temperatures and light intensities associated with their later dates (Table 1), however their m, values are consistent the same size particles in other experiments. For limonene SOA (Fig. 4a–c), filled symbols (undenuded particles) show that  $m_r$ s increase from 1.47 to 1.55 as the particle diameters increase up to ~320 nm. After this, they drop off somewhat (for HC/NO<sub>x</sub> = 6.9–13, Fig. 4b and c).  $\alpha$ -Pinene (Fig. 4d–f,) shows the same  $m_r$ trend with diameter; refractive indices increase from 1.43 to 1.54 as the diameter of particles increase when the HC/NO<sub>x</sub> ratio was 28–33 (Fig. 4d). For the lower HC/NO<sub>x</sub> ratio experiments,  $m_r$ s drop off somewhat as particles grow above 370 nm (Fig. 4e and f).

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Decreasing  $m_r$ s for larger particles toward the end of the experiments may result from shifts in the composition of the condensing material, or possibly changes in particle composition caused by heterogeneous or in-particle reactions (i.e., aging).

For both limonene and  $\alpha$ -pinene SOA generated at high or intermediate HC/NO<sub>x</sub> ratios, the TD changes their  $m_{\rm r}$ , in most cases to that of the corresponding particle size earlier in the experiment (Fig. 4, indicated with ovals and arrows). For example, for limonene HC/NO<sub>x</sub> = 14 (Fig. 4b and c), after heating, ~370 nm particles were reduced by 50–80 nm while their  $m_{\rm r}$ s were increased by about 0.03 from 1.48–1.5 to 1.5–1.54. For HC/NO<sub>x</sub> = 13, as the ~320 nm particles shrank by 80–100 nm while their  $m_{\rm r}$ s decreased from 1.51–1.53 to 1.4–1.48. Both cases match the values for this size of SOA before heating.  $\alpha$ -Pinene SOA (Fig. 4e and f) behaves similarly. The observed TD behavior is consistent with a simple model for aerosol growth in which more volatile species are deposited on lower volatility species as the experiments progress; heating appears to simply reverse the process. This behavior is consistent with the notion that condensed species do not mix effectively, but instead layer material on as they grow, or with the notion Lee et al. (2011) that heating particles may break down oligomers, which evaporate.

In contrast to mid and high  $NO_x$  photooxidations, for both limonene and  $\alpha$ -pinene at low  $NO_x$  (HC/NO<sub>x</sub> = 33), thermodenuding produces particles with  $m_r$ s that are lower than those of the same size earlier in the experiment (Fig. 4a and d). For limonene HC/NO<sub>x</sub> = 32–33 (Fig. 4a) SOA, heating reduced the size of limonene SOA by 30–100 nm with concomitant changes in the  $m_r$ s from 1.48–1.52 to  $\sim$  1.38–1.48.  $\alpha$ -Pinene SOA from HC/NO<sub>x</sub> = 28–33 (26 August and 21 September , Fig. 4d) behaves similarly. The  $m_r$  values for the denuded particles are lower than those of the same size undenuded particles by 0.03–0.06. The results suggest significantly different chemical composition of SOA generated under low NO<sub>x</sub> conditions.

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### 3.4.2 Ozonolysis

Figure 5 shows refractive indices of SOA generated from  $\alpha$ -pinene and limonene ozonolysis with and without an OH scavenger. As particles grow, mrs increase from about 1.39 to 1.52. The OH scavenger had no discernible effect on  $m_r$ s for either terpene, consistent with results of Kim et al. (2010) for  $\alpha$ -pinene SOA at somewhat higher mass concentrations. The 18 November  $\alpha$ -pinene data has lower refractive indices (1.39-1.45) than the other experiments, a phenomenon we have observed before for particles generated at lower temperatures (18-20°C) (Kim et al., 2010). Salo et al. (2012) show observations suggesting ~ 20 °C is an important transition temperature for the chemical composition of  $\alpha$ -pinene SOA.

For all limonene ozonolysis SOA, as well as  $\alpha$ -pinene/O<sub>3</sub>/no OH scavenger, retrieved m<sub>r</sub>s for the thermodenuded particles match the same sized particles before heating (Fig. 5). However, for  $\alpha$ -pinene SOA with OH scavenger (Fig. 5b, 20 August and 18 October), m<sub>r</sub>s of thermodenuded SOA (TD temperature of 65-85°C) had slightly higher m<sub>r</sub>s than the undenuded particles (1.49–1.55 compared to 1.48–1.5). This suggests that although the MFR is independent of the experimental conditions (Sect. 3.3.2 and Fig. 3), the chemical composition of these particles after evaporation is different. Consistent with this, Cappa and Wilson (2011) and Huffman et al. (2009) observed mass spectra of SOA generated from  $\alpha$ -pinene/ozone/no OH scavenger does not change despite significant mass loss due to evaporation over the temperature range of 23–170 °C. However, for  $\alpha$ -pinene/O<sub>3</sub> with OH scavenger SOA, Kostenidou et al. (2009) showed that, the fraction of  $CO_2^+$  ( $m/z^{-1}$  44) of SOA increased by 37.5% when particles are denuded at 60-70°C. However, given the differences in particle generation conditions (e.g., tube vs. chamber, oxidation time) between studies, there may be explanations other than the presence of OH scavenger. More controlled experiments relating on refractive index with chemical composition are needed.

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### **Atmospheric implications**

Changing the refractive index from 1.4 to 1.5 decreases asymmetry factor by 0.067, which in turn changes radiative forcing by at least 12-19% for non-absorbing aerosol (Andrews et al., 2006; Kim et al., 2010; Marshall et al., 1995). Figure 6 shows the ranges of  $m_r$ s determined from chamber studies by our group together with results from the literature. Horizontal lines indicate the values used in coupled models with detailed treatment of aerosols (Kinne et al., 2003; Pere et al., 2011; Zaveri et al., 2010). The wide ranges for each experiment result from the changing chemical composition as the particles are growing and aging. It is clear that SOA  $m_r$ s are highly variable and dependent on a number of factors, including HC/NO<sub>x</sub> ratios, mass, size, aging and temperature, which determine the chemical composition of SOA. In an effort to retrieve the  $m_r$ s of SOA closely resembling ambient aerosol, this work explores the effect of parent hydrocarbon ( $\alpha$ -pinene, limonene and toluene) and oxidation chemistry (with and without OH scavenger for ozonolysis, different HC/NO<sub>x</sub> ratios for photooxidation) on the volatility and related refractive indices of SOA, accomplished by removing high volatility components using thermodenuder. The above factors as well as a minor contribution from RH and temperature appear to determine the volatility as well as related  $m_r$ s of the SOA.

More work is clearly needed before SOA optical properties are well understood. Based on results to date, however some recommendations can be made. From a global perspective, biogenic aerosol is generated most commonly under low NO<sub>x</sub> conditions, therefore the median  $m_r$ s of for  $\alpha$ ,  $\beta$  and limonene generated at low NO<sub>x</sub>, 1.44, may be the best value to use. The refractive index of anthropogenic SOA is generally higher, suggesting a representative value of around 1.55.

Supplementary material related to this article is available online at: http://www.atmos-chem-phys-discuss.net/13/1949/2013/ acpd-13-1949-2013-supplement.pdf.

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Acknowledgements. H. K. gratefully acknowledges the support of a UCLA dissertation year fellowship. The authors are indebted to B. Barkey (UCLA) who is responsible for the design and development of the polar nephelometer as well as the refractive index retrieval algorithms and ongoing helpful discussions. The authors also appreciate assistance with experiments from W. Choi and M. Kuang.

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Table 1. Initial conditions, temperatures, relative humidities and results of the photo oxidation experiments.

	Initial Conditions						Results				
Expt.	Hydrocarbon	HC (ppb)	NO <sub>x</sub> (ppb)	HC/NO <sub>x</sub> (ppbC ppb <sup>-1</sup> )	Temp (°C) <sup>1</sup>	RH (%) <sup>1</sup>	ΔHC (ppb)	# density (#/cm <sup>3</sup> ) <sup>2</sup>	Size mode (nm) <sup>2</sup>	Mass <sup>3</sup> (μg m <sup>3</sup> )	Yield <sup>4</sup> (%)
18-Aug	α-pinene	150	230	6.5	39–33	17–23	150	255	573	9–44	5.9
22-Aug	$\alpha$ -pinene	152	110	14	37–39–33	20-17-24	148	599	514	19-103	14
21-Sep	$\alpha$ -pinene	142	50	28	33–36	28-25	138	3740	359	18-107	16
26-Aug	$\alpha$ -pinene	153	47	33	39–46–42	18–13–15	130	1540	429	22-118	17
8-Aug	Limonene	208	300	6.9	37–32	16-20	206	3692	429	96-287	35
4-Aug	Limonene	140	98	14	36–40	15-18	138	1833	400	34-195	35
16-Nov	Limonene	157	120	13	24–26	38-34	144	3523	359	32-214	37
2-Aug	Limonene	130	41	32	38–42	15-18	128	2470	300	11-219	43
16-Aug	Limonene	130	39	33	35–39–34	22-18-24	130	2795	372	14–275	47
29-Aug	Toluene	2760	1350	15	42-35	18–26	990	5460	445	74–365	4.0
24-Aug	Toluene	2770	620	32	39–35	20-24	820	3090	429	100-320	11

<sup>&</sup>lt;sup>1</sup>Initial and final temperature and relative humidity.

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<sup>&</sup>lt;sup>2</sup> Final aerosol number concentration and size mode. These values have not been adjusted for wall losses.

<sup>&</sup>lt;sup>3</sup>Mass concentration in the chamber was determined from the SMPS measured size distribution in the chamber over the period for which meaningful PN measurements were made. Particle density was assumed to be 1.2 g cm<sup>3</sup> for  $\alpha$ -pinene, 1.25 g cm<sup>3</sup> for limonene and 1.24 g cm<sup>3</sup> for toluene.

<sup>&</sup>lt;sup>4</sup>Calculated from peak measured aerosol mass and the corresponding quantity of reacted hydrocarbon, corrected for wall losses. Because of the uncertainties in the measurement from GC (±3%) and measurement of SMPS (±10%) yields are uncertain to  $\pm 10\%$ .

**Table 2.** Initial conditions, temperatures, relative humidities and results of the ozonolysis experiment.

Initial conditions						Results					
Run	Hydrocarbon	HC (ppb)	Ozone (ppb)	Temp (°C) <sup>1</sup>	RH (%)	ΔHC (ppb)	# density <sup>2</sup> (#/cm <sup>3</sup> )	Size mode <sup>2</sup> (nm)	Mass <sup>3</sup> (μg m <sup>3</sup> )	Yield <sup>4</sup> (%)	
20-Aug	$\alpha$ -pinene <sup>5</sup>	143	500	26–27	33–31	107	1080	478	28–230	46	
18-Oct	$\alpha$ -pinene <sup>5</sup>	150	500	23-26	34-36	130	2780	414	39-271	44	
29-Sep	$\alpha$ -pinene	170	500	23-28	38-30	156	5820	322	37-271	40	
10-Nov	$\alpha$ -pinene	160	500	23-22	24-25	160	25600	217	71-349	45	
18-Nov	$\alpha$ -pinene	126	500	18–20	43-39	118	17400	233	34-215	39	
31-Aug	Limonene <sup>5</sup>	167	500	25-27	38-35	167	1280	300	98-579	78	
23-Nov	Limonene	198	500	20-22	41-35	198	33000	209	83-614	72	
24-Oct	Limonene	150	500	21–23	42-40	150	18800	250	96–454	72	

 $<sup>^{1,2,3,4}</sup>_{-}$  Same as footnotes 1–4 in Table 1.

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<sup>&</sup>lt;sup>5</sup>Cyclohexane was added in 50 fold excess compared to the hydrocarbon to suppress OH formation.



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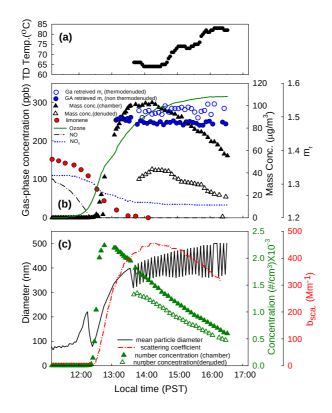
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**Fig. 1.** Profile of the  $\alpha$ -pinene photooxidation experiment performed on 22 August, with initial  $\alpha$ -pinene, 152 ppb; NO<sub>v</sub>, 110 ppb. (a) Temperature profile of TD (b) hydrocarbon, NO, NO<sub>v</sub>, O<sub>3</sub>, aerosol mass concentration and GA determined real refractive index of SOA from chamber and thermodenuder (c) time evolution of SOA particle number concentrations, mean diameters of SOA from chamber and thermodenuder and scattering coefficient ( $\beta_{sca}$ ).



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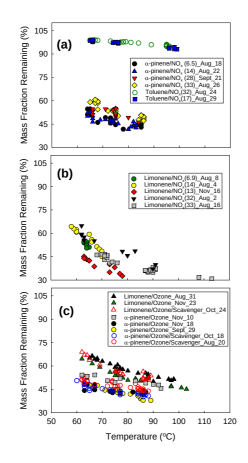
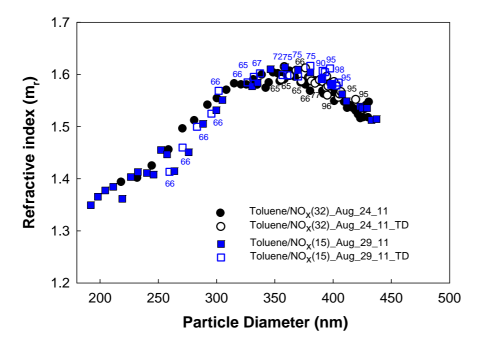


Fig. 2. Mass fraction remaining after passing through the TD as a function of TD temperature for SOA formed by photooxidation of (a) toluene and  $\alpha$ -pinene and (b) limonene at different HC/NO<sub>x</sub> ratios, and (c) ozonolysis SOA. All data corrected for experimentally determined TD losses (Fig. S3).



**Fig. 3.** Retrieved refractive indices for thermodenuded (open symbols) and undenuded (filled symbols) toluene SOA formed by photooxidation of toluene at different  $HC/NO_x$  ratios. Numbers indicate the TD temperature (°C).

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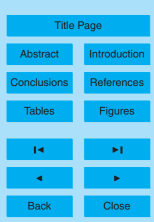




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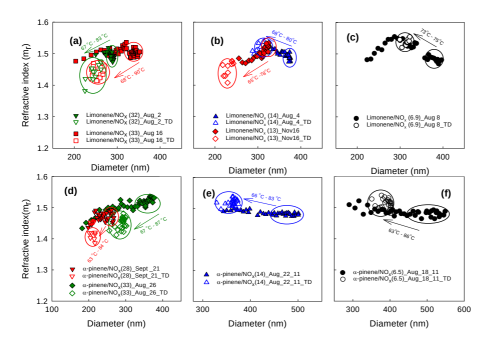
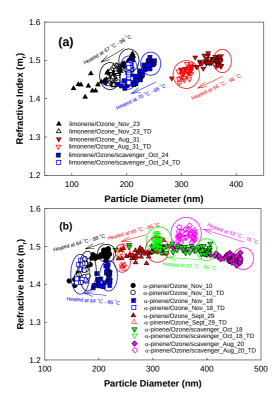


Fig. 4. Retrieved refractive indices for thermodenuded (open symbols) and undenuded (filled symbols) SOA formed by photooxidation of limonene (upper panel; (a-c) and  $\alpha$ -pinene (lower panels; (d-f) at different HC/NO<sub>x</sub> ratios. TD temperatures are indicated.



**Fig. 5.** Retrieved refractive indices for thermodenuded (open symbols) and undenuded (filled symbols) SOA formed by ozonolysis of **(a)**  $\alpha$ -pinene and **(b)** limonene with and without scavenger.

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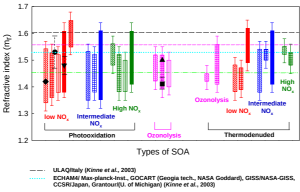
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**Fig. 6.** Comparisons of refractive indices of thermodenuded and undenuded SOA studied in previous studies (Kim et al., 2010, 2012). Ranges of refractive index for each SOA result from changing chemical composition as the particles are growing or aging; error bars are  $\pm 0.15$  for Kim et al. (2010, 2012) studies. Barkey et al. (2007) has  $\pm 0.15$ , Lang-Yona et al. (2010) has  $\pm 0.05$  and Nakayama et al. (2010) has  $\pm 0.04$  uncertainties, respectively. Uncertainties of Schnaiter et al. (2005) study is not available. Horizontal lines indicate SOA or organic aerosol refractive indeces assumed by different current aerosol/climate models. Black symbols represent literature values of SOA generated using same hydrocarbon and oxidation chemistries denoted in the figure. The phenol data is not published and generated at HC/NO<sub>x</sub> = 35–41. HC/NO<sub>x</sub> ratios are 20–33, 13–19 and 6.3–11 for low, intermediate and high NO<sub>x</sub>, respectively.

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