Real-time, controlled OH-initiated oxidation of biogenic secondary organic aerosol

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Abstract. The chemical complexity of atmospheric organic aerosol (OA) requires novel methods for characterization of its components and description of its atmospheric processing-induced transformations. We present the first field deployment of the Toronto Photooxidation Tube (TPOT), a field-deployable flow reactor for the controlled exposure of ambient aerosol to OH radicals. The system alternates between sampling of (1) (unreacted) ambient aerosol, (2) aerosol exposed to UV light and subjected to a ∼4 to 10°C temperature increase, and (3) aerosol that is oxidized by OH (in addition to the aforementioned UV exposure/temperature increase). This allows both characterization of the aging process and classification of aerosol in terms of its volatility and reaction-based properties. Summer-time measurements by an aerosol mass spectrometer coupled to the TPOT were performed in the remote forest of western Canada, resulting in aerosol dominated by biogenic secondary organic aerosol. Volatilization/UV exposure resulted in an approximately 10 to 25 % decrease in organic mass and resulted in a slight increase in oxygenation. OH oxidation resulted in a further organic mass decrease (additional ∼25 %) and yielded an aerosol with O:C values comparable to those characteristic of low volatility, highly oxygenated OA. Most OH-induced changes occurred within ∼3 day-equivalents of atmospheric processing, with further reactions generally proceeding at a greatly reduced rate. Positive matrix factorization (PMF) analysis of the TPOT data yielded five factors. One factor is related to primary biomass burning organic aerosol, while the others describe oxygenated organic aerosol (OOA) components in terms of reactivity and volatility: (1) volatile and reactive; (2) non-volatile and reactive; (3) non-volatile and reactive early-generation product; (4) non-volatile and non-reactive product. This PMF classification of aerosol components directly in terms of reactivity and volatility is enabled by the TPOT-modulated perturbation of aerosol composition, and is not otherwise accessible. The particle-phase reaction end products have mass spectra similar to the low-volatility oxygenated organic aerosol (LV-OOA) factors widely reported in the literature, providing supporting evidence for aged organic aerosol formation from OH-driven oxidation processes.

1 Introduction

Atmospheric aerosols have important effects on climate, visibility, and human health. However, quantification of such effects remains highly uncertain due in part to challenges in characterizing the organic aerosol (OA) fraction. Such difficulties include the large number of atmospheric organic compounds, many of which cannot presently be identified or isolated, wide range of emissions sources, and myriad pathways for atmospheric OA production and/or processing.

OA can be directly emitted (primary organic aerosol, POA) from sources such as biomass burning and anthropogenic combustion. Alternatively, volatile organic compounds (VOCs) may be emitted by biogenic sources, biomass burning, or anthropogenic activities, and are then oxidized in the atmosphere to form lower-volatility products that partition to the particle phase, forming secondary organic aerosol (SOA) (Kanakidou et al., 2005; de Gouw and Jimenez, 2009; Hallquist et al., 2009). POA and SOA from different sources can mix and become further oxidized by heterogeneous reaction with oxidizing species such as OH radicals. Further,
a significant fraction of ambient OA, whether POA or SOA, is semivolatile and may partition back to the gas phase in response to dilution and/or temperature changes, become further oxidized through gas-phase reactions, and then re-enter the particle phase. All the while, the preexisting OA is subject to continuous injections of fresh POA, SOA, and SOA precursors, which in turn undergo the atmospheric processing described above. The resulting OA is a highly complex mixture formed from disparate sources and affected by a wide array of atmospheric processing mechanisms.

Several strategies have been introduced to simplify, isolate, and/or deconvolve these sources and processes. Factor analysis techniques applied to aerosol composition are frequently used for source apportionment. Such approaches include chemical mass balance (e.g. Hidy and Friedlander, 1971; Schauer et al., 1996; Hannigan et al., 2005; Chow et al., 2007), which represents the data as a linear combination of predetermined, static factors, and positive matrix factorization (PMF) (e.g. Paatero and Tapper, 1994; Zhao and Hopke, 2006; Lanz et al. 2007; Ulbrich et al., 2009), which utilizes measurement uncertainties and does not require a priori knowledge of the factor profiles. Hybrid approaches, such as PMF with one or more fixed factors, have also been utilized (Lanz et al., 2008), as have custom models based on known tracers, variable factors, etc. (Wahlin 2003; Zhang et al., 2005; Christensen et al., 2006). For all of these approaches, a central challenge is the source complexity and constantly changing nature (via gas/particle partitioning and chemical reactivity) of the measured aerosol. Descriptions of SOA generation and/or aging are particularly challenging. For example, PMF of data from the Aerodyne aerosol mass spectrometer (AMS), typically describes SOA as a linear combination of semivolatile and low-volatility oxygenated organic aerosol, however more detailed information about reactivity, volatility, sources, etc. is unavailable in the absence of other integrated measurements, such as VOCs (Slowik et al., 2010) or with an online thermomdenuder (Huffman et al., 2009).

In contrast, laboratory-based methods potentially allow for controlled aging processes and emission sources, but can be challenging to extrapolate to the atmosphere. Smog chambers provide a laboratory-based method for controlled exposure of model compounds or selected source emissions to oxidizing species. Common model compounds studied include isoprene, α-pinene, or other terpenes as surrogates for biogenic emissions, and aromatic species such as toluene as surrogates for anthropogenic combustion. Alternatively, source emissions such as wood burning and diesel vehicles may be injected directly into the smog chamber, and then aged under controlled conditions (Grieshop et al., 2009; Mentel et al., 2009; Chirico et al., 2010). Such analysis of model compounds and specific sources allows for isolation of particular reaction systems, however simplification of the reactive system and/or increased concentrations relative to ambient conditions make it challenging to directly extrapolate such results to the atmosphere.

The methods described above have shown that oxidative aging tends to move OA from a less oxygenated, semivolatile state to a more oxygenated, low-volatility state. However, the timescales over which this occurs and the associated mechanisms remain unclear. Further, in ambient studies such changes must generally be inferred from complex datasets, in which it is difficult to segregate the relative effects of different sources, atmospheric reactions, and partitioning. Here we employ a complementary approach to the characterization of atmospheric aging, namely oxidation of directly sampled ambient aerosol by OH radicals using the Toronto Photooxidation Tube (TPOT). This retains the real-world sources and complexity of ambient measurements while allowing a controlled aging environment. Similar flowtube systems have previously been used to measure the reactivity of ambient urban aerosol (George et al., 2008), estimate the SOA-forming potential model compounds (Kang et al., 2011), and characterize the evolution of laboratory biomass burning emissions (Cubison et al., 2011). They have also been applied to studies of oxidative aging of laboratory aerosol (George et al., 2007; Kroll et al., 2009; George and Abbatt, 2010; Kessler et al., 2010; Lambe et al., 2011a, b). In the TPOT, aging is conducted in a flowtube with a residence time on the order of minutes and the system rapidly switches between reacted and unreacted conditions, which enables highly time-resolved measurements of aerosol aging. The resulting mass spectral changes due to oxidative processing are characterized in terms of selected m/z and using the PMF analysis technique. Here we discuss deployment of the TPOT during summer in a forested region known for high levels of biogenic aerosol formation without significant anthropogenic pollution. We address the following questions: (1) Does OH oxidation lead to mass loss of biogenic organic aerosol? (2) What is the character of the mass spectral response, and what insight does it provide regarding the oxidation mechanism? (3) Can PMF analysis demonstrate how different fractions of the biogenic aerosol respond to OH oxidation?

2 Materials and methods

2.1 Sampling location

Measurements were conducted as part of the Whistler Aerosol and Cloud Study (WACS 2010, 15 June to 28 July 2010). The TPOT system was deployed at the Raven’s Nest station (1320 m a.s.l.) on Whistler Mountain in Whistler, BC, Canada (50.08° N, 122.95° W). Whistler is located in the remote Canadian forest, approximately 120 km north of Vancouver. The Raven’s Nest site was located in the forest, approximately 450 m below treeline. A detailed overview of the WACS 2010 campaign is presented elsewhere (Macdonald et al., 2012). Organic aerosol in the Whistler area has
previously been shown to be strongly influenced by biogenic emissions during the late spring and early summer (e.g. Leaitch et al., 2009; Sun et al., 2009; Schwartz et al., 2010; Takahama et al., 2011; Leaitch et al., 2011). Later in the summer, regional forest fire emissions can also influence the site (McKendry et al., 2011; Takahama et al., 2011). The sampling period for WACS 2010 was chosen to maximize biogenic influences, while minimizing biomass burning ones. As discussed in the results section, this strategy was mostly successful, with the exception of a regional biomass burning event during the last few days of the study. A large biogenic event with organic aerosol levels approaching 6.5 µg m⁻³ was observed from July 6 to 10, while significant local formation of SOA from monoterpenic oxidation observed in the diurnal cycles of 13 to 21 July (Macdonald et al., 2012).

The TPOT was housed in an unused single-story restaurant at the Raven’s Nest site. The sampling inlet was located approximately 0.5 m above the roof of the building. Air was continuously sampled through this inlet at 1100 sccm, passing through 3 m of 6 mm o.d. stainless steel tubing before being introduced into the TPOT. The sampling lines were electrochemically coated with amorphous silicon (SilcoTek, Bellefonte, PA, USA) to minimize adsorptive losses of volatile organic compounds. Controlled OH-initiated oxidation was performed in the TPOT system (see Sect. 2.2), and the resulting particle composition was measured by aerosol mass spectrometry (see section 2.3).

2.1.1 Toronto Photooxidation Tube (TPOT)

The TPOT is conceptually based on flow reactor systems previously described and characterized in the literature (George et al., 2007; Lambe et al., 2011a), however the system used in the present study has been significantly upgraded in terms of reactor geometry (higher volume/surface ratio, improved injection/sampling port configuration) and material (most glass and copper surfaces replaced with silicon-coated steel). A schematic of the system is shown in Fig. 1. All surfaces in the system that contact ambient air are coated with the amorphous silicon material described in Sect. 2.1, with the exception of a Pyrex mixing volume and valve systems, discussed below. Sampled ambient air (1100 sccm), a humidified flow (N₂ passed through a bubbler) maintaining the entire TPOT system at 40 % RH, and a mixed N₂/O₃ flow are combined in a Pyrex mixing volume (residence time ~1 min). The N₂/O₃ flow is either irradiated by a lamp at 185 nm to produce O₃, or bypasses the lamp system. The N₂/O₂ path is controlled by an automatic 4-way valve system that switches between conditions every 12 min. To maintain system stability, flow is continuously maintained through both the lamp and lamp-bypass lines, with one line introduced into the TPOT and the other exhausted to a pump. Ozone concentrations are monitored in the exhaust line using a Model 202 Ozone Analyzer (2B Technologies, Inc., Boulder, CO, USA).

Fig. 1. Schematic diagram of the TPOT system. “4-w vlv” and “3-w vlv” denote the automatic 4-way and manual 3-way valves, respectively.

After the mixing chamber, particles pass into the reaction zone. This consists of a silicon-coated stainless steel tube (11 cm o.d. × 29 cm length) with 4-point injection and sampling configurations on the entrance and exit flanges. A UV lamp emitting at 254 nm is located on the center axis of the tube, and is surrounded by quartz housing. A ~20 slpm flow of air is maintained between the lamp and the housing to prevent lamp overheating and minimize lamp-induced temperature increases in the flowtube. Residence time in the reaction zone is approximately 1 min, and the surface area-to-volume ratio is approximately 1/3 cm⁻¹. Alternatively, a manual 3-way valve can be used to send particles through a bypass flow tube system that is identical in all ways to the reaction zone, except that the quartz housing contains no lamp. After passing through the reaction zone or reaction bypass, particles were sampled by an aerosol mass spectrometer (C-ToF-AMS, see section 2.3), as well as a differential mobility analyzer coupled to a condensation particle counter and a cloud condensation nuclei counter, as discussed elsewhere (Wong et al., 2011). The transit time from the TPOT exit to the AMS is approximately 30 s.

The OH concentration in the TPOT is controlled by varying the O₃ concentration in the N₂/O₂ flow. Steady-state OH concentrations are determined by calibration experiments in which the decay of methyl-ethyl-ketone (MEK) is measured using a time-of-flight proton transfer reaction mass spectrometer (ToF-PTRMS, Ionicon Analytik, Innsbruck, Austria). This allowed formation of an empirical relationship between O₃ concentration and OH exposure. Because the ToF-PTRMS was not routinely available, reactor stability was periodically monitored by introducing 145 nm bis(2-ethylhexyl) sebacate (C₂₉H₅₀O₄, BES) particles into the TPOT immediately before or after an MEK calibration at the same conditions. BES is an organic liquid with well-characterized heterogeneous OH reactivity (George et al., 2011).
2007). This calibration allowed the generation of an empirical relationship between the MEK-calibrated OH exposure (and O$_3$ concentration) and BES reaction as described by the fractional increase in AMS $m/z$ 44. These BES experiments were used to confirm the stability of the O$_3$ concentration/OH exposure relationship, not to perform a priori estimation of the OH exposure. Such an estimate would be complicated by factors such as instrument-to-instrument differences in fragmentation patterns, differences in the kinetics between gas-phase and heterogeneous reactions, etc. Control experiments indicated negligible reactivity with O$_3$ at TPOT concentrations (40 to 1100 ppbv) (see Fig. S2). OH exposures calculated from the MEK calibrations were reproducible to within $\sim$30%.

The temperature within the reaction zone was monitored by thermocouple mounted on the inner wall of the lamp housing, thereby approximating the closest surface to the lamp encountered by the ambient aerosol. During WACS 2010, the lamp subjected the sampled air to a temperature increase of $\sim$4°C relative to the indoor temperature at the sampling enclosure. However, it is possible that the aerosol does not reach room temperature before entering the TPOT, and that the heating is instead described by the difference between outdoor and TPOT temperatures. The time series of this relationship is plotted in the Supplement (Fig. S1) and yields a mean temperature difference of 10.4 $\pm$2.1°C.

### 2.2 Aerosol mass spectrometer

The Aerodyne time-of-flight aerosol mass spectrometer (C-ToF-AMS, Aerodyne Research, Inc., Billerica, MA, USA) provides the non-refractory composition of particles with vacuum aerodynamic diameter between approximately 60 and 600 nm (Drewnick et al., 2005). Instrument characteristics and operating/analysis procedures are described in detail in the literature (e.g. Jayne et al., 2000; Allan et al., 2003; Canagaratna et al., 2007). Particles are continuously sampled from atmosphere through a 100 μm critical orifice ($\sim$100 cm$^3$ min$^{-1}$) into an aerodynamic lens ($\sim$2 torr), which focuses the particles into a narrow beam. The particles impact on a resistively-heated tungsten surface (600°C, 10$^{-7}$ torr), where they flash vaporize. The resulting gas ionized by electron impact (70 eV) and the ions are detected by a time-of-flight mass spectrometer (ToFwerk AG, Thun, Switzerland). The AMS also has particle sizing capabilities, but these were not utilized during the present study due to mechanical problems. Mass spectra were recorded with 1 min time resolution.

During normal operation, the particle beam is alternately blocked, yielding the instrument background, and unblocked. The difference between these two measurements is the mass spectrum of the sampled particles, together with signal from the major gases in air (primarily N$_2$, O$_2$, H$_2$O, Ar, and CO$_2$). Contributions from these gases, including fragment and isotopic ions, are removed from the particle spectra through the ToF-AMS Toolkit v1.51 data analysis package (D. Sueper, University of Colorado-Boulder, Boulder, CO, USA) for Igor Pro v6.22A (Wavemetrics, Inc., Lake Oswego, OR, USA) based on the mass spectral fragmentation analysis algorithm developed by Allan et al. (2003). These algorithms are also used to quantitatively separate organic ion signals from other species. Gaseous fragmentation patterns were characterized using regular filter measurements to zero the particle signal. A collection efficiency of 0.5 is assumed throughout the AMS transmission window. However, the analysis herein utilizes normalized values or ratios and is therefore unaffected by this assumption.

### 2.3 Positive matrix factorization

The AMS organic mass spectral time series was analyzed using positive matrix factorization (PMF) (Paatero and Tapper, 1994; Paatero 1997). PMF represents the AMS time series as a linear combination of static factor profiles and their time-dependent intensities according to the matrix equation $X = GF + E$, where $X$ is the data matrix (i time points x j m/z), the $G$ matrix contains the factor time series matrix (i time points x p factors), the $F$ matrix contains the factor mass spectra (p factors vs. j m/z) and $E$ is the residual matrix (i time points x j m/z). PMF requires as inputs the AMS mass spectral time series and associated uncertainties, which are calculated according to Allan et al. (2003). The first application of PMF to AMS datasets was conducted by Lanz et al. (2007), and many other studies have followed.

For the TPOT data during WACS 2010, the AMS dataset $X$ consists of 20778 time points (i.e. 1-min mass spectra) and 270 m/z. A minimum error corresponding to measurement of a single ion was applied to the dataset and m/z assumed in AMS data analysis procedures to be proportional to m/z: 44 were downweighted as described in Ulbrich et al. (2009). No further downweighting was applied to the dataset. PMF allows the user to determine the number of factors in the output solution; here solutions containing 1 to 12 factors were investigated. Matrix rotations were explored by varying the $f_{Peak}$ parameter from $\sim$2 to 2. The selected solution (see Sect. 3.3) was a 5-factor solution at $f_{Peak} = -0.5$. The value of the object function $Q$, calculated as the sum of the squares of the uncertainty-scaled residuals, was 1.13 times the theoretically-expected value. The average residual, calculated as the mean of $|\text{residual}|/\text{signal}$, was 0.083. 5-factor solutions with $f_{Peak}$ values between $-0.5$ and 0 were not convergent using the standard criteria, and the negative $f_{Peak}$ solutions provided more clearly distinguished factors than the positive $f_{Peak}$ ones. Convergence could be achieved for $f_{Peak}$ values between $-0.5$ and 0 only by significantly relaxing the convergence criteria, which yielded solutions with significantly higher $Q$-values; these solutions were therefore rejected. Each factor in the 5-factor solution could be interpreted meaningfully, making it preferable to lower-order solutions. Higher order solutions provided uninterpretable
factors and/or split/mixed factors, which frequently indicate an excessive number of factors (Ulbrich et al., 2009). The possibility of local minima in the PMF solution was explored by initiating the PMF algorithm from 150 different random starting points (“seeds”); no solutions with significant differences from the selected solution were found.

3 Results and discussion

The Whistler site is in a forested region with strong monoterpene emissions. A summary of the major results from the WACS 2010 campaign is provided elsewhere (Macdonald et al., 2012). During the periods of TPOT sampling, the aerosol can mostly be classified as biogenic SOA resulting from gas-phase oxidation of monoterpenes. The primary exception to this classification occurred near the end of the campaign (26 to 27 July), when the site was influenced by transported aerosol from regional forest fires. Strong diurnal cycles of organic mass, biogenic VOC emissions and oxidizing species are observed, which can be attributed to both upslope/downslope flow and local chemical reactions.

As discussed in Sect. 2.2, aerosol passing through the TPOT is subjected to one of four conditions: (1) unreacted ambient aerosol (ambient condition, “Amb”); (2) exposure to O$_3$ (ambient + O$_3$ condition, “AmbO$_3$”); (3) exposure to UV light (ambient + UV condition, “AmbUV”); and (4) exposure to O$_3$ and light, resulting in exposure to OH radicals (ambient + UV + OH condition, “AmbUVOH”). Control experiments comparing conditions (1) and (2) to determine the effects of O$_3$ exposure showed no changes above detection limit to either the total organic mass or individual m/z within the spectrum. A sample control experiment of this type is shown in the Supplement (Fig. S2) and the R$^2$ between the spectra is 0.9995. Therefore, from this point we will treat conditions (1) and (2) as equivalent and refer to both as the “Amb” condition. For the AmbUV condition, we cannot distinguish the effects of volatilization and irradiation by 254 nm light. We further note that although only a 4 $^\circ$C temperature increase was measured for the AmbUV condition, some larger local increases in temperature may occur within the TPOT, especially near the quartz tube. Effects of volatilization/UV exposure and OH oxidation on the mass spectra are treated separately below in Sects. 3.1 and 3.2. Because repeated perturbation of the ambient aerosol by the TPOT forces mass spectral variability based on particle volatility/UV reactivity and OH reactivity, factor analysis describes the particle composition in terms of these properties. This is illustrated via PMF analysis of the TPOT AMS data in Sect. 3.3. The duration of Amb, AmbUV, and AmbUVOH periods varies throughout the study in response to changes in selected OH exposure and observed ambient aerosol composition. Listings of the averaging periods used are presented in the supplement (Table S1 for Amb, Table S2 for AmbUV and AmbUVOH in the Supplement).

3.1 UV exposure

Figure 2 shows a sample time series of TPOT data, with ambient (“Amb”), ambient + 254 nm UV light (“AmbUV”), and ambient + UV light + OH (“AmbUVOH”) conditions labelled. The figure shows a decrease in organic mass on the order of 15 % between the Amb and AmbUV conditions. There are two possible explanations for this mass loss: (1) volatilization of semivolatile organics due to heating induced by the UV lamp; or (2) photolysis of OA species such as peroxides. Although the sampled aerosol is diluted by a humidified N$_2$O$_2$ flow within the TPOT system, this dilution takes place before the particles are split into Amb and AmbUV pathways and therefore does not contribute to the observed mass loss. Wall loss experiments were conducted in the reaction zone with the UV lamp on using size-selected bis(2-ethylhexyl) sebacate (“BES”, C$_{26}$H$_{50}$O$_4$) particles. These experiments showed approximately 100 % transmission for 100 to 300 nm particles, and approximately 90 % transmission for 50 nm particles. Losses through the 3-way switching valve were not calculated, however here the pathways taken by the Amb and AmbUV pathways are identical. It is therefore extremely unlikely that particle losses significantly contribute to the observed mass loss. Finally, we note that the comparison of Amb and AmbUV periods requires interpolation of the Amb condition over a 3 to 4 hour time interval (see Tables S1 and S2 in the Supplement). While this introduces some scatter into the Amb/AmbUV comparison, because the sampling site is influenced primarily by daily temperature/boundary layer cycles and major changes in weather/air mass origin (as opposed to local sources), the interpolation is not expected to bias the comparison.

The relative effects of photolysis and volatilization on OA mass loss between Amb and AmbUV conditions cannot be deconvolved in the current dataset. Note that the AmbUV condition represents an approximate temperature increase of only $\sim$4 $^\circ$C (TPOT vs. indoor air temperature) to $\sim$10 $^\circ$C (TPOT vs. outdoor air temperature). If this mass loss was caused by TPOT heating, it would indicate that the particles
are quite volatile. Conversely, if the mass loss were due to photolysis, it would indicate a large fraction of photoreactive species such as peroxides. Figure 3a shows the fraction of mass lost as a function of $f_{44}$ over the entire campaign, where $f_{44}$ is defined as the fraction of organic mass occurring at $m/z$ 44. (Averaging periods for the data points in this and similar figures are reported in Table S1.) For each point in Fig. 3a, the $\text{Org}_{\text{Amb}}$ is calculated as the mean of a single ambient measurement period, as denoted by one of the labelled periods in Fig. 2. The corresponding $\text{Org}_{\text{AmbUV}}$ data is drawn from the average of the two nearest flanking AmbUV periods (12 min). The signal at $m/z$ 44 is dominated by the CO$_2^+$ ion, which in the AMS is frequently observed from organic acids (e.g. Duplissy et al., 2011). $f_{44}$ can be used to estimate the molecular O:C ratio (Aiken et al., 2008).

Over the entire study, the AmbUV condition causes a loss of 10 to 25% of the organic signal. As shown in Fig. 3a, the fraction of organics lost decreases with increasing $f_{44}$, suggesting that particle volatility and/or photolysis is inversely related to oxygenation. This trend with volatility is consistent with previous studies (e.g. Huffman et al., 2009). There is no dependence of $\text{Org}_{\text{AmbUV}}/\text{Org}_{\text{Amb}}$ on the temperature difference between the TPOT and outdoor air. While the correlation shows significant scatter, the trend is evident. Note also that some volatilized OA may recondense on the particles, so volatility effects on the mass loss in Fig. 3a will be somewhat suppressed. Sample mass spectra comparing the Amb and AmbUV conditions are shown in Fig. S3 in the Supplement.

3.2 OH oxidation

Exposure of aerosol to OH radicals (i.e. the AmbUVOH condition) causes a further decrease in organic mass relative to $\text{Org}_{\text{AmbUV}}$ (see Fig. 2). As shown in Fig. 3b, the magnitude of this decrease depends primarily on OH exposure, with the extent of oxidation of the unprocessed particle (as represented by $f_{44}$) having little effect. (Averaging periods for this and similar figures are given in Table S2.) A crucial feature of this plot is that an organic mass increase beyond experimental uncertainties is not observed for any OH exposure. This suggests that chemistry in the TPOT is dominated by heterogeneous reactions and/or gas-phase reactions of semivolatile species partitioning from the particle phase, and that SOA formation is not significant. This is consistent with the flowtube intercomparison study of Lambe et al. (2011a), which showed the SOA yield from a system decreases with an increasing ratio of wall surface area to volume. Peak monoterpane concentrations during the period of TPOT sampling were on the order of 0.5 ppbv (Macdonald et al., 2012); at these levels, significant TPOT SOA formation is not expected (Lambe et al., 2011a). SOA yields also decrease with decreasing VOC concentrations; for a relatively pristine site such as Whistler, TPOT SOA production is negligible.

As shown in Fig. 2, TPOT conditions also slightly affect the inorganic species. Notably, sulfate and ammonium increase for the AmbUVOH condition, while nitrate decreases. The increase in sulfate is likely due to oxidation of gas-phase SO$_2$ to H$_2$SO$_4$. This H$_2$SO$_4$ is then rapidly neutralized by ammonium, resulting in a corresponding increase of particulate ammonium. The decrease in nitrate may be due either to consumption of organonitrates, whose major mass fragments occur at the same $m/z$ as inorganic nitrate, or to reaction of HNO$_3$ with OH.

Figure 4 describes the changes in the AMS mass spectra due to volatilization and OH oxidation using the framework developed by Ng et al. (2010). Ambient SOA typically falls within the triangle denoted by the solid lines in the figure, with more aged (and typically less volatile) SOA lying in the upper left. PMF analyses of AMS spectra often resolve two SOA-related factors, termed low-volatility and semi-volatile oxygenated organic aerosol (LV-OOA and SV-OOA), which respectively fall in the upper left and lower sections of this triangle. As shown in the figure, the ambient particles begin in the SV-OOA region of the triangle. Volatilization (AmbUV condition) moves them only...
slightly upwards and to the left. However, a strong effect is seen from OH oxidation. At the lowest OH exposure levels ($\sim 1.5 \times 10^{11}$ molec cm$^{-3}$ s), only a slight movement towards the upper left is observed. However, the highest OH exposures ($\sim 1.5 \times 10^{12}$ molec cm$^{-3}$ s) move the composition well into the LV-OOA region of the triangle. These OH exposures correspond to approximately 0.85 and 10 day-equivalents of atmospheric aging, respectively, assuming a 24-h mean OH concentration of $2 \times 10^{6}$ molec cm$^{-3}$. Note, however, that significant aging takes place on shorter timescales; an OH exposure of $\sim 5.8 \times 10^{11}$ molec cm$^{-3}$ s ($\sim 3.7$ day-equivalents) already causes movement into the LV-OOA region.

While the framework utilized in Fig. 4 provides a simple overview of the aging process, additional information is available from consideration of the entire mass spectrum. Fig. 5 shows the effects of low and high OH exposures on the intensities of all organic m/z up to m/z 125. Changes in the mass spectra are plotted as the ratio of spectra obtained under the AmbUVOH condition to the AmbUV condition (the AmbUV and AmbUVOH spectra are plotted independently in Fig. S4 in the Supplement). Spectra are not normalized, so a value of 1 indicates no change in the mass measured at that m/z, while values $>1$ or $<1$ indicate mass increases or decreases, respectively. The figures show decreases at all m/z except for m/z 44, 45, and m/z dependent on 44. (In the AMS data processing algorithm, the organic signal at m/z 16 through 21 and 28 are assumed to be in constant proportion to m/z 44 (Allan et al., 2003; Aiken et al., 2008). The apparent increases at some higher m/z for the low OH condition occur at m/z having too low signal for the apparent increases to be reliable. The lone exception occurs at m/z 45. Approximately 10% of the signal at this ion results from the CO$_2^+$ isotope, and the rest is probably dominated by the HCO$_2^+$ ion, which would also suggest the formation of organic acids (see Fig. S5).

While decreased mass due to OH oxidation is evident throughout the spectra shown in Fig. 5, the fractional decrease varies significantly by m/z. For example, at the high OH exposure, m/z 43 decreases by 37%, while m/z 105 decreases by 47%. Generally, the fractional decrease is larger for fragments at higher m/z and for fragments characteristic of hydrocarbons or slightly oxygenated hydrocarbons. The change in mass for selected fragments (m/z 43, 44, 55, and 91) and total organic mass is shown as a function of OH exposure in Fig. 6. The trends shown in Fig. 5 are also evident in this figure, with for example m/z 91 showing a stronger decrease than m/z 43. However, Fig. 6 also provides additional insight into the rate of heterogeneous oxidation. The rate of mass decrease slows considerably after $\sim 3$ day-equivalents of oxidation (OH exposures $\sim 6.9 \times 10^{11}$ molec cm$^{-3}$ s), as do the changes in composition.

Fitting the m/z 44 production to the function $y = 1 + a(1-e^{-kt})$ yields a kinetic rate constant of $3.0 \times 10^{-12}$ cm$^3$ molec$^{-1}$ s$^{-1}$. Rate constants for the reaction of other m/z, as well as for PMF factors (see Sect. 3.3), are given in Table 1. While significant uncertainties exist because of the scarcity of TPOT data at low OH exposures, this value indicates considerably faster m/z 44 production than that observed for bis(2-ethylhexyl) sebacate (C$_{26}$H$_{50}$O$_4$), a long-chain saturated hydrocarbon with two ester groups ($k = 5.58 \times 10^{-13}$ cm$^3$ molec$^{-1}$ s$^{-1}$, George et al., 2007). The TPOT m/z 44 production rate is quite similar to the production of primary products from BES oxidation ($k = 2.8 \times 10^{-12}$ cm$^3$ molec$^{-1}$ s$^{-1}$), suggesting that m/z 44 (characteristic of carboxylic acids) is a primary product of the ambient aerosol oxidation, rather than requiring multiple
steps to be produced. This is also consistent with the lack of increase at any \(m/z\) other than 44 in Fig. 6. However, we cannot rule out the possibility that an increase in an ion due to OH oxidation may be offset by a corresponding decrease by a different ion at the same nominal \(m/z\).

Oxidation of biogenic SOA in this study has some similarities to heterogeneous OH oxidation of urban aerosol. For the urban aerosol, \(m/z\) 44 enhancement of up to 20% was observed (George et al., 2008), while a 25% increase was observed for biogenic SOA (see Fig. 6). For both aerosol types, production of \(m/z\) 44 occurs relatively rapidly, with most of the production occurring in less than \(\sim\)3 day-equivalents of OH exposure. This suggests a similar rate for fragmentation reactions. Here a fragmentation reaction is defined as a reaction adding oxygen with a net carbon loss, while a functionalization reaction adds oxygen with no net carbon loss (Kroll et al., 2009). However, the total biogenic SOA mass decreased even at low exposures, quickly reaching its maximum decrease of \(\sim 15\%\) in less than \(\sim 3\) day-equivalents of OH exposure. In contrast, decrease in urban OA mass was observed until \(\sim 8\) day-equivalents of exposure, although a similar OA decrease (30%) was observed for the highest exposures (George et al., 2008). The similar OA loss at high exposures suggests similar functionalization/fragmentation branching ratios for the two aerosol types. However, the lower OH exposures needed to induce OA loss for the biogenic aerosol indicates that fewer fragmentation reactions are required, perhaps because the biogenic SOA has a product distribution weighted towards more volatile species.

Further insight into the mechanistic changes to biogenic SOA associated with heterogeneous OH oxidation can be used by investigation of the location of the OA in the Van Krevelen space, that is the molecular H:C ratio as a function of the O:C ratio (Heald et al., 2010). Because the AMS instrument used for the current study is capable only of unit mass resolution measurements, the O:C ratio is estimated using the parameterization of Aiken et al. (2008) based on the fraction of organic mass contained at \(m/z\) 44. The H:C ratio is estimated using the parameterization of Ng et al. (2011a), which is based on the organic mass fractions at \(m/z\) 43. The Ng et al. (2011a) parameterization has been evaluated for SOA only, and only for OA where \(0.05 < f_{43} < 0.25\) and \(f_{44} > 0.06\). The \(f_{43}\) and \(f_{44}\) conditions are satisfied for the current dataset (\(f_{43}\) minimum/maximum = 0.067/0.093; \(f_{44}\) minimum/maximum = 0.10/0.23). The organic aerosol composition during periods of TPOT operation was likely dominated by SOA (Macdonald et al., 2012) with the exception of 26 to 27 July, when the site was influenced by local biomass burning emissions.

Figure 7a shows the Van Krevelen plot (estimated molecular H:C vs. O:C ratios) for the AmbUV and AmbUVOH conditions (open and closed symbols, respectively), with the data points colored by OH exposure. Corresponding AmbUV and AmbUVOH conditions are connected by a line, the slope of which is plotted in Fig. 7b as a function of the OH exposure. Points that are possibly influenced by the 26 to 27 July biomass burning plume are labeled (and discussed in Sect. 3.3). Similar to Fig. 6, most chemical changes to the aerosol already occur for OH exposures below \(6.9 \times 10^{11}\) molec cm\(^{-3}\) s (\(\sim 3\) day-equivalents of oxidation). The slope of the Van Krevelen plot has previously been interpreted in terms of changes in organic functionality (e.g. Heald et al., 2010; Ng et al., 2011a). In the present study, Van Krevelen slopes of approximately \(-0.1\) and \(-0.2\) are observed for low and high OH exposures, respectively. The combination of these slopes with the observed lack of increase in any \(m/z\) other than \(m/z\) 44 (see Figs. 5 and 6) suggests the changes in the aerosol

### Table 1. Kinetic rate constants for total organics, selected \(m/z\), and PMF factors with OH.

<table>
<thead>
<tr>
<th>Reacted species</th>
<th>(K_{OH}) (cm(^3) molec(^{-1}) s(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Organics</td>
<td>(4.35 \times 10^{-12})</td>
</tr>
<tr>
<td>(m/z) 43</td>
<td>(9.18 \times 10^{-13})</td>
</tr>
<tr>
<td>(m/z) 55</td>
<td>(1.78 \times 10^{-12})</td>
</tr>
<tr>
<td>(m/z) 57</td>
<td>(1.83 \times 10^{-12})</td>
</tr>
<tr>
<td>(m/z) 91</td>
<td>(2.88 \times 10^{-12})</td>
</tr>
<tr>
<td>(m/z) 105</td>
<td>(3.40 \times 10^{-12})</td>
</tr>
<tr>
<td>OOA(_R),(_V)</td>
<td>(3.26 \times 10^{-12})</td>
</tr>
<tr>
<td>OOA(_R),(_NV)</td>
<td>(1.92 \times 10^{-11})</td>
</tr>
<tr>
<td>(m/z) 44</td>
<td>(2.29 \times 10^{-12})</td>
</tr>
<tr>
<td>OOA(_P),(_NV)</td>
<td>(4.35 \times 10^{-12})</td>
</tr>
</tbody>
</table>
are driven by significant fragmentation coupled with formation of COOH groups. Given the \( \sim 10\% \) uncertainty in the H:C parameterization (Ng et al., 2011a) and the assumptions discussed above, the obtained Van Krevelen slopes are consistent with known \( \alpha \)-pinene oxidation mechanisms. For example, one pathway for the oxidation of \( \alpha \)-pinene (\( \text{C}_{10}\text{H}_{16} \)) proceeds to the 1st-generation product \( \text{cis-pinonic acid} (\text{C}_{10}\text{H}_{16}\text{O}_3, \text{slope} = 0) \), then to \( \text{cis-pinic acid} (\text{C}_{9}\text{H}_{14}\text{O}_3, \text{slope} = -0.3) \), while subsequent oxidation to 3-methyl-1,2,3-butaneetricarboxylic acid (MBTCA) follows a slope of \(-0.2\) (Szmigielski et al., 2007). This oxidation pathway is plotted in Fig. 7a together with the TPOT data, and the two processes have similar slopes.

### 3.3 PMF analysis

Positive matrix factorization was conducted on the TPOT AMS organic mass spectra. The analyzed dataset includes the Amb, AmbUV, and AmbUVOH conditions. When applied to an AMS mass spectral time series, PMF uses temporal correlations between \( m/z \) to resolve factors. Temporal variability in the mass spectrum is therefore required for PMF analysis. In conventional AMS datasets (i.e. direct ambient sampling), this variability is governed by factors such as transport, source locations, atmospheric reactions, etc. In contrast, mass spectral variability in the TPOT dataset is governed by volatilization and OH oxidation. Therefore, separation of factors in the TPOT PMF is governed by properties of particle components such as volatility, reactivity, and reaction product formation. This provides a useful framework for discussing particle properties and aging that would not otherwise be accessible.

A 5-factor solution was selected for the TPOT PMF. This solution provides a set of factors that can each be meaningfully interpreted. Higher-order solutions include “split” factors, which can indicate an excessive number of factors (Ulbrich et al., 2009) and/or factors that cannot be meaningfully interpreted from the available data. An excerpt of the factor time series from the 5-factor solution for a typical day is shown in Fig. 8a, while the time series from a short period influenced by a local biomass burning plume at the end of the study is shown in Fig. 8b. Only excerpts of the entire time series are shown to make visible the effects of TPOT conditions, and a sample of the Amb, AmbUV, and AmbUVOH conditions are shown as shaded regions. Residuals corresponding to the excerpted time series, and mass spectral residuals for the entire campaign are shown in Fig. S6 in the Supplement. The five factors include a biomass burning organic aerosol (BBOA) factor and four oxygenated organic aerosol (OOA) factors. In addition to the PMF factors, Fig. 8 shows the time series of acetonitrile (measured by a PTR-MS) and the AMS signal at \( m/z 39 \), which is related both to potassium and organics.

Figure 8b shows a strong increase in the BBOA factor coinciding with arrival of the plume at the sampling site. In addition to the AMS BBOA factor, the effects of the plume could be observed visually and by smell, through a spike in gas-phase acetonitrile and AMS \( m/z 39 \) and connected by meteorology to known local forest fires (Macdonald et al., 2012). Because of the short temporal duration of biomass burning influences at the site, it is likely that PMF is unable to retrieve a “pure” BBOA factor. This is suggested by high residuals at \( m/z 60 \), during the biomass burning period. This \( m/z \) is a characteristic fragment of cellulose pyrolysis products such as levoglucosan. The organic mass fraction
at m/z 60 ($f_{60}$) is typically elevated above background levels during periods of biomass burning influence (Cubison et al., 2011). In the present study, $f_{60}$ reaches a maximum of 0.72 % during the biomass burning periods (compared to background values of 0.2 ± 0.05 %), while the $f_{60}$ value for the BBOA factor itself is only 0.51 %, indicating that the BBOA factor does not fully represent the variability of this important biomass burning tracer. Note further that even during the burning period, the BBOA factor never accounts for more than 80 % of the total OA, so the discrepancy between measured and PMF-reconstructed $f_{60}$ is larger than indicated by the BBOA spectrum. Additionally, slight increases in the BBOA factor are observed during periods uninfluenced by biomass burning (e.g., in Fig. 8a), while the acetonitrile and m/z 39 concentrations are more than an order of magnitude...
lower. All these observations suggest that the BBOA factor is slightly mixed with other sources.

The other four factors can all be classified as subtypes of oxygenated organic aerosol (OOA) and are distinguished by their volatility and reactive behavior. We denote these factors as OOA\textsubscript{R,V}, OOA\textsubscript{R,NV}, OOA\textsubscript{RP,NV}, and OOA\textsubscript{P,NV}, where the first subscript describes reactivity (\textquoteleft\textquoteleft R\textquoteleft\textquoteleft = consumed by reaction with OH, \textquoteleft\textquoteleft RP\textquoteleft\textquoteleft = may react with or be produced by OH, \textquoteleft\textquoteleft P\textquoteleft\textquoteleft = produced by reaction with OH), and the second subscript describes volatility (\textquoteleft\textquoteleft V\textquoteleft\textquoteleft = volatile in the TPOT, \textquoteleft\textquoteleft NV\textquoteleft\textquoteleft = non-volatile in the TPOT). Note that volatility is operationally defined from the \textasciitilde 4 °C increase in temperature from the TPOT lamp, as discussed in Sect. 3.1. Thus, for example, OOA\textsubscript{R,V} is both reactive and volatile. This is evident in Fig. 8 from the decreased OOA\textsubscript{R,V} mass going from the Amb to AmbUV conditions, and further decrease from AmbUV to AmbUVOH. The figure shows that OOA\textsubscript{R,V} is the only factor affected by the Amb/AmbUV transition. However, OH exposure causes a strong decrease in OOA\textsubscript{R,NV}, and produces OOA\textsubscript{P,NV}. Depending on the OH exposure, OOA\textsubscript{RP,NV} may be produced (right-hand side of Fig. 8a and b), unaffected (left-hand side of Fig. 8b), or reacted away (not shown here but discussed later in conjunction with Fig. 10).

The PMF factor mass spectra are shown in Fig. 9. Among the OOA factors, the \textit{f}_{44} value (which correlates with O:C), is at its minimum for the volatile component OOA\textsubscript{R,V} (\textit{f}_{44} = 0.016). Within the NV components, \textit{f}_{44} increases as the factors move from reactants to products (\textit{f}_{44}(OOA\textsubscript{R,NV}) = 0.10; \textit{f}_{44}(OOA\textsubscript{RP,NV}) = 0.18; \textit{f}_{44}(OOA\textsubscript{P,NV}) = 0.24). This provides direct evidence linking O:C with both volatility and reaction progress. A second trend in the mass spectra is the prevalence of fragments at higher \textit{m}/\textit{z} for the more reactive and volatile species. As shown in the Supplement (Figs. S7 and S8 in the Supplement), the OOA\textsubscript{R,V} and OOA\textsubscript{R,NV} spectra are both quite similar to semi-volatile oxygenated organic aerosol SV-OOA spectra resolved from conventional PMF analyses, while the OOA\textsubscript{R,NV} and OOA\textsubscript{P,NV} spectra are more similar to previously reported LV-OOA (Ng et al., 2011b). The BBOA factor shows somewhat elevated signal at \textit{m}/\textit{z} 60 and 73, which both individually constitute \textasciitilde 0.5 % of the total organic signal. These \textit{m}/\textit{z} result from fragmentation of cellulose pyrolysis products such as levoglucosan and are frequently used as tracers for biomass burning influence in AMS spectra. The contributions of these two \textit{m}/\textit{z} to the BBOA spectrum are larger by approximately a factor of 2 or more than their contributions to any other factor.

Figure 10 shows the effect of OH exposure on TPOT PMF factor mass concentrations. The BBOA, OOA\textsubscript{R,V}, OOA\textsubscript{R,NV}, and OOA\textsubscript{RP,NV} are shown as the ratio of mass concentrations for the AmbUVOH to AmbUV conditions. For OOA\textsubscript{P,NV}, the difference between AmbUVOH and AmbUV mass concentrations is instead shown because of the sometimes low AmbUV OOA\textsubscript{P,NV} concentrations, which lead to very high and scattered AmbUVOH/AmbUV ratios. The top panel shows a decrease in BBOA, OOA\textsubscript{R,V} and OOA\textsubscript{R,NV} mass for all OH exposures. OOA\textsubscript{R,V} and OOA\textsubscript{R,NV} are shown to be quite reactive, although there is a great deal of scatter in the measurements. Reaction at this initial stage may depend strongly on details of the particle composition. Similar to the behavior of individual \textit{m}/\textit{z} shown in
Fig. 11. Ambient organic aerosol composition (i.e. Amb condition) in terms of TPOT PMF factors. The stacked factor time series equal the total organic mass.

Fig. 6, most of the decrease in OOA\textsubscript{R,V} (~75 % consumed) and OOA\textsubscript{R,NV} (~50 % consumed) occurs within ~3 day-equivalents of photochemical aging. The reactivity of these factors suggests that ambient SV-OOA, which has a similar mass spectrum, may also react quite quickly in the atmosphere. Aging-induced reductions in SV-OOA-like species have previously been observed for airmasses in southwestern Ontario, Canada (Slowik et al., 2011).

Somewhat different behavior is observed for BBOA. While ~35 % of BBOA is consumed within ~3 day-equivalents, this value continues to increase, reaching ~60 % or more after ~6 day-equivalents of exposure. This indicates that BBOA is not a fully conserved tracer for the AMS, consistent with both the spectral changes in Fig. 6 and ambient studies of biomass burning plumes (Capes et al., 2008). The figure does not show evidence for significant decreases in BBOA for less than ~1 day-equivalent of aging.

A further contrast is observed for the OOA\textsubscript{RP,NV} factor. Here an increase in mass on the order of ~20 % is observed during approximately the first 3 day-equivalents of aging. However, higher exposures yield no net change relative to the unreacted concentrations. This suggests that OOA\textsubscript{RP,NV} can be considered an early- or intermediate-stage reaction product, but itself reacts given sufficiently high OH exposures. On the other hand, while OOA\textsubscript{P,NV} forms rapidly, with most production occurring, within ~3 day-equivalents of aging, there is no evidence for its consumption within the range of exposures studied, suggesting that the end product of particle-phase oxidation resembles OOA\textsubscript{P,NV}. As noted previously, OOA\textsubscript{P,NV} is spectrally similar to AMS LV-OOA spectra. Figure 10 thus supports the current understanding that atmospheric aging pushes aerosols of varying composition towards an LV-OOA-like state (Ng et al., 2010; Jimenez et al., 2009). The figure also indicates that heterogeneous oxidation and/or gas-phase oxidation of organic material partitioning between the particle and gas phase can significantly affect both primary and secondary aerosol components on atmospherically-relevant timescales.

The TPOT PMF analysis discussed above can also provide insight into the composition and volatility/reactivity-based properties of the (unreacted) ambient aerosol. Figure 11 shows the TPOT PMF representation of particle composition for 19 and 21 July. These days take place during the early and late stages, respectively, of a period of increasing temperatures, biogenic emissions, and biogenic SOA. On 19 July (lower panel), when the temperature and biogenic emissions are lower (daily maximum temperature ~19\textdegree C), there is lower total organic mass and changes to the composition during the day are negligible. In contrast, 21 July (maximum temperature ~23 \textdegree C) shows a 50–100 % increase...
in the total organic mass and changes in composition relative to those on 19 July at the same time of day. Peak concentrations of organic aerosol, biogenic VOCs, and OH radicals all occur during mid to late afternoon (Macdonald et al., 2012), and the strong diurnal patterns observed for these species are consistent with local production. Figure 11 shows that the increase in organic aerosol mass during the day on 21 July caused by local production primarily yields increases in the OOA$_{RP,NV}$ and OOA$_{RP,NV}$ components. OOA$_{RP,NV}$-like compounds may also be produced, but the increased midday temperatures could prevent them from significantly partitioning to the particle. A small increase in OOA$_{RP,NV}$ during the middle of the day on 21 July is also evident. While highly oxygenated, LV-OOA-like factors such as OOA$_{RP,NV}$ and OOA$_{RP,NV}$ are frequently associated with long-range transport, the evidence presented in Figs. 10 and 11 suggests that they may also form during rapid local production in some cases. This is consistent with rapid formation of highly oxidized SOA observed in polluted airmasses in southwestern Ontario, Canada (Slowik et al., 2011).

Coupling the reactivity/volatility-based description of ambient aerosol composition shown in Fig. 11 with the PMF factor reactivity characterization in Fig. 10 enables a detailed description of the present aerosol composition. Using such reactivity and volatility coordinates makes for easier comparison to the framework presented by Jimenez et al. (2009) for the aging of organic aerosols. It also enables projection of the future changes in composition during the particle’s atmospheric lifetime (albeit neglecting future SOA formation from gas-phase precursors due to TPOT SOA yield properties). Accessing such a description requires online manipulation of the aerosol composition by a system such as the TPOT; the information is not directly accessible using conventional ambient measurements. This suggests that controlled manipulation of ambient aerosol may be a useful tool for characterizing particle properties and predicting their evolution during the course of atmospheric aging. Such an approach has been applied to laboratory-generated SOA from $\alpha$-pinene ozonolysis (at concentrations many times higher than ambient) (George and Abbatt, 2010), yielding reactant- and product-like PMF factors (see Figs. S8 and S9 in the Supplement). The mass spectrum of the reactant-like factor is quite similar to OOA$_{RV}$, however its volatility is unknown. Interestingly, the product-like factor mass spectrum is similar to OOA$_{RP,NV}$ rather than OOA$_{RP,NV}$. This is consistent with the trend towards decreased OH-induced spectral changes for $\alpha$-pinene ozonolysis SOA relative to the biogenic SOA; for example, George and Abbatt (2010) observed little effect of OH on $m/z$ 43, while for biogenic SOA a significant decrease is observed (see Fig. 6). Further experiments are required to fully elucidate the effects of the initial particle composition on reaction products and pathways.

The general approach herein, in which PMF analysis is performed on organic aerosol subject to forced changes in the chemical and/or physical environment (e.g. temperature or relative humidity), could be applied more widely to better determine the nature of organic particulates. However, this approach does add to the complexity of the data analysis. The present analysis is simplified by the dominance of a single source class (biogenic SOA). Other locations may have more source complexity, however it is not necessary to apply PMF to an entire dataset at once. Rather, periods characteristic of particular source/aerosol types can be analyzed separately (e.g. if biomass burning periods were more prevalent in the current study, separate TPOT PMF analyses could be performed on biogenic and biomass burning-dominated periods). For each period segregated in this way, the UV/oxidation-based variability remains, and these properties would likely still be reflected in the retrieved PMF factors.

4 Conclusions

We present the first field deployment of the Toronto Photooxidation Tube (TPOT), a field-deployable flow reactor for the controlled OH oxidation of ambient aerosol. The deployment took place during summer in a forested location strongly influenced by biogenic organic aerosol formation during warm periods. Aerosols were continuously sampled, alternating between three conditions: (1) unreacted ambient aerosol (“Amb”); (2) ambient aerosol UV light and subjected to a $\sim$4 to 10$^\circ$C temperature increase (“AmbUV”); and (3) ambient aerosol additionally subjected to OH exposures ranging from approximately $2 \times 10^{10}$ to $2 \times 10^{12}$ molec cm$^{-3}$s (“AmbbUVOH”), corresponding to up to $\sim$10 day-equivalents of atmospheric photochemical aging. Particle mass and composition were monitored with a time-of-flight aerosol mass spectrometer (C-ToF-AMS). Due in large part to the geometry of the TPOT system, significant SOA generation was not observed and the dominant reactive processes were instead heterogeneous oxidation and/or gas-phase reaction of semivolatile species partitioning from the particle to the gas phase.

The AmbUV condition caused a reduction in the organic mass on the order of 10 to 25%, depending on the oxygenation of the unperturbed aerosol. Reaction with OH caused further decreases in the organic mass on the order of up to an additional 25%. Similar decreases were observed throughout the mass spectrum, with the exception of $m/z$ 44 ($CO_2^+$ ion) and $m/z$ 45.

Positive matrix factorization analysis of the TPOT AMS organic mass spectra yielded a primary biomass burning factor and four oxygenated organic aerosol factors directly related to reactivity and volatility. This classification of aerosol components is enabled by the TPOT-modulated perturbation of aerosol composition, and is not otherwise accessible. Analysis of the PMF factors and individual $m/z$ both indicate rapid formation of highly oxygenated organic aerosol, likely with a strong contribution from organic acid functional
groups. Evidence for early/intermediate-stage OH reaction products is also observed. OH oxidation significantly affects particle composition even for exposures corresponding to < 1 day-equivalent of atmospheric aging, and most reaction with biogenic SOA appears complete after ~3 day-equivalents. The particle-phase reaction end products have mass spectra similar to the low-volatility oxygenated organic aerosol (LVOOA) factors widely reported in the literature (e.g. Zhang et al., 2005; Lanz et al., 2007; Jimenez et al., 2009; Ulbrich et al., 2009), thus providing supporting evidence that aged organic aerosol does arise through OH-driven oxidation processes. This general approach of PMF analysis of organic aerosol subject to forced changes in the chemical and/or physical environment (e.g. temperature or relative humidity), could be applied more widely to better determine the nature of organic particulates.

Supplementary material related to this article is available online at: http://www.atmos-chem-phys.net/12/9775/2012/acp-12-9775-2012-supplement.pdf.

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References


J. G. Slowik et al.: Real-time, controlled OH-initiated oxidation of biogenic secondary organic aerosol


