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Identifying precursors and aqueous organic aerosol formation pathways during the SOAS campaign

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Abstract. Aqueous multiphase chemistry in the atmosphere can lead to rapid transformation of organic compounds, forming highly oxidized, low-volatility organic aerosol and, in some cases, light-absorbing (brown) carbon. Because liquid water is globally abundant, this chemistry could substantially impact climate, air quality, and health. Gas-phase precursors released from biogenic and anthropogenic sources are oxidized and fragmented, forming water-soluble gases that can undergo reactions in the aqueous phase (in clouds, fogs, and wet aerosols), leading to the formation of secondary organic aerosol (SOAAO). Recent studies have highlighted the role of certain precursors like glyoxal, methylglyoxal, glycolaldehyde, acetic acid, acetone, and epoxides in the formation of SOAAQ. The goal of this work is to identify additional precursors and products that may be atmospherically important. In this study, ambient mixtures of watersoluble gases were scrubbed from the atmosphere into water at Brent, Alabama, during the 2013 Southern Oxidant and Aerosol Study (SOAS). Hydroxyl (OH•) radical oxidation experiments were conducted with the aqueous mixtures collected from SOAS to better understand the formation of SOA through gas-phase followed by aqueous-phase chemistry. To-

tal aqueous-phase organic carbon concentrations for these mixtures ranged from 92 to 179 μ M-C, relevant for cloud and fog waters. Aqueous OH-reactive compounds were primarily observed as odd ions in the positive ion mode by electrospray ionization mass spectrometry (ESI-MS). Ultra high-resolution Fourier-transform ion cyclotron resonance mass spectrometry (FT-ICR-MS) spectra and tandem MS (MS–MS) fragmentation of these ions were consistent with the presence of carbonyls and tetrols. Products were observed in the negative ion mode and included pyruvate and oxalate, which were confirmed by ion chromatography. Pyruvate and oxalate have been found in the particle phase in many locations (as salts and complexes). Thus, formation of pyruvate/oxalate suggests the potential for aqueous processing of these ambient mixtures to form SOA_{AQ}.

1 Introduction

Aqueous multiphase chemistry has the potential to alter the climate-relevant properties and behavior of atmospheric aerosols. It is well established that a major pathway for secondary organic aerosol (SOA) formation is via the partitioning of semi-volatile products of gas-phase photochemical reactions into preexisting organic particulate matter (Seinfeld and Pankow, 2003). Semi-volatile partitioning theory is widely used to model SOA (Odum et al., 1996; Seinfeld and Pankow, 2003; Donahue et al., 2006). However differences between organic aerosol mass/properties predicted via this formation mechanism and those measured in the atmosphere suggest that other processes (e.g., aqueous chemistry) may also contribute (Foley et al., 2010; Hallquist et al., 2009).

Recent studies have highlighted the importance of water-soluble organic gases (WSOGs), liquid water, and condensed-phase reactions to SOA formation and properties (Ervens et al., 2011; Monge et al., 2012; Carlton and Turpin, 2013). Biogenic and anthropogenic gas-phase precursors are oxidized to form WSOGs such as glyoxal, methylglyoxal, glycolaldehyde, and acetone (Spaulding et al., 2003). These WSOGs are too volatile to form SOA through absorptive partitioning, but they can undergo aqueous reactions in clouds, fogs, and wet aerosols to form low-volatility products and "aqueous" SOA (SOA_{AO}) (Blando and Turpin, 2000; Ervens et al., 2004, 2008, 2011; Kroll et al., 2005; Liggio et al., 2005; Lim et al., 2005; Heald et al., 2006; Loeffler et al., 2006; Sorooshian et al., 2006; Volkamer et al., 2006, 2007; De Haan et al., 2009a; El Haddad et al., 2009; Rossignol et al., 2014). Inclusion of aqueous chemistry of clouds, fogs, and wet aerosols in models and experiments helps to explain discrepancies in atmospheric observations of SOA that are not explained by semi-volatile partitioning theory, particularly high atmospheric O/C ratios; enrichment of organic aerosol aloft; and formation of oxalate, sulfur-, and nitrogencontaining organics and high-molecular-weight compounds (Kawamura and Ikushima, 1993; Kawamura et al., 1996; Crahan et al., 2004; Kalberer et al., 2004; Herrmann et al., 2005; Altieri et al., 2006; Carlton et al., 2006; Heald et al., 2006; Volkamer et al., 2007, 2009; Nozière and Cordova, 2008; De Haan et al., 2009b; El Haddad et al., 2009; Galloway et al., 2009; Shapiro et al., 2009; Lim et al., 2010; Lin et al., 2010; Nozière et al., 2010; Perri et al., 2010; Sareen et al., 2010; Schwier et al., 2010; Sorooshian et al., 2010; Sun et al., 2010; Ervens et al., 2011, 2013; Lee et al., 2011; Tan et al., 2012; He et al., 2013; Gaston et al., 2014; Ortiz-Montalvo et al., 2014). Although uncertainties are large, modeling studies show that SOAAQ is comparable in magnitude to "traditional" SOA (Carlton et al., 2008; Fu et al., 2008, 2009; Gong et al., 2011; Myriokefalitakis et al., 2011; Liu et al., 2012; Lin et al., 2014). However, SOAAO precursors and their chemical evolution remain poorly understood.

Much of what we know about aqueous chemistry leading to SOA_{AQ} formation is derived from laboratory studies with single precursors hypothesized to be important; however, the most important precursors for SOA_{AQ} formation in the ambient environment may remain unidentified. A small number of studies conducted with ambient mixtures have provided insights into the pathways of SOA_{AQ} formation. For example, photochemical oxidation of aerosol filter samples and cloud water from Whistler, British Columbia, suggest that watersoluble organic compounds of intermediate volatility (e.g., *cis*-pinonic acid) can be important precursors for SOA_{AQ} (Lee et al., 2012). Pyruvic acid oxidation experiments on Mt. Tai, China cloud water suggested a slowing of pyruvic acid oxidation presumably due to competition for OH radicals with the complex dissolved cloud water organics (Boris et al., 2014). However, further ambient measurements are needed to identify precursors important for ambient SOA_{AQ} formation in atmospheric waters.

This work reports, for the first time, results of aqueous OH radical oxidation experiments conducted in ambient mixtures of water-soluble gases. Ambient mixtures were collected in the southeastern US during the Southern Oxidant and Aerosol Study (SOAS) in the summer of 2013; experiments were used to identify water-soluble gases that may serve as precursors of atmospheric aqueous SOA. This region has experienced an overall cooling trend in surface temperature over the second half of the twentieth century, compared to the warming trend observed elsewhere in the US (Robinson et al., 2002; Goldstein et al., 2009; Portmann et al., 2009). Biogenic sources dominate emissions in this region with varying degrees of impact from anthropogenic sources. Measurements by Nguyen et al. (2014) and model results by Carlton and Turpin (2013) indicate the significance of anthropogenic aerosol liquid water (ALW) in this region and support a role for ALW in SOAAQ formation. In the southeastern US, photochemistry and abundant liquid water coexist, making it an ideal location to study SOAAO formation through gas-phase followed by aqueous-phase chemistry. The objective of this work is to identify WSOGs important to SOA_{AO} formation. Since OH oxidation experiments were conducted in dilute solution, we will also identify products expected through cloud/fog processing of ambient WSOG mixtures. Products may differ in aerosols, where solute concentrations are higher, and radical-radical chemistry and acid-catalyzed reactions (e.g., epoxide ring-opening reactions yielding tetrols and organosulfates from isoprene epoxydiol, IEPOX) are important. We expect that aqueous chemistry in clouds, fogs, and wet aerosols is a sink for reactants identified herein and that this work will motivate laboratory studies and chemical modeling of newly identified aerosol/cloud precursors.

2 Methods

Samples of ambient water-soluble mixtures collected in mist chambers during the SOAS field study were used to conduct controlled aqueous OH radical oxidation experiments. Mass spectral techniques were used to tentatively identify compounds with decreasing abundance in aqueous OH oxidation experiments, but not in control experiments with the aim of identifying new aqueous chemistry precursors for further study. Ion chromatography (IC) was used to identify selected products.

2.1 Mist chamber field sampling at SOAS (in Brent, AL)

Water-soluble gases were scrubbed from filtered ambient air at the Centerville ground site in Brent, AL, during SOAS. Samples were collected from 1 June to 14 July 2013 from 1 m above the sampling station roof through a 1.3 cm OD Teflon inlet (approximately 1.7 m in length). Four mist chambers (Anderson et al., 2008a, b; Dibb et al., 1994; Hennigan et al., 2009) were operated in an air-conditioned trailer (indoor temperature: $25 \,^{\circ}$ C) at $25 \,\text{Lmin}^{-1}$ in parallel for 4 h, typically 2–3 times each day between 07:00 and 19:00 CDT. Particles were removed by passing the ambient air through a pre-baked quartz fiber filter (QFF) (Pall, 47 mm) prior to introduction into the mist chamber.

The mist chambers were operated with 25 mL of $17.5 \pm 0.5 M\Omega$ ultra-pure water; additional water was added during the run to replace water lost by evaporation. Samples from all four mist chambers were composited daily and frozen in 35–40 mL (experiment-sized) aliquots. Field water blanks were collected concurrently from the same water supply, transported, and stored with the samples. Total organic carbon (TOC) concentrations ranged from 45 to 180 μ M-C (Supplement Table S1). At the beginning of the study, mist chambers were baked at 500 °C for 4 h. Prior to and at the end of a sampling day, each mist chamber was cleaned using a 5 min deionized (DI) water wash step.

Based on daily forecast predictions, certain days were selected for intensive sampling (Table S1). Intensive sampling during SOAS was conducted on days when high levels of isoprene, sulfate, and NO_x were predicted by the National Center for Atmospheric Research (NCAR) using the FLEXible PARTicle dispersion model (FLEXPART) (Stohl et al., 2005) and the Model for Ozone and Related Chemical Tracers (MOZART) (Emmons et al., 2010). In general, mist chamber samples on intensive sampling days had higher organic content (TOC = 92–179 μ M-C), and hence we focused our experiments on those days and included two additional days from the non-intensive period that had high TOC values (Table 1).

2.2 Aqueous OH radical oxidation in a cuvette chamber

Ambient SOAS field samples were exposed to OH radicals in a custom-built photochemical temperature-controlled $(25 \,^{\circ}C)$ quartz cuvette reaction chamber. Ten screw-capped quartz cuvettes (Spectrocell Inc., Oreland, PA) containing 3 mL of sample were placed equidistant around a 254 nm mercury lamp (Heraeus Noblelight, Inc. Duluth, GA) housed in a quartz sheath (Ace Glass Inc., Vineland, NJ). A solar spectrum lamp was not used because the objective was to produce OH radicals by H₂O₂ photolysis, rather than to mimic tropospheric photolysis. The chamber was protected from ambient light by covering in aluminum foil. OH radicals $(1.25 \times 10^{-2} \,\mu\text{M} \text{ [OH] s}^{-1})$ were generated in situ by photolysis of 125 µM H₂O₂, added to each cuvette prior to inserting the lamp. While we can calculate the OH production rate from hydrogen peroxide photolysis $(1.25 \times 10^{-2} \,\mu\text{M})$ $[OH] s^{-1}$, the concentration of OH in the reaction vessel depends also on the reactivity of the organics. If the WSOG mix behaves similarly to glyoxal, OH concentrations would be on the order of 10^{-12} M (similar to (Tan et al., 2009). Cuvettes were removed at t = 10, 20, 30, 40, 60, 80, 100, 120,and 150 min, and any remaining H2O2 was destroyed by addition of 36 μ L of 1 % catalase (Sigma; 40 200 units mg⁻¹). A duplicate cuvette was removed at $t = 40 \min$ to calculate method precision. The following control experiments were performed: (1) sample + H_2O_2 , (2) sample + UV, (3) $H_2O_2 + UV$, and (4) field water blank + OH. Replicate experiments were performed on selected samples. Ambient conditions for sample collection are given in Table 1 for samples used in experiments.

2.3 Analytical methods

Samples and field water blanks from all collection days were characterized by TOC analysis (Shimadzu 5000A) and electrospray ionization mass spectrometry (ESI-MS; HP -Agilent 1100). IC (Dionex ICS 3000) was used to analyze organic anions and track the formation of products and intermediates. Samples at each reaction time were analyzed by ESI-MS in positive and negative ion modes to identify precursors and products. Selected samples were also analyzed by ultra-high-resolution electrospray ionization Fourier-transform ion cyclotron resonance mass spectrometry (ESI-FT-ICR-MS) and tandem MS (MS-MS) on a Thermo-Finnigan LTQ-XL at Woods Hole Oceanographic Institute, MA, to determine elemental composition and extract structural information on precursors. Analytical details and quality control measures have been described previously (Perri et al., 2009). Briefly, the ESI quadrupole mass spectrometer was operated in positive and negative ion modes over a mass range of 50–1000 amu. In the negative ion mode, the mobile phase consisted of 1 : 1 methanol to 0.05 % formic acid in water; in the positive ion mode, it consisted of 0.05~%formic acid in water. The fragmentor and capillary voltages for the ESI-MS were set at 40 and 3000 V (nitrogen drying gas; 10 L min⁻¹; 350 °C), respectively. Nitrobenzoic acid in the negative ion mode and caffeine in the positive ion mode were used as mass calibrants. Standard mixtures were analyzed with each experimental sequence: acetic acid, pyruvic acid, nitric acid, succinic acid, tartaric acid, ammonium sulfate, and oxalic acid in the negative ion mode and glyoxal, methylglyoxal, and glycolaldehyde in the positive ion mode.

ESI ionization efficiency varies with sample mix and over time. However, the mass spectra of the mist chamber samples

Collection	Collection	μМ ТОС	<i>T</i> (°C)	RH (%)	O ₃ (ppbv)
date	time		all day	all day	all day
			(coll. time)	(coll. time)	(coll. time)
11 Jun 2013	7.00 am-7.00 pm	139.5	22-32	53–99	9.9–38.2
			(23–32)	(53–98)	(11.3–38.2)
12 Jun 2013	7.00 am-7.00 pm	179.7	23-33	48–94	13.1-41.8
			(23–33)	(48–94)	(13.1-41.8)
15 Jun 2013	7.00 am-7.00 pm	117.0	17-31	45–94	11.6-53.4
			(18–31)	(45–90)	(11.6–53.4)
16 Jun 2013	7.00 am-7.00 pm	108.2	22-32	53-93	2.6-41.7
			(22–32)	(53-84)	(23.1-41.7)
20 Jun 2013	8.00 am-5.00 pm	131.5	20-30	55–98	4.8-52.2
			(21-30)	(59–94)	(6.2–42.8)
21 Jun 2013	10.00 am-6.00 pm	104.4	20-30	50-93	16.6-45.2
			(25-30)	(50-78)	(30.3–45.2)
29 Jun 2013	7.00 am-7.00 pm	92.0	21-31	43-100	16.3-53.7
			(22–31)	(43–100)	(16.3–53.7)
30 Jun 2013	7.00 am-7.00 pm	98.7	20-30	38-100	1-53.5
			(20–30)	(38–100)	(1–53.5)

Table 1. Sample collection dates for which experiments were run and their sampling conditions. Temperature, relative humidity, and ozone ranges are shown both for the entire day and collection time period. Total organic carbon (TOC) accuracy and precision are verified with potassium hydrogen phthalate (KHP) standards to be better than 5 % (Perri et al., 2009).

were similar across experimental days, and the variability in the glyoxal, methylglyoxal, and glycolaldehyde standard ESI signals were 4–7% across analysis days, suggesting that ion abundance trends (Fig. S1–S2 in the Supplement) will reflect concentration trends. Six injections were averaged for each sample, and data were retained for each ion abundance greater than 0 within 95% confidence intervals. Ions were considered to be above detection limits when peaks were greater than the average plus 3 standard deviations of the water blank.

Organic acids were measured by IC (IonPac AS11-HC column; 30 °C, AG11-HC guard column) with conductivity detection (35 °C), using a Milli-Q water eluent and the potassium hydroxide (KOH) gradient method. For oxalate, the method precision is 22 %, calculated as a pooled coefficient of variation (CV) from pairs of cuvettes removed at t = 40 min. The analytical precision for oxalate is 19 % (pooled CV) based on replicate analysis of 30 % of samples. Analytical accuracy for oxalate is 7 %. The limit of detection (LOD) for oxalate by this protocol has been previously determined to be 0.1 μ M (Perri et al., 2009).

Samples from 15 and 30 June 2013 were analyzed using ultra-high-resolution FT-ICR-MS in the positive ion mode using 1:1 methanol and water as the mobile phase at $4\,\mu\text{L}\,\text{min}^{-1}$, capillary temperature of 260 °C, and 3.8–4.2 kV spray voltage. Weekly analysis of standards (caffeine, peptide MRFA, Ultramark, sodium dodecyl sulfate (SDS), and sodium taurocholate) verified the mass accuracy <2 ppm. Previously pyruvic acid and peroxyhemiacetal standards analyzed with the same protocol were within

2–10 ppm. Five precursor masses were isolated (isolation width: m/z = 2) and fragmented by collision-induced dissociation (CID) (normalized collision energy: 26–33 %) with helium in the ion trap (IT) and infrared multiphoton dissociation (IRMPD) with a CO₂ laser. Elemental composition (within ± 1 ppm) and double-bond equivalents of ions were calculated by the Midas Molecular Formula Calculator (v1.1). No restrictions were placed on the number of carbon, hydrogen, oxygen, nitrogen, sodium, and sulfur atoms included in the molecular formula calculations.

Compounds are detected in the ESI-MS by forming cluster ions with hydrogen, sodium, or ammonium in the positive ion mode; compounds are sometimes hydrated with water or methanol. In the negative ion mode, ions are deprotonated.

3 Results and discussion

OH oxidation experiments at concentrations relevant to cloud/fog water were conducted on samples collected on 11, 12, 15, 16, 20, 21, 29, and 30 June 2013, days on which TOC was highest, ranging from 92 to $179 \,\mu$ M-C in samples.

3.1 Precursors in SOAS samples

The concentration dynamics in experiments conducted with the eight daily composites were similar. Positive ions at m/z 125, 129, 143, 173, and 187 exhibited reactant-like trends (Fig. 1; 30 June sample + OH), showing decreasing signal intensity with increasing exposure to OH. These ions disappeared after 40 min of oxidation. Together, these ions ac-

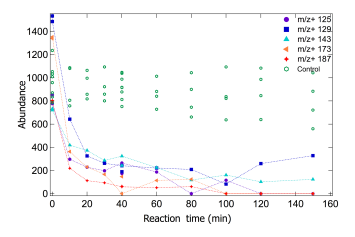


Figure 1. Positive ions (ESI-MS) exhibiting precursor-like trends during aqueous OH radical oxidation experiments with the ambient mixtures collected on 30 June. All days show similar trends, with all 5 reactant masses showing statistically significant decreasing trends as compared to the control experiments. Controls (sample + UV, sample + H₂O₂) shown for m/z 187; other masses show similar trends.

count for roughly 30% of the total ion current in the positive mode in the experiment samples. In control experiments, the abundance of these ions did not change over time, as illustrated in Fig. 1 (sample + UV and sample + H₂O₂) for the positive ion at m/z 187 in samples collected on 15 and 30 June. Hence, reaction with UV or H₂O₂ alone does not explain the decreasing signals in the presence of OH radicals. In control experiments where we generated OH radicals in field water blanks, these ions were not observed, confirming that they are not contaminants from the water source. Experiments conducted on all sample days showed the same reactants decreasing with exposure to OH, indicating that the water-soluble organics captured from the ambient daytime air in the mist chambers varied little across the study.

Elemental formulae assigned to precursor ions by Midas with corresponding MS–MS fragmentation data for the ions from 15 and 30 June samples are shown in Table 2. Both sampling days showed similar fragmentation spectra, consistent with the presence of the same parent compounds on both days, despite potential differences in the air mass on these days. MS–MS spectra were not obtained under the acquisition conditions of this work for positive ions at m/z 143, 129, and 125. Proposed structures for the positive ions at m/z 187 and 173 based on MS–MS data are shown in Figs. 2 and 3. The O / C ratio for the reactant masses range from 0.1 to 0.8. In the following sections, we discuss individual reactants observed by ESI-MS. The discussion below is focused on the molecular formulas, rather than the detected ions, i.e., the parent compounds with Na⁺ or H⁺.

m/z 187: this is the ion with the highest m/z observed by ESI-MS in the positive ion mode that follows a precursor-like trend (Figs. 1 and 2). The composition C₇H₁₆O₄Na is

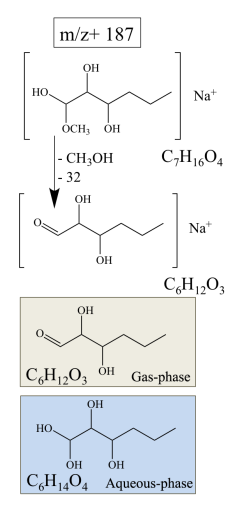


Figure 2. Proposed structure for the positive ion at m/z 187. The top structure is the parent compound detected as a reactant in the ESI-MS; the following structures show the MS–MS fragments. This compound would take the forms shown in the shaded boxes if present in atmospheric air and water.

assigned by the Midas Molecular Formula Calculator based on the exact mass 187.0942 from FT-ICR MS, corresponding to the composition $C_7H_{16}O_4$ for the neutral compound. The MS–MS shows loss of methanol, to give a product ion at m/z 155.0680, corresponding to the neutral molecular formula, $C_6H_{12}O_3$, consistent with expectations for a sodium ion complex of a dihydroxy hemiacetal in methanol solution, shown in Fig. 2. In the absence of methanol, this compound would appear hydrated with water as the $C_6H_{14}O_4$ tetrol, as shown in the blue box in Fig. 2. The corresponding gas-phase compound is shown in the tan box in Fig. 2.

E-2-hexenal and Z-3-hexenal are unsaturated aldehydes that have frequently been detected during field studies and are emitted to the atmosphere from vegetation due to leaf wounding (O'Connor et al., 2006). The gas-phase oxidation of these two green-leaf volatiles, as shown in Fig. 4a and b,

Precursor peak using ESI-MS	Positive mode m/z (using FT-ICR)	[M+Na] ⁺ or [M+H] ⁺	Mol. wt.	Double-bond equivalents
187	187.0942	C7H16O4Na	164.1043	0
	155.0680	$C_6H_{12}O_3Na$	132.0786	1
173	173.0782	C ₆ H ₁₄ O ₄ Na	150.0887	0
	141.0523	C ₅ H ₁₀ O ₃ Na	118.0625	1
	129.0524	$C_4H_{10}O_3Na$	106.0625	0
143	143.0676	C ₅ H ₁₂ O ₃ Na	120.0781	0
129	129.0520	$C_4H_{10}O_3Na$	106.0625	0
125	125.096	C ₈ H ₁₃ O	124.0883	3

Table 2. Elemental formulas assigned to precursor ions using ESI-FT-ICR MS in the positive ionization mode and Midas Molecular Formula

 Calculator. MS–MS fragmentation data are also shown.

could explain the presence of $C_6H_{12}O_3$ in the gas phase and $C_6H_{14}O_4$ in the aqueous phase (Fig. 2).

m/z 173: on most sampling days this reactant mass has the highest abundance in the positive mode ESI mass spectra (Supplement Fig. S1). Similar to other reactant peaks, it reacts away within the first 40 min of exposure to OH in the cuvette chamber (Fig. 1). The Midas-suggested molecular formula for this parent ion (m/z 173.0782) and its two fragment ions at m/z 141.0523 and 129.0524 are C₆H₁₄O₄, C₅H₁₀O₃, and C₄H₁₀O₃, respectively (a reactive parent ion with the formula C₄H₁₀O₃ was also observed and is discussed below).

A likely structure for positive mode m/z 173 is shown in Fig. 3. In this case the compound is proposed to be a C₅H₁₀O₃ aldehyde in the gas phase (tan box in Fig. 3) and a C₅H₁₂O₄ tetrol in water (blue box in Fig. 3). In the FT-ICR-MS it is seen hydrated with methanol. The parent ion at m/z 173 loses methanol to form C₅H₁₀O₃ (m/z 141.0523), and it also loses C₂H₄O to form C₄H₁₀O₃ (m/z 129.0524). The aqueous oxidation precursor observed as m/z 173 and tentatively identified as the C₅H₁₀O₃ aldehyde shown in tan in Fig. 3 could be derived from another green-leaf volatile. Specifically, C₅H₁₀O₃ is consistent with the gas-phase oxidation product of (*E*)-2-methyl-2-butenal, another green-leaf volatile (Fig. 4c) (Jiménez et al., 2009; Lanza et al., 2008).

Positive ions at m/z 143, 129, and 125: no fragments were observed for these reactants under conditions of MS–MS acquisition in this work. The Midas-predicted molecular formulae for the ions at m/z 143.0676, 129.0520, and 125.096 are C₅H₁₂O₃, C₄H₁₀O₃, and C₈H₁₂O, respectively. Interestingly, the reactant detected at m/z 129 has the same mass as a fragment of the parent ion at m/z 173 discussed earlier (and the structure of the C₄H₁₀O₃ fragment shown in Fig. 3 is a possible structure for m/z 129).

Methodological limitations: in this work, we aim to collect the ambient mix of water-soluble gases into water at concentrations comparable to those found in clouds and fogs with the purpose of simulating cloud/fog-relevant OH oxida-

tion chemistry and identifying previously unrecognized precursors of aqueous chemistry. Below we discuss limitations with respect to our ability to collect and store these aqueous mixtures and with respect to our ability to identify the OHreactive compounds collected.

Mist chamber collection times (4 h) were selected with the aim of collecting ambient mixtures of water-soluble gases near Henry's law equilibrium. Two pieces of evidence suggest that gas-aqueous partitioning of the WSOGs is close to Henry's law equilibrium in our samples. In previous testing conducted at a different east coast location that is a recipient of long-range transport (i.e., central New Jersey), we found that WSOG concentrations in the mist chamber leveled off after 1-3 h of ambient sampling, suggesting that the collected WSOG mixture approaches Henry's law equilibrium over these collection times. This is consistent with the one measurement we have of breakthrough at SOAS, where we ran two mist chambers in series and found TOC concentrations within ± 11 % of each other. These measurements suggest that a representative mixture of the water-soluble gases entering the mist chambers was collected.

However, WSOGs can be lost during sampling and storage through (1) losses in tubing and by adsorption to the QFF during collection; (2) reactions in the mist chamber during collection with water-soluble ambient oxidants capable of penetrating the inlet (e.g., ozone); and (3) losses during storage post collection. The QFF removes particles upstream of the mist chambers. In the early stages of sampling, on the clean filter, adsorption of gases on the filter will reduce the concentrations of gases sampled by the mist chamber until these gases reach gas-phase-adsorbed-phase equilibrium. Using glyoxal as a WSOG surrogate and the work of Mader and Pankow (2001), we predict that the measured WSOG in the mist chamber will be depleted for less than 2% of our sampling time (after $< 0.1 \text{ m}^3$) due to loss to the filter. Thus, we expect water-soluble gases to penetrate through the QFF very efficiently for collection in the mist chamber wa-

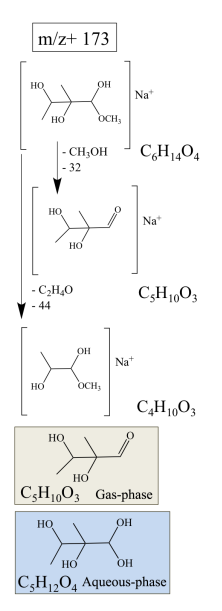


Figure 3. Proposed structure for the positive ion at m/z 173. The top structures in each panel are the parent compound detected as a reactant in the ESI-MS; the following structures show the MS–MS fragments. This compound would take the forms shown in the shaded boxes if present in atmospheric air and water.

ter. Losses to Teflon inlets and chamber walls (Krechmer et al., 2016) can be significant and variable and may reduce the number of species we are able to collect and identify in this work. While OH radicals are unlikely to penetrate the inlet, ozone might. Thus some ozone could be scrubbed by the mist chambers and could result in oxidation of some unsaturated WSOGs during collection.

Though many organics are stable when stored frozen in water, IEPOX does not survive extended storage in water (confirmed with our organic synthesis collaborator). We expect this to be the case for isoprene hydroxyhydroperoxide

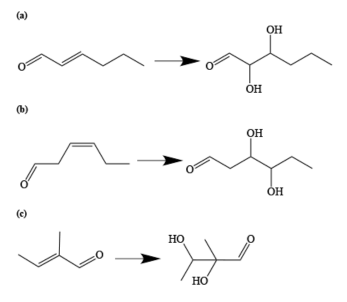


Figure 4. Gas-phase oxidation of (a) *E*-2-hexenal, (b) *Z*-3-hexenal, and (c) (*E*)-2-methyl-2-butenal. The diols are proposed to result from hydrolysis of the corresponding epoxides, which may be generated by attack of OH followed by addition of O_2 , 1,5-H transfer, and elimination of OH (Crounse et al., 2012; Peeters et al., 2001) or by reaction with RO₂. (Stark, 1997).

(ISOPOOH) also. ISOPOOH is an OH oxidation product of isoprene, which is further oxidized by OH under low-NO conditions to form isomeric IEPOX (Paulot et al., 2009). IEPOX and ISOPOOH were present in the gas phase during the SOAS campaign (Nguyen et al., 2015). They have relatively high Henry's law constants (i.e., $H_{L,IEPOX} = 2.7 \times 10^6 \text{ M} \text{ atm}^{-1}$). IEPOX was readily detected in ambient samples spiked with 3000, 300, and 30 µM of IEPOX, indicating that it can be ionized in our sample matrix. (Authentic *trans-* β -IEPOX, which is the predominant isomer of IEPOX, was synthesized for this purpose; Zhang et al., 2012.) However, it was not found in our ambient samples since it is not stable when stored in water.

Together, the ions discussed herein account for 30% of the total ion abundance. The remaining 70% did not exhibit a clearly decreasing trend during OH oxidation experiments. The remaining 70% could represent compounds that are stable with respect to OH oxidation in water or that react too slowly to be observed under the experimental conditions used. However, it should be recognized that some watersoluble OH-reactive compounds might not have been identifiable as reactive because they were intermediates of multiple precursors. We have observed this phenomenon previously in aqueous OH oxidation experiments conducted with mixed aldehyde standards (unpublished). In those experiments, several modest concentration increases and decreases were observed in experiments and model predictions for some intermediates. Together, these methodological limitations suggest that additional OH-reactive water-soluble gases were likely present that we were unable to identify.

3.2 Product formation during aqueous oxidation experiments

Figure 5 shows significant formation of oxalate and pyruvate in OH radical experiments conducted with all samples but not during the control experiments (sample + UV; sample + H₂O₂). Pyruvate peaks around 60–80 min, which is earlier than the oxalate peak at 100–120 min (Fig. 5). Acetate/glycolate (which co-elute in the IC) also forms in at least some samples and reacts away in the presence of OH (Fig. S3). Sulfate and nitrate concentrations remained constant throughout the experiment as measured by the IC. While there may be many sources of oxalate, aqueous OH radical oxidation of pyruvate in the aqueous phase is known to form oxalate at dilute (cloud-relevant) concentrations (Carlton et al., 2006). Aqueous acetate oxidation is also a source of oxalate (Tan et al., 2012). The concentration dynamics are consistent with a role for these compounds in the formation of oxalate in the ambient mixtures although the mechanisms by which pyruvate and acetate formed are not well constrained in these experiments. These observations suggest that oxalate, pyruvate, and acetate/glycolate can form in ambient mixtures of water-soluble gases in the southeastern US in the presence of clouds/fogs and oxidants. Pyruvate and oxalate have been observed primarily in the particle phase in the atmosphere (Saxena and Hildemann, 1996; Limbeck et al., 2001; Yao et al., 2002; Kawamura et al., 2003; Martinelango et al., 2007). Moreover, modeling studies of oxalate, the most abundant dicarboxylic acid in the atmosphere, suggest that aqueous chemistry is a large contributor of oxalate formation globally, making it a good tracer for SOAAO formed in clouds and fogs (Myriokefalitakis et al., 2011). Above- versus below-cloud measurements also support this (Sorooshian et al., 2010). Thus, the experiments suggest that aqueous oxidation of ambient (southeastern US) water-soluble mixtures at cloud/fog-relevant concentrations has the potential to form material that remains in the particlephase species after droplet evaporation, i.e., SOAAO. However, the atmospheric prevalence of particle-phase oxalate can only be explained by the formation of salts and complexes, since oxalic acid is volatile and the volatility of oxalate salts is orders of magnitude lower than that of oxalic acid (Ortiz-Montalvo et al., 2014; Paciga et al., 2014). The aerosol at the SOAS ground site was acidic (campaign average pH \sim 0.94) (Guo et al., 2015), and as a consequence oxalic acid may remain largely in the gas phase in this environment but may eventually react on more basic surfaces, e.g., coarse particles. Note that we expect the products of aqueous chemistry in wet aerosols to be different from those in clouds and fogs because of the extremely high (molar) solute concentrations in wet aerosols (Surratt et al., 2007; Noziere

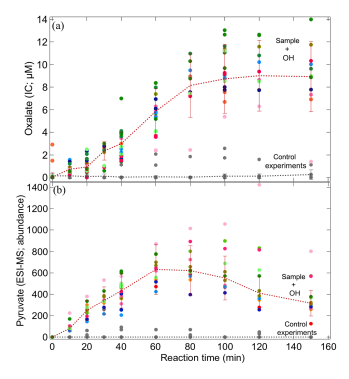


Figure 5. (a) Oxalate (by IC) for all OH radical oxidation experiments conducted with ambient samples (Table 1). (b) Abundance of the negative ion at m/z 87 (pyruvate) as observed in the ESI-MS when the ambient SOAS samples are exposed to OH. Error bars represent the pooled coefficient of variation calculated across experimental days. Note that oxalate and pyruvate are formed in all samples in the presence, but not the absence, of OH. Gray points represent control experiments (11 June sample + UV, 11 June sample + H₂O₂, 30 June field water blank + OH).

et al., 2008; Galloway et al., 2009; Lim et al., 2010; Sareen et al., 2010; Nguyen et al., 2012).

3.3 Atmospheric implications

We have tentatively characterized several water-soluble OHreactive species collected at an isoprene-rich photochemically active location in the southeastern US. In several cases compounds with the same elemental composition were measured in the gas phase by high-resolution time-of-flight chemical ionization mass spectrometry (HRToF-CIMS). The tentative structures for the proposed reactants are consistent with formation from green-leaf volatiles and isoprene oxidation. Aqueous OH oxidation under dilute conditions (TOC of ca. 100 µM) relevant to fogs and clouds produced oxalate and pyruvate, suggesting that cloud/fog processing of these compounds (and subsequent neutralization or complexation) is a potential source of SOA. The reactants characterized in this work are precursors for aqueous chemistry and are potentially important SOAAO precursors in all atmospheric waters, i.e., clouds, fogs, and wet aerosols. The aqueous chemistry of these precursors is poorly understood and warrants further study.

4 Conclusions

We observed formation of pyruvate, oxalate, and acetate/glycolate during OH oxidation experiments conducted with ambient mixtures of WSOG from the southeastern US. The formation of these highly oxygenated organic acids indicates a potential for SOA_{AQ} formation (e.g., upon neutralization with NH₃, metal complexation, or heterogeneous reaction on course dust/salt particles). Given the acidity of SOAS fine particles, we think it is unlikely that oxalate will be found in substantial quantities in the fine aerosol at the SOAS ground site.

We tentatively characterized several water-soluble reactive precursors which undergo aqueous chemistry and SOA_{AQ} formation in wet aerosols, clouds, and fogs at this location. High-resolution mass spectrometric analyses suggest precursors had O / C ranging from 0.125 to 0.80, and some are tentatively gas-phase oxidation products of green-leaf volatiles. No distinct difference was seen in the aqueous oxidation of ambient samples collected across days during the SOAS field campaign. Further work involving organic synthesis, aqueous OH oxidation of authentic standards, and mass spectral analyses with pre-separation are likely to yield further insights into the aqueous chemistry of these compounds in the future.

5 Data availability

Included are sampling and TOC data for all collection days (Table S1); time series of ESI-MS abundances for m/z 125, 129, 143, 173, and 187 (Fig. S1); discussion and graphical comparisons with chemical ionization mass spectrometry (CIMS) (Fig. S2), and concentration dynamics for acetate/glycolate during aqueous OH experiments and control experiments (Fig. S3). The material is available free of charge via the Internet at http://pubs.acs.org.

The Supplement related to this article is available online at doi:10.5194/acp-16-14409-2016-supplement.

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