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Laboratory studies of the aqueous-phase oxidation of polyols: submicron particles vs. bulk aqueous solution

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Abstract. Oxidation in the atmospheric aqueous phase (cloud droplets and deliquesced particles) has received recent attention as a potential pathway for the formation of highly oxidized organic aerosol. Most laboratory studies of aqueous-phase oxidation, however, are carried out in bulk solutions rather than aqueous droplets. Here we describe experiments in which aqueous oxidation of polyols (water-soluble species with chemical formula C$_n$H$_{2n+2}$O$_n$) is carried out within submicron particles in an environmental chamber, allowing for significant gas–particle partitioning of reactants, intermediates, and products. Dark Fenton chemistry is used as a source of hydroxyl radicals, and oxidation is monitored using a high-resolution aerosol mass spectrometer (AMS). Aqueous oxidation is rapid, and results in the formation of particulate oxalate; this is accompanied by substantial loss of carbon to the gas phase, indicating the formation of volatile products. Results are compared to those from analogous oxidation reactions carried out in bulk solution. The bulk-phase chemistry is similar to that in the particles, but with substantially less carbon loss. This is likely due to differences in partitioning of early-generation products, which evaporate out of the aqueous phase under chamber conditions (in which liquid water content is low), but remain in solution for further aqueous processing in the bulk phase. This work suggests that the product distributions from oxidation in aqueous aerosol may be substantially different from those in bulk oxidation experiments. This highlights the need for aqueous oxidation studies to be carried out under atmospherically relevant partitioning conditions, with liquid water contents mimicking those of cloud droplets or aqueous aerosol.

1 Introduction

A large fraction of atmospheric organic aerosol (OA) is known to be secondary in nature, formed by the oxidation and subsequent condensation of gas-phase precursors. However, the sources and formation mechanisms for the most oxidized fraction of OA remain highly uncertain. Ambient measurements indicate that the most highly oxidized OA has an average carbon oxidation state of between 0 and +1 (Daumit et al., 2013; Jimenez et al., 2009; Kroll et al., 2011) and can form rapidly over short timescales (approximately 1 to 3 days; Jimenez et al., 2009; Volkamer et al., 2006). However, the organic aerosol formed in laboratory studies is typically substantially less oxidized than this (Donahue et al., 2012; Heald et al., 2010; Kroll et al., 2011; Qi et al., 2010), and model studies generally have difficulty reproducing the same level of oxidation in such short timescales (Dzepina et al., 2009, 2011; Hodzic et al., 2010; Lee-Taylor et al., 2011).

Aqueous-phase oxidation has received considerable attention as a potential formation pathway for highly oxidized organic aerosol (Ervens et al., 2011; Lim et al., 2010). The oxidation of organic species in the aqueous phase has recently been investigated in the laboratory for a range of water-soluble species, including small carbonyls (Altieri et al., 2008; Carlton et al., 2007; Kirkland et al., 2013; Perri et al., 2009, 2010), isoprene and its oxidation products (Altieri et al., 2006; Kameel et al., 2013; Liu et al., 2012a; Renard et al., 2013; Zhang et al., 2010), and phenolic compounds (Smith et al., 2014; Sun et al., 2010). Such bulk-phase studies have clearly demonstrated that aqueous-phase oxidation, when it occurs, can lead to the rapid formation of highly oxidized organic species.

The importance of such oxidation processes relies critically on partitioning (Donahue et al., 2014): a compound
will undergo aqueous-phase oxidation only if it is primarily partitioned to the atmospheric aqueous phase (cloud droplets or aqueous submicron particles), rather than the gas or condensed-organic phase. The partitioning between the gas and aqueous phases is determined by a compound’s effective Henry’s law constant ($H^*$) (or alternatively its saturation vapor concentration over water ($c^*_{aq}$); Ervens et al., 2011), as well the liquid water content (LWC) of the air mass, as shown in Eq. (1):

$$f_{aq} = \frac{(LWC) H^* RT}{10^{12} + (LWC) H^* RT} = \frac{LWC/c^*_{aq}}{1 + LWC/c^*_{aq}},$$  

in which $f_{aq}$ is the equilibrium fraction in the aqueous phase, LWC is in units of $\mu$g m$^{-3}$, $H^*$ is in units of M atm$^{-1}$, and $c^*_{aq}$ is in units of $\mu$g m$^{-3}$ and is equal to $10^{12} H^{-1} R^{-1} T^{-1}$ (where $R = 0.0821$ L atm mol$^{-1}$ K$^{-1}$ and $T$ is temperature in K). By Eq. (1), the partitioning of a single compound (with a fixed $H^*$ or $c^*_{aq}$) can be very different for different values of LWC. Atmospheric LWCs can span many orders of magnitude, ranging from $\sim 1$ to $100 \mu$g m$^{-3}$ for aqueous aerosol, and $\sim 10^5$ to $10^6 \mu$g m$^{-3}$ for cloud droplets (Ervens et al., 2011), while a bulk aqueous solution has a LWC on the order of $10^{12} \mu$g m$^{-3}$ (the density of water). Therefore, $f_{aq}$ for a given compound will also vary by many orders of magnitude for these systems, as illustrated in Fig. 1. Many compounds that are considered to be “water-soluble” for bulk solutions ($H^*$ of 1–1000 M atm$^{-1}$) will not actually partition significantly into aqueous submicron particles, or even into cloud droplets. This could have an important influence on the resulting chemistry, and suggests that a bulk solution may not always accurately represent aqueous processing under atmospheric conditions.

Some recent bulk-solution experiments have begun to examine the role of gas–particle partitioning under lower-LWC conditions by atomizing or nebulizing the bulk aqueous solution as it undergoes aqueous oxidation (Lee et al., 2011a, b; Liu et al., 2012b; Ortiz-Montalvo et al., 2012; Zhao et al., 2012). While these studies do indeed show loss of the most volatile species formed, leaving behind only low-volatility condensed-phase products, the chemistry that forms these products still takes place in the bulk aqueous solution, with limited partitioning into the gas phase. To our knowledge, only two studies (Nguyen et al., 2013; Volkamer et al., 2009) have examined oxidation chemistry within aqueous droplets themselves, allowing for gas–particle partitioning that mimics what may occur in the atmosphere. These studies found enhanced uptake and aerosol yield from glyoxal (Volkamer et al., 2009) and glycolaldehyde (Nguyen et al., 2013) in the presence of aqueous submicron particles; results from these experiments have not been explicitly compared to those in which oxidation was carried out in bulk solutions.

Here we describe laboratory studies of the oxidation of water-soluble organic species in the aqueous phase, with experiments conducted both within bulk aqueous solutions and within submicron aqueous particles. The goal of these experiments is to compare the oxidation chemistry under very different partitioning conditions. We focus on the oxidation of polyols with the chemical formula $C_nH_{2n+2}O_n$ (with one hydroxyl group on each carbon atom). Polyols with four or more carbon atoms have exceedingly high values of $H^*$ ($>10^{16}$ M atm$^{-1}$, Sander, 1999), ensuring they will be present in the aqueous phase even at the low LWC in our chamber; thus any observed partitioning will involve reaction intermediates and products only. To our knowledge, this work is the first direct comparison of aqueous oxidation in submicron particles and in bulk aqueous solution.

2 Experimental methods

Two sets of experiments are conducted: (1) bulk oxidation, in which reactions take place within a bulk aqueous solution of $\sim 0.5$ L volume, and (2) chamber oxidation, in which reactions take place within submicron aqueous particles in an environmental chamber. Both sets of experiments use dark Fenton chemistry (Fe$^{2+}$ and hydrogen peroxide) as an aqueous-phase source of oxidants. While there is some uncertainty as
to the exact oxidant-forming mechanisms in Fenton systems, in part due to the complex iron speciation, it is likely that hydroxyl radicals (OH) are the predominant oxidant formed in our experiments (Ma et al., 2006; Southworth and Voelker, 2003). Fe\(^{2+}\) reacts with hydrogen peroxide (H\(_2\)O\(_2\)) to produce OH and Fe\(^{3+}\), which is subsequently converted back to Fe\(^{2+}\) via reaction with other species:

\[
\text{Fe}^{2+} + \text{H}_2\text{O}_2 \rightarrow \text{Fe}^{3+} + \text{HO}^- + \text{OH}, \\
\text{Fe}^{3+} + \text{H}_2\text{O}_2 \rightarrow \text{Fe}^{2+} + \text{HO}_2 + \text{H}^+, \\
\text{Fe}^{3+} + \text{HO}_2 \rightarrow \text{Fe}^{2+} + \text{O}_2 + \text{H}^+. 
\]

(R1) (R2) (R3)

Alternatively, Fe\(^{3+}\) can form complexes with water or organic acids; these pathways compete with the regeneration of Fe\(^{2+}\) and can eventually slow the production of OH:

\[
\text{Fe}^{3+} + \text{H}_2\text{O} \rightarrow \text{Fe(OH)}^{2+} + \text{H}^+, \\
\text{Fe}^{3+} + \text{L}^- \rightarrow \text{Fe(L)}^{2+} \quad (\text{L} = \text{organic ligand}). 
\]

(R4) (R5)

A more detailed treatment of this chemistry is described elsewhere (Arakaki and Faust, 1998; Faust and Hoigne, 1990; Hoffmann et al., 1996; Ma et al., 2006; Nguyen et al., 2013; Ou et al., 2008). In both sets of experiments, we start with a mixture of a polyol and Fe\(^{2+}\), and initiate the oxidation chemistry with the addition of H\(_2\)O\(_2\). The use of Fenton chemistry to initiate oxidation within aqueous submicron particles was first demonstrated in a recent glycolaldehyde uptake study by Nguyen et al. (2013); the present experiments differ from that work in that we focus on dark Fenton chemistry, using the addition of H\(_2\)O\(_2\) rather than exposure to UV lights to initiate the reaction.

### 2.1 Bulk aqueous oxidation

For bulk oxidation experiments, the general technique of Lee et al. (2011a) is used, in which an atomizer serves both as the reactor containing the aqueous solution and as the method for aerosolizing the solution, enabling online analysis by aerosol instrumentation. Figure 2a shows a simple schematic of the experimental setup. Reactions are carried out in a 1 L borosilicate reaction bottle, and the solution is continuously atomized using a constant output atomizer (TSI, model 3076) with a backing pressure of 30 psi of pure compressed air (Aadco 737-13A/C with methane reactor), giving 2.5–3 LPM aerosol output flow. The atomizer output is passed through an empty 1 L bottle to remove excess liquid water, then through a diffusion dryer, and finally into the aerosol instruments, described in Sect. 2.3.

A major difference between the present experiments and previous bulk oxidation experiments (Lee et al, 2011a, b, 2012; Liu et al., 2012b; Ortiz-Montalvo et al., 2012; Zhao et al., 2012) is the oxidation scheme: here OH radicals are generated via dark Fenton chemistry rather than photolysis of hydrogen peroxide, avoiding the use of 254 nm lights. The initial atomization solution contains iron(II) sulfate heptahydrate (99 + %) and a polyol (glycerol, 99.5 + %, erythritol, 99 + %, adonitol 99 + %, mannitol 98 + %, or volemitol, all Sigma-Aldrich) fully dissolved in Milli-Q water; concentrations are given in Table 1. Prior to H\(_2\)O\(_2\) addition, this solution is atomized into the aerosol mass spectrometer (AMS) (see below) to characterize the reactants and ensure they are inert with respect to each other. To initiate Fenton chemistry, hydrogen peroxide (30 % w/w in water, Alfa Aesar) is added and the solution is shaken to facilitate mixing. After ~3 h, once the reaction appears to have gone to completion, the solution is exposed to UV light for additional oxidation (via photo-Fenton chemistry) and/or direct photolysis, using four external black lights (Sylvania BL-350 ECO, output 300–400 nm). Photo-Fenton reactions regenerate Fe\(^{2+}\) and can produce additional OH:

\[
\text{Fe}^{3+} (\text{OH}^-) + \text{hv} \rightarrow \text{Fe}^{2+} + \text{OH}, \\
\text{Fe}^{3+} (\text{L}^-) + \text{hv} \rightarrow \text{Fe}^{2+} + \text{L}. 
\]

(R6) (R7)

### 2.2 Chamber oxidation

Chamber experiments are conducted to allow for aqueous oxidation in submicron particles, providing partitioning conditions representative of deliquesced atmospheric aerosol. The chamber oxidation setup is shown in Fig. 2b. The MIT chamber is a 7.5 m\(^3\) Teflon (0.13 mm PFA) bag within a temperature-controlled environmental room. Two banks of 24 black lights (Sylvania BL-350 ECO) on opposite sides of the chamber provide UV irradiation when needed. The chamber is run as a semi-batch reactor with 5 LPM pure air added to balance instrument sample flows. A fraction of this air is sent through a bubbler containing Milli-Q water in order to maintain a relative humidity of between 67 and 79 %, ensuring that all sulfate-containing particles are aqueous. Prior to reaction, seed particles composed of iron(II) sulfate (FeSO\(_4\)) and a polyol are introduced into the chamber via atomization. The liquid water content in the chamber is not directly measured, but is estimated to be in the range of ~20 to ~150 \(\mu\)g m\(^{-3}\) prior to oxidation, based on concentrations of particulate water measured by the AMS. These values are broadly consistent with liquid water contents estimated by assuming the particles have a similar hygroscopic growth factor as particles of ammonium sulfate (Seinfeld and Pandis, 2006). Concentrations of the atomizer solution are given in Table 1; it should be noted that, within the aqueous particles themselves, concentrations will be substantially higher than these values, since the solution is concentrated upon atomization by a factor of ~1000 (based on measured mass loadings and estimated LWC).

Seed particles are allowed to mix in the chamber for 1 h prior to addition of H\(_2\)O\(_2\), which initiates the reaction. H\(_2\)O\(_2\) is not added directly to the atomizer solution, since this would initiate oxidation in the bulk solution prior to atomization, and so is instead introduced in the gas phase, from which it subsequently partitions into the aqueous phase. Gas-phase H\(_2\)O\(_2\) is introduced by sending 1–1.5 LPM air through
a two-neck flask containing 6.0 µL of 30% aqueous H$_2$O$_2$ solution and into the chamber via a Teflon line; full evaporation of the H$_2$O$_2$ takes approximately 30 min. Assuming all the H$_2$O$_2$ enters the chamber and partitions into the aqueous particles according to Henry’s law (with $H^* = 10^5$ M atm$^{-1}$; Sander, 1999), this gives 270 ppb H$_2$O$_2$ in the gas phase, and 27 mM in the aqueous particles. This aqueous concentration is higher than in the bulk experiments, but the concentrations of Fe$^{2+}$ and polyol are also higher in the particles than in the bulk solution. However, because some H$_2$O$_2$ is expected to be lost to surfaces, such as the Teflon inlet tubing and chamber walls, these H$_2$O$_2$ concentrations likely represent upper limits. After completion of the dark Fenton chemistry, the black lights are turned on for additional photolytic and/or oxidative chemistry, as in the bulk experiments. Under dark conditions, gas-phase oxidation is unlikely; however, during irradiation the photolysis of H$_2$O$_2$ will lead to the formation of some gas-phase OH; as discussed below, this is unlikely to affect the observed chemistry.

2.3 Aerosol analysis

Aerosol size distribution, volume, and number density are monitored using a scanning mobility particle sizer (SMPS; TSI Inc.). Chemical composition of non-refractory particulate matter is measured using a high-resolution time-of-flight AMS (Aerodyne Research Inc.). This allows for the measurement of chemical families, such as total organic, sulfate, and ammonium (Canagaratna et al., 2007; DeCarlo et al., 2006), as well as elemental ratios of the organic species, most importantly oxygen-to-carbon (O/C) and hydrogen-to-carbon (H/C) (Aiken et al., 2007, 2008). All data reported here were taken with the instrument operated in V mode. Because the amount of sulfate per particle is constant, abundances of all reported ions and families are normalized to sulfate, accounting for possible changes in collection efficiency (CE), variations in atomizer output, dilution in the chamber or atomizer, and loss of particles to surfaces. Some sulfate signal (10–25%) is measured in the “closed” mass spectra, suggesting that FeSO$_4$ does not flash-vaporize at the temperature of the AMS vaporizer (~550–600 °C). However, this fraction does not vary significantly over any experiment, suggesting the AMS “diff” signal is sufficient for this normalization. In most chamber experiments the aerosol is not sent through a dryer prior to analysis, though drying appears to have no effect on measured elemental ratios or sulfate-normalized concentrations. For chamber oxidation experiments, we also monitor temperature and relative humidity (Vaisala), NO$_x$ (Horiba Inc.), and ozone (2B Tech).

3 Results and discussion

A series of control experiments were carried out to verify that aqueous-phase, dark Fenton chemistry was indeed responsible for any chemistry observed. No reaction was observed between the polyols and FeSO$_4$ in the absence of H$_2$O$_2$, nor between the polyols and H$_2$O$_2$ in the absence of FeSO$_4$, confirming that both Fe$^{2+}$ and H$_2$O$_2$ are necessary for oxidation to occur. Furthermore, no chemical changes were observed
upon addition of H$_2$O$_2$ or exposure to UV when a chamber experiment (using erythritol-FeSO$_4$ particles) was conducted under dry conditions (RH < 4%). The sulfate-normalized organic signal gradually decayed over the course of this experiment, but this is likely a result of gradual evaporation of organic material as the chamber air is diluted. The lack of a reaction under such dry conditions confirms that the chemistry described below indeed takes place only in the aqueous phase.

### 3.1 Bulk oxidation of erythritol

Results for the bulk oxidation of erythritol (C$_4$H$_{10}$O$_4$) are shown in Fig. 3. Figure 3a shows the aerosol mass spectrum of the aqueous erythritol solution, taken over a 30 min period immediately prior to the addition of H$_2$O$_2$. Figure 3b shows the time traces of key AMS ions and species (all normalized to sulfate). Once H$_2$O$_2$ is added, oxidation occurs immediately. Key ions associated with erythritol (m/z 29, 39, 61, 73, 91) begin to decay rapidly, presumably by reaction with OH. Using a $k_{OH}$ of $1.9 \times 10^9$ M$^{-1}$ s$^{-1}$ (Herrmann et al., 2010), this decay is consistent with an initial aqueous OH concentration of $4.8 \times 10^{-12}$ M, which is within an order of magnitude of the average OH concentrations estimated for ambient deliquesced particles (Herrmann et al., 2010). The OH concentration, estimated from the time dependence of the polyol concentration (see Supplement), drops over the course of the experiment, reaching a final OH exposure of $2.7 \times 10^{-9}$ M s. Sulfate-normalized organic signal initially decreases, likely due to the formation and evaporation of compounds that are more volatile than erythritol, but then gradually rises, presumably from the addition of oxygen-containing functional groups that increase the molecular weight and reduce the volatility of the organic species. Ammonium (NH$_4^+$) also rises, likely due to uptake of ammonia from the laboratory air; this is consistent with acidification of the solution, which is known to occur upon initiation of dark Fenton chemistry (Nguyen et al., 2013). This is accompanied by an instantaneous change in the appearance of the atomizer solution from colorless to yellow-orange, consistent with the formation of Fe(OH)$_2^{2+}$ (Zuo and Hoigne, 1992).

Rapid growth of the CO$_2^+$ ion in the AMS is observed upon oxidation (Fig. 3b). This highly oxidized ion fragment, which typically indicates the presence of carboxylic acids, is the most abundant ion in the product mass spectra, shown in Fig. 3c. Figure 3d shows the elemental ratios of the particular organic species as a function of time; the rapid rise in O/C and drop in H/C upon addition of H$_2$O$_2$ is also consistent with rapid oxidation of the organic species. It should be noted that the initial H/C and O/C measured by the AMS do not match the true values for erythritol. This is a result of using the default elemental analysis correction factors, which have been determined empirically for an ensemble of species rather than individual compounds (Aiken et al., 2008); however, errors in the absolute elemental ratios are not expected to affect the observed trends in H/C and O/C, nor the overall conclusions of this work. The rapid oxidation of dissolved organic species observed here is broadly consistent with the findings of other bulk oxidation studies (Altieri et al., 2006, 2008; Carlton et al., 2007; Kirkland et al., 2013; Lee et al., 2011a, b; 2012; Liu et al., 2012a; Perri et al., 2009; Sun et al., 2010; Tan et al., 2009, 2010; Zhang et al., 2010; Zhao et al., 2012).

The AMS mass spectrum of the oxidation products closely resembles that of oxalate (C$_2$O$_4^{2-}$) (Mensah et al., 2011; Takegawa et al., 2007), which has been shown to be a major product in other aqueous-phase oxidation systems (Carlton et al., 2007; Perri et al., 2009; Tan et al., 2010). A pathway...
Figure 3. Results for the oxidation of erythritol within the bulk aqueous phase: (a) AMS spectra of unreacted erythritol; (b) sulfate-normalized mass concentrations of total organic (Org), ammonium (NH₄⁺), CO₂⁺, and key ions associated with erythritol (m/z 29, 39, 61, 73, 91) as a function of reaction time; (c) AMS spectra of oxidation products; and (d) oxygen-to-carbon and hydrogen-to-carbon ratios as a function of reaction time. Results in (a, c, d) are from high-resolution mass spectrometric analysis. All traces in (b) are from unit mass resolution, except NH₄⁺ and CO₂⁺, which are high-resolution traces.

for the formation of oxalic acid from the aqueous oxidation of ethylene glycol (a C2 diol) has been shown by Tilgner and Herrmann (2010) and mechanisms for the formation of oxalic acid from larger diacids have been described by Ervens et al. (2004). We did not observe the formation of organo-nitrogen compounds, which were major reaction products in the uptake of glyoxal (Galloway et al., 2009; Nozière et al., 2009) and glycolaldehyde (Nguyen et al., 2013). In those studies, imidazoles and other N-containing products were formed via nucleophilic attack of ketones and aldehydes by ammonia (Galloway et al., 2009; Nozière et al., 2009); such reactions are unlikely in the present system given the lack of carbonyl moieties in the polyols studied here.

The formation of oxalate (a highly oxidized C2 compound) from erythritol (a less oxidized C4 compound) likely occurs via multiple generations of oxidation. Individual intermediate ions (not shown) have significantly different temporal behavior, with some growing in and decaying before others, suggesting complex chemistry with intermediate products formed at different stages of oxidation. The varying rates of decay of erythritol ions in Fig. 3b (e.g., m/z 61 vs. 91) provide further evidence that multigenerational oxidation is occurring in this system.

Upon exposure to UV, the organic ions decay still further, and the total organic signal decreases dramatically, in a short period of time (lifetime ∼ 36 min; see Fig. S1a, Supplement). This is likely a result of either further oxidation by OH generated from photo-Fenton chemistry (Reaction R6) or direct photolysis (Reaction R7) to form small volatile species that cannot be detected by the AMS. This provides additional evidence for oxalate, since Fe(III) oxalato complexes are known to rapidly photolyze, a process that has been suggested as an important sink of atmospheric oxalate (Sorooshian et al., 2013). Given the abundance of iron in the system, our results suggest that the main condensed-phase oxidation product is oxalate, present as an iron oxalato complex.

3.2 Chamber oxidation of erythritol

Results for the oxidation of erythritol in aqueous particles in the chamber are shown in Fig. 4. As in the bulk experiment, dark Fenton chemistry within submicron particles in the chamber was found to lead to the rapid decay of erythritol and formation of oxidized products. A maximum OH concentration of 3.5 × 10⁻¹² M is reached between 30 and 40 min, and subsequently decreases to give a final OH exposure of 4 × 10⁹ M s; these values are similar (within a factor of 2) to those of the bulk experiments.
Figure 4. Same as Fig. 3, but for the oxidation of erythritol within submicron particles (chamber experiments): (a) AMS spectra of unreacted erythritol; (b) sulfate-normalized mass concentrations of total organic (Org), ammonium (NH$_4^+$), CO$_2^+$, and key ions associated with erythritol (m/z 29, 39, 61, 73, 91) as a function of reaction time; (c) AMS spectra of oxidation products; and (d) oxygen-to-carbon and hydrogen-to-carbon ratios as a function of reaction time.

The initial organic mass spectrum of unreacted erythritol in the chamber (Fig. 4a) closely matches that of erythritol in the bulk (Fig. 3a). The observed chemical changes in the two oxidation systems, as described by the time traces of key AMS ions from the chamber experiment (Fig. 4b), the product mass spectrum (Fig. 4c), and changes to O/C and H/C (Fig. 4d) are also similar to the bulk oxidation results. The primary difference is that the total organic signal decreases substantially more than in the bulk, with no subsequent increase, discussed in detail below. As in the bulk experiment, a rapid loss of organic mass is observed upon UV irradiation (Fig. S1b in the Supplement). This loss is unlikely to arise from oxidation by gas-phase OH (formed from H$_2$O$_2$ photolysis): the rapid loss of the organic species (lifetime $\sim$ 7 min) would require an OH mixing ratio of $>6 \times 10^7$ molec cm$^{-3}$ (assuming a $k_{OH}$ of $<3 \times 10^{-11}$ cm$^3$ molec$^{-1}$ s$^{-1}$), far greater than can be produced by H$_2$O$_2$ photolysis. Instead, the rapid loss of organic mass probably arises from the direct photolysis of Fe(III) oxalato complexes, as in the bulk oxidation experiment.

Our finding of rapid oxidation by Fenton chemistry in the absence of UV light differs from the results of Nguyen et al. (2013), who found oxidation to occur only upon UV irradiation (photo-Fenton chemistry). In that work, the H$_2$O$_2$ was added directly to the atomizer solution (along with FeSO$_4$), and so dark Fenton chemistry may have gone to completion prior to the introduction of the organic species (glycolaldehyde) in the gas phase. By contrast, in the present experiments, H$_2$O$_2$ is added last, so that all OH is produced in the presence of the aerosol-phase organic compound. Because of such differences, and the differences in the organic species studied, results from the two chemical systems are difficult to compare directly. Nonetheless, our results are broadly consistent with those of Nguyen et al. (2013), in that they indicate that aqueous-phase oxidation in submicron particles can lead to the rapid formation of highly oxidized organic aerosol.

3.3 Oxidation of other polyols

Similar results are seen for the oxidation of most of the other polyols, in both the bulk phase and the chamber. The one exception is the oxidation of glycerol (a C3 polyol), which behaved differently in both the bulk and chamber experiments, presumably due to its high volatility. In the glycerol bulk experiments, the measured organic mass actually increased after oxidation, likely because the less volatile oxidation products evaporated less than glycerol upon atomization; in the chamber experiments, the LWC was too low for any measurable particulate glycerol. However, the C5–C7 polyols (adonitol, mannitol, and volemitol) all behaved similarly to
erythritol upon oxidation, with decreases in the intensity of the polyol ions and H/C, increases in the CO\(_2^+\) ion intensity and O/C, and product mass spectra resembling that of oxalate (Figs. S2–S4 in the Supplement). Estimated OH concentrations and exposures were also similar between the bulk and chamber cases (see Fig. S5 in the Supplement). As in the erythritol case, exposure to UV led to a rapid decrease in all organic ion signal intensity, as well as a decrease in total organic mass. In each case, the fraction of organic material remaining after oxidation was smaller for the chamber experiment than for the bulk experiment.

Shown in Fig. 5 is the fraction of carbon remaining in the condensed phase (\(f_C\)) after reaction in the bulk solution or deliquesced particles for each polyol (except glycerol, for which meaningful values of \(f_C\) could not be determined). This is calculated from the changes in organic mass concentration and elemental ratios:

\[
f_C = \frac{[\text{Org}]_i}{[\text{Org}]_f} \left( \frac{16O/C_i + H/C_i + 12}{16O/C_f + H/C_f + 12} \right), \tag{2}\]

in which [Org] is the average high-resolution sulfate-normalized mass concentration of total organic, and O/C and H/C are the oxygen-to-carbon and hydrogen-to-carbon ratios, respectively, of the organic species, with the subscripts \(i\) and \(f\) denoting initial and final conditions (before and after H\(_2\)O\(_2\) addition), respectively. For example, in the case of erythritol oxidation, 51% of the initial carbon remained after bulk oxidation, but only 16% remained after oxidation in aqueous aerosol in the chamber. For all bulk experiments, between 48 and 58% of the original carbon remained in the condensed phase after oxidation, with no obvious dependence on carbon number. However, a clear trend is observed for the chamber experiments, with \(f_C\) increasing with the size of the polyol.

### 3.4 Differences between chamber and bulk experiments

The above observations – that a larger fraction of carbon remains in the condensed phase for the bulk experiments than for the chamber experiments, and that this difference gets smaller with increasing carbon number – can be explained either by differences in reactivity or differences in partitioning between the two systems. The submicron particles in the chamber have higher reactant concentrations and a greater ionic strength than the dilute bulk solution, and also require dissolution of gas-phase H\(_2\)O\(_2\) prior to oxidation; these differences may cause the observed differences in oxidation kinetics. Furthermore, the availability of gas-phase H\(_2\)O\(_2\) in the chamber that can partition into the submicron particles could allow for a longer production of OH. However, the ratio of polyol to Fe\(^{2+}\) is also greater in the chamber, so the OH may react away more quickly. These complex effects are difficult to quantify, but could result in different chemical reactions and degrees of oxidation between the two systems.

![Figure 5](image-url)
Figure 6. Simplified mechanism to explain observed differences between oxidation in the bulk and in submicron particles. The polyol reacts to form intermediates with lower $H^+$ (higher volatility over water), which remain fully in the aqueous phase during bulk experiments, but partition to the gas phase (red arrows) during chamber experiments. This partitioning, which occurs to a greater extent for smaller polyol precursors, competes with further aqueous oxidation, and has the effect of lowering the product (oxalate) yield. Upon exposure to ultraviolet light, the Fe(III) oxalato complex photolyzes, forming small gas-phase products.

The present results confirm those of previous studies showing that aqueous-phase oxidation is an efficient pathway for the rapid formation of highly oxidized material. However, when oxidation occurs within submicron particles, the fraction of carbon remaining in the condensed phase is substantially smaller than in the bulk oxidation experiments, implying that the formation of highly oxidized OA by aqueous chemistry may be somewhat less important than bulk-phase experiments suggest. Bulk oxidation experiments may not accurately simulate the chemistry that takes place in the atmospheric aqueous phase, due to large differences in LWC (and therefore partitioning) between the bulk and atmospheric droplets or particles. This difference points to the importance of running aqueous-phase oxidation experiments under atmospherically relevant partitioning conditions. This is analogous to chamber studies of secondary organic aerosol (SOA) formation via partitioning into the condensed-organic phase, which to be representative of atmospheric conditions must be run at low total organic mass concentrations ($c_{OA}$) (Presto and Donahue, 2006). A major difference between such “traditional” SOA chamber experiments and aqueous-phase oxidation, however, is that LWC can vary by a great deal more than $c_{OA}$ during atmospheric processing, since they do not include the effects of partitioning of reaction intermediates. Further, the experiments with glycerol oxidation demonstrate that in bulk-phase experiments it is possible to oxidize compounds that would not actually be present in the atmospheric aqueous phase under most conditions. Results from the oxidation of larger polyols indicate that differences in LWC between bulk and chamber oxidation result in significant differences in partitioning, which in turn can affect the chemistry. For experiments like ours with a single product, this affects only the final yield. However, when multiple oxidation products are formed, with intermediates of varying volatility, changing the LWC may also alter the product distributions.

This study underscores the need for conducting oxidation experiments at atmospherically relevant liquid water contents, and extending the existing suite of bulk experiments...
to additional partitioning conditions. Oxidizing previously studied compounds (small carbonyls, acids, isoprene oxidation products, etc.) within aqueous submicron particles would provide valuable information on the role of partitioning and LWC on the formation of highly oxidized OA for a wider range of water-soluble organic compounds. Further, the large differences in partitioning between deliquesced particles and clouds (Fig. 1) suggest that a similar set of experiments is a useful next step for the accurate study of atmospheric cloud processing. Oxidation experiments involving actual cloud droplets, with LWCs much higher than those of aqueous particles but still far lower than that of the bulk aqueous phase, would improve our understanding of this potentially important source of oxidized organic aerosol.

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