Effects of Precursor Concentration and Acidic Sulfate in Aqueous Glyoxal—OH Radical Oxidation and Implications for Secondary Organic Aerosol

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Previous experiments demonstrated that aqueous OH radical oxidation of glyoxal yields low-volatility compounds. When this chemistry takes place in clouds and fogs, followed by droplet evaporation (or if it occurs in aerosol water), the products are expected to remain partially in the particle phase, forming secondary organic aerosol (SOA). Acidic sulfate exists ubiquitously in atmospheric water and has been shown to enhance SOA formation through aerosol phase reactions. In this work, we investigate how starting concentrations of glyoxal (30–3000 µM) and the presence of acidic sulfate (0–840 µM) affect product formation in the aqueous reaction between glyoxal and OH radical. The oxalic acid yield decreased with increasing precursor concentrations, and the presence of sulfuric acid did not alter oxalic acid concentrations significantly. A dilute aqueous chemistry model successfully reproduced oxalic acid concentrations, when the experiment was performed at cloud-relevant concentrations (glyoxal <300 µM), but predictions deviated from measurements at increasing concentrations. Results elucidate similarities and differences in aqueous glyoxal chemistry in clouds and in wet aerosols. They validate for the first time the accuracy of model predictions at cloud-relevant concentrations. These results suggest that cloud processing of glyoxal could be an important source of SOA.

Introduction

Organic aerosols affect visibility, health, and global climate (1, 2). Current models underestimate organic aerosol concentrations in the free troposphere, suggesting there is a missing secondary organic aerosol (SOA) formation mechanism (3, 4). There is ample evidence suggesting that fog and cloud processing contributes to global SOA budget (5–8); however, our knowledge about the overall importance of fog and cloud processing to SOA formation is quite limited.

Fog and cloud processing, which is the dominant source of atmospheric sulfate, has been hypothesized to be a substantial source of SOA globally (9, 10). Briefly, reactive organic precursors are oxidized in the gas phase to form water-soluble products. These products readily partition into cloud droplets and react further with aqueous oxidants to form low-volatility compounds. Some reactions occur only in the aqueous phase (e.g., Russell mechanism of peroxy radicals), and these reactions may lead to products not seen in gas phase chemistry. Upon cloud droplet evaporation, these low-volatility organics remain at least in part in the particle phase (e.g., 90% for oxalic acid), forming SOA (11). SOA could form through similar aqueous reactions in aerosol water as well.

Glyoxal is a common α-dicarbonyl formed in the atmospheric oxidation of biogenic and anthropogenic precursors, with a global source of 45 Tg a−1 (12). UV photolysis and reaction with the hydroxyl radical (OH) are primary gas phase loss processes for glyoxal (13). Glyoxal concentrations in the Mexico City atmosphere are found significantly below model predictions, suggesting existence of missing sinks (14). Recent field uptake of glyoxal by clouds and wet aerosols could plausibly explain this observation (12). Glyoxal can readily enter droplets because of a high effective Henry’s law constant (Hen > 3 × 10^15 M atm^−1 at 25 °C) and fast uptake rate (15, 16). Typical aqueous glyoxal concentrations vary from a few µM in rainwater to 276 µM in fogwater (17). Concentrations in wet aerosols could be several orders of magnitude higher. Aqueous oxidation of glyoxal forms low-volatility compounds, including oxalic acid and larger multifunctional products (18). In fact, aqueous formation from glyoxal and other carbonyl compounds helps explain the atmospheric abundance of oxalic acid and presence of oligomers in aerosols and clouds. Recent modeling suggests SOA through the aqueous reaction pathway is comparable in magnitude to other SOA formation pathways, though uncertainties are large (12, 19).

Sulfur(VI) contributes to the acidity of cloud droplets. Several studies suggest that sulfuric acid may participate in aerosol phase oligomerization reactions, including aldol condensation and hemiacetal—acetal formation (20, 21). However, the effect of acidic sulfate on SOA production through cloud processing has not been examined.

The kinetics of aqueous glyoxal OH radical oxidation have been studied by Carlton et al., but glyoxal concentrations in those experiments were 1–3 orders of magnitude higher than typical cloud and fog conditions (18). In this work, we investigate how the presence of acidic sulfate and starting concentrations of glyoxal affect product formation in bulk aqueous glyoxal OH radical experiments conducted at cloud-relevant pH. Some organic acids not previously identified in glyoxal oxidation were observed in these experiments as a result of improved analytical resolution. Reaction vessel kinetic modeling captured well product formation at cloud-relevant concentrations. Further chemical mechanism development is needed to improve SOA predictions from glyoxal in wet aerosols.

Experimental Section

Batch Reactions. Batch aqueous reactions of glyoxal and OH radical with and without sulfuric acid were conducted in a 1 L glass reaction vessel as described in detail elsewhere (22). The effect of two factors, glyoxal concentration and sulfuric acid concentration, on the production of low volatility compounds was studied. Experimental conditions are provided in Table S1 of the Supporting Information. Initial glyoxal concentrations were 30, 300, and 3000 µM, and H2SO4 concentrations were 0, 280, and 840 µM. The hydroxyl radical
(3 \times 10^{-12} \text{ M to } 6 \times 10^{-12} \text{ M, estimated}) was formed continuously by photolysis of H_2O_2, using a monochromatic (254 nm) mercury lamp (Heraeus Noblelight, Inc., Duluth, GA). Thus the OH radical concentration remained relatively constant, while the concentration of glyoxal decreased to 0 during experiments. All experiments were conducted at 25 ± 2°C in duplicate. The pH varied from 5.2 to 2.1, decreasing with increasing H_2SO_4 and with increasing reaction time. Samples (10 mL) were taken at similar time points in each experiment with 10% duplicates. A 20 µL aliquot of a 1% catalase solution was added to each sample immediately to destroy H_2O_2. Batch reaction samples were analyzed within 12 h of collection by ion chromatography. Selected samples were also analyzed by mass spectral methods. One glyoxal experiment was performed with real-time mass spectral analysis as described in analytical methods.

The following control experiments were also conducted: glyoxal + UV, glyoxal + H_2SO_4, glyoxal + H_2O_2 ± H_2SO_4, H_2O_2 ± H_2SO_4, and mixed standard + UV. The mixed standard contained oxalic, glyoxylic, malonic, succinic, formic, and glycolic acids. Pure water was also sampled from the reaction vessel and analyzed like samples.

The 30 µM glyoxal experiments are cloud relevant, and 300 µM experiments could represent some heavily polluted fogs. Glyoxal concentrations could reach extremely high values (~1–10 M) during cloud evaporation and in aerosol water. Examination of the chemistry with increasing glyoxal concentration provides some insights into differences between the aqueous photochemistry of glyoxal in cloud and aerosol water.

**Online Experiments.** ESI-MS online experiment (1000 µM glyoxal + 5 mM H_2O_2 + UV) was run as described by Perri et al. (23). The isocratic pump continuously delivered reaction solution from reaction vessel into ESI-MS at 0.11 mL/min, and the binary pump delivered mobile phase at 0.11 mL/min. Samples were analyzed in the negative ionization mode. Discrete samples were frozen for IC analysis.

**Analytical Methods.** Carboxylic acids were quantified by ion chromatography (IC) (ICS-3000, Dionex, Sunnyvale, CA) with an IonPac AS11-HC column (30 µm), AG11-HC guard column (Dionex, Sunnyvale, CA), and conductivity detector (35°C). A photodiode array detector provided additional product validation. Monovalent anions such as glycolate (5.9 min), formate (6.8 min), and glyoxylate (9.7 min) are only weakly retained. Bivalent ions such as succinate (20.4 min), malonate (21.5 min), and oxalate (24.6 min) elute as monovalent ions. Trivalent ions such as citrate are strongly retained and elute even later. Acetate and glycolate (5.9 min) and succinate and malate (20.4 min) as well as malonate and tartrate (21.5 min) coelute.

Fresh samples from batch experiments were analyzed by electrospray ionization mass spectrometry (ESI-MS) (HP-Agilent 1100) as described previously (24). ESI-MS is a soft ionization method that does not fragment ions. Carboxylic acids are detected in the negative ionization mode as molecular weight minus one ion because of the loss of an acidic proton. Aldehydes and alcohols are detected in the positive mode. Glyoxal is detected as m/z 117 and 131 as previously reported. The ion m/z 117 was used to qualitatively represent glyoxal in this work. A frozen sample (~20 °C) taken 30 min into the experiment (3000 µM glyoxal + OH radical; experiment 13, Table S1 of the Supporting Information) was analyzed by Fourier transform ion cyclotron resonance (FT-ICR) ESI-MS (Thermo-Finnigan LTQ-XL, Woods Hole Oceanographic Institute Mass Spectrometer Facility) to determine the elemental formulas of products from 95–500 amu (mass resolution 100–750 k) as described by Perri et al. (23).

Selected samples were analyzed for total organic carbon, and hydrogen peroxide was measured in organic control experiments (H_2O_2 ± H_2SO_4 + UV). Analytical details are provided in the Supporting Information.

**Kinetic Modeling.** Aqueous glyoxal photooxidation was modeled using a mechanism based on Lim et al. (25) (Table 1) and the differential solver, FACSIMILE (AEA Technology, Oxfordshire, U.K.). The H_2O_2 photolysis rate (k = 1.1 \times 10^{-4} s^{-1}, reaction 1, Table 1) was fitted by simulating the H_2O_2 concentration in H_2O_2 + UV control experiments with reactions 1–5 (Table 1; Figure S1 of the Supporting Information). Modeled OH radical concentrations during the experiments were on average 3 \times 10^{-12} – 6 \times 10^{-12} M (Table S1 of the Supporting Information).

**Quality Assurance/Quality Control (QA/QC).** The expected products of glyoxal + OH radical are glyoxylic and oxalic acids. Neither glyoxal nor oxalic acid degraded in the presence of H_2O_2 alone (mixed standard + H_2O_2 control experiment). However, glyoxylic acid + H_2O_2 formed formic acid. To evaluate the effectiveness of catalase in destroying H_2O_2 in samples, we treated some samples from the mixed standard (pH 3.2) + H_2O_2 control experiment (10 mL) with 20 µL of a 1% catalase solution. Some H_2O_2 was still present in these samples after 5 min of contact with catalase. We believe the sample acidity inhibited catalase performance. The reaction of glyoxylic acid + H_2O_2 is slow compared to glyoxylic acid + OH radical (Table 1) in the reaction vessel. However, in samples OH radicals are instantly depleted. The destruction of H_2O_2 by catalase in these samples was not fast enough to prevent conversion of glyoxylic acid to formic acid prior to sample analysis in batch experiments. This explains the low glyoxylic acid concentrations measured by Carlson et al. (18) and suggests that formic and glyoxylic acid concentrations measured in batch experiments described here do not accurately reflect concentrations in the reaction vessel.

Data quality for organic acids is presented in detail in Table S2 and the text of the Supporting Information. Recoveries are near 100%, except for glyoxylic acid (86.5%). With the addition of H_2O_2, recoveries were unchanged for glycolic acid, malonic acid, succinic acid, and oxalic acid. However, glyoxylic acid disappeared, and formic acid concentration increased. Precision is better than 5% for all quantified organic acids.

**Results and Discussion**

**Online Results.** ESI-MS online analysis (Figure 1) demonstrates qualitative agreement with predictions (25, 26) and batch aqueous glyoxal + OH radical experiments conducted previously (18). Specifically, glyoxylic acid (m/z 73) is the first generation product. Oxalic acid (m/z 89) increased rapidly as glyoxylic acid decayed. The time profile of oxalic acid, quantified by IC, matches the abundance of m/z 89 (ESI-MS) very well. In agreement with previous batch experiments (18), a large number of additional ions were observed, including m/z 103, 117, 133, and 149, which will be discussed later. In previous batch experiments, the rapid appearance of formic acid (measured by IC) was taken to suggest that glyoxal + OH radical forms formic acid directly. As discussed in the previous section, we now know that glyoxylic acid reacts with residual H_2O_2 to form formic acid in collected samples from batch experiments. Because the ESI-MS cannot measure formic acid, this online experiment cannot be used to verify or refute the possibility that formic acid is formed directly from glyoxal.

**Effect of Sulfuric Acid Addition.** The addition of sulfuric acid had little effect on the oxalic acid production (Figure 2). Sulfuric acid appears to enhance oxalic acid decay slightly in the late stage of 30 µM experiments, and a slight suppression of oxalic acid production might occur at the
### Aqueous Reactions and Rate Constants in Glyoxal + OH Radical Model

| Reaction                                                                 | Rate Constant (M⁻¹ s⁻¹) | Footnote |
|--------------------------------------------------------------------------|--------------------------|----------|
| 1 H₂O₂ + hv → 2OH                                                       | 1.1 × 10⁻⁴              | b        |
| 2 OH + H₂O₂ → HO₂ + H₂O                                                | 2.7 × 10⁷               |          |
| 3 HO₂ + H₂O₂ → OH + H₂O + O₂                                            | 3.7                     |          |
| 4 HO₂ + HO₂ → H₂O₂ + O₂                                                | 8.3 × 10⁵               |          |
| 5 OH + HO₂ → H₂O + O₂                                                  | 7.1 × 10⁹               |          |
| 6 GLY + OH(→O₂) → GLYAC + HO₂                                           | 1.1 × 10⁹               |          |
| 7 GLYAC + OH → OXLAC + HO₂ + H₂O                                        | 3.62 × 10⁸              |          |
| 8 GLYAC⁻ + OH → OXLAC⁻ + HO₂ + H₂O                                     | 2.9 × 10³               |          |
| 9 OXLAC + OH → 2CO₂ + 2H₂O                                             | 1.4 × 10⁵               | c        |
| 10 OXLAC⁻ + OH → CO₂ + CO₂⁻ + 2H₂O                                    | 4.7 × 10⁷               |          |
| 11 OXLAC²⁻ + OH → CO₂+CO₂⁻ + OH⁻                                      | 7.7 × 10⁶               |          |
| 12 H₂O → H⁺ + OH⁻                                                      | 1.0 × 10⁻⁰              |          |
| 13 HO₂ → H⁺ + O₂⁻                                                      | 1.6 × 10⁻⁵              |          |
| 14 GLYAC ↔ H⁺ + GLYAC⁻                                                 | 3.47 × 10⁻⁴             |          |
| 15 OXLAC ↔ H⁺ + OXLAC⁻                                                 | 5.67 × 10⁻²             |          |
| 16 OXLAC⁻ ↔ H⁺ + OXLAC²⁻                                               | 5.42 × 10⁻⁵             |          |
| 17 CO₂⁻ + O₂ → O₂⁻ + CO₂                                               | 2.4 × 10⁹               |          |
| 18 GLYAC + H₂O₂ → HCO₂H + CO₂ + H₂O                                    | 1.0 × 10⁸               |          |
| 19 HCO₂H + OH → CO₂ + HO₂ + H₂O                                        | 2.4 × 10⁹               |          |
| 20 HCO₂⁻ + OH → CO₂⁻ + H₂O                                            | 1.77 × 10⁻⁴             |          |
| 21 HCO₂H ↔ H⁺ + HCO₂⁻                                                  | 1 × 10⁻¹⁰              |          |
| 22 OH + O₂⁻ → OH⁻ + O₂                                                 | 1 × 10⁻⁷               |          |
| 23 HCO₃ + OH → CO₃⁻ + H₂O                                             | 6.5 × 10⁻⁸              |          |
| 24 CO₃⁻ + O₂ → CO₃⁻ + O₂                                               | 1.5 × 10⁵               |          |
| 25 CO₃⁻ + HCO₂⁻ → HCO₃⁻ + CO₂⁻                                        | 8 × 10⁵                 |          |
| 26 CO₃⁻ + H₂O₂ → HCO₃⁻ + HO₂                                           | 4.3 × 10⁻⁷             |          |
| 27 CO₃⁺(H₂O) ↔ H⁺ + HCO₃⁻                                              | 5.6 × 10⁴               |          |
| 28 HCO₃⁻ ↔ H⁺ + CO₃⁻                                                   | 4.69 × 10⁻¹¹           |          |

*Reactions are taken from Lim et al. (29) and references therein except where footnoted. GLY = glyoxal, GLYAC = glyoxylic acid, OXLAC = oxalic acid, OH = OH radical. Dissociation rate constants (kₐ, M⁻¹ s⁻¹) are calculated from the equilibrium constant (Kₑq, M) and association rate constants (kₐ, M⁻¹ s⁻¹) by kₐ = Kₑq × kₛ. The reaction between oxalic acid and dissolved oxygen was removed from the initial mechanism as no reaction was observed in control experiments. The reaction was measured by Leitzke et al. (34) and was not included by Lim et al.*

### Effect of Precursor Concentration

Increasing precursor concentrations resulted in a nonlinear decrease in the mass of oxalic acid (at maximum) per mass of glyoxal reacted, from 136% in 3000 μM experiments to 94% and 38% in 300 and 30 μM experiments, respectively (Figure 2, Table S1 of the Supporting Information). Also, the oxalic acid production was slower in 3000 μM experiments compared to lower concentration experiments. Most but not all of the decrease in the yield of oxalic acid with increasing precursor concentration is captured by the model results and reflects the fact that H₂O₂ concentrations are also much higher in the higher concentration experiments, whereas OH radical concentrations are only slightly larger. At the higher H₂O₂ concentrations, the formation of formic acid from glyoxylic acid and H₂O₂ competes with the formation of oxalic acid from glyoxylic acid and OH radical in the reaction vessel, reducing oxalic acid production. Oxalic acid did not form in the beginning of 3000 μM experiments. Formation of organo-sulfur compounds is possible but is beyond the scope of this work.
control experiments (glyoxal + UV, glyoxal + H₂O₂). In glyoxal + OH radical experiments, glyoxal was completely consumed after 40 min as indicated by ESI-MS positive mode analysis (Supporting Information), and oxalic acid increased to a maximum concentration 40–70 min into the reaction. Peak oxalic acid concentrations were higher than reported previously (18). We believe that is because large differences in the void volume between samples and standards in the previous analysis (conducted by HPLC-UV−vis, not IC) altered the baseline near the oxalic acid peak.

The dilute aqueous chemistry model (Table 1) successfully reproduced oxalic acid time profiles in 30 µM experiments conducted without H₂SO₄ (Figure 2). Total organic carbon (TOC) analysis (Figure 3) shows that the model is capable of predicting the organic carbon content of the reaction vessel in 30 and 300 µM experiments. This suggests that oxalic acid production from glyoxal can be predicted at cloud-relevant concentrations without considering the formation of higher molecular weight products.

At higher concentrations (3000 µM), measured oxalic acid concentrations were lower and peaked later than the model predicted, and expected organic carbon accounted for less than 60% of the measured TOC in 40 and 60 min samples. A similar observation has been made for glycoaldehyde and OH radical (23). The ESI-MS negative mode spectrum of the 3000 µM glyoxal + OH radical products (20 min) is more complex and contains more higher molecular weight compounds than the spectrum obtained 20 min into the 30 µM experiment (Figure 4). In the mass range from 50 to 500 amu, m/z 188, 271, and 279 were the median ions present in the 30, 300, and 3000 µM experiments, respectively. A total of 76, 161, and 204 ions were detected, respectively, in...
the 30, 300, and 3000 \(\mu\)M experiments. The negative mode spectrum of a mixed standard that included all expected products and precursors was quite simple, suggesting that the complexity seen in samples was not an artifact of the electrospray ionization process (Figure 4 insert; glyoxal, H\(_2\)O\(_2\), glyoxylic acid, oxalic acid, formic acid, and glycolic acid, each at 200 \(\mu\)M; these products are \(20-300 \mu\)M in 3000 \(\mu\)M glyoxal experiments at 20 min). This complexity was not seen in control experiments, indicating that the higher molecular weight products are only formed in the presence of the OH radical. The larger number and complexity of higher molecular weight products with increasing precursor concentration could explain the gap between measured TOC and carbon in predicted products and is consistent with the possibility that formation of higher molecular weight products or oligomers plays an increasingly important role as concentrations increase. The formation of higher molecular weight products could explain the lower measured oxalic acid concentrations and slower production rate relative to predictions in the 3000 \(\mu\)M experiments. The trends exhibited in Figures 2–4 suggest that it will be necessary to account for the formation of higher molecular weight products in order to accurately predict oxalic acid formation (and aqueous-phase SOA formation) from glyoxal + OH radical in aerosol water, where glyoxal concentrations can be 3 orders of magnitude greater than those in our highest experiments.

Additional Carboxylic Acids and Oligomer Formation. IC chromatograms of 30 and 300 \(\mu\)M experiments are relatively simple, only showing organic acids predicted by the explicit dilute aqueous chemistry model. In contrast, several additional peaks not found in control experiments showed clear growth and decay in IC analysis of 3000 \(\mu\)M experiments (Figure S3 of the Supporting Information). Specifically, small peaks with the same retention time as malonic plus tartaric acid and succinic plus malic acid standards were observed. A peak with the retention time expected for mesoxalic acid was also found. Several small peaks that eluted after 30 min are consistent with the presence of tricarboxylic acids on the basis of the separation mechanism of the column. Additionally, the presence of compounds with the exact elemental formulas as succinic acid (\(m/z\) 117.0193, C\(_4\)H\(_5\)O\(_4\)), malonic acid (\(m/z\) 103.0036, C\(_4\)H\(_4\)O\(_4\)), malic acid (\(m/z\) 133.0142, C\(_4\)H\(_5\)O\(_5\)), tartaric acid (149.0097, C\(_4\)H\(_6\)O\(_6\)), and mesoxalic acid (\(m/z\) 116.98294, C\(_3\)H\(_3\)O\(_5\)) in the FT-ICR mass spectra provides strong support for the formation of these compounds in 3000 \(\mu\)M experiments. These ions were not found in analyses of mixed standards or in control experiments. Figure 5 shows time profiles for malonic plus tartaric acids (quantified as malonic acid) and succinic plus malic acid (quantified as succinic acid). The relatively high concentrations of these compounds in 3000 \(\mu\)M experiments suggest that compounds with more than two carbon atoms could be important products in glyoxal oxidation at the considerably higher concentrations observed in aerosol water. The carbon balance for 3000 \(\mu\)M experiments is improved by including malonic acid and succinic acid, but the carbon in the sum of quantified species...
Similar to methylglyoxal and glycolaldehyde, repeated addition of a subunit (s1) with the elemental formula C₃H₄O₂ results in differences of 12, 14, and 16 amu, respectively (Figure 4). 

+OH radical samples exhibit a “haystack” pattern with mass and molecular weight 72.0211 to organic acids was found oxidation of methylglyoxal and glycoaldehyde (23, 27). As previously reported for methylglyoxal (24), pyruvic acid (22), and glycolaldehyde (23), the mass spectra of 3000 μM glyoxal + OH radical samples exhibit a “haystack” pattern with mass differences of 12, 14, and 16 amu, respectively (Figure 4). Similar to methylglyoxal and glycolaldehyde, repeated addition of a subunit (s₁) with the elemental formula C₃H₄O₂ and molecular weight 72.0211 to organic acids was found (Table 2). These products did not form in the absence of OH radicals. The formation of oligoesters by repeated addition of s₁ was verified by MS-MS in methylglyoxal experiments (27). Oligoesters might form through esterification (condensation) reactions or radical—radical reactions, suggesting that s₁ could be a radical or a compound with alcohol and acid functionalities. Unlike glyoxal oligomers formed through hemiacetal formation, we expect that oligomers from OH radical reactions will be irreversibly formed. In addition, in this work, the repeated addition of subunits with molecular formulas C₉H₁₅O₄ (s₂) and C₈H₁₃O₄ (s₃) molecular weights 116.0110 and 132.0058, respectively) to parent compounds could also be seen in the FT-ICR-MS analysis of 3000 μM glyoxal + OH radical samples (Tables 2,3). For example, oxalic acid (C₂H₂O₄) was found to add 1–4 different or same subunits, forming C₇H₁₄O₄–C₉H₁₅O₄, C₇H₁₄O₄–(C₃H₄O₂)₂.

### TABLE 2. Oligomer Series Found by FT-ICR-MS Negative Ionization Mode in a Sample Taken at 30 min Reaction Time (Experiment 13)*

| parent acid | succinic acid | m/z | malic acid | m/z | tartaric acid | m/z |
|-------------|---------------|-----|-----------|-----|--------------|-----|
|             | chemical formula |     | chemical formula |     | chemical formula |     |
| s₁          | C₃H₄O₂        | 117.01929 | C₈H₁₅O₄ | 133.01422 | C₆H₁₂O₁₀ | 149.00907 |
| s₂          | C₄H₆O₄        | 189.04051 | C₈H₁₈O₄ | 205.0254   | C₆H₁₂O₁₀ | 221.03029 |
| s₃          | C₄H₆O₄        | 233.0303  | C₉H₁₅O₄ | 249.0252   | C₆H₁₂O₁₀ | 265.02011 |
|             | C₆H₁₀O₃       | 249.0252 | C₉H₁₅O₄ | 265.02011 | C₆H₁₂O₁₁ | 281.01495 |
| 2s₁         | C₁₁H₁₈O₁₀     | 305.05123 | C₁₀H₁₇O₉ | 277.05649 | C₁₀H₁₇O₁₀ | 293.05128 |
| s₁ + s₂     | C₁₁H₁₇O₁₁     | 321.04690 | C₁₁H₁₇O₁₀ | 321.04609 | C₁₁H₁₇O₁₁ | 337.04124 |
| s₁ + s₃     | C₁₁H₁₇O₁₁     | 321.04690 | C₁₁H₁₇O₁₀ | 321.04609 | C₁₁H₁₇O₁₁ | 337.04124 |
| s₂ + s₁     | C₁₁H₁₇O₁₂     | 349.0412  | C₁₁H₁₇O₁₁ | 349.07748 | C₁₁H₁₇O₁₂ | 365.03687 |
| 2s₂         | C₁₁H₁₇O₁₃     | 365.03604 | C₁₁H₁₇O₁₁ | 365.03604 | C₁₁H₁₇O₁₂ | 381.03092 |
| s₂ + s₃     | C₁₁H₁₇O₁₂     | 365.03604 | C₁₁H₁₇O₁₁ | 365.03604 | C₁₁H₁₇O₁₂ | 397.02577 |
| s₃ + s₂     | C₁₁H₁₇O₁₂     | 377.0724  | C₁₁H₁₇O₁₁ | 377.0724  | C₁₁H₁₇O₁₂ | 397.02577 |
| 3s₂         | C₁₂H₁₉O₁₃     | 421.0624  | C₁₂H₁₇O₁₁ | 421.0624  | C₁₂H₁₇O₁₂ | 409.06211 |
| s₁ + 2s₁    | C₁₂H₁₉O₁₄     | 421.0624  | C₁₂H₁₇O₁₁ | 421.0624  | C₁₂H₁₇O₁₂ | 409.06211 |
| s₁ + s₁ + s₂| C₁₁H₁₇O₁₃     | 437.05727 | C₁₂H₁₇O₁₁ | 437.05727 | C₁₂H₁₇O₁₂ | 425.05733 |

* Products formed from parent acids are categorized by addition of subunits: s₁ denotes C₃H₄O₂, s₂ denotes C₄H₄O₄, and s₃ denotes C₄H₅O₅. All compounds are shown as an anion with one negative charge via losing a proton during the ionization process. (Around 45% of the ion abundance in this sample is accounted for by products in this table.).
C$_2$H$_4$O$_3$–(C$_6$H$_4$O$_2$)$_n$ etc. It should be noted that oligomeric products from these two subunits account for only a portion of the FT-ICR-MS signal. About 45% of the ion abundance in the 30 min sample is accounted for by the products listed in Tables 2 and 3. The higher molecular weight carboxylic acids and oligomers identified by IC and/or FT-ICR-MS could partially account for the missing carbon in the 3000 µM experiment. This work suggests that product complexity and oligomer formation becomes increasingly important as precursor concentrations increase from those typically seen in clouds to those typically seen in aerosol water.

This work provides insights pertaining to the aqueous OH radical oxidation of glyoxal at cloud- and aerosol-relevant concentrations. At cloud-relevant concentrations the Lim et al. (25) dilute aqueous chemistry model successfully predicts oxalic acid formation and total carbon. As precursor concentrations increase, SOA prediction is complicated by the increasing formation of higher molecular weight products; peak oxalic acid concentrations occur later in the reaction sequence, and higher molecular weight products, including >C2 organic acids and oligomers form. These products are also expected to contribute to SOA. In wet aerosols, where glyoxal also reacts with species other than the OH radical, glyoxal chemistry could be even more complex (26–31). Thus, more research is needed to build a chemical model capable of accurately predicting SOA formation from glyoxal in wet aerosols. This work suggests acidic sulfate has only a small effect on oxalic acid production at cloud- and fog-relevant aerosols. This work suggests acidic sulfate has only a small effect on oxalic acid production at cloud- and fog-relevant aerosols. This work suggests acidic sulfate has only a small effect on oxalic acid production at cloud- and fog-relevant aerosols. This work suggests acidic sulfate has only a small effect on oxalic acid production at cloud- and fog-relevant aerosols. This work suggests acidic sulfate has only a small effect on oxalic acid production at cloud- and fog-relevant aerosols. This work suggests acidic sulfate has only a small effect on oxalic acid production at cloud- and fog-relevant aerosols. This work suggests acidic sulfate has only a small effect on oxalic acid production at cloud- and fog-relevant aerosols.

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Supporting Information Available

Analytical details, QA/QC, measurement of H$_2$O$_2$ photolysis (Figure S1), decay of glyoxal (Figure S2), and IC chromatogram of a sample (Figure S3). This material is available free of charge via the Internet at http://pubs.acs.org.

Literature Cited

(1) Kanakidou, M.; Seinfeld, J. H.; Pandis, S. N.; Barnes, I.; Dentener, F. J.; Facchini, M. C.; Van Dingenen, R.; Ervens, B.; Nenes, A.; Nielsen, C. I.; Swietlicki, E.; Putaud, J. P.; Balkanski, Y.; Fuzzi, S.; Horst, J.; Moortgat, G. K.; Winterhalter, R.; Myhre, C. E. L.; Tsigardis, K.; Vignati, E.; Stephanou, E. G.; Wilson, J. Organic aerosol and global climate modelling: A review. Atmos. Chem. Phys. 2005, 5, 1053–1123.

(2) Poschl, U. Atmospheric aerosols: Composition, transformation, climate, and health effects. Angew. Chem., Int. Ed. 2005, 44 (46), 7520–7540.

(3) Volkamer, R.; Jimenez, J. L.; San Martini, F.; Dreppa, K.; Zhang, Q.; Salcedo, D.; Molina, M. J.; Warneke, C.; Holloway, J. S.; Neuman, J. A.; Flocke, F. M.; Seinfeld, J. H.; Concentrations and sources of organic carbon aerosols in the free troposphere over North America. J. Geophys. Res. 2006, 111, D23S47, DOI:10.1029/2006JD0067705.

(4) Heald, C. L.; Jacob, D. J.; Turquety, S.; Hudman, R. C.; Weber, R. J.; Sullivan, A. P.; Peltier, R. E.; Atlas, E. L.; de Gouw, J. A.; Warneke, C.; Holloway, J. S.; Neuman, J. A.; Flocke, F. M.; Seinfeld, J. H.; Concentrations and sources of organic carbon aerosols in the free troposphere over North America. J. Geophys. Res. 2006, 111, D23S47, DOI:10.1029/2006JD0067705.

(5) Chebbi, A.; Carlier, P. Carboxylic acids in the troposphere, occurrence, sources, and sinks: A review. Atmos. Environ. 1996, 30 (24), 4233–4249.

(6) Heald, C. L.; Jacob, D. J.; Turquety, S.; Hudman, R. C.; Weber, R. J.; Sullivan, A. P.; Peltier, R. E.; Atlas, E. L.; de Gouw, J. A.; Warneke, C.; Holloway, J. S.; Neuman, J. A.; Flocke, F. M.; Seinfeld, J. H.; Concentrations and sources of organic carbon aerosols in the free troposphere over North America. J. Geophys. Res. 2006, 111, D23S47, DOI:10.1029/2006JD0067705.

(7) Blando, J. D.; Turpin, B. J. Secondary organic aerosol formation in cloud and fog droplets: A literature evaluation of plausibility. Atmos. Environ. 2000, 34, 1623–1632.

(8) Gelencser, A.; Varga, Z. Evaluation of the atmospheric significance of multiphase reactions in atmospheric secondary organic aerosol formation. Atmos. Chem. Phys. 2005, 5, 2823–2831.

(9) Lim, H.-J.; Ervens, B. Atmospheric oxalic acid and SOA production by acid-catalyzed particle-phase reactions. Atmos. Chem. Phys. 2007, 7, 814–817.

(10) Gelencser, A.; Varga, Z. Evaluation of the atmospheric significance of multiphase reactions in atmospheric secondary organic aerosol formation. Atmos. Chem. Phys. 2005, 5, 2823–2831.

(11) Limbeck, A.; Puxbaum, H.; Otter, L.; Scholes, M. C. Semivolatile behavior of dicarboxylic acids and other polar organic species at a rural background site (Nylsvley, RSA). Atmos. Environ. 2001, 35, 1853–1862.

(12) Fu, T.-M.; Jacob, D. J.; Wittrock, F.; Burrows, J. P.; Vrekoussis, M.; Henze, D. K. Global budgets of atmospheric glyoxal and methylglyoxal, and implications for formation of secondary organic aerosols. J. Geophys. Res. 2008, 113, D15303, DOI: 10.1029/2007JD009505.

(13) Feierabend, K. J.; Zhu, L.; Talukdar, R. K.; Burkholder, J. B. Rate coefficients for the OH + HCHO/OH (glyoxal) reaction (J. Phys. Chem. A 2008, 112 (1), 73–82).

(14) Volkamer, R.; San Martini, F.; Molina, L. T.; Salcedo, D.; Jimenez, J. L.; Molina, M. J. A missing sink for gas-phase glyoxal in Mexico City: Formation of secondary organic aerosol. Geophys. Res. Lett. 2007, 34, L19B07, DOI:10.1029/2007GL030752.

(15) Schweitzer, F.; Magli, L.; Mirabel, P.; George, C. Uptake rate measurements of methanesulfonic acid and glyoxal by aqueous droplets. J. Phys. Chem. A 1998, 102 (3), 593–600.

(16) Ip, H. S. S.; Huang, X. H. H.; Yu, J. Z. Effective Henry’s law constants of glyoxal, glyoxylic acid, and glycolic acid. Geophys. Res. Lett. 2009, 36, L01B02, DOI:10.1029/2008GL036212.

(17) Munger, J. W.; Jacob, D. J.; Daube, B. C.; Horowitz, L. W. Formaldehyde, glyoxal, and methylglyoxal in air and clouds at a rural mountain site in central Virginia. J. Geophys. Res. 1995, 100 (D5), 9325–9333.

(18) Carlton, A. G.; Turpin, B. J.; Altieri, K. E.; Seitzinger, S.; Reff, A.; Lim, H.-J.; Ervens, B. Atmospheric oxalic acid and SOA production from glyoxal: Results of aqueous photooxidation experiments. Atmos. Environ. 2007, 41 (35), 7588–7602.

(19) Carlton, A. G.; Turpin, B. J.; Altieri, K. E.; Seitzinger, S. P.; Mathur, R.; Roselle, S. J.; Weber, R. I. CMAC model performance enhanced when in-cloud secondary organic aerosol is included: Comparisons of organic carbon predictions with measurements. Environ. Sci. Technol. 2008, 42 (23), 8798–8802.

(20) Jang, M.; Czoschke, N. M.; Lee, S.; Kamens, R. M. Heterogeneous atmospheric aerosol production by acid-catalyzed particle-phase reactions. Science 2002, 298, 814–817.

(21) Inuma, Y.; Müller, C.; Böge, O.; Gnauk, T.; Herrmann, H. The formation of organic sulfate esters in the limonene ozonolysis secondary organic aerosol (SOA) under acidic conditions. Atmos. Environ. 2007, 41 (27), 5571–5583.
(22) Carlton, A. G.; Turpin, B. J.; Lim, H.-J.; Altieri, K. E.; Seitzinger, S., Link between isoprene and secondary organic aerosol (SOA): Pyruvic acid oxidation yields low volatility organic acids in clouds. Geophys. Res. Lett. 2006, 33.

(23) Perri, M. J.; Seitzinger, S.; Turpin, B. J. Secondary organic aerosol production from aqueous photooxidation of glycolaldehyde: Laboratory experiments. Atmos. Environ. 2009, 43 (8), 1487–1497.

(24) Altieri, K. E.; Carlton, A. G.; Lim, H.-J.; Turpin, B. J.; Seitzinger, S. P. Evidence for oligomer formation in clouds: Reactions of isoprene oxidation products. Environ. Sci. Technol. 2006, 40 (16), 4956–4960.

(25) Lim, H. J.; Carlton, A. G.; Turpin, B. J. Isoprene forms secondary organic aerosol through cloud processing: Model simulations. Environ. Sci. Technol. 2005, 39, 4441–4446.

(26) Ervens, B.; Feingold, G.; Frost, G. J.; Kreidenweis, S. M. A modeling study of aqueous production of dicarboxylic acids: 1. Chemical pathways and speciated organic mass production. J. Geophys. Res. 2004, 109, D15205, DOI:10.1029/2003JD004387.

(27) Altieri, K. E.; Seitzinger, S. P.; Carlton, A. G.; Turpin, B. J.; Klein, G. C.; Marshall, A. G. Oligomers formed through in-cloud methylglyoxal reactions: Chemical composition, properties, and mechanisms investigated by ultra-high resolution FT-ICR mass spectrometry. Atmos. Environ. 2008, 42 (7), 1476–1490.

(28) Noziere, B.; Dziedzic, P.; Cordova, A. Products and kinetics of the liquid-phase reaction of glyoxal catalyzed by ammonium ions (NH₄⁺). J. Phys. Chem. A 2008, 113 (1), 231–237.

(29) Shapiro, E. L.; Szprengiel, J.; Sareen, N.; Jen, C. N.; Giordano, M. R.; McNeil, V. F. Light-absorbing secondary organic material formed by glyoxal in aqueous aerosol mimics. Atmos. Chem. Phys. 2009, 9 (7), 2289–2300.

(30) De Haan, D. O.; Tolbert, M. A.; Jimenez, J. L., Atmospheric condensed-phase reactions of glyoxal with methylamine. Geophys. Res. Lett. 2009, 36.

(31) Galloway, M. M.; Chhabra, P. S.; Chan, A. W. H.; Surratt, J. D.; Flagan, R. C.; Seinfeld, J. H.; Keutsch, F. N. Glyoxal uptake on ammonium sulphate seed aerosol: Reaction products and reversibility of uptake under dark and irradiated conditions. Atmos. Chem. Phys. 2009, 9 (10), 3331–3345.

(32) Corrigan, A. L.; Hanley, S. W.; De Haan, D. O. Uptake of glyoxal by organic and inorganic aerosol. Environ. Sci. Technol. 2008, 42 (12), 4428–4433.

(33) Liggio, J.; Li, S.-M.; McLaren, R. Heterogeneous reactions of glyoxal on particulate matter: Identification of acetals and sulfate esters. Environ. Sci. Technol. 2005, 39 (6), 1532–1541.

(34) Leitzke, A.; Reisz, E.; Flyunt, R.; von Sonntag, C. The reactions of ozone with cinnamic acids: Formation and decay of 2-hydroxyperoxy-2-hydroxyacetic acid. J. Chem. Soc., Perkin Trans. 2 2001, (5), 793–797.