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Real-Time Studies of Iron Oxalate-Mediated Oxidation of Glycolaldehyde as a Model for Photochemical Aging of Aqueous Tropospheric Aerosols

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Abstract

The complexation of iron (III) with oxalic acid in aqueous solution yields a strongly absorbing chromophore that undergoes efficient photodissociation to give iron (II) and the carbon dioxide anion radical. Importantly, iron (III) oxalate complexes absorb near-UV radiation ($\lambda > 350$ nm), providing a potentially powerful source of oxidants in aqueous tropospheric chemistry. Although this photochemical system has been studied extensively, the mechanistic details associated with its role in the oxidation of dissolved organic matter within aqueous aerosol remain largely unknown. This study utilizes glycolaldehyde as a model organic species to examine the oxidation pathways and evolution of organic aerosol initiated by the photodissociation of aqueous iron (III) oxalate complexes. Hanging droplets (radius 1 mm) containing iron (III), oxalic acid, glycolaldehyde, and ammonium sulfate (pH ~ 3) are exposed to irradiation at 365 nm and sampled at discrete time points utilizing field-induced droplet ionization mass spectrometry (FIDI-MS). Glycolaldehyde is found to undergo rapid oxidation to form glyoxal, glycolic acid, and glyoxylic acid, but the formation of high molecular weight oligomers is not observed. For comparison, particle-phase experiments conducted in a laboratory chamber explore the reactive uptake of gas-phase glycolaldehyde onto aqueous seed aerosol containing iron and oxalic acid. The presence of iron oxalate in seed aerosol is found to inhibit aerosol growth. These results suggest that photodissociation of iron (III) oxalate can lead to the formation of volatile oxidation products in tropospheric aqueous aerosols.
Introduction

Tropospheric aqueous-phase chemistry plays a key role in the aging of dissolved organics in the atmosphere. Aqueous-phase processing may occur as a result of “dark” reactions such as acid catalysis, hydration, and oligomerization or may result from photochemical excitation of dissolved light-absorbing species. One of the key components directing the aging of dissolved organics is the availability of oxidative species such as the hydroxyl radical (OH) and hydrogen peroxide ($H_2O_2$), which may be present by direct generation in solution or by uptake from the gas phase. Many laboratory studies have investigated secondary organic aerosol (SOA) production initiated by photolysis of dissolved $H_2O_2$. There is increasing evidence, however, that processes involving the photolysis of photoactive organic and organometallic compounds may also be important sources of highly reactive aqueous oxidants.

Transition metal ions present in cloudwater or aqueous aerosol are known to undergo photoinitiated electron transfer processes that can be a significant source of oxidative species. Iron, the most abundant transition metal ion in tropospheric particles, can be found in aerosol particles originating from sea spray, mineral dust, and anthropogenic emissions. Measured concentrations in fog waters range from micromolar or less in rural areas to tens of micromolar in highly polluted environments. As a result of these significant concentrations, iron can play an important role in aqueous-phase tropospheric oxidation of organics. The reactions of dissolved iron-hydroxy complexes, or the Fenton reactions, are a well-characterized source of atmospheric oxidants. The direct reaction of hydrogen peroxide with iron (III) yields...
hydroxyl radicals in the dark Fenton reaction (1), whereas the photo-Fenton reaction yields hydroxyl radicals by photolysis of the iron (III) hydroxy complex (2-3).

\[ \text{Fe}^{2+} + \text{H}_2\text{O}_2 \rightarrow \text{Fe}^{3+} + \text{OH}^+ + \text{OH}^- \]  
(1)

\[ \text{Fe}^{3+} + 2\text{H}_2\text{O} \leftrightarrow \text{Fe(OH)}^2+ + \text{H}_3\text{O}^+ \]  
(2)

\[ \text{Fe(OH)}^2+ \rightarrow \text{Fe}^{3+} + \text{OH}^+ \]  
(3)

Several groups have recently explored the impact of iron photochemistry on SOA formation. Chu and co-workers studied the effects of iron (II) sulfate and iron (III) sulfate seed aerosol on SOA formation at 50% relative humidity (RH) and found that iron (II) complexes inhibited SOA formation, whereas iron (III) had little effect on SOA formation. They attributed the effect of iron (II) to the reduction of organic condensate by the iron species and subsequent disruption of oligomerization processes. Nguyen and co-workers investigated the uptake and oxidation of glycolaldehyde in aqueous aerosol containing hydrogen peroxide and iron complexes. They found that the photo-Fenton reaction significantly increased the degree of oxidation in aerosol particles when compared with \( \text{H}_2\text{O}_2 \) photolysis alone (ratio of O/C=0.9 with iron and \( \text{H}_2\text{O}_2 \) vs. O/C=0.5 with \( \text{H}_2\text{O}_2 \)).

Although laboratory experiments have demonstrated that photo-Fenton chemistry can lead to significant oxidation of organics, the role of such processes in atmospheric aerosol is not clear, as the reactions are highly dependent on pH, iron concentration, and the concentration of other ligands that readily complex with iron. Notably, the formation of stable complexes between iron and dicarboxylate ligands can have a significant impact on the photochemical reaction pathways. Complexation between the oxalate anion, one of the most abundant low-molecular weight organic compounds found in aqueous aerosol, and iron has long been known to generate significant yields of oxidative species by photochemical reduction of iron. Initially studied as a chemical actinometer, seminal work by Zuo and Hoigné demonstrated...
that photolysis of iron-oxalate complexes could lead to the generation of oxidative species under atmospherically relevant conditions.\textsuperscript{25}

The proposed pathway for the photochemical generation of oxidative species from iron (III) oxalate complexes is shown in reactions 4-10 below. Depending on the pH, ionic strength, and concentration of iron and oxalate, three complexes are formed: \( \text{Fe(C}_2\text{O}_4\text{)}^+ \), which has low photochemical reactivity,\textsuperscript{38} and \( \text{Fe(C}_2\text{O}_4\text{)}_2^- \) and \( \text{Fe(C}_2\text{O}_4\text{)}_3^{3-} \), both of which undergo photochemical reduction of iron with high quantum yield.\textsuperscript{31, 39} The mechanism of photochemical dissociation of \( \text{Fe(C}_2\text{O}_4\text{)}_3^{3-} \) has been investigated in detail, with two groups presenting evidence for either intermolecular electron transfer (4a) or intramolecular electron transfer (4b) as the primary reaction pathway.\textsuperscript{40-43} The mechanism of dissociation of \( \text{Fe(C}_2\text{O}_4\text{)}_2^- \) has not been studied extensively but is also expected to yield the oxalate anion radical, which rapidly dissociates to \( \text{CO}_2 \) and \( \text{CO}_2^{•-} \) (reactions 5-6).\textsuperscript{25} The subsequent reaction of \( \text{CO}_2^{•-} \) with \( \text{O}_2 \) leads to the eventual formation of \( \text{H}_2\text{O}_2 \) (reactions 7-9), which can then interact directly with \( \text{Fe}^{2+} \) (reaction 1) or with \( \text{FeC}_2\text{O}_4 \) (reaction 10) to produce \( \text{OH} \).\textsuperscript{44}

\[
\begin{align*}
\text{Fe(C}_2\text{O}_4\text{)}_3^{3-} & \xrightarrow{\text{hv}} \text{Fe(C}_2\text{O}_4\text{)}_2^- + \text{C}_2\text{O}_4^{•-} \quad (4a) \\
\text{Fe(C}_2\text{O}_4\text{)}_2^- & \xrightarrow{\text{hv}} \text{Fe(C}_2\text{O}_4\text{)}_2^- + 2\text{CO}_2^{•-} \quad (4b) \\
\text{Fe(C}_2\text{O}_4\text{)}_2 & \xrightarrow{\text{hv}} \text{Fe}^{2+} + \text{C}_2\text{O}_4^{2-} + \text{C}_2\text{O}_4^{•-} \quad (5) \\
\text{C}_2\text{O}_4^{•-} & \rightarrow \text{CO}_2^{•-} + \text{CO}_2 \quad (6) \\
\text{CO}_2^{•-} + \text{O}_2 & \rightarrow \text{CO}_2 + \text{O}_2^{•-} \quad (7) \\
\text{O}_2^{•-} + \text{H}_3\text{O}^+ & \rightarrow \text{HO}_2^\text{•} + \text{H}_2\text{O} \quad (8) \\
2\text{HO}_2^\text{•} & \rightarrow \text{H}_2\text{O}_2 + \text{O}_2 \quad (9) \\
\text{FeC}_2\text{O}_4 + \text{H}_2\text{O}_2 & \rightarrow \text{FeC}_2\text{O}_4^+ + \text{OH}^\text{•} + \text{OH}^- \quad (10)
\end{align*}
\]

Although the mechanism and kinetics of dissociation of iron (III) oxalate complexes have been examined in detail, much less information is available on the oxidation of atmospherically
relevant compounds by this photochemical system. Zuo and Zhan found that the presence of iron (III) oxalate complexes increases the rate of oxidation of sulfur dioxide compared to iron alone under atmospherically relevant conditions.\textsuperscript{45} In field sampling studies, Sorooshian et al. found an inverse correlation between concentrations of dissolved iron and oxalate in stratocumulus cloudwater above the northeastern pacific ocean, suggesting that ferrioxalate photochemistry may play an important role in determining cloudwater composition under certain conditions.\textsuperscript{22} These studies serve as initial demonstrations of the potential importance of such reactions in aqueous tropospheric chemistry, but further laboratory and field studies are necessary to better discern the complex interplay of reactions influencing the oxidation of dissolved organic compounds.

Glycolaldehyde serves as an excellent model system to study the influence of liquid-phase ferrioxalate photochemistry on the oxidation of dissolved organics. A major product of isoprene oxidation, glycolaldehyde is produced with an estimated global flux greater than 42 Tg C yr\textsuperscript{-1}.\textsuperscript{13, 46, 47} The abundance of glycolaldehyde, along with its high solubility,\textsuperscript{48} means that its oxidation processes may play a significant role in the formation of SOA in aqueous aerosol.

This study explores the photochemically initiated reaction pathways involved in the oxidation of aqueous glycolaldehyde in the presence of ferrioxalate complexes. Real-time studies of photochemical reactions in microliter droplets utilizing field-induced droplet ionization mass spectrometry (FIDI-MS)\textsuperscript{49, 50} provide insight into the changes in chemical composition of this model system upon UV irradiation. Complementary particle-phase environmental chamber experiments\textsuperscript{51, 52} examine the evolution of iron-doped aqueous aerosol in the presence of gas-phase glycolaldehyde under simulated atmospheric conditions.
Experimental

Field-Induced Droplet Ionization Mass Spectrometry

The FIDI-MS source employed in this study is based upon an initial design described by Grimm and co-workers. The general design and features of the updated source are described here, with further details given in the supporting information. A hanging droplet of 1.5-2 mm in diameter (2-4 µL) is suspended on the end of a stainless steel capillary between two parallel plate electrodes separated by 6.3 mm (Figure 1c). The parallel plates are mounted to a translation stage to allow alignment of an aperture in the electrically grounded plate with the atmospheric pressure inlet of an LTQ-XL mass spectrometer (Thermo-Fisher, Waltham, MA). The capillary is mounted on a separate translation stage to provide for placement of the droplet midway between the two plates in alignment with the inlet of the LTQ-XL. A droplet is formed from liquid fed through the capillary using a motorized syringe pump. Mass spectrometric sampling of the hanging droplet is accomplished by application of a pulsed high voltage (3-5 kV, 100 ms duration) to the back parallel plate and to the suspended capillary at half the magnitude applied to the back plate to maintain field homogeneity between the electrodes. When a sufficiently high voltage is applied, the electrical forces overcome the surface tension of the droplet, resulting in the bipolar ejection of highly charged progeny droplets of less than 1 µm in diameter from opposite ends of the suspended droplet. Charged droplets of a specific polarity enter the transfer capillary of the mass spectrometer, resulting in the detection of gas-phase ions in a manner similar to electrospray ionization (ESI). Sampling of either positive or negative ions
is achieved by switching the polarity of the high voltage applied to the back plate and capillary. The pulsed high voltage is controlled by a custom power supply and software described elsewhere. 

Figure 1. Experimental apparatus for studying photochemistry utilizing FIDI-MS. The primary components of the system are depicted schematically in (a) and shown as utilized in the laboratory in (b). A CCD camera image of the FIDI region is shown in (c).

The entire FIDI source is mounted within a custom enclosure that allows for control of the environment in which the droplet is suspended, as shown in Figure 1a,b. The enclosure is equipped with a fused silica window to allow for the study of photochemical reactions and for visual droplet monitoring during experiments. To study the photochemistry of suspended droplets in the absence of oxygen, the chamber was flushed with nitrogen at a flow rate of 1.2
L/min for ~10 minutes prior to sampling, and the flow of nitrogen was maintained throughout the experiment; the sample was also degassed by bubbling with nitrogen. For all other experiments undertaken in this work, the enclosure was left open to laboratory air. Photochemical reactions in the hanging droplet are accomplished by irradiation with 365 nm light generated by a fiber optic-coupled LED (FCS-0365-000, Mightex Systems, Pleasanton, CA). The light is focused onto the droplet using a collimating lens (74-UV, Ocean Optics, Dunedin, FL) coupled to a spherical focusing lens. A radiant flux of 760 ± 5 µW was measured at the location of the hanging droplet with use of a thermal power sensor (S302C, ThorLabs, Newton, NJ) coupled to a power and energy meter (PM100USB, Thorlabs). This measurement yields an upper limit of $1.4 \times 10^{15}$ photons cm$^{-2}$ s$^{-1}$ on the photon flux encountering the hanging droplet, approximately one order of magnitude greater than the solar actinic flux at 365 nm with a 30° solar zenith angle.

Photochemical experiments employing FIDI-MS were performed in solutions containing 0.5 mM H$_2$SO$_4$, 100 µM NH$_3$, 50 µM FeCl$_3$, 250 µM oxalic acid, and 250 µM glycolaldehyde. Control experiments were also performed by excluding iron, oxalic acid, or glycolaldehyde from the mixture. In a typical experiment, a droplet of the solution was formed on the end of the capillary and allowed to rest for 1 minute prior to exposure to 365 nm radiation. Individual droplets were then sampled by application of a high voltage pulse after 0-3 min of irradiation. The presented spectra are averaged from four experiments on discrete droplets at each time point. Between experiments, the sample line was rinsed thoroughly with 10 mM H$_2$SO$_4$ to prevent the accumulation of iron precipitates.

**Aerosol Chamber Instrumentation**

Aerosol size distributions were measured with a custom-built scanning mobility particle sizer (SMPS) consisting of a differential mobility analyzer (model 3081, TSI, Shoreview, MN)
coupled to a condensation particle counter (model 3010, TSI). Bulk aerosol composition was monitored by a high-resolution time-of-flight Aerodyne Aerosol Mass Spectrometer (HR-AMS). The principles and details of the HR-AMS are described elsewhere.\textsuperscript{58,59} Prior to experiments, the instrument’s ionization efficiency was calibrated with 350 nm $\text{NH}_4\text{NO}_3$ particles. The instrument was operated in V-mode, and data were analyzed following the recommendations of Aiken et al. and Canagaratna et al.\textsuperscript{60,61}

Volatile organic compounds (VOCs) were monitored by a custom-built chemical ionization mass spectrometer (CIMS) with CF$_3$O$^-$ reagent ions.\textsuperscript{62,63} The instrument was primarily operated in negative mode, which allows for the detection of analyte [R] and reagent clusters \([R\cdot\text{CF}_3\text{O}]^-.\) For strongly acidic species \([H\cdot X],\) the transfer product \([H\cdot X\cdot F]^-\) is detected. For select species such as glycolaldehyde, the instrument was operated in tandem mass spectrometry (MS/MS) mode.\textsuperscript{63} In this arrangement, a parent ion selected by the first quadrupole is exposed to elevated pressure of N$_2$ and fragmented into daughter ions that correspond to a unique compound at a common unit mass. Here, the daughter ion of the m/z 145/85 cluster is taken to reflect the glycolaldehyde signal.

Gas-phase concentrations of O$_3$ and NO + NO$_2$ (NO\textsubscript{x}) were measured by Horiba (model APOA-360) and Teledyne monitors (model T200), respectively. The temperature and RH were monitored by Vaisala probes.

**Aerosol Chamber Experimental Protocol**

Experiments were performed in the Caltech dual 24 m$^3$ Teflon chambers.\textsuperscript{51,52} Prior to each experiment, the chamber was flushed with air scrubbed by activated carbon, silica gel, Purafil Media, and molecular sieves to achieve a particle background < 10 cm$^{-3}$, O$_3$ background < 10 ppb, and NO\textsubscript{x} background < 5 ppb. Once cleaned, chamber air was humidified to $\geq 80\%$ relative
humidity (RH) using a water circulator coupled to a Nafion membrane humidifier. Aerosols were injected into the chamber through a wet-walled humidifier; consequently, particles are assumed to be deliquesced. All experiments were conducted at 25°C. Further details of the aerosol chamber experimental protocol can be found in the supporting information.

All experiments were performed in the presence of aerosol and gas-phase glycolaldehyde at low-NO$_x$ and low-HO$_x$ conditions (Table 1). Typical gas-phase oxidants (e.g. O$_3$, OH, NO) were not purposely injected into the chamber, and the resulting chemistry is therefore attributed to glycolaldehyde photolysis and interactions between volatilized glycolaldehyde and particles.

Over the course of an experiment, NO$_x$ levels never rose above the detection limits of the Teledyne monitors. Ozone rose slightly (~10 ppb) after photochemistry was induced, suggesting that low, pre-existing NO$_x$ concentrations were present in the chamber.

**Table 1.** Experimental Conditions Summarizing Atomizer Solution Composition ([Fe]$_0$, [Oxalate]$_0$, pH), Initial Glycolaldehyde Concentrations, and Chamber Relative Humidity.$^a$

<table>
<thead>
<tr>
<th>Exp. Number and Description$^b$</th>
<th>[Fe]$_0$ (µM)</th>
<th>[Oxalate]$_0$ (µM)</th>
<th>pH</th>
<th>[Gly]$_0$ (ppb)</th>
<th>RH (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. No Seed</td>
<td>0</td>
<td>0</td>
<td>---</td>
<td>85</td>
<td>88</td>
</tr>
<tr>
<td>2. AS+Gly</td>
<td>0</td>
<td>500</td>
<td>3</td>
<td>83</td>
<td>85</td>
</tr>
<tr>
<td>3. 0.005% Fe + Gly</td>
<td>2</td>
<td>500</td>
<td>3</td>
<td>97</td>
<td>90</td>
</tr>
<tr>
<td>4. 0.03% Fe + Gly</td>
<td>10</td>
<td>500</td>
<td>3</td>
<td>93</td>
<td>87</td>
</tr>
<tr>
<td>5. 0.3% Fe+ Gly</td>
<td>100</td>
<td>500</td>
<td>3</td>
<td>81</td>
<td>94</td>
</tr>
<tr>
<td>6. 5% Fe + Gly</td>
<td>2000</td>
<td>500</td>
<td>3</td>
<td>100</td>
<td>92</td>
</tr>
</tbody>
</table>

$^a$ All atomizer solutions contain 0.015 M ammonium sulfate (AS) and sulfuric acid added to achieve a pH of 3

$^b$ Aerosol iron content is represented as the percentage of total inorganic aerosol mass
The aerosol seed composition was varied to investigate the impact of iron interactions on the uptake and particle-phase oxidation of glycolaldehyde (Table 1). Varying concentrations of iron (II) sulfate and oxalic acid were added to atomizer solutions of 0.015 M ammonium sulfate. The pH of the solutions was adjusted with 10 wt% sulfuric acid to an initial pH of 3 as measured by an Orion pH meter (Thermo-Fisher, Waltham, MA). We expect the pH to quickly change since the photochemistry of ferrioxalate systems is known to consume and generate hydronium ions. Solutions were prepared fresh daily to avoid the slow formation of iron (III) oxide from the reaction of dissolved iron (II) with oxygen. Experiments in the absence of iron were performed first to avoid the possibility of iron contamination within the atomizer, on chamber walls, and along sampling lines.

Iron concentrations were chosen to reflect a range of iron content observed in atmospheric water. Iron mass fractions in aerosol ranging from 3.5-7.7% have been reported, and iron solubilities in atmospheric waters are reported to range between 0.018 and 22%. For example, measurements on ambient cloud water samples found soluble iron concentrations as high as 0.57 µM, and concentrations up to 1.3 µM were observed in samples impacted by shipping emissions. Based on this wide range of soluble iron content, we doped our atomizer solutions with 2, 10, 100 or 2000 µM of iron, which correspond to inorganic weight fractions of 0.005, 0.03, 0.3 and 5%, respectively. Assuming that the aerosol exhibits hygroscopic properties similar to pure ammonium sulfate, we expect atomizer solutions to concentrate by approximately two orders of magnitude; thus, these conditions reflect systems analogous to wet aerosol as opposed to the dilute cloud water systems described above. Given the high concentrations of oxalate observed in atmospheric waters (up to ~0.12 µg m$^{-3}$), particles were also doped with oxalate concentrations comparable to that of iron (500 µM). We limit our interpretation of chamber data...
to trends associated with iron doping. Future studies may consider the effect of oxalate in
comparison to other organic acids.

Photochemical Box Model

A photochemical box model is employed to investigate glycolaldehyde losses due to gas-phase
processes. Equations describing glycolaldehyde photolysis and reaction with OH are outlined in
the supporting information. The photochemical model was initiated with a glycolaldehyde
concentration of 85 ppb (Exp. 1, Table 1).

Results and Discussion

Photochemistry of Iron (III) Oxalate Complexes and Glycolaldehyde by FIDI-MS

Figure 2 shows the FIDI-MS spectra of hanging droplets containing iron (III) oxalate
complexes and glycolaldehyde exposed to irradiation at 365 nm for 0-3 minutes. The HSO$_4^-$
anion and its dimer are the most prominent ions in the spectrum due to the high concentration of
sulfate in solution (0.5 mM) utilized to achieve a pH of ~3. Oxalic acid (Ox) is also detected
prior to irradiation (0 min) as a monomer, homodimer, and heterodimer with hydrogen sulfate
(m/z 89, 179, and 187, respectively), and the heterodimer of glycolaldehyde hydrate
(Glycol+H$_2$O) with hydrogen sulfate is also observed at m/z 175. The Fe(C$_2$O$_4$)$_2^-$ ion, predicted
by equilibrium calculations to be the dominant complex of iron (III) in the sample (Figure S4), is
detected at m/z 232. Upon irradiation, the Fe(C$_2$O$_4$)$_2^-$ complex and oxalic acid ions are rapidly
depleted, and new ions are observed at m/z 173, 175, and 191. These photochemical oxidation
products are assigned to glycolic acid (glyc, m/z 173), glyoxylic acid (glyox, m/z 171), and
glyoxal dihydrate (m/z 191), all observed as adducts with hydrogen sulfate. Formic acid is also
observed as a dimer with hydrogen sulfate at m/z 143 with low intensity after 3 min of irradiation. It is possible that these ions represent the formation of organosulfate compounds rather than noncovalent complexes, but the free glycolic acid and glyoxylic acid ions (m/z 73 and 75, respectively), also increase in intensity over the course of the experiment, suggesting that the observed species are likely adducts formed during gas-phase ion generation. The formation of these product ions is not observed in control experiments in which iron, glycolaldehyde, or oxalic acid is excluded from the reaction mixture (Figures S5, S6).

**Figure 2.** Monitoring the photochemistry of aqueous glycolaldehyde and iron (III) oxalate complexes by FIDI-MS. The left column shows the full spectrum, and the right column highlights the region where glycolaldehyde and its oxidation products are detected (10x zoom). The depletion of Fe(C$_2$O$_4$)$_2^-$ and oxalic acid (Ox) are clearly observed over the course of three
minutes, and the oxidation of glycolaldehyde (glycol) to glycolic acid (glyc), glyoxylic acid (glyox), and glyoxal is observed.

Scheme 1. Oxidation Pathway of Glycolaldehyde in the Presence of Iron (III) Oxalate Complexes.

The observed chemistry of the glycolaldehyde-ferrioxalate systems is summarized in Scheme 1. As described in reactions 4-10, photodissociation of iron oxalate complexes leads to the formation of hydroxyl radicals that oxidize glycolaldehyde to form glyoxal and glycolic acid. Further oxidation yields formic acid and glyoxylic acid from glycolaldehyde and glycolic acid, respectively. Glyoxylic acid may then undergo one final oxidation process to regenerate oxalic acid, whereas formic acid oxidation leads to the formation of carbon dioxide. Alternatively,
hydrated glyoxal, detected prominently in FIDI-MS experiments in agreement with previous equilibrium measurements,\textsuperscript{66-68} can be oxidized to glyoxylic acid hydrate and subsequently to oxalic acid.\textsuperscript{4} The branching ratios of each pathway are challenging to determine within the current experiment, as the signal intensity in FIDI, similar to ESI, does not depend linearly on the molecule concentration in solution and thus provides only semi-quantitative measurement.\textsuperscript{54}

The observed oxidation pathways are similar to those observed by Perri and co-workers for the oxidation of aqueous glycolaldehyde by hydroxyl radicals produced by H$_2$O$_2$ photolysis initiated by a 254 nm Hg lamp.\textsuperscript{13} They also observed higher molecular weight oligomers in significant abundance and identified malonic and succinic acid as two of the major oligomerization products. Further study by Ortiz-Montalvo and co-workers identified malic and tartaric acids as additional oligomerization products.\textsuperscript{69} Similarly, in a study of aerosol formation from reactive uptake of glycolaldehyde on seed aerosol doped with iron and H$_2$O$_2$, Nguyen and co-workers proposed that oligimerization products of glycolaldehyde contributed to the high observed O/C ratios of aerosol organics.\textsuperscript{26} In contrast, the FIDI-MS experiments presented in this study did not detect the formation of any oligomers in significant abundance. The comparatively high flux of oxidative species produced by the photodissociation of iron (III) oxalate complexes may favor sequential oxidation over oligomerization under the conditions utilized in this study.

\textit{Photochemistry of Iron Oxalate and Glycolaldehyde in Deoxygnated Environment}

The photochemistry of the iron (III) oxalate and glycolaldehyde system was also investigated in the absence of oxygen by purging both the sample solution and the reaction chamber with nitrogen gas. As shown in Figure 3, little change in the composition of the droplets is observed during irradiation under such conditions. In contrast to the nearly complete depletion of oxalate in the presence of oxygen, the ions associated with oxalic acid only marginally decrease in
intensity over the course of the reaction. In addition, the intensity of the Fe(C_2O_4)_2^- is significantly diminished after 1 min of irradiation, and little oxidation of glycolaldehyde is observed.

Figure 3. Photochemistry of deoxygenated solutions of iron (III) oxalate and glycolaldehyde. The full spectrum is shown on the left, and the right column shows a 10x zoom on the region of detection for glycolaldehyde and its oxidation products. In the absence of dissolved oxygen, little oxidation of glycolaldehyde is observed, and oxalate is not depleted through the course of the experiment. The Fe(C_2O_4)_2^- ion is significantly depleted after 1 min of irradiation.

The absence of glycolaldehyde oxidation products in deoxygenated solutions is readily explained by the necessity of oxygen for the generation of hydroxyl radicals as detailed in reactions 4-10. The small amount of oxidation observed is attributed to trace concentrations of
oxygen gas in the enclosed reaction environment. The presence of oxygen is also necessary for the regeneration of iron (III) following photo-catalyzed reduction. Weller and co-workers recently measured a quantum yield of 0.8 ± 0.1 for iron (II) formation under conditions similar to those utilized in this study, indicating that iron (III) should be rapidly depleted in the absence of oxygen.\(^{31}\) Consistent with this result, we do not observe the \(\text{Fe}(_2\text{C}_2\text{O}_4)_2^-\) ion following irradiation in a nitrogen atmosphere. Similarly, little change in the intensity of oxalate ions is observed, as only 20-30\% of the oxalic acid can be dissociated before all of the iron present is reduced. Zuo and Hoigne also observed that nearly all of the iron (III) was reduced to iron (II) rapidly in deoxygenated solutions, whereas in oxygenated solution a steady state was reached once approximately 70\% of the iron (III) was reduced.\(^{25}\)

**Chamber Experiments**

Figure 4 summarizes the evolution of organic aerosol and gas-phase glycolaldehyde concentrations during chamber photooxidation experiments. Organic mass is presented as the ratio of \(\text{Org}/\text{SO}_4\) to minimize uncertainties in particle wall loss, HR-AMS collection efficiencies, and line losses. Across all experiments, the glycolaldehyde signal decays after the initiation of photochemistry. The glycolaldehyde decay rate is comparable among all experiments. Glycolaldehyde will photolyze in the Caltech chamber with a photolysis rate constant of \(j_{\text{Gly}} = 2.1 \times 10^{-6} \text{ s}^{-1}\). Furthermore, glycolaldehyde photolysis leads to reactive products, such as formaldehyde, that promote OH formation and enhance glycolaldehyde decay.\(^{70}\) In experiments absent of seed, glycolaldehyde decay is well-represented by a simple photochemical model (see supporting information); thus, the bulk of glycolaldehyde decay is attributable to non-heterogeneous processes.
For seeded experiments doped with low concentrations of iron (Exp. 2 and 3), the loss of glycolaldehyde after 1000 min is comparable to that of Exp. 1. In contrast, seeded experiments doped with high concentrations of iron (Exp. 4-6) exhibit an additional ~10% decrease in glycolaldehyde. These modest increases in glycolaldehyde decay may reflect enhanced losses due to particle-phase oxidation in the presence of iron. Gas-particle partitioning in systems such as glycolaldehyde and glyoxal, which participate in dark particle-phase reactions to form diols, hemiacetals, and amines/imines,\textsuperscript{26, 71} is kinetically limited.\textsuperscript{1, 26, 72, 73} Enhanced heterogeneous reactions would increase the rate of VOC partitioning and consequently deplete gas-phase concentrations. We note, however, that these processes likely play a small role in comparison to glycolaldehyde photolysis.

Figure 4. Measurements of (a) aerosol Org/SO\(_4^{2-}\) ratios and (b) gas-phase glycolaldehyde during chamber experiments. Organic aerosol formation is suppressed when iron is present in the seed.
aerosol (a). Enhanced particle-phase iron concentrations lead to greater glycolaldehyde decay (b); however, most of the glycolaldehyde decay is determined by gas-phase photochemical processes (No Seed, Model).

The impact of ferrioxalate photochemistry is most pronounced on organic aerosol concentrations. Figure 4a illustrates organic aerosol growth for systems doped with various concentrations of iron. In experiments absent of iron, organic aerosol is observed to increase upon the initiation of photochemistry. As demonstrated in Figure S2, this growth (and simultaneous gas-phase glycolaldehyde decay) stops when lights are turned off. In the supporting information, we demonstrate that ~50% of the decay in gas-phase glycolaldehyde is likely attributable to the reaction with OH. Glycolaldehyde reacts with OH in the gas phase to form glyoxal at high yields (~22%). Using the photochemical box model, we estimate that 3.5 ppb of glyoxal could have formed as a result of glycolaldehyde photochemistry. Previous studies have shown that glyoxal efficiently forms SOA on ammonium sulfate seed under dark and irradiated conditions. Kampf et al. observed 0.75 µg m⁻³ of SOA formation in the presence of low glycolaldehyde concentrations (~10 ppb). The authors performed these experiments under conditions similar to those described here. We hypothesize that a significant fraction of SOA formed during iron-free experiments could have resulted from glyoxal partitioning; however, quantification of glyoxal SOA is beyond the scope of this study.

In systems doped with iron, organic aerosol growth is inhibited. Even for the lowest concentrations of iron, we observe no noticeable increase in organic aerosol concentrations. This growth inhibition may result from a number of processes. First, glycolaldehyde was likely oxidized in the particle according to the reactions outlined by Scheme 1 to form CO₂ and other volatile species. This pathway is supported by FIDI-MS experiments in which glyoxal, glycolic
acid, and glyoxylic acid form shortly after irradiation. Second, it is possible that oligomerization processes were interrupted due to the oxidation of hydrated glyoxal monomers and/or glycolaldehyde monomers. As discussed for FIDI-MS experiments, glyoxal diols can be oxidized by OH to form oxalic acid via the glyoxylic acid pathway. This oxidation could shift oligomer equilibrium and consequently lead to a loss of SOA.

The proposed chamber mechanism is consistent with known heterogeneous reactions for glyoxal and glycolaldehyde. It is notable that several oxidation pathways can lead to the formation of oxalic acid. With additional oxalic acid, ferrioxalate complexes would likely reform and subsequently lead to the recycling of hydroxyl radicals. We hypothesize that this process could set up a catalytic cycle that would ultimately lead to the destruction of SOA precursors (such as glyoxal diols). This inhibition mechanism is consistent with the lack of SOA formed in any experiment performed with iron-containing aerosols.

Previous studies have demonstrated that iron chemistry has notable impacts on organic aerosol consumption and/or inhibition. Using dark Fenton chemistry as a source of OH, Daumit et al. oxidized suspended particles of polyols and observed substantial loss of carbon to the gas phase. The authors demonstrated that the product distributions in suspended particles were substantially different than those observed in bulk solutions due to the differences in liquid water content. Chu et al. conducted conventional SOA yield experiments in the presence of inorganic seed and found substantial reduction in SOA formation in systems containing FeSO₄. The authors attribute these observations to iron reduction reactions and interruption of dark accretion reactions; however, losses are also likely to occur as a result of photooxidation of iron-organic complexes.
The studies described above were conducted using seed composed of pure FeSO$_4$. The results presented here indicate that even a trace amount of iron (0.005 wt%) has the potential to strongly impact aerosol composition. Ferrioxalate chemistry is likely to apply to other aqueous systems where oxalic acid is produced in large abundance. In addition, these results demonstrate that oxidation products in aqueous organic systems, such as oxalate formed by the OH + glycolaldehyde mechanism, may participate in the recycling of oxidizing species and subsequent oxidation of dissolved organic material. Further laboratory investigations into the role of iron (III) oxalate photochemistry in reactive uptake, along with additional field measurements, are necessary to better constrain the role of such complexes in aqueous tropospheric photochemistry.

**Supporting Information.** Materials used in experiments, design of FIDI-MS apparatus, environmental chamber protocol, predicted equilibrium complexes of iron, FIDI-MS control experiments, and details of the photochemical box model. This material is available free of charge via the Internet at http://pubs.acs.org.

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