Photochemical degradation of iron(III) citrate/citric acid aerosol quantified with the combination of three complementary experimental techniques and a kinetic process model

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Received: 29 July 2020 – Discussion started: 10 August 2020
Revised: 3 November 2020 – Accepted: 9 November 2020 – Published: 13 January 2021

Abstract. Iron(III) carboxylate photochemistry plays an important role in aerosol aging, especially in the lower troposphere. These complexes can absorb light over a broad wavelength range, inducing the reduction of iron(III) and the oxidation of carboxylate ligands. In the presence of O2, the ensuing radical chemistry leads to further decarboxylation, and the production of \( \cdot \)OH, HO2, peroxides, and oxygenated volatile organic compounds, contributing to particle mass loss. The \( \cdot \)OH, HO2, and peroxides in turn reoxidize iron(II) back to iron(III), closing a photocatalytic cycle. This cycle is repeated, resulting in continual mass loss due to the release of CO2 and other volatile compounds. In a cold and/or dry atmosphere, organic aerosol particles tend to attain highly viscous states. While the impact of reduced mobility of aerosol constituents on dark chemical reactions has received substantial attention, studies on the effect of high viscosity on photochemical processes are scarce. Here, we choose iron(III) citrate (\( \text{Fe}^{III}(\text{Cit}) \)) as a model light-absorbing iron carboxylate complex that induces citric acid (CA) degradation to investigate how transport limitations influence photochemical processes. Three complementary experimental approaches were used to investigate kinetic transport limitations. The mass loss of single, levitated particles was measured with an electrodynamic balance, the oxidation state of deposited particles was measured with X-ray spectromicroscopy, and HO2 radical production and release into the gas phase was observed in coated-wall flow-tube experiments. We observed significant photochemical degradation with up to 80 % mass loss within 24 h of light exposure. Interestingly, we also observed that mass loss always accelerated during irradiation, resulting in an increase of the mass loss rate by about a factor of 10. When we increased relative humidity (RH), the observed particle mass loss rate also increased. This is consistent with strong kinetic transport limitations for highly viscous particles. To quantitatively compare these experiments and determine important physical and chemical parameters, a numerical multilayered photochemical reaction and diffusion (PRAD) model was developed that treats chemical reactions and the transport of various species. The PRAD model was tuned to simultaneously reproduce all experimental results as closely as possible and captured the essential chemistry and transport during irradiation. In particular, the photolysis rate of \( \text{Fe}^{III} \), the reoxidation rate of \( \text{Fe}^{II} \), HO2 production, and the diffusivity of O2 in aqueous \( \text{Fe}^{III}(\text{Cit}) / \text{CA} \) system as function of RH and \( \text{Fe}^{III}(\text{Cit}) / \text{CA} \) molar ratio could be constrained. This led to satisfactory agreement within model
uncertainty for most but not all experiments performed. Photochemical degradation under atmospheric conditions predicted by the PRAD model shows that release of CO$_2$ and repartitioning of organic compounds to the gas phase may be very important when attempting to accurately predict organic aerosol aging processes.

1 Introduction

Photochemistry in the atmosphere (either in the gas phase or in the particle phase) plays an important role in aerosol aging processes. Photochemically produced free radicals in the gas phase (mainly OH) can be taken up by aerosol particles, inducing multiphase chemistry. However, uptake is limited by the collision rate and condensed phase molecular transport when diffusion coefficients are sufficiently low, which restricts chemical reactions to the near-surface region of the particle. In contrast, photochemically generated radicals in aerosol particles can be present throughout their bulk due to light penetrating their whole volume. Radical production induced by a direct photochemical reaction occurs when the energy of light quanta is high enough (mostly the UV part of the solar spectrum) to cause bond cleavage or rearrangement in a molecule. In the lower troposphere where UV light intensity is low, indirect photochemistry initiated by near UV and visible radiation may become significant. Important indirect photochemical processes are transition metal complex photochemistry and photosensitized processes (Corral Arroyo et al., 2018; George et al., 2015). This work focuses on iron carboxylate-catalyzed photochemistry due to its abundance and reactivity in the atmosphere.

Iron is the most abundant transition metal in the earth’s crust. Wind erosion is the main source of iron in the atmosphere, but anthropogenic activities such as industrial processes, traffic, and combustion processes can also potentially release iron in particulate form (Deguillaume et al., 2005). Depending on parameters such as temperature, pH value, ionic strength, and the concentration of involved substances, iron can combine with inorganic or organic ligands to form complexes (Deguillaume et al., 2005; Faust and Hoigne, 1990; Kieber et al., 2005). Iron can be found complexed with low molecular weight inorganic species such as the hydroxide anion (OH$^-$), sulfate (SO$_4^{2-}$), and sulfite (SO$_3^{2-}$) (Brandt and van Eldik, 1995; Hofmann et al., 1991; Weschler et al., 1986). Quantifying iron atmospheric processing and solubility is of global importance, especially for nutrient input into the world’s oceans (Hamilton et al., 2019; Kanakidou et al., 2018). Heterogeneous chemistry involving particulate iron and SO$_2$ can result in sulfate formation and increase aerosol loading (Grgić et al., 1998, 1999; Grgić, 2009). Additionally, iron photochemical processing in aerosol particles, fog droplets, and cloud water is an important radical source (Bianco et al., 2020; Abida et al., 2012) and sink for organic compounds (Weller et al., 2014, 2013; Herrmann et al., 2015). Organic compounds are a major component in atmospheric aerosol particles and have received more and more attention as potential ligands for iron(III) complexation. For instance, humic-like substances derived from water-soluble organic compounds have been reported to be strong chelating ligands with iron(III) (Dou et al., 2015; Kieber et al., 2003; Okochi and Brimblecombe, 2002; Willey et al., 2000). Oxalate and other carboxylates have been identified to be important ligands for iron(III) because they are available in sufficient amounts (Chebbi and Carlier, 1996; Kahnt et al., 2014; Kawamura et al., 1985) and the carboxylate groups are acidic enough to dissociate and chelate with iron(III) at atmospheric pH values (Okochi and Brimblecombe, 2002). Field studies have confirmed that soluble iron is mostly in complexes with carboxylate functions (Tapparo et al., 2020; Tao and Murphy, 2019).

In atmospheric aqueous phases, iron normally exists in oxidation states (II) and (III) and they can convert into each other via redox cycling. The ratio between iron(II) and iron(III) in aerosol particles is quite variable and depends on several factors such as the presence of light, oxidizing compounds, and ligands. For example, Grgić et al. (1999) reported that the concentration ratio of iron(II) to iron(III) varied between 0.9 and 3.1 in urban aerosol particles with a size range of 0.4–1.6 µm. Iron(III) carboxylate complexes [Fe$^{III}$ (OOC–R)$_2$$^{2+}$] are well-known photoactive compounds (Wang et al., 2012; Weller et al., 2013, 2014). They can easily get excited by light in the UV–VIS range, inducing ligand-to-metal charge transfer (LMCT) (Cie´sla et al., 2004), which is an inner-sphere electron transfer (i.e., the electron transfer occurs via a covalently bound bridging ligand) from the carboxylate group to the iron. Investigations using time-resolved transient spectroscopy reported the formation of long-lived radical complexes, [Fe$^{II}$ (OOC–R)$_2$$^{2+}$], with lifetimes on the order of a millisecond, followed by the dissociation to the organic radical R–COO$^-$ and an Fe$^{II}$ aqua complex (Feng et al., 2007; Glebov et al., 2011; Pozdnyakov et al., 2009; Zhang et al., 2009):

\[
[\text{Fe}^{III}(\text{OOC}–\text{R})]^{2+} + h\nu \rightarrow [\text{Fe}^{III}(\text{OOC}–\text{R})]^{2+*} \quad (R1)
\]

\[
[\text{Fe}^{III}(\text{OOC}–\text{R})]^{2+*} \rightarrow [\text{Fe}^{II}(\text{OOC}–\text{R})]^{2+} \quad (R2)
\]

\[
[\text{Fe}^{II}(\text{OOC}–\text{R})]^{2+} \rightarrow \text{Fe}^{2+} + \text{R}–\text{COO}^- \quad (R3)
\]

\[
\text{R}–\text{COO}^- \text{ will decarboxylate almost instantaneously (} k_{R3} \approx 10^9–10^{12} \text{ s}^{-1} \text{) (Abel et al., 2003; Bockman et al., 1997; Hilborn and Pincock, 1991):}
\]

\[
\text{R}–\text{COO}^- \rightarrow \text{R} + \text{CO}_2.
\]

The alkyl radical R$^-$ will react rapidly with dissolved O$_2$, producing a peroxy radical with $k_{R4} \approx 2 \times 10^9 \text{ M}^{-1} \text{s}^{-1}$ (von Sonntag and Schuchmann, 1991):

\[
\text{R}^- + \text{O}_2 \rightarrow \text{RO}_2.
\]
Subsequent reactions of R and RO_2 are specific depending on the type of ligand and its substitution.

In this work we investigated iron(III) citrate ([Fe^{III}(OOCCH_2)_2C(OH)(COO)], in short Fe^{III}(Cit)), as a model species to better understand iron carboxylate photochemistry in atmospheric aerosol particles. Our Fe^{III}(Cit) system undergoes LMCT reactions in the same way as countless other iron(III) carboxylate compounds (Ciesla et al., 2004; Weller et al., 2013, 2014). Its photochemical reaction scheme is well established in both solution (Abida et al., 2012; Faust and Zepp, 1993; Pozdnyakov et al., 2012) and solid states (Abrahamson et al., 1994). Citric acid (CA) is an established proxy for oxygenated atmospheric organic matter with its thermodynamic properties, water diffusivity, and viscosity having been well studied (Lienhard et al., 2012, 2014; Song et al., 2016). For these reasons, it is a valid and reliable proxy for atmospheric iron carboxylate photochemical processes.

As schematically described in Fig. 1, Fe^{III}(Cit) absorbs light up to 500 nm, inducing LMCT, followed by immediate decarboxylation of the central carboxyl group since the hydroxyl group adjacent to a carboxyl group facilitates decarboxylation (Weller et al., 2013). In the presence of O_2, oxidants such as HO_2 and H_2O_2 will be produced, which can oxidize Fe^{II} back to Fe^{III} via Fenton reactions (Fenton, 1894) with additional oxidant production. Fe^{III} then combines with another CA molecule in this aqueous system, closing the photocatalytic cycle, in which iron acts as a catalyst for CA degradation. In addition, the generation of reactive oxygen species (ROS) and peroxo radicals leads to further decarboxylation and more production of oxygenated volatile organic compounds (OVOCs) (e.g., acetone) (Pozdnyakov et al., 2008; Wang et al., 2012). Therefore, this photodegradation process is potentially an important sink of carboxylate groups in the troposphere.

We expect that the chemistry subsequent to the initial photochemical reaction steps in the aerosol phase will be significantly altered by diffusion limitations when Fe^{III}(Cit) particles mixed with CA attain a high viscosity. As viscosity increases, molecular diffusion coefficients tend to decrease (Koop et al., 2011) and therefore photochemical cycling will also be slow. Increasing water content is expected when the relative humidity (RH) increases and will effectively plasticize particles (Koop et al., 2011) leading to better mixing of all compounds within the particles and faster photochemical cycling when compared with lower RH. However, these effects have been investigated in dark systems but not in photochemical systems (Berkemeier et al., 2016; Shiraiwa et al., 2011; Shiraiwa and Seinfeld, 2012; Steimer et al., 2015a). In order to better understand this system and how it reacts to RH, we used a triad of photochemical experiments including electrodynamic balance (EDB) measurements, scanning transmission X-ray microscopy coupled with near-edge X-ray absorption fine structure (STXM/NEXAFS) spectroscopy, and a coated-wall flow-tube (CWFT) experiments to investigate how particle size, mass, and indicators of chemical composition change during photochemical processes. In this work we mostly focus on the humidity dependence of this photochemical degradation, while Alpert et al. (2021) focus on the impacts on ROS and the fate of free radicals during this photochemical degradation.

To perform a quantitative comparison of these experiments and determine their relevant properties, a numerical multi-layered photochemical reaction and diffusion (PRAD) model was developed that treats chemical reactions and the transport of various species. In addition, we will use the PRAD model to simulate photochemical aging processes under atmospheric conditions. In the following we briefly discuss our experimental approaches in Sect. 2 and include a detailed explanation of the PRAD model in Sect. 2.5. A comparison between the experimental results and the PRAD model is presented in Sect. 3. Finally, we discuss the impact and atmospheric importance of kinetic limitations to photochemical degradation in Sect. 4.

2 Methods

2.1 Solution preparation

CA (≥ 99.5 %) and iron(III) citrate tribasic monohydrate (18 %–20 % Fe basis) were purchased from Sigma-Aldrich. Iron(II) citrate (Fe^{II}(HCit)) was purchased from Dr. Paul Lohmann GmbH KG. Dilute aqueous solutions of Fe^{III}(Cit)/CA and Fe^{II}(HCit)/CA were made in ultrapure water (18 MΩ cm⁻¹, Milli-Q). Since Fe^{III}(Cit) only
dissolves slowly in water, CA solutions with suspended Fe$^{III}$\((\text{Cit})\) crystals were sonicated for at least 24 h. The same dissolving procedure was also applied to the Fe$^{II}$\((\text{HCit})\) powders. Note that all procedures were done under red light illumination because Fe$^{III}$\((\text{Cit})\) is light sensitive. The molar ratio between Fe$^{III}$\((\text{Cit})\) and CA was different for each experimental method used in this study. For EDB, STXM/NEXAFS, and CWFT experiments, stock solutions were prepared with molar ratios of 0.05, 1.0, and 0.07, respectively.

### 2.2 Bulk property measurements by EDB

We used an EDB to measure the mass loss in single, levitated particles under irradiation. The experimental setup has been described previously (Steimer et al., 2015b). In short, an electrically charged aqueous particle (radius $\sim 10 \mu$m) is injected into an EDB. The balance is of the double ring design (Davis et al., 1990) with a high AC voltage applied to the two parallel electrode rings and a DC voltage across the hyperbolic endcaps. The DC field compensates the gravitational force of the particle and is used as a measure for the mass of the particle. The EDB is placed in a three-wall glass chamber with a cooling liquid (ethanol) pumped through the two inner walls and an insulation vacuum between two outer walls to control the temperature ($T$) at the location where the particle levitates. The RH within the chamber is regulated by adjusting the ratio of a dry and a humidified gas flow through the chamber. In the experiments described in this work, we used a typical total flow of 40 sccm and set the total pressure inside the cell at $8 \times 10^4$ Pa.

The spherical particles were characterized by two methods based on Mie-resonance spectroscopy: (i) a narrow band-width tunable diode laser (TDL, tuning range 765–781 nm) was used to determine the refractive index and radius simultaneously with high-precision (Steimer et al., 2015b); and (ii) simultaneously, a broad-band LED centered around 640 nm was used to illuminate the particle. The backscatter signal from the LED was recorded using a spectrograph with a slow-scan back-illuminated charge-coupled device (CCD) array detector to follow the resonance wavelength shift of the particle (Zardini et al., 2006). If we assume the refractive index stays constant during the experiment, the radius change of a particle is easily calculated from this resonance wavelength shift:

$$\frac{r(t)}{r_0} = \frac{\lambda_0}{\lambda_0 + \Delta\lambda(t)} = 1 + \frac{\Delta\lambda(t)}{\lambda_0}. \tag{1}$$

We illuminated particles to induce a photochemical reaction with either a CW diode laser emitting at 375 nm (LuxX 375-20, Omicron Laserage) or a frequency-doubled diode laser emitting at 473 nm (gem 473, Laser Quantum). At wavelengths of 375 and 473 nm, Fe$^{III}$\((\text{Cit})\) is reported to have a molar absorptivity of 796 and 60.7 M$^{-1}$ cm$^{-1}$, respectively (Pozdnyakov et al., 2008).

In a typical EDB experiment, we let the particle equilibrate to RH and $T$ in a pure O$_2$ gas phase for up to 10 h in the dark before irradiation. Exemplary raw data of an experiment at 46% RH and 293.5 K are shown in Fig. 2. The measured DC voltage compensating for the gravitational force, as well as the radius of the particle deduced from low-resolution Mie-resonance spectroscopy, decreased dramatically during illumination in the first 18 h, with more than half of the initial mass lost to the gas phase. Note that the radius and mass loss rates increased as seen in Fig. 2c. We assumed the refractive index and density of the particle did not change upon photochemistry and therefore the mass loss calculated from the DC voltage could be directly compared with size change by calculating the par-
particle mass remaining ratio,

$$\frac{m(t)}{m_0} = \left( \frac{r(t)}{r_0} \right)^3,$$

where $m_0$ is the particle mass prior to irradiation. Mass loss derived from both ways independently is shown in Fig. 2d and reveal that there is a little difference between the mass loss up to $t \approx 65 000\,s$, corresponding to when $\frac{m(t)}{m_0} \approx 0.4$. Therefore, the refractive index and density are mostly governed by those of aqueous CA up until half the particle mass is lost. The total mass loss over 24 h irradiation is more significant and drops by 80 % for the particular experiment shown in Fig. 2. In addition, we observed the mass loss rate was initially $\sim 1.3\,\%\,h^{-1}$ and increased to $\sim 14\,\%\,h^{-1}$ when 40 % to 60 % of the initial mass was lost. This mass loss acceleration is discussed in further detail with the help of the PRAD model simulations in the Sect. 3.1. At $t \approx 65 000\,s$, the mass loss slowed down considerably when we observed a distortion in the Mie-resonance pattern (Fig. 2b and video in the Supplement). The distortion may be attributable to partial crystallization of iron citrate in the particle, which would explain the slowing photochemical degradation.

### 2.3 Chemical characterization by STXM/NEXAFS

STXM/NEXAFS measurements were performed at the PolLux endstation located at the Swiss Light Source (SLS) to obtain the Fe oxidation state of particles between 0.2 and 2 µm in diameter (Flechsig et al., 2007; Frommherz et al., 2010; Raabe et al., 2008). Particles containing Fe(III)(Cit)/CA were nebulized from aqueous solution with a mole ratio between Fe(III)(Cit):CA of 1:1. They were dried in air at RH < 30 % and impacted onto silicon nitride membranes mounted in portable sample holders. The sample holders were transported to the endstation in an evacuated container and shielded against ambient light. Once there, they were mounted in the PolLux environmental microreactor (Huthwelker et al., 2010) and kept under a total pressure of 1.5 × 10^4 Pa, $T = 293.5\,K$, RH = 40, 50 or 60 %, with a controlled gas flow. Further details of sample preparation have been previously reported (Alpert et al., 2019; Huthwelker et al., 2010; Steimer et al., 2014). The microreactor was mounted in a vacuum chamber for in situ STXM/NEXAFS analysis. When desired, the microreactor could operate in vacuum conditions without a gas flow. The transmission of X-ray photons through the particles was measured and converted to optical density, $OD = -\ln(I/I_0)$, where $I$ and $I_0$ are the transmitted and incident photon flux as a function of X-ray energy. The Fe L-edge absorption was probed over the X-ray energy range of 700–735 eV. Figure 3 shows an example of NEXAFS spectra of Fe(III)(Cit)/CA particles before (orange) and after (red) irradiation with UV light. X-ray energy calibration was consistently performed using FeCl$_2$ and compared with the existing literature for FeCl$_2$ and FeCl$_3$ salts (Moffet et al., 2012) and a mixture of xanthan gum and FeCl$_2$ oxidized by O$_3$ (Alpert et al., 2019). The peak absorption for iron(II) and iron(III) are at X-ray energies of 708.3 and 710.0 eV. We were capable of resolving peaks separated by 0.4 eV at the Fe L-edge. Ferrous and ferric iron peaks are separated by 1.7 eV and thus clearly distinguishable. Following a previous procedure (Alpert et al., 2019), we imaged particles at these two energies to determine the OD ratio between them. The fraction of Fe(III) out of total Fe, $\beta$, was then determined using the parameterization from Moffet et al. (2012). It is important to note that the X-ray energy absorption peaks observed for FeCl$_2$ and FeCl$_3$ were identical for FeCl$_2$ mixed with xanthan gum either unexposed or exposed to O$_3$ (Alpert et al., 2019). However, we have found that these peaks shifted by about +0.4 eV, possibly due to the strong complexation with CA. Small shifts in energy can occur depending on the chemical environment surrounding Fe atoms (Garvie et al., 1994; Moffet et al., 2012). In agreement with Alpert et al. (2019), the peak absorption energies for our particles were independent of RH from 0 % to 60 % within ±0.2 eV. When calculating $\beta$, we always imaged particles at 708.3 and 710.0 eV.
2.4 **HO$_2$** production determined by CWFT

The HO$_2$ release upon irradiation of a Fe$^{III}$(Cit)/CA thin film was measured by scavenging HO$_2$ with an excess of NO in a CWFT reactor (Duran glass, inside diameter 1.2 cm, length 50 cm). The film was composed of Fe$^{III}$(Cit)/CA, and deposited inside the tubular glass flow tube with a thickness between 0.15 and 0.2 $\mu$m and an error of about 20%. Details of the film preparation have been described previously (Corral Arroyo et al., 2018; González Palacios et al., 2016). Seven UV lamps (UV-A range, Philips Cleo Effect) were mounted surrounding the glass reactor held at 298.15 K. The total light output between 300 and 590 nm was 210 W m$^{-2}$. The flows of N$_2$, O$_2$, and NO passing through the reactor were controlled. The NO concentration during CWFT photochemical experiments was in excess ($>10^{13}$ molecules cm$^{-3}$) to efficiently scavenge 99% of HO$_2$ produced by the film. The concentration of NO was tracked by a chemiluminescence detector (Ecophysics CLD 77 AM). In an example CWFT experiment at RH = 29.3%, a clear NO loss was observed when the UV lights were switched on as shown in Fig. 4, which was due to the release of HO$_2$ radicals into the gas phase and reaction of NO with HO$_2$ forming NO$_2$ and OH: OH is then scavenged by NO producing HONO. The production of HONO was routinely checked as described in González Palacios et al. (2016). Therefore, the production rate of HO$_2$, $P_{HO_2}$, was calculated from the loss rate of NO assuming a 2:1 ratio to HO$_2$ conversion:

$$P_{HO_2} = \frac{[NO] \times \text{flow}}{2S_{\text{film}}}. \quad (3)$$

where [NO] is the loss of gas-phase concentration of NO in molecules cm$^{-3}$, flow is the volumetric gas flow in the CWFT in cm$^3$s$^{-1}$, and $S_{\text{film}}$ is the surface area of the film in cm$^2$.

2.5 **Development of the photochemical reaction and diffusion (PRAD) model**

We developed a PRAD model to interpret our experiments and to understand any feedback between transport limitations and photochemistry, especially under low RH conditions, corresponding to high viscosity of the particle phase. The PRAD model consists of two modules: a detailed chemical process module, treating equilibria and chemical reactions, and a transport module handling the physical transport of all species (including diffusion in the aqueous phase as well as gas–particle phase partitioning). Conceptually, the PRAD model relies on the kinetic model framework for aerosol surface chemistry and gas–particle interactions (Pöschl et al., 2007), which is similar, for example, to the KM-GAP model (Shiraiwa et al., 2012). Numerically, the PRAD model uses a Euler forward step method as explained in detail below, while KM-GAP solves coupled differential equations. In passing, there are alternative approaches; for example, Kinetoscope (Houle et al., 2015) does not integrate sets of coupled differential equations to predict the time history of a chemical system. Instead, it uses a general stochastic algorithm to propagate a reaction.

As illustrated in Fig. 5, the PRAD model divides a spherical droplet into a number of shells, $n$, which exchange molecules after each chemical time step. Shell thickness and the number of shells were adjusted to enable the resolution of steep concentration gradients within a reasonable computation time. The volume of each shell was constant instead of the thickness and the shells become thinner and thinner from the center to the surface of the particle. For each shell and at each time step, we first calculated the composition using the thermodynamic equilibria of the Fe$^{III}$(Cit)/CA system, as listed in Table 2. The Newton–Raphson method (Burden and Faires, 2011) was then used to calculate the turnover and the concentration of products and reactants over time for the chemical reactions also listed in Table 2 with a fixed time step of 0.2 s. After each time step, diffusion of all species between the shells and the evaporation of products (or condensation of the reactant O$_2$) were computed. The time step, $\Delta t$, for physical transport processes was determined dynamically to ensure both numerical stability and computational efficiency.

For each species, the molar flux from shell $i$ to the next shell $i+1$ was calculated as

$$f_i = -4\pi r_i^2 D_i \frac{dc}{dr} \bigg|_{r=r_i} = -4\pi r_i^2 D_i \frac{c_{i+1} - c_i}{0.5(r_{i+1} - r_{i-1})}; \quad \forall i \in \{1, 2, \ldots, n-1\},$$

where $D_i$ is the liquid phase diffusion coefficient of the corresponding species. Shell $i$ extends from $r_{i-1}$ to $r_i$, while shell $i + 1$ extends from $r_i$ to $r_{i+1}$, with $r$ being the distance from the particle center. In Eq. (4), $c$ is the molar concentration in
each shell of the aqueous particle, defined as

\[ c_i = \frac{N_i}{V_i}; \; \forall i \in \{1, 2, \ldots, n\}, \]  

(5)

where \( N_i \) is moles of a particular species in shell \( i \), and \( V_i \) is the total volume of shell \( i \).

At the outermost shell \( n \), the gas–particle phase partitioning of each species was determined by the modified Raoult’s law. The flux from shell \( n \) into the gas phase, \( f_n \), was calculated to be

\[ f_n = -4 \pi r_n^2 D_g \frac{dc}{dr} \bigg|_{r=r_n} = -4 \pi r_n^2 D_g \frac{c_n - c_n^g}{r_n} \]

\[ = -4 \pi r_n D_g \frac{P_{\text{partial}} - P_{\text{vapor}}}{RT}, \]  

(6)

where \( D_g \) is the gas phase diffusion coefficient of the corresponding species, and \( R \) is the gas constant. \( P_{\text{partial}} \) is the partial pressure of the species in the gas phase at infinite distance from the particle. Based on Henry’s law, vapor pressure of the species \( P_{\text{vapor}} \) is defined as

\[ P_{\text{vapor}} = \frac{c_n}{H}, \]  

(7)

where \( c_n \) is the molar concentration in the outermost shell \( n \), and \( H \) is the Henry’s law coefficient. Substituting Eq. (7) into Eq. (6) leads to

\[ f_n = -4 \pi r_n D_g \left( \frac{P_{\text{partial}}}{RT} - \frac{c_n}{HR RT} \right). \]  

(8)

We calculated the partial pressure of H\(_2\)O from RH, and took the partial pressure of O\(_2\) based on the total pressure and the fraction of O\(_2\) gas flow used in experiments. For other species, \( P_{\text{partial}} \) is negligible and assumed to be zero. So, for all species other than H\(_2\)O and O\(_2\), \( f_n \) simplifies to

\[ f_n = -4 \pi r_n D_g \left( 0 - \frac{c_n}{HR RT} \right) = 4 \pi r_n D_g \frac{c_n}{HR RT}. \]  

(9)

Based on the calculated fluxes, the change in moles, \( \Delta N_i \), of each species in \( \Delta t \) was given by

\[ \Delta N_i = (f_{i-1} - f_i) \Delta t; \; \forall i \in \{1, 2, \ldots, n\}, \]  

(10)

from which the concentration and corresponding shell thickness using the molar volume of each species were recalculated for the next time step.

The chemical loss rate of O\(_2\) was treated in the liquid phase diffusion module instead of in the chemical module, since the loss can be very fast and its lifetime can be smaller than 10\(^{-6}\) s. If O\(_2\) loss due to reaction was determined in the chemical module, a chemical time step of 10\(^{-6}\) s would result in extensive computational time beyond what is reasonable for this study. Therefore, within each liquid phase diffusion time step, in addition to the physical transport, the chemical loss of O\(_2\) molecules was calculated in each shell:

\[ \frac{dN_i}{dt} = -kO_2 N_i; \; \forall i \in \{1, 2, \ldots, n\}. \]  

(11)

Integration yields

\[ N_i = N_i(e^{-kO_2 \Delta t}); \; \forall i \in \{1, 2, \ldots, n-1\}, \]  

(12)

where \( kO_2 \) is the total chemical reaction rate of O\(_2\). In the outermost shell \( n \), O\(_2\) molar concentration is assumed to be at steady state all times, meaning that the chemical loss of O\(_2\) is always compensated by the condensation of O\(_2\) from RH, and took the chemical module, a chemical time step of 10\(^{-6}\) s would result in extensive computational time beyond what is reasonable for this study. Therefore, within each liquid phase diffusion time step, in addition to the physical transport, the chemical loss of O\(_2\) molecules was calculated in each shell:

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(11)

Integration yields

\[ N_i = N_i(e^{-kO_2 \Delta t}); \; \forall i \in \{1, 2, \ldots, n-1\}, \]  

(12)
We have parameterized aqueous and gas phase diffusion coefficients, $D^g_t$ and $D^g_q$, respectively, for all species $j$ given in Appendix A1 and A2. There are several other assumptions and approximations made to adapt this model to the aqueous Fe$^{III}$ (Cit) / CA system:

1. We always set water activity in the particle phase in equilibrium with the gas phase, since the RH of the gas phase did not change during each experiment.

2. The bulk accommodation coefficients of all species were assumed to be 1.

3. Neglecting the influence of Fe$^{III}$ (Cit), the water activity was taken from that of CA ($\alpha_w^{CA}$), which was determined from the mass fraction of CA, $W_{CA}$, by Lienhard et al. (2012):

$$\alpha_w^{CA} = \frac{1 - W_{CA}}{1 + q \cdot W_{CA} + r \cdot W_{CA}^2},$$  \hspace{1cm} (19)

with

$$q = -3.16761 + 0.019397 T - 4.02725 \times 10^{-5} T^2, \hspace{1cm} (20)$$

and

$$r = 6.59108 - 0.05294 T + 1.06028 \times 10^{-4} T^2. \hspace{1cm} (21)$$

The water activity of citrate ($\alpha_w^{Cit}$) was calculated using the same equation

$$\alpha_w^{Cit} = \frac{1 - W_{Cit}}{1 + q \cdot W_{Cit} + r \cdot W_{Cit}^2},$$  \hspace{1cm} (22)

where the mass fraction of citrate $W_{Cit}$ was treated in two fractions: citrate without Fe and citrate containing Fe. The water activity of an aqueous 1 M Fe$^{III}$ (Cit) solution was determined using a water activity meter (AquaLab water, Model 3B, Decagon Devices, USA) at room temperature. We found this water activity corresponds to that of a 0.81 M aqueous CA solution. Therefore, for calculating water activity, all iron-containing citrate complexes (either Fe$^{II}$ or Fe$^{III}$) were treated with a reduced concentration: namely with a factor of 0.81 of the corresponding citrate molarity. Hence, the overall amount of citrate was calculated as

$$N_{Cit}^0 = N_{Cit} + 0.81 N_{FeCit},$$  \hspace{1cm} (23)

accordingly and $W_{Cit}$ was determined using

$$W_{Cit} = \frac{M_{Cit} \cdot N_{Cit}^0}{M_{Cit} \cdot N_{Cit}^0 + M_{H_2O} \cdot N_{H_2O}},$$  \hspace{1cm} (24)

where $M_{Cit}$ and $M_{H_2O}$ are the molar mass of CA and water, respectively. For other species $j$, the contribution to the water activity is proportional to their molar volume ($MV_j$), so that in total,

$$\alpha_w = \alpha_w^{CA} \times \alpha_w^{Cit} \times \frac{MV_{H_2O}}{MV_{H_2O} + \sum_j MV_j}. \hspace{1cm} (25)$$

4. Fe$^{III}$ (Cit) photolysis, decarboxylation, and oxidation of the alcohol group in presence of O$_2$ yields the compounds O=CH$_2$(COOH)$_2$ or O=CH$_2$(COOH)$_2$, which are C$_5$ species. We assumed that half of the C$_5$ species undergo photochemical reactions to produce CO$_2$ and compounds with 2–4 carbon atoms, C$_4$, C$_3$ and C$_2$ (see Reactions R10–R14 shown in Table 2), all of which are capable of being released to the gas phase depending on their solubility.

5. We estimated the quantum yield in Reactions (R1) and (R2) in Table 2, as $\Phi = 1.0$ at $\lambda = 375$ nm and $\Phi = 0.002$ at $\lambda = 473$ nm (Dou et al., 2019), and we parameterized $\Phi$ as a function of wavelength, $\lambda$:

$$\Phi = \frac{e^{-0.145(\lambda - 430)}}{1 + e^{-0.145(\lambda - 430)}}. \hspace{1cm} (26)$$

In total, the PRAD model includes 13 equilibria and 17 chemical reactions among 32 species, as well as their condensed phase diffusivities and Henry’s law coefficients. Some of these parameters are known from previous studies (see Tables 1 and 2 for references), while others are not known and difficult to estimate. For instance, even though absorption spectra of Fe$^{III}$ (Cit) have been measured in aqueous solution (Pozdnyakov et al., 2012), the corresponding quantum yield has not, which leaves the photolysis rate of Fe$^{III}$ (Cit), $j$, unknown. Also, there are no data reported on the diffusivity of O$_2$ in aqueous CA solutions, and the chemical reaction rate of the oxidation of the Fe$^{III}$ citrate complex by O$_2$ is quite uncertain (Gonzalez et al., 2017). In order to find the optimal parameter set, we compared the experimental data of the three setups taken under well-controlled conditions with model predictions and tuned the unknown parameters manually.

We restricted our tuning of the parameters to reach satisfactory agreement with all experimental data simultaneously. The equilibrium constants and rate coefficients that were tuned are indicated in Table 2 (the sensitivity of the PRAD model results to a few of its parameters is shown in Appendix A5). The parameters were adjusted in a wide and acceptable range until a good representation of our data could be obtained. For example, the fraction of iron(III) in a photoactive complex (equilibrium E5 in Table 2) must have been high enough to reproduce STXM/NEXAFS observations that iron could be reduced to low levels as seen in Fig. 7 described below. In comparison, equilibrium constant of E7 must have been much lower than equilibrium constant of E5 so that the amount of iron(III) in a nonphotoactive complex was small compared to being in complex with citrate. As another example, oxidation of Fe$^{2+}$ (Reactions R5–R8 in Table 2) is fairly well referenced, and therefore we adjusted the rate of Reaction (R9) until the model reoxidation rates matched those observed. Tuning of individual bulk diffusion coefficients for all species was not attempted. Instead, we
Table 1. Liquid phase diffusivity factors (normalized to water) and Henry’s law coefficients (Sander, 2015) of major species in the Fe\(^{III}\)(Cit) photochemistry system.

| Number | Name                  | Formula                                      | \(l_j^a\) | \(H_0\) (M atm\(^{-1}\))\(^b\) | \(Q\)\(^b\) |
|--------|-----------------------|----------------------------------------------|-----------|---------------------------------|------------|
| 1      | water                 | H\(_2\)O                                      | 1         | 1                               | 1          |
| 2      | cit total             | –                                             | –         | –                               | –          |
| 3      | ferric (Fe\(^{III}\)) total | –                                           | –         | –                               | –          |
| 4      | ferrous (Fe\(^{II}\)) total | –                                           | –         | –                               | –          |
| 5      | citric acid (CA)      | (CH\(_2\)COOH\(_2\))\(_2\)C(OH)(COOH)/H\(_3\)Cit | 1.20 \times 10\(^{-6}\) | infinite | 10 000 |
| 6      | dihydrogen citrate    | (CH\(_2\)COOH\(_2\))\(_2\)C(OH)(COOH)/H\(_2\)Cit\(^-\) | 1.20 \times 10\(^{-6}\) | infinite | 10 000 |
| 7      | hydrogen citrate      | (CH\(_2\)COOH\(_2\))C(OH)(CH\(_2\)COO)(COO\(_2\))\(^-\)/HCit\(^2\)\(^-\) | 1.20 \times 10\(^{-6}\) | infinite | 10 000 |
| 8      | citrate               | C(OH)(CH\(_2\)COO\(_2\))\(_2\)(COO\(_3\))\(^-\)/Cit\(^3\)\(^-\) | 1.20 \times 10\(^{-6}\) | infinite | 10 000 |
| 9      | Fe\(^{III}\)(Cit)(OH\(_-)\)  | 3.92 \times 10\(^{-7}\) | infinite | 10 000 |
| 10     | Fe\(^{III}\)(HCit\(_+)\)  | 5.04 \times 10\(^{-7}\) | infinite | 10 000 |
| 11     | ferrous citrate       | Fe\(^{II}\)(HCit)                          | 5.04 \times 10\(^{-7}\) | infinite | 10 000 |
| 12     | ferric citrate        | Fe\(^{III}\)(Cit)                          | 5.04 \times 10\(^{-7}\) | infinite | 10 000 |
| 13     | ferric ion            | Fe\(^{3+}\)                                 | 3.78 \times 10\(^{-5}\) | infinite | 10 000 |
| 14     | Fe\(^{III}\)(OH\(_2\))\(^2+\) | 1.18 \times 10\(^{-5}\) | infinite | 10 000 |
| 15     | ferrous ion           | Fe\(^{2+}\)                                 | 3.78 \times 10\(^{-5}\) | infinite | 10 000 |
| 16     | hydrogen ion          | H\(^+\)                                     | –         | –                               | –          |
| 17     | hydroxide ion         | OH\(^-\)                                    | –         | –                               | –          |
| 18     | hydroperoxy radical   | HO\(_2\)                                    | 1.13 \times 10\(^{-4}\) | 4 \times 10\(^4\) | 5900  |
| 19     | superoxide radical    | O\(_2\)\(^-\)                               | –         | –                               | –          |
| 20     | radicals              | OH\(_-\)C(CH\(_2\)COO\(_2\))\(_2\)+OH\(_-\)C(CH\(_2\)COOH\(_2\)) | 2.69 \times 10\(^{-6}\) | infinite | 10 000 |
| 21     | hydroxyl radical      | OH\(_-)\)                                   | –         | –                               | –          |
| 22     | hydrogen peroxide     | H\(_2\)O                                    | 1.07 \times 10\(^{-4}\) | 8.3 \times 10\(^5\) | 7400  |
| 23     | oxygen                | O\(_2\)                                     | 1.20 \times 10\(^{-4}\) | 3.5 \times 10\(^{-2}\) | 1500  |
| 24     | carbon dioxide        | CO\(_2\)                                    | depends on \(a_w\) and \(T\) | 3.4 \times 10\(^{-1}\) | 2400  |
| 25     | acetone               | CH\(_3\)COCH\(_3\)                         | 1.18 \times 10\(^{-5}\) | 30                             | 4600  |
| 26     | unkn. prod C\(_4\)   | C\(_4\)                                      | 5.30 \times 10\(^{-6}\) | 1 \times 10\(^5\) | 6000  |
| 27     | O=C(CH\(_2\)COO\(_2\))\(_2\)+O=C(CH\(_2\)COOH\(_2\)) | 1.95 \times 10\(^{-6}\) | infinite | 8000  |
| 28     | acetic acid           | CH\(_3\)COOH                                | 3.23 \times 10\(^{-5}\) | 4.1 \times 10\(^3\) | 6300  |
| 29     | Fe\(^{III}\)([O=C(CH\(_2\)COO\(_2\))\(_2\)]   | 7.22 \times 10\(^{-7}\) | infinite | 10 000  |
| 30     | unkn. prod C\(_5\)   | C\(_5\)                                      | 2.69 \times 10\(^{-6}\) | 1 \times 10\(^7\) | 8000  |
| 31     | unkn. prod C\(_5\)   | C\(_5\)\(_-\)stable                        | 2.69 \times 10\(^{-6}\) | 5 \times 10\(^8\) | 8000  |

\(^a\) \(l_j^a\) is a factor of the diffusion coefficient of each species \(j\) normalized to that of water. \(^b\) Henry’s law is described as a function of temperature \(T\):

\[
H = H_0e^{Q/T}
\]
simplified the representation of diffusion coefficients using a parameterization as a function of molar mass, as described in Appendix A1. The two constants in Eq. (A8) and the two constants in Eq. (A3) were tuned, resulting in the absolute diffusion coefficients shown in Fig. A1. Henry’s law coefficients for gasses were tuned, however purposefully set at values higher than expected for pure water or highly dilute aqueous solution. This was inspired by previous studies regularly reporting solubility of, e.g., O₂ and CO₂ higher in a variety of organic liquids than water (Fogg, 1992; Battino et al., 1983). It is important to note that the result of this tuning does not mean that we found the global minimum in the parameter space (see, e.g., (Berkemeier et al., 2017)). A thorough search for a global minimum for our model with 16 tuning parameters for chemistry, four tuning parameters (and our parameterization) for diffusion, and nine tuning parameters for solubility is computationally very expensive and beyond the scope of this paper. However, for our purpose here, namely modeling typical timescales of photochemical degradation of organic aerosol under atmospheric conditions (see Sect. 3.5), the PRAD model framework should allow sufficiently accurate predictions. In other words, we expect similar mass degradation in atmospheric particles due to the fact that many other relevant iron carboxylate compounds undergo LMCT similarly as to our model system (Weller et al., 2013, 2014). Additionally, if a particular system requires parameter values that significantly differ from ours, the PRAD model framework itself should still be valid. Note that careful evaluation is needed when picking a single parameter of the PRAD model for use in another context. Comparisons of the refined model with our experimental data are shown in the next section.

3 Comparisons between experimental measurements and model simulations

3.1 The effect of RH on photocatalytic degradation efficiency

We performed experiments with single, levitated particles under continuous UV irradiation (375 nm) in pure O₂ at different RH to assess the effects of RH on the photocatalytic cycle shown in Fig. 1 and tested the model performance under these conditions. Qualitatively, a continuous decrease of particle mass and size is expected to occur due to evaporation of volatile products, as shown in Fig. 2. Figure 6 shows the fraction of particle mass remaining with the irradiation time at three different RH calculated from resonance wavelength shifts (Eqs. 1 and 2). Clearly, particle mass was lost to the gas phase with time due to the evaporation of photochemical products and, similarly to Fig. 2, all data show a very significant acceleration of mass loss with time. After tuning some of the parameters of the model as further discussed below, the PRAD model simulations reproduces our data with a very similar trend and magnitude over all, which gives us confidence that the PRAD model captures the essential chemistry and transport during irradiation. (How particle mass evolves subsequently until 80 % mass loss in both experiments and models is shown in Fig. A4.) However, the model is not able to capture the full degree of acceleration of the degradation rate, as it does not attempt to include the complete multigenerational oxidation chemistry at the level of individual components after initial radical production.

The degradation processes were faster at higher RH. At lower RH, the particle was expected to be more viscous, diffusion coefficients were expected to be lower, products were generated at a lower rate, and volatile products moved more slowly to the surface to evaporate. More importantly, O₂ taken up by the particle from the gas phase diffused more slowly into the bulk of the particle at lower RH. Thus, less HO₂ and H₂O₂ formed and less Fe²⁺ could be reoxidized from the surface to the center of the particle. The observed gradient in the Fe³⁺ fraction, β, and the modeled gradients in O₂ and ROS in the particle have been shown with radial profiles in Alpert et al. (2021). This resulted in fewer photochemically active Fe³⁺ complexes available for photocatalytic degradation. The characteristic degradation time shortened by a factor of 5.5 when RH increased from 46 % and 61 %, which demonstrates that photochemical cycling is highly sensitive to the microphysical conditions. The diffusivity of O₂ must have significantly impacted reoxidation rates. In addition, the diffusion coefficients of both Fe²⁺ and Fe³⁺ species increases with RH. Therefore, the molecular transport between both iron and oxygen reactants increases causing a highly nonlinear trend in increasing mass loss with increasing RH.

3.2 Determination of iron(III) reduction rate and iron(II) reoxidation rate by STXM/NEXAFS

In STXM/NEXAFS experiments, the freshly prepared Fe³⁺(Cit) mixed with CA at x = 1.0 particles were irradiated to determine the Fe³⁺(Cit) photolysis rate, as shown in Fig. 7. Each experimental data point is the average Fe³⁺ fraction from 16–36 individual particles. Fitting an exponential function, β = β₀e⁻ᵏᵇ consortium, yields β₀ = 0.93 ± 0.09 and a first-order decay rate of 𝑗_𝑜𝑏𝑠 = 0.08 ± 0.01 s⁻¹. The LED power at the sample was measured to be 5.9 ± 0.6 mW in total and had a Gaussian spectral profile between 361 and 374 nm at full width half maximum. When mounting the UV fiber optics and collimator lens for multiple samples, the illuminated area had a circle equivalent diameter of 5±1.5 mm. Using the absorption cross section calculated from the molar attenuation coefficient (Pozdnyakov et al., 2008), Φ = 1.0 and propagating all uncertainties yields a photochemical reaction rate of 𝑗_𝑎𝑙𝑐 = 0.20 ± 0.12 s⁻¹, which is in agreement with 𝑗_𝑜𝑏𝑠. This implies that assuming a quantum yield of 1 at these UV wavelengths is reasonable. This calculated value for the photochemical reaction rate was used in the PRAD model for
Table 2. Compilation of equilibria, chemical reactions, and corresponding rate constants in Fe$^{III}$ (Cit) photochemistry system.

| Number | Reactions | $K_{eq}/k_r/\sigma$ | Sources |
|--------|-----------|---------------------|---------|
| E1     | $H_2O \rightleftharpoons OH^- + H^+$ | $1 \times 10^{-14}$ M | |
| E2     | $H_2Cit \rightleftharpoons H_2Cit^- + H^+$ | $7.5 \times 10^{-3}$ M | Martell and Smith (1982) |
| E3     | $H_2Cit^- \rightleftharpoons HCit^2^- + H^+$ | $1.7 \times 10^{-5}$ M | Martell and Smith (1982) |
| E4     | $HCit^2^- \rightleftharpoons Cit^{3-} + H^+$ | $4.0 \times 10^{-7}$ M | Martell and Smith (1982) |
| E5     | $Fe^{3+} + Cit^{3-} \rightleftharpoons Fe^{III}(Cit)$ | $1.58 \times 10^{13}$ M$^{-1}$ | tuning parameter |
| E6     | $Fe^{3+} + Cit^3- + H_2O \rightleftharpoons$ | $8.35 \times 10^{4}$ M$^{-1}$ | tuning parameter |
|        | $Fe^{III}(Cit)(OH)^- + H^+$ |                      |         |
| E7     | $Fe^{3+} + HCit^{2-} \rightleftharpoons Fe^{III}(HCit)^+$ | $2.51 \times 10^{7}$ M$^{-1}$ | tuning parameter |
| E8     | $Fe^{2+} + HCit^{2-} \rightleftharpoons Fe^{II}(HCit)$ | $1.935 \times 10^{10}$ M$^{-1}$ | tuning parameter |
| E9     | $Fe^{3+} + H_2O \rightleftharpoons Fe^{III}(OH)^{2+} + H^+$ | $4.57 \times 10^{-3}$ M | Smith and Martell (1976) |
| E10    | $O_2^- + H^+ \rightleftharpoons HO_2$ | $6.3 \times 10^{4}$ M$^{-1}$ | Bielski et al. (1985) |
| E11    | $Fe^{2+} + O=O=C(CH_2COO)^{2-} \rightleftharpoons$ | $2 \times 10^{3}$ M$^{-1}$ | tuning parameter |
|        | $Fe^{II}[O=C(CH_2COO)]_2$ |                      |         |
| E12    | $2H^+ + OH-C(CH_2COO)^{2-} \rightleftharpoons$ | $1.5 \times 10^{6}$ M$^{-2}$ | tuning parameter |
|        | $OH-C(CH_2COO)H_2$ |                      |         |
| E13    | $2H^+ + O=C(CH_2COO)^{2-} \rightleftharpoons$ | $1.5 \times 10^{6}$ M$^{-2}$ | tuning parameter |
|        | $O=C(CH_2COOH)H_2$ |                      |         |
| R1     | $Fe^{III}(Cit) + hv \rightarrow$ | $3.0 \times 10^{-18}$ (at 375 nm) or $2.3 \times 10^{-19}$ | Pozdnyakov et al. (2012) |
|        | $Fe^{2+} + OH-C(CH_2COO)^{2-} + CO_2$ | | |
| R2     | $Fe^{III}(Cit)(OH)^- + hv \rightarrow$ | | |
|        | $Fe^{2+} + OH-C(CH_2COO)^{2-} + OH^- + CO_2$ | | |
| R3     | $OH-C(CH_2COO)^{2-} + O_2 \rightarrow$ | $1 \times 10^{6}$ M$^{-1}$ s$^{-1}$ | Hug et al. (2001) |
|        | $O=C(CH_2COO)^{2-} + O_2^- + H^+$ | | |
| R4     | $HO_2 + HO_2 \rightarrow$ | depends on $a_w$ | tuning parameter |
|        | $H_2O_2 + O_2$ | | |
| R5     | $Fe^{2+} + O_2^- (+2H^+) \rightarrow Fe^{3+} + H_2O_2$ | $1 \times 10^{7}$ M$^{-1}$ s$^{-1}$ | Rush and Bielski (1985) |
| R6     | $Fe^{2+} + HO_2 (+H^+) \rightarrow Fe^{3+} + H_2O_2$ | $1.2 \times 10^{6}$ M$^{-1}$ s$^{-1}$ | Rush and Bielski (1985) |
| R7     | $Fe^{2+} + H_2O_2 \rightarrow Fe^{3+} + OH + OH^-$ | $76$ M$^{-1}$ s$^{-1}$ | Walling (1975) |
| R8     | $Fe^{2+} + OH \rightarrow Fe^{III}(OH)^{2+}$ | $4.3 \times 10^{8}$ M$^{-1}$ s$^{-1}$ | Christensen and Sehested (1981) |
| R9     | $Fe^{II}(HCit) + O_2 \rightarrow Fe^{III}(Cit) + HO_2$ | $0.05$ M$^{-1}$ s$^{-1}$ | tuning parameter |
| R10    | $C_5 + hv \rightarrow$ | $1 \times 10^{-21}$ cm$^2$ | tuning parameter |
|        | $C_3 + 2CO_2$ | | |
| R11    | $C_8 + hv \rightarrow$ | $1 \times 10^{-20}$ cm$^2$ | tuning parameter |
|        | $C_3 + CO_2$ | | |
| R12    | $C_5 + hv \rightarrow$ | $1 \times 10^{-22}$ cm$^2$ | tuning parameter |
|        | $C_2 + C_3$ | | |
| R13    | $C_4 + hv \rightarrow$ | $1 \times 10^{-21}$ cm$^2$ | tuning parameter |
|        | $C_2 + C_2$ | | |
| R14    | $C_5 + hv \rightarrow$ | $1 \times 10^{-20}$ cm$^2$ | tuning parameter |
|        | $C_4 + CO_2$ | | |
| R15    | $C_{stable}$ | 0.5 | tuning parameter |
| R16    | radical self-reaction | 0 | tuning parameter |
analyzing the STXM/NEXAFS experiments shown in Fig. 7. We note that our estimate of the photochemical reduction rate and agreement with data is based on the reproducibility of setting up our optical system in the X-ray vacuum chamber. In Fig. 8 (discussed below), this setup procedure was performed for RH = 40 % and RH = 50–60 % independently, and still, iron reduction was in agreement with model predictions in the first minutes after UV light was switched off. The width of the red shading in Fig. 7 is large and the UV-fiber setup was likely the largest source of error for \( j \). A better estimate would require repeat measurements as a function of RH, e.g., to elucidate any systematic uncertainty on iron reduction reactions due to viscosity changes. However, this was not possible as usage of the X-ray beam for STXM/NEXAFS experiments was limited to a few days to complete all experiments.

In a different set of experiments, we irradiated particles using a much lower power setting with \( j = 2.2 \times 10^{-3} \text{ s}^{-1} \) in a mixed He and O\(_2\) atmosphere and at a fixed RH for 15 min to reduce Fe\textsuperscript{III} to Fe\textsuperscript{II}. The UV light was then switched off to allow reoxidation in the dark while measuring \( \beta \) over time. Figure 8 shows \( \beta \) as a function of time at RH = 40 %, 50 %, and 60 %. Clearly, the Fe\textsuperscript{III} fraction increased more significantly more slowly with time at drier conditions. While particles were observed to reoxidize to \( \beta = 0.7 \) within 2 h at 60 % RH and expected to be completely reoxidized within about 6 h according to the PRAD model simulations, no significant reoxidation occurred on this timescale for the particles exposed to only 40 % RH. Modeling the reoxidation with the PRAD model yields very satisfactory agreement indicating that the diffusivity parameterizations of the model are capturing the RH dependence of the molecular transport in the viscous matrix.

3.3 Determination of iron(II) reoxidation rate with single, levitated particles using EDB

As the experiments with single, levitated particles yield only bulk properties and not the oxidation state of iron citrate directly, we designed a dedicated experimental procedure to indirectly determine the reoxidation rate of Fe\textsuperscript{II}. We used multiple irradiation and reoxidation repetitions as shown schematically in Fig. 9. Initially, we exposed a newly injected Fe\textsuperscript{III} (Cit)/CA aqueous particle to blue laser irradiation (473 nm, 4 W cm\(^{-2}\)) in pure N\(_2\) for 500 s to ensure all Fe\textsuperscript{III} was reduced through photolysis (Reactions R1 and R2 listed in Table 2). As previously described, these reactions
led to CO$_2$ production with subsequent loss of CO$_2$ to the gas phase, which was observed as a shift in the Mie-resonance wavelength. This shift is shown in Fig. 2b and on an enlarged scale in Fig. 10. After irradiation, we switched the gas flow from N$_2$ to O$_2$ in the dark, and Fe$^{II}$ was oxidized back to Fe$^{III}$ over time in this period, either by ROS (Reactions R5–R8) or directly by O$_2$ (Reaction R9). After a defined time spent in O$_2$, the gas flow was switched back to N$_2$ followed by irradiation to repeat the photolysis step done initially. The ratio of the Mie-resonance wavelength shift of the two photolysis steps was set to be proportional to the ratio of reoxidized Fe$^{III}$/Fe$_{tot}$. These two steps (i.e., photolysis in N$_2$ and reoxidation in O$_2$) were repeated several times, but between each irradiation the particle was exposed to O$_2$ for different time periods. Following this procedure we intended to map out the characteristic time for reoxidation at various RH.

An example of the corresponding raw data (293.5 K and 48 % RH) and retrieved oxidation state is shown in Fig. 10. Black circles indicate the first irradiation step with a fresh Fe$^{III}$(Cit)/CA particle, and red circles indicate the irradiation step that followed the particle after exposure to O$_2$ for 45 min in the dark. Clearly, the resonance wavelength decreased more during the first irradiation than the second. Therefore, we can conclude unambiguously that the Fe$^{III}$(Cit)/CA particle initially had more Fe$^{III}$ than what could be reoxidized in O$_2$ for 45 min. Quantitative scaling, however, requires knowledge of the initial Fe$^{III}$ fraction. Our experiments showed that long exposure (tens of hours) to O$_2$ yielded larger Mie-resonance shifts than those of the initial photolysis of the freshly prepared particle. This indicated that the initial Fe$^{III}$ fraction was less than 1.0. Hence, we accordingly normalized the Fe$^{III}$ fraction to the data at long (> 15 h) exposure times. For the experiment included in Fig. 10, for example, the initial Fe$^{III}$ fraction of the particle was 0.76, indicating that the particle was partially reduced during sample preparation. After the particle was totally photoreduced, exposure to O$_2$ for 45 min did not reoxidize all reduced Fe$^{II}$ to Fe$^{III}$, only 0.36 Fe$^{III}$ was recovered (as shown by two horizontal dashed lines in Fig. 10).

Another set of experiments was done by starting with freshly injected Fe$^{II}$(HCit)/CA particles instead of Fe$^{III}$(Cit)/CA. The only difference in experimental procedure, compared to what is described above, is that there is no first irradiation step. Instead, the particle is initially exposed to O$_2$ in the dark for a certain time interval, during which Fe$^{II}$ is oxidized only directly by O$_2$ (Reaction R9).
Figure 11. Fe$^{\text{III}}$ fraction after different time intervals of recovery in O$_2$ from EDB observations. Fe$^{\text{III}}$(Cit) / CA (molar ratio of 0.05) particles at 293.5 K with 24 % RH (red diamonds), 48 % RH (green triangles), and 65 % RH (blue squares). Black dots: experiments with Fe$^{\text{II}}$(HCit) / CA particles at 48 % RH. Lines: red (24 % RH), black (48 % RH), and blue (65 % RH) are the Fe$^{\text{III}}$ fractions predicted using the PRAD model; the shaded areas indicate model output assuming ±2 % RH uncertainty in the EDB experimental conditions.

Afterwards, the same irradiation and recovery procedures as reported above were used. The equivalent Fe$^{\text{III}}$ fractions of all experiments are shown in Fig. 11. Any uncertainty in the normalization of an individual experiment will cause a corresponding uncertainty in the normalized mass loss, which made estimating the uncertainty for individual data points impossible. Nevertheless, from the complete data set, it was evident that with longer time intervals in O$_2$, more Fe$^{\text{III}}$ was recovered. At 48 % and 65 % RH, about 10 and 3 h exposure to O$_2$, respectively, was sufficient for all Fe$^{\text{II}}$ to be reoxidized, while at 24 % RH, the recovery even after 25 h was not yet complete. The general trend is consistent with our observation in the STXM/NEXAFS experiments (Sect. 3.2) and can be attributed to molecular diffusion limitations at lower RH: it takes more time for O$_2$ to diffuse into the particle and for Fe$^{\text{II}}$ to diffuse out to the surface of the particle to react with O$_2$. It was also evident from these data that Fe$^{\text{II}}$ reoxidized by O$_2$ is as important as Fe$^{\text{II}}$ reoxidized by radicals and peroxides as there is no significant difference between the experiments starting from Fe$^{\text{II}}$(HCit) / CA compared to those with Fe$^{\text{III}}$(Cit) / CA. In addition, it indicates that both iron(II) and iron(III) can act as a photocatalyst as long as iron(II) can be oxidized to iron(III), which was also confirmed by Grgi´c et al. (1999).

By tuning the direct oxidation rate of Fe$^{\text{II}}$(HCit) by O$_2$ (Reaction R9) and the diffusivity of O$_2$, we are able to model the recovery rate at different RH using the PRAD model as shown in Fig. 11. There is satisfactory agreement for the larger RH, but significant underestimation of reoxidation for the experiments at 24 % RH. Our model requires the reaction rate coefficient of Reaction (R9) to be 0.05 M$^{-1}$s$^{-1}$, which is a factor of 60 smaller than the value that Gonzalez et al. (2017) estimated from their model (3 ± 0.7 M$^{-1}$s$^{-1}$). The liquid phase diffusivity of O$_2$ in our model is 7.1 × 10$^{-19}$ m$^2$s$^{-1}$ at 24 % RH, 1.9 × 10$^{-15}$ m$^2$s$^{-1}$ at 48 % RH, and 2.1 × 10$^{-14}$ m$^2$s$^{-1}$ at 65 % RH. These diffusivities of O$_2$ are 2–4 orders of magnitude smaller than those of CO$_2$ determined in Dou et al. (2019). However, we need to stress that in the model, some of the iron-related complex equilibrium constants, their diffusion coefficients, and the Henry’s law coefficient of O$_2$ at different water activities are highly uncertain as well, yielding to a significant uncertainty in the determination of O$_2$ diffusivity. For example, if the solubility of O$_2$ was less than what our parameters predict now, a larger O$_2$ diffusivity would be consistent with our data. In addition, with the total gas flow used in our experiments, it takes about 8 min for replacing the entire EDB gas volume from N$_2$ to O$_2$, and from the response of the particle to flow condition change, we estimate an interval of about 30 min to reach full equilibration to the new gas phase conditions. Therefore, the lifetime of organic radicals needs to be reconsidered. In the PRAD model, we do not take the radical–radical self-reactions (Reaction R16) into account, which may turn out to be a significant sink for the radicals. However, it should be pointed out that the parameter set we have now is a good compromise with additional constraints from STXM/NEXAFS and CWFT experiments.

Another approximate approach to analyze the data of Fig. 11 is to use the analytical solutions for a reacto-diffusive kinetic regime. Here, O$_2$ taken up from the gas phase by a particle remains confined to a very thin layer below its surface compared to its size provided it reacts reasonably fast with the organic components. Under these conditions there are always pairs of reaction rate and diffusion constants representing the experiments equally well (Alpert et al., 2019; Steimer et al., 2014). In the reacto-diffusive framework, with a constrained reaction rate and Henry’s law coefficient of O$_2$, the diffusion coefficient of O$_2$ can be estimated to be 3.6 × 10$^{-16}$ and 4.4 × 10$^{-15}$ m$^2$s$^{-1}$ at 48 % and 65 % RH, respectively (details are given in Appendix A4). These are both 1 order of magnitude less than the values from PRAD model prediction, but still consistent with each other when considering all uncertainties.

3.4 HO$_2$ production measured by CWFT experiments

The CWFT experiment allows us to investigate another aspect of the photochemistry of the Fe$^{\text{III}}$(Cit) / CA system. According to Reactions (R1)–(R3) shown in Table 2, the HO$_2$ radical is produced upon irradiation and will partition to the gas phase. Figure 12a shows the RH dependence of HO$_2$ production, $P_{\text{HO}_2}$, from thin films in the CWFT experiment. We observed that $P_{\text{HO}_2}$ increased with RH when the RH was in-
Dotted and dashed blue lines were determined from parameterization of HO\textsubscript{2} production from continued photochemistry of Fe\textsuperscript{III}(Cit) / CA (molar ratio of 0.07) films as a function of RH in CWFT experiments. Dotted and dashed blue lines were determined from parameterization of k\textsubscript{5} as a function of RH (Corral Arroyo et al., 2018) and k\textsubscript{5} = 8.3 × 10\textsuperscript{5} M\textsuperscript{-1}s\textsuperscript{-1} in dilute aqueous solution (Bielski et al., 1985), respectively. (b) The maximum HO\textsubscript{2} first-order loss rate determined from the result of the maximum HO\textsubscript{2} concentration times HO\textsubscript{2} self-reaction rate, k\textsubscript{5}, which was adjusted to match observations in panel (a) (black dots), or times k\textsubscript{5} parameterized as a function of RH for our PRAD model (Eq. A11) with confidence interval of 95 % (red line and shading), or times k\textsubscript{5} from Corral Arroyo et al. (2018) (dotted blue line), or k\textsubscript{5} = 8.3 × 10\textsuperscript{5} M\textsuperscript{-1}s\textsuperscript{-1} (Bielski et al., 1985) (dashed blue line).

Figure 12. (a) Measured (black dots, experimental uncertainty of 3 % smaller than symbol size) and modeled (red line and shaded area) HO\textsubscript{2} production from continued photochemistry of Fe\textsuperscript{III}(Cit) / CA (molar ratio of 0.07) films as a function of RH in CWFT experiments.

3.5 Photochemical degradation under atmospheric conditions

The PRAD model was also developed to be used in more general particle systems. After establishing a parameter set for the PRAD model framework that satisfactorily explains the experimental data obtained with three complementary experimental techniques over a wide parameter range, we used the model to predict photochemical degradation of organic aerosol particles containing carboxylate complexes. For example, an aqueous organic acid particle with a dust inclusion containing iron may exhibit a low enough pH to dissolve part of the iron of the dust inclusion and form photoreactive iron carboxylate complexes (George et al., 2012, 2015). If we take the PRAD model as being representative for such a class of particles, we may estimate the degradation of the organic mass by volatilization of products to the gas phase. Figure 13 shows examples of such PRAD model predictions: we assume that the organic mass of the particle is composed of aqueous Fe\textsuperscript{III}(Cit) / CA with a molar ratio of the latter being 0.01. Here, panel (a) illustrates size dependence and panel (b) illustrates RH dependence for particles exposed to air at 293.5 K and an actinic flux at 30° zenith angle (Madronich et al., 1995). After 12 h exposure to sunlight, the organic mass of the particle (here, CA) was photochemically processed to CO\textsubscript{2} and smaller compounds of high volatility yielding a combined mass loss to the gas phase of up to 50 % depending on size and RH.

To better understand the importance of mass loss in this generalized system, we make a simple and modest comparison of mass loss in Fig. 13 on the order of 20 % over 5 h with ambient mass accumulation measured in the field. Our results are equal to a mass loss rate of about 0.4 µg m\textsuperscript{-3} (air) h\textsuperscript{-1} assuming an aerosol population with an organic mass of ~ 10 µg m\textsuperscript{-3} (air) undergoing iron carboxylate photochemistry. This rate is much higher than the observed organic mass accumulation in ambient air masses due to photochemical aging during atmospheric transport at about 0.06 µg m\textsuperscript{-3} h\textsuperscript{-1} or 6 µg m\textsuperscript{-3} over 4 d (Zaveri et al., 2012; Moffet et al., 2012). This implies that the mass loss rates are fast enough to affect the balance between aerosol mass accumulation and loss. Hence, photochemical degradation may be very significant in
iron-containing organic aerosol, perhaps even more important than degradation through reactions with gas phase oxidants.

Note that our model does not fully capture the acceleration of degradation as discussed in Sect. 3.1. Therefore, we argue that the degradation presented in Fig. 13 represents a lower limit of the expected degradation under atmospheric conditions. Clearly, panel (a) indicates that smaller particles degrade significantly faster than larger ones. However, the size dependence is more complex than the expected simple square law if reactions were purely limited by condensed phase diffusion. For particles with a radius larger than 50 nm, the time to repartition 10% of the mass to the gas phase depends almost linearly on size for these specific conditions. Panel (b) shows that photochemical processes are faster at higher RH corresponding to less viscous particles. The relative mass loss after 12 h irradiation between 20% and 60% RH is almost an order of magnitude larger for the highest RH compared to dry conditions because of transport limitations at low RH. However, it should be noted that the relative mass loss at higher humidities (between 40% and 60% RH) is quite similar. Here, the photochemical degradation is barely limited by condensed phase diffusivity, but by iron availability in the particle. These simulations show the potential of photochemical degradation under atmospheric conditions; a systematic study exploring the whole range of atmospheric conditions is beyond the scope to this work.

4 Conclusions

We used three complementary experimental techniques to characterize the impact of reduced mobility of aerosol constituents on photochemical degradation in highly viscous particles. As an atmospherically relevant model system, we chose aqueous Fe\textsuperscript{III}(Cit) / CA particles. These three experimental techniques investigated specific aspects of this photochemical reaction system. In EDB experiments, we measured the mass loss relating to the continual production and loss of CO\textsubscript{2} and other volatile products. We observed very significant condensed phase degradation and strong acceleration of the degradation rate with time. Further studies are needed to quantify all atmospheric implications, but our study suggests that photochemistry in iron-containing organic aerosol will lead to a significant repartitioning of condensed phase mass to the gas phase. We used STXM/NEXAFS to directly measure iron oxidation state in situ with an environmental microreactor. These experiments yielded valuable information about where iron photochemical reduction and reoxidation reactions took place, namely only very close to the surface, and it allowed us to characterize to what degree iron compounds diffused inside single particles. We showed that O\textsubscript{2} uptake and diffusion into a particle is a limiting factor considering the reactions required to produce species with an oxidative potential. In addition, we found that the direct O\textsubscript{2} reaction with iron(II) organic complexes does occur and generates radicals inside the particle. Flow-tube experiments performed on thin Fe\textsuperscript{III}(Cit) films showed continuous production of HO\textsubscript{2}, revealing a radical source inside the particles driven by photochemistry.

All data were used to constrain equilibrium and kinetic parameters as well as reaction rate coefficients in a new PRAD model with sufficient complexity to allow comparison with data of all experiments simultaneously. In particular, we were able to constrain the photolysis rate of Fe\textsuperscript{III} due to the use of various light sources with various spectral intensities, while capturing the photochemical reduction. In addition, we determined the HO\textsubscript{2} production rate and its first-order loss rate, and the diffusivity of O\textsubscript{2} in aqueous Fe\textsuperscript{III}(Cit) / CA system as a function of RH and Fe\textsuperscript{III}(Cit) / CA molar ratio with a choice of O\textsubscript{2} related reaction rate coefficient and O\textsubscript{2} Henry’s law coefficient.

Although a systematic study exploring the whole range of atmospheric conditions was beyond the scope to this work, there are some aspects of the PRAD model and certain parameters that we argue are reliable and pertinent to atmospheric aerosol photochemistry. First, coefficients in the
The chemical evolution of the organic species resulting from the continual photochemical oxidation was not the scope of the present study. A separate study focusing on individual OVOCs and condensed phase products is currently underway that will allow us to better constrain the chemical regimes and the evolution of the oxidation state of the organic fraction with time. This will also allow us to assess more details of organic peroxy radical chemistry that help explain the observed “missing” HO₂ sink in this system. Furthermore, testing the PRAD model with different organic carboxylate ligands is desirable to broaden its applicability.

Using the PRAD model for predicting photochemical degradation for iron-containing organic aerosol under atmospheric conditions let us conclude that this pathway of repartitioning condensed phase mass to the gas phase is important and its regional and global impact should be investigated in further modeling studies. The PRAD model may serve as a basic framework for the chemistry and transport of compounds in single particles for such studies.
**Appendix A**

### A1 Parameterization of $D_j^l(x, T, a_{aw})$

The liquid phase diffusion coefficients, $D_j^l(x, T, a_{aw})$, where \( j \) is an index for all species, depend on RH, \( T \) and the molar ratio, \( x \), between Fe$^{III}$(Cit) and CA. $D_j^l(x, T, a_{aw})$ was scaled with the diffusion coefficient of water in a Fe$^{III}$(Cit)/CA aqueous system, $D_{H_2O}^l(x, T, a_{aw})$, using a scaling factor, \( f_j \), following

\[
l_j^l = \frac{D_j^l(x, T, a_{aw})}{D_{H_2O}^l(x, T, a_{aw})}, \tag{A1}
\]

\( D_j^l(x, T, a_{aw}) \) has not been previously determined for aqueous Fe$^{III}$(Cit)/CA although solution viscosity has and was found to be higher than aqueous CA solutions when \( x > 0.05 \) (Alpert et al., 2021), implying slower molecular transport. Lienhard et al. (2014) reported the diffusion coefficient of water in aqueous CA without iron (i.e., \( x = 0 \)), $D_{H_2O}^{l\text{CA(aq)}}(T, a_{aw})$. In order to determine \( D_j^l(x, T, a_{aw}) \) in the PRAD model for a single experiment with a fixed value of \( x \), $D_{H_2O}^l$ was scaled with $D_{H_2O}^{l\text{CA(aq)}}(T, a_{aw})$ using another factor, \( f_s \), following

\[
D_{H_2O}^l(x, T, a_{aw}) = D_{H_2O}^{l\text{CA(aq)}}(T, a_{aw}) f_s(x), \tag{A2}
\]

where

\[
\log f_s(x) = -0.7106 e^{-\frac{x}{2}}. \tag{A3}
\]

The diffusion coefficient of CO$_2$ at \( T = 20^\circ\text{C} \) and \( x = 0.05 \), $D_{CO_2}^s$, was independently measured by Dou et al. (2019) as a function of \( a_{aw} \). We parameterized $D_{CO_2}^s$ as

\[
D_{CO_2}^s = \left(D_{CO_2}^s(a_{aw} = 1)\right)^{a_{aw}} \cdot \left(D_{CO_2}^s(a_{aw} = 0)\right)^{1-a_{aw}}, \tag{A4}
\]

where $D_{CO_2}^s(a_{aw} = 0) = 1.19 \times 10^{-16} \text{m}^2\text{s}^{-1}$ is the diffusion coefficient of CO$_2$ at \( T = 20^\circ\text{C}, x = 0.05, \) and \( a_{aw} = 0 \). The \( T \) dependent diffusion coefficient of CO$_2$ in water in Eq. (A4) is

\[
D_{CO_2}^s(a_{aw} = 1) = D_0 \left(\frac{T}{T_{ks}} - 1\right)^m, \tag{A5}
\]

where $D_0 = 1.39 \times 10^{-8} \text{m}^2\text{s}^{-1}$, $T_{ks} = 227.0$ K, and $m = 1.7094$. In the exponent terms of Eq. (A4),

\[
\alpha = e^{(1-a_{aw})^2(A+B a_{aw})}, \tag{A6}
\]

where $A = 0.2824$ and $B = -1.8086$. In order to introduce a \( x \) dependence of $D_{CO_2}^1$ for various experiments reported here, \( f_s \) from Eq. (A3) for water was applied following

\[
D_{CO_2}^1(x, T, a_{aw}) = D_{CO_2}^s(T, a_{aw}) f_s(x). \tag{A7}
\]

For all other species (excluding H$_2$O and CO$_2$), $D_j^l(x, T, a_{aw})$ was determined using Eq. (A1) with

\[
\log l_j^l = -0.7710 M_j^{\frac{1}{2}} - 1.4732, \tag{A8}
\]

where, \( M_j \), is molar mass. The diffusion coefficients of H$_2$O, CO$_2$, O$_2$, and CA as a function of \( a_{aw} \) at \( x = 0.05 \) and 1.0 are plotted in Fig. A1. Generally, \( D_j^l(x, T, a_{aw}) \) always decreases when RH or \( T \) is lowered. \( D_j^l(x, T, a_{aw}) \) decreases when \( x \) increases beyond 0.05, but remains relatively constant otherwise. One caveat to using Eqs. (A1)–(A8) to calculate \( D_j^l \) is that mass loss will ultimately lead to an increase in \( x \), however, the PRAD model keeps \( D_j^l \) fixed throughout the course of a model run. Since we observed increasing mass loss rates over time, any decrease in \( D_j^l \) leading to slower chemical cycling due to increasing \( x \) was likely a minor effect. Moreover, the product distribution and any effect on diffusion coefficients was unknown, so further time-resolved adjustments to \( D_j^l \) were not considered. We suggest future studies investigate how molecular transport changes over the photochemical lifetime of iron carboxylate complexes.

### A2 Parameterization of $D_j^g$

The gas phase diffusivity of each species \( j \), $D_j^g$, was approximated via its molar mass ($M_j$) compared to that of water (M$_{H_2O}$),

\[
D_j^g = D_{H_2O}^g \sqrt{\frac{M_{H_2O}}{M_j}}, \tag{A9}
\]
with

\[ D_{K}^{\text{H}_{2}\text{O}} = 0.211 \left( \frac{T}{T_0} \right)^{1.94} \left( \frac{P_0}{p} \right), \]  

(A10)

where \( T_0 = 273.15 \text{ K} \), \( p_0 = 1013.25 \text{ mbar} \), and \( D_{K}^{\text{H}_{2}\text{O}} \) is in \( \text{cm}^2 \text{s}^{-1} \) (Pruppacher and Klett, 2010).

A3 Parameterization of \( k_5 \)

Based on the measurement of \( \text{HO}_2 \) production as a function of RH using CWFT experiments, the HO2 self-reaction rate (Reaction R4 in Table 2), \( k_5 \), was adjusted so the PRAD model would exactly reproduce the data. We parameterized \( k_5 \) as a third degree polynomial function of RH (%):

\[
\log k_5 = -2.854 \times 10^{-5} \text{RH}^3 + 0.0024 \text{RH}^2 + 0.1087 \text{RH} - 0.05018,
\]

(A11)

as shown in Fig. A2.

A4 \( \beta \) estimated by a reacto-diffusive framework

It is well known that multiphase reactions can follow a reacto-diffusive kinetic regime (Alpert et al., 2019; Steimer et al., 2014). For reacto-diffusive limitations as observed in Fig. 11, the Fe\textsuperscript{III} fraction as a function of time was calculated as

\[
\beta(t) = 1 - (1 - Kt)^2,
\]

(A12)

with the assumption that \( \beta_0 = 0 \), i.e., there was no Fe\textsuperscript{III} at \( t = 0 \):

\[
K = \frac{3(H_{O_2} D_{O_2}^{1/2}) p_{O_2} \sqrt{k_{O_2}}}{\sqrt{[Fe_{tot}] dp}}.
\]

(A13)

where \( HO_2 \) and \( D_{O_2}^{1/2} \) are the Henry’s law coefficient and diffusion coefficient, respectively, for \( O_2 \) in the Fe\textsuperscript{III}(Cit)/CA matrix, \( p_{O_2} \) is the pressure of \( O_2 \), \( k_{O_2} \) is the total reaction rate of \( O_2 \), and \( d_p \) is the diameter of the single particle. \( \beta = \frac{t}{T} \) when

\[
I_\frac{1}{2} = \frac{1 - \sqrt{1 - \frac{1}{e}}}{K} = \frac{1 - \sqrt{1 - \frac{1}{e}}}{\sqrt{[Fe_{tot}dp]} 3(H_{O_2} D_{O_2}^{1/2}) p_{O_2} \sqrt{k_{O_2}}}
\]

(A14)

In a typical EDB experiment, \( d_p = 20 \mu\text{m} \), \( H_{O_2} = 3.5 \times 10^{-2} \text{ M atm}^{-1} \), \( p_{O_2} = 8 \times 10^4 \text{ Pa} = 0.789 \text{ atm} \), and \( k_{O_2} = 0.05 \text{ M}^{-1} \text{s}^{-1} \). \([Fe_{tot}] \) is 0.3192, 0.2763, and 0.2345 M at 24 %, 48 %, and 65 % RH, respectively. From EDB data points in Fig. 11, we estimated that \( \frac{t}{T} \) of Fe\textsuperscript{III} can be fully recovered after around 1.7 h at 48 % RH and 0.45 h at 65 % RH. Thus, the diffusion coefficient of \( O_2 \) can be estimated to be \( 3.6 \times 10^{-16} \) and \( 4.4 \times 10^{-15} \text{ m}^2 \text{s}^{-1} \) at 48 % and 65 % RH, respectively, which are both 1 order of magnitude less than the values from PRAD model prediction, but still consistent with each other when considering all uncertainties. It should be noted that in Eq. (A14), \( HO_2 \sqrt{D_{O_2}^{1/2} k_{O_2}} \) is actually the constraint, thus any uncertainty in \( HO_2 \) or \( k_{O_2} \) can change \( D_{O_2}^{1/2} \).

A5 Sensitivity of the PRAD model to various model parameters

As discussed in Sect. 2.5, we performed manual tuning of the PRAD model parameters to reach satisfactory agreement with all experimental data simultaneously. To show the sensitivity of the PRAD model results to a few of its parameters, Fig. A3 shows again the data of the photocatalytic degradation experiment at 46 % RH described in Sect. 3.1. In addition, we show the output of the model as well as model outputs obtained by varying one of the parameters by the indicated factors while keeping all other parameters constant. Clearly, the sensitivities of the model output to varying these parameters are very different: while the model output is quite sensitive to varying the rate constant for the oxidation of Fe\textsuperscript{II}(HCit) (Reaction R9 in Table 2) as well as to oxygen diffusivity, the sensitivity to the equilibrium constant E8, Fe\textsuperscript{II} + HCit\textsuperscript{2−} = Fe\textsuperscript{II}(HCit), is significantly smaller and the model is basically insensitive to the equilibrium constant E5, Fe\textsuperscript{III} + Cit\textsuperscript{3−} = Fe\textsuperscript{III}(Cit). In other words, this experiment alone allows us to constrain Reaction (R9) or oxygen diffusivity as long as the other parameter is known, but does not allow us to constrain the constants for the two equilibria.

A6 The full EDB data set showing the RH effect on photocatalytic degradation efficiency

Figure 6 in the main text shows the photochemical degradation up to a mass remaining ratio of 0.6 and compares its temporal evolution with the PRAD model output. Here, we
Figure A3. Black: mass remaining data versus time of the photodegradation experiment discussed in Sect. 3.1; RH 46 %. Red lines: model outputs obtained by varying a single parameter with factors: 10 and 0.1 (dashed), 5 and 0.2 (dotted), 2 and 0.5 (dash-dotted), 1 (solid). In panel (a), the equilibrium constant of E8 is varied and in panel (b) the equilibrium constant of equilibrium E5 is varied. Panel (c) shows the sensitivity to the reaction constant (Reaction R9, see Table 2) and panel (d) shows the sensitivity to oxygen diffusivity (Table 1).

Figure A4. Fe^{III} (Cit)/CA (molar ratio 0.05) particle mass change with irradiation time (log scale) at different RH: 33 % (red), 46 % (green), and 61 % (blue). The irradiation wavelength was 375 nm, its intensity was 0.25 W cm\(^{-2}\), and the experimental temperature was 293.5 K. Thick lines are EDB experimental data. Thin lines are PRAD model outputs (with ±2 % RH uncertainty shown as the shaded area).

show in Fig. A4 for completeness the full experimental data set. Clearly seen is the shift to longer time scales with decreasing RH and that the PRAD model is no longer capable of accurately capturing the mass loss once about 20 % of the initial mass is lost. However, we may conclude that even for mass remaining ratios lower than 0.5, the experimental data show a very significant shift of the degradation time scales with RH, but the total photochemical mass loss remains of the same order of magnitude, independent of RH.
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