A comparison of the chemical sinks of atmospheric organics in the gas and aqueous phase

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Abstract. Photochemical reactions represent the main pathway for the removal of non-methane volatile organic compounds (VOCs) in the atmosphere. VOCs may react with hydroxyl radical (OH), the most important atmospheric oxidant, or they can be photolyzed by actinic radiation. In the presence of clouds and fog, VOCs may partition into the aqueous phase where they can undergo aqueous photolysis and/or reaction with dissolved OH. The significance of direct aqueous photolysis is largely uncertain due to the lack of published absorption cross sections and photolysis quantum yields. In light of this, we strive to identify atmospherically relevant VOCs where removal by aqueous photolysis may be a significant sink. The relative importance of different photochemical sinks is assessed by calculating the ratios of the removal rates inside air parcels containing cloud and fog droplets. This relative approach provides useful information in spite of the limited aqueous photolysis data. Results of this work should help guide researchers in identifying molecules that are the most likely to undergo aqueous OH oxidation and photolysis. For example, we find that out of the 27 atmospherically relevant species investigated, the removal of glyceraldehyde and pyruvic acid by aqueous photolysis is potentially an important sink. We also determine the relative magnitudes of these four chemical sinks for the set of relevant organic compounds.

1 Introduction

Organic compounds play a key role in the chemistry of the atmosphere. Global emissions of non-methane volatile organic compounds (VOCs) total approximately 1300 Tg C yr\(^{-1}\) (Atkinson and Arey, 2003; Goldstein and Galbally, 2007). It is estimated that 130–270 Tg C yr\(^{-1}\) of emitted VOCs are removed by dry and wet deposition, while the remaining 1030–1170 Tg C yr\(^{-1}\) are transformed by chemical reactions. Oxidation by hydroxyl radical is the most important atmospheric sink for many VOCs (Andreae and Crutzen, 1997). Gas phase photolysis can also be important for certain classes of compounds including carbonyls (Moortgat, 2001), peroxides (Lee et al., 2000), and organic nitrates (Atkinson, 1990). In the presence of clouds and fog, VOCs may partition into the aqueous phase to various extents depending on their solubility. As in the gas phase, OH is the main oxidant for the transformation of aqueous organics in cloud and fog droplets (Ervens et al., 2011). Direct photolysis in the aqueous phase can also be an important sink depending on the chemical functionality of the species (Vione et al., 2006; Graedel and Goldberg, 1983).

Given the complexity of possible VOC photo-oxidation mechanisms and products, we strive to classify compounds by the rate at which they are removed by gaseous OH oxidation, gaseous photolysis, aqueous OH oxidation, and direct aqueous photolysis. We develop a framework to compare these four rates of removal for a subset of compounds identified as likely contributors to the water-soluble fraction of atmospheric particles (Saxena and Hildemann, 1996). Figure 1 is a schematic of our analysis showing the four chemical sinks under investigation and the dimensionless parameters we derive to represent the behavior of the selected VOCs. Direct liquid photolysis is parameterized into a photolysis rate constant that depends on the actinic flux of solar radiation, the quantum yield of photolysis, and the absorption cross section of the absorbing species. In contrast to the wealth of photochemical studies of organic molecules in the gas-phase, in organic solvents, glassy solids, and rare-gas matrices,
measurements of photolysis quantum yields of organic compounds in water are more limited. The photochemistry treatise by Calvert and Pitts (1966) describes a few early aqueous photolysis studies of organic compounds, most of which were done under 254 nm irradiation. Only a handful of additional photolysis quantum yield studies in aqueous solutions under irradiation conditions that are relevant for the lower atmosphere (\( \lambda > 280 \text{ nm} \)) have appeared since then. Examples include photolysis of a number of organic pesticides (Vione et al., 2006; Wan et al., 1994), aromatic carbonyls (Ledger and Porter, 1972), substituted phenols (Czaplicka, 2006; Albinet et al., 2010), substituted benzyl derivatives (Zimmerman and Sandel, 1963), small aldehydes without correction for hydration (Hirshberg and Farkas, 1937), dicarbonyls such as biacetyl (Faust et al., 1997), hydrogen peroxide (Chu and Anastasio, 2005; Goldstein et al., 2007), methyl peroxide (Epstein et al., 2012), and pyruvic acid (Leemakers and Vesley, 1963; Larsen and Vaida, 2012; Guzmán et al., 2006, 2007). This data set is too meager for the development of a conceptual treatment of quantum yields or even an educated estimation for the selected atmospherically relevant compounds. In the absence of aqueous absorption cross section and quantum yield data for most organics, we aim to identify systems where direct aqueous photolysis is potentially a significant sink. While photolysis of certain organic compounds may generate OH radical in the gaseous or aqueous phase (Zellner et al., 1990; Epstein et al., 2012; Faust and Allen, 1993; Faust, 1994; Chu and Anastasio, 2005; Monod et al., 2007), we investigate direct photolysis and OH oxidation independently. Direct photolysis and indirect photolysis (reaction with oxidants formed by photolytic reactions) are linked by the potential generation of OH in the photolysis of certain organic compounds. However, OH yields from direct photolysis of most organic compounds are uncertain and depend on the photolytic mechanism. With published OH rate constants for oxidation in the gas and aerosol phase, along with absorption cross section and quantum yield data for photolysis in the gas phase, we can estimate the significance of these four chemical sinks to better understand the chemical fate of water soluble atmospheric organics. We stress that this is not a modeling study – our goal is to help guide researchers in identifying molecules that are the most likely to undergo aqueous OH oxidation and/or direct photolysis.

We ignore several other VOC chemical sinks to simplify our analysis and focus on the competition between oxidation by OH and photolysis. For example, alkenes are susceptible to oxidation by ozone (Bailey, 1978) and NO3 (Atkinson, 1991) but we are not investigating any olefinic compounds. On a global scale, organics may react with Cl atoms, however this is thought to be a minor VOC loss process (Atkinson and Arey, 2003). For specific compounds under certain conditions, aqueous reactions with dissolved sulfur dioxide in cloud and fog droplets may be significant (Finlayson-Pitts and Pitts, 2000). In addition, organic compounds can take part in various isomerization, oligomerization, and hydrolysis reactions, which are often catalyzed by inorganic constituents of the droplets such as acids. Biological activity may also lead to the destruction of dissolved organics in cloud droplets. Microbes can drive the oxidation of carbonaceous compounds during periods with low dissolved OH concentrations in cloud droplets (Vaitilingom et al., 2010). We also neglect potential photosensitization processes from contaminants dissolved in atmospheric waters (Vione et al., 2006).

2 Methods

Out of the oxygenated organic compounds identified by Saxena and Hildemann (1996) we choose a subset of 27 compounds for which there is the required experimental data published in the literature. These selected compounds contain a range of functionalities: monocarboxylic acids, dicarboxylic acids, aldehydes, ketones, keto-carboxylic acids, ethers, and peroxides. While a similar analysis for a series of nitrogen-containing organic compounds is beyond the scope of this manuscript, we also investigate methyl nitrate, the simplest organic nitrate. With the exception of tartaric acid and malic acid, all compounds are either intermediate-volatility organic compounds (IVOCs) or volatile organic compounds (VOCs); the majority of their mass partitions in the vapor phase as opposed to the condensed phase. Tartaric acid and malic acid are low-volatility organic compounds (LVOCs); their mass will partition to the condensed phase in almost all atmospheric conditions (Donahue et al., 2009). The oxygen to
carbon ratio of the selected compounds ranges from 0 to 2. A full list of the compounds used and their corresponding references is presented in the Supplement. All of the IVOCs and VOCs are delivered to cloud and fog droplets by gaseous diffusion. However, tartaric acid and malic acid, as components of particulate matter, may also be delivered to the aqueous phase by particle scavenging, known to be very efficient in clouds (Limbeck and Puxbaum, 2000). For each of the compounds studied, we assume that the partitioning between the gaseous and aqueous phase is in a local state of equilibrium. Because of the variability of transport processes and competing reactions, equilibrium may not be completely established in all cases (Winiwarter et al., 1994; Audiffren et al., 1998).

For example, complete equilibrium is unlikely when water droplets of different sizes and pHs are present or in the case of an air mass with a variable liquid water content. Henry's law may not hold in the presence of irreversible chemical reactions that are fast enough to prevent the establishment of equilibrium (Finlayson-Pitts and Pitts, 2000), in droplets that are not well mixed (Finlayson-Pitts and Pitts, 2000), clouds that have been recently formed (Chaumerliac et al., 2000), or droplets that are not sufficiently dilute (Seinfeld and Pandis, 1998). In polluted environments, organics may enter cloud or fog droplets through the dissolution of organic aerosol particles potentially resulting in aqueous concentrations that exceed equilibrium values. However, for atmospheric gases at typical concentrations, Henry’s law reasonably approximates equilibrium with cloud or fog droplets (Seinfeld and Pandis, 1998), and our equilibrium-based calculations should therefore provide a good first-order estimate for the magnitude of aqueous phase processes. In the presence of high ion concentrations, Henry’s law constants can be modified with Satchenow coefficients (De Bruyn et al., 1995; Kolb et al., 1997). However, in order to streamline the analysis and maintain its generality, we assume that each of the compounds studied do not interact with each other or other inorganic compounds typically found in cloud or fog droplets.

2.1 Comparison of aqueous and gaseous photolysis

For atmospheric organics, the implications of direct aqueous photolysis (Bateman et al., 2011; Schwarzenbach et al., 2005; Nguyen et al., 2012) are elusive due to the lack of literature data. However, the direct gaseous photolysis of atmospheric organics is well-studied (Atkinson et al., 2006; Sander et al., 2011). We strive to exploit the likely similarities in these processes in order to constrain the magnitude of aqueous phase photolysis by comparison with gas phase photolysis. The photolysis rate constant, \( J \), is calculated with the following integral:

\[
J = \int F_\lambda(\lambda) \cdot \phi(\lambda) \cdot \sigma(\lambda) \cdot d\lambda
\]

where \( F_\lambda \) is the actinic flux, \( \phi \) is the quantum yield of photolysis, and \( \sigma \) is the absorption cross section, all functions of wavelength, \( \lambda \). In the absence of aqueous quantum yields and/or absorption cross sections, we can constrain the ratio of the gas phase and aqueous phase rate constants for a particular compound using educated assumptions on how its photochemistry is affected by dissolution in water. The actinic flux can be enhanced slightly inside a cloud droplet compared to the interstitial space (Finlayson-Pitts and Pitts, 2000). The maximum enhancement in the average actinic flux, calculated in Mayer and Madronich (2004), from work originally detailed in Madronich (1987) and Ruggaber et al. (1997) is 1.26 and 1.33, respectively. The solvent has the potential to change the absorption cross sections upon dissolution, known as solvatochromism (Marini et al., 2010), however, this effect is likely insignificant for the set of compounds that we are investigating. Xu et al. (1993) finds that the absorption cross section shifts by about 10 nm to the blue and the intensity increases by 15–35 % for a series of small ketones when they are dissolved in water. For hydrogen peroxide (Graedel and Weschler, 1981) and methyl peroxide (Epstein et al., 2012), the absorption cross sections of the gas and aqueous phases are similar. The magnitude of the change in quantum yield is far less certain. For direct aqueous photolysis, the quantum yield is generally lower than the corresponding gas-phase value because energy dissipation due to collisions with the solvent molecules (cage effect) depresses the quantum yield in solution (Calvert and Pitts, 1966; Farkas and Hirshberg, 1937). However, in certain cases the yield may actually increase if the electronically excited molecule can undergo a reaction with solvent molecules. Few experimental measurements of aqueous quantum yields exist, making attempts to constrain the change in quantum yield upon dissolution difficult. To obtain an upper estimate of aqueous phase photolysis, we will assume that the quantum yield in solution is less than or equal to the gas-phase quantum yield.

To constrain the ratio of the rate of removal due to gaseous and aqueous photolysis we consider a system with an air parcel containing liquid water droplets and a pollutant \( x \), which is in equilibrium between the gaseous and aqueous phase. The parcel has a liquid water content (LWC) defined in units of liquid water per volume of air. The ratio of the rate of gaseous photolysis and aqueous photolysis for species \( x \) is defined by the parameter \( Z \):

\[
Z = \frac{\frac{d n_{x+\nu}}{dt}}{\frac{d n_{x+\nu}}{dt}} = \left(R \cdot T \cdot LWC_v \cdot k_{Rx}\right)^{-1} \left(\frac{J_{\text{gas}}}{J_{\text{aq}}}\right)_{x+\nu}
\]

where \( n_{x+\nu} \) is the moles of species \( x \) undergoing photolysis in the gas or aqueous phase in a given volume of air, \( t \) represents time, \( R \) is the gas constant, \( T \) is temperature, \( LWC_v \) is the volume based liquid water content in units of volume of liquid water per volume of air, \( k_{Rx} \) is the effective Henry’s law constant for species \( x \) in units of molar concentration per pressure, \( J_{\text{gas}} \) and \( J_{\text{aq}} \) are the photolysis rate constants of species \( x \) in the gaseous and aqueous phase, respectively. The ratio of the rates does not depend on the concentration
of species \( x \) because we assume that the gaseous and aqueous forms are in equilibrium. Derivations of all significant equations are presented in Appendix A.

Equation (2) is a model for systems where the species of interest does not hydrate or dissociate in the aqueous phase. However, photolabile carbonyls hydrate in the aqueous phase to form the corresponding gem-diol (Bell and McDougall, 1960). Using a generic mono-aldehyde as an example, we can write:

\[
\text{RCCHO} + \text{H}_2\text{O} \xleftrightarrow{K_{\text{hyd}}} \text{RC(OH)}_2
\]  

(3)

where \( K_{\text{hyd}} \) is the hydration equilibrium constant. We define the variable \( \alpha_N \) as the fraction of the species that is present in form \( N \) upon dissolution. For a monocarbonyl, there may

be two forms in solution: the free molecule and the hydrated molecule. For a small dicarbonyl, there may be three forms in solution: the free molecule, the partially hydrated molecule, and the fully hydrated molecule. For a mono-carbonyl:

\[
\alpha_{\text{free}} = (K_{\text{hyd}} + 1)^{-1} \quad \text{and} \quad \alpha_{\text{hyd}} = (K_{\text{hyd}}^{-1} + 1)^{-1}
\]  

(4)

where \( \alpha_{\text{free}} \) is the fraction of the free molecule in solution and \( \alpha_{\text{hyd}} \) is the fraction of the hydrated molecule in solution. Aldehydes hydrate to a larger extent than ketones. For most ketones, only a small fraction of molecules exist in the gem-diol form (Bell and Gold, 1966). This has significant implications for photolysis as gem-diols do not absorb actinic UV radiation, and therefore are not photolabile in the atmosphere. We make a simplifying assumption that if we normalize the photolysis rate constant of the free form of species \( x \) to unity, the photolysis rate constant of the fully hydrated form will be null, and in the special case of a dicarbonyl, the normalized photolysis rate constant of the partially hydrated form will be half of the free form because one of the photoactive carbonyls has been converted to non-photoactive gem-diol form. We define this normalized photolysis rate constant as \( \beta_N \):

\[
\beta_{\text{free}} = 1, \quad \beta_{\text{hyd}} = 0,
\]

in the case of a dicarbonyl \( \beta_{\text{part-hyd}} = 1/2 \)  

(5)

Therefore, in the case of a hydrating carbonyl, \( x \), we can penalize the aqueous rate constant by the following factor, \( \gamma_x \):

\[
\gamma_x = \sum_N \alpha_N \cdot \beta_N
\]  

(6)

where \( \gamma_x \leq 1 \). Using this factor in Eq. (2) allows us to continue using the effective Henry’s law constant of species \( x \) despite the fact that \( x \) may hydrate into multiple forms in solution:

\[
Z = \frac{\frac{dC_{\text{gas}}}{dt}}{\frac{dC_{\text{aq}}}{dt}} = (R \cdot T \cdot \text{LWC}_V \cdot k_{H_x})^{-1} \left( \frac{j_{\text{gas}}^{x+h_v}}{j_{\text{aq}}^{x+h_v}} \gamma_x \right) \]  

(7)

Because \( \gamma_x \) is always less than unity, the maximum effective photolysis rate is reduced in the aqueous phase for all carbonyls.

Finally, Henry’s Law constants are strongly dependent on temperature. We perform calculations of \( Z \) at 298 K where most constants are measured to ensure a straightforward analysis. However, in Appendix B, we describe a way of correcting these equations for the temperature effects.

### 2.2 Comparison of aqueous and gaseous OH oxidation

The majority of atmospheric organics are vulnerable to OH oxidation in both the gaseous and aqueous phase. We aim to elucidate the relative significance of each of these processes for our selected set of compounds by calculating the ratio of the rate of OH oxidation in the gas and aqueous phase. The most significant difference between this analysis and the previous discussion is that the rate constants due to OH oxidation in the aqueous (Ervens et al., 2003b; Buxton et al., 1988; Monod et al., 2005) and gaseous phases (Sander et al., 2011; Atkinson et al., 2006) have been experimentally determined for many atmospheric organics. Even if gas phase OH oxidation rate constants have not been measured, structure activity relationships are well developed, and provide accurate rate constant estimates (Atkinson, 1987, 1988; Kwok and Atkinson, 1995). We define \( W \) as the ratio of the rate of gas phase OH oxidation of \( x \) and the rate of aqueous phase OH oxidation of \( x \). For non-ionizable species:

\[
W = \left( \frac{dC_{\text{gas}}}{dt} \right)_{k_{x+\text{OH}}} \quad \left( \frac{dC_{\text{aq}}}{dt} \right)_{k_{x+\text{OH}}} = \frac{dC_{\text{gas}}}{dC_{\text{aq}}} = \frac{k_{x+\text{OH}}^{\text{gas}}}{k_{x+\text{OH}}^{\text{aq}}} \]  

(8)

Where \( n_{x+\text{OH}} \) represents the moles of \( x \) undergoing oxidation by OH in the gas or aqueous phase, \( k_{x+\text{OH}} \) is the rate constant for OH oxidation with \( x \) in the aqueous or gas phase, \( C_{\text{gas}}^{\text{gas}} \) is the gas phase concentration of OH (moles of OH per volume of gas), and \( C_{\text{aq}}^{\text{aq}} \) is the aqueous phase concentration of OH (moles of OH per volume of water). When using this equation, the units of \( k_{x+\text{OH}} \) and \( C_{\text{OH}} \) must be consistent so that the ratios of rate constants and concentrations are unitless.

Equation (8) can be used for species that hydrate because hydroxyl radical rate constants are typically tabulated for a solution containing all aqueous forms of the parent molecule (Ervens et al., 2003b; Monod et al., 2005). However, for species that ionize, we must use a modified version of Eq. (8) to account for the dissimilar aqueous OH rate constants of the ionized and native form. With measured \( pK_a \) values, we can calculate the fraction of the acid HA that is present in the ionized form \( (f_{A^-}) \) and the native form \( (f_{HA}) \) at a given solution pH.

\[
f_{A^-} = \left( 1 + 10^{pK_a-\text{pH}} \right)^{-1} \quad f_{HA} = \left( 1 + 10^{\text{pH}-pK_a} \right)^{-1}
\]  

(9)
For a given pH and with OH rate constants for reaction with the acid, $k_{HA+OH}$, and its anion, $k_{A^-+OH}$, we can write $W$ as:

$$W = \frac{d^n_{HA+OH}}{dt^n} = \left( R \cdot T \cdot LWC_v \cdot k_{H_A} \right)^{-1} \left( h_{HA} \cdot k_{HA+OH} + f_{A^-} \cdot k_{A^-+OH} \right) \left( \frac{C_{gas}^{HA+OH}}{C_{OH}^{gas}} + \frac{C_{aq}^{HA+OH}}{C_{OH}^{aq}} \right)$$

where $C_{OH}^{gas}$ is the gas phase OH concentration as a function of temperature and pressure, $Y$ is a function of the solar zenith angle, SZA:

$$Y(SZA) = \frac{d^n_{HA+OH}}{dt^n} = \frac{k_{gas}^{HA+OH} \cdot C_{OH}(SZA)}{d^n_{HA+OH}}$$

where $C_{OH}(SZA)$ is the gas phase OH concentration as a function of the solar zenith angle in units that correspond to the units of $k_{A^-+OH}$. This concentration is described by the empirical power-law function relating the OH concentration to the photolysis rate of $O_3 \rightarrow O(1D)$ described in Rohrer and Berresheim (2006).

For species with measured absorption cross sections as a function of temperature and quantum yields as a function of temperature and/or pressure, we can go one step further and examine how $Y$ varies with atmospheric height and solar zenith angle.

$$Y(SZA, z) = \frac{d^n_{HA+OH}}{dt^n} = \frac{k_{gas}^{HA+OH}(T) \cdot C_{OH}(SZA, T)}{d^n_{HA+OH}}$$

where the change in temperature at a given height in the atmosphere, $z$, is given by:

$$-\frac{dT}{dz} = \Gamma$$

The parameter $\Gamma$ represents the lapse rate, for which we use the average value in the troposphere, 6.5 K km$^{-1}$ (Wallace and Hobbs, 2006). The actinic flux as a function of solar zenith angle and wavelength is predicted with the Tropospheric Ultraviolet and Visible (TUV) Radiation Model (ACD, Edition 4.4).

### 3 Results

#### 3.1 Comparison of aqueous and gaseous photolysis

Our comparison of aqueous and gaseous photolysis with Eqs. (2) and (7) provide an upper estimate to the extent of in-cloud photolysis for the following reasons. We perform this analysis using an air parcel at 298 K with a liquid water content of 0.5 g m$^{-3}$, approximately the largest value that is routinely measured in the atmosphere based on measurements of stratus and cumulus clouds (Seinfeld and Pandis, 1998; Hobbs, 1993). Aqueous processes are more significant in wetter air parcels as a larger fraction of each organic resides in the aqueous phase. The results of our analysis are not highly sensitive to our reasonable maximum LWC selection. If we chose a value of 0.25 g m$^{-3}$ instead, maximum rates of gaseous photolysis relative to rates of aqueous photolysis would only be doubled. This is unlikely to influence our conclusions, which are based on the order-of-magnitude estimates. In addition, aqueous quantum yields may be significantly depressed by solvent effects, thus slowing down the rate of aqueous photolysis relative to gaseous photolysis.

Figure 2 shows the results of this analysis for a series of atmospherically relevant, non-hydrating species, specifically peroxides and methyl nitrate. The ratio of gas and aqueous rate constants are shown on the axis of abscissas and the effective Henry’s law constants are shown on the axis of ordinates. Colors and diagonal isopleths show the magnitude of $Z$. Molecules are labeled in order of increasing $Z$. References for the Henry’s law parameters are presented in the Supplement. The x-axis intentionally spans a large range to provide an indication of the sensitivity of this analysis. Actual values should lie somewhere to the right ($J_{aqueous} < J_{gas}$) of our upper bound estimations.

Of the four molecules included in Fig. 2, only hydrogen peroxide has the potential for an appreciable aqueous photolysis removal rate relative to gas phase photolysis. With actual experimental data on the absorption cross sections and quantum yields of gaseous (Sander et al., 2011) and aqueous hydrogen peroxide (Chu and Anastasio, 2005), we can pinpoint the location of this molecule on the x-axis more precisely. The two inward arrows adjacent to the hydrogen peroxide estimation point (1) on Fig. 2 detailing the experimentally measured range in $J_{gas}/J_{aqueous}$ from a solar zenith angle of 0°–90° shows that the actual value is moderately below our upper-bound estimate. Gelencsér and Varga (2005) estimate that multi-phase reactions become important relative to gas-phase photo-oxidation processes when the effective Henry’s law constant is greater than $10^3$ M atm$^{-1}$. We find a similar threshold value in our analysis of the photolysis of non-hydrating species.

The series of atmospherically relevant, hydrating species is presented in Fig. 3. When experimental measurements of the Henry’s Law constants were unavailable (malonic acid, succinic acid, maleic acid, lactic acid, tartaric acid, and
glyceraldehyde), group contribution estimations were used (Saxena and Hildemann, 1996). Aqueous photolytic removal is potentially important in several of these species due to their large effective Henry’s law parameters. Upper-bound estimates are not all centered about the $J_{\text{gas}} / J_{\text{aqueous}} = 1$ line due to hydration into non-photolabile gem-diols in the aqueous phase. For example, at low concentrations, formaldehyde is almost exclusively in diol form (Dong and Dasgupta, 1986; Le Botlan et al., 1983). Therefore, formaldehyde (molecule #13 in Fig. 3) is represented by the arrow pointing towards $J_{\text{gas}} / J_{\text{aqueous}} = \infty$. Aqueous photolytic removal of glyceraldehyde ($Z \leq 1.0 \times 10^{-4}$) is potentially very significant. Aqueous removal of pyruvic acid ($Z \leq 2.7 \times 10^{-1}$) is likely significant and aqueous removal of gluteraldehyde ($Z \leq 6.2$), methyl glyoxal ($Z \leq 15$), glyoxal ($Z \leq 20$), glycolaldehyde ($Z \leq 22$), and glyoxylic acid ($Z \leq 29$) has the potential to be somewhat important. Ethers and compounds with only carboxylic acid functionality were not analyzed because they are not photolabile under typical tropospheric conditions (Chebbi and Carlier, 1996; Atkinson and Arey, 2003; Finlayson-Pitts and Pitts, 2000).

The y-axis of Figs. 2 and 3 provides a visual sensitivity analysis of our assumption of Henry’s law equilibrium.

For specific atmospheric conditions, previous researchers quantify the departure from equilibrium as the ratio of the observed aqueous phase concentration of species “x” and the predicted aqueous phase concentration with Henry’s law from the observed gas phase concentration (Winiwarter et al., 1994; Leriche et al., 2000):

$$\mathcal{R} = \frac{C_{\text{aq}}^{x}}{k_{Hx} \cdot C_{\text{gas}}^{x}} \quad (14)$$

where the units of $k_{Hx}$ correspond to the units of $C_{\text{gas}}^{x} / C_{\text{aq}}^{x}$ as to make $\mathcal{R}$ unitless. For specific atmospheric conditions where $\mathcal{R}$ values are known, one can multiply the ordinate values by these factors to obtain a more realistic estimate of $Z$. Since $\mathcal{R}$ is a function of local conditions, we only use the equilibrium values calculated from Henry’s law to make our analysis as general as possible.
3.2 Comparison of aqueous and gaseous OH oxidation

The ratio of the hydroxyl radical concentration in the gas and aqueous phase is a key uncertainty in the evaluation of \( W \) (the ratio of the rate of gas phase OH oxidation to aqueous phase OH oxidation). We use this OH ratio as one of our free parameters to allow for a sensitivity analysis, but present an educated guess to better constrain our results. For the gaseous OH concentration, we use the upper end of the measured and theoretical estimates suggested in Seinfeld and Pandis (1998), \( C_{\text{OH}}^{\text{daytime}} = 1 \times 10^7 \text{ molecules cm}^{-3} \) and \( C_{\text{OH}}^{\text{nighttime}} = 2 \times 10^5 \text{ molecules cm}^{-3} \). For the aqueous OH concentration in a cloud or fog droplet, we use the recommended values in the review by Ervens et al. (2011) that were determined with theoretical models (Ervens and Volkamer, 2010; Ervens et al., 2003a): \( C_{\text{OH}}^{\text{daytime}} = 10^{-13} \text{ M} \) and \( C_{\text{OH}}^{\text{nighttime}} = 10^{-14} \text{ M} \). The OH concentrations that we use are similar to the values assumed in Gelencsér and Varga (2005).

Figure 4 shows a graphical representation of Eq. (8) for a series of peroxides and ethers in an air parcel with a liquid water content of 0.5 g m\(^{-3}\) at 298 K. The ratio of the gas and aqueous phase OH concentrations (moles of OH per volume of air/moles of OH per volume of liquid), which we will refer to as the “OH ratio”, are shown on the x-axis. The product of the ratio of the gas and aqueous OH rate constants and the inverse Henry’s law constant (atm M\(^{-1}\)) is represented on the y-axis. Colors along with the diagonal isopleths indicate the magnitude of \( W \). Each species is represented by a labeled horizontal line bound by our daytime and nighttime estimates in the OH ratio. Species are labeled in order of increasing \( W \). A line showing the OH ratio in Henry’s Law equilibrium (rarely realized in practice because of the high OH reactivity) is also presented for perspective. All of the rate constants are referenced in the Supplement. Based on this analysis, we find that OH oxidation of hydrogen peroxide is important in the aqueous phase. Gas-phase OH oxidation strongly dominates the removal of ethyl peroxide, methyl peroxide, methyl tert-butyl ether, dimethyl ether, and ethyl ether.

Figure 5 is a continuation of this evaluation for a series of carbonyls under the same conditions. Aqueous oxidation dominates the removal of glyoxal compared to gas phase oxidation under the conditions of high LWC. Also, aqueous reaction with OH is likely a significant sink in the removal of methyl glyoxal compared to gas phase oxidation. Removal of formaldehyde by aqueous oxidation may be more important depending on the OH ratio. Gaseous oxidation dominates the removal of acetone, 2-butanone, acetaldehyde, butanal, and 8-propanol compared to aqueous oxidation by OH.

Figure 6 shows a similar analysis for a series of carboxylic acids using Eq. (10) without color contours for simplicity. Molecules are represented by labeled rectangles instead of

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**Fig. 4.** The ratio of the rate of gas phase OH oxidation to the rate of aqueous phase oxidation for a series of peroxides and ethers in an air parcel with a liquid water content of 0.5 g m\(^{-3}\) at 298 K. Labeled horizontal lines are bound by differences in \( C_{\text{OH}}^{\text{gas}}/C_{\text{OH}}^{\text{aqueous}} \) (defined as the moles of OH per volume of air/the moles of OH per volume of liquid) during different times of the day. The line demarcating OH at Henry’s law equilibrium is shown on the left side of the plot. Species are labeled in order of increasing \( W \): (1) hydrogen peroxide, (2) ethyl peroxide, (3) methyl peroxide, (4) methyl tert-butyl ether, (5) dimethyl ether, and (6) ethyl ether.

**Fig. 5.** The ratio of the rate of gas phase OH oxidation to the rate of aqueous phase oxidation for a series of carbonyls in an air parcel with a liquid water content of 0.5 g m\(^{-3}\) at 298 K. Labeled horizontal lines are bound by differences in \( C_{\text{OH}}^{\text{gas}}/C_{\text{OH}}^{\text{aqueous}} \) (defined as the moles of OH per volume of air/the moles of OH per volume of liquid) during different times of the day. The line demarcating OH at Henry’s law equilibrium is shown on the left side of the plot. Species are labeled in order of increasing \( W \): (1) glyoxal, (2) methyl glyoxal, (3) formaldehyde, (4) acetone, (5) 2-butanone, (6) acetaldehyde, (7) butanal, and (8) 8-propanol.
Fig. 6. The ratio of the rate of gas phase OH oxidation to the rate of aqueous phase oxidation for a series of acids in an air parcel with a liquid water content of 0.5 g m\(^{-3}\) at 298 K. Labeled rectangles are bound by differences in \(c_{OH}^{gas}/c_{OH}^{aqueous}\) (defined as the moles of OH per volume of air/the moles of OH per volume of liquid) during different times of the day and the pH of the cloud or fog droplet ranging between 2 and 6. The line demarcating OH at Henry’s law equilibrium is shown on the left side of the plot. Species are labeled in order of increasing equilibrium is shown on the left side of the plot. Species are labeled in order of increasing \(W\) at \(pH = 6\): (1) tartaric acid (R,R), (2) malic acid, (3) malonic acid, (4) lactic acid, (5) succinic acid, (6) pyruvic acid, (7) formic acid, (8) propanoic acid, (9) glyoxylic acid, (10) 2-methyl propanoic acid, (11) pentanoic acid, (12) hexanoic acid, (13) acetic acid, (14) 3-methyl butanoic acid, and (15) 2,2-dimethyl 1-propanoic acid.

The vertical coordinate on Figs. 4–6 can be simplified because gas and aqueous OH rate constants are related for individual species. Most rate constants lie within a factor of 3 of each other, and a line relating log \(k_{OH}^{gas}\) and log \(k_{OH}^{aqueous}\) (Haag and Yao, 1992). In light of this relationship, the dimensionless number, DN, introduced in this work, is a reasonable predictor for the magnitude of \(W\) for our range of assumed OH ratio where:

\[
DN = \frac{1}{10} (LWC \cdot R \cdot T \cdot k_H)^{-1}
\]

Figure 7 shows the predictive power of DN for the series of compounds investigated during daytime OH concentrations in an air parcel with a LWC = 0.5 g m\(^{-3}\) at 298 K. When DN \(\gg 1\), gaseous OH oxidation is more important than aqueous phase oxidation. Aqueous phase oxidation is more important when DN \(\ll 1\). Both reactions are potentially significant when DN is on the order of unity. This relationship holds with air parcels containing different liquid water contents or parcels with nighttime OH concentrations; a multi-panel plot showing \(W\) as a function of DN under different conditions is presented in the Supplement.
3.3 Comparison of gaseous OH oxidation and photolysis

For a given species, the ratio of the rate of gas phase OH oxidation to the rate of gaseous photolysis, $Y$, is dependent on the solar zenith angle, temperature, and pressure (see Eq. 12). We first investigate a series of 12 compounds with published quantum yield and absorption cross section measurements as a function of SZA at 298 K and 1 atm using Eq. (11). The references for these values are presented in the Supplement. Figure 8 shows how $Y$ varies with SZA for each of the selected compounds at sea level and 298 K. The width of the red swath is due to the uncertainty in the empirical function relating OH concentration to the rate of O$_3$ photolysis. The dashed line indicates where the rate of gas phase OH oxidation is equal to the rate of gaseous photolysis, $Y = 1$.

For formaldehyde, glyoxal, methyl glyoxal, and hydrogen peroxide, the relative importance of each of these removal processes is dependent on the SZA and the local OH concentration.

We also investigated this ratio as a function of SZA and altitude assuming a standard tropospheric lapse rate of $6.5 \text{ K km}^{-1}$ (Wallace and Hobbs, 2006) and a scale height of 7.5 km for four compounds with published quantum yields and absorption cross section measurements as a function of temperature and pressure and OH rate constants as a function of temperature. See the Supplement for these references. In this case, the actinic flux was predicted as a function of wavelength, SZA, and altitude with a surface albedo of 0.154 and an ozone column of 300 Dobson units. Figure 9 shows a complicated dependence of $Y$ on altitude and SZA for each of the four species investigated. This analysis at the bounds of the OH concentration uncertainty is presented in the Supplement.
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4 Atmospheric implications

With knowledge of the three ratios, W, Z, and Y, which characterize the significance of each of the four chemical sinks investigated, we can calculate the relative contribution of each of the sinks for a specific compound. The results of this analysis are presented in Fig. 10 for compounds with a full set of requisite data in an air mass with a LWC = 0.05 g m$^{-3}$ (characteristic of light clouds and fogs) and a LWC = 0.5 g m$^{-3}$ (characteristic of very dense clouds) at 298 K with SZA = 20°. The length of each bar shows the best-guess estimation of the chemical sink corresponding to its color. In the case of aqueous photolysis, the cyan bars show an upper bound estimate of its contribution to the removal of each specific species at 298 K. One of the major conclusions of this analysis is that even in conditions where aqueous processes are most favorable, LWC = 0.5 g m$^{-3}$, removal by aqueous photolysis remains a minor sink of the species investigated in Fig. 10. The only compounds with an appreciable aqueous photolysis contribution are hydrogen peroxide and methyl glyoxal. (However, aqueous photolysis may be important for more structurally complicated compounds that do not appear in Fig. 10. Published OH rate constants, gaseous absorption cross sections, and quantum yields are available only for relatively simple compounds.) At lower temperatures, the increased solubility of cloud and fog droplets slightly enhances this contribution, but even in an air mass with a LWC = 0.5 g m$^{-3}$ at 273 K, liquid photolysis of hydrogen peroxide accounts for less than 20% of its removal. However, photolysis of aqueous H$_2$O$_2$ is a significant source of aqueous OH; its rate of production is on the order of the rate of droplet scavenging from the gas phase (Zellner et al., 1990). A complete analysis detailing the magnitude of each of the chemical sinks at 273 K for species with published Henry’s Law temperature dependences is presented in Appendix B.

Our analysis of the ratio of gas phase photolysis and aqueous phase photolysis can help guide the selection of compounds for future experiments. Aqueous photolysis absorption cross sections and quantum yields are not well established in the literature, but may be important parameters for specific compounds. Glyceraldehyde and pyruvic acid were not included in Fig. 10 due to the lack of published gaseous absorption cross sections and quantum yields. However, based on the results of Fig. 3, we expect aqueous photolysis to be a significant sink for these compounds. For compounds that were not analyzed in this manuscript, the potential significance of aqueous photolysis can be determined with knowledge of the Henry’s law constant and the hydration equilibrium constants if hydration occurs in the aqueous phase.

For most acids in wet clouds, aqueous phase OH oxidation is more important that gas phase OH oxidation. For atmospherically relevant compounds without published aqueous rate constants, experiments designed to measure these values may prove to be fruitful. Since the ratio of the gas and aqueous phase OH removal rates is a strong function of LWC and the Henry’s Law constant, one can determine additional species for which experimental measurements are warranted.

We have chosen to treat aqueous phase photolysis and oxidation separately, but in certain cases, photolytic cleavage may produce OH and other free radicals that will react with the organic precursor. This will effectively increase the quantum yield of photolysis for the removal of the precursor beyond what would be measured if the liberated OH was being quenched by a competing reaction. However, in the absence of photo-catalytic processes or chain reactions, we expect this potential enhancement to be small (at most a factor of 2 or 3) as in the case of methyl peroxide (Epstein et al., 2012).

As indicated in Fig. 9, the ratio of gas phase oxidation to gas phase photolysis is a complicated function of altitude and SZA. Gas phase absorption cross section and quantum yield measurements at 298 K and 1 atm may not provide enough information to describe the atmospheric behavior of certain organics especially at higher SZA. For example, photolysis of methyl glyoxal, and hydrogen peroxide relative to oxidation is enhanced at high latitudes (SZA > 60°) and altitudes compared to the ratio at 1 atm. In the case of methyl glyoxal at solar zenith angles < 60°, photolysis relative to oxidation is suppressed at higher altitudes. In light of these complicated dependencies, we recommend caution when extrapolating room temperature and 1 atm measurements of photolysis quantum yields and absorption cross sections.

We have assumed that the organic molecules in our analysis do not interact with each other. While this assumption is
Fig. 10. Estimated magnitudes of each of the 4 chemical sinks investigated for compounds with sufficient published data. All acids are assumed to be in cloud or fog droplets at a pH = 6. Calculations were performed with a solar zenith angle of 20°. The right panel details selected species in a parcel with a LWC = 0.5 g m⁻³ at 298 K and represents the upper limit for the contribution of aqueous processes in all situations at 298 K. The left panel shows the contribution of each of the four sinks in an air parcel with a more realistic LWC = 0.05 g m⁻³ at 298 K. The contributions of aqueous processes in this panel are also an upper estimate for the selected LWC.

needed to constrain the complexity of interacting molecules and draw conclusions, it may not be appropriate in all conditions. For example, in a polluted environment, organics may compete for aqueous phase OH radicals in a cloud or fog droplet. A more complicated analysis is required to capture these effects.

Appendix A

Equation derivations

A1 Derivation of Z

We start with the rate laws for photolysis in the gas and aqueous phase:

$$\frac{dC_{x+h\nu}^{\text{gas}}}{dt} = -J_{x+h\nu}^{\text{gas}} C_{x}^{\text{gas}}$$  \hspace{1cm} (A1)

$$\frac{dC_{x+h\nu}^{\text{aq}}}{dt} = -J_{x+h\nu}^{\text{aq}} C_{x}^{\text{aq}}$$  \hspace{1cm} (A2)

where $C_{x}^{\text{gas}}$ is the gas phase concentration of species x and $C_{x}^{\text{aq}}$ is the aqueous phase concentration of x. We transform concentration units into a molar basis with the volume of the air parcel ($V_{\text{air}}$) and the volume of water inside the air parcel ($V_{\text{H}_2\text{O}}$).

$$\frac{dn_{x+h\nu}^{\text{gas}}}{dt} = V_{\text{air}} \cdot \frac{dC_{x+h\nu}^{\text{gas}}}{dt}$$  \hspace{1cm} (A3)

$$\frac{dn_{x+h\nu}^{\text{aq}}}{dt} = V_{\text{H}_2\text{O}} \cdot \frac{dC_{x+h\nu}^{\text{aq}}}{dt}$$  \hspace{1cm} (A4)

We define Z as the ratio between these two molar rates:

$$Z = \frac{\frac{dn_{x+h\nu}^{\text{gas}}}{dt}}{\frac{dn_{x+h\nu}^{\text{aq}}}{dt}}$$  \hspace{1cm} (A5)

The ratio of air and water volumes can be simplified with a volumetric cloud liquid water content (LWCᵥ, volume of H₂O/volume of air parcel):

$$Z = \frac{V_{\text{air}}}{V_{\text{H}_2\text{O}}} \cdot \frac{\frac{dC_{x+h\nu}^{\text{gas}}}{dt}}{\frac{dC_{x+h\nu}^{\text{aq}}}{dt}}$$  \hspace{1cm} (A6)
Fig. A1. Estimated magnitudes of each of the 4 chemical sinks investigated for compounds with sufficient published data. All acids are assumed to be in cloud or fog droplets at a pH = 6. Calculations were performed with a solar zenith angle of 20°. The right panel details selected species in a parcel with a LWC = 0.5 g m\(^{-3}\) at 273 K and represents the upper limit for the contribution of aqueous processes in all situations at 273 K. The left panel shows the contribution of each of the 4 sinks in an air parcel with a more realistic LWC = 0.05 g m\(^{-3}\) at 273 K. The contributions of aqueous processes in this panel are also an upper estimate for the selected LWC.

where the volumetric liquid water volume is defined as:

\[
LWC_v = \frac{LWC}{\rho_{H_2O}} \quad (A7)
\]

The density of water is represented by \(\rho_{H_2O}\) and LWC is the liquid water content in units of mass of water per volume of air parcel. We now introduce the Henry’s law constant to describe the ratio of the aqueous phase concentration of \(x\) (\(C_{aQ}^x\)) and the partial pressure of \(x\) (\(p_{x}^{gas}\)).

\[
k_H = \frac{C_{aQ}^x}{p_{x}^{gas}} \quad (A8)
\]

Converting pressure to concentration units with the ideal gas constant and temperature yields the equilibrium ratio of gas and aqueous phase concentration of \(x\):

\[
\frac{C_{x}^{gas}}{C_{x}^{aq}} = \frac{1}{k_H RT} \quad (A9)
\]

Our final working equation to describe the ratio of the rate of gas phase photolysis to the rate of aqueous phase photolysis is:

\[
Z = \frac{1}{LWC_v \cdot k_H \cdot R \cdot T} \left( \frac{J_{x+h\nu}^{gas}}{J_{x+h\nu}^{aq}} \right) \quad (A10)
\]

A2 Derivation of \(W\)

We start with the rate laws for oxidation of species \(x\) in the gas and aqueous phase by OH:

\[
\frac{dC_{x+OH}^{gas}}{dt} = -k_{x+OH}^{gas} C_{x}^{gas} C_{OH} \quad (A11)
\]

\[
\frac{dC_{x+OH}^{aq}}{dt} = -k_{x+OH}^{aq} C_{x}^{aq} C_{OH} \quad (A12)
\]

Converting to molar units:

\[
\frac{dn_{x+OH}^{gas}}{dt} = V_{air} \cdot \frac{dC_{x+OH}^{gas}}{dt} \quad (A13)
\]

\[
\frac{dn_{x+OH}^{aq}}{dt} = V_{H_2O} \cdot \frac{dC_{x+OH}^{aq}}{dt} \quad (A14)
\]

We define \(W\) as the ratio between the loss rate due to gas phase oxidation and the loss rate due to aqueous oxidation:

\[
W \equiv \frac{\frac{dn_{x+OH}^{gas}}{dt}}{\frac{dn_{x+OH}^{aq}}{dt}} \quad (A15)
\]
We define the ratio of these rates in molar units as

\[ W = \frac{V_{\text{air}}}{V_{\text{H}_2\text{O}} \cdot \frac{dC_{x+\text{OH}}^{\text{gas}}}{dr}} \cdot \frac{k_{x+\text{OH}}^{\text{gas}} C_{x}^{\text{gas}}}{k_{x+\text{OH}}^{\text{aq}} C_{x}^{\text{aq}}} \]  

(A16)

Inserting the Henry’s law constant to describe the ratio of gas and aqueous phase concentrations yields the final working equation:

\[ W = \frac{1}{\text{LWC}_v \cdot k_H \cdot R \cdot T} \cdot \frac{k_{x+\text{OH}}^{\text{gas}} C_{x}^{\text{gas}}}{k_{x+\text{OH}}^{\text{aq}} C_{x}^{\text{aq}}} \]  

(A17)

### A3 Derivation of Y

We start with rate laws describing gas phase oxidation by OH and gas phase photolysis

\[ \frac{dc_{x+\text{OH}}^{\text{gas}}}{dr} = -k_{x+\text{OH}}^{\text{gas}} C_{x}^{\text{gas}} C_{x}^{\text{gas}} \]  

(A18)

\[ \frac{dc_{x+\text{hv}}^{\text{gas}}}{dr} = -J_{x+\text{hv}}^{\text{gas}} C_{x}^{\text{gas}} \]  

(A19)

We define the ratio of these rates in molar units as Y:

\[ Y = \frac{dc_{x+\text{hv}}^{\text{gas}}}{dr} = \frac{dc_{x+\text{OH}}^{\text{gas}}}{dr} \]  

(A20)

Since both rates describe the gas phase, the ratio of molar rates is equal to the ratio of concentration-based rates:

\[ \frac{dc_{x+\text{hv}}^{\text{gas}}}{dr} = \frac{dc_{x+\text{OH}}^{\text{gas}}}{dr} \]  

(A21)

Inserting Eqs. (A18) and (A19) into Eq. (A21) yields our final working equation for Y:

\[ Y = \frac{k_{x+\text{OH}}^{\text{gas}} C_{x}^{\text{gas}}}{J_{x+\text{hv}}^{\text{gas}}} \]  

(A22)

### Appendix B

#### Chemical sink analysis at different temperatures

Z as a function of temperature can be calculated from Z at 298 K. Both Henry’s Law constants and for carboxyls, hydration equilibrium constants are sensitive to temperature. When temperature dependent Henry’s Law data are available, they are typically fit to the function (Sander et al., 2011):

\[ \ln k_{H_x} = A + B \cdot T^{-1} + C \cdot \ln(T) \]  

(B1)

where A, B, and C are fitting parameters. Z calculated at T1 can be adjusted to temperature T2 with the unitless correction factor, CF:\n
\[ \text{CF}_{T_1 \rightarrow T_2} = e^{B \left( \frac{1}{T_1} - \frac{1}{T_2} \right) + C \cdot \ln \left( \frac{T_1}{T_2} \right)} \]  

(B2)

Hydration equilibrium constants are also a function of temperature. For a di-carbonyl, Z calculated at T1 can be adjusted to temperature T2 with the unitless correction factor, CF_{Khyd}, which accounts for this temperature dependence:

\[ \text{CF}_{T_1 \rightarrow T_2} = e^{\frac{\Delta S_{\text{hyd}}}{R} \left( \frac{1}{T_1} - \frac{1}{T_2} \right) + 1} \]  

(B3)

where \( \Delta S_{\text{hyd}} \) is the change in entropy upon hydration and \( \Delta H_{\text{hyd}} \) is the change in enthalpy upon hydration. Z at T2 is the product of the correction factors and Z at T1:

\[ Z(T_2) = \text{CF}_{T_1 \rightarrow T_2} \cdot \text{CF}_{K_{\text{hyd}}} \cdot \text{CF}_{Z(T_1)} \]  

(B4)

The factor W can also be corrected for other temperatures by accounting for differences in both the Henry’s Law parameter and the ratio of the gaseous and aqueous rate constants. W calculated at T1 can be adjusted to its value at T2:

\[ W(T_2) = \text{CF}_{T_1 \rightarrow T_2} \cdot \text{CF}_{Z(T_1)} \cdot W(T_1) \]  

(B5)

where CF_{k} accounts for the temperature dependence on the rate constants:

\[ \text{CF}_{T_1 \rightarrow T_2} = \exp \left[ \left( \frac{1}{T_2} - \frac{1}{T_1} \right) \left( \frac{E_A^{\text{aq}}}{R} - \frac{E_A^{\text{gas}}}{R} \right) \right] \]  

(B6)

and \( E_A^{\text{aq}} \) is the activation energy of the OH oxidation reaction in the aqueous phase and \( E_A^{\text{gas}} \) is the activation energy of the gas phase reaction.

Using Eqs. (B4) and (B5), we can predict Z and W factors at 273 K from the values calculated at 298 K. The correction factor used to account for the changing gas solubility parameters at different temperatures, CF_{Hlaw}, is dependent on the fitting parameters used to correlate Henry’s Law parameters with temperatures (see Eq. B2). For most compounds, these fitting parameters are available in the literature (Sander et al., 2011; Sander, 1999). Typically C is small, and can be safely set to 0. For the compounds investigated, B ranges from 4700 to 7700 K, yielding correction factors of 0.04 to 0.004 going from temperatures of 298 to 273 K. This correction factor is significant and could change the magnitude of Z by a few orders of magnitude while changes in the percent hydrated have a relatively small effect. Therefore, with the lack of temperature dependent hydration equilibrium data for many relevant compounds and the relatively small effect on Z, we only address changes due to increased Henry’s Law parameters at depressed temperatures.
W can also be corrected by CF, which captures the temperature dependence in the gas and aqueous OH rate constants. However, the availability of temperature dependent rate data for the compounds investigated is limited. Fortunately, CF tends to be on the order of 1 for reasonably small temperature changes. For example, when the activation energy of the aqueous and gaseous reactions differ by 2500 K, (an upper estimate from a selection of published OH rate constants for several water soluble compounds) CF = 0.5. Overestimating or underestimating our value of W by not correcting by this factor will not significantly change the results of our analysis. The lack of published activation energies and the limited influence temperature changes have on CF motivate the assumption that CF is unity. Y may also depend on temperature. Absorption cross section and quantum yield data are extremely limited, but for the four species for which we determined Y as a function of SZA and altitude, Y was not a strong function of temperature. Therefore, we assume that between 298 K and 273 K the magnitude of Y will not change as to affect the results of our analysis.

Figure A1 shows the magnitude of each chemical sink for the compounds with published Henry’s Law temperature dependences at 273 K. In general, aqueous processes play a more significant role due to their enhanced solubility, but for the species investigated, aqueous photolysis still remains a minor sink.

Appendix C

Nomenclature and units

Variable, typical units, and definition are presented in their order of appearance in the text.

- \( J \) [s\(^{-1}\)] photolysis rate constant
- \( F_a \) [cm\(^{-2}\) s\(^{-1}\) nm\(^{-1}\)] actinic flux
- \( \sigma \) [cm\(^2\)] absorption cross section
- \( \Phi \) [unitless] quantum yield of photolysis
- \( \lambda \) [nm] wavelength of light
- \( \text{LWC} \) \( \left[ \frac{\text{mol}_{H_2O}}{m_{\text{air}}} \right] \) Liquid water content in terms of the mass of water per volume of air
- \( Z \) [unitless] ratio of the rate of gas phase photolysis to the rate of aqueous phase photolysis
- \( n_x \) [mol] moles of species \( x \)
- \( R \) [L atm \( \text{mol} \text{K}^{-1} \)] or \( \frac{J}{\text{mol} \text{K}^{-1}} \) ideal gas constant
- \( T \) [K] temperature
- \( k_{Hx} \) \( \left[ \frac{\text{mol}_{H_2O}}{\text{atm}} \right] \) Henry’s Law constant of species \( x \)
- \( J_{x+h\nu}^{\text{gas}} \) [s\(^{-1}\)] photolysis rate constant of species \( x \) in the gas phase
- \( J_{x+h\nu}^{\text{aq}} \) [s\(^{-1}\)] photolysis rate constant of species \( x \) in the aqueous phase
- \( t \) [s] hydration equilibrium constant
- \( \alpha_N \) [unitless] mole fraction of form \( N \) in the aqueous phase upon hydration
- \( \alpha_{\text{free}} \) [unitless] mole fraction of the free form (unhydrated) in solution at equilibrium
- \( \alpha_{\text{hyd}} \) [unitless] mole fraction of the hydrated form in solution at equilibrium
- \( \beta_N \) [unitless] normalization constant for the photolysis rate constant of form \( N \)
- \( \beta_{\text{free}} \) [unitless] normalization constant for the photolysis rate constant of the free form
- \( \beta_{\text{hyd}} \) [unitless] normalization constant for the photolysis rate constant of the hydrated form
- \( \beta_{\text{part,hyd}} \) [unitless] normalization constant for the photolysis rate constant of the partially hydrated form
- \( \gamma_x \) [unitless] constant used to penalize the aqueous photolysis rate of species \( x \) for the presence of non-photolabile gem-diol groups
- \( W \) [unitless] ratio of the rate of gas phase reaction with OH to the rate of aqueous phase reaction with OH
- \( k_{x+OH}^{\text{gas}} \) \( \left[ \frac{\text{mol}^{-1}}{\text{mol}^{-1}} \right] \) gas phase rate constant for the reaction of \( x \) with OH
- \( k_{x+OH}^{\text{aq}} \) \( \left[ \frac{\text{mol}^{-1}}{\text{mol}^{-1}} \right] \) aqueous phase rate constant for the reaction of \( x \) with OH

LWC \( \left[ \frac{\text{mol}_{H_2O}}{m_{\text{air}}} \right] \) volume based liquid water content (water volume per volume of air)
C_{OH}^{\text{gas}} \left[ \text{mol} / \text{m}^3 \right] \quad \text{concentration of OH in the gas phase}

C_{OH}^{\text{aq}} \left[ \text{mol} / \text{m}^3 \right] \quad \text{concentration of OH in the aqueous phase}

C_x^{\text{gas}} \left[ \text{mol} / \text{m}^3 \right] \quad \text{concentration of species } x \text{ in the gas phase}

C_x^{\text{aq}} \left[ \text{mol} / \text{m}^3 \right] \quad \text{concentration of species } x \text{ in the aqueous phase}

f_A \quad \text{mole fraction of the acid that is in the ionized form at equilibrium}

f_{HA} \quad \text{mole fraction of the acid that is in the non-ionized form at equilibrium}

Y \quad \text{ratio of the rate of gas phase reaction with OH to the rate of gas phase photolysis}

SZA \quad \text{deg latitude} \quad \text{solar zenith angle}

z \quad [\text{km}] \quad \text{altitude}

Γ \quad [\text{Kkm}^{-1}] \quad \text{average lapse rate}

Θ \quad [\text{unitless}] \quad \text{ratio of the observed aqueous phase concentration to the predicted aqueous phase concentration using Henry’s law and the observed gas phase concentration}

DN \quad [\text{unitless}] \quad \text{parameter (defined in this work) that indicates the ratio of the rate of gas phase reaction with OH relative to the rate of aqueous phase reaction with OH}

CF_{Hlaw}^{T_1 \rightarrow T_2} \quad [\text{unitless}] \quad \text{correction factor used to adjust } Z \text{ and } W \text{ due to the temperature dependance of Henry’s Law constants}

CF_{hyd}^{T_1 \rightarrow T_2} \quad [\text{unitless}] \quad \text{correction factor used to adjust } Z \text{ and } W \text{ due to the temperature dependance of hydration equillibrium}

ΔS_{hyd} \quad [\text{Jmol}^{-1} \text{K}^{-1}] \quad \text{change in entropy upon hydration}

ΔH_{hyd} \quad [\text{Jmol}^{-1}] \quad \text{change in enthalpy upon hydration}

CF_k^{T_1 \rightarrow T_2} \quad [\text{unitless}] \quad \text{correction factor used to adjust } W \text{ due to the temperature dependance of OH rate constants}

E_A^{\text{aq}} \quad [\text{Jmol}^{-1}] \quad \text{activation energy of species } x \text{ in the aqueous phase}

E_A^{\text{gas}} \quad [\text{Jmol}^{-1}] \quad \text{activation energy of species } x \text{ in the gas phase}

ρ_{H_2O} \quad [\text{kg} \text{H}_2\text{O} / \text{m}^3] \quad \text{the density of water}

V_{air} \quad [\text{m}^3 / \text{air}] \quad \text{air volume}

V_{H_2O} \quad [\text{m}^3 / \text{H}_2\text{O}] \quad \text{water volume}

Supplementary material related to this article is available online at: http://www.atmos-chem-phys.net/12/8205/2012/acp-12-8205-2012-supplement.pdf.

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