Secondary organic aerosol formation in cloud droplets and aqueous particles (aqSOA): a review of laboratory, field and model studies

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Abstract

Progress has been made over the past decade in predicting secondary organic aerosol (SOA) mass in the atmosphere using vapor pressure-driven partitioning, which implies that SOA compounds are formed in the gas phase and then partition to an organic phase (gasSOA). However, discrepancies in predicting organic aerosol oxidation state, size and product (molecular mass) distribution, relative humidity (RH) dependence, color, and vertical profile suggest that additional SOA sources and aging processes may be important. The formation of SOA in cloud and aerosol water (aqSOA) is not considered in these models even though water is an abundant medium for atmospheric chemistry and such chemistry can form dicarboxylic acids and “humic-like substances” (oligomers, high-molecular-weight compounds), i.e., compounds that do not have any gas phase sources but comprise a significant fraction of the total SOA mass. There is direct evidence from field observations and laboratory studies that organic aerosol is formed in cloud and aerosol water, contributing substantial mass to the droplet mode.

This review summarizes the current knowledge on aqueous phase organic reactions and combines evidence that points to a significant role of aqSOA formation in the atmosphere. Model studies are discussed that explore the importance of aqSOA formation and suggestions for model improvements are made based on the comprehensive set of laboratory data presented here. A first comparison is made between aqSOA and gas-SOA yields and mass predictions for selected conditions. These simulations suggest that aqSOA might contribute almost as much mass as gasSOA to the SOA budget, with highest contributions from biogenic VOC emissions in the presence of anthropogenic pollutants (i.e., NOx) at high relative humidity and cloudiness. Gaps in the current understanding of aqSOA processes are discussed and further studies (laboratory, field, model) are outlined to complement current data sets.
1 Introduction

A role for organic aerosols (OA) in the climate system has been recognized over several decades and much progress has been made in characterizing OA sources, transformations and radiative impacts. Globally, about 20% of the total OA mass is directly emitted (primary organic aerosols, POA) (Kanakidou et al., 2005; Spracklen et al., 2011). Traditionally, secondary organic aerosol (SOA) formation is thought to occur by gas phase photochemistry followed by absorption of semi-volatile products into preexisting OA mass (“gasSOA”, Fig. 1) (Pankow, 1994a, b; Odum et al., 1996). Recently, it has been recognized that a substantial fraction (50–75%, Shrivastava et al., 2006) of POA is semivolatile, evaporates when the plume becomes more dilute, and is then available in the gas phase to take part in photochemical reactions (Fig. 1) (Robinson et al., 2007). This material is like SOA in some ways and like POA in others; its atmospheric concentrations can be reduced with primary PM controls, but it has the physicochemical properties of SOA. Whereas previous models tended to underestimate SOA mass (e.g., DeGouw et al., 2005; Heald et al., 2006; Volkamer et al., 2006; Jimenez et al., 2009), the treatment of POA as semivolatile improved OA mass prediction (e.g., Hodzic et al., 2010; Lee-Taylor et al., 2011), but the temporal variability (e.g., Murphy and Pandis, 2009; Jathar et al., 2011) and vertical profile of OA remained poorly captured (Carlton et al., 2008). While currently it is thought that these semivolatile compounds add to SOA mass by further oxidation in the gas phase and re-condensation into particulate organic matter, it is at least as likely that this oxidized material, as well as other water-soluble products of gas-phase photochemistry, dissolve into an aqueous phase where it forms SOA through further oxidation (aqSOA, Fig. 1).

More than a decade ago, it was hypothesized that organic compounds are oxidized in the aqueous phase of cloud and fog droplets and products remain in the particle phase upon water evaporation (Blando and Turpin, 2000). More recently, it was hypothesized that chemical processes in the aqueous phase of hygroscopic particles (aerosol water) can also efficiently contribute to aqSOA mass (Volkamer et al., 2007).
Aqueous phase processes start with water-soluble, polar precursors and form SOA that is more oxygenated than gasSOA as they lead to functionalization or accretion (e.g., acid formation, oligomerization) rather than to breakage of the carbon structure as in gas phase reactions. Laboratory studies have shown that at low, cloud-relevant aqueous phase concentrations, organic acids (e.g., oxalate) are formed from small aldehydes and related compounds. At the higher solute concentrations associated with aerosol water, “high-molecular-weight compounds” (HMWC) tend to preferentially form (Lim et al., 2010). As discussed below, the organic fractions of atmospheric aerosol particles have properties consistent with such aqueous laboratory studies.

While for gasSOA, the amount of preexisting organic mass determines the extent to which condensable species are added to particles (“partitioning theory”, Odum et al., 1996), aqSOA formation depends on the amount of cloud, fog or aerosol water. Globally, aerosol water exceeds dry aerosol mass (ammonium, sulfate, nitrate) by a factor of 2–3 (Meng et al., 1995; Wang et al., 2008); thus, the accessibility of semivolatile (or low volatility), water-soluble species to an aqueous phase might be greater than to an organic phase. Freshly emitted, hydrophobic particles are converted into hygroscopic, water-containing particles within several hours (∼10’s km) (Ervens et al., 2010; Riemer et al., 2010); thus, regions with only hydrophobic particles with very little aerosol water are very limited.

AqSOA formation will not only affect the total SOA mass predictions but also impacts aerosol properties that translate into the aerosol direct and indirect effects in more ways than just increased aerosol mass burden. Condensation of low volatility species (gasSOA) and volume phase reactions (aqSOA) modify aerosol size distributions in different ways, the former adding mass to the condensation mode and, with the latter adding mass to the droplet mode (John et al., 1990). High-resolution analytical techniques capable of molecular characterization of OA mass suggest that a substantial fraction of SOA is composed of material that is more highly oxidized (higher oxygen-to-carbon (O/C) ratio) than gasSOA formed in dry smog chamber experiments (Aiken et al., 2008; Ng et al., 2010), whose yields are the basis for predicting SOA in models. Models that
include the prediction of O/C ratios often underestimate the degree of oxygenation of SOA which points to a biased prediction of the product distribution and related SOA properties (e.g., Jathar et al., 2011). SOA formation through aqueous chemistry could explain the high O/C ratios as observed in atmospheric organic aerosol. The O/C ratio can be related to SOA hygroscopicity which affects interaction with water-vapor, influencing haze and cloud drop formation, and the scattering/absorbing of radiation. There is some evidence that aqSOA can comprise light absorbing organics (e.g., Gelencser et al., 2003; Shapiro et al., 2009). The addition of different types of organics (water-soluble aqSOA, as opposed to more hydrophobic gasSOA) also might lead to phase separation within a single particle (Bertram et al., 2011). Because aqSOA is more oxygenated and water-soluble, we also expect its atmospheric lifetime to be shorter than gasSOA as it is more readily removed by wet deposition (Fu et al., 2008, 2009).

A host of organic acids has been observed in ambient particles, with oxalate globally ubiquitous throughout the tropospheric column (e.g., Kawamura and Sakaguchi, 1999; Kawamura et al., 2003). The atmospheric dynamics of oxalate suggests that it is predominantly secondary, and yet its abundance has not been explained by gas phase chemistry. Large multifunctional organic compounds including oligomeric SOA constituents and organics that include heteroatoms (e.g., sulfur, nitrogen; Blando et al., 1998; Romero and Oehme, 2005; Surratt et al., 2007a; Stone et al., 2009) comprise the largest component of OA (Fuzzi et al., 2001). Due to their structural similarities with humic substances, these compounds are often called “humic-like substances (HULIS)”; however, it has been noted that their similarities to humic compounds are limited (Graber and Rudich, 2006) and thus in the present review these compounds are rather termed HMWC. Substantial amounts of HMWC have been observed to form in biomass burning plumes (Miyazaki et al., 2009b), rural areas (Lin et al., 2010b) and urban environments (Krivacsy et al., 2008). While it seems possible that HMWC might be also formed in an organic phase of particles, the high polarity and water-solubility of this material rather suggests their formation in an aqueous phase (Fuzzi et al., 2001). Based on their atmospheric dynamics these compounds are believed to be
predominantly secondary and not directly emitted, but their formation processes are not captured by gasSOA models. Their observation in a wide variety of environments suggests the ubiquitous occurrence of aqueous phase processing.

In this review, we document the substantial evidence that aqueous phase chemistry as observed in laboratory studies (Sect. 2) is important in the atmosphere. Evidence of aqSOA from field studies together with model predictions is discussed (Sect. 3). For selected precursors and conditions, predicted aqSOA and gasSOA yields and masses are compared (Sect. 4). Finally, major uncertainties in determining aqSOA formation are assessed which should be considered in future studies (Sect. 5).

2 Laboratory experiments of aqSOA formation

2.1 Photochemical aqueous phase reactions

2.1.1 Carbonyl compounds (≤C₅)

The OH radical represents the main oxidant of organic compounds in the aqueous phase of cloud droplets (Ervens et al., 2003a). Kinetic data for aqueous phase reactions with OH are available or can be estimated based on structure-reactivity-relationships (e.g., Ervens et al., 2003b; Herrmann, 2003; Monod et al., 2005; Monod and Doussin, 2008; Minakata et al., 2009; Herrmann et al., 2010). Most OH reactions with organic compounds are near the diffusion limit in the aqueous phase \((k = 10^9 - 10^{10} \text{ M}^{-1} \text{ s}^{-1})\). OH concentration of \(\sim 10^{-13} \text{ M}\) and \(10^{-12} \text{ M}\) in cloud and aerosol water, respectively, have been estimated based on model studies (Ervens et al., 2003a; Ervens and Volkamer, 2010) since there are no measurements in ambient aerosol and/or cloud water available. Thus, a first-order rate constant \((k [\text{OH}])\) of \(< 10^{-3} \text{ s}^{-1}\) in cloud water and \(< 10^{-2} \text{ s}^{-1}\) in aerosol water can be estimated for converting water-soluble compounds into aqSOA mass. This estimate is based on the assumption of identical transport parameters into, on and inside of cloud droplets and aqueous particles.
C₂ and C₃ carbonyl compounds (e.g., glyoxal, glycolaldehyde, glyoxylic acid, glycolic acid, methylglyoxal, pyruvic acid) represent first- and/or second-generation gas-phase oxidation products of both anthropogenic (e.g., aromatics, acetylene) and biogenic (isoprene) compounds. First generation products of isoprene include methacrolein (MACR) and methylvinyl ketone (MVK), which are less soluble but formed with higher yields. Henry’s law constants (K_H) for the C₂–C₃ carbonyl compounds are in the range of 10³–10⁵ M atm⁻¹; those for MVK and MACR are much smaller in neutral or moderately acidic solution (K_H(MVK) = 41 M atm⁻¹, K_H(MACR) = 6.5 M atm⁻¹ (Iraci et al., 1999)), which suggests that only a small fraction of these compounds are present in the aqueous phase of particles.

The chemistry of carbonyl compounds in the aqueous phase differs from that in the gas phase in part because these compounds hydrate. Aqueous-phase product formation also depends on the concentrations of organics in solution. Glyoxal has been used as a model compound to explore these differences. The aqueous OH-radical oxidation of glyoxal upon hydration has been discussed in recent detailed studies (Tan et al., 2009; Ervens and Volkamer, 2010; Lim et al., 2010). In brief, in dilute solutions (resembling solute concentrations as in cloud water, e.g., Igawa et al., 1989; Munger et al., 1995) with [Glyoxal]_aq < 1 mM, the OH radical oxidation of glyoxal yields glyoxylic acid, which is further oxidized to oxalic acid (Karpe Vel Leitner and Doré, 1997) (Table 1). Carlton et al. (2007) first observed HMWC in glyoxal photooxidation experiments at organic concentrations that were higher than in cloud water but likely less than in aerosol water ([Glyoxal]_aq ≤ mM) (Tan et al., 2009). For methylglyoxal and its direct oxidation product, pyruvic acid, a qualitatively similar product pattern has been found at organic cloud-water concentrations with the formation of small acids (oxalic, glyoxylic, acetic) (Tan et al., 2010). At higher concentrations (mM), the same transition to HMWC formation is seen from both pyruvic acid and methylglyoxal (Altieri et al., 2008; Perri et al., 2009) with oligomeric series with fragment patterns that differ by ΔM = 72 (C₃H₄O₂) (Tan et al., 2010, 2011).
Aqueous phase oxidation experiments of isoprene, MACR and MVK in the presence of OH show very similar product patterns. An explicit mechanism suggests that the oxidation of these C5 carbonyl compounds is closely intertwined with the smaller, more soluble ones that can account for a large fraction of the aqSOA observed in laboratory studies; HMWC are assumed to be intermediates in this reaction scheme (Zhang et al., 2010; Huang et al., 2011). In ambient aerosol particles and cloud droplets, the observed bulk processes will have only a minor importance due to the limited solubility of these aqSOA precursors; however, in the same study it was discussed that surface reactions could occur and contribute more to organic processing and aqSOA formation. Aqueous oxidation experiments with MACR and MVK show that small acids comprise up to 8.8 % and 23.8 %, respectively, of the aqSOA yield (defined as mass concentration of aerosol after 7 h, related to the initial aqueous phase MACR and MVK concentrations, Zhang et al., 2010). The value for MACR is in good agreement with experiments over longer time scales that reported aqSOA yields of 2–12 % with highest values after 22 h (ElHaddad et al., 2009).

The direct reaction of MVK and MACR with ozone is probably not a competitive pathway as compared to the much faster OH-initiated reactions, but it might act as an important source of oxidants (e.g., H2O2) (Chen et al., 2008). While H2O2 itself reacts only slowly with organic carbonyl compounds (Stefan and Bolton, 1999), it might be a precursor for OH (either by photolysis, or by Fenton reaction with Fe2+ or Cu+) and thus could provide an additional radical source in the aqueous phase that could start OH-initiated processes.

It has been shown that SOA formation rates in laboratory aerosol experiments are 2–3 orders of magnitude faster than those that were predicted based on the OH reaction using uptake and reaction parameters as for dilute aqueous solutions (Ervens and Volkamer, 2010) which suggests that the overall aqSOA production rate (per volume of water) is higher in aerosol than in cloud water. While this difference in glyoxal reactivity cannot be solely ascribed to faster kinetics due to ionic strength effects (Herrmann, 2003), it is rather possible that additional radical pathways in aqueous particles occur.
that are largely suppressed in dilute aqueous phase, but lead to accretion products of organic radicals at higher organic concentrations. Since observed aqSOA formation both scales with bulk aerosol water mass and particle surface, this additional pathway has been parameterized by either a bulk aqueous phase process or a surface-limited process (Ervens and Volkamer, 2010). Data from bulk aqueous phase experiments have been fitted to develop detailed reaction mechanisms for the possible formation and recombination of organic radicals of both glyoxal and methylglyoxal (Lim et al., 2010; Tan et al., 2010) (Table 1).

Experiments in the absence of OH, but under UV light, show similar products from pyruvic acid suggesting that intermediate organic radicals can be also formed by direct photolysis (Guzman et al., 2006). Products in that study included C$_6$ and C$_7$ carboxylic acids which could act as precursors for smaller acids. Solutions of both oxalic and pyruvic acid also showed the formation of HMWC with oligomeric structures in the presence of ozone and UV light (Grgic et al., 2010). In those experiments, it was suggested that pyruvic acid acts as a photosensitizer that accelerates the oxidation of oxalic acid leading to oligomer formation, i.e., indicating an additional role of carbonyl compounds in (or on) the aqueous phase that is not captured by current kinetic models.

All aforementioned experiments were performed in pure water and the only difference between “cloud-water” and “aerosol-water” experiments was the organic aqSOA precursor concentration. However, in atmospheric aerosol water the concentrations of inorganic solutes (e.g., ammonium, sulfate) are much higher and thus aqueous ammonium sulfate solutions might better represent the reaction medium for wet aerosols. Photooxidation of glyoxal, methylglyoxal, glycolaldehyde, methacrolein, α-pinene, and isoprene in an aqueous sulfate solution yields organosulfate species. It is proposed that these species form from reactions involving sulfate radicals (SO$_4^-$) formed in the presence of OH (Noziere et al., 2010b; Perri et al., 2010).

\[ \text{HSO}_4^- + \text{OH} \rightarrow \text{SO}_4^- + \text{H}_2\text{O} \]  

The rate constant of this reaction \((k = 3.5 \times 10^5 \text{ M}^{-1} \text{s}^{-1}, \text{Tang et al., 1988})\) increases
with decreasing pH, which might explain trends of increased organosulfate yields under more acidic conditions in ambient aerosol particles (Edney et al., 2005; Surratt et al., 2007b; Zhang et al., 2011). The molecular weight distribution observed in these laboratory experiments suggests that not only the sulfates of the organic parent compounds but also HMWC with sulfate groups are formed (Noziere et al., 2010b). Experiments in ammonium sulfate solutions in the absence of OH sources but under UV illumination also lead to organosulfates, suggesting additional sources of the sulfate radical (Galloway et al., 2009).

Oligomers that are formed upon photooxidation of limonene by ozone and could contribute to gasSOA by absorption into an organic phase can be further photooxidized in the aqueous phase (Bateman et al., 2011). These experiments demonstrate that the aqueous phase can change the fate and properties of compounds typically considered to be gasSOA products by acting as a SOA sink whereas in organic particles the further oxidation is likely hindered due to diffusion limitations of reactants.

2.1.2 Aromatic compounds

Monocyclic aromatic compounds can be considered as proxies for anthropogenic (e.g., benzene, toluene, xylene) and biomass burning emissions (e.g., lignin). Compounds such as benzene, toluene, xylene react in the gas phase and form gasSOA with highest yields at low NO\textsubscript{x} conditions (Ng et al., 2007). At high NO\textsubscript{x} levels, the formation of carbonyl products (glyoxal, methylglyoxal) is favored which suggests different sensitivities of gasSOA and aqSOA towards NO\textsubscript{x}. As described above, these compounds are precursors of aqSOA but not gasSOA. Photochemical aqueous phase experiments suggest that ring-retaining aromatic oxidation products from phenol and its methoxylated derivates guaiacol and syringol might contribute to aqSOA as they form aromatic dimer products in high yields (near unity, related to the initial aqueous phase concentration) by recombination of phenoxy radicals (Sun et al., 2010). The total aromatic concentrations in these studies (100 µM) resemble those as found from wood smoke markers in winter fog (Sagebiel and Seiber, 1993); however, at other locations much
smaller phenol concentrations are generally observed (Anastasio et al., 1997). Similar results as for methoxyphenols were found in laboratory experiments with other phenolic, vanillyl and syringyl compounds (Chang and Thompson, 2010) and 3,5-dihydroxy benzoic acid (Gelencser et al., 2003). For these latter compounds aqueous reaction times of more than one day were needed to detect pronounced signals of ring-opening products (Hoffer et al., 2004a). Although ring-cleavage products of aromatics will follow reactions as discussed in Sect. 2.1.1, ring-retaining products will add more mass on a mass-to-mass basis and could also affect the optical properties of SOA due to their structural similarities to soot (Sect. 2.4.2).

It should be noted that the atmospheric reaction time scales will depend on the availability of organics and reactants (oxidants) in aerosol and cloud water; as the quantitation of OH radicals in the aqueous laboratory experiments is usually not performed, the comparison of reaction rates from different experiments might not be meaningful. A more useful comparison of the importance of different pathways can only be obtained by using multiphase process models that predict aqSOA formation using the rate constants as summarized in Tables 1 and 2.

2.2 Dark reactions of carbonyl compounds

While photochemical (oxidative) reactions are irreversible, several thermal (dark) reactions form products reversibly, i.e., their reactions can be expressed as equilibria, in form of effective Henry's law constants $K_H^*$ (Kroll et al., 2005b; Galloway et al., 2009; Ip et al., 2009). These constants are often higher by several orders of magnitude than the physical solubility ($K_H$) and effective Henry’s law constants that include hydration ($K_{H,\text{Hydr}}$). The observed gas/particle partitioning, $K_H^*$, of glyoxal has been interpreted as a product of $K_{H,\text{Hydr}}$ and the equilibrium constant for oligomerization $K_{\text{olig}}$ (Ervens and Volkamer, 2010).

$$K_H^* = K_{H,\text{Hydr}} (1 + K_{\text{olig}})$$

(2)
$K_{\text{olig}}$, and thus, $K_H^*$, are only valid for reversible (dark) oligomerization and not for irreversible photochemical reactions (Ervens and Volkamer, 2010; Lim et al., 2010). A more recent study discusses that the enhanced uptake of glyoxal is not due to oligomerization but due to a shift of the partitioning of the glyoxal dihydrate which is more stabilized in ionic solutions (Yu et al., 2011). The hydration and oligomerization of glyoxal, methylglyoxal and other carbonyl compounds are thermodynamically favorable (Barsanti and Pankow, 2004, 2005; Kua et al., 2008; Krizner et al., 2009) but their kinetic formation is slow with rate constants of $k_{\text{hydration}} < 0.01 \text{s}^{-1}$ (Creighton et al., 1988; Krizner et al., 2009) and $k_{\text{olig}} = 100 \text{M}^{-1} \text{s}^{-1}$ (glyoxal), respectively (Ervens and Volkamer, 2010). Overall, these products contribute little mass to the total predicted aqSOA budget since (i) their formation rate constants are small, (ii) the largest observed molecules only contain 3-4 glyoxal subunits and thus are substantially smaller than oligomers from photochemical reactions and (iii) they are formed reversibly and thus their major fraction will evaporate together with water. However, glyoxal oligomerization has recently been shown to contribute significantly to nanoparticle growth (Wang et al., 2010a, 2011b).

Methylglyoxal and other carbonyl compounds that form enol structures can undergo aldol condensation reactions. While aldol condensation products (e.g., for methylglyoxal) are thermodynamically favorable (Barsanti and Pankow, 2005), both experimental and theoretical studies have shown that rate constants of aldol condensation are too low to be important for significant aqSOA formation under typical atmospheric conditions ($0.074 \text{M}^{-1} \text{s}^{-1} < k_{\text{aldol}} < 15 \text{M}^{-1} \text{s}^{-1}$ (Casale et al., 2007; Minerath and Elrod, 2009); $k_{\text{aldol}} < 3 \text{M}^{-1} \text{s}^{-1}$, Noziere et al., 2006).

While neither aldehyde reactions with water nor the self-reaction (including aldol condensation) are predicted to efficiently form aqSOA, interactions of aldehydes with inorganic compounds such as ammonium leading to imidazoles have been suggested to be thermodynamically and kinetically favored (Kua et al., 2011). Several laboratory experiments confirm the reaction of glyoxal and methylglyoxal with ammonium (Noziere et al., 2009; Shapiro et al., 2009) and with monomethyl amine and amino acids in
the absence of photochemical oxidants (De Haan et al., 2009a, c, 2010). Oligomeric products composed of only C, H and O in those experiments were explained by NH$_4^+$ catalysis (Noziere et al., 2009, 2010a). Empirical expressions of the rate constant of the glyoxal/ammonium reaction show a decrease with increasing pH (Noziere et al., 2009; Yu et al., 2011), i.e., opposite to the trend for organosulfate formation (Sect. 2.1.1).

Reactions between ammonium (amines, amino acids) and (methyl)glyoxal are likely to occur in evaporating cloud droplets, i.e. after glyoxal and methylglyoxal dissolve into the relatively large aqueous volume of cloud droplets and as water evaporation increases the concentrations of the nitrogen containing solutes (i.e. mostly ammonium). This mechanism implies that the reaction between volatile, water-soluble organics has to occur faster than the drop evaporation (De Haan et al., 2009b; De Haan et al., 2010). The time scales of these reactions are much longer than for photochemical reactions ($k = 0.3 \text{ M}^{-1} \text{s}^{-1}, k < 0.12 \text{ M}^{-1} \text{s}^{-1}, k \leq 5 \times 10^{-6} \text{s}^{-1}$ for reactions of glyoxal with amines, amino acids and methylglyoxal reaction with ammonium, respectively, Table 2). Thus, it can be expected that their contribution to aqSOA is small; however, the particular structure and properties of the products (e.g., imidazoles) might be possibly used as aqSOA tracers.

### 2.3 Caveats of laboratory experiments

Laboratory studies as described in the previous sections represent idealized systems, i.e., the aqueous solutions do not reflect the complex mixture of organic and inorganic aerosol constituents as found in cloud droplets and aerosol particles. In addition, some of the experiments were performed under conditions that are not necessarily atmospherically relevant. However, only such conditions and the simplicity of the reaction systems allow the identification of products and mechanisms by the applied analytical methods. While the concentration of a single compound might be relatively small, the sum of all (organic) compounds might add up to several M and organic radicals are likely to form from the reaction of OH radical with this complex mixture, creating a more complex array of HMWC than one would see in experiments with a single compound
While a homogenously mixed aqueous phase might be a good proxy for cloud water, it seems likely that in aerosol particles different phases exist whose solutes might not efficiently interact with each other. The smaller abundance of different solutes in laboratory solutions might prevent the formation of different phases (e.g., separation of hydrophobic compounds, water-soluble organics and inorganic ions; Marcolli and Peter, 2005; Zuend et al., 2008; Bertram et al., 2011; Smith et al., 2011) and thus maximize the interactions of reactants within the condensed phase. In an organic phase, the formation of HMWC slows down with increasing product formation since the condensed phase becomes more viscous (“glassy”) and prevents efficient reactant uptake into and transport within this phase (Virtanen et al., 2010; Ziemann, 2010; Pfrang et al., 2011). In contrast, an aqueous aerosol phase containing oxidized and hygroscopic compounds might allow more efficient mixing and oxidation (Cappa et al., 2011). Phase separation and the presence of surface active compounds are likely to affect transport (evaporation, uptake, diffusion) and reaction rates in ambient particles in ways that are not captured in the simple mixtures used in laboratory experiments.

Laboratory experiments that simultaneously measured gas and particle phase concentrations of glyoxal and methylglyoxal upon oxidation of aromatic compounds have shown that their particulate fraction is 4–5 orders of magnitude higher than predicted based on thermodynamic equilibrium (Healy et al., 2008; 2009). The reason for this enhanced partitioning is not fully clear; however, it implies that chemical and/or physical effects in aqueous seed aerosol decrease the effective vapor pressure and lead to higher organic concentrations than in pure water. Some of these deviations might be due to analytical issues or sampling artifacts (Turpin et al., 2000). At least some extraction and chemical analysis methods lead to the “break-up” of oligomers and the particulate fraction might reflect not only the carbonyls but also their oligomers. Thus, laboratory experiments with high organic concentrations seem to be a good proxy for aerosol water.
While in laboratory experiments, the aqueous phase OH is usually produced from H$_2$O$_2$ photolysis, a major source of the OH radical in cloud water is the uptake from the gas phase (Ervens et al., 2003a). Model and laboratory studies have shown that as much as about 1/3rd of the OH radical found in clouds is formed from aqueous reactions (Arakaki and Faust, 1998). Aqueous phase reactants that form OH radical (e.g., ferric ion (Fenton reaction) (Arakaki and Faust, 1998), H$_2$O$_2$ and nitrate (Zellner et al., 1990) are present in considerably higher concentrations in particles, likely enhancing the photochemical formation of OH in the condensed phase. Evidence of efficient OH and H$_2$O$_2$ formation in particles has been recently reported at urban and rural sites in the San Joaquin Valley (Shen and Anastasio, 2011; Shen et al., 2011). Hydrogen peroxide levels in aerosol water exceed predictions based on Henry’s law equilibrium by up to two orders of magnitude since aqueous phase processes represent a continuous, significant source of H$_2$O$_2$ (Arellanes et al., 2006; Wang et al., 2010b). In addition, enhanced rates of organic oxidation reactions in aerosol water lead to a faster recycling of OH. This, and uncertainties in OH radical sinks, make concentrations of OH radicals in aerosol water highly uncertain. Different transport processes into and inside aerosol particles, high solute concentrations, chemical complexity and uncertain oxidant levels complicate efforts to understand and simulate aqueous chemistry in ambient aerosols.

2.4 Aerosol properties of laboratory-derived aqSOA

2.4.1 CCN properties: surface tension $\sigma$ and hygroscopicity $\kappa$ ($\alpha$O/C ratio)

Products of the dark reaction of ammonium, in the absence of oxidants, with glyoxal and with glyoxal-methylglyoxal mixtures show a distinct surface tension $\sigma$ suppression as compared to pure water ($\Delta\sigma \sim 20–35\%$) at molar carbon concentrations due to the oligomeric character of the products with hydrophobic and hydrophilic entities (Sareen et al., 2009; Noziere et al., 2010b; Schwier et al., 2010). It has been speculated that an efficient condensation of surface active gas phase species on deliquesced aerosol particles might be sufficient to form a surface active film that suppresses the
surface tension (Romakkaniemi et al., 2011); however, no observed data can support this mechanism yet.

For aqSOA formation to be substantial, precursor compounds must have high water-solubilities (and/or high concentrations); this suggests that aqSOA will necessarily have high polarity and O/C ratios. While gasSOA usually has O/C ratios of 0.3–0.5 (Aiken et al., 2008; Massoli et al., 2010; Lambe et al., 2011), aqSOA products have higher O/C ratios since the precursors are already highly oxygenated: \(\text{O/C}_{\text{glyoxal}} = 1 - 2\) (depending on hydration), \(\text{O/C}_{\text{methylglyoxal}} = 1.5\), leading to aqSOA with \(\text{O/C} = 2\) (oxalate) in cloud-relevant solutions, and \(\text{O/C} \sim 1\) at aerosol-relevant concentrations (Altieri et al., 2008; Perri et al., 2009). Dark reactions yield products with O/C ratios of 0.5–1.5 (Noziere et al., 2009; Sareen et al., 2009; Shapiro et al., 2009) which is on average lower than those observed in photochemical aqueous phase experiments but still higher than the average O/C ratios of chamber-derived gasSOA.

The hygroscopicity of particles – represented by the hygroscopicity parameter \(\kappa\) (Peters and Kreidenweis, 2007) – correlates with the O/C ratio (Jimenez et al., 2009; Lambe et al., 2011). Extrapolating this linear \(\kappa\)-O/C relationship to products found in glyoxal and methylglyoxal aqSOA experiments suggests that aqSOA will be more hygroscopic than gasSOA, with \(\kappa\) of \(\sim 0.15-0.6\), i.e. with an upper limit that is comparable to ammonium sulfate. This prediction agrees well with the hygroscopic growth factor (GF) measurements of MACR-derived aerosol aqSOA of GF \(\sim 1.4\) at RH \(= 90\%\) (Michaud et al., 2009). Extrapolating the reported GF-O/C (at RH \(= 95\%\)) correlation by Jimenez et al. (2009) to RH \(= 90\%\) yields approximately \(\kappa_{\text{aqSOA}} \sim 0.2\).

Sensitivity model studies of CCN activity over ranges of \(\kappa\) and \(\sigma\) suggest that (i) in cloud water and dilute aerosol water as encountered near cloud base where droplet activation occurs, organic solute concentrations are much lower (\(\sim \mu\text{M}-0.1\text{mM}\)) and \(\sigma\) suppression due to organics is likely negligible and (ii) high molecular weight and reduced surface tension offset each other resulting in CCN properties for aqSOA similar to those of ammonium sulfate (Ervens et al., 2005; Padro et al., 2010). Thus, while the physicochemical properties of aqSOA in terms of CCN ability might be approximated...
by inorganic solutes, the addition of hygroscopic aqSOA mass, in addition to sulfate, to the droplet mode of an aerosol population will increase particle sizes and thus has implications in terms of direct and indirect aerosol effects.

### 2.4.2 Optical properties

The semidirect and direct aerosol effects due to light absorption by organics (“brown carbon”) are not – or only very crudely – included in climate models. Several reactions discussed in Sects. 2.1 and 2.2 yield light-absorbing products that could contribute to atmospheric brown carbon (Sareen et al., 2009; Shapiro et al., 2009). Experiments of glyoxal with ammonium yield yellow-brownish solutions that stem from the formation of C-N containing products (Debus, 1858; Galloway et al., 2009; Sareen et al., 2009; Shapiro et al., 2009; Trainic et al., 2011). Absorption measurements show absorption at $\lambda = 209$ nm immediately after mixing the compounds (Noziere et al., 2009) and a band that builds up over several hours at $\lambda = 290$ nm. Spectra from the same reaction system over longer time scales revealed absorption between 300–400 nm after a few hours of reaction and structure-less absorption spectra after several days. The refractive index $n$ for products of the glyoxal and ammonium reaction depends on the particle size but can be described within a range of values $n = (1.57 + 0 \, i)$ to $n = (1.7 + 0.02 \, i)$ (Trainic et al., 2011).

Photochemical experiments with aromatic compounds (phenol and related compounds) also showed a shift in absorption to longer wavelengths (from 350–$\sim650$ nm) whereas the precursors show distinct absorption at $\lambda \sim300$ nm (Gelencser et al., 2003; Hoffer et al., 2004b; Chang and Thompson, 2010). The evolution of the absorption at larger wavelengths has been explained by the formation of larger conjugated oligomers that exhibit soot-like structures. Such structures cannot be formed in the gas phase in an oxygen-containing atmosphere. Product spectra resembled the spectra of atmospheric HULIS; correlations were observed between absorbing organic mass and RH and the soluble extracts from ambient aerosols associated with biomass burning and SOA (Hecobian et al., 2010), implying formation of aqSOA (Sect. 3.4.2). Even though
the data sets for optical properties of aqSOA compounds are sparse, the qualitative observations summarized here suggest that small aqSOA contributions to the total aerosol mass might have distinct effects on the aerosol radiative properties. However, light-absorption is not unique to aqSOA, since also some gasSOA products absorb UV light (e.g., nitroaromatics; Jacobson, 1999; Jaoui et al., 2008; Nakayama et al., 2010) at smaller wavelengths and with a stronger wavelengths dependence (Ångstrom exponent) than soot (Kirchstetter et al., 2004). The particle size to which light-absorbing organic carbon is added will depend on the formation process (gasSOA, aqSOA) and impacts the overall absorbing/scattering properties of aerosol populations. AqSOA is generally added to larger particles (droplet mode) whereas gasSOA rather adds to smaller particles.

3 Observations and model studies of aqSOA

3.1 Missing SOA source in current models

Many gasSOA models fail to reproduce total SOA mass and/or specific properties (De-Gouw et al., 2005; Heald et al., 2005; Volkamer et al., 2006; Hodzic et al., 2010). While an underestimate of SOA mass can imply an underestimate of precursor concentrations (emissions) or gasSOA yields, underprediction of the O/C ratio, correlation with RH, and correlation with aqueous phase chemistry tracers (e.g., sulfate) infer a role for aqSOA.

Water-soluble organic carbon (WSOC) in the free troposphere above North America correlates with sulfate and methanol (Heald et al., 2006). Since sulfate and methanol are known to form predominantly in the aqueous phase and WSOC is predominantly SOA in locations with little biomass burning, the observed correlation between WSOC and aqueous phase tracers suggests that WSOC might be largely composed of aqSOA. In East Asia, SOA predictions were low by ~50 % for the boundary layer but low by a factor 10–100 in the free troposphere (Heald et al., 2005), suggesting a role for
3.2 Evidence of aqSOA based on aerosol size distributions

Ambient aerosol populations often show two distinct submicron modes (<0.2 µm and 0.5–1 µm) where the larger (droplet) mode is formed from the smaller (condensation) mode through volume-phase reactions in clouds and wet aerosols (e.g., sulfate formation) (Hering and Friedlander, 1982; John et al., 1990; Meng and Seinfeld, 1994; Kerminen and Wexler, 1995). An abundance of highly oxidized organics and organosulfates in the droplet mode has been observed in several locations (e.g., Pickle et al., 1990; Blando et al., 1998). A shift in the carbonyl size distribution to a larger median at high RH has also been observed (Maria et al., 2004). Size resolved measurements of OA, sulfate and nitrate in Los Angeles during marine influence and humid conditions show an abundance of low-volatility oxygenated organics, sulfate and nitrate in the droplet mode (Filho et al., 2003). These observations provide strong evidence that the organic aerosol at this time was either (1) formed in the aqueous phase or (2) present in condensation mode particles that grow into the droplet mode by aqueous sulfate and/or nitrate formation. The fact that the highest OA mass concentrations were observed when the OA was present in the droplet mode suggests the former, an aqSOA formation process. During periods of clear dry air and intense photochemical activity condensation mode OA dominated, characteristic of the gasSOA route. The modality of SOA reflects its formation mechanisms that have been observed and predicted for sulfate (Hoppel et al., 1994; Feingold and Kreidenweis, 2000). GasSOA models do not reproduce the observed bimodality of organic aerosol mass (Pandis et al., 1993).
3.3 AqSOA formation in cloud droplets

3.3.1 Observation and prediction of oxalate as a proxy of aqueous phase processing

It is well established that globally the majority (70%) of atmospheric sulfate is formed in cloud droplets and only a minor fraction originates from condensation of \( \text{H}_2\text{SO}_4 \) on particles or from formation in aerosol water (Sievering et al., 1992; Seinfeld and Pandis, 1998). Enhanced sulfate concentrations are found in the cloud-processed droplet mode. In such particles, small dicarboxylic acids are often found internally mixed with sulfate (Yao et al., 2003; Yu et al., 2005; Huang et al., 2006; Sorooshian et al., 2006), pointing to their efficient formation during cloud-processing. Oxalate, the smallest dicarboxylic acid (salt) is formed from aqueous oxidation of several precursor compounds (Warneck, 2003; Ervens et al., 2004), and it has been found to be the most abundant individual organic compound in organic aerosol (Neusüss et al., 2000).

Oxalate can be routinely measured by ion chromatography of integrated filter samples, or equivalent online systems (e.g. particle-in-liquid-sampler (PILS), Orsini et al., 2003), along with inorganic ions, providing robust data sets for a wide variety of atmospheric conditions. Comparisons of box model results with measurements show that chemical aqueous phase processes in cloud water can explain enhanced in-cloud and above-cloud oxalate-to-sulfate ratios compared to below-cloud ratios (Sorooshian et al., 2007). During the ICARTT experiments flights over Ohio and surrounding areas, inorganic ions and oxalate were measured in cumulus clouds with thicknesses between 500–700 m. In model calculations, it was assumed that sulfate formation could be appropriately reproduced and thus predicted sulfate/oxalate ratios were compared to measurements (Sorooshian et al., 2006). The general agreement of the absolute sulfate and oxalate masses in this study as well as their ratios gave confidence in the understanding of in-cloud oxalate formation. During CARMA I, a field campaign designed to study marine stratocumulus off the coast of Monterey, CA, oxalate predictions were higher by a factor of 10 (210 ng m\(^{-3}\)) than the observed ones (Crahan et
al., 2004). The discrepancy was ascribed to missing wet deposition or particle removal from the clouds; while not considered by these authors, chemical losses (oxidation by OH or photolysis of metal complexes) may also contribute to oxalate losses (e.g., Ervens et al., 2004; Kawamura et al., 2010). A global modeling study concluded that in-cloud formation could explain the atmospheric abundance of oxalate (Myriokefalitakis et al., 2011).

While the vapor pressure of oxalic acid does not suggest it, roughly 90% of oxalate is found in the particle phase (Baboukas et al., 2000; Yao et al., 2002). This behavior can be explained by salt or complex formation (ammonium, potassium, metal ions (Furukawa and Takahashi, 2011)); increased oxalate concentrations in (alkaline) dust and biomass burning particles confirm this conclusion since at higher pH, salt formation is favored (Yao et al., 2002; Sullivan and Prather, 2007; Moffett et al., 2008). Back trajectories of dust particles, internally mixed with oxalate, revealed that these particles were not cloud-processed but that they were aged due to condensation of soluble species onto the dust particle surface. Hygroscopic material on aged particles provides liquid water on the particles (aerosol water) (Yao et al., 2002) and thus might also explain oxalate in the coarse mode of marine particles which cannot be explained by in-cloud processes (Rinaldi et al., 2011). Because the vapor pressure of oxalic acid is much higher than that of oxalate salts, this species can be transported in the gas phase (regardless of how it forms), and react with anions in dust and sea salt particles.

The total contribution of oxalic acid to the total OA fraction usually does not exceed a few percent (e.g. Kawamura et al., 1996b, 2003) and is only one of many products formed through aqueous chemistry. However, the numerous data sets for oxalate and the availability of detailed chemical mechanisms of its formation pathways make it a good tracer to test our understanding of aqueous phase processing of small organic molecules.
3.3.2 Observed and predicted larger (>C$_2$) aqSOA products

Larger dicarboxylic acids have been found internally mixed with oxalate, which suggests that their formation pathways are connected (e.g., Yao et al., 2003; Yu et al., 2005; Huang et al., 2006; Sorooshian et al., 2006). Specifically, correlations have been observed of oxalate with sulfate, malonate (C$_3$) and succinate (C$_4$) in aerosol particles (e.g., Kawamura et al., 2000, 2003; Narukawa et al., 2003; Jung and Kawamura, 2011). Large dicarboxylic acids can form smaller acids including oxalate through aqueous oxidation. In addition, >C$_2$ diacids can form from smaller compounds through aqueous organic radical-radical reactions in the high organic concentrations found in wet aerosols. While a motor vehicle source of small dicarboxylic acids has been implied in early studies (Kawamura and Kaplan, 1987; Kawamura et al., 1996a), more recent studies do not find any correlation of dicarboxylic acids with traffic markers (NO, NO$_2$) (Martinelango et al., 2007). In addition to dicarboxylic acids, recent size-resolved measurements also show a predominance of HMWC in the droplet mode (Lin et al., 2010b), suggesting concurrent formation of HMWC through aqueous chemistry. Correlations of HMWC with potassium imply the formation in biomass burning plumes that include aged aerosol particles (Lin et al., 2010a).

A related measure of the amount of highly oxygenated OA is the signal of the mass-to-charge ratio $m/z = 44$ (CO$_2^+$) as obtained from aerosol mass spectrometer (AMS) measurements, which is proportional to the number of acid functionalities in the organic mass fraction. A recent study has shown that cloud droplet residuals (at the Pacific Coast and in continental air masses) show highest correlations with $m/z = 44$, the oxalate mass and the total organic acid mass to the total organic mass as compared to aerosol masses that were not processed by clouds (Sorooshian et al., 2010). A related study for a data set in the Gulf of Mexico shows that aqSOA formation in clouds can clearly alter the vertical profile of the particulate WSOC distribution. Such studies corroborate conclusions regarding enhanced organic aerosol concentrations in regions of cloud outflow (Peltier et al., 2008).
The implementation of in-cloud aqSOA formation into a regional air quality model (CMAQ) to simulate SOA formation over northeastern US resulted in a much better agreement between predictions and aircraft measurements of WSOC (Carlton et al., 2008), despite a very crude estimate of glyoxal conversion into aqSOA (using a fixed yield of 4%). Not only did the bias in the total predicted SOA mass to observations decrease (from −64% to −15%), but the model also captured better the temporal variability in OA, especially for flights through clouds. Using the same model (CMAQ) to simulate SOA formation in the same region, but using an explicit aqueous phase mechanism to predict organic acid formation in clouds from glyoxal and methylglyoxal, aqSOA formation was predicted to account for an overall SOA mass increase of 27% for rural scenarios and 7% for anthropogenically influenced areas (Chen et al., 2007). In the same study it was also found that aqSOA formation in clouds was quite uniform at the surface and in free troposphere layer (0.28 µg m\(^{-3}\) and 0.25 µg m\(^{-3}\), respectively). These authors concluded that aqSOA formation might be in general more important for biogenically-impacted regions since glyoxal and methylglyoxal are formed in higher yields from isoprene than from anthropogenic emissions. Of course, anthropogenic emissions can alter gas phase chemistry (e.g., through NO\(_x\)), oxidant production and aerosol water concentrations (i.e., through sulfate formation), all of which are important to aqSOA formation. Note both of these modeling studies neglected aqSOA formation in aerosol water.

### 3.4 AqSOA formation in aerosol water

#### 3.4.1 AqSOA tracer compounds

HMWC is mainly formed in aerosol water as opposed to cloud water as the rates of self-reactions of organics (oligomerization, accretion reactions) are negligible in dilute (cloud) water (Sect. 2). Because of the different types of reactions in concentrated and dilute aqueous phase, aqSOA mass formation rates [ng m\(^{-3}\)gas] do not scale with the liquid water content (LWC) but have high overall rates in both aqueous regimes. This
transition from dilute to concentrated aqueous phase chemistry is unique to aqSOA formation. Sulfate has only negligible formation rates [ng m$^{-3}$] in the small volume of aerosol water and thus its formation rates in aerosols are much smaller than in clouds (Sievering et al., 1992; Meng and Seinfeld, 1994). Mass formation in aerosol water results in a similar size modification of an aerosol population (mass addition to the droplet mode) as cloud processing (Maria et al., 2004) and thus, the existence of HMWC together with sulfate or other cloud-derived material points to aqSOA formation in the aqueous phase.

Many compounds such as di- and oxocarboxylic acids, esters and organosulfur compounds that do not have any (efficient) gas phase sources have been found in rain and cloud water samples (Graedel and Weschler, 1981; Blando and Turpin, 2000); their presence in droplets does not necessarily imply that they have been formed in this dilute aqueous phase. The presence of aldehyde-sulfur (IV) adducts (e.g., hydroxy-methane sulfonate) in fog, clouds and aerosols has been interpreted as tracers of aqueous phase chemistry as the corresponding adduct formation does not occur in the gas phase (Munger et al., 1986; Olson and Hoffmann, 1989; Dixon and Aasen, 1999; Lee et al., 2003; Whiteaker and Prather, 2003).

Evidence from laboratory studies suggests that HMWC with acid, sulfur or ester functionalities can be formed in the aerosol aqueous phase and might then dissolve in droplets. In rain and fog water samples, several hundreds of compounds in a mass range of 50<$m/z<$500 were identified by ESI-FT-ICR-MS (Electrospray ionization coupled with Fourier transform ion cyclotron resonance mass spectrometry) showing series of oligomers with compounds that differ by mass fragments of C$_3$H$_4$O$_2$ ($\beta$-hydroxy ketones) (Altieri et al., 2009; Mazzoleni et al., 2010), i.e., the same fragments that have also been identified in laboratory experiments pointing to the oligomerization of carboxylic and hydroxycarboxylic acids (Tables 1 and 2) (Altieri et al., 2008; Yasmeen et al., 2010). HMWC in atmospheric particles comprise a substantial fraction of the water-soluble organic matter (Fuzzi et al., 2001; Krivacsy et al., 2008; Stone et al., 2009) which was originally attributed to primary sources (Likens et al., 1983).
primary and secondary HMWC are polyacidic, the latter comprise lower average molecular weight and lower aromatic moiety content (Graber and Rudich, 2006). While it is possible that a fraction of this organic matter is formed in different liquid (organic) fractions of an aerosol particle, the high polarity and number of carboxylic acid groups of the identified compounds (Fuzzi et al., 2001; Salma and Lang, 2008), their agreement with laboratory-derived aqSOA products (Sect. 2), and correlation with clearly aqueous-phase-derived species and with the amount of aerosol/cloud liquid water contents suggest that they are mainly formed in droplets and/or aqueous particles. The total bulk masses range from 2–100 µg m\(^{-3}\) and up to 60 % of the water-soluble organic matter in ambient and biomass burning aerosol, respectively (Fuzzi et al., 2002; Lin et al., 2010b).

3.4.2 Correlation of WSOC with RH

The observed increase in particle-phase water-soluble organic carbon (WSOC\(_p\)) at high RH may also point to aqSOA formation. In Atlanta, the fraction of total WSOC (gas + particulate) that is present in particles (F\(_p\)) increases sharply at high RH, suggesting a positive correlation with aerosol water (Fig. 2a, b). No relationship was readily observed between F\(_p\) and particle organic mass (Fig. 2c), which would be expected from the gasSOA formation route since gasSOA forms through partitioning of semivolatile species into particulate organic matter (Hennigan et al., 2008a,b, 2009). This peak in F\(_p\) coincides with particulate nitrate and thus the formation of both particulate components might be initiated by HONO.

Based on the measured WSOC gas/particle partitioning, an effective Henry’s law constant \(K_{H^*} \sim 10^9 \text{ M atm}^{-1}\) for total WSOC was derived (Hennigan et al., 2009). While not pertaining to a specific compound, this value exceeds all values that are known for the physical solubility of common atmospheric organic compounds and is even higher than any derived from studies that suggest reversible oligomerization (Sect. 2.2); values of this order of magnitude have only been observed for glyoxal/aqSOA partitioning in photochemical laboratory experiments where the application of \(K_{H^*}\) strictly does not
apply (Volkamer et al., 2009). However, inferring that the observed particulate WSOC mass is indeed photochemically and thus irreversibly formed is somewhat speculative based on the current data set as the bulk OC measurements do not give any information on organic speciation. However, in additional studies organic acids and HMWC have been found to be major components of this fraction which might hint to the presence of photochemically formed oxidation products (Sullivan and Weber, 2006a, b; Miyazaki et al., 2009a).

Correlations between WSOC$_p$, nitrate and particle water were also observed in Mexico City during the MILAGRO experiment (Hennigan et al., 2008). As often observed, WSOC$_p$ and nitrate peaked shortly following sunrise when RH is high and photochemical processes begin. A similar trend was observed during the Environmental Protection Agency (EPA) Southern Oxidants Study (SOS) PM$_{2.5}$ Supersite Project in Atlanta (Lee et al., 2003). Internally mixed organic and nitrate particles showed a maximum in the morning when RH was high. A smaller maximum of organics was observed in the afternoon associated with internally mixed organic and sulfate particles.

As the day progressed during MILAGRO, concentrations of nitrate and WSOC$_p$ decreased due to boundary layer expansion and evaporation of some fraction of newly formed secondary nitrate and WSOC as RH approached the daily minimum in early afternoon. Only roughly 35% of the fresh WSOC$_p$ appeared to be volatile, a fraction considerably less than that for nitrate, which resulted in an increasing WSOC/nitrate ratio, indicating a significant irreversible net aqSOA production in aerosol water (Hennigan et al., 2008a). For the same field experiment, AMS measurements provided the O/C ratio as a proxy of the degree of oxygenation of the organic aerosol mass. While this ratio agrees well with data from other ambient aerosol samples for the first two weeks (0.3 < O/C < 0.9, Ng et al., 2010) it is clearly increased for the time period where elevated RH was observed (Fig. 2d, Fast et al., 2007; Hodzic et al., 2010). The increase of the O/C ratio with RH (Fig. 2e) suggests that not only oxygen is added to preexisting molecules (e.g., heterogeneous oxidation of aerosol components with little change in particle mass, only $\Delta$O) but rather that additional organics are taken up into
the increased aqueous volume and further oxidized into aqSOA products that exhibit high O/C ratios (cf Sect. 2.4.1). The predicted trend from a regional model that only includes gasSOA formation shows the opposite behavior (Fig. 2e; Hodzic et al., 2010), which supports once more that gasSOA models do not capture the RH dependence of SOA formation. At the same location (during the MCMA field experiment), the comparison of modeled and observed gas phase glyoxal revealed a ‘missing sink’ of glyoxal as a highly water-soluble, volatile compound together with an unexpected source of SOA (Volkamer et al., 2007) which gave the first observational evidence of aqSOA formation in aqueous particles. Estimates based on this mismatch between modeled and observed glyoxal gas phase concentrations, suggest that in Mexico City aqSOA formation from glyoxal in aqueous particles could contribute substantially to total SOA (up to 15\%) (Volkamer et al., 2007; Dzepina et al., 2011).

The O/C ratio is directly related to the ratio of total organic mass to organic carbon (OM/OC) as oxygen is the most abundant heteroatom in organic molecules (Pang et al., 2006). In many model studies, a constant OM/OC ratio is applied to convert from measured OC to total organic aerosol mass although it has been shown that it greatly varies depending on the composition of the OM fraction (Turpin and Lim, 2001). For HMWC, a ratio of OM/OC = 1.8 has been determined (Salma et al., 2008). A similarly elevated OM/OC ratio for total OM has been reported for the Southeast US (Simon et al., 2010). While it is possible that this result might be influenced by the assumptions made in this study, this finding is also consistent with a large role for aqSOA in this high RH, photochemically-active location (Hennigan et al., 2009) (Fig. 2a–c).

Measurements in LA, during the CalNex experiment (http://www.esrl.noaa.gov/csd/calnex/), show the opposite trend in terms of WSOC partitioning (F_p), correlation with RH and OC, as compared to the Southeast US (Fig. 2b,c versus 2g,h). No relationship is observed between F_p and RH, but unlike in Atlanta, a correlation is observed between F_p and OC mass, in agreement with the partitioning theory of gasSOA formation. In Los Angeles, this points to much less significant aqSOA contribution, even though RH shows a similar temporal profile with even higher values in LA. One conclusion from
the comparison of LA and Southeast US data might be that the higher emissions of biogenic compounds make the southeastern region more conducive to aqSOA routes as compared to the LA basin. OA measurements at very different locations are not sufficiently comprehensive to elaborate on any conclusions. In Cairo, a dry, highly anthropogenically influenced location, the formation of water-insoluble SOA has been observed (Favez et al., 2008) which might add to evidence that not only the SOA precursor, but also the meteorological conditions, and thus the type of chemical process (aqueous or gas phase) determines the formation and distribution of SOA. Studies also show that products of aqueous photochemical reactions can better reproduce the mass spectra of aged secondary organic aerosols in Whistler, Canada, an area that was highly influenced by biogenic emissions (Lee et al., 2011). Firm conclusions on the relative role of precursor compounds and non-chemical effects to the contributions of gasSOA and aqSOA cannot be drawn based on these sparse data sets and warrant future field and model studies.

3.4.3 Discussion of model approaches to predict aerosol aqSOA

While several models include some representation of cloud aqSOA formation, data sets of reaction parameters for aerosol aqSOA formation are much less constrained. No comprehensive module has been developed to date that predicts aerosol aqSOA formation based on the reaction parameters summarized in Tables 1 and 2. However, in the following we present the current attempts to estimate the role of the aqueous phase of aerosols and discuss their caveats that could be improved based on the most recent understanding of aerosol aqSOA processes.

Based on vapor pressures, activity coefficients and solubility, it has been predicted that in the South Coast Air Basin of California less than 30% of condensable gasSOA precursors partition to the aqueous phase of particles as opposed to the (hydrophobic) organic phase (Griffin et al., 2003; Chang et al., 2010). However, these studies did not take into account more volatile water-soluble organics that might dissolve into the aqueous phase (e.g., glyoxal) for further processing. Similar to the trends as identified
in LA (Fig. 2f–h), in that model study no (or an inverse) trend between aqSOA and total OA mass was observed.

Extending a very detailed cloud chemistry model to aerosol water conditions, it was concluded that the aqueous phase formation rates \([\text{M s}^{-1}]\) of mono- and multifunctional organic acids by the OH radical are high in both cloud droplets and aqueous particles (Tilgner and Herrmann, 2010). However, appreciable aqSOA mass formation was only predicted in cloud droplets since the absolute mass conversion \([\text{ng m}^{-3}\text{gas}]\) in aerosol water is small. Similar qualitative conclusions were drawn from global model studies where an irreversible uptake of glyoxal and methylglyoxal into cloud and aerosol water using a reactive uptake parameter \(\gamma = 0.0029\) (based on laboratory studies by Liggio et al., 2005a, b) (Sect. 2) was assumed (Fu et al., 2008, 2009). In these studies, it was concluded that aqSOA formation could enhance the total SOA burden by 60 % and that globally 64 % of aqSOA in the free troposphere is non-biogenic. A reactive uptake coefficient \(\gamma\) implies a surface-limited process whereas the most efficient (photochemical) aqSOA formation processes are suggested to occur in the bulk aqueous phase (Sect. 2.1). In addition, a surface-limited process would rather be at odds with measured size distributions that show clear processing of droplet mode particles (Hersey et al., 2011). The minor importance of aqSOA formation in aerosol water in these studies is due to the application of identical reaction parameters for cloud and aerosol water. Since LWC in clouds is significantly larger than in aerosols \((\sim 0.1 < \text{LWC(cloud)} \text{[g m}^{-3}] < \sim 1; \sim 1 < \text{LWC(aerosol)} \text{[µg m}^{-3}] < \sim 100\), the same chemical bulk reactions will always yield higher aqSOA masses in cloud droplets. However, laboratory studies clearly show that the application of identical reaction parameters for both aqueous regimes is not appropriate since high solute concentrations in aerosol water open additional chemical pathways (oligomerization) that lead to different aqSOA products (Sect. 2).

Global aqSOA prediction in aerosol water applying reversible aqSOA formation from glyoxal \((K_H^* = 2.6 \times 10^7 \text{M atm}^{-1})\) (Myriokefalitakis et al., 2008) certainly represents a lower bound of aqSOA since photochemical processes might form aqSOA mass more efficiently and irreversibly (Sect. 2.1). A combination of aqSOA formation in particles
(using $\gamma = 0.0029$) and an in-cloud parameterization similar to Ervens et al. (2008) was applied to estimate the aqSOA burden over the continents (Stavrakou et al., 2009). This study concluded that glyoxal loss on aerosol particles is at least as efficient as on cloud droplets or even up to 60% higher. A recent detailed model study of global oxalate formation by aqueous processes in cloud and aerosol water predicted that more than 90% of this compound is formed in clouds whereas it was assumed, based on laboratory experiments (Lim et al., 2010), that the photochemical glyoxal oxidation in aerosol water has an oxalate yield of 20% (Myriokefalitakis et al., 2011).

The summary of these studies shows that none of the models fully captures all conversions of water-soluble organics in aerosol water as summarized in Tables 1 and 2. Most reaction parameters for aerosol aqSOA formation are available for glyoxal and methylglyoxal. However, not only small carbonyl compounds but also alcohols, carboxylic acids, organic peroxides and larger, semivolatile compounds might act as precursors for aqSOA (Fig. 1). While they might alternatively participate in gasSOA formation, their partitioning and further processing in aerosol water will lead to different mass distribution and aerosol properties.

4 Comparison of gasSOA and aqSOA mass predictions

4.1 AqSOA formation in cloud droplets vs aerosol particles

The model approaches summarized above do not allow a comprehensive comparison of the importance of cloud and aerosol aqSOA formation. In order to provide a first estimate on the extent to which cloud and aerosol chemistry yield aqSOA mass, we simulated these processes in a parcel model. The same model was applied previously to evaluate aqSOA formation in clouds, i.e., with RH $\geq 100\%$ and LWC $\geq 0.01$ g m$^{-3}$ (Ervens et al., 2004, 2008). Here the model has been extended to account for aerosol aqSOA formation during periods when RH $< 100\%$. Model simulations were performed over four hours using repeated 1-h trajectories that prescribe meteorological
parameters (temperature, pressure, RH). RH as a function of time in the applied trajectories is shown in Fig. 3a for the simulation time between 2.5 and 3.5 h.

Photochemical rate constants for glyoxal and methylglyoxal as derived from experiments at high organic concentrations (Volkamer et al., 2009; Ervens and Volkamer, 2010; Lim et al., 2010; Tan et al., 2010) as well as the reaction of glyoxal with ammonium have been included (Tables 1 and 2; reactions marked with (*)). Note that the lumped photochemical rate constants used here do not describe a detailed mechanism but are first-order rate constants that have been fitted to the observed temporal aqSOA mass profiles in laboratory experiments. Comparisons of aqSOA mass formation rates using these rate constants and the detailed mechanisms by Lim et al. (2010) and Tan et al. (2010a) that describe the same reaction system, show good agreement.

In Fig. 3b the predicted cloud aqSOA mass is shown as the sum of oxalic, glyoxylic, glycolic, and pyruvic acid masses that is assumed to remain in the particle phase upon cloud evaporation. If the air parcel is inside the cloud, cloud aqSOA masses of up to 2 µg m$^{-3}$ are predicted; during cloud processing and evaporation part of this mass is oxidized and/or evaporates (Ervens et al., 2008). After 3 cloud cycles, about 1 µg m$^{-3}$ cloud aqSOA is predicted to remain in the particles at RH $\sim$ 75%. A continuous mass increase of about 1 µg m$^{-3}$ aerosol aqSOA formed during the time when the parcel is outside of clouds is predicted after the same simulation time for all trajectories (differently colored traces in Fig. 3c). The approximately equal amount of aqSOA predicted to form in cloud droplets and aerosol water seem at odds as the aerosol LWC is smaller by several orders of magnitude than cloud LWC. However, as pointed out in Sect. 2, there is a transition between dilute (cloud) and concentrated (aerosol) aqueous phase chemistry. Thus, even small concentrations in cloud water can lead to significant mass formation rates (related to gas phase concentrations [ng m$^{-3}$ gas]) whereas in aerosol water oligomerization reactions lead to higher overall aqSOA formation [Ms$^{-1}$] resulting in similar overall aqSOA mass production rates [ng m$^{-3}$ gas].

In general, the relative contributions of cloud and aerosol aqSOA to the total SOA depend on the time scales air parcels are exposed to supersaturated (clouds) and
subsaturated (aerosols) conditions. However, the results in Fig. 3 show that on a comparable time scale aqSOA formation in clouds and aerosol water might be equally efficient. It has been shown that dicarboxylic acids formed in clouds might increase the total organic aerosol mass by up to 10% (Sorooshian et al., 2010), in agreement with the fraction of these acids to total WSOC. However, the processing time in aerosol water might be much longer (hours-days) than processing in cloud water, as aqueous aerosols do not undergo activation and evaporation cycles of several minutes, but instead can be exposed to cycles of elevated RH through repeated diurnal temperature cycles or changes in altitude. This suggests that the vast majority of aqSOA may form in aerosol water and be comprised of products with higher carbon number than their precursors.

4.2 Partitioning of aqSOA and gasSOA compounds

4.2.1 Concepts of partitioning

Whereas gasSOA mass is a function of preexisting organic aerosol mass, gasSOA compounds can absorb into, the amount of aqSOA material that can be dissolved in aerosol (or cloud) water depends on the LWC. The formation and partitioning of gasSOA species between the gas and particle phase is described by either the traditional approach that assumes two oxidation products P1 and P2 (2-product model; Odum et al., 1996) or by a more recent approach that includes also semivolatile products and distributes the oxidation products over logarithmically spaced “volatility bins” according to their saturation pressure (volatility basis set (VBS); Donahue et al., 2006). Both approaches ascribe stoichiometric coefficients to the predicted (generic) gasSOA products and the amount of mass that partitions to the particle phase \( F_{\text{particle}} \) depends on the saturation vapor pressures \( C^* \) (= inverse of the partitioning coefficient \( K_p \)) and the
amount of organic material in the absorbing aerosol ($m_{\text{org}}$):

$$C^* = K_p^{-1} = \frac{F_{\text{gas}} \cdot m_{\text{org}}}{F_{\text{particle}}} [\mu g m^{-3}]$$  

(3)

Particle phase reactions in aqueous or organic phases might further decrease the vapor pressure of the gasSOA compounds and thus enhance the partitioned fraction, i.e., moving it into a “lower volatility bin” which implies a higher partitioned fraction.

The fraction of the total water-soluble aqSOA precursor and product masses that dissolves in aerosol or cloud water depends on the LWC. The Henry’s law constant $K_H$ (or $K_H^*$, Sect. 2.2) represents the ratio of aqueous and gas phase concentration; thus, the product of $K_H$ and LWC represents a measure that includes the same quantities as $C^*$ whereas $m_{\text{org}}$ is replaced by the water mass $m_{\text{water}}$. In order to differentiate $C^*$ for gasSOA which is a function of $m_{\text{org}}$, and $C^*$ for aqueous phase species, we will refer to the latter to as $C^*_{\text{aq}}$. Thus, for both gasSOA and aqSOA the partitioned fraction $F_p$ (similar to Fig. 2) of predicted SOA products can be estimated for assumed organic and water masses, respectively, by

$$F_p(\text{gasSOA}) = \frac{m_{\text{org}}/C^*}{1 + m_{\text{org}}/C^*}$$  

(4)

$$F_p(\text{aqSOA}) = \frac{LWC/C^*_{\text{aq}}}{1 + LWC/C^*_{\text{aq}}}$$  

(5)

Such predicted fractions $F_p$ do not give any information about the actual SOA mass as it does not include information on the amounts formed from the different precursors (stoichiometric coefficients for gasSOA and time scales for aqSOA). However, the purpose of comparing $F_p(\text{gasSOA})$ and $F_p(\text{aqSOA})$, as done in the following section, is rather to provide a sense of the ability to which precursors and oxidation products might partition.
4.2.2 Comparison of partitioned gasSOA and aqSOA fractions

Organic and water masses may be on the same order of magnitude in ambient aerosol, and, under the assumption that both water and organic matter are equally accessible to condensing species, a comparison of the partitioning of individual compounds can be made by applying Eqs. (3–5). In order to show this concept, the partitioning of gasSOA products from benzene, toluene, xylene, and isoprene is shown in Fig. 4a and expressed as the fraction $F_p$ of gasSOA species in the particle phase. We show results for both gasSOA approaches (2-Product model, VBS) using $K_p$ values for $P_1$ and $P_2$ from Henze and Seinfeld (2006), Henze et al. (2007), and Ng et al. (2007) and $C^*$ values from Tsimpidi et al. (2010). The lower-volatility products from all precursors partition to a significant extent to the particle phase (30–60 % for $m_{\text{org}} = 1 \mu\text{g m}^{-3}$; >60 % for $m_{\text{org}} = 10 \mu\text{g m}^{-3}$). Predictions using the VBS reveal that some gasSOA products from aromatics might partition to a high extent to the particle phase for any assumed $m_{\text{org}}$. However, both aromatics and isoprene also produce products with $1000 > C^* > 1$ which result in $F_p < 5 \%$. It should be noted that such compounds might get further processed (“aerosol ageing”) and thus decrease their vapor pressure which will move them to smaller $C^*$ values.

Figure 4b shows the partitioning ($F_p$) of aqSOA products in aerosol water. The highest $F_p$ value (30 % and 5 % for $\text{LWC} = 10 \mu\text{g m}^{-3}$ and $1 \mu\text{g m}^{-3}$, respectively) is predicted based on the observations of the partitioning of total WSOC in Atlanta (Sect. 3.4.2). This fraction might also include photochemically formed products. Photochemically formed glyoxal oxidation products in aerosol water might partition to a few percent (<10 %) into aerosol water whereas the reversibly formed glyoxal oligomeric products as predicted from effective Henry’s law constants by Ip et al. (2009) might not contribute to aerosol mass to a significant extent even if relatively high aerosol LWC is available. Based on the Henry’s law constant, oxalic acid might only partition to $\sim 10 \%$ into aerosol water. Observations have shown that oxalate concentrations in the gas phase are negligible and have ascribed enhanced partitioning to complex and/or salt
formation which is not captured by the “solubility concept” used in Fig. 4b (Furukawa and Takahashi, 2011). In addition, it should be noted that the application of Henry’s law to aerosol water might not be strictly valid since aerosol water does not represent an ideal solution. In a more thorough approach, other thermodynamic parameters such as activity coefficients and fugacity should be taken into account.

In cloud water, $F_p$ for all species is much higher and even species with $K_H = 10^5 \text{ M atm}^{-1}$ might partition to a significant extent to the aqueous phase. Such species might include acids (at enhanced pH), and small monoaldehydes. However, these species will not necessarily contribute to aqSOA mass upon cloud evaporation as they are too volatile. For comparison, two inorganic aerosol compounds (HNO$_3$ and NH$_3$) are added to Fig. 4b as well. While they will be nearly completely dissolved in cloud water, aerosol water does not represent a significant sink for their total budgets. This estimate does not include the formation of salts in the particle phase which will shift the effective $F_p$ of HNO$_3$/NO$_3^-$ and NH$_3$/NH$_4^+$ to more efficient partitioning.

### 4.3 Predicted formation of gasSOA and aqSOA from common precursors

#### 4.3.1 Definition of gasSOA and aqSOA yields

The partitioning of individual species does not give any information about the absolute SOA mass as it neglects reaction times and stoichiometric coefficients. Smog chamber experiments report gasSOA yields as the mass ratio of formed SOA mass and the consumed precursor hydrocarbon (HC) mass for a given set of experimental conditions (absorbing aerosol mass, oxidant levels).

$$Y(\text{SOA}) = \frac{m(\text{SOA})}{\Delta [\text{HC}]}$$  \hspace{1cm} (6)

In yield parameterizations according to the 2-product model, it is assumed that the condensable gasSOA mass can be represented by two proxy compounds that are formed with stoichiometric factors $\alpha_1$, $\alpha_2$ and have partitioning coefficients $K_{p1}$, and $22335$
K_{p2}, respectively (Odum et al., 1996). The gasSOA yield is constant for a given set of experimental conditions (precursor, NO_{x} level, mass loading etc.) since chemical loss processes on dry SOA are slow (Molina et al., 2004; Petters et al., 2006) as they only occur on the particle surface and not in the bulk. NO_{x} levels in the gas phase determine the fate of peroxy radicals and the product distribution which, in turn, affect the amount of formed condensable species, with preferential peroxide formation at low NO_{x} levels (Fig. 5). Parameter sets for \alpha_1, \alpha_2, K_{p1}, and K_{p2} for various precursors and NO_{x} conditions are included in current gasSOA models.

Most aqueous phase laboratory experiments are performed in bulk aqueous phase, i.e. reported “aqSOA yields” refer to the consumed precursor aqueous phase compound (e.g., Carlton et al., 2007; El-Haddad et al., 2009; Lim et al., 2010; Perri et al., 2010). Maximum aqSOA yields of up to 100\% from aqueous phase glyoxal (Lim et al., 2010) and nearly 70\% from aqueous phase methylglyoxal (Tan et al., 2010) have been reported; aqSOA yields for aqueous MACR and MVK are significantly smaller with 2–12\% and 5–10\%, respectively (ElHaddad et al., 2009; Zhang et al., 2010). In all experiments, aqSOA yields were observed to be time-dependent with an initial increase and a later decrease due to oxidation of aqSOA products to CO_{2}. Unlike reported gasSOA yields, these reported laboratory-derived aqSOA yields (even for a given processing time) cannot directly be implemented into multiphase models as they only refer to the conversion of the aqueous phase species and thus represent upper limits of the yields related to the gas phase species. Aquous phase concentrations are usually not predicted by models that exceed studies on a process level. AqSOA precursors (such as glyoxal, methylglyoxal, MVK, MACR) also have loss terms in the gas phase (oxidation, photolysis) that are suppressed in bulk laboratory experiments. In the atmosphere, the overall aqSOA yield, related to the gas phase concentration of a precursor, depends on the relative strength of gas phase loss terms and uptake and further processing in the aqueous phase. A more thorough assessment of the amount of aqSOA formed from specific precursors can only be obtained by simulating multiphase systems that take into account gas and aqueous phase processes simultaneously.
While low NO\textsubscript{x} levels often lead to higher production rates of low volatility compounds and gasSOA products, high NO\textsubscript{x} levels preferentially form high volatility, highly water-soluble species (Fig. 5). However, on larger (spatial, temporal) scales similar concentrations of carbonyl compounds (glyoxal, methylglyoxal) have been predicted under all NO\textsubscript{x} conditions since slow conversions of peroxides and organic nitrates might yield similar products (Fu et al., 2009). Thus, for aqSOA yields a clear separation into “high NO\textsubscript{x}” and “low NO\textsubscript{x}” conditions is not as obvious as it is for gasSOA and on large scales similar aqSOA yields in both scenarios can be expected. In summary, the time dependence of aqSOA reactions in the aqueous phase, and the time dependence of the NO\textsubscript{x} dependent formation of water-soluble aqSOA precursors to NO\textsubscript{x} concentrations make it difficult to compare aqSOA and gasSOA yields on a common basis. In addition, the partitioning of low-volatility or semivolatile species between gas, organic and aqueous phases depends on the abundance of the respective particulate phases. Despite these complex interactions, we will give in the following section some rough estimates of SOA yields and mass predictions from box model calculations using the data in Tables 1 and 2 (processes marked with (*)) in order to provide some guidance to identify conditions under which aqSOA formation is likely to be important.

### 4.3.2 Comparison of gasSOA and aqSOA yields

Cloud aqSOA yields (mass of aqSOA per mass of isoprene reacted in the gas phase) from multiphase isoprene oxidation over several hours (i.e. relatively short time scales) have been parameterized as a function of NO\textsubscript{x} levels, cloud contact time and cloud LWC (Ervens et al., 2008). These yields ranged from 3–9 % for high NO\textsubscript{x} levels and 0.3–0.7 % for low NO\textsubscript{x} levels. GasSOA yields show the opposite trends with NO\textsubscript{x} levels and range from 1–4 % (Kroll et al., 2005a; Henze and Seinfeld, 2006).

A similar approach has been taken here to compare aerosol aqSOA yields and gas-SOA yields for various precursors: a box model has been initialized with a single precursor compound (benzene, toluene, xylene or isoprene) and a constant OH concentration of 10\textsuperscript{6} cm\textsuperscript{-3}. GasSOA products comprise two products (P1, P2) according to the
2-product model. In addition to these condensable products, all of the selected precursors form glyoxal and methylglyoxal (except benzene) in the gas phase and eventually form aqSOA in the aqueous phase:

\[
\text{Precursor} + \text{OH} \rightarrow \alpha_1 P_1 + \alpha_2 P_2 + \text{glyoxal} + \text{methylglyoxal} \rightarrow \text{gasSOA forming aqSOA}
\]  

(7)

GasSOA yields can be calculated based on the 2-product model or the VBS based on

\[
Y\text{(gasSOA)} = \sum_{i=1,n} \left[ m_{\text{org}} \frac{\alpha_i K_{p,i}}{1 + m_{\text{org}} \cdot K_{p,i}} \right]
\]  

(8)

where \(n\) is either 2 (2-product model) or represents the number of different volatility bins oxidation products are distributed to (VBS). NO\textsubscript{x} dependent glyoxal and methylglyoxal yields have been determined for aromatics and isoprene in (dry) smog chamber experiments (Spaulding et al., 2003; Nishino et al., 2010). Total glyoxal and methylglyoxal yields from isoprene are less constrained as these species are not only first-order but also second-order products that are formed from the major oxidation products MACR and MVK (Galloway et al., 2011).

High-NO\textsubscript{x} smog chamber experiments of isoprene oxidation have shown that RH does not change the total SOA mass but significantly impacts the product distribution due to higher yields of glycolaldehyde and hydroxyacetone at high RH (Nguyen et al., 2011). In this study, no differences in methylglyoxal levels were detected and the impact on glyoxal was not evaluated. Under low NO\textsubscript{x} conditions, a decrease in SOA yields of a factor 2–3 under high RH was observed (Zhang et al., 2011). This trend is in agreement with the higher yields of carbonyl compounds under high NO\textsubscript{x} conditions that might result in more aqSOA formation. Aqueous phase chemistry is not directly dependent on NO and NO\textsubscript{2} as they have low solubility and thus their role as aqueous phase oxidants is minor; the major influence of NO\textsubscript{x} on aqSOA yields lies in the preceding gas
phase chemistry that determines the concentration levels of aqSOA precursors. In both studies, it was found that at high RH, contributions of organonitrates increased whereas at low RH, ester formation was predominant. A similar trend of SOA yields with RH was observed for SOA formation from α-pinene; however, in this study, experiments were only performed at 2% ≤ RH ≤ 50% (Cocker et al., 2001a). Parikh et al. (2011) inferred a significant contribution of aqSOA to total SOA from aromatics based on box model simulations of chamber experiments over a wide range of RH with highest SOA yields at high RH. These results seem at odds with earlier results of the same reaction system that did not find any effect of RH on SOA yields form aromatics (Cocker et al., 2001b); however, in these studies the range of RH was limited (≤ 50%) and thus the amount of aerosol water on the ammonium sulfate seed aerosol was minimal.

The box model applied here includes Eq. (7) for benzene, toluene, xylene, and isoprene as representative SOA precursors. For gasSOA yield predictions, the same module as in a previous SOA model study has been used (Bahreini et al., 2009); a multiphase mechanism as depicted in Eq. (7) was assumed for an aerosol population that consisted of an of 2 µg m⁻³ organic mass and 20 µg m⁻³ aerosol water. The dependence of gas phase carbonyl yields on NOₓ levels are reflected in the trends of aqSOA yields with higher yields at high NOₓ, in contrast to gasSOA yields (Fig. 5). However, it should be cautioned here again, that the carbonyl yields are based on chamber experiments over relatively short time scales that might not capture the conversion of ‘low NOₓ products’ into carbonyl compounds (Fig. 5). The results in Fig. 6 show that benzene likely does not form high amounts of aqSOA for the assumed conditions. The data for aqSOA formation from phenol is not included in the current estimate even though in laboratory experiments it has been observed that ring-retaining products from aqueous phase phenol oxidation might form additional aqSOA (Sun et al., 2010) (Sect. 2.1.2). For toluene and xylene, methylglyoxal yields are higher than those for glyoxal; however, the Henry's law constant of methylglyoxal is much smaller than that of glyoxal and thus its transport into the aqueous particles is slower resulting in smaller aqSOA yields. Thus, the overall aqSOA yields from toluene and xylene (aqSOA yield at low
NO\textsubscript{x} 1.9\%, 3.5\%; aqSOA yield at high NO\textsubscript{x}: 2.3\%, 2.3\%) are mostly from glyoxal.

The aqSOA yields from isoprene are comparable to gasSOA yields for the conditions chosen here. Direct contributions of MVK and MACR to aqSOA are not included in the current model approach. These latter products are not very water-soluble but have shown to form aqSOA once dissolved (ElHaddad et al., 2009; Noziere et al., 2010b; Zhang et al., 2010). Their vapor pressure is also reasonably low that they might form gasSOA and thus are partially captured by P1 and P2. It should be noted that in case of efficient removal of MACR and MVK by the aqueous phase, gasSOA yields as derived from (dry) smog chamber experiments might be smaller and the numbers in Fig. 6 might represent an overestimate for gasSOA. This overlap in low volatility and water-solubility makes it difficult to derive accurate yields. The trends in our yield predictions follow those from a previous model study where significant aqSOA formation from isoprene was predicted (Couvidat and Seigneur, 2011). However, both this latter study and our study assume identical partitioning parameters ($\alpha_i$, $K_{p,i}$) for isoprene under high and low RH conditions although recent chamber experiments suggest that at high RH aqSOA formation might play a role and this cannot be fully parameterized by Eq. (8) (Nguyen et al., 2011).

4.3.3 Comparison of gasSOA and aqSOA mass predictions

The comparison of gasSOA and aqSOA yields provided here describes the relative propensity of a compound to form gasSOA versus aqSOA. However, the absolute importance of gasSOA versus aqSOA pathways to the total SOA budget depends also on the availability of liquid water and the relative availability of the various precursors. In Table 3, four sets of conditions are summarized representing a wide range of biogenic/anthropogenic concentrations, and dry/humid scenarios. These conditions represent averages of observations even though they do not reflect the variability as observed in these regions. We do not attempt to predict the exact observed SOA masses in these regions as the simplicity of a box model does not allow the simulation of dynamic (e.g., transport, mixing, dilution) and detailed chemical processes (e.g.,
photochemistry). We rather use the input data in Table 3 to compare gasSOA versus aqSOA formation in the context of different VOC and RH conditions.

Figure 7 shows the predicted gasSOA and aqSOA masses for the four cases specified in Table 3. We show gasSOA predictions for both the 2-product model and the VBS; in agreement with other studies, VBS gives higher gasSOA predictions as also semivolatile compounds are taken into account. In anthropogenic influenced areas at high RH (scenarios I and III), the contribution of aqSOA to the total predicted SOA mass is about 30–50% despite the relatively high glyoxal and methylglyoxal yields from aromatics. In the biogenically-dominated scenario (II) SOA formation from gas and aqueous phase pathways are roughly equally important. The opposite trend of carbonyl (glyoxal, methylglyoxal, hydroxyacetone, glycolaldehyde) and gasSOA yields could have a significant impact on SOA predictions. Whereas under high NO_x conditions the gasSOA yields from isoprene are small, the importance of aqSOA increases. Several studies have pointed to the role of anthropogenic compounds (NO_x) in oxidation of biogenic carbon emissions that lead to SOA formation (Weber et al., 2007; Carlton et al., 2010; Spracklen et al., 2011). However, such interactions are underestimated by current gasSOA approaches. At very low RH (scenario IV), none of the chosen precursors will significantly contribute to aqSOA formation due to the negligible amount of aerosol water. This latter result is in agreement with a model study in Houston, TX, where reasonable agreement between predicted and observed (airborne) SOA masses where obtained based on gasSOA yields (Bahreini et al., 2009; Zhang and Ying, 2011). In summary, Fig. 7 highlights the sensitivities of selected regions for aqSOA formation that should be explored in more detail in future studies. These studies suggest that aqSOA formation will be most important in humid regions that are influenced by biogenic emissions.
5 Outlook: knowledge gaps and needed research

5.1 Laboratory studies of aqueous and multiphase processes

Aqueous phase experiments in highly concentrated organic and inorganic aqueous solutions have provided a relatively well-constrained set of kinetic data and chemical mechanisms for OH radical oxidation and other processes of carbonyl compounds in dilute and concentrated solutions (cloud- and aerosol water-like, respectively). However, several mysteries remain which affect the accuracy of aqSOA model prediction. Several studies suggest that the partitioning of glyoxal and methylglyoxal (and likely related compounds) into aqueous seed aerosol is much higher than in pure water. This increased uptake was assumed to be related to oligomerization which is associated with relatively long time scales until equilibrium is reached; however, the dependence of different seed aerosol compositions on observed $K_H^*$ and kinetic data in seed aerosols could not be explained (Ervens and Volkamer, 2010). A recent study suggests that the shift of glyoxal partitioning towards the particle phase is due to a stabilization of the glyoxal dihydrate (Yu et al., 2011). However, this effect is only described qualitatively and awaits a better quantification and numerical description to be implemented in models. Similar to enhanced partitioning of organics, several studies point to enhanced levels of reactive oxygen species in particles (Hasson and Paulson, 2003; Arellanes et al., 2006; Shen et al., 2011). The reason for these deviations from thermodynamic equilibrium ($K_H$) is not clear and related modeling assumptions of oxidant transport into and within aerosol particles and the resulting oxidant levels are a significant source of uncertainty in aqSOA prediction.

Water-soluble organic and inorganic compounds might form separate phases within an aerosol particle (Zuend et al., 2008; Bertram et al., 2011). Such effects will affect the availability of reactants of inorganic/organic cross-reactions that lead to organosulfates or organo-nitrogen compounds. It seems unfeasible for the near future to develop and implement detailed thermodynamic models that take such effects into account. However, laboratory experiments should be designed to examine the extent to which...
such phase separation indeed occurs and affects aqSOA formation rates.

Most laboratory experiments are designed as homogeneous aqueous phase experiments and thus observed aqSOA formation are interpreted as bulk reaction parameters. Some multiphase experiments that investigate the uptake and further processing of aqSOA precursors into aqueous aerosols, have shown evidence that aqSOA formation could also occur at the gas/particle interface (Ervens and Volkamer, 2010; Grgic et al., 2010; Huang et al., 2011). Size-resolved measurements of processed aerosol distributions will give evidence of the importance of either type of process since the relative contributions of bulk versus surface aqSOA formation will determine the size distribution of the predicted SOA. In general, there is a lack of experiments of SOA formation at RH above the deliquescence RH or even above the CCN activation threshold (droplet formation) of seed aerosol. Thus, aerosol and cloud chambers that allow the characterization of the chemical multiphase system are a promising tool to elucidate further details of aqSOA formation (e.g., Wang et al., 2011a).

5.2 Identification of additional aqSOA precursors

While our discussion of aqSOA formation is mostly focused on glyoxal and methylglyoxal, other small compounds undergo similar processes (e.g., glycolaldehyde, pyruvic acid, other small acids, peroxides, and alcohols). While in laboratory studies it has been shown that such compounds can indeed form HMWC, it is likely that also other highly functionalized compounds with longer carbon chains (>C3) are taken up into cloud or aerosol water and become processed to form HMWC or acids. While the contributions of larger dicarboxylic acids in cloud water are small as compared to the total OC masses, the contributions of their oligomers in aerosols might be considerable.

5.3 Role of parameters that affect gas phase chemistry: RH and NOx

Most smog chamber experiments are performed under (nearly) dry conditions and thus yields of gas phase products of SOA precursors are usually reported for those
conditions. However, as shown in a recent experiment, the amount of water vapor can affect the product distribution from isoprene oxidation (Nguyen et al., 2011). While coincidentally the SOA yields under high and low RH were the same in these experiments, the product distribution is different which will have impact on the properties of SOA mass (solubility, optical properties etc.). The few available studies that address the role of NOx on SOA formation under high RH suggest the formation of aqSOA (Nguyen et al., 2011; Zhang et al., 2011). However, more systematic studies are needed to identify the product distributions from a greater variety of precursors in order to refine both the gasSOA parameters (α, Kp) and aqSOA yields.

5.4 Identification of tracers of ambient aqSOA

While oxalate is a good tracer for aqSOA formation, its formation rate in aerosol water might be small due to the preference of accretion reactions. Organosulfates that can only be ascribed to aqSOA processes have been identified in ambient samples (Froyd et al., 2010; Hatch et al., 2011; Kristensen and Glusius, 2011; Olson et al., 2011). These compounds only comprise a small fraction of total OC; however, their formation pathways are unique and thus point to aqSOA formation. It is likely that specific HMWC, patterns of HMWC or subunits that add multiple times to HMWC, identified in rain or fog water (Altieri et al., 2009; Mazzoleni et al., 2010), can be used to deduce the aqueous processing of aerosol particles. One value in identifying such tracers is that they could potentially be used to estimate aqSOA contributions via receptor modeling techniques to test our understanding of underlying formation mechanisms using detailed process aqSOA models, like oxalate prediction was used to test the understanding of aqSOA in a recent global model (Myriokefalitakis et al., 2011). Such work would benefit from the measurement of aqSOA tracers in regions where aqSOA formation is likely, such as in regions of cloud outflow or forested humid regions.

As discussed in Sect. 3.2, aqSOA formation modifies aerosol size distributions in a different way than gasSOA condensation with pronounced mass addition to the droplet mode. Thus, size-resolved, instead of only bulk, OA measurements (e.g. AMS spectra)
should be able to provide insights into SOA formation pathways. AqSOA products have in general low volatility and thus might significantly contribute to low volatility oxygenated organic aerosol (LV-OOA). Specific compounds, such as oxalate, even exceed the range of carbon oxidation state and O/C ratio as covered by LV-OOA (Kroll et al., 2011). AMS data are often characterized by means of positive matrix factorization (PMF) which identifies types of aerosol based on similarities in spectra (Ulbrich et al., 2009); available spectra in scenarios where aqSOA formation can be expected should be revisited to identify features that might point to aqSOA formation.

5.5 Evaluation of gasSOA versus aqSOA formation in more comprehensive models

The time scales of gasSOA versus aqSOA formation in different scenarios could be compared by using the framework by Kerminen and Wexler (1995) that gives general equations for aqueous phase and condensation processes as a function of ambient parameters. Such a comparison can be used to evaluate the relative importance of such processes and to identify areas and conditions for which either SOA source might be important.

Several models have included cloud aqSOA formation and have compared its efficiency to gasSOA formation; however, the implementation of the full, detailed chemistry as listed in Tables 1 and 2 represents a major computational burden and thus simpler expressions are needed. Whereas first attempts have been made to parameterize cloud aqSOA yields from isoprene (Ervens et al., 2008), this expression cannot be used to simulate aqSOA in anthropogenic scenarios or in aerosol water. Thus, a more general expression for water-soluble, direct aqSOA precursors (e.g., glyoxal, methylglyoxal) would be desirable.
The reason for the lack of comprehensive model studies of aerosol aqSOA formation is the missing comprehensive data sets for the underlying chemical processes. The parameters in Tables 1 and 2 provide such data sets and can be used in process models. Upon such model simulations, appropriate and simple expressions can be developed for use in large scale models. Thus, in terms of model applications, the development of simple but robust expressions that parameterize the formation of aerosol aqSOA is sorely needed. These expressions should be carefully tested against laboratory measurements and models that allow well-constrained detailed chemistry modules, i.e. in box or parcel models that are restricted in terms of microphysical/dynamic processes but are able to include many chemical species and processes.

6 Summary and conclusions

The current review summarizes evidence that secondary organic aerosol forms in the aqueous phase of cloud/fog/rain droplets and in aerosol water as was postulated about a decade ago (Blando and Turpin, 2000). While this former “plausibility study” focused mostly on processes in dilute aqueous phase (droplets), many recent laboratory experiments and ambient aerosol samples suggest that more complex chemical reactions can occur in aerosol water forming accretion products such as oligomers. Kinetic data from such laboratory experiments for photochemical and dark reactions are summarized. Model results using this data that compare aqSOA formation in cloud and aerosol water suggest that over comparable time scales both sources are comparably important. Considering that individual cloud droplets only exist for several minutes whereas water-containing aerosol particles reside in the atmosphere over hours or days might suggest a more significant role for aerosol aqSOA formation. Comparisons of aqSOA formation in aerosol water to gasSOA formation suggest that aqSOA formation contributes substantially to the total SOA mass, in particular in regions of high RH, biogenic influence and high NOx conditions. However, explorative box model calculations performed here are very restricted in their large scale predictability.
Major uncertainties in current aqSOA estimates are identified and suggestions are made for further laboratory experiments and model development activities. We also identify a need for aqSOA tracer compounds to be used for receptor models in order to test our understanding of the chemical processes summarized based on laboratory studies. It can be expected that aqSOA will gain in importance in future climate as elevated temperature and higher RH increase carbonyl yields and thus aqSOA yields will likely increase (Hennigan et al., 2008c; Slowik et al., 2010; Hoyle et al., 2011).

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1983.


Petters, M. D., Prenni, A. J., Kreidenweis, S. M., DeMott, P. J., Matsunaga, A., Lim, Y. B., and


Table 1. Summary of experimental conditions, kinetic data and products from photochemical experiments leading to aqSOA.

<table>
<thead>
<tr>
<th>Organic reactant concentration</th>
<th>Oxidant (Source)</th>
<th>Inorganic solutes</th>
<th>Rate constant</th>
<th>Products</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glyoxal</td>
<td>1 mM OH (Pulse radiolysis)</td>
<td>–</td>
<td>$1.1 \times 10^9 \text{ M}^{-1} \text{s}^{-1}$ (*)</td>
<td>Formic, acetic, glyoxylic, acetic acid</td>
<td>Buxton et al. (1997)</td>
</tr>
<tr>
<td></td>
<td>0.5 mM OH (0.5 mM H$_2$O$_2$)</td>
<td>–</td>
<td>–</td>
<td>Oligomers</td>
<td>Karpel Vel Leitner and Döré (1997)</td>
</tr>
<tr>
<td></td>
<td>2 mM OH (10 mM H$_2$O$_2$)</td>
<td>–</td>
<td>–</td>
<td>Glyoxylic, oxalic, acids Oligomers</td>
<td>Carlton et al. (2007)</td>
</tr>
<tr>
<td></td>
<td>0.03 mM 0.3 mM 3 mM OH</td>
<td>–</td>
<td>$1.1 \times 10^8 \text{ M}^{-1} \text{s}^{-1}$</td>
<td>Succinic, oxalic, malonic, malic, tartaric, mesoxalic acid - Transition from “mainly acids” to “mainly oligomers” - Oligomers ($\Delta = \text{C}_2\text{H}_2\text{O}_3$)</td>
<td>Tan et al. (2009); Lim et al. (2010)</td>
</tr>
<tr>
<td></td>
<td>~0.1 – 1 M OH (~20 ppm C$_2$H$_2$O$_3$)</td>
<td>~6 M (NH$_4$)$_2$SO$_4$</td>
<td>$0.8 \text{ s}^{-1} &lt; k &lt; 7 \text{ s}^{-1}$ (*)</td>
<td>Fitted $k$ in order to explain observed SOA mass in excess to OH reaction; $k$ scales with OH concentration with highest value at $[\text{OH}] = 10^7 \text{ cm}^{-3}$</td>
<td>Volkamer et al. (2009); Ervens and Volkamer (2010)</td>
</tr>
<tr>
<td></td>
<td>~0.1 – 1 M UV</td>
<td>~6 M (NH$_4$)$_2$SO$_4$</td>
<td>Glycolic acid sulfate, C-N compounds</td>
<td>Galloway et al. (2009)</td>
<td></td>
</tr>
<tr>
<td>Glycolaldehyde</td>
<td>1 mM OH (5 mM H$_2$O$_2$)</td>
<td>–</td>
<td>Succinic, malonic, oxalic, glycolic, glyoxylic acids</td>
<td>Perri et al. (2009)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1 mM OH (5 mM H$_2$O$_2$)</td>
<td>1 mM H$_2$SO$_4$</td>
<td>Organosulfates (esters)</td>
<td>Perri et al. (2010)</td>
<td></td>
</tr>
</tbody>
</table>
Table 1. Continued.

<table>
<thead>
<tr>
<th>Methylglyoxal</th>
<th>0.1–4 mM</th>
<th>OH (0.3 µM) (0.5 mM H₂O₂)</th>
<th>–</th>
<th>7 × 10⁸ M⁻¹ s⁻¹</th>
<th>–</th>
<th>Ervens et al. (2003b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 mM</td>
<td>OH (10 mM H₂O₂)</td>
<td>–</td>
<td>Glyoxyl, pyruvic, oxalic acids and oligomers with C₃H₄O₂ subunits</td>
<td>Altieri et al. (2008); Tan et al. (2010)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\[ k = 4 \text{ s}^{-1} (*) \]

Fitted \( k \) based on observed aqSOA formation in laboratory studies Tan et al. (2010); scales with [OH] with highest value at [OH] = 10⁷ cm⁻³

This study

For fitting, cf. procedure described for glyoxal by Ervens and Volkamer (2010)

Pyruvic acid

<table>
<thead>
<tr>
<th>1–12 mM</th>
<th>OH (0.3 µM) (0.5 mM H₂O₂)</th>
<th>–</th>
<th>–</th>
<th>Ervens et al. (2003b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>10 mM</td>
<td>OH (20 mM H₂O₂)</td>
<td>–</td>
<td>Glyoxyl, acetic acid</td>
<td>Altieri et al. (2006)</td>
</tr>
<tr>
<td>5 mM, 10 mM</td>
<td>OH (20–100 mM H₂O₂)</td>
<td>–</td>
<td>Glyoxyl, acetic acid</td>
<td>Carlton et al. (2006)</td>
</tr>
<tr>
<td>100 mM</td>
<td>UV</td>
<td>–</td>
<td>C₆, C₇ carboxylic acids</td>
<td>Guzman et al. (2006)</td>
</tr>
<tr>
<td>5 mM, 10 mM</td>
<td>OH (100 mM, 20 mM H₂O₂)</td>
<td>–</td>
<td>Glyoxyl, acetic acid</td>
<td>Carlton et al. (2006)</td>
</tr>
<tr>
<td>Pyruvic + Oxalic acid</td>
<td>100 mM</td>
<td>UV</td>
<td>–</td>
<td>C₆, C₇ carboxylic acids</td>
</tr>
<tr>
<td>5 mM each</td>
<td>O₃ (UV light)</td>
<td>–</td>
<td>Pyruvic acid dimer and Pyruvic/formic acid-dimer formation over ~5 h (photosensitized)</td>
<td>Grgic et al. (2010)</td>
</tr>
</tbody>
</table>
Table 1. Continued.

<table>
<thead>
<tr>
<th></th>
<th>MACR</th>
<th>OH (8 mM H₂O₂)</th>
<th>–</th>
<th>5.8×10⁹ M⁻¹ s⁻¹ methylglyoxal, formaldehyde, hydroxycetone and acetic acid/acetate Pyruvic, glyoxylic acid Oligomer series with hydroxyl carbonyl and carboxylic acids</th>
<th>Liu et al. (2009) El-Haddad et al. (2009)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0.05 mM</td>
<td>OH (0.4 mM H₂O₂)</td>
<td>–</td>
<td>3.7 M (NH₄)₂SO₄ or 1 M Na₂SO₄</td>
<td>≥ 10⁻⁵ s⁻¹ 2.5×10⁹ M⁻¹ s⁻¹ Organosulfates 150 &lt; M &lt; 300</td>
</tr>
<tr>
<td></td>
<td>2–5 mM</td>
<td>SO₄²⁻ (UV)</td>
<td>–</td>
<td>–</td>
<td>Glyoxal, Methylglyoxal, Pyruvic, Acetic, Oxalic, malonic acids, HMWC</td>
</tr>
<tr>
<td></td>
<td>100–500 mM</td>
<td>SO₄²⁻ (UV)</td>
<td>–</td>
<td>–</td>
<td>Organosulfates</td>
</tr>
<tr>
<td></td>
<td>MVK</td>
<td>OH (2 mM H₂O₂)</td>
<td>–</td>
<td>8×10⁸ M⁻¹ s⁻¹</td>
<td>Light absorbing dimer (phenoxy radical)</td>
</tr>
<tr>
<td></td>
<td>Isoprene</td>
<td>11 mM SO₄²⁻</td>
<td>–</td>
<td>–</td>
<td>Organosulfates</td>
</tr>
<tr>
<td></td>
<td>α-pinene</td>
<td>150 mM SO₄²⁻</td>
<td>–</td>
<td>–</td>
<td>Organosulfates</td>
</tr>
<tr>
<td></td>
<td>3,5-dihydroxy benzoic acid</td>
<td>0.02 mM OH (0.1 mM H₂O₂ + 0.05 mM FeCl₃)</td>
<td>–</td>
<td>–</td>
<td>Small organic acids, phenolic dimers and higher oligomers formed from phenoxy radicals</td>
</tr>
<tr>
<td></td>
<td>Phenol</td>
<td>0.1 mM OH (0.1 mM H₂O₂)</td>
<td>–</td>
<td>–</td>
<td>Absorbing products (λ = 500 nm) Spectra structure depending on organic acids</td>
</tr>
<tr>
<td></td>
<td>Guaiacol</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Syringol</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Phenols, Vanillyls, Syringols</td>
<td>10 mM OH (1 mM H₂O₂)</td>
<td>–</td>
<td>–</td>
<td></td>
</tr>
</tbody>
</table>

(*) These rate constants are used in the box model studies in Sect. 4.
Table 2. Summary of experimental conditions, kinetic data and products from dark (non-photochemical) reactions leading to aqSOA.

<table>
<thead>
<tr>
<th>Organic reactant concentration</th>
<th>Reactant/Solutes</th>
<th>Rate constant</th>
<th>Products</th>
<th>Comments</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glyoxal ~0.1 M</td>
<td>5 M Amm Sulf</td>
<td>100 M$^{-1}$s$^{-1}$; 0.1 M$^{-1}$s$^{-1}$</td>
<td>Fitted rate constant for glyoxal oligomerization (forward/back reaction)</td>
<td>Volkamer et al. (2009); Ervens and Volkamer (2010)</td>
<td></td>
</tr>
<tr>
<td>1 M Amine</td>
<td></td>
<td>0.3 M$^{-1}$s$^{-1}$</td>
<td>Imine oligomers</td>
<td>(De Haan et al., 2009c)</td>
<td></td>
</tr>
<tr>
<td>0.1 – 1 M Amino acids</td>
<td></td>
<td>4.5 x 10$^{-4}$ &lt; $k$ [M$^{-1}$s$^{-1}$] &lt;0.12</td>
<td>Acetal oligomers, imidazoles</td>
<td>Depending on amino acid</td>
<td>De Haan et al. (2009a)</td>
</tr>
<tr>
<td>NH$_4^+$</td>
<td></td>
<td>2x 10$^{-10}$ exp(1.5$\cdot$n$_{NH4}$) exp(2.5$\cdot$ pH) M$^{-1}$s$^{-1}$ (*)</td>
<td>Iminium intermediates, imidazoles</td>
<td>Empirical expression based on laboratory studies</td>
<td>Noziere et al. (2009)</td>
</tr>
<tr>
<td>1.5 M NH$_4^+$ (1 M)</td>
<td></td>
<td>(2.01±0.4) x 10$^{-12}$ M$^{-1}$s$^{-1}$</td>
<td>Imidazoles</td>
<td>Yu et al. (2011)</td>
<td></td>
</tr>
<tr>
<td>Methylglyoxal</td>
<td></td>
<td>–</td>
<td>Imidazoles, iminium</td>
<td>Shapiro et al. (2009)</td>
<td></td>
</tr>
<tr>
<td>2 M NH$_4^+$</td>
<td>≤ 5 x 10$^{-6}$ M$^{-1}$s$^{-1}$ (*)</td>
<td>Oligomeric amino compounds and organosulfates</td>
<td>Time scale of product formation ~1 h</td>
<td>Sareen et al. (2009)</td>
<td></td>
</tr>
<tr>
<td>1 mM (NH$_4$)$_2$SO$_4$, Na$_2$SO$_4$, or H$_2$SO$_4$</td>
<td></td>
<td>Oligomers of hydro ketones (C$_3$H$_4$O$_3$)</td>
<td>Time scale of product formation ~1 h</td>
<td>Yasmineen et al. (2010)</td>
<td></td>
</tr>
<tr>
<td>Methylglyoxal</td>
<td></td>
<td>–</td>
<td>Imidazoles</td>
<td>Kua et al. (2011)</td>
<td></td>
</tr>
<tr>
<td>2 M (total) 3.1 M (NH$_4$)$_2$SO$_4$</td>
<td></td>
<td>Light absorbing hemi acetal, aldol condensation products</td>
<td>Theoretical calculations of prediction of thermodynamic kinetic feasibility</td>
<td>Schwier et al. (2010)</td>
<td></td>
</tr>
<tr>
<td>MACR</td>
<td>~ μM O$_3$</td>
<td>0.003 s$^{-1}$</td>
<td>Hydroxymethylhydroperoxide, HOCH$_2$OH, methylglyoxal</td>
<td>Pseudo-first order $k$ with MACR in excess ($r = 5$ min)</td>
<td>Chen et al. (2008)</td>
</tr>
<tr>
<td>MVK</td>
<td>≤ 3 M$^{-1}$s$^{-1}$</td>
<td>Aldol condensation products</td>
<td>Aldol condensation products</td>
<td>Noziere et al. (2006)</td>
<td></td>
</tr>
<tr>
<td>2–5 mM (~ μM O$_3$)</td>
<td>4 x 10$^{-10}$ M$^{-1}$s$^{-1}$</td>
<td>Pyruvic, malonic, oxalic acid, methylglyoxal, glyoxal, and their oligomers</td>
<td>Theoretical estimate based on generic epoxide</td>
<td>Minerath and Elrod (2009)</td>
<td></td>
</tr>
<tr>
<td>Epoxide hydrolysis</td>
<td>–</td>
<td>0.074 &lt; $k$ [M$^{-1}$s$^{-1}$] &lt;15</td>
<td>Aldol condensation products</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(*) These rate constants are used in the box model studies in Sect. 4.
Table 3. Initial conditions for box model studies to evaluate gasSOA and aqSOA mass formation, Eq. (7). Model results are presented in Fig. 7.

<table>
<thead>
<tr>
<th></th>
<th>RH [%]</th>
<th>Dry mass [µg m(^{-3})]</th>
<th>Organic fraction</th>
<th>Benzene [ppb]</th>
<th>Toluene [ppb]</th>
<th>Isoprene [ppb]</th>
<th>Region typical for air mass</th>
</tr>
</thead>
<tbody>
<tr>
<td>I.</td>
<td>70</td>
<td>24</td>
<td>0.4</td>
<td>1</td>
<td>3</td>
<td>0</td>
<td>Los Angeles Hersey et al. (2011)</td>
</tr>
<tr>
<td>II.</td>
<td>90</td>
<td>1</td>
<td>0.75</td>
<td>0</td>
<td>0</td>
<td>2</td>
<td>Amazon Martin et al. (2010)</td>
</tr>
<tr>
<td>III.</td>
<td>80</td>
<td>15</td>
<td>0.5</td>
<td>0.6</td>
<td>3</td>
<td>0</td>
<td>Po Valley Steinbacher et al. (2005); Carbone et al. (2010)</td>
</tr>
<tr>
<td>IV.</td>
<td>45</td>
<td>15</td>
<td>0.67</td>
<td>0.2</td>
<td>0.3</td>
<td>0.4</td>
<td>Houston, TX Bahreini et al. (2009)</td>
</tr>
</tbody>
</table>
Fig. 1. Schematic of gasSOA and aqSOA formation pathways in the gas and aqueous phases of the atmosphere. Dashed arrows denote oxidation reactions.
Fig. 2. Observations of organic carbon or related parameters at different locations. 
Upper panel: Atlanta: (a) Averaged diurnal evolution of temperature, RH and WSOC fraction in the particle phase (\(F_p\)); (b) \(F_p\) as a function of RH; (c) \(F_p\) as a function of total organic carbon 
Middle panel: Mexico City: (d) Temporal evolution of RH and O/C ratio (AMS data and model prediction; Hodzic et al. (2010)); (e) Observed and predicted O/C ratio as a function of RH 
Bottom panel: Los Angeles: (f) Averaged diurnal evolution of temperature, RH and WSOC fraction in the particle phase (\(F_p\)); (g) \(F_p\) as a function of RH; (h) \(F_p\) as a function of total organic carbon.
Fig. 3. Results from parcel model studies using six trajectories (differently colored traces) that differ by temporal evolution of meteorological parameters (temperature, pressure, RH, in-cloud time etc) The chemical mechanism includes reaction parameters for glyoxal and methylglyoxal as listed in Tables 1 and 2 (processes marked with (*)). (a) Time profiles of RH for six different trajectories (color-coded) (b) predicted aqSOA formation in cloud water (c) Predicted aqSOA formation in aerosol water. The prediction of similar aqSOA amounts from both sources is a coincidental result and will strongly depend on time scales parcels spend in/outside of clouds, ambient RH, and cloud LWC.
Fig. 4. Particulate fraction $F_p$ as predicted based on different SOA approaches (a) gasSOA is predicted based on 2-product model or VBS and displayed as a function of the partitioning constant $K_p$ and saturation concentration $C^*$ for different amount of organic aerosol mass. (b) aqSOA is shown as a function of $K_H$ (or $K_{H^*}$). This constant is converted to $K_{p,aq}$ and $C_{aq}^*$ which corresponds to $K_p$ ($C^*$) for gasSOA but relates to water instead of absorbing organic material (Sect. 4.2.2). Values are shown for two different aerosol LWC; trends for cloud LWC are only shown as line.
Fig. 5. Fate of peroxy radicals in the gas phase in different NO\textsubscript{x} scenarios. Dashed arrows depict slow pathways that might only occur on large (spatial, temporal) scales.
Fig. 6. Comparison of gasSOA and aqSOA yields from various precursors in the presence of [OH] = 10^6 cm^{-3} over 12 h. GasSOA yields are based on smog chamber experiments under low (red)/high (blue) NOx conditions and are calculated for an assumed organic aerosol mass of 10 µg m^{-3}. AqSOA yields are derived from box model studies using kinetic data (Tables 1 and 2, data with *) and are derived for 20 µg m^{-3} aerosol water.
Fig. 7. Comparison of predicted gasSOA and aqSOA masses from various precursors in the presence of [OH] = 10^6 cm\(^{-3}\) over 12 h for the conditions as specified in Table 3.