simpleGAMMA – a reduced model of secondary organic aerosol formation in the aqueous aerosol phase (aaSOA)

J. L. Woo and V. F. McNeill

Department of Chemical Engineering, Columbia University, New York, New York, 10027, USA

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Correspondence to: V. F. McNeill (vfm2103@columbia.edu)

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Abstract

There is increasing evidence that the uptake and aqueous processing of water-soluble volatile organic compounds (VOCs) by wet aerosols or cloud droplets is an important source of secondary organic aerosol (SOA). We recently developed GAMMA (Gas–Aerosol Model for Mechanism Analysis), a zero-dimensional kinetic model that couples gas-phase and detailed aqueous-phase atmospheric chemistry for speciated prediction of SOA and organosulfate formation in cloudwater or aqueous aerosols. Results from GAMMA simulations of SOA formation in aerosol water (McNeill et al., 2012) indicate that it is dominated by two pathways: isoprene epoxydiol (IEPOX) uptake followed by ring-opening chemistry (under low-NO$_x$ conditions) and glyoxal uptake. This suggested that it is possible to model the majority of aqueous aerosol phase SOA mass using a highly simplified reaction scheme. We have therefore developed a reduced version of GAMMA, simpleGAMMA. Close agreement in predicted aaSOA mass is observed between simpleGAMMA and GAMMA under all conditions tested (between pH 1–4 and RH 40–80 %) after 12 h of simulation. simpleGAMMA is computationally efficient and suitable for coupling with larger-scale atmospheric chemistry models.

1 Introduction

Quantifying the sources of tropospheric aerosol material is important for accurate modeling of air quality and climate. In-situ processes leading to the formation of new organic aerosol material, collectively known as secondary organic aerosol (SOA) formation, are poorly constrained in atmospheric chemistry models (Hallquist et al., 2009; Kanakidou et al., 2005). Disagreement between model results and observations of the quantity, oxidation state, and location of organic aerosols in the atmosphere has suggested an incomplete representation of SOA formation pathways in models (Heald et al., 2005; Jimenez et al., 2009). In the past decade, the uptake of water-soluble volatile organic compounds (VOCs) into cloud droplets or aerosol water, followed by aqueous-phase
chemical processing, has received increased attention as a possibly important source of SOA (Blando and Turpin, 2000; Ervens et al., 2011). It is thought to be especially significant in the case of isoprene-derived SOA formation. This is because most of the gas-phase oxidation products of isoprene, are, like isoprene itself, highly volatile, however some, like glyoxal (GLYX), isoprene-derived epoxydiols (IEPOX) (Paulot et al., 2009; Surratt et al., 2010), and methacrylic acid epoxide (MAE) (Lin et al., 2013), are water-soluble. These species also undergo reactive processing in the aqueous phase of cloud droplets or aerosols, increasing their uptake from the gas-phase.

Despite mounting evidence that aqueous atmospheric chemistry is a significant source of SOA, aqueous aerosol and cloudwater SOA formation is yet not widely represented in 3-D atmospheric chemistry and air quality models. This is due, in part, to the challenges of bridging scales between the detailed information generated by laboratory experiments, and simplified representations suitable for implementation in 3-D models, which can afford to carry relatively few chemical tracers. Including cloudwater organic chemistry in large scale atmospheric chemistry models has improved agreement with observations (Carlton et al., 2008; Liu et al., 2012; Myriokefalitakis et al., 2011), but aqueous aerosol processes are just beginning to be represented (Knote et al., 2014; Lin et al., 2014; Pye et al., 2013).

Previously, we developed GAMMA (Gas–Aerosol Model for Mechanism Analysis), a zero-dimensional kinetic model that couples gas- and detailed aqueous aerosol-phase chemistry for speciated prediction of SOA and organosulfate formation in the aqueous aerosol phase under ambient or laboratory conditions (McNeill et al., 2012; Woo et al., 2013). GAMMA represents aaSOA formation in terms of bulk aqueous uptake followed by aqueous-phase reaction (Schwartz, 1986). GAMMA includes IEPOX chemistry following Eddingsaas et al. (2010), and uses the effective Henry’s Law constant, $H^*$, constrained by aerosol chamber studies (Sumner et al., 2014) to describe glyoxal uptake and dark reactions, as well as detailed photochemical organosulfate formation and brown carbon formation from glyoxal, methylglyoxal, and acetaldehyde (Woo et al., 2013). For more information regarding other specific mechanisms included
in GAMMA, as well as rate constants for these reactions and other physical parameters, the reader is referred to McNeill et al. (2012) (including the Supporting Information) and Woo et al. (2013).

Simulations using GAMMA indicate that the IEPOX pathway dominates aaSOA formation, leading to up to \( \sim 0.9 \mu \text{g m}^{-3} \) of SOA mass under conditions typical of the rural SE USA (McNeill et al., 2012). Pye and coworkers predicted similar mean concentrations (0.6–1.0 \( \mu \text{g m}^{-3} \)) of IEPOX-derived SOA mass for the SE USA in Summer 2006, using CMAQ with a surface reactive uptake formulation of IEPOX aaSOA formation (Pye et al., 2013). In urban (high-NO\(_x\)) environments, aaSOA is primarily formed via glyoxal uptake (McNeill et al., 2012).

This predominance of two aaSOA formation pathways involving relatively few species, compared to the total number of aqueous compounds tracked by GAMMA, suggests that it is possible to model the majority of aqueous aerosol phase SOA mass using a highly simplified reaction scheme, which is computationally efficient and suitable for coupling with larger-scale atmospheric chemistry models. GAMMA has therefore been used as a guide to develop a reduced mechanism for aaSOA formation, simpleGAMMA. simpleGAMMA reduces the total number of tracked aqueous species from 140 to 4 (glyoxal, IEPOX, 2-methyltetrol, and IEPOX organosulfate), with 2 species partitioning between the gas and aqueous aerosol phases (glyoxal and IEPOX), and a single aqueous-phase chemical process (reactive uptake of IEPOX), compared to 118 in GAMMA.

2 simpleGAMMA: model description

As in GAMMA, the time evolution of the aqueous aerosol phase concentration \( (C_i, \text{ in mol L}^{-1}) \) of a given chemical species \( i \) is described in simpleGAMMA by the following differential equation (Schwartz, 1986):

\[
\frac{dC_i}{dt} = \text{rate of change of } C_i
\]
\[
\frac{dC_i}{dt} = \frac{k_{mt,i}}{RT} P_i - \frac{k_{mt,i}}{H_i^*RT} C_i + \sum_k r_{ik,aq}
\]

(1)

Here, \(P_i\) is the gas-phase partial pressure of species \(i\), \(H_i^*\) is the effective Henry's Law constant, \(R\) is the universal gas constant, and \(T\) is temperature. The rates \(r_{ik,aq}\) represent chemical reactions in the aerosol phase that can act as sources or sinks for a given species. \(k_{mt,i}\) is the gas-aerosol mass transfer coefficient for species \(i\), given by:

\[
k_{mt,i} = \frac{1}{\frac{R^2}{3D_{g,i}} + \frac{4R}{3\omega_i\alpha_i}}
\]

(2)

where \(R\) is the aerosol particle radius, \(D_{g,i}\) is the gas-phase diffusion coefficient, \(\omega_i\) is the thermal velocity, and \(\alpha_i\) is the accommodation coefficient. A suitable gas-phase chemical mechanism should be employed, and the loss or gain of species to/from the aerosol phase should be accounted for, following, for example:

\[
\frac{dP_i}{dt} = -k_{mt,i} a_L P_i + \frac{k_{mt,i} a_L}{H_i^*} C_i + \sum_j r_{ij,\text{gas}} + E_i - D_i
\]

(3)

where \(a_L\) is the aerosol aqueous liquid volume fraction (cm\(^3\) cm\(^{-3}\) of air), \(r_{ij,\text{gas}}\) is the rate of gas phase reaction \(j\) that species \(i\) participates in, and \(E_i\) and \(D_i\) are the emission and deposition rates of species \(i\), respectively.

Note that simpleGAMMA is a reduced version of the *aqueous-phase mechanism* of GAMMA (McNeill et al., 2012). The gas-phase mechanism of GAMMA was not changed because it is intended that simpleGAMMA take gas phase concentration fields as inputs from an external source, i.e. from field measurements or from existing models of atmospheric chemistry, which have gas-phase chemical mechanisms but lack representations of aqueous aerosol phase SOA formation. For tests reported here, we ran
simpleGAMMA with the full gas phase mechanism of GAMMA, following Eqs. (1)–(3). A full description of the gas and aqueous phase mechanisms of GAMMA, the simulation conditions, and results can be found in McNeill et al. (2012). We note that, subsequent to the publication of McNeill et al. (2012), the gas and aqueous phase chemistry of methacrylic acid epoxide (MAE) was introduced to the full version of GAMMA following Lin et al. (2013). The predicted contribution of this pathway to aaSOA was minor compared to IEPOX and glyoxal, consistent with the findings of Pye et al. (2013). Therefore, it is not included in simpleGAMMA.

The processes leading to aaSOA formation in simpleGAMMA are a subset of those represented in GAMMA, and they were selected with the goal of minimizing the number of aqueous-phase tracers and species being exchanged between the gas and aerosol phases, while maximizing the aaSOA mass captured compared to that as predicted by GAMMA after 12 h of simulated chemistry, assuming no initial aerosol-phase organic mass. The detailed comparison of GAMMA and simpleGAMMA output under a range of typical environmentally relevant conditions can be found in the following section. The aqueous phase species tracked in simpleGAMMA are: IEPOX, glyoxal, 2-methyltetrol, and IEPOX organosulfate. Mass transfer between the gas and aerosol phases only occurs for IEPOX and glyoxal. The effective Henry’s Law constants ($H^*$) and accommodation coefficients used to describe uptake for these species are given in Table 1. Note that these $H^*$ values have been updated based on advances in the literature since McNeill et al. (2012), but the model intercomparisons performed in this study were performed using the McNeill et al. (2012) $H^*$ values, for consistency.

Reversible hydration and oligomerization chemistry of glyoxal in the aqueous phase is captured using $H^*$ (Schwartz, 1986), and therefore those processes are not represented explicitly in either GAMMA or simpleGAMMA. The aqueous processing of IEPOX to form 2-methyltetrols (tetrol) and and IEPOX organosulfate (IEPOXOS) is represented as one reactive process following a simplified version of the mechanism of Eddingsaas et al. (2010):

$$\text{IEPOX(aq)} \rightarrow (1 - \beta)\text{tetrol} + \beta\text{IEPOXOS} \quad (4)$$
Here, we apply a value for the branching ratio, $\beta$, of 0.4, which is an estimate based on the measurements of Eddingsaas and coworkers for the most concentrated bulk solution they studied. The rate constant for the reaction, $k_1$, is a function of proton activity and nucleophile concentrations again following Eddingsaas et al. (2010). We have modified the formula to include the possible protonation of IEPOX (aq) by ammonium as observed by Nguyen et al. (2014)

$$k_1 = k_{H^+} a_{H^+} + k_{SO_4^{2-}} \left[ SO_4^{2-} \right] a_{H^+} + k_{HSO_4^-} \left[ HSO_4^- \right] + k_{NH_4^+} \left[ NH_4^+ \right]$$

(5)

Here, $a_{H^+}$ is the H$^+$ activity, $k_{H^+} = 5 \times 10^{-2}$ s$^{-1}$, $k_{SO_4^{2-}} = 2 \times 10^{-4}$ M$^{-1}$ s$^{-1}$, and $k_{HSO_4^-} = 7.3 \times 10^{-4}$ M$^{-1}$ s$^{-1}$. The ammonium rate constant, $k_{NH_4^+}$, was calculated using GAMMA and the results of the chamber study of Nguyen et al. (2014) to be $1.7 \times 10^{-5}$ M$^{-1}$ s$^{-1}$.

The architecture of the simpleGAMMA program is similar to that of GAMMA (McNeill et al., 2012). simpleGAMMA and GAMMA were originally written in MATLAB (MathWorks, Inc.), utilizing the initial value ODE solver ode15s.m, but simpleGAMMA is also available in Python and Fortran. Required input parameters for simpleGAMMA are: gas phase concentration fields for IEPOX and glyoxal, aerosol pH, aerosol size distribution or volume-weighted average aerosol diameter, aerosol liquid water content, and aerosol sulfate and bisulfate concentrations. The test simulations in this study were for the same conditions as the high-NO$_x$ and low-NO$_x$ scenarios in McNeill et al. (2012). As in that study, the seed aerosols were assumed here to be composed of ammonium sulfate, following the size distribution of Whitby (1978), with aerosol loadings of 4.0 µgm$^{-3}$ (rural conditions, following Tanner et al., 2009) or 20 µgm$^{-3}$ (urban conditions, following Jimenez et al., 2003). Initial aerosol composition was determined by E-AIM outputs for the defined initial pH and RH values.
3 Results

3.1 Low-NO<sub>x</sub> (rural) conditions

Similar to what was observed in McNeill et al. (2012), under simulated rural (low-NO<sub>x</sub>) environments, both GAMMA and simpleGAMMA predict that aaSOA is dominated by IEPOX and its aerosol-phase reaction products. The evolution of aaSOA mass as predicted over 12 h of dawn-to-dusk simulation under low-NO<sub>x</sub> conditions using GAMMA and simpleGAMMA is shown in Fig. 1 (for aerosol pH = 1 and RH 65%). The pie charts compare the aaSOA composition predicted by both models at 3, 6, 9, and 12 h of simulation. The accumulation of aaSOA under low-NO<sub>x</sub> conditions is highly dependent on the formation of gas-phase IEPOX. Although identical gas-phase mechanisms and initial conditions were used in this model intercomparison, the gas-phase chemistry, especially gas-phase OH, is perturbed by the differences in the aqueous-phase mechanisms and gas-aerosol mass transfer between the two models. IEPOX formation is relatively slow, resulting in differences in predicted aaSOA composition between the two models in early simulation hours as IEPOX levels build in the gas phase. After six hours of simulation time the high molecular masses of IEPOX and its substitution products overtake and dominate the aaSOA composition.

Figure 2 shows the total aaSOA mass predicted by GAMMA and simpleGAMMA for 12 h of simulation under low-NO<sub>x</sub> conditions, with varying aerosol pH and RH. Like GAMMA, simpleGAMMA predicts maximum aaSOA formation under low-NO<sub>x</sub> conditions when aerosol pH is low and RH is low (but not so low as to cause aerosol efflorescence). This is because in-particle processing of IEPOX is initiated by protonation, so conditions which maximize the in-particle proton concentration yield the highest IEPOX processing. Close agreement (to within 15%) exists between aaSOA mass predicted by GAMMA and by simpleGAMMA for RH > 50%, and for all RH values for aerosol pH > 2.0. For RH ≤ 45%, the highly efficient in-particle IEPOX chemistry at low pH (pH < 2.0) leads to larger discrepancies (up to 40%) between the two models.
3.2 High-NO\textsubscript{x} (urban) conditions

aaSOA mass as predicted by GAMMA is dominated by “dark” uptake of glyoxal under high-NO\textsubscript{x} conditions (McNeill et al., 2012). Gas-phase IEPOX formation is expected to be minor in this regime (Paulot et al., 2009). A comparison of evolved aaSOA mass and composition under high-NO\textsubscript{x} conditions as predicted by GAMMA and simpleGAMMA can be seen in Fig. 3. Figure 4 shows total aaSOA mass predicted by the two models after 12 h of simulation under high-NO\textsubscript{x} conditions, with varying aerosol pH and RH. Close agreement in predicted total aaSOA mass exists between simpleGAMMA and GAMMA for all relative humidity and pH values tested. Like GAMMA, under high NO\textsubscript{x} conditions simpleGAMMA predicts increasing aaSOA formation with increasing RH (and therefore increasing aerosol liquid water content), and no pH dependence, consistent with glyoxal dark uptake being the dominant aaSOA formation mechanism. GAMMA predicts some contribution to aaSOA mass by photochemical production of succinic acid (vis. Fig. 3); since glyoxal is the dominant precursor for succinic acid formation, this leads to only a small difference in overall predicted aaSOA mass between GAMMA and simpleGAMMA, which does not include in-particle photochemistry.

3.3 Computational performance

The goal of simpleGAMMA is to faithfully represent aaSOA formation with a low number of tracers, in order to simplify the implementation of aqueous aerosol SOA formation in 3-D models (by coupling the gas-phase schemes of those models with simpleGAMMA). However, simpleGAMMA is also computationally faster than GAMMA when run as a box model, as described in this study, due to the reduced number of tracers and reactions in the aqueous phase (recall that the gas phase mechanisms of GAMMA and simpleGAMMA were identical for purposes of this study). In ten simulations with starting aerosol pH 1 and 65\% ambient RH, computational run-time for simpleGAMMA under low-NO\textsubscript{x} conditions spanned between 10–12 s for 12 h of simulation, compared to 33–42 s for GAMMA. These runs were performed on an Intel Core i7-3520M CPU.
Discussion and outlook

The agreement between GAMMA and simpleGAMMA indicate that this reduced framework can be useful to represent aaSOA mass formation over a variety of relevant ambient conditions. Coupling of simpleGAMMA with regional and global scale 3-D atmospheric chemistry models is currently underway (Jathar et al., 2014).

While we have demonstrated good agreement between simpleGAMMA and GAMMA, the limitations of GAMMA also apply to simpleGAMMA; for example, neither model includes a treatment of oxidative aging of aaSOA at this time due to a lack of kinetic and mechanistic data. The only sources of aqueous-phase OH in GAMMA are HOOH photolysis or Henry’s Law transfer of OH from the gas phase. Therefore we, like others (Ervens et al., 2014; Waxman et al., 2013), observed OH-limited chemistry in the aqueous aerosol phase using GAMMA, and this informed the simpleGAMMA formulation. For this reason, simpleGAMMA is not recommended for the treatment of aqueous SOA formation in cloudwater, which is not OH limited and is dominated by aqueous phase photochemistry. The role of UV light in aaSOA formation by glyoxal is unresolved (Galloway et al., 2009, 2011; Kampf et al., 2013; Volkamer et al., 2009). A recent data analysis study using GAMMA (Sumner et al., 2014) suggested a possible role for photo-enhanced chemistry in aaSOA formation by glyoxal involving organic photosensitizers such as fulvic acid (Monge et al., 2012). This chemistry can be represented in simpleGAMMA by including irreversible glyoxal uptake with \( \gamma \sim 10^{-3} \) during sunlit hours, consistent with Fu et al. (2008), who based their representation on the experiments of Liggio et al. (2005), and with Waxman et al. (2013).
We previously predicted using GAMMA that glyoxal is the main contributor to aqueous aerosol-phase “brown carbon” formation by carbonyl-containing VOC precursors (Woo et al., 2013). Following that work, it is straightforward to track the formation of light-absorbing glyoxal derivatives in simpleGAMMA, with concentration-dependent aerosol light absorption calculated in post-processing. However, we note that fast photobleaching of aerosol brown carbon formed via this pathway has been demonstrated, limiting its potential impact on atmospheric chemistry and climate (Lee et al., 2014; Sareen et al., 2013; Woo et al., 2013).

Code availability

For more information and to access the simpleGAMMA program, please visit mcneill-lab.org/gamma or contact V. Faye McNeill, vfm2103@columbia.edu. simpleGAMMA was originally written in MATLAB (MathWorks, Inc.) and is also available in Python and Fortran.

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References


Table 1. Gas-aerosol mass transfer parameters in simpleGAMMA.

<table>
<thead>
<tr>
<th>Species</th>
<th>Effective Henry's Law constant, $H^*$ (Matm$^{-1}$)</th>
<th>Accomodation Coefficient, $\alpha$</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>IEPOX</td>
<td>$3 \times 10^7$</td>
<td>0.02</td>
<td>McNeill et al. (2012), Nguyen et al. (2014)</td>
</tr>
<tr>
<td>GLYX</td>
<td>$2.7 \times 10^7$</td>
<td>0.023</td>
<td>Herrmann et al., (2005), Sumner et al. (2014)</td>
</tr>
</tbody>
</table>
Figure 1. Comparison of simpleGAMMA and GAMMA under low-NO$_x$ conditions, pH1, RH 65%.
Figure 2. Comparison of predicted aaSOA after 12 h of simulated time as a function of RH at pH 1 (left) and pH at 45 and 65% RH (right, dotted and solid lines respectively), low-NO\textsubscript{x} conditions.
Figure 3. Comparison of high-NO$_x$ simpleGAMMA and high-NO$_x$ GAMMA, pH1, RH65.
Figure 4. Comparison of predicted aaSOA after 12 h of simulated time with respect to RH at pH 1 (left) and with respect to pH at 65 % RH (right), high-NOx conditions.