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Evaluation of the atmospheric significance of multiphase reactions in atmospheric secondary organic aerosol formation

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Abstract

In a simple conceptual cloud-aerosol model the mass of secondary organic aerosol (SOA) that may be formed in multiphase reaction in an idealized scenario involving two cloud cycles separated with a cloud-free period is evaluated. The conditions are set to those typical of continental clouds, and each parameter used in the model calculations is selected as a mean of available observational data of individual species for which the multiphase SOA formation route has been established. In the idealized setting gas and aqueous-phase reactions are both considered, but only the latter is expected to yield products of sufficiently low volatility to be retained by aerosol particles after the cloud dissipates. The key variable of the model is the Henry-constant which primarily determines how important multiphase reactions are relative to gas-phase photooxidation processes. The precursor considered in the model is assumed to already have some affinity to water, i.e. it is a compound having oxygen-containing functional group(s). As

a principal model output an aerosol yield parameter is calculated for the multiphase SOA formation route as a function of the Henry-constant, and has been found to be significant already above H~10³ M atm⁻¹. Among the potential precursors that may be eligible for this mechanism based on their Henry constants, there are a suite of oxygenated compounds such as primary oxidation products of biogenic and anthropogenic hydrocarbons, including, for example, pinonaldehyde. Finally, the analogy of multiphase SOA formation to in-cloud sulfate production is exploited.

1. Introduction

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Organic aerosol particles, in spite of their increasingly important role in key atmospheric processes such as radiative transfer or cloud formation, still constitute a puzzle to the atmospheric science community. Since their chemistry remains largely unresolved on the level of individual organic species, our understanding on their sources is still incomplete. To assess the role of human activities on these important atmospheric con-

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stituents with the help of more robust atmospheric models, much better insights into organic aerosol sources are needed. Source apportionment of size-resolved aerosol has been perfected using the concept of atmospheric organic tracers mainly in urban or metropolitan areas, especially in Los Angeles (Schauer et al., 1996). However, outside

- the immediate source regions, with increasing contribution from biogenic secondary aerosol sources for which the tracer approach does not work, source apportionment becomes vague and organic aerosol sources remain largely unknown. Therefore, over most of the troposphere, including rural and remote areas in the boundary layer and in the free troposphere where the contribution of organic aerosol to aerosol mass con-
- centrations has been shown to be significant (Novakov et al., 1997) we know very little about the origin of this constituent. On the other hand, it is these regions and not source areas which determine a large part of the global effects of atmospheric aerosol, therefore better knowledge on sources of organic aerosol would be critically needed.

The first step towards a better understanding of sources which would also allow more thorough validation of global atmospheric models would be a more reliable assessment of the contributions of primary and secondary sources to organic aerosol mass concentrations. In atmospheric science, primary aerosol particles are those which are released directly from mostly surface sources, such as soot particles from combustion or sea salt particles by bubble bursting from the oceans. In contrast, secondary parti-

- cles form in the atmosphere by gas-to-particle transformations following (or preceding) photochemical reactions of directly emitted gaseous or volatile precursors. Among the major inorganic constituents of tropospheric fine aerosol, such as sulfate, nitrate or ammonium, secondary processes are of utmost importance. In contrast, on the global scale organic aerosol mass concentrations have been thought to be dominated by pri-
- ²⁵ mary emissions, mostly from fossil fuel combustion and biomass burning (Penner et al., 2001). Although it is not straightforward to translate annual global emission inventories to relative share of primary versus secondary organic aerosol in the volume of the troposphere, a recent global model based on our current understanding of emissions and secondary organic aerosol (SOA) formation predicted that over the entire

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troposphere primary organic aerosol predominates (Chung and Seinfeld, 2002). On the other hand, regional scale SOA estimates predicted more significant relative contributions of SOA for rural and background regions (Castro et al., 1999; Brown et al., 2002). The most frequently used method to estimate the extent of SOA formation is

- ⁵ based on the evolution of OC/EC ratio. In this concept SOA formation is manifested in the increase in OC/EC ratio of fine aerosol from source areas to rural and background regions (Turpin and Huntzicker, 1991). The method assumes no bias for the determination of elemental carbon (EC) which is considered as a conservative tracer for primary combustion-generated OC emissions. Although these statements are at
- ¹⁰ least questionable, uncertainties are generally understood to arise in the estimation of a characteristic primary OC/EC ratio (Strader et al., 1999; Lim and Turpin, 2002). Using this approach, even urban areas showed relative high SOA contributions in summer (mean 46% in Atlanta, but at times as much as 88% of the 1-h average OC concentrations) (Lim and Turpin, 2002). On an annual basis, at a receptor site in the South
- ¹⁵ Californian Air Basin under strong anthropogenic influence, between 27% and 38% of the organic carbon was assigned to be of secondary origin (Gray, 1986). Furthermore, recent studies in more pristine regions such as national parks indicate an even higher share of SOA (between 42 and 98% for the period of July–October in Texas) (Brown et al., 2002). In Portugal, SOA contribution to fine OC was estimated to be 45% and 68–78% for winter and summer, respectively (Castro et al., 1999).

The general understanding of atmospheric SOA formation is that it mainly occurs by the photochemical reactions of volatile organic compounds (VOCs) and the subsequent condensation of low volatility reaction products onto pre-existing aerosol particles. Among the VOCs involved in SOA formation biogenic unsaturated compounds predominate (Griffin et al., 1999), the primary mechanism of gas-to-particle transfor-

mation is absorptive partitioning into the organic phase (Odum et al., 1996). Very recently there has been mounting evidence that other processes such as heterogeneous and multiphase reactions in the troposphere could induce significant SOA formation in addition to the abovementioned mechanism. The distinction between het-

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erogeneous and multiphase reactions which are constrained to the surface of solid particles and take place in the bulk of hydrometeors, respectively, follows the recommendations by Ravishankara (1997). Heterogeneous reactions are expected to result in a drastic reduction in the equilibrium vapor pressure of the SOA products thereby in-

- ⁵ creasing aerosol yield (Jang and Kamens, 2001). Acid-catalyzed carbonyl chemistry on aerosol particles, which are essentially interactions between the organic and inorganic components of tropospheric aerosol, includes hydration, hemiacetal and acetal formation, aldol condensation, and polymerization in the aerosol phase (Jang et al., 2002). Recent smog-chamber experiments in combination with laser desorption/ionisation
- ¹⁰ mass spectrometry measurements have shown that semi-volatile photooxidation products of 1,3,5-trimethylbenzene tend to polymerize in the aerosol phase in reactions catalyzed only by the semi-volatile acidic reaction products (Kalberer et al., 2004). Another recent smog-chamber study on the effect of acidic seeds on α -pinene ozonolysis has inferred that acidity promotes SOA formation and increases aerosol yield by up to
- 40% (linuma et al., 2004). The first laboratory evidence for the irreversible formation of polymeric species in heterogeneous reactions on sulfuric acid has been presented recently for isoprene (Limbeck et al., 2003). The conclusions drawn from these bulk experiments were that isoprene, whose SOA formation has been thought previously to be negligible (Pandis et al., 1991), may contribute to SOA formation through this
 mechanism. On the molecular level, the possibility of isoprene photooxidation in acid catalyzed reactions has been supported by the identification of two diastereoisomers of a polyol with a C₅ isoprene skeleton, 2-methylthreitol and 2-methylerythritol in forest aerosol (Claeys et al., 2004a).

Although it is well established that in the troposphere a major part of sulfate is formed ²⁵ in reactions in cloud droplets multiphase organic chemistry is still in its infancy. This might be surprising because the possibility of VOC oxidation in cloud and fog droplets was first raised more than 20 years ago (Graedel and Goldberg, 1983). Aqueousphase OH reactions in cloud and fog droplets were proposed as pathways for the oxidation of aldehydes to their corresponding carboxylic acids (Chameides, 1984; Graedel

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et al., 1986; Cho and Carmichael, 1986; Seinfeld and Pandis, 1989). The first laboratory evidence for multiphase polymerization reactions of lignin pyrolysis products in the troposphere has been presented very recently (Gelencsér et al., 2003; Hoffer et al., 2004). The in-cloud formation of oxalic acid from acetylene and ethene in the marine atmosphere has been quantitatively evaluated in a simple model by Warneck (2003). The calculated production rate has been found to largely account for available observations of oxalic acid mass concentrations in the Central Pacific (Kawamura and Sakaguchi, 1999). The suggested oxidation scheme has received additional support from the detection of intermediates glyoxal and glyoxylic acid in marine precipitation

- (Sempere and Kawamura, 1996). The proposed mechanism which includes gas-phase photooxidations, gas-to-particle transfer of intermediates driven by further photooxidation reactions in the aqueous phase, clearly represents a new route for SOA formation previously not considered in atmospheric models. This is because partitioning of semivolatile intermediates into the aerosol phase would be insignificant (therefore negligible
- ¹⁵ SOA mass formation would be inferred by the traditional SOA mechanism) without the aqueous-phase reactions which consume the hydrated species thereby maintaining a flux from the gas phase to the droplet phase. Furthermore, upon the dissipation of the cloud the end-product oxalic acid is not expected to return to the gas phase because of its very low volatility (Clegg et al., 1996). The in-cloud mass production of C_2 - C_6 di-
- ²⁰ carboxylic acids and pyruvic acid has also been calculated in a cloud parcel model for clean and polluted continental conditions (Ervens et al., 2004a). The volatile precursors considered were toluene, ethylene, cyclohexene, and isoprene, and the model was run for several cloud cycles. The organic mass additions for the clean and polluted scenario were found to be ~150 ng m⁻³ and ~400 ng m⁻³, respectively. It should be noted,
 ²⁵ however, that in this case the total liquid water content was not allowed to fall below 1 mg kg⁻¹ after any cloud cycle (i.e. the droplets were not allowed to return to ambient aerosol state). This inevitably means that a substantial fraction of semi-volatile organic species were retained between cloud cycles which would have otherwise evaporated, therefore the model may not be the best quantitative approach for the abovementioned

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new SOA formation mechanism.

In this paper we adopt a simple conceptual model for cloud processing to evaluate the potential significance of the new SOA formation mechanism in the global atmosphere and to constrain the range of precursors that might be eligible for this mecha-5 nism. This effort pursues the tempting analogy of sulfate production in clouds in the

realm of organic chemistry, i.e. whether multiphase processes could induce significant SOA mass production in excess of the conventional mechanism, perhaps even from organic precursors previously not considered to be significant in SOA formation.

2. Conceptual model for multiphase SOA formation

- The conceptual model uses a fixed and idealized atmospheric scenario to help follow the course of multiphase SOA formation and evaluate its atmospheric significance. Being a conceptual model, it does not rely on the specific physical-chemical properties of any given organic photooxidation system, including reaction rates, equilibrium constants, and equilibrium vapor pressure of precursors and products. However, the selection of these key parameters is constrained by observed properties of several re-
- action systems of atmospheric relevance. This simple approach helps keep track of the process itself without being lost in the extreme complexity of atmospheric multiphase systems. Advanced multiphase chemistry models that have recently become available have just started to evaluate the in-cloud production of low molecular weight dicar-
- ²⁰ boxylic acids as abundant low volatility aerosol constituents (Warneck, 2003; Ervens et al., 2004a). In spite of the existence of such models, our approach may help assess the significance of this new formation pathway relative to the conventional SOA formation mechanism that has long been included in atmospheric models of regional and global scale. Furthermore, it may allow us to explore the analogy with the well-established in-
- cloud sulfate production in order to shed light on the potential global importance of the process. And last but not least our model may promote further studies on this field by constraining the range of compounds that might be eligible for this new SOA formation

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route.

Our conceptual model starts with a volatile organic precursor that already has some affinity for water. Such compounds may either be directly emitted by natural and anthropogenic sources or formed as first generation products by the photooxidation of VOCs.

- ⁵ The list of potential precursors can be found in the excellent review by Saxena and Hildemann (1996). Although of the two source types the latter is clearly predominant in most environments, it is outside the scope of this paper to include gas-phase formation of the precursor. The model starts with an air parcel containing the precursor at a fixed overall mixing ratio of 10 ppt in which a cloud with a total liquid water content (LWC)
- ¹⁰ of 0.3 g m^{-3} consisting of $10 \,\mu\text{m}$ droplets is allowed to form at a temperature of 288K. The cloud is assumed to persist for 1 hour than dissipates and the compounds formed are either vented to the gas-phase or remain in the aerosol phase depending on their equilibrium vapor pressure through absorptive partitioning. This cloudless period lasts for 8 hours than a cloud with the same parameters forms again for 1 additional hour. ¹⁵ In the model, aerosol formation is evaluated for the overall process consisting of three
- elementary stages.

As regards mass balance, the system is considered to be closed, i.e. the precursor is not replenished by photochemical reactions in the gas-phase. While this constraint may not be suitable for precursors formed from long-lived gaseous species such as

- acetylene or from those whose emission sources are widely distributed, it may be reasonable for photooxidation products of highly reactive volatile organic compounds. In any case, this approach places a lower limit for the yield of cloud chemical processes. The upper limit would be to assume an open system, i.e. steady-state gas-phase mixing ratio of the precursor throughout the duration of the entire cloud cycle. In the cloud
- the equilibrium of the precursor between the gas and droplet phase is established according to Henry's law. Henry's constant turns out to be the key variable in the process which primarily determines the importance of multiphase SOA formation with respect to gas-phase photooxidation. At t=0 photooxidation is allowed to start both in the gas and droplet phase assuming steady-state oxidant concentrations in both phases. For

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OH radical concentrations steady-state values of 2.5×10^{6} cm⁻³ and 5×10^{-13} M are set for the gas and aqueous phase, respectively (Warneck, 2003; Lelieveld and Crutzen, 1991).

- Second-order gas-phase reaction rate for all species is assumed to be 1×10⁻¹¹ cm³ molecule⁻¹ s⁻¹, derived as a mean from experimental rate constants of various oxygenated compounds (Calogirou et al., 1999; Glasius et al., 2000; Hallquist et al., 1997; Atkinson, 1989). Similarly, for aqueous-phase reaction rates a single value of 5×10⁸ M⁻¹ s⁻¹ is assumed for all species (Dutot et al., 2003). Mass transfer from the gas phase is accounted for using "global reaction rate expression" by Ravishankara (1997). While the product of the gas-phase photooxidation reaction is not considered
- any further it is assumed that the equilibrium vapor pressure and Henry constant of the first generation product of the aqueous-phase reaction has decreased by a factor of 100 and increased by a factor of 1000, respectively, relative to those of the precursor. These assumptions are based on the properties of intermediates in relevant
- in-cloud reactions leading to the formation of dicarboxylic acids as reported by Ervens et al. (2004a) and Warneck (2003). This first generation reaction product as intermediate in the process is assumed to react further in the aqueous phase yielding a second generation product of low volatility. Gas-phase reaction of this intermediate is also considered as a sink but its product is deemed too volatile to contribute to aerosol mass.
- ²⁰ For simplicity, the rates of the reactions are assumed to be the same as those of the precursor. The second generation product of the aqueous-phase reaction is assumed to have an equilibrium vapor pressure by a factor of 10⁶ lower than that of the initial precursor based on relevant references (Ervens et al., 2004a; Warneck, 2003).

The duration of the cloud event is set to 1 h, after which the cloud dissipates and equilibrium between the gas and aerosol phase is established for both the precursor and products. This process releases the precursor and a large fraction of its first-generation product back to the gas-phase. The equilibrium between the gas and aerosol phase is evaluated by the absorptive partitioning theory by Pankow (1994). Here the key parameter is the equilibrium vapor pressure which is assumed to be 5 Pa at 288K for the

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precursor. The activity coefficient (ξ_i) and the mass concentration of total suspended particulate matter are assumed to be unity and 1 μ g m⁻³, respectively.

Then follows a cloud-free period of 8 h during which only gas-phase reactions proceed. Then another cloud cycle occurs which lasts for another 1 h, in-cloud equilibrium is reestablished and aqueous-phase reactions resume. The cloud parameters are the same as in the initial case. The process ends with the dissipation of the cloud and the re-establishment of equilibrium between the gas and aerosol based on the principles above.

As the principal model output the overall SOA formation per unit volume of air is calculated as the sum of the carbon mass concentrations of primary and secondary products that remain in the aerosol phase at the end. The mass concentrations of all species are expressed on a carbon basis, therefore no mass increase is assumed for the first and second generation products. Furthermore, it is important to stress that ambient aerosol state is considered here, unlike the advanced model by Ervens

- et al. (2004a) which does not allow liquid water content to fall below 1 mg m^{-3} after any cloud cycle. Another key output of the model is the aerosol yield (*Y*) of the entire process, which is defined as the ratio of the mass of aerosol formed to the mass of the precursor reacted, exactly the same as for interpreting smog-chamber experiments (Hoffmann et al., 1997). In addition, the model allows the calculation of the relative
- ²⁰ contributions of gas- and aqueous phase reactions over the entire process. Note that the total depletion of precursor is less than the combined yields of gas-phase and multiphase reactions since in both cases consecutive reactions are considered. The model outputs are expressed as a function of Henry's constant which is the only variable in this conceptual model. This model helps evaluate the role of Henry's constant in multi-
- ²⁵ phase SOA formation, assess the magnitude of the process and compare it to in-cloud sulfate formation. Furthermore, with the range of Henry' constants constrained, potential precursors can be selected from the compilation by Saxena and Hildemann (1996) which, when combined with the retrospective approach to gas-phase formation mechanisms by Pun et al. (2000), may allow the identification of primary VOCs that might

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3. Results and discussion

The total mass of aerosol carbon formed in unit volume of air in the two 1-h cloud events separated with an 8-h cloudless period is shown as a function of the Henry-constant
of the precursor, with the set parameters detailed in the model description. In the same figure, two limiting cases are also depicted: the low bound (in terms of aerosol mass production) when the rate of the gas-phase reaction is increased and that of the aqueous-phase reaction is decreased by a factor of 2 each; and the high bound vice versa. These constrain a range for the given scenario which could give reasonable
estimates provided that the other assumptions (e.g. reduction of Henry-constants and equilibrium vapor pressures from precursors to first- and second-generation products) are valid.

All curves in Fig. 1 show a steep increase starting already above $H \sim 10^3$ M atm⁻¹, and a plateau from less than $H \sim 10^6 \,\text{M}\,\text{atm}^{-1}$ which indicates that during the cloud events virtually all species reside in the droplets and being oxidized there. It is also obvious 15 that the production rate is very sensitive to Henry constant in a relatively confined range between $H \sim 10^3$ and 5×10^4 M atm⁻¹, but its magnitude is also strongly dependent on the rates of the gas-phase and agueous-phase reactions. In other terms, it is primarily the Henry-constant which determines whether multiphase aerosol production is possible or not. If yes, the reaction rates will determine its magnitude and significance 20 relative to gas-phase photooxidation processes. From the model it is also possible to calculate aerosol yield (Y) as a function of the logarhytm of the Henry-constant. It is shown in Fig. 2, together with the low and high bound estimates specified above. The aerosol yield of the precursor shows a wide span from being negligible to very significant in almost the same logH range as absolute aerosol mass concentrations in 25 Fig. 1. Remember that the aerosol yield specifies how much aerosol is formed (on a

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carbon mass basis) relative to the fraction of precursors reacted - it is therefore a di-

rect measure of the importance of aerosol formation versus any other chemical sinks. Additionally, it should be noted that the value of Y refers to the actual atmospheric scenario modeled; it may be substantially different if other scenarios were set. With this in mind, aerosol yields can be used to link cloud-processing to the much better known

 gas-phase photooxidation mechanisms, since the precursors considered here are firstor second order reaction products of gas phase processes. Knowing the stoichiometry of gas-phase photooxidation of a parent VOC to its oxygenated product – which is actually the precursor of cloud-processing –, and the aerosol yield of the precursor, one should be able to assess the importance of multiphase SOA formation from the given
 parent compound, at least for the scenario considered.

The minimum value of Henry constant that is needed to furnish non-negligible secondary organic aerosol formation by cloud processes in the given scenario is about H~10³ M atm⁻¹. This constrains the range of potential precursors that might be eligible for this mechanism. One should remember that the parameters of this conceptual ¹⁵ model are non-specific, therefore for any individual species it can only be regarded as very rough indication for the likelihood that such multiphase processes might or might not be important. Verification of potential pathways should involve experimental evidence that consecutive aqueous-phase oxidation reactions do occur under conditions relevant in tropospheric cloud water, identification and determination of the physico-²⁰ chemical properties of the products formed and the determination of the rates of the

reactions. Preferably, potential pathways should be evaluated in more advanced multiphase chemistry models such CAPRAM (Hermann et al., 2000), or those developed by Ervens et al. (2004a, b) and Warneck (2003).

Among potential candidates, glyoxal and glycolaldehyde, formed by the gas-phase ²⁵ photooxidation of acetylene and ethene, respectively, both turn out be important precursors for multiphase secondary aerosol formation having effective Henry-constants 3.4×10^5 and 4.14×10^4 M atm⁻¹, respectively (Betterton and Hoffmann, 1988). Alternatively, glyoxal may also be formed by the photooxidation of toluene (Ervens et al., 2004a). In fact, the significance of these in-cloud chemical processes has already been

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firmly established in a recent model study on the formation of oxalic acid in the marine atmosphere (Warneck, 2003). In general, multifunctional organic compounds may appear to be potential precursors for multiphase aerosol production. For example, hydroxyacetone which is produced by isoprene photooxidation (Yu et al., 1995) may fall in this category based on its Henry constant (2×10⁴ M atm⁻¹) (Klotz et al., 1999). However, 5 no information on the mechanism of aqueous-phase photooxidation of hydroxyacetone has been available in the literature. Similarly, pinonaldehyde, the primary product of the gas-phase reactions of α -pinene with ozone and OH also seems to be eligible for this mechanism based on its estimated Henry constant of $H \sim 9 \times 10^3 \,\text{M} \,\text{atm}^{-1}$ (Meylan and Howard, 1991). Other potential bifunctional precursors include 3-hydroxypropanoic 10 acid $(H \sim 3.7 \times 10^7 \text{ M atm}^{-1})$ which may be formed from (Z)-3-hexen-1-ol (leaf alcohol) emitted by vegetation (Pun et al., 2000). The photooxidation of unsaturated oxygencontaining compounds (e.g. 3-methyl-3-buten-1-ol) emitted in substantial amounts by plant species in Europe (König et al., 1995) is expected to yield hydroxy carbonyls (Grosjean et al., 1993) which might also be aerosol precursors through the multiphase 15 route.

On the contrary, methacrolein, a primary photooxidation product of isoprene, whose aqueous phase transformation to 2,3-dihydroxymethacrylic acid by hydrogen peroxide has been successfully demonstrated in the laboratory (Claeys et al., 2004b), has far too low Henry constant (6.5 M atm⁻¹) (Iraci et al., 1998) to support multiphase aerosol formation against gas-phase photooxidation processes. However, a gas-phase photooxidation product of methacrolein, methacrylic acid, which has also been shown to yield 2,3-dihydroxymethacrylic acid upon reaction with hydrogen peroxide in the aqueous phase (Claeys et al., 2004b), may turn out to be a potential precursor for multiphase

²⁵ SOA formation based on its moderate Henry constant of 2×10³ M atm⁻¹ (Meylan and Howard, 1991).

Finally, one important issue remains, namely how multiphase SOA formation relates to sulfate formation in in-cloud processes. Gas-phase SO₂, on account of its acid-base properties and chemical interactions with other soluble species such as formaldehyde

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(Klippel and Warneck, 1978), is a highly soluble gas which has an effective Henry constant well within the range that has been shown to be important in multiphase oxidation processes. Secondly, gas-phase oxidation rate of SO₂ by OH is lower than those of most organic compounds, while typical aqueous phase oxidation of SO_2 – though the mechanism of oxidation is rather complex and pH-dependent – are about an order of 5 magnitude faster than that of most organic species at pH typical of cloud water (Warneck, 1988; Dutot et al., 2003). But most importantly, SO₂ oxidation produces the essentially non-volatile sulphuric acid in a single step while SOA formation requires at least two consecutive reactions to yield low-volatility multifunctional products that remain in the aerosol phase after the dissipation of the cloud. All these factors make 10 multiphase SOA formation less effective per unit mass concentration of the precursor than in-cloud sulfate production. Nevertheless, considering the vast amounts of VOCs that are emitted by vegetation and oxidized in the troposphere this process may still represent an important route for organic aerosol mass production. Last but not least this mechanism may bring into focus new volatile species that are not primarily impor-

this mechanism may bring into focus new volatile species that are not primarily important in terms of mass emission relative to natural hydrocarbons but have high aerosol yields through the multiphase route.

4. Conclusions

This study establishes multiphase photooxidation as a potentially significant source of secondary organic aerosol (SOA) with the help of a simple conceptual cloud-aerosol chemistry model using a fixed and idealized atmospheric scenario. The model is based on the assumption that aqueous reactions of organic species with OH radicals yield an essentially non-volatile product in two consecutive reaction steps. In the model overall aerosol production is evaluated as a function of the Henry constant of the precursor.

The results show that there is a limit in aqueous solubility of organic photooxidation products or of directly emitted oxygenated compounds above which their contribution to SOA formation via in-cloud oxidation processes needs to be critically evaluated. This

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limit corresponds to Henry constant of about 10³ M atm⁻¹ which qualifies many bifunctional compounds and carbonyls of larger molecular weight such as pinonaldehyde as potential precursors for the process. For compounds that has qualified the aerosol yield of an idealized scenario involving two cloud cycles separated with a cloudless period varies from about 1% to over 70%, strongly depending on the relative rates of gas-phase and aqueous phase photooxidation reactions. While ample data have been available for the former, unfortunately very little is known about the mechanisms and rates of aqueous-phase organic oxidation reactions. Although multiphase SOA formation is clearly a less efficient process than in-cloud sulfate production per unit mass concentration of the precursor, it may significantly contribute to atmospheric SOA production on a global scale. However, much better understanding of cloud organic chem-

duction on a global scale. However, much better understanding of cloud organic chemistry is needed before the significance of this process can be assessed with a tolerable level of certainty.

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Fig. 1. Aerosol mass production of multiphase reactions per unit volume of air as a function of the logarhytm of the Henry constant of the precursor for the idealized scenario considered (1 h in cloud – 8 h cloudless – 1 h in cloud). Initial mixing ratio of the precursor is 10 ppt C. Middle curve: with $k_{gas}=2.5\times10^{-5}$ cm³ molecule⁻¹ s⁻¹, $k_{aqueous}=2.5\times10^{-4}$ M⁻¹ s⁻¹; upper curve: with $k_{gas}=1.25\times10^{-5}$ cm³ molecule⁻¹ s⁻¹, $k_{aqueous}=5\times10^{-4}$ M⁻¹ s⁻¹; lower curve: with $k_{gas}=5\times10^{-5}$ cm³ molecule⁻¹ s⁻¹, $k_{aqueous}=1.25\times10^{-4}$ M⁻¹ s⁻¹.

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Fig. 2. Aerosol yield of multiphase reactions as a function of the logarhytm of the Henry constant of the precursor for the idealized scenario considered (1 h in cloud – 8 h cloudless – 1 h in cloud). Middle curve: with $k_{gas} = 2.5 \times 10^{-5} \text{ cm}^3$ molecule⁻¹ s⁻¹, $k_{aqueous} = 2.5 \times 10^{-4} \text{ M}^{-1} \text{ s}^{-1}$; upper curve: with $k_{gas} = 1.25 \times 10^{-5} \text{ cm}^3$ molecule⁻¹ s⁻¹, $k_{aqueous} = 5 \times 10^{-4} \text{ M}^{-1} \text{ s}^{-1}$; lower curve: with $k_{gas} = 5 \times 10^{-5} \text{ cm}^3$ molecule⁻¹ s⁻¹, $k_{aqueous} = 1.25 \times 10^{-4} \text{ M}^{-1} \text{ s}^{-1}$; lower curve: with $k_{gas} = 5 \times 10^{-5} \text{ cm}^3$ molecule⁻¹ s⁻¹, $k_{aqueous} = 1.25 \times 10^{-4} \text{ M}^{-1} \text{ s}^{-1}$.