Modeling secondary organic aerosol formation from isoprene oxidation under dry and humid conditions

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Abstract

A new model for the formation of secondary organic aerosol (SOA) from isoprene was developed. This model uses surrogate molecular species (hydroxy-hydroperoxides, tetrols, methylglyceric acid, organic nitrates) to represent SOA formation. The development of this model used available experimental data on yields and molecular composition of SOA from isoprene and methacrolein oxidation. This model reproduces the amount of particles measured in smog chambers under both low-NO_x and high-NO_x conditions. Under low-NO_x conditions, the model reproduces the transitional formation of hydroxy-hydroperoxides particles, which are photolyzed and lead to SOA mass decrease after reaching a maximum. Under high-NO_x conditions, particles are assumed to be formed mostly from the photo-oxidation of a PAN-type molecule derived from methacrolein (MPAN). This model successfully reproduces the complex NO_x-dependence of isoprene oxidation and suggests a possible yield increase under some high-NO_x conditions. Experimental data correspond to dry conditions (RH<10%). However, particles formed from isoprene are expected to be highly hydrophilic, and isoprene oxidation products would likely partition between an aqueous phase and the gas phase at high humidity in the atmosphere. The model was extended to take into account the hydrophilic properties of SOA, which are relevant under atmospheric conditions, and investigate the effect of particulate liquid water on SOA formation. An important increase in SOA mass was estimated for atmospheric conditions due to the hydrophilic properties. Experiments should be conducted to confirm the results of this study, which have implications for SOA modeling.

1 Introduction

Atmospheric fine particles are known to have effects on health, atmospheric visibility, materials and climate. Among those particles, particulate organic matter (POM) represents a large fraction of the particulate mass, typically between 20 and 60% (Kanakidou...
et al., 2005; Yu et al., 2007; Zhang et al., 2007a). Those particles are either primary (directly emitted as particles) or secondary (particles formed by chemical reactions in the atmosphere). In the latter case, secondary organic aerosols (SOA) are formed by the gas-to-particle partitioning of the oxidation products of volatile organic compounds (VOCs). VOCs are emitted by both anthropogenic and biogenic sources. However, SOA formation is poorly understood due to the numerous and complex chemical phenomena involved (successive VOC oxidation steps, NO\textsubscript{x} chemistry, compounds partitioning between several phases, SOA degradation, interactions between compounds in the particle, oligomerization).

The complexity of SOA formation makes chemical modeling of Particulate Matter (PM) difficult. To model SOA formation, most air quality models use simple parameterizations based on yields estimated from smog chamber measurements conducted under specific conditions, which can be different from atmospheric conditions (for example, oxidation under dry conditions). Those parameterizations rarely represent the effects of the NO\textsubscript{x} level or oligomerization, whereas those effects have been shown to greatly affect the level of SOA predicted (Ng et al., 2007a,b; Pun and Seigneur, 2007). Therefore, it is necessary to develop a parameterization that gathers all those phenomena.

Isoprene (2-methylbutadiene) is the biogenic VOC, which has the largest emission rate of all the non-methane VOCs, estimated at 600 Tg yr\textsuperscript{-1} (Guenther et al., 2006). Until recently, isoprene was believed not to be a major SOA precursor, despite its large emission flux, due in part to the high volatility of its first generation oxidation products (such as methacrolein and methyl vinyl ketone). Furthermore, some smog chamber studies initially showed that no significant aerosol growth was observed from isoprene photooxidation under high-NO\textsubscript{x} conditions (Pandis et al., 1991). However, during field studies in Amazonia, compounds like tetrals with the same carbon skeleton as isoprene were identified (Claeys et al., 2004; Edney et al., 2005), suggesting that isoprene could in fact contribute to SOA formation. SOA formation was then confirmed by several chamber studies (Kroll et al., 2006; Surratt et al., 2006; Ng et al., 2008).
Therefore, it is important to take isoprene-SOA into account and the effect of humidity on the isoprene-SOA partitioning to properly model SOA formation. Chamber experiments led to the identification of species composing isoprene SOA like tetrols and methylglyceric acid (Surratt et al., 2006; Kleindienst et al., 2009). As those products are rather small organic molecules (number of carbon atoms ≤5) and with numerous functional groups, they are expected to be highly hydrophilic. Then, those compounds would likely partition between an aqueous phase and the gas phase at high humidity in the atmosphere. It has been shown that taking into account the hydrophilic properties of isoprene SOA increases greatly the amount of SOA formed (Pun, 2008), whereas the parameterization of Zhang et al. (2007b) used currently in most air quality models does not take those effects into account.

The objective of this work is to develop a new model for the formation of SOA from isoprene, which takes into account NO$_x$ chemistry, the hydrophilic properties of molecular species and oligomerization. First, a model is developed based on results from experiments conducted under dry conditions. Next, the model is extended to humid conditions by taking into account gas-aqueous phase partitioning.

2 Model development

2.1 Method

To develop the model for SOA formation from isoprene oxidation, the experimental results of Kroll et al. (2006) and Surratt et al. (2006) were used. Those experiments were done in the Caltech experimental chamber under both low-NO$_x$ and high-NO$_x$ conditions. A complete description of the facility is given by Cocker III et al. (2001). The gas-phase chemistry was simulated with RACM2 (Goliff and Stockwell, 2008), which includes a mechanism for isoprene oxidation. This mechanism has been shown to perform well for oxidant formation (Kim et al., 2010). The mechanism was modified to take into account the formation of the surrogate species to represent SOA formation.
as described below. The ROS2 algorithm (Verwer et al., 1999) was used to solve the chemical kinetic equations.

The experiments of Kroll et al. (2006) and Surratt et al. (2006) used blacklights, which emit in the ultraviolet between 300 and 400 nm with a maximum at 354 nm (Kroll et al., 2006). Under those conditions, nitrate radical (NO$_3$) and ozone (O$_3$) photolysis hardly occurs (Carter et al., 2005) and their photolysis rates are almost negligible. The rate given by Kroll et al. (2006) was used for H$_2$O$_2$ photolysis, which leads to hydroxy radical (HO) formation in the chamber. The photolysis rate for NO$_2$ photolysis was chosen so that the maximum ozone concentration matches the results from Kroll et al. (2006). O$_3$, NO$_3$ and HONO photolysis rates were then calculated using the photolysis rates relative to NO$_2$ given by Carter et al. (2005).

As those experiments were done under dry conditions (RH<5%), partitioning between the gas phase and an organic liquid phase was assumed. The partitioning was calculated using the model of Pankow (1994a,b), which defines the absorption equilibrium constant ($K_{p,i}$) according to Raoult’s law as:

\[
K_{om,i} = \frac{F_{i,om}}{A_i M_o} = \frac{760RT}{\text{MW}_{om} 10^6 \zeta_i p_{L,i}^0}
\]  

(1)

where $A_i$ is the gas-phase concentration of compound $i$ (ng m$^{-3}$), $F_{i,om}$ is the concentration of the compound $i$ in the absorbing organic phase (ng m$^{-3}$), $M_o$ is the absorbing organic mass concentration (µg m$^{-3}$), $R$ is the ideal gas constant (8.206 $\times$ 10$^{-5}$ m$^3$ atm mol$^{-1}$ K$^{-1}$), $T$ is the temperature (K), MW$_{om}$ is the mean molecular weight of the absorbing organic phase (g mol$^{-1}$), $\zeta_i$ is the activity coefficient of compound $i$ in the organic phase, and $p_{L,i}^0$ is the saturation vapor pressure of the absorbed compound (torr). The thermodynamic model UNIFAC (UNIversal Functional group Activity Coefficient) was used to calculate activity coefficients (Fredenslund et al., 1975). The missing UNIFAC parameters (for the functional groups nitrate and hydroperoxide) were taken from Comernolle et al. (2009).
Kleindienst et al. (2009) measured the effective enthalpies of vaporization of SOA for both low-NO$_x$ and high-NO$_x$ conditions. Those values were used in this study: a value of 38.4 kJ mol$^{-1}$ for compounds formed under low-NO$_x$ conditions (tetrols, hydroxy-hydroperoxides) and a value of 43.2 kJ mol$^{-1}$ for compounds formed under high-NO$_x$ conditions (methyl-glyceric acid, organic nitrates).

The model development for SOA formation via oxidation of isoprene by NO$_3$ and HO is presented next. Although SOA formation occurs via oxidation of isoprene by O$_3$ (Kamens et al., 1982; Kleindienst et al., 2007; Nguyen et al., 2010), this pathway was not taken into account here because of insufficient quantitative and mechanism information to develop a model at this point.

### 2.2 Oxidation of isoprene by NO$_3$

The results of Ng et al. (2008) were used to model oxidation of isoprene by NO$_3$. In their study, a C$_5$-hydroxy-trinitrate (C$_5$H$_9$N$_3$O$_{10}$) compound (structure of the compound shown in Table 1) was found to be predominant in the “typical experiment” of Ng et al. (2008) (experiment relevant to atmospheric conditions). Therefore, it was used as a surrogate compound for SOA formation for oxidation by NO$_3$.

Isoprene reacts with NO$_3$ to form the surrogate ISON in RACM2 (which groups all compounds formed from the oxidation of isoprene with NO$_3$) by this reaction:

\[
\text{ISO} + \text{NO}_3 \Rightarrow \text{ISON} \tag{R1}
\]

(with a kinetic constant $k=6.61 \times 10^{-13} \text{ molecule}^{-1} \text{ cm}^3 \text{ s}^{-1}$). However, there is no oxidation of this compound by NO$_3$ in RACM2. The following reaction was added to RACM2 to take into account the formation of C$_5$-hydroxy-trinitrate:

\[
\text{ISON} + \text{NO}_3 \Rightarrow \ldots \Rightarrow \alpha_{\text{NO}_3} \text{C}_5\text{H}_9\text{N}_3\text{O}_{10} \tag{R2}
\]

where $\alpha_{\text{NO}_3}$ is the stoichiometric coefficient of the C$_5$-hydroxy-trinitrate. A kinetic constant of $7.0 \times 10^{-14} \text{ molecule}^{-1} \text{ cm}^3 \text{ s}^{-1}$ was taken from Rollins et al. (2009).
The method of Odum et al. (1996) with aerosol yields was used by Ng et al. (2008) with two surrogate compounds to estimate SOA formation parameters. Here, the method was used but with only one surrogate compound because one surrogate compound is sufficient to reproduce the results of Ng et al. (2008) as shown in Fig. 1. A stoichiometric yield $\alpha_{NO_3}$ of 0.074 was estimated with a saturation vapor pressure of $1.12 \times 10^{-6}$ torr.

It should be noted that the formation of C$_5$H$_9$N$_3$O$_{10}$ may be a minor pathway for SOA formation in the atmosphere because there is probably not enough NO$_3$ in the atmosphere to form much C$_5$-hydroxy-trinitrate (C$_5$-dihydroxy-dinitrate is more likely). Reaction (R2) could then overestimate SOA formation from ISON oxidation. However, this reaction could be used to deduce whether or not oxidation of isoprene by NO$_3$ can lead to a significant quantity of SOA. In this study, this reaction was used to make sure that NO$_3$ does not contribute significantly to SOA formation in environmental chambers.

2.3 Oxidation of isoprene by HO: formation of methyl-tetrols and hydroxy-hydroperoxides under low-NO$_x$ conditions

Methyl-tetrols and hydroxy-hydroperoxides are expected to be the two main compounds formed from the photo-oxidation of isoprene under low-NO$_x$ conditions. Whereas tetrols have been explicitly identified in chamber experiments (Kleindienst et al., 2009), the structure of hydroxy-hydroperoxides is still unknown and has not been conclusively identified (Kroll et al., 2006). Nevertheless, it is expected to be a key-component for SOA formation (Kroll et al., 2006; Surratt et al., 2006) and Surratt et al. (2006) confirmed that peroxides constitute a large fraction of SOA under low-NO$_x$ conditions. Under the conditions of Kroll et al. (2006), the RO$_2$ (organic radical) chemistry is dominated by RO$_2$+HO$_2$ reactions. Hydroxy-hydroperoxides are expected to be formed by those reactions. In RACM2, the formation of the radical (noted ISOP) from the oxidation of isoprene by HO, can lead to the formation of first-generation hydroxy-hydroperoxides (noted ISHP) by the following chemical reactions:
ISO + HO ⇒ ISOP \hspace{1cm} (R3)

ISOP + HO₂ ⇒ ISHP \hspace{1cm} (R4)

However, the surrogate compound ISHP is too volatile to form SOA (the group contribution method SIMPOL.1 (Pankow and Asher, 2008) gives a saturation vapor pressure of 6.09×10⁻³ torr). Only one of the isoprene double bonds has been oxidized to form ISHP. If hydroxy-hydroperoxides SOA are formed, it can be assumed that ISHP is a key intermediate in their formation. So, it was assumed that hydroxy-hydroperoxides are formed from the oxidation of the ISHP double bond by HO and a subsequent RO₂ + HO₂ reaction:

\[
\text{ISHP} + \text{HO} \Rightarrow \ldots \Rightarrow \ldots + \alpha \text{BiPER} + \beta \text{BiDER} \hspace{1cm} (R5)
\]

where BiPER is a dihydroxy-dihydroperoxide, which may undergo photolysis, BiDER is another product with structure unknown, which is not photolyzed, \( \alpha \) and \( \beta \) are respectively the stoichiometric coefficients of BiPER and BiDER. Because of lack of data, molar mass and molecular structure of BiDER were assumed to be those of tetrals but the saturation vapor pressure was chosen to be different from the tetral saturation vapor pressure. A rate constant of \( 3.0 \times 10^{-11} \text{ molecule}^{-1} \text{ cm}^{3} \text{ s}^{-1} \) was chosen for Reaction (R5) (kinetics supposed similar to the kinetics of the MACR+HO reaction in RACM2).

After reaching a maximum, the SOA mass was observed to decrease rapidly in the experiments of Kroll et al. (2006) under low-NOₓ conditions. The decrease stopped immediately when the lights were turned off. It indicates that the loss is due to photolytic activity and not to wall losses. To our knowledge, rapid loss of SOA has not been observed in the photo-oxidation of another compound.

The mechanism for SOA loss is not clear. The loss of SOA could be due to either gas-phase reactions (consumption of the partitioning compounds may drive equilibrium away from the particle phase) or particle-phase reactions. For example, SOA loss could
be due to gas-phase reaction with HO or photolysis. Kroll et al. (2006) assumed that the compounds accountable for the rapid SOA loss may be organic hydroperoxides as this loss is only seen under low-NO\textsubscript{x} conditions.

Because of the uncertainties of the SOA loss mechanism, modeling the loss by a global BiPER degradation kinetics (degradation into both particulate and gas phases) was chosen:

BiPER $\Rightarrow$ Unknown Products \hspace{1cm} (R6)

Tetrols are assumed to be formed from the oxidation of diols which can be formed from RO\textsubscript{2}+RO\textsubscript{2} reactions. Under low-NO\textsubscript{x} conditions, RO\textsubscript{2}+RO\textsubscript{2} reactions become predominant when concentrations of isoprene become high but can contribute slightly to SOA formation in Kroll et al. (2006). RACM2 does not take into account the diol formation from isoprene. However, Ruppert and Becker (2000) give a yield for diol formation between 7.1% and 9.3%. A yield of 8.0% was assumed here. Based on data of Ruppert and Becker (2000), a rate constant for the oxidation of diols by HO of $13.0\times10^{-11}$ molecule\textsuperscript{-1} cm\textsuperscript{3} s\textsuperscript{-1} was chosen. Under the conditions of Ruppert and Becker (2000), diols (noted here DIOL) can be expected to be formed from an ISOP+ISOP reaction, as follows:

\begin{align*}
2 \text{ISOP} & \Rightarrow 2 \text{MACR} + \text{HCHO} + \text{HO}_2 + 0.16 \text{DIOL} \hspace{1cm} (R7) \\
\text{The ISOP+ISOP reaction has been taken from P"oschl et al. (2000) and DIOL formation was added to this reaction (the yield was multiplied by two because two radicals from isoprene react in this reaction). ISOP can also be oxidized by radicals MO\textsubscript{2} and ACO\textsubscript{3}. The following reactions have been added to RACM2:} \\
\text{ISOP} + \text{MO}_2 & \Rightarrow \ldots + 0.16 \text{DIOL} \hspace{1cm} (R8) \\
\text{ISOP} + \text{ACO}_3 & \Rightarrow \ldots + 0.16 \text{DIOL} \hspace{1cm} (R9) \\
\text{DIOL} + \text{HO} & \Rightarrow \gamma \text{BiMT} + \text{HO} \hspace{1cm} (R10)
\end{align*}
where BiMT is the surrogate compound for tetrols and \( \gamma \) is the corresponding stoichiometric coefficient of BiMT. The group contribution method SIMPOL.1 gives similar saturation vapor pressures for tetrols and \( C_5 \)-hydroxy-trinitrate. Accordingly, the saturation vapor pressure of tetrols was taken identical to that of \( C_5 \)-hydroxy-trinitrate. A value for \( \gamma \) of 0.16 was used (assuming that the second oxidation has the same yield as the oxidation of the first double bond).

In the atmosphere, concentrations of ISOP are not typically sufficient for Reaction (R8) to be dominant. Inside environmental chambers, Reactions (R8) and (R9) probably do not produce high quantity of DIOL due to the low concentrations of radicals \( MO_2 \) and \( ACO_3 \). However, those radicals are likely to be in higher concentrations in the atmosphere and BiMT could be formed by this pathway.

The stoichiometric coefficients \( \alpha \) and \( \beta \), the reaction rate of degradation of BiPER (R6) and the saturation vapor pressures of BiPER and BiDER were selected to minimize the error between modeled and experimental results by a least square method for both maximal and final mass of SOA. A mean temperature of 25°C was assumed. However, to be able to solve the system, one of the parameters has to be set. According to Kroll et al. (2006), the SOA loss is about 0.006–0.018 min\(^{-1}\). A mean value of 0.012 min\(^{-1}\) was then taken for Reaction (R6). Optimization gives the values of the parameters presented in Table 2. Results of the comparison between modeled and measured SOA formation are shown in Table 3 and Fig. 2. Figure 3a shows the results of a simulation for 63.6 ppb of isoprene.

The model reproduces well the results from Kroll et al. (2006) but there are still uncertainties, in particular in the mechanism for SOA degradation and in the values for the yield and the saturation vapor pressure of methyl-tetrols. Under the conditions of Kroll et al. (2006), tetrols only represent a small part of SOA (simulation for the oxidation of 63.6 ppb of isoprene gives 8% of tetrols). But tetrols would represent a larger fraction of SOA with a larger quantity of oxidized isoprene because the ISOP+ISOP reaction would then be competitive with the ISOP+HO\(_2\) reaction. This is confirmed by a simulation done with 500 ppb where tetrols represent about 40% of SOA formed (Fig. 3b).
The pathway by which the SOA loss occurs should be clarified (particle-phase reaction, gas-phase reaction, photolysis).

2.4 Oxidation of isoprene by HO: formation of methyl-glyceric acid under high-NO\textsubscript{x} conditions

Under high-NO\textsubscript{x} conditions, Surratt et al. (2006) showed that methyl-glyceric acid (MGA) and its nitrate derivative can oligomerize and form substantial amount of SOA. Recent studies showed that MGA is formed from the oxidation of MPAN (PAN-type molecule derived from the oxidation of methacrolein) (Surratt et al., 2010; Chan et al., 2010) and that MGA strongly depends on the initial \([\text{NO}_2]/[\text{NO}]\) ratio (because MPAN formation is favored under high NO\textsubscript{2} conditions).

To model SOA formation under high-NO\textsubscript{x} conditions, we assumed that SOA formation comes mainly from MGA and its nitrate derivative formation via MPAN oxidation. Currently, there is not enough information to distinguish MGA formation from its nitrate derivative formation in the model. However, experiments by Surratt et al. (2006) show that the two molecules are formed in equivalent quantities when isoprene is oxidized under high-NO\textsubscript{x} conditions. We assumed that MPAN oxidation leads to the same overall mass (particle+gas) of MGA and its nitrate derivative. It means that there is 1.375 (molecular weight ratio of the two compounds) times as much MGA as nitrate:

\[ \text{MPAN} + \text{HO} \Rightarrow \ldots + 1.375 \lambda \text{BiMGA} + \lambda \text{BiNGA} \quad \text{(R11)} \]

where BiMGA represents MGA, BiNGA represents its nitrate derivative and \(\lambda\) is the stoichiometric coefficient for BiNGA. However, when methacrolein is directly oxidized, a higher quantity of nitrate derivative is formed. This fact could be due to faster MPAN formation when methacrolein is directly oxidized: more NO\textsubscript{x} is available to react with MPAN, which leads to higher nitrate derivative concentrations. When isoprene is oxidized, methacrolein is formed first and MPAN formation occurs latter in the oxidation process. Then, there is more time for NO\textsubscript{x} to be consumed to form HNO\textsubscript{3}. As the
nitrate derivative has a higher molecular weight than MGA and a priori a similar saturation vapor pressure, the hypothesis made could lead to underestimation of the SOA mass in environmental chamber under some conditions.

To take into account oligomerization, a simple parameterization was used:

\[ K_{\text{oligo}} = \frac{A_{\text{oligomer}}}{A_{\text{monomer}}} \]  

(2)

where \( K_{\text{oligo}} \) represents the ratio of the oligomer mass \( (A_{\text{oligomer}}) \) to the monomer mass \( (A_{\text{monomer}}) \). With this simple parameterization, an effective partitioning coefficient can be calculated by:

\[ K_{\text{eff,om},i} = K_{\text{om},i}(1 + K_{\text{oligo}}) \]

(3)

where \( K_{\text{om},i} \) is the absorption equilibrium constant of the monomer. It was calculated by using the saturation vapor pressures obtained with the SIMPOL.1 method for MGA and its nitrate derivative. The same parameter \( K_{\text{oligo}} \) was applied for both BiMGA and BiNGA.

The parameters \( \lambda \) and \( K_{\text{oligo}} \) were selected to minimize the error between the model and the results from Kroll et al. (2006) and Surratt et al. (2006) by a least-square method. A mean temperature of 298 K and a relative humidity of 5% were assumed. Optimization gives \( \lambda = 0.046 \) and \( K_{\text{oligo}} = 64.2 \). Results are presented in Table 4. Figure 4 shows the results of a simulation for 42.7 ppb of isoprene, 98 ppb of NO and 31 ppb of NO\(_2\). The slight difference between the BiMGA and BiNGA concentrations is due to different activity coefficients.

The model successfully reproduces the results of Kroll et al. (2006) within the uncertainties of the experiments except for one experiment, (the lowest quantity of NO\(_x\), the model gives a result \( 7.0 \mu g m^{-3} \) close to the upper value \( 6.9 \mu g m^{-3} \)). This slight overestimation could be attributed to a calculated quantity of radical HO\(_2\) too important or the fact that some reactions (the radical formed from ISHP+HO could react with NO and forms less SOA) are missing in the gas-phase mechanism. For high isoprene
experiments, the model gives relative errors from 5 to 20%, but the model is less sensitive to NO\textsubscript{x} conditions than observed in the experiments. This lack of sensitivity of the model may be due to the fact that the BiMGA/BiNGA ratio in the model does not vary with NO\textsubscript{x} conditions. It may also explain the underestimation of SOA formed when methacrolein is directly oxidized (which leads to a lower BiMGA/BiNGA ratio (Surratt et al., 2006)).

The reactions added to RACM2 for SOA formation from isoprene are listed in Table 5. The first six reactions dominate under low-NO\textsubscript{x} conditions whereas the last two reactions dominate under high-NO\textsubscript{x} conditions.

3 Influence of parameters

3.1 Influence of NO\textsubscript{x} level and NO\textsubscript{2}/NO ratio

Based on the experiments of Kroll et al. (2006), the SOA yield seems to be maximal for an oxidation of 300 ppb of NO\textsubscript{x} and seems to decrease for higher concentrations. The recent study of Chan et al. (2010) shows that under high-NO\textsubscript{x} conditions, the SOA yield strongly depends on the [NO\textsubscript{2}]/[NO] ratio due to MPAN chemistry (MPAN formation is favored under high NO\textsubscript{2} conditions).

To study the impact of the NO\textsubscript{x} level, the amount of SOA formed was calculated as a function of initial NO\textsubscript{x} concentrations (Fig. 5) for different [NO\textsubscript{2}]/[NO] initial ratios (the ratio may change with time). According to the model results, the amount of SOA decreases strongly only for low [NO\textsubscript{2}]/[NO] ratio and is below the amount of SOA formed in no-NO\textsubscript{x} conditions for very high concentrations of NO\textsubscript{x} (more than 500 ppb of NO\textsubscript{x} for 45 ppb of oxidized isoprene and an initial [NO\textsubscript{2}]/[NO] ratio of 0.1).

For higher [NO\textsubscript{2}]/[NO] ratios, it seems that the SOA yield is higher under high-NO\textsubscript{x} conditions than under low-NO\textsubscript{x} conditions. For an initial [NO\textsubscript{2}]/[NO] ratio of 5, the quantity of SOA formed appears insensitive to the NO\textsubscript{x} quantity. However, in reality, a stronger sensitivity may be observed if the BiMGA/BiNGA ratio were sensitive to NO\textsubscript{x} concentrations.
For a [NO₂]/[NO] ratio of 1, the final SOA yield (when there is no more SOA growth or loss) was calculated as a function of initial isoprene concentration for several initial concentrations of NOₓ (Fig. 6). For the no-NOₓ simulation, the yield increases and stabilizes at around 3.75%. In this case, a typical SOA yield curve is obtained. For high-NOₓ simulations, the evolution of the yield is more complex, because of the MPAN chemistry. Yields can vary from 5 to 7% if more than 100 ppb of isoprene is oxidized. For 300 ppb of NOₓ, the yield decreases slightly between 300 and 400 ppb of oxidized isoprene and increases for higher concentrations of isoprene.

It is difficult to deduce a general pattern for isoprene SOA formation in presence of NOₓ because of the complex effects of the gas-phase chemistry. Depending on NOₓ concentrations and the [NO₂]/[NO] ratio, the yield can be lower or as much as twice higher than the no-NOₓ SOA yield. However, for atmospheric conditions (NOₓ and isoprene concentrations generally inferior to 100 ppb), the yield should be higher under high-NOₓ conditions than under low-NOₓ conditions, when SOA formation occurs in an organic phase.

3.2 Influence of temperature

Figure 7 shows the effect of temperature on the amount of SOA formed for different [NO₂]/[NO] ratios. In all cases, a decrease of the temperature by 20 °C leads to an increase of SOA mass by about 2 µg m⁻³. This result can be explained by an increase in SOA volatility when the temperature increases. The temperature affects SOA partitioning more than the kinetics of the reactions leading to the formation of SOA. However, the activation energy was not used for all the reactions due to a lack of data and the chemical kinetics could be in reality more sensitive to temperature.
4 Extension of the model to humid conditions

The model described above applies to dry conditions. However, it is important to implement in an air quality model an SOA model that covers the full range of atmospheric conditions including high humidity conditions. Consequently, a parameterization for SOA formation must take into account humid conditions. Moreover, the SOA compounds produced from isoprene photooxidation may be highly hydrophilic due to their structural composition, i.e., short compounds (number of carbons ≤5) with several functional groups. Under humid conditions, in presence of both an aqueous phase and an organic liquid phase, isoprene SOA surrogates may partition mostly between the aqueous phase and the gas phase rather than the organic phase and the gas phase.

For the extension to humid conditions, a parameterization for the partitioning between the gas phase and the aqueous phase was added to the model described above. It was assumed that all surrogate SOA compounds can partition between the gas phase and any liquid phase (aqueous phase and organic phase). Surratt et al. (2010) showed that under low-NO$_x$ conditions, an epoxidiol intermediate should be formed from Reaction (R5) with a high yield (about 75%). This epoxidiol can lead to methyltetrol and other components formation in the presence of an acid aerosol. However, the study of aqueous-phase reaction (Ervens et al., 2008) was considered outside the scope of this work and was not modeled here.

4.1 Parameterization of the partitioning between an aqueous phase and the gas phase

To extend the model to humid conditions, the partitioning between the gas phase and the aqueous phase is calculated with a modified Henry’s law (Pun et al., 2002):

$$H_i = \frac{A_i \zeta_i}{M_i \text{LWC}_p}$$  \hspace{1cm} (4)
with $H_i$ the Henry's law constant of compound $i$ (µM atm$^{-1}$), $p_i$ the partial pressure of compound $i$ in the gas phase (atm), LWC the liquid water content ($L_{\text{water}}/m^3$ of air), $M_i$ the molar mass of compound $i$ and $\zeta_i$ the activity coefficient of compound $i$ in the aqueous phase defined by reference to the infinite dilution. $\zeta_i$ can be computed by using the UNIFAC model:

$$\zeta_i = \frac{\gamma_i}{\gamma_i^\infty}$$  \hspace{1cm} (5)

where $\gamma_i$ is the activity coefficient of the compound $i$ in the aqueous phase computed with UNIFAC and $\gamma_i^\infty$ is the activity coefficient of compound $i$ at infinite dilution computed with UNIFAC.

Different methods can be used to determine the Henry's law constant. Functional group methods give very high value for the Henry's law constant of BiMT ($2.7 \times 10^{16}$ M atm$^{-1}$ and $1.66 \times 10^{16}$ M atm$^{-1}$ with the methods of Meylan and Howard (2000) and Suzuki et al. (1992), respectively). On the contrary, the HENRYWIN bond contribution method of Meylan and Howard (2000, 1991) predicts a lower value for BiMT: $2.45 \times 10^6$ M atm$^{-1}$. However, according to the data available in Raventos-Duran et al. (2010), the HENRYWIN bond contribution method seems to underestimate the Henry’s constant for compounds with several hydroxy groups (for example, for propane-1,3-diol, this method predicts a Henry’s law constant of $6.31 \times 10^3$ M atm$^{-1}$ whereas the experimental value is $1.0 \times 10^6$ M atm$^{-1}$). SPARC online (Hilal et al., 2008) seems then to provide the best estimation for Henry’s constants ($3.38 \times 10^{10}$ M atm$^{-1}$ for BiMT and $5.25 \times 10^8$ M atm$^{-1}$ for BiMGA). However, SPARC online does not provide algorithms to calculate the Henry’s law constants of the other compounds.

We chose to calculate the other Henry’s law constants by reference to either BiMT or BiMGA using the following equation:

$$H_i = \frac{\gamma_j^\infty p_j^o}{\gamma_i^\infty p_i^o} H_j$$  \hspace{1cm} (6)
where \( j \) is the compound of reference (BiMT or BiMGA) and \( i \) is the compound for which Henry’s law constant is wanted. The Henry’s law constants for BiPER, BiDER and \( \text{C}_5\text{H}_9\text{N}_3\text{O}_{10} \) were calculated by reference to BiMT. The Henry’s law constant for BiNGA was calculated by reference to BiMGA. The saturation vapor pressures used in the model for dry conditions were used for the calculation. The calculated values are presented in Table 6.

### 4.2 Influence of an aqueous phase on SOA formation

The partitioning of the surrogate compounds under atmospheric conditions was calculated for four different cases. In the first case, SOA partition between the gas phase and an organic phase constituted by Primary Organic Aerosols (POA). A mean value of 5 µg m\(^{-3}\) for POA concentrations was chosen as follows: 40% of \( \text{C}_{23}\text{H}_{47}\text{COOH} \), 5% of \( \text{C}_8\text{H}_{17}\text{CH}=\text{CHC}_7\text{H}_{14}\text{COOH} \), 15% of 4-(2-propio)-syringone, 12% of \( \text{C}_{29}\text{H}_{60} \) and 28% of 2-carboxybenzoic acid (EPRI, 1999). In the second case, SOA partition between the gas phase and an aqueous phase. A value of 50 µg m\(^{-3}\) was used in the second case. The aqueous aerosol was supposed to be too acid for BiMGA and BiNGA to dissociate. In the third case, SOA can condense on 5 µg m\(^{-3}\) of POA and on 50 µg m\(^{-3}\) of liquid water. In the fourth case, surrogates can condense on an organic phase assumed to be ideal (activity coefficients equal to 1).

Under no-NO\(_x\) conditions ([NO]=[NO\(_2\])=0.0), no substantial amount of SOA is formed in the first case (maximum of SOA formed is 0.27 µg m\(^{-3}\)) whereas a significant amount is formed in the second case. The low amount of SOA formed in the first case is due to low affinity of the surrogate compounds formed under those conditions (mainly BiPER and BiDER) with POA. In the second and third case, the same amount of SOA is formed (because all SOA compounds seem to partition only in the aqueous phase). Figure 8 shows the evolution of SOA formed for the second and third cases. A substantial amount of isoprene is formed (1.6 µg m\(^{-3}\) at the end of the oxidation and a maximum at 6.7 µg m\(^{-3}\)). The amount of SOA formed is even superior to the case where SOA would condense on an ideal organic phase, which gives much
greater concentrations than the non-ideal organic phase case. This result shows the importance of taking into account the non-ideality of particles for SOA partitioning. Furthermore, SOA formed from isoprene photooxidation under low-NO$_x$ conditions are not likely to condense on POA (due to non-ideality of the organic phase), it would more likely condense efficiently on an aqueous phase. It should be noted that, whereas surrogates do not condense on POA, they condense almost entirely on the aqueous phase (only 1% of BiPER remains in the gas phase).

Under high-NO$_x$ conditions ([NO]=[NO$_2$]=50 ppb), all the components formed are not highly hydrophilic. Figure 9 shows the quantity of SOA formed for the different cases under high-NO$_x$ conditions. There is still a lower quantity of SOA (0.16 µg m$^{-3}$) formed when SOA condense on POA than when they condense on an aqueous phase (0.59 µg m$^{-3}$). But, contrary to the low-NO$_x$ conditions, the third case shows that a significant part of the surrogates may condense on the organic phase. The fourth case (ideal organic phase) strongly increases the mass of SOA. The difference between low-NO$_x$ and high-NO$_x$ conditions is due to lower Henry’s law constants of the surrogate species, which dominate under high-NO$_x$ conditions. Whereas, the measured SOA yield was higher for dry conditions under high-NO$_x$ conditions than under low-NO$_x$ conditions, it is possible that in the atmosphere under high-NO$_x$ conditions, less SOA could be formed due to lower water solubility of the corresponding SOA. The increase in SOA mass under humid conditions is commensurate with the results obtained by Pun (2008) who estimates, using a different approach, an increase by a factor of five of SOA mass by taking into account the hydrophilic properties of SOA from isoprene.

To explain the absorption of surrogates in the organic phase in high-NO$_x$ but not in low-NO$_x$, the distribution of the SOA (formed from the oxidation of 10 ppb of isoprene with 50 ppb of NO and NO$_2$) between the organic and aqueous phases for different $M_{\text{water}}/M_{\text{organic}}$ ratios is shown in Fig. 10. BiMT, BiPER and BiDER (compounds formed under low-NO$_x$ conditions) are only present in the aqueous phase and do not condense significantly in the organic phase. BiMGA (formed under high-NO$_x$) does not condense on the aqueous as efficiently as the surrogates formed under low-NO$_x$
conditions. However, for a $M_{\text{water}}/M_{\text{organic}}$ ratio of 1, 60% of particulate BiMGA is in the aqueous phase and around 95% for a ratio of 10. When, the $M_{\text{water}}/M_{\text{organic}}$ ratio is superior to 1 in the atmosphere, BiMGA should partition preferably between the aqueous phase and the gas phase. BiNGA and $C_5$-hydroxy-trinitrate seem to be hydrophobic because even for a $M_{\text{water}}/M_{\text{organic}}$ ratio of 10, they condense preferably on the organic phase. The condensation of SOA on the organic phase under high-NO$_x$ conditions is then due to the hydrophobic properties of BiNGA (only a small quantity of $C_5$-hydroxy-trinitrate is formed under the conditions considered here).

AEC (Pun et al., 2002, 2006) is a model for SOA formation that takes into account the hydrophilic and hydrophobic properties of surrogates. It distinguishes compounds into two types: hydrophilic compounds (which are absorbed only in an aqueous particle) and hydrophobic compounds (which are absorbed only in an organic particle). In AEC, a compound cannot condense on both phases; we investigate here the uncertainty associated with this formulation. According to the distribution of the surrogates, BiMT, BiPER, BiDER and BiMGA may be considered hydrophilic compounds and BiNGA and $C_5H_9N_3O_{10}$ hydrophobic compounds. Under low-NO$_x$ conditions, as the compounds formed are highly hydrophilic and condense almost entirely on organic particles, the hypothesis of AEC does not change the results significantly. Figure 11 shows the evolution of SOA formed with time for the high-NO$_x$ case, assuming that surrogates are only hydrophilic or only hydrophobic. The difference between the case where all compounds can partition between all the phases (see Fig. 8) and the case with the hypothesis of AEC is greater for high-NO$_x$ conditions because BiMGA and BiNGA (major components formed under high-NO$_x$ conditions) are neither totally hydrophilic or hydrophobic. In this case, the mass of SOA is underestimated by the AEC hypothesis by 11%.
4.3 Influence of oligomerization and particle pH on partitioning

SOA surrogates have been separated between hydrophobic and hydrophilic compounds, however, calculated partitioning can change whether oligomerization is taken into account in the computation or not. We assumed that esterification does not occur in the aqueous phase, although Altieri et al. (2008) observed formation of oligoesters in an aqueous phase. Additional absorption of acidic SOA species (BiMGA and BiNGA) occurs by dissociation in an aqueous phase. For a pH superior to $pK_a$ ($pK_a$=4 estimated for BiMGA with SPARC online), BiMGA and BiNGA would, therefore, dissociate and would be more efficiently absorbed. On the other hand, oligomerization in the organic phase could be less important in the atmosphere due to less concentrated monomers. The amount of SOA formed (for 10 ppb of isoprene, 50 ppb of NO, 50 ppb of NO$_2$, 5 µg m$^{-3}$ of POA and 50 µg m$^{-3}$ of liquid water) was simulated to investigate the sensitivity of the system to additional absorption due to acid dissociation and oligomerization.

To investigate the effect of potential oligomerization in the aqueous phase and pH, an effective Henry’s law constant for BiMGA and BiNGA was used:

$$H_{\text{eff},i} = H_i (1 + K_{\text{oligo}})$$  \hspace{1cm} (7)

where $K_{\text{oligo}}$ is the same constant as the one in Eq. (3) (it supposes that additional absorption due to dissociation and oligomerization occurs with the same extent in both phase). Figure 12a shows the evolution of the SOA mass. Additional absorption of BiMGA and BiNGA greatly increases the amount of SOA formed (1.7 µg m$^{-3}$ against 0.68 µg m$^{-3}$ without additional absorption). Moreover, there is almost no condensation on POA: all the condensation occurs on the aqueous phase. Figure 13a shows the distribution of the surrogates between the two condensed phases for different $M_{\text{water}}/M_{\text{organic}}$ ratios. For a $M_{\text{water}}/M_{\text{organic}}$ ratio of 1, almost all the particulate BiMGA is in the aqueous phase and nearly 80% of the particulate BiNGA is in the aqueous phase. Contrary to the case where there is no additional absorption in the aqueous phase (see Fig. 10), BiNGA can then be considered hydrophilic.
To investigate the possible absence of oligomerization in the organic phase, \( K_{\text{oligo}}=0 \) was used. Figure 12b shows the evolution of the SOA mass without oligomerization in the organic phase. The final SOA mass is reduced by 16% (0.57 µg m\(^{-3}\) without oligomerization against 0.68 µg m\(^{-3}\) with oligomerization). There is no significant absorption of the surrogates on the organic phase. Figure 13b shows the distribution of the surrogates between the two phases. As in the previous case, BiNGA partitions more efficiently between the aqueous phase and the gas phase and can be considered hydrophilic.

Whether BiNGA condenses on POA or an aqueous phase depends on potential additional absorption in the aqueous phase by potential oligomerization and dissociation and on the extent of oligomerization in the organic phase. Most of the components can be classified as hydrophilic (BiMT, BiPER, BiDER and BiMGA) but BiNGA can be considered either as hydrophobic or hydrophilic, depending on oligomerization in the organic phase or additional absorption in the aqueous phase.

### 4.4 Sensitivity of the model to Henry’s law constants

One of the main uncertainties of the model is the values of Henry’s law constants. The constants used here were estimated using the saturation vapor pressure and the activity coefficient at infinite dilution; however, as discussed above, other methods may lead to quite different results. For that reason, a sensitivity study was conducted to evaluate the impact of those uncertainties on liquid-gas partitioning.

Under low-NO\(_x\) conditions, the partitioning is not very sensitive to the values. If Henry’s law constants are divided by 2, 1.76 µg m\(^{-3}\) of SOA are formed instead of 1.83 µg m\(^{-3}\) (for 50 µg m\(^{-3}\) of liquid water and 10 ppb of initial isoprene). If the Henry’s law constants are divided by 10, 0.92 µg m\(^{-3}\) is formed (i.e. half of SOA previously formed). On the contrary, when Henry’s law constants are multiplied by 10, the quantity of SOA does not change significantly (1.86 µg m\(^{-3}\) instead of 1.83 µg m\(^{-3}\)) because most of the SOA species are already present in the condensed phase. So, for BiMT, BiPER and BiDER, the amount of SOA formed does not change significantly for a small
decrease of the Henry’s law constants or for an important increase of the Henry’s law constant. Of course, the impact of Henry’s law constant values is more important for lower quantity of liquid water, but even at 5 µg m\(^{-3}\) of liquid water, Henry’s law constant values have a small impact on SOA formation: 1.46 µg m\(^{-3}\) of SOA for Henry’s law constants divided by two, and 1.86 µg m\(^{-3}\) for Henry’s constants multiplied by ten instead of 1.73 µg m\(^{-3}\).

Under high-NO\(_x\) conditions, the partitioning is more sensitive. However, it was shown previously that using effective Henry’s law constants taking into account additional absorption due to oligomerization and dissociation greatly influences the partitioning of the surrogates. The main uncertainty for BiMGA and BiNGA partitioning is then the extent of the additional absorption rather than the actual Henry’s law constants for BiMGA and BiNGA.

For C\(_5\)H\(_9\)N\(_3\)O\(_{10}\) formed by oxidation of isoprene by NO\(_3\) radicals, as the component seems hydrophobic, the value of its Henry’s law constant does not impact its partitioning noticeably.

5 Conclusions

A model for SOA formation from isoprene oxidation that takes into account different chemical regimes and the hydrophilic properties of SOA has been developed. For low-NO\(_x\) conditions, SOA was found to be highly hydrophilic and it is likely that components formed would condense on an aqueous phase rather than on an organic phase. For high-NO\(_x\) conditions, methyl glyceric acid (BiMGA) was found to be hydrophilic whereas the nitrate derivative (BiNGA) can be considered as hydrophilic or hydrophobic depending on the hypothesis formulated.

A major conclusion is the importance of the hydrophilic properties of SOA formed from isoprene oxidation. Because the parameterizations derived from smog chambers experiments apply to dry conditions, SOA formation could be underestimated in current models.
This SOA model can be implemented in an air quality model to evaluate the amount of isoprene-SOA formed in the atmosphere. However, several questions remain before implementing this model. Are all the pathways described here important for SOA formation? The formation of C₅-hydroxy-trinitrate may not be relevant in the atmosphere and in any case, it could be a minor pathway for the formation of SOA in the atmosphere. If 3-D simulations show that those compounds do not contribute significantly to the SOA burden, this pathway could be removed from the model. Can the number of surrogates be reduced? Some surrogates like BiMT and BiPER have similar thermodynamic properties (same molecular structure and high Henry’s law constants), and they could perhaps be merged into a single surrogate.

It should be noted that the main uncertainties of the model concern the high-NOₓ regime. However, isoprene is emitted by biogenic sources in rural areas where concentrations of NOₓ are low. Most of SOA should therefore, be formed under low-NOₓ conditions and the impact of the uncertainties of the high-NOₓ regime should be limited.

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Zhang, Q., Jimenez, J., Canagaratna, M., Allan, J., Coe, H., Ulbrich, I., Alfarrà, M., Takami, A.,
### Table 1. Properties of the selected SOA surrogate species.

<table>
<thead>
<tr>
<th>Surrogate species</th>
<th>Calculated saturation vapor pressure(^a) (torr)</th>
<th>Selected saturation vapor pressure(^b) (torr)</th>
<th>Formula</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methyl-tetrol</td>
<td>(7.35 \times 10^{-7}) at (298) K</td>
<td>(1.12 \times 10^{-6}) at (298) K</td>
<td><img src="image" alt="Methyl-tetrol structure" /></td>
</tr>
<tr>
<td>dihydroxy-dihydroperoxide</td>
<td>(4.07 \times 10^{-7}) at (298) K</td>
<td>(2.61 \times 10^{-6}) at (298) K</td>
<td><img src="image" alt="Dihydroxy-dihydroperoxide structure" /></td>
</tr>
<tr>
<td>(\text{C}_5)-hydroxy-trinitrate</td>
<td>(3.6 \times 10^{-7}) at (293) K</td>
<td>(1.12 \times 10^{-6}) at (293) K</td>
<td><img src="image" alt="C5-hydroxy-trinitrate structure" /></td>
</tr>
<tr>
<td>Methyl-glyceric acid</td>
<td>(1.40 \times 10^{-5}) at (298) K</td>
<td>(1.40 \times 10^{-5}) at (298) K</td>
<td><img src="image" alt="Methyl-glyceric acid structure" /></td>
</tr>
<tr>
<td>Nitrate organic acid</td>
<td>(1.39 \times 10^{-5}) at (298) K</td>
<td>(1.39 \times 10^{-5}) at (298) K</td>
<td><img src="image" alt="Nitrate organic acid structure" /></td>
</tr>
</tbody>
</table>

\(^a\) The saturation vapor pressures were calculated using the SIMPOL.1 method (Pankow and Asher, 2008).

\(^b\) Modified for a better fitting between results from the model and experiments.
Table 2. Values of the optimized parameters used for low-NO\textsubscript{x} conditions.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\alpha$</td>
<td>0.28</td>
</tr>
<tr>
<td>$\beta$</td>
<td>0.030</td>
</tr>
<tr>
<td>BiPER degradation</td>
<td>$2.0 \times 10^{-4}$ molecule$^{-1}$ cm$^3$ s$^{-1}$</td>
</tr>
<tr>
<td>BiPER saturation vapor pressure</td>
<td>$2.6 \times 10^{-6}$ torr</td>
</tr>
<tr>
<td>BiDER saturation vapor pressure</td>
<td>$4.1 \times 10^{-7}$ torr</td>
</tr>
</tbody>
</table>
Table 3. Comparison of measured\(^a\) and modeled SOA formation (\(\Delta M_o\)) for low-NO\(_x\) conditions: maximum \(\Delta M_o\) and final \(\Delta M_o\).

<table>
<thead>
<tr>
<th>Initial isoprene (ppb)</th>
<th>Measured (\Delta M_o) (max) (µg m(^{-3}))</th>
<th>Calculated (\Delta M_o) (max) (µg m(^{-3}))</th>
<th>Measured (\Delta M_o) (final) (µg m(^{-3}))</th>
<th>Calculated (\Delta M_o) (final) (µg m(^{-3}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>90.0</td>
<td>27.0 ± 0.5</td>
<td>26.9</td>
<td>9.3 ± 0.4</td>
<td>8.6</td>
</tr>
<tr>
<td>46.1</td>
<td>13.5 ± 0.6</td>
<td>11.5</td>
<td>3.8 ± 0.5</td>
<td>3.4</td>
</tr>
<tr>
<td>23.0</td>
<td>2.3 ± 0.5</td>
<td>2.8</td>
<td>0.6 ± 0.3</td>
<td>0.5</td>
</tr>
<tr>
<td>12.2</td>
<td>0.7 ± 0.1</td>
<td>0.07</td>
<td>0.3 ± 0.1</td>
<td>0.02</td>
</tr>
<tr>
<td>63.6</td>
<td>17.8 ± 0.5</td>
<td>18.0</td>
<td>5.0 ± 0.5</td>
<td>5.6</td>
</tr>
<tr>
<td>29.4</td>
<td>6.2 ± 0.8</td>
<td>5.1</td>
<td>2.2 ± 0.5</td>
<td>1.3</td>
</tr>
<tr>
<td>47.8</td>
<td>11.1 ± 0.5</td>
<td>12.2</td>
<td>3.0 ± 0.4</td>
<td>3.7</td>
</tr>
<tr>
<td>41.6</td>
<td>8.4 ± 0.4</td>
<td>9.8</td>
<td>2.4 ± 0.5</td>
<td>2.9</td>
</tr>
</tbody>
</table>

\(^a\) Measured values taken from Kroll et al. (2006).
Table 4. Comparison of measured and modeled SOA formation ($\Delta M_o$) for high-NO$_x$ conditions.

<table>
<thead>
<tr>
<th>Initial reagent</th>
<th>Initial NO (ppb)</th>
<th>Initial NO$_2$ (ppb)</th>
<th>Measured $\Delta M_o$ (µg m$^{-3}$)</th>
<th>Calculated $\Delta M_o$ (µg m$^{-3}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kroll et al. (2006) (isoprene)</td>
<td>46.7</td>
<td>242</td>
<td>24</td>
<td>$6.3 \pm 1.0$</td>
</tr>
<tr>
<td></td>
<td>43.5</td>
<td>496</td>
<td>30</td>
<td>$2.9 \pm 1.2$</td>
</tr>
<tr>
<td></td>
<td>49.1</td>
<td>51</td>
<td>27</td>
<td>$5.6 \pm 1.3$</td>
</tr>
<tr>
<td></td>
<td>42.7</td>
<td>337</td>
<td>68</td>
<td>$4.6 \pm 1.0$</td>
</tr>
<tr>
<td></td>
<td>42.0</td>
<td>708</td>
<td>37</td>
<td>$1.7 \pm 1.1$</td>
</tr>
<tr>
<td>Surratt et al. (2006) (isoprene)</td>
<td>500</td>
<td>827</td>
<td>34</td>
<td>74</td>
</tr>
<tr>
<td></td>
<td>500</td>
<td>759</td>
<td>112</td>
<td>73</td>
</tr>
<tr>
<td></td>
<td>500</td>
<td>805</td>
<td>87</td>
<td>104</td>
</tr>
<tr>
<td></td>
<td>500</td>
<td>825</td>
<td>80</td>
<td>111</td>
</tr>
<tr>
<td></td>
<td>500</td>
<td>891</td>
<td>74</td>
<td>95</td>
</tr>
<tr>
<td>Surratt et al. (2006) (methacrolein)</td>
<td>500</td>
<td>791</td>
<td>60</td>
<td>181</td>
</tr>
<tr>
<td></td>
<td>500</td>
<td>898</td>
<td>30</td>
<td>197</td>
</tr>
</tbody>
</table>
Table 5. Reactions and products added to RACM2.

<table>
<thead>
<tr>
<th>Reactions</th>
<th>kinetic rate parameters (molecule$^{-1}$ cm$^3$ s$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ISHP+HO$\Rightarrow$...+0.28BiPER+0.030BiDER</td>
<td>$3.0 \times 10^{-11}$</td>
</tr>
<tr>
<td>BiPER$\Rightarrow$Unknown Products</td>
<td>$2.0 \times 10^{-4}$</td>
</tr>
<tr>
<td>2ISOP$\Rightarrow$2MACR+HCHO+HO$_2$+0.16DIOL</td>
<td>$2.0 \times 10^{-12}$</td>
</tr>
<tr>
<td>ISOP+MO$_2$$\Rightarrow$...+0.16DIOL</td>
<td>$3.40 \times 10^{-14}$ exp(221/T)</td>
</tr>
<tr>
<td>ISOP+ACO$_3$$\Rightarrow$...+0.16DIOL</td>
<td>$8.40 \times 10^{-14}$ exp(221/T)</td>
</tr>
<tr>
<td>DIOL+HO$\Rightarrow$0.16BiMT+HO*</td>
<td>$1.30 \times 10^{-10}$</td>
</tr>
<tr>
<td>ISON+NO$_3$$\Rightarrow$0.074C$_5$H$_9$N$<em>3$O$</em>{10}$</td>
<td>$6.61 \times 10^{-13}$</td>
</tr>
<tr>
<td>MPAN+HO$\Rightarrow$...+0.063BiMGA+0.046BiNGA</td>
<td>$3.20 \times 10^{-11}$</td>
</tr>
</tbody>
</table>

* Oxidant HO is added as both reactant and product in order not to modify HO prediction by RACM2.
Table 6. Calculated Henry’s law constants for SOA surrogate species.

<table>
<thead>
<tr>
<th>Surrogate species</th>
<th>$H_i$ (M atm$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BiMT</td>
<td>$3.30 \times 10^{10}$</td>
</tr>
<tr>
<td>C$_5$H$_9$N$<em>3$O$</em>{10}$</td>
<td>$4.75 \times 10^{6}$</td>
</tr>
<tr>
<td>BiMGA</td>
<td>$5.25 \times 10^{8}$</td>
</tr>
<tr>
<td>BiNGA</td>
<td>$3.73 \times 10^{7}$</td>
</tr>
<tr>
<td>BiPER</td>
<td>$8.09 \times 10^{9}$</td>
</tr>
<tr>
<td>BiDER</td>
<td>$8.91 \times 10^{10}$</td>
</tr>
</tbody>
</table>
Fig. 1. SOA yield data and yield curve for isoprene-NO$_3$ reaction using one surrogate compound.
Fig. 2. SOA growth versus isoprene reacted. Dots correspond to measurements from Kroll et al. (2006). Lines correspond to results from the model.
Fig. 3. Evolution of the calculated mass of SOA formed from isoprene photooxidation as a function of time (a) for an initial concentration of 63.6 ppb of isoprene (b) for an initial isoprene concentration of 500 ppb.
Fig. 4. Evolution of the calculated mass of SOA formed from the photooxidation of isoprene as a function of time for initial concentrations of 42.7 ppb of isoprene, 98 ppb of NO and 31 ppb of NO$_2$. 
Fig. 5. Evolution of calculated SOA mass with NO\textsubscript{x} for 45 ppb of oxidized isoprene and different initial [NO\textsubscript{2}]/[NO] ratios.
Fig. 6. Evolution of calculated yield with isoprene concentration for different NOx concentrations with a \([\text{NO}_2]/[\text{NO}]\) ratio of 1.
Fig. 7. Evolution of SOA with NO\textsubscript{x} for 45 ppb of oxidized isoprene at different temperatures for: (a) [NO\textsubscript{2}]/[NO] ratio of 0.1, (b) [NO\textsubscript{2}]/[NO] ratio of 1, (c) [NO\textsubscript{2}]/[NO] ratio of 5.
Fig. 8. Effect of an aqueous phase on SOA formation under no-NO$_x$ conditions for 10 ppb of initial isoprene concentration for different cases. Case 1: Condensation on 5 µg m$^{-3}$ of POA. Case 2: Condensation on 50 µg m$^{-3}$ of liquid water. Case 3: Condensation on 5 µg m$^{-3}$ of POA and on 50 µg m$^{-3}$ of liquid water. Case 4: Condensation on 5 µg m$^{-3}$ of an ideal organic phase.
Fig. 9. Effect of an aqueous phase on SOA formation under high-NOx conditions ([ISO]=[NO]=[NO2]=50 ppb). Case 1: Condensation on 5 µg m\(^{-3}\) of POA. Case 2: Condensation on 50 µg m\(^{-3}\) of liquid water. Case 3: Condensation on 5 µg m\(^{-3}\) of POA and on 50 µg m\(^{-3}\) of liquid water. Case 4: Condensation on 5 µg m\(^{-3}\) of an ideal organic phase.
Fig. 10. Distribution of the absorbed surrogates between the aqueous phase and the organic phase for different $M_{\text{water}}/M_{\text{organic}}$ ratios if oligomerization only happens in the organic phase.
Fig. 11. Simulated SOA concentration in aqueous and organic phases for 10 ppb of isoprene and high-NO$_x$ conditions ([NO]=[NO$_2$]=50 ppb) assuming all the surrogates are hydrophilic except for BiNGA and C$_5$H$_9$N$_3$O$_{10}$. 
**Fig. 12.** Evolution with time of the mass of SOA formed: (a) with additional absorption in the aqueous phase (acid dissociation and oligomerization) and (b) with no additional absorption in the aqueous phase and no oligomerization in the organic phase.
Fig. 13. Distribution of the absorbed surrogates between the aqueous phase and the organic phase for different $M_{\text{water}}/M_{\text{organic}}$ ratios: (a) with additional absorption in the aqueous phase (acid dissociation and oligomerization) and (b) with no additional absorption in the aqueous phase and no oligomerization in the organic phase.