Authors’ Response to Reviews of

GENerator of reduced Organic Aerosol mechanism (GENOA v1.0): An automatic generation tool of semi-explicit mechanisms

Zhizhao Wang, Florian Couvidat, Karine Sartelet

Reviewer # 1

1. General comments

RC: This paper discusses procedures developed to reduce the size of large gas/aerosol mechanisms to greatly reduced mechanisms that give predictions of secondary organic aerosol (SOA) that agree with those of the larger mechanism in a selected set of environmental conditions to within a specified tolerance. Because of the complexity of the atmospheric reactions or organic compounds, it is necessary to use reduced mechanisms in practical airshed model applications, but most reduced mechanisms used in airshed model applications were developed primarily focused on accurate ozone predictions. However, SOA predictions are much more affected by the chemical complexity of organic reactions than predictions of ozone, and developing methods to reduce mechanisms for without significantly affecting SOA predictions is an important research priority in atmospheric chemical mechanism development.

In fact, work with GECKO suggest that multi-generation mechanisms may be necessary for reliable SOA predictions, and even the MCM, which greatly lumps reactions of 2nd and higher generation products, may be too reduced for this application. However, the use of MCM as an example is sufficient to illustrate the method, and as discussed below MCM may be about the largest mechanism that could be reduced using the method discussed in this work, given current computer capabilities. In any case, without some way to reduce these huge mechanisms, we have no choice to continue to rely on the empirical and parameterized SOA models that are adjusted to fit SOA yields measured environmental chamber data without consideration of the actual chemistry and how different chemical conditions in the atmosphere affect SOA yields. If we had suitable reduction methods for SOA predictions, then use of SOA models that are based on actual chemistry might become practical.

AR: The authors would like to first thank Dr. William Carter for his detailed comments and insightful suggestions. Many of these suggestions are constructive and have been taken into account for the further development of the GENOA algorithm.

We have carefully considered the comments and revised the manuscript accordingly. Please note that the line and section numbers mentioned in the response correspond to the version of the manuscript before revision.

RC: The method discussed here involves use of a 3D grid model representing a large continental domain and various seasons, with SOA calculated using the large mechanism to be reduced as the starting point, and then uses sets of 0D scenarios derived from selected grid cells and times during the grid model simulation in an algorithm to develop the reduced mechanism. This has the disadvantage that it requires a full 3D calculation
The focus of this paper is a specific software package developed for mechanism reduction, which they call GENOA, though the main interest of this paper from a scientific perspective is the method itself. The name of the software is somewhat misleading because it is not actually a mechanism generator, but instead is a method to reduce existing mechanisms, potentially including those developed by actual mechanism generators such as GECKO.

The role of GENOA is more like a "generator of reduced mechanisms" than as a "generator of mechanisms". In our opinion, the term "generator" might be better than other terms such as "producer" or "reducer". However, it is true that the term "generator" could still be misleading. Thus, GENOA is always referred to as
a "generator of reduced mechanisms," rather than a "generator" alone, as GENOA is designed to generate mechanisms from reduction, not from scratch.

RC: This could potentially be a useful tool for the atmospheric chemistry research community, if suitably documented and made publicly available. The fact that they include a users manual along with the Supplementary Information suggests that this is the intention of the authors. If it has sufficient flexibility, it could possibly be used for other criteria besides SOA predictions and perhaps even for other applications besides atmospheric modeling, such as, for example, combustion modeling or reducing large liquid-phase mechanisms. However, the users manual does not contain sufficient information to actually run the model for general applications, other than duplicating the results given in this paper. In particular, it would need information on how to interface this with output of existing 3D air quality models.

AR: Thank you for the positive comments. Certainly, one of the goals of GENOA is to build the reduction algorithm that can be used for other modellers and even for other applications besides SOA prediction.

As explained previously, the monthly diurnal profiles of hourly meteorological data (e.g., temperature, relative humidity), and hourly concentrations of oxidant, radical, and other inorganic species were extracted from 3-D results. The user can, therefore, simply compute these profiles from 3-D results for GENOA reduction.

We have added the following explanation to the manual:

**GENOA user's manual, section 4, page 15:**

For example, to run GENOA over a specific domain and using specific 3D model results, the user has to construct the files described in section 2.2.2 to provide the monthly profiles for different variables.

RC: It is worth pointing out that the reduced mechanisms developed using this are strictly speaking reliable to give predictions to within the desired tolerances only for the airshed conditions used in its development. They developed this example using the conditions of all of Europe for a whole year, but it may be more practical to use this to optimize mechanisms for specific urban scenarios, for use in regulatory modeling. To be useful for this, the users manual would need to be improved so it can be used with other 3D models and modeling scenarios.

Although this paper is reasonably well written, it does have areas where improvements are needed before it is accepted for publication, and I have some suggestions. These are given below in approximate order of importance.

AR: In the example of BCARY reduction, the reduced "Rdc." mechanism was trained under training conditions that covered both high and low NOx regimes. The training condition "ADD2", for instance, received high NO concentrations from Milan, and thus is considered a typical urban condition. Consequently, the "Rdc." mechanism could be adequate for use over Europe or in specific urban scenarios, as it was developed under both low and high-NOx conditions and efficiently reproduces the SOA concentrations in most metropolises (e.g., Paris, Berlin).

Compared to the "Rdc." mechanism, undoubtedly, more reduction could be achieved if the SOA mechanism was trained specifically for urban scenarios using an urban-specific training dataset. In other words, as pointed out by the reviewer, training with fewer and more targeted conditions will optimize the resulting SOA mechanism (more accurate and smaller), which is intended for a small domain or specific scenarios.
2. Improvements need to be done

RC: It looks relative changes in SOA concentrations are used as the criteria to test a reduction approach. This means that 5% error in a grid cell where almost no SOA is formed is given equal weights to 5% error in a grid cell with high SOA, where the model prediction is relatively more important. Wouldn’t absolute error be a better criterion, or at least among the criteria employed? Shouldn’t there be a cutoff to remove cells with very low SOA, or was this incorporated implicitly by the choice of testing scenarios?

AR: Thank you for such an insightful suggestion. We used relative errors in the evaluation of the reduction, because they are sensitive to small changes and it is easy to set universal criteria for conditions with a wide range of SOA concentrations. Because sesquiterpene has high SOA yields, all 0-D simulations in the paper resulted in SOA concentrations consistently exceeding 1 $\mu g/m^3$ (with an initial BCARY concentration of 5 $\mu g/m^3$). Therefore, for BCARY reduction, there may be no need for a cut-off concentration.

We have added the information to the paper:

\textit{Section 2.3, line 267:}

The initial BCARY concentration is taken equal to 5 set to five $\mu g/m^{-3}$ in order to ensure high SOA production the (SOA concentration is always greater than one $\mu g/m^{-3}$ at all evaluated conditions at all conditions).

However, for other precursors that have SOA yields lower than sesquiterpene, a cut-off concentration may be required for conditions with low SOA production, where large relative errors can occur as a result of low SOA concentrations.

The user manual has been updated to inform the user that there are several options that can be taken to avoid this issue:

\textit{GENOA user’s manual, section 4, page 15:}

It should be noted that reduced mechanisms may cause large errors when there is a low SOA concentration. The reason might not be directly linked to reduction of the mechanism performance, but to the evaluation criterion, based on relative errors, which naturally has large variations when the absolute value is small. There are several solutions to resolve this uncertainty:

- The user can set high initial concentrations of the studied SOA precursor in order to ensure high SOA production.
- The user may specify a threshold SOA concentration for condition selection in the training and pre-testing datasets.
- The user may also evaluate reductions using a different type of error.

RC: The 3D model simulations of continental Europe employing the chemistry-transport model CHIMERE was used as the standard against which the reductions were compared. However, CHIMERE mechanism is not exactly the same as MCM, and I could not find an indication of whether the “CHIMERE surrogate for sesquiterpene” (line 198) is exactly the same as the MCM b-caryophyllene mechanism as used in this work. If that is the case, it should be stated explicitly when CHIMERE is first mentioned, since I couldn’t find such a
statement in Lanzafame et al (2022). If it is not the case, then justification needs to be given as to whether this is an appropriate standard against which to test the reductions.

AR: As detailed in reply to one of the general comments, we have not used 3-D but 0-D SOA simulations as a reference to evaluate the reductions. In the 0-D simulation, the concentration profiles of ozone, radicals, and inorganics, as well as environmental parameters, were derived from the 3-D CHIMERE simulations.

In CHIMERE v2020r1, there are several gas-phase chemical mechanisms embedded. We have used the MELCHIOR2 mechanism [Derognat et al., 2003], which contains 120 reactions and less than 80 lumped species. Evidently, it is an implicit gas-phase mechanism that is highly simplified compared to the explicit MCM mechanism.

The species HUMULE in CHIMERE (the MELCHIOR2 mechanism) does not refer to a specific sesquiterpene, but to a lumped class that includes all sesquiterpene species. As we use "HUMULE" concentrations only to determine whether sesquiterpenes are present at a specific location and select several conditions for varying sesquiterpene concentrations in the pre-testing dataset, there is no consistency issue.

Relevant modifications have been done to section 2.2 line 19 of the paper, as mentioned previously.

RC: It is not clear to me whether the concentrations of the inorganic species such as OH, O3, HO2, etc were constrained to be exactly what was calculated by CHIMERE in the 0D models for the selected locations, or if they were calculated using the mechanisms using boundary or initial conditions somehow obtained from CHIMERE. If the former (which I presume to be the case) this should be stated explicitly, and if the latter than more detail about the inputs to the 0D models need to be provided, if only in an Appendix.

AR: Sorry for the ambiguity. As mentioned in reply to the previous comment, the concentrations of certain species (i.e., ozone, radicals, and other inorganics) are constrained to be exactly what was calculated in the 3-D CHIMERE simulations for the selected conditions (locations + month). The concentrations of other species resulting from sesquiterpene degradation are simulated using the 0-D SSH-aerosol model. As the CHIMERE concentrations represent 24-hour monthly averages, the diurnal profiles are repeated for five days in 0-D simulations (the simulation time is five days to consider adequately SOA formation and aging processes).

We have added the explanation of how the coupling is generally resolved between VOC species and oxidants in modeling:

Section 1, line 36:

Along with To complete implicit gas-phase mechanisms, implicit SOA mechanisms have been developed (Kim et al., 2011), which model the SOA formation specifically without modifying the concentrations of ozone and major radicals. In 3-D modeling, implicit SOA mechanisms or parameterizations are usually added to implicit gas-phase mechanisms, conserving the oxidant chemistry of the implicit gas-phase mechanism.

Implicit SOA mechanisms are often established based on experimental data from smog chamber experiments to represent the formation and evolution of SOA, such as the two-product empirical SOA model (Odum et al., 1996) and the volatility basis set (VBS) that splits VOC oxidation products into a uniform set of volatility "bins" Donahue et al., 2006).

And the focus of our semi-explicit SOA mechanisms:
The generated semi-explicit mechanisms are designed to preserve the accuracy of explicit mechanisms for SOA formation, while keeping the number of reactions/species low enough to be suitable for large-scale modeling, particularly in 3-D AQMs. The focus of the semi-explicit mechanism is solely on the accurate modelling of SOA. Because ozone, major radicals, and other inorganics are also affected by inorganic and other VOC chemistry, their concentrations are not tracked with the semi-explicit mechanism. Instead, they are simulated using existing implicit gas-phase chemical mechanisms.

The computation of stoichiometric coefficients and rate constants for new lumped species (as shown in Figure 2) based on concentrations and lifetimes need to be discussed. The use of "average produced concentrations from five-day 0D simulations" needs more discussion than as a footnote in the table. Presumably the [HO2], [NO], etc. concentrations used to compute the lifetimes are also averaged? What are the 5 0D simulations they use to compute the average? The later discussion indicates they use 8 scenarios for initial training, and more than that when they are close to being finished.

To ensure coherency, all reduction parameters are calculated based on five-day 0-D simulation results of training conditions. For example, in the BCARY reduction, the lifetime of inorganics (e.g., HO2, NO) is calculated as the average of 16 simulations (two five-day simulations starting at 0h/12h for eight training conditions). Considering that inorganic concentrations repeat the diurnal profiles of 3-D simulations, the five-day average is equivalent to the daily average. This is different from the concentrations of sesquiterpene oxidation products, where a five-day average could take into account both formation and aging processes.

For clarity, the following explanation has been added to the paper:

As detailed in Table 3, \( f_w \) is computed as a function of chemical lifetime \( \tau \) following the computation of Seinfeld and Pandis (2016), and the reference concentrations \( C_r \) that are the arithmetic mean concentrations of a set from five-day 0D simulations of calculated from 0-D simulations using the explicit VOC mechanism under [HO2, NO]. Both \( \tau \) and \( C_r \) are based on averages of simulations across all training conditions.

Unless stated otherwise, a 5 day simulation is performed for each condition starting at midnight (0 h) and noon (12 h) for each condition, focusing on taking into account both the daytime and nighttime chemistry. All 0-D simulations are run for five days in order
to consider adequately SOA formation and aging processes.

RC: The maps indicate that the reduction errors are the greatest in Southern Europe than in the North. Is that because there is more secondary SOA predicted for the South, which should be more photochemically reactive? It might be useful to show a relationship between SOA level predicted and the reduction error for the various grid cells or scenarios.

AR: In order to illustrate the relationship between testing errors and SOA levels, we have added the SOA concentration map (Fig. 1) to the main paper, which corresponds to the results in the error map (Figure 3 of the paper):

![Error map showing geographic distribution of testing results](image1)

**Figure 1:** Error map that describes the geographic distribution of (a) error and (b) average SOA concentration of the testing results in July and August simulated using the "Rdc." mechanism. The total number of conditions displayed is 4,717 out of 12,159 that were tested. The results of all testing conditions are shown in Appendix C for reference.

We can see that SOA concentrations are generally higher under Nordic and North African conditions (The same initial BCARY concentration is set (5 µg/m³ for all conditions). As only one photolysis reaction is preserved in the "Rdc." mechanism (248 are preserved in the original MCM), the "Rdc." mechanism is probably not be very sensitive to photochemical reactivity. As can be seen from the bar plot in Fig. 5 in the paper, the testing errors in July and August over Southern Europe are actually close to 3 %, although they seem to be higher (placed in the error category between 3 % and 6 %) than others. In northern Africa, the high error might be partially attributed to the photochemical reactivity, but the main cause is still the high temperatures and dry climate.

It should also be noted that although some conditions on the map appear to have large errors (e.g., in northern Africa), the results may still be acceptable as long as they adhere to the pre-testing criteria (average error ≤ 3 %, and maximum error ≤ 20 %). As the average SOA concentration map of all testing results may be too complex to interpret, it is not included in the main paper but in Appendix C for reference (Fig. 2).

RC: On line 169 they state that "jumping" is restricted to cases where the intermediate to be removed results in the formation of only a single compound. However, this is not the case for the examples they show on Table 4, where each intermediate forms more than one compound. (Reactions 14 and 15 in Table 5 reflect merged competing reactions, and are not explicit.) In fact, any rapidly reacting intermediates that have
Figure 2: Error maps of (a) error and (b) average SOA concentration of the testing results simulated using the "Rdc." mechanism on all (i.e., 12 159) testing conditions.

Several reactions that form different products, can be "jumped" without affecting simulation results as long as their reactions are either unimolecular or with O2, so lifetimes would not vary with conditions. This is the case almost all of the alkoxy radicals in explicit atmospheric mechanisms. This should be pointed out in the discussion. The main approximation with "jumping" compounds with several reactions would that it would not deal with temperature effects if some of the competing reactions have different temperature dependences in the full model. Whether this is acceptable for their continental model could be determined by their algorithm.

AR: The jumping example shown in Table 4 of the paper (line 169) is indeed a mistake, as we mixed up the concept of jumping with a reduction process "pre-reduction" that are applied to the original explicit mechanism. We have added one section describing the pre-reduction (with an example of Table 1) to the main paper:

Section 2, line 103, before Section "Reduction strategies":

A pre-reduction process is conducted on the original MCM mechanism before it is used as the reference mechanism for the reduction. This process skips very fast reactions (i.e., reaction rate of $1.0 \times 10^6$ corresponding to a lifetime of 1 µs) to avoid numerical problems. For computational efficiency, it also combines elementary reactions with the same reactants into combined reactions with non-integer stoichiometric coefficients.

An example is shown in Table 1, where the original MCM reactions No. 1 to 7 have first been merged into the combined reactions No. 8 to 10. The pre-reduction compacts the reaction list (from 1 626 to 1 242 reactions), improving the reduction efficiency. The pre-reduction also skips two biradicals (i.e., BCALOOA and CH2OOF), that are extremely reactive and disintegrate instantaneously with a kinetic rate coefficient of $10^6$ s$^{-1}$. As a result, the reactions No. 8 to 10 can then be repented by one reaction No. 11, whose kinetic rate coefficient corresponds to that of the reaction producing the skipped species (in this case, the ozonolysis of BCAL, reaction No. 9).

We have also rephrased the jumping section with an new example (Table 2).
Table 1: Reactions before and after pre-reduction, where MCM species BCALOOA and CH2OOF are skipped over by their degradation products.

<table>
<thead>
<tr>
<th>No.</th>
<th>Reaction</th>
<th>Kinetic rate coefficient $^b$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$\text{BCAL} + \text{O}_3 \rightarrow \text{BCALOOA} + \text{HCHO}$</td>
<td>$1.1 \times 10^{-16} \times 0.670$</td>
</tr>
<tr>
<td>2</td>
<td>$\text{BCAL} + \text{O}_3 \rightarrow \text{BCLKET} + \text{CH2OOF}$</td>
<td>$1.1 \times 10^{-16} \times 0.330$</td>
</tr>
<tr>
<td>3</td>
<td>$\text{BCALOOA} \rightarrow \text{BCALOO}$</td>
<td>$1.0 \times 10^6 \times 0.500$</td>
</tr>
<tr>
<td>4</td>
<td>$\text{BCALOOA} \rightarrow \text{C146O2} + \text{OH}$</td>
<td>$1.0 \times 10^6 \times 0.500$</td>
</tr>
<tr>
<td>5</td>
<td>$\text{CH2OOF} \rightarrow \text{CH2OO}$</td>
<td>$1.0 \times 10^6 \times 0.370$</td>
</tr>
<tr>
<td>6</td>
<td>$\text{CH2OOF} \rightarrow \text{CO}$</td>
<td>$1.0 \times 10^6 \times 0.500$</td>
</tr>
<tr>
<td>7</td>
<td>$\text{CH2OOF} \rightarrow \text{HO}_2 + \text{CO} + \text{OH}$</td>
<td>$1.0 \times 10^6 \times 0.130$</td>
</tr>
<tr>
<td>8</td>
<td>$\text{BCAL} + \text{O}_3 \rightarrow 0.67 \text{BCALOOA} + 0.33 \text{BCLKET}$</td>
<td>$1.1 \times 10^{-16}$</td>
</tr>
<tr>
<td></td>
<td>+ $0.33 \text{CH2OOF} + 0.67 \text{HCHO}$</td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>$\text{BCALOOA} \rightarrow 0.5 \text{BCALOO} + 0.5 \text{C146O2} + 0.5 \text{OH}$</td>
<td>$1.0 \times 10^6$</td>
</tr>
<tr>
<td>10</td>
<td>$\text{CH2OOF} \rightarrow 0.37 \text{CH2OO} + 0.63 \text{CO} + 0.13 \text{HO}_2 + 0.13 \text{OH}$</td>
<td>$1.0 \times 10^6$</td>
</tr>
<tr>
<td>11</td>
<td>$\text{BCAL} + \text{O}_3 \rightarrow 0.5 \text{BCALOO} + 0.5 \text{C146O2} + 0.37 \text{CH2OO}$</td>
<td>$1.1 \times 10^{-16}$</td>
</tr>
<tr>
<td></td>
<td>+ $\text{BCLKET} + \text{HCHO} + 0.13 \text{HO}_2$</td>
<td></td>
</tr>
<tr>
<td></td>
<td>+ $0.63 \text{CO} + 1.1 \text{OH}$</td>
<td></td>
</tr>
</tbody>
</table>
As shown in Table 4, the ozonolysis of BCAL results in the formation of the excited Criegee biradical compounds BCALOOA and CH2OOF. These two biradicals are extremely reactive and disintegrate instantaneously with a kinetic rate coefficient of $10^{15} \text{ s}^{-1}$. As a result of the jumping strategy, three MCM reactions (reactions Nos. 13, 14, and 15 in Table 4) are reduced to one reaction R5, whose kinetic rate coefficient corresponds to that of the reaction producing the skipped species (in this case, the ozonolysis of BCAL, reaction No. 13).

As shown in Table 4, the alkoxy radicals BCALOO formed from ozonolysis of BCAL (reaction No. 11 in Table 1) is jumped over to its only destruction product BCLKET. Consequently, the reactions No. 12 to 16 are removed, and reaction No. 11 is updated to reaction No. 17. Currently, the jumping strategy is considered when the destruction of a single compound (to be jumped) results in the production of a single compound (jumping). The difference in carbon numbers between reduced species cannot exceed three in order to prevent significant differences in organic mass before and after jumping.

Table 2: Reactions before and after the jumping strategy, where MCM species BCALOO is jumped over by its degradation product BCLKET.

<table>
<thead>
<tr>
<th>No.</th>
<th>Reactions</th>
<th>Kinetic rate coefficient</th>
</tr>
</thead>
<tbody>
<tr>
<td>12</td>
<td>BCALOO + CO $\rightarrow$ BCLKET</td>
<td>$1.2 \times 10^{-15}$</td>
</tr>
<tr>
<td>13</td>
<td>BCALOO + NO $\rightarrow$ BCLKET + NO2</td>
<td>$1.0 \times 10^{-14}$</td>
</tr>
<tr>
<td>14</td>
<td>BCALOO + NO2 $\rightarrow$ BCLKET + NO3</td>
<td>$1.0 \times 10^{-15}$</td>
</tr>
<tr>
<td>15</td>
<td>BCALOO + SO2 $\rightarrow$ BCLKET + SO3</td>
<td>$7.0 \times 10^{-14}$</td>
</tr>
<tr>
<td>16</td>
<td>BCALOO $\rightarrow$ BCLKET + H2O2</td>
<td>$1.4 \times 10^{-17} \times [\text{H}_2\text{O}]$</td>
</tr>
<tr>
<td></td>
<td>BCAL + O3 $\rightarrow$ 1.5 BCLKET + 0.5 C146O2</td>
<td></td>
</tr>
<tr>
<td></td>
<td>$+$ 0.37 CH2OO + HCHO</td>
<td>$1.1 \times 10^{-16}$</td>
</tr>
<tr>
<td></td>
<td>$+$ 0.13 HO$_2$ + 0.63 CO + 1.13 OH</td>
<td></td>
</tr>
</tbody>
</table>

Currently, only jumping to a single compound was done, as doing jumping to multiple compounds may not affect significantly the reduction of BCARY-SOA mechanism. Furthermore, compounds involved in several reactions with different temperature effects cannot be jumped over if they lead to multiple compounds without neglecting the temperature effect. After removing some reactions or some species, some compounds can become a candidate for "single" jumping. At the end of the reduction, only two compounds remain that could be candidate for jumping to multiple compounds. However, these two compounds (C133O and BCBOO) compounds are involved in several reactions with different temperature effects and therefore cannot be jumped over.

**RC:** The example of lumping they present in Table 2 gives a rather trivial reduction example that should not really require an algorithm. The chemical lifetimes of the 3 peroxy radical species being lumped are the same for the major loss processes, with the only difference being the RO$_2$+RO$_2$ reactions that make almost no contribution (see Figure 2), so the "weighting ratios" should actually be very close the stoichiometric coefficients in the explicit mechanism (reaction "0" in Table 1), and the simulation results should also be the very close. An additional example where the lumping may not be as obvious, and which may potentially be more dependent
on environmental conditions, should also be given. This would aid the discussion of the lumping approach and give the reader a better understanding of the lumping approach and its general utility and potential drawbacks. Or is the kind of peroxy radical lumping illustrated in Table 2 the only type of lumping they actually did? If so, this should be stated.

AR: Thank you for the suggestion. Typically, lumping occurs between compounds sharing similar chemical pathways or molecular structures. While the example of lumping in Table 2 is trivial, it serves to illustrate how lumping merges reactions and build new surrogates.

Taking into account the reviewer’s comments, two additional examples from the BCARY reduction have been added to appendix:

In Appendix C:

Besides the example shown in Sect. 2.2.3, two additional examples have been added from the BCARY reduction: one illustrates the lumping of two similar compounds formed by different reactions, and the other illustrates the lumping of two more distinct compounds. The first example is the MCM species C1313NO3 and C152NO3 (Fig. 3). These two species come from different reactions. The molecular structures of both compounds are similar (they contain organic nitrates, aldehydes, and alcohols), but C152NO3 contains an additional carboxylic acid where C1313NO3 contains an aldehyde. The corresponding reactions before and after lumping are summarized in Table 3 where the new surrogate “mC1313NO3” is built from C1313NO3 with a weighting ratio of 83% and C152NO3 with a weighting ratio of 17%. As a result of this lumping, the average error increase under training conditions is 0.001 % (the tolerance is 0.01 %).

Another example of lumping is the MCM species BCALBOC and C1310OH (Fig. 4). Unlike the previous example, these two species are more distinct. According to MCM, BCALBOC are generated through \( \text{O}_3 \)-initiated reactions, while C1310OH are generated through high-generation oxidations. There is less similarity in the structures or chemical reactions of the two molecules. MCM contains the OH reaction of BCALBOC, and the \( \text{O}_3 \) and OH reactions of C1310OH. However, this reduction was accepted since lumping them only increased the average error by 0.01 % under training conditions (the tolerance was 1 %). The new surrogate ”mBCALBOC” is constructed from BCALBOC with a weighting ratio of 98 % and C1310OH with a weighting ratio of 2 %.

As C1310OH has a low weighting ratio, the lumping would be substituted by replacing (a special case of lumping), where the weighting ratio of BCALBOC is set to 100 % and of C1310OH is set to 0 %. In that case, instead of forming a new surrogate, C1310OH is replaced by BCALBOC. In BCARY reduction, this type of replacing was not used, but it can be activated by the user by setting the weighting ratio threshold.

RC: I don’t understand the difference between "removing species" and "removing elementary-like reactions". Strictly speaking, all species that are removed should be done by renormalizing yields from competing
Table 3: Reactions related to the reduction of MCM species C1313NO3 and C152NO3 via lumping.\(^a\)

<table>
<thead>
<tr>
<th>Reactions before lumping</th>
<th>Kinetic coefficient</th>
</tr>
</thead>
<tbody>
<tr>
<td>Production</td>
<td></td>
</tr>
<tr>
<td>C1313O2 + NO → C1313NO3</td>
<td>KRO2NO × 0.134</td>
</tr>
<tr>
<td>C152O2 + NO → C152NO3</td>
<td>KRO2NO × 0.136</td>
</tr>
<tr>
<td>Destruction</td>
<td></td>
</tr>
<tr>
<td>C1313NO3 + OH → C116CHO + HCHO + NO2</td>
<td>5.59 × 10(^{-11})</td>
</tr>
<tr>
<td>C152NO3 + OH → BCLKBOC + HCHO + NO2</td>
<td>1.58 × 10(^{-11})</td>
</tr>
<tr>
<td>Reactions after lumping</td>
<td></td>
</tr>
<tr>
<td>Production</td>
<td></td>
</tr>
<tr>
<td>C1313O2 + NO → 0.134 mC1313NO3 + 0.866 C1313O + 0.866 NO2</td>
<td>KRO2NO</td>
</tr>
<tr>
<td>C152O2 + NO → 0.136 mC1313NO3 + 0.864 C152O + 0.864 NO2</td>
<td>KRO2NO</td>
</tr>
<tr>
<td>Destruction</td>
<td></td>
</tr>
<tr>
<td>mC1313NO3 + OH → C116CHO + HCHO + NO2</td>
<td>5.59 × 10(^{-11}) × 0.82945</td>
</tr>
<tr>
<td>mC1313NO3 + OH → BCLKBOC + HCHO + NO2</td>
<td>5.59 × 10(^{-11}) × 0.17055</td>
</tr>
</tbody>
</table>

\(^a\) 0.82945 is the exact weighting ratio of C1313NO3 and 0.17055 is the exact weighting ratio of C152NO3.

Figure 4: Molecular structures of the MCM species BCALBOC (left) and C1310OH (right).
Table 4: Reactions related to the reduction of MCM species BCALBOC and C1310OH via lumping.\(^a\)

<table>
<thead>
<tr>
<th>Reactions before lumping</th>
<th>Kinetic coefficient</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Production</strong></td>
<td></td>
</tr>
<tr>
<td>BCOOA → BCALBOC</td>
<td>1.0 × 10^6 × 0.15</td>
</tr>
<tr>
<td>C1310O2 → C1310OH</td>
<td>2.5 × 10^{-13} × [RO2] × 0.2</td>
</tr>
<tr>
<td><strong>Destruction</strong></td>
<td></td>
</tr>
<tr>
<td>BCALBOC + O3 → BCBOOA + HCHO</td>
<td>1.1 × 10^{-16} × 0.670</td>
</tr>
<tr>
<td>BCALBOC + O3 → BCLKBOC + CH2OOF</td>
<td>1.1 × 10^{-16} × 0.330</td>
</tr>
<tr>
<td>BCALBOC + OH → C152O2</td>
<td>6.98 × 10^{-11}</td>
</tr>
<tr>
<td>C1310OH + OH → C1310CO + HO2</td>
<td>6.2 × 10^{-11}</td>
</tr>
<tr>
<td><strong>Reactions after lumping</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Production</strong></td>
<td></td>
</tr>
<tr>
<td>BCOOA → mBCALBOC</td>
<td>1.0 × 10^6 × 0.15</td>
</tr>
<tr>
<td>C1310O2 → mBCALBOC</td>
<td>2.5 × 10^{-13} × [RO2] × 0.2</td>
</tr>
<tr>
<td><strong>Destruction</strong></td>
<td></td>
</tr>
<tr>
<td>mBCALBOC + O3 → BCBOOA + HCHO</td>
<td>1.10 × 10^{-16} × 0.670 × 0.97675</td>
</tr>
<tr>
<td>mBCALBOC + O3 → BCLKBOC + CH2OOF</td>
<td>1.10 × 10^{-16} × 0.330 × 0.97675</td>
</tr>
<tr>
<td>mBCALBOC + OH → C1310CO + HO2</td>
<td>6.20 × 10^{-11} × 0.97675</td>
</tr>
<tr>
<td>mBCALBOC + OH → C152O2</td>
<td>6.98 × 10^{-11} × 0.023251</td>
</tr>
</tbody>
</table>

\(^a\) 0.97675 is the exact weighting ratio of BCALBOC and 0.023251 is the exact weighting ratio of C1310OH.
pathways if the reactions are lumped, which is the same as converting them to "elementary-like reactions", and then removing the less important reactions. Without renormalization of the other branching ratios, the removal would cause an unnecessary loss of mass or, worse, an artificial radical sink. An artificial radical sink may not affect SOA predictions if the concentrations of the inorganics are constrained in the test simulations (and thus would pass the tests in the reduction algorithm discussed here if that were the case), but the reduced mechanism would be unsuitable for predicting radical levels if used in a full model. Even small artificial radical sinks may have non-negligible effects on the radical chain reactions involved in many photooxidation mechanisms.

AR: It is true that the reduction with the current removing strategies may have an impact on the formation of inorganic radicals. However, as indicated in several replies previously, we develop GENOA primarily for SOA concentrations and not for the concentrations of inorganic radicals. Since inorganic radical formation and destruction are still simulated with implicit gas-phase chemical mechanisms, the removing strategies used in BCARY reduction are acceptable, as long as SOA concentrations are reproduced.

RC: Section 2.2.3 describes the 12,159 conditions used for the testing datasets. It should be stated in that section that Figure D1 has a map of these locations. What are the 4,717 conditions mapped on Figure 6?

AR: These 4,717 conditions (out of the total 12,159 testing conditions) are the testing conditions during the summer period from July to August. As the map of all testing conditions is difficult to read, we present only the testing results of the two months in the main paper. The testing results of all conditions can be found in the Appendix B for reference.

The locations of testing conditions have also been added in the main paper:

Section 2.2.3, line 253:

Fig. B1 indicates the locations the testing dataset as well as the testing results for BCARY reduction.

RC: Are any of the lumping criteria listed around Lines 134-143 options that can be varied by the user? Define "extremely short lifetimes". Do they allow lumping compounds that react with different atmospheric species or types of reaction (e.g., with OH, O₃, NO, etc, by photolysis, or unimolecular), but happen to have comparable lifetimes for most scenarios?

AR: Yes, the lumping criteria listed around lines 134-143 can be modified by the user. We have summarized the user-chosen criteria into Table 5, which will be available in an Excel file in the supplementary material.

For intermediate compounds, such as biradicals (e.g., MCM species BCAOO), which degrade rapidly to closed shell molecules, jumping is considered to be more appropriate than lumping. In BCARY reduction, we checked the species type and ruled biradical them out of lumping as they have a low lifetime (approximately a few microseconds). In the paper, the term "extremely short lifetimes" is used to indicate that intermediate compounds are better suited for jumping since they have a short lifetime.
### Table 5: The user-chosen reduction options and parameters applied to BCARY reduction.

<table>
<thead>
<tr>
<th>Category</th>
<th>Meaning (and notion if mentioned in the paper)</th>
<th>Threshold</th>
</tr>
</thead>
<tbody>
<tr>
<td>reduction evaluation</td>
<td>$\epsilon_{\text{ref}}$, the maximum error compared to the reference mechanism under training conditions</td>
<td>$\leq 1%$ to $10%$</td>
</tr>
<tr>
<td>reduction evaluation</td>
<td>$\epsilon_{\text{pre}}$, the maximum error compared to the previous validated mechanism under training conditions</td>
<td>$\leq 1%$ to $10%$</td>
</tr>
<tr>
<td>reduction evaluation</td>
<td>$\epsilon_{\text{ave}\text{-pre-testing}}$, the maximum error compared to the reference mechanism under pre-testing conditions</td>
<td>$\leq 3%$</td>
</tr>
<tr>
<td>reduction evaluation</td>
<td>$\epsilon_{\text{max}\text{-pre-testing}}$, the maximum error compared to the reference mechanism under pre-testing conditions</td>
<td>$\leq 25%$</td>
</tr>
<tr>
<td>reduction strategy</td>
<td>reduction error calculation method</td>
<td>see paper</td>
</tr>
<tr>
<td>removal strategy</td>
<td>searching order in species/reaction lists</td>
<td>high-to-low generation</td>
</tr>
<tr>
<td>removing reactions</td>
<td>$B_{\text{rm}}$, Maximum hourly branching ratio of the reaction</td>
<td>$\leq 5%$ to $100%$</td>
</tr>
<tr>
<td>jumping</td>
<td>carbon number difference</td>
<td>$\leq 3$</td>
</tr>
<tr>
<td>jumping</td>
<td>minimum $P_{\text{sat}}$ of species considered to be jumped</td>
<td>$\geq 1.0 \times 10^{-6}$ atm</td>
</tr>
<tr>
<td>lumping</td>
<td>order of species list: find lumping for condensables</td>
<td>order by $P_{\text{sat}}$</td>
</tr>
<tr>
<td>lumping</td>
<td>weighting ratio $f_{\text{w}}$, calculation method</td>
<td>see paper</td>
</tr>
<tr>
<td>lumping</td>
<td>functional group check I: restrict to species with same groups</td>
<td>7 Types, see paper</td>
</tr>
<tr>
<td>lumping</td>
<td>functional group check II: do not participate lumping</td>
<td>ROO species</td>
</tr>
<tr>
<td>lumping</td>
<td>mass difference</td>
<td>$\leq 100$ g mol$^{-1}$</td>
</tr>
<tr>
<td>lumping</td>
<td>lifetime difference</td>
<td>$\leq 10$-fold</td>
</tr>
<tr>
<td>lumping</td>
<td>carbon number difference</td>
<td>$\leq 2$</td>
</tr>
<tr>
<td>lumping</td>
<td>weighting ratio threshold by which a species is considered to be replaced by another</td>
<td>$\leq 0%$</td>
</tr>
<tr>
<td>replacing</td>
<td>Minimum mass of species participating in replacing</td>
<td>$&gt; 100$</td>
</tr>
<tr>
<td>replacing</td>
<td>carbon number difference</td>
<td>$\leq 3$</td>
</tr>
<tr>
<td>Aerosol-oriented treatment</td>
<td>number of condensables to activate aerosol-oriented treatments</td>
<td>$\leq 20$</td>
</tr>
<tr>
<td>0-D simulation</td>
<td>simulation starting times</td>
<td>0 h &amp; 12 h</td>
</tr>
</tbody>
</table>
The term "extremely short lifetimes" has been rephrased to avoid confusion:

Section 2.1.1, line 141:

- No lumping for intermediate compounds with extremely short lifetimes (e.g., Lumping is not considered for biradicals (ROO)), as they are candidate for jumping instead that degrade rapidly into closed shell molecules, as jumping is considered to be more appropriate for these compounds.

RC: *It probably would be appropriate to point out that the MCM mechanism they use as the starting point has already been reduced to some extent. It already "jumps" carboncentered radicals by replacing them by the peroxy radical formed when they react. It also combines elementary reactions with the same reactants into combined reactions with noninteger stoichiometric coefficients. Removing species from such reactions should not be done without renormalizing coefficients representing the competing processes, as discussed above.*

AR: As previously explained, the rationale behind the "removing species" strategy is justified by the fact that we do not intend to track the destruction or production of inorganic radicals. The section entitled "Pre-reduction" has been added to the main paper (in section 2, line 103, as detailed previously), which describes the treatments on the original MCM mechanism before training.

RC: *Figure 2 and Table A1 use "RRNO3" to indicate the fraction of peroxy radicals reacting with NO3. The term "RNO3" should be used instead, to be consistent with the terminology for the other peroxy loss processes. I found this confusing when I first saw this.*

AR: The term "$R_{RNO3}$" was adopted, because the other term "$R_{NO3}$" had already been used for the reacting ratio of NO3 with SOA precursor. To avoid confusion, we have changed "$R_{RNO3}$" to "$R_{NO3–RO2}$", along with other reacting ratios showing the reactivity with peroxy radicals (RO2):

Section 2.2.1, line 222:

- The reacting ratios of RO2 species with NO (R$^{NO2←NO}$), HO2 (R$^{HO2←HO2}$), NO3 (R$^{NO3←SO4}$), and other RO2 species (R$^{RO2←RO2}$), whose sum equals 1, indicate the relative reactivity of successive reactions with RO2 species.

3. Suggestions

RC: *It would be better if the four items in the reduction strategy were discussed in the order they are actually applied in the algorithm. "Lumping" and "replacing" is discussed first, but actually "removing" and "jumping" is done first, which is appropriate.*

AR: Thank you for the helpful suggestion. Reduction strategies have been reordered in section 2.1 as follows: "removing", "jumping", "lumping" and "replacing".

RC: *It would be useful to include a table of options or parameters that control the reduction process, indicating which can be changed by the user, and the value(s) of the parameters used in the examples in this work. This would be particularly relevant if the intention is to make this software available for use by other researchers.*

AR: Thank you for the suggestion. A table of reduction options and parameters applied to BCARY reduction has been added to an Excel file (Table 5), as part of the supplementary material.
RC: The geographical and meteorological conditions for the 80D scenarios used for the training dataset are summarized on Table 5, and a map of their locations is shown on Figure C1 in an Appendix. The map showing the locations is useful and should be in the main paper, and if the points are shown along with the “errors” on Figure 6, it would take no added journal space and also indicate whether there is an relationship between the errors and the locations of the training scenarios. It also would be helpful for Table 5 to have columns showing the amounts of SOA calculated, the major daytime loss process for b-caryophyllene, and at least the day and night fractions that peroxy radicals react with NO. These are the most important chemical differences in the scenarios, and including this on the table would give this information in a more compact and perhaps clearer manner than on Figure 2. Figure 2 is also useful, but it is difficult to read and could be formatted better.

AR: Thank you for the helpful suggestions. We have put together the locations of the training and pre-testing datasets in one figure (Fig. 5), and have attached it to the main paper (The figure number corresponds to the new numbering in the revised paper):

Section 2.2.2, line 243:
The locations of the training and pre-testing conditions are presented in the Fig. 3.

Figure 5: Locations of the training (see legends) and pre-testing (white scattered dots) datasets used in the BCARY reduction.

We have also added two columns to the training dataset information table (Table 5 of the paper), including average simulated SOA concentrations and average NO reactivity ratios with proxy radicals ($R_{RO2-NO}$, indicating the chemical regimes (high/low NOx)). Considering that figure 2 of the paper may be sufficient to illustrate the major daytime loss process for BCARY (the reactivity of BCARY with O3, OH, or NO3 at 12h shown in the left panel), we decided to not repeat this information in the table. On the right half panel of Figure 2, the $R_{RO2-NO}$ ratio at 0h and 12h indicates the fractions of NO that react with peroxy radicals at day and at night, respectively. Since they are consistent with the day and night fractions that peroxy radicals react with NO, the additional information is not added to the table (daily average NO ratio would be enough). For clarity, the digits in Table 5 of the paper have also been changed for temperature and relative humidity.

Meanwhile, Table 5 of the paper has been revised as follows (Table 6):
### Table 6: Geographic and meteorological conditions of the training dataset

<table>
<thead>
<tr>
<th>Condition Name</th>
<th>Lat °N</th>
<th>Lon °E</th>
<th>Time</th>
<th>TEMP</th>
<th>RH %</th>
<th>R_{NO} b</th>
<th>SOA µg/m³</th>
</tr>
</thead>
<tbody>
<tr>
<td>OH NO</td>
<td>36.0</td>
<td>15.4</td>
<td>Jul.</td>
<td>299</td>
<td>679</td>
<td>60</td>
<td>4.1</td>
</tr>
<tr>
<td>OH HO₂</td>
<td>32.0</td>
<td>-9.4</td>
<td>Jul.</td>
<td>295</td>
<td>777</td>
<td>20</td>
<td>6.1</td>
</tr>
<tr>
<td>NO₃ NO</td>
<td>40.25</td>
<td>-3.4</td>
<td>Jul.</td>
<td>302</td>
<td>928</td>
<td>69</td>
<td>4.4</td>
</tr>
<tr>
<td>NO₃ HO₂</td>
<td>32.0</td>
<td>36.6</td>
<td>Aug.</td>
<td>302</td>
<td>738</td>
<td>29</td>
<td>5.7</td>
</tr>
<tr>
<td>O₃ NO</td>
<td>69.0</td>
<td>33.8</td>
<td>Jan.</td>
<td>260</td>
<td>284</td>
<td>99</td>
<td>5.2</td>
</tr>
<tr>
<td>O₃ HO₂</td>
<td>68.0</td>
<td>18.2</td>
<td>Dec.</td>
<td>265</td>
<td>789</td>
<td>25</td>
<td>4.6</td>
</tr>
<tr>
<td>ADD1</td>
<td>41.5</td>
<td>-14.2</td>
<td>Dec.</td>
<td>288</td>
<td>876</td>
<td>20</td>
<td>5.5</td>
</tr>
<tr>
<td>ADD2</td>
<td>45.75</td>
<td>9.0</td>
<td>Dec.</td>
<td>279</td>
<td>858</td>
<td>100</td>
<td>4.4</td>
</tr>
</tbody>
</table>

a from left to right: name, latitude, longitude, time period, average temperature, average relatively humidity, daily average NO reacting ratio, simulated total SOA concentration of the training conditions.

b the daily average NO reacting ratio is calculated out of the RO₂ reactivity of NO, HO₂, NO₃, and RO₂. Conditions with high R_{NO} ratio are considered as in high NOₓ regime.

c the initial concentration of BCARY is 5 µg/m³.

**RC:** It would also be useful to include a map showing locations where the 150 testing scenarios were derived. This may cause too much clutter if added to Figure 6, but should at least be in the Appendix. There should also be a map showing the SOA levels calculated by the CHIMERE model for the European domain. Perhaps it could be combined with the figure in the Appendix showing the locations of the testing scenarios, though it may be useful enough to be in the main text.

**AR:** Thank you for the suggestions. As explained in one previous reply, we have combined the locations of the pre-testing dataset and the training dataset in one figure to the main paper. Meanwhile, the maps showing the average SOA concentrations of the testing conditions in summer (July and August) and all times (Fig. 1 and Fig. 2) have been added to the main paper and to Appendix B, respectively.

**RC:** Figure 4 would be more useful if it included some brief indication of what kinds of compounds the model species refer to.

**AR:** Thank you for the suggestion. We have added Figure 6 to Appendix C, which includes the molecular structures of MCM species kept in the final reduced mechanism. Since merged species (names starting with "m", nine species in total) contain complex functional groups and can be surrogates for multiple MCM compounds, they are difficult to classify. Therefore, we have added Table 7 to Appendix B, which lists the names of the "Rdc." species with corresponding lumped species in original MCM. Particularly for condensables, an Excel file (Fig. 7) has also been added to the supplementary materials, which contains the molecular structures of "Rdc." condensables in UNIFAC structural group format.
Figure 6: The molecular structures of the MCM species that are mentioned in this paper.
<table>
<thead>
<tr>
<th>Name of the UNIFAC groups</th>
<th>Index</th>
<th>NBCOOH</th>
<th>CI41CO2H</th>
<th>BCKSOZ</th>
<th>C133CO</th>
<th>C132OOH</th>
<th>C131PAN</th>
</tr>
</thead>
<tbody>
<tr>
<td>alkane CH3</td>
<td>0</td>
<td>3.00E+00</td>
<td>1.89E+00</td>
<td>2.90E+00</td>
<td>8.20E-01</td>
<td>2.97E-02</td>
<td>2.00E+00</td>
</tr>
<tr>
<td>alkane CH2</td>
<td>1</td>
<td>5.00E+00</td>
<td>4.69E+00</td>
<td>3.10E+00</td>
<td>1.64E+00</td>
<td>5.73E-02</td>
<td>4.00E+00</td>
</tr>
<tr>
<td>alkane CH</td>
<td>2</td>
<td>2.00E+00</td>
<td>1.89E+00</td>
<td>2.00E+00</td>
<td>8.19E-01</td>
<td>1.70E-02</td>
<td>2.00E+00</td>
</tr>
<tr>
<td>alkane C</td>
<td>3</td>
<td>1.00E+00</td>
<td>9.45E-01</td>
<td>9.99E-01</td>
<td>4.10E-01</td>
<td>1.49E-02</td>
<td>1.00E+00</td>
</tr>
<tr>
<td>methanol CH2OH</td>
<td>5</td>
<td>5.63E-02</td>
<td>5.24E-04</td>
<td>5.85E-01</td>
<td>9.85E-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>methanol CH2OH</td>
<td>6</td>
<td>6.99E-04</td>
<td>9.27E-07</td>
<td>3.09E-04</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>methanol COH</td>
<td>7</td>
<td>1.42E-04</td>
<td>5.30E-04</td>
<td>6.96E-05</td>
<td>3.02E-04</td>
<td></td>
<td></td>
</tr>
<tr>
<td>calcohol between two alcohols OHCH2OH</td>
<td>9</td>
<td>4.09E-04</td>
<td>2.04E-05</td>
<td>1.22E-04</td>
<td>1.21E-03</td>
<td></td>
<td></td>
</tr>
<tr>
<td>calcohol between two alcohols OHCHOH</td>
<td>10</td>
<td>2.72E-04</td>
<td>1.30E-05</td>
<td>1.22E-04</td>
<td>6.05E-04</td>
<td></td>
<td></td>
</tr>
<tr>
<td>calcohol between two alcohols OHCOH</td>
<td>11</td>
<td>6.99E-04</td>
<td>6.48E-06</td>
<td>6.10E-05</td>
<td>3.02E-04</td>
<td></td>
<td></td>
</tr>
<tr>
<td>calcohol in tails of alcohol OHCHOH</td>
<td>12</td>
<td>1.11E-01</td>
<td>1.34E-03</td>
<td>1.18E+00</td>
<td>2.27E+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>calcohol in tails of alcohol OHCHOH</td>
<td>13</td>
<td>1.11E-01</td>
<td>2.34E-03</td>
<td>6.06E-01</td>
<td>3.47E+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>calcohol in tails of alcohol OHCOH</td>
<td>14</td>
<td>1.05E-05</td>
<td>1.05E-03</td>
<td>1.00E-02</td>
<td>1.32E+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>calcohol in tails of alcohol OHCOH</td>
<td>15</td>
<td>5.47E-02</td>
<td>5.24E-04</td>
<td>5.90E-01</td>
<td>9.85E-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>alkene CH2=C</td>
<td>18</td>
<td>1.00E+00</td>
<td>1.00E+00</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>alkene CH=O</td>
<td>19</td>
<td>5.27E-06</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>alcohol OH</td>
<td>26</td>
<td>5.61E-02</td>
<td>1.05E-03</td>
<td>5.90E-01</td>
<td>9.85E-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ketone CH3CHO</td>
<td>29</td>
<td>9.92E-01</td>
<td>1.01E-01</td>
<td>9.95E-01</td>
<td>6.96E-01</td>
<td>1.00E+00</td>
<td></td>
</tr>
<tr>
<td>ketone CH2CHO</td>
<td>30</td>
<td>1.24E-01</td>
<td>9.99E-01</td>
<td>2.17E+00</td>
<td>6.69E-01</td>
<td>1.00E+00</td>
<td></td>
</tr>
<tr>
<td>aldehyde CHO</td>
<td>31</td>
<td>1.42E-02</td>
<td>1.01E-01</td>
<td>5.42E-03</td>
<td>1.88E-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ester CH3CCOO</td>
<td>32</td>
<td>5.30E-03</td>
<td>3.75E-03</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ether CHO</td>
<td>36</td>
<td>8.99E-01</td>
<td>2.96E-01</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>carboxylic acid COOH</td>
<td>37</td>
<td>9.39E-01</td>
<td>4.10E-01</td>
<td>1.85E-02</td>
<td></td>
<td></td>
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<tr>
<td>nitrate CHONO2</td>
<td>40</td>
<td>1.00E+00</td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td>hydroperoxycarboxyl CH2OH-CH</td>
<td>42</td>
<td>2.59E-04</td>
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<tr>
<td>hydroxyperoxycarboxyl CH-CH</td>
<td>43</td>
<td>1.68E-01</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>hydroxyperoxycarboxyl CO-CH</td>
<td>44</td>
<td>1.00E+00</td>
<td>8.32E-01</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>hydroxyperoxycarboxyl CH-OC</td>
<td>52</td>
<td>8.99E-01</td>
<td>2.96E-01</td>
<td></td>
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<tr>
<td>peroxyacyl nitrate PAN</td>
<td>54</td>
<td>1.00E+00</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Figure 7: UNIFAC functional group decomposition for the condensable species in the "Rdc." mechanism.
Table 7: The new surrogates in the "Rdc." mechanism and the corresponding lumped species in the original MCM mechanism. Noted that the "Rdc." surrogates may also go through other reductions (e.g., jumping) that does not affect their structure.

<table>
<thead>
<tr>
<th>Rdc. surrogate</th>
<th>lumped MCM species</th>
</tr>
</thead>
<tbody>
<tr>
<td>mBCSOZ</td>
<td>BCSOZ, BCAL, BCKET</td>
</tr>
<tr>
<td>mC141CO2H</td>
<td>C141CO2H, C143CO, C1310CO, BCALCCO, C143OH, BCCOH, BCAOH</td>
</tr>
<tr>
<td>mBCALO2</td>
<td>BCALO2, C146O2, C142O2, BCKAO2, C147O2</td>
</tr>
<tr>
<td>mBCKSOZ</td>
<td>BCKSOZ, BCLKET, BCALOH, BCKBCO, BCKAOH, BCSOZOH</td>
</tr>
<tr>
<td>mC131CO3</td>
<td>C131CO3, C141CO3, C1211CO3, C137CO3</td>
</tr>
<tr>
<td>mC131O2</td>
<td>C131O2, C144O2, C143O2, BCLKAO2, C152O2, BCLKCO2</td>
</tr>
<tr>
<td>mC132OOH</td>
<td>C132OOH, BCSOZOOH, C133OOH, C146OOH, C147OOH, C1313OOH, BCLKBOOH, BCLKAOOH, C152OOH, C145OOH, C148OOH, C144OOH, BCALOOH, BCKBOOH, C151OOH</td>
</tr>
<tr>
<td>mC133O2</td>
<td>C133O2, C1313O2</td>
</tr>
<tr>
<td>mC133CO</td>
<td>C133CO, C131CO2H, C148CO, C145OH, C1313OH, BCLKBOH, BCLKAOH, C152OH, C151OH, C147OH, BCLKACO, C148OH, C1211CO2H</td>
</tr>
</tbody>
</table>

References


Authors’ Response to Reviews of
GENerator of reduced Organic Aerosol mechanism (GENOA v1.0): An automatic generation tool of semi-explicit mechanisms
Zhizhao Wang, Florian Couvidat, Karine Sartelet

Reviewer # 2

RC: The paper discusses the development of a semi-explicit reduced organic aerosol mechanism for sesquiterpenes (Bcary), with the aim for employing it in air quality and large scale models. GENOA mechanism is based on the widely used near-explicit Master chemical mechanism (MCM). The mechanism used different strategies namely, lumping, replacing, jumping an removing to reduce the MCM scheme. The reduction procedure is tested under various environmental conditions (RH, temp etc.), resulting in a final reduced mechanism (Rdc.) suitable to simulate SOA. The simulated SOA using Rdc has low average error when compared to the near-explicit MCM scheme. This is a well thought out work, with suitable implications to better reproduce SOA in large scale and air quality models. I would therefore, recommend the publication of this work after the authors have answered the following questions:

AR: We would like to thank reviewer # 2 for the positive comments and constructive suggestions, which are much useful to improve the manuscript. We have carefully considered all of these comments and revised the manuscript accordingly. Please note that the line and section numbers mentioned in the response correspond to the version of the manuscript before revision.

1. General comments

RC: The main question is why did the authors chose sesquiterpenes? Why not isoprene or monoterpenes? The motivation to use sesquiterpenes should be highlighted.

AR: Thank you for the questions. Sesquiterpene was selected because it is a well-known source of SOA, and its formation mechanism is well documented. Sesquiterpene is also an ideal candidate for model development and demonstration of the reduction methodology. The oxidation products of sesquiterpene are less volatile and tend to condense more readily than those of lighter molecules such as monoterpen and isoprene.

We have added the explanation to the main paper:

Section 1, line 72:

The application of GENOA to the MCM degradation scheme of β-caryophyllene (BCARY) (Jenkin et al., 2012) is described in Sect. 3. The β-caryophyllene species is selected because it widely serves as a benchmark for modeling the tropospheric chemistry and SOA formation relevant to...
selected for investigation and demonstration of the GENOA algorithm, because it is one of the most abundant and representative sesquiterpenes (SQT) (e.g., Li et al., 2015; Xavier et al., 2019). and its degradation has been evaluated in chamber simulations (Jenkin et al., 2012). The BCARY scheme in the STOCHEM-CRI mechanism (Khan et al., 2017) and. Sesquiterpenes are a well-known source of SOAs (Hellén et al., 2020; Tasoglou and Pandis, 2015 and their degradation mechanisms (as BCARY) is well documented in the experiment data of Tasoglou and Pandis (2015) are also used for evaluation for near-explicit MCM mechanism (Jenkin et al., 2012). Studies have also compared BCARY SOA yields simulated using the MCM mechanism to chamber data (e.g., Xavier et al., 2019). BCARY is therefore an ideal candidate for model development and demonstration of the reduction methodology. In this paper, the near-explicit MCM BCARY degradation scheme serve as a reliable benchmark for GENOA. The experiment data from Tasoglou and Pandis (2015); Chen et al. (2012) are also compared to the newly developed reduced mechanism in Appendix A.

RC: Since GENOA is a semi-explicit mechanism, can it be used with any box or air quality model?

AR: Yes. The semi-explicit SOA mechanisms generated by GENOA can be added to the implicit gas-phase mechanism to model SOA in any box or air quality model. In the case of the sesquiterpene SOA, the "Rdc." mechanism adds 14 organic species including six condensables to SOA models, which is computationally feasible even for global 3-D modeling.

Generally, implicit SOA mechanisms applied to 3-D models are often based on the surrogate approach (e.g., Odum’s two-product) or the Volatility Basis Set. They are added to implicit gas-phase mechanisms without altering the pathways of oxidants. The same approach is adopted for the semi-explicit SOA mechanisms. The reduced SOA mechanism includes gas-phase chemical mechanism and aerosol properties for condensable species, which can be added to the implicit gas-phase mechanism in air quality models, without altering the pathways of ozone and major radicals.

For example, the 3-D CHIMERE model can simulate SOA using the implicit gas-phase mechanism MELCHIOR2 and SOA mechanism H2O [Couvidat et al., 2018]. With the reduced SOA mechanisms, SOAs in CHIMERE are simulated by MELCHIOR2, the reduced SOA mechanisms generated by GENOA, and H2O for other SOA precursors not covered by our mechanisms.

In the paper, We have added the explanation of how the coupling is generally resolved between VOC species and oxidants in 3-D models:

Section 1, line 36:

Along with To complete implicit gas-phase mechanisms, implicit SOA mechanisms have been developed (Kim et al., 2011), which model the SOA formation specifically without modifying the concentrations of ozone and major radicals. In 3-D modeling, implicit SOA mechanisms or parameterizations are usually added to implicit gas-phase mechanisms, conserving the oxidant chemistry of the implicit gas-phase mechanism.

Implicit SOA mechanisms are often established based on experimental data from smog chamber experiments to represent the formation and evolution of SOA, such as the two-product empirical SOA model (Odum et al., 1996) and the volatility basis set (VBS) that splits VOC oxidation products into a uniform set of volatility "bins" Donahue et al., 2006).

We have also added the following statements about our semi-explicit mechanisms:
Section 2, line 80:

The generated semi-explicit mechanisms are designed to preserve the accuracy of explicit mechanisms for SOA formation, while keeping the number of reactions/species low enough to be suitable for large-scale modeling, particularly in 3-D AQMs. The focus of the semi-explicit mechanism is solely on the accurate modelling of SOA. Because ozone, major radicals, and other inorganics are also affected by inorganic and other VOC chemistry, their concentrations are not tracked with the semi-explicit mechanism. Instead, they are simulated using existing implicit gas-phase chemical mechanisms.

RC: Although comparison has been made against MCM, the performance of a model can be made by comparing it against existing experimental SOA yields. There has been quite a lot of published experimental Bcary SOA yield experiments. I would suggest the authors to discuss GENOA derived SOA yields in comparison to these experiments.

AR: As GENOA is used to reduce the MCM mechanism, we compared the mechanisms before and after the reduction to ensure that the generated SOA mechanism preserves the performance of the explicit mechanism on SOA formation.

We have also compared SOA yields simulated by MCM and "Rdc." mechanisms to the experimental data from [Tasoglou and Pandis, 2015] and [Chen et al., 2012] in Appendix A. As shown in Fig. A1 of Appendix A, the results of the "Rdc." mechanism (noted as "Rdc.") are in good agreement with the experimental data (noted as "Tasoglou," and "chen.") and the results of the MCM mechanism ("v1b2"). Moreover, as now mentioned in the paper, [Xavier et al., 2019] has already performed some evaluations of BCARY SOA from the MCM mechanism against to chamber data.

RC: Is GENOA a carbon number conserving mechanism. Its is not clear from the manuscript if the mechanism is carbon conserving or not? If it is not then how do the authors justify it?

AR: Strictly speaking, the carbon number is not conserved explicitly during the reduction process. However, GENOA is designed to provide a good estimation of the contribution of the different functional groups. Carbon number is constrained in reduction by lumping (the difference in the carbon number between lumped species cannot exceed 2), jumping and replacing (the difference cannot exceed 3). For lumping and replacing, there is also a restriction on the total mass (< 100 µg/m³). For removing, there are no restrictions on either mass or carbon number.

Consequently, as we always constrain the total SOA concentration with strict error criteria (i.e., $\epsilon_{ref}$ and $\epsilon_{pre}$), the OM/OC, H/C, and O/C ratios are well reproduced (see Fig. 9 in the paper), so the carbon number should be also be well reproduced. If necessary, a specific restriction on conserving the carbon number can be added by the user in the reduction.

2. Specific questions

RC: L13-15: Motivation -> Although the health and climate effects of aerosols are introduced in every paper, the authors should maybe consider to explain these in a few words or a sentence. Also, it would be nice to explain why there is a need to improve the SOA representation in AQMs.

AR: Thank you for the suggestion. We have rephrased the motivation as follows:
Atmospheric aerosols have garnered considerable attention due to their undesirable effects on human health (Breyssse et al., 2013) and climate (Seinfeld et al., 2016; McNeill, 2017), and as such, they must be well represented in models of atmospheric chemistry, climate, and human health: they change the earth’s radiation balance and cloud formation (Ramanathan et al., 2001; McNeill, 2017); they trigger a wide variety of acute and chronic diseases (Breyssse et al., 2013). Because the effects of aerosols on health depend on their size and composition (Schwarze et al., 2006), adequate representations of the aerosol composition, mass, and number concentrations are required in air quality models (AQMs).

Besides being directly emitted, aerosols can be secondary, i.e., formed in the atmosphere through chemical reactions and gas-particle mass transfer as secondary aerosols. Secondary-Based on the chemical composition, they can be further divided into secondary inorganic aerosol (SIA) and secondary organic aerosol (SOA). SOA, which represents a significant fraction of aerosols (e.g., Gelencsér et al., 2007; Hallquist et al., 2009), and, is largely formed by the condensation of the oxidation products from the degradation of volatile organic compounds (VOCs) (Kanakidou et al., 2005). As SOA formation involves multiple processes such as the emission of SOA precursor gases, VOC gas-phase chemistry, gas-to-particle partitioning (Kanakidou et al., 2005; Hallquist et al., 2009), there are great complexity and uncertainty to accurately predict SOA formation with the simplified representations currently used in air quality models (Porter et al., 2021).

Section 1, line 18:

The chemistry of VOC degradation is comprehensively described in explicit atmospheric models. The state of knowledge on VOC chemistry can be reflected by explicit chemical mechanisms, such as which contain all known essential reaction pathways in VOC degradation. For instance, Jenkin et al. (1997); Saunders et al. (2003) developed the near-explicit Master Chemical Mechanism (MCM) (Jenkin et al., 1997; Saunders et al., 2003), which details which describes detailed gas-phase chemical processes related to VOC oxidation, or. Another example is the Generator for Explicit Chemistry and Kinetics of Organics in the Atmosphere (GECKO-A) (Aumont et al., 2005), which uses a prescribed protocol to assign complete reactions pathways and kinetic data to the degradation of VOCs. Explicit mechanisms represent the state of knowledge current understanding of atmospheric chemistry, including information about reaction pathways, kinetics data, and chemical structures (which may be used to deduce thermodynamic properties based on structure-activity relationships).

Another example is the Generator for Explicit Chemistry and Kinetics of Organics in the Atmosphere (GECKO-A) (Aumont et al., 2005), which uses a prescribed protocol to assign complete reaction pathways and kinetic data to the degradation of VOCs. Explicit mechanisms represent the state of knowledge current understanding of atmospheric chemistry, including information about reaction pathways, kinetics data, and chemical structures (which may be used to deduce thermodynamic properties based on structure-activity relationships).
Explicit mechanisms are mainly used in the box model for solving the multigeneration oxidation of multitudinous VOCs (e.g., Yuan et al., 2018). While the near explicit MCM mechanism has been applied to three-dimensional models (e.g., Ying and Li, 2011; Li et al., 2015), explicit mechanisms are generally too computationally intensive to be used in 3-D air quality models because of the large number of species involved. Hence, lumped mechanisms are used in 3-D air quality model because of the large number of species involved..


AR: Changed.

RC: L34-35: Are all the above mechanisms (lumped, CB05, MCM, GECKO-A) developed primarily for ozone simulation? Also it would be good to give examples of a few model species.

AR: Because grand-level ozone is one of the most important air pollutants, most gas-phase chemical mechanisms are accompanied by reactions with small changes or in favor of other reductions. These mechanisms were primarily developed for ozone simulation and reflect only the most significant chemical phenomena occurring in the atmosphere, where VOCs are represented by Implicit gas-phase mechanisms were developed and validated to simulate the concentrations of oxidants and other conventional air pollutants such as ozone and NO2. In these mechanisms, VOCs have been grouped into a limited number of model species because of computational considerations, and the SOA formation is usually not considered.

RC: L55: “suitable to”

AR: Revised.

RC: L112-113: Why is this order used in the reduction strategy for BCARY? How would any other order influence the reduction strategy?

AR: This reduction order was the most effective among all the reduction orders we tested. As each validated reduction can affect the subsequent reductions, reductions with small changes or in favor of other reductions are preferred to be run first. Hence, we adopted this reduction order:

First, the strategies of removing reactions (deleting trivial reactions) and jumping (jumping over negligible species) are tested, which trim the scheme for further reduction. In the following step, lumping and replacing (extension of lumping) are applied, which results in a significant merge of both reaction pathways and species. Finally, the strategy of removing species is adopted, following removing gas-particle partitioning for condensable species that cannot be removed with removing species.
We have added the explanation to the paper:

**Section 2.1, line 112:**

For the BCARY reduction, the reduction strategies are employed in the following order: removing reactions, jumping, lumping, replacing, removing species, and finally removing gas-particle partitioning.

The reduction strategies are ordered based on their potential influences on the mechanism. The first applied strategies, removing reactions and jumping, trim trivial reactions and species without altering the properties of the species. They are followed by lumping and replacing (as an extension to lumping), which refine the mechanisms considerably by merging the species and reactions involved. Afterwards, the "removing species" strategy attempts to delete all merged and unmerged species. Finally, the strategy of removing gas-particle partitioning is applied in order to remove the partitioning of condensable species, which cannot be removed by removing species. This current order has been tested and found to be efficient for the BCARY mechanism, but it can be changed by the user along with other user-chosen parameters shown in Table 5.

---

**RC:**  *Table 1: Typo in reaction 8. I think it is supposed to say 0.2 BCBOH instead of 0.2 BCAOH.*

**AR:** Actually, both BCAO2 and BCBO2 (Fig. 1) form the same compound BCAOH (Fig. 2) through the self- and cross-reactions of peroxy radicals (RO2-RO2 reaction). As a consequence, BCBOH does not exist in MCM mechanism.

![Figure 1: The molecular structures of the MCM species BCAO2 (left) and BCBO2 (right).](image1)

**Figure 1:** The molecular structures of the MCM species BCAO2 (left) and BCBO2 (right).

![Figure 2: The molecular structure of the MCM species BCAOH.](image2)

**Figure 2:** The molecular structure of the MCM species BCAOH.

**RC:** *Table 2: Typo in lumped reaction R2: 0.753 * (fw,a BCANO3 + fw,b BCBNO3 + fw,c BCCNO3). I would guess the factor is 0.247 instead of 0.753.*

**AR:** Corrected.

**RC:** *Table 2: Cr,b and Cr,c are not defined. It should be defined similar to Cr,a for better clarity.*

**AR:** Thanks for the suggestion. The definitions of Cr,b and Cr,c have been added to Table 2.
RC:  **L119:** It would be much clear to write it as “In this example, a total of 12 chemical reactions involving three organic compounds are reduced to five reactions (4 lumped (R1-4) + 1 surrogate (R0)).”

AR: Thanks for the suggestion. The sentence has be rephrased:

**Section 2.1.1, line 119:**

In this example, a total of **12 chemical reactions** (No. 17 to 29) involving three organic compounds are reduced to **four reactions with one new surrogate** five reactions (production reaction R2 and four destruction reactions R3 to R6 of the new surrogate).

RC: **L122-123:** How are the BCARY isomers undergoing similar reactions with HO2, NO and NO3? Are the authors referring to the R1-4 in the lumped scheme. Please make this clear to the readers.

AR: Here we wanted to point out that MCM species BCAO2, BCBO2, and BCCO2 share similar structures and properties. The fact that they are isomers (which is not necessary for lumping) may explain why all of them reacted with the same species (HO2, NO, NO3, and other peroxy radicals (RO2)). As long as species meets the lumping criteria, they can be merged together via lumping, and their reactions with different oxidants are also be lumped accordingly.

In order to avoid ambiguity, we have rephrased the sentence as follows:

**Section 2.1.1, line 122:**

As demonstrated in the tables, the organic compounds BCAO2, BCBO2, and BCCO2 from the original MCM scheme are the peroxy radicals formed from the OH-initiated oxidation of β-caryophyllene (Table 2). **Because these isomers undergo similar reactions with HO2, NO, NO3, they may conform to the lumping standard.** It is evident from their structures (shown in fig. C1) that they are isomers and may share similar chemical properties. When applying the lumping strategy, BCAO2, BCBO2, and BCCO2 are merged into a new surrogate named "mBCAO2" (Table 3).

RC:  **L122: Why is Cr,a,b,c an arithmetic mean of 5 day simulations? I.e was this 5 day period selected?**

AR: A five-day simulation period is chosen for calculating $C_{r,a,b,c}$ and all other reduction parameters, as a compromise between a shorter period that may not reflect the aging of SOA, and a longer period that may less adequately address the SOA formation and be computational expensive. As the concentrations of ozone, radicals, inorganics and environmental parameters (e.g., temperature, relative humidity) are extracted from the 3-D CHIMERE simulations (24-hour monthly averages). Their diurnal profiles are repeated for five days in 0-D simulations.

To clarify, the following explanation has been added for the settings of 0-D simulation:

**Section 2.3, line 268:**

Unless stated otherwise, a 5-day simulation is performed. Two simulations are performed for each condition starting at midnight (0 h) and noon (12 h) for each condition, focusing on taking into account both the daytime and nighttime chemistry. All 0-D simulations are run for five days in order to consider adequately SOA formation and aging processes.

RC:  **L129: kinetic -> kinetics**
AR: Corrected.

RC: L130: weighting -> weighing
AR: Corrected.

RC: L135: What is this specific behavior?
AR: Here we wanted to point out that different types of compounds may be involved in different types of reactions. A radical, for instance, may be better lumped with another radical than a condensable compound. A PAN compound has a decomposition reaction that does not have other compounds. Therefore, a restriction on certain structural groups was applied to the BCARY reduction.

The term “specific behavior” has been rephrased for clarity:

Section 2.1.1, line 135:
Due to specific chemical behavior, compounds with specific structural groups sharing common chemical behavior may be more appropriately merged together. Thus, compounds containing the following functional groups can only participate in lumping with compounds containing the same groups: peroxyacetyl nitrates (PAN), organic nitrates (RONO2), organic radicals (R), oxy radicals (RO), peroxy radicals (RO2), carboxylic acids (RC(O)OH), percarboxylic acids (RC(O)OOH).

RC: L162: Aren’t alkoxy radicals are RO.
AR: Corrected.

RC: L179: Can the authors explain what the maximum hourly branching ratio is?
AR: We have added the definition of the maximum hourly branching ratio to the main paper:

Section 2.1.4, line 178:
There is no particular restriction to exclude species from the reduction attempt via the strategy of removing compounds or removing gas-particle partitioning. However, for removing reaction, a threshold for its maximum hourly branching ratio under training conditions on the branching ratio of the reaction is applied to the reduction. The branching ratio is defined as the ratio of the destruction rate of one reaction to the sum of the destruction rates of all reactions of the targeted species. In the BCARY reduction, a maximum branching ratio ($B_{rm}$) is applied to the reduction defined as a restriction criterion. All reactions with hourly branching ratio (averaged over the training conditions) under this value (reactions that are likely to a minimal effect on SOA formation) are considered as candidate for removal.

RC: L185: Do the authors mean that that after one loop of reduction (as shown in Figure 1), the subsequent reduction is carried out in the reverse order? And why is saturation vapor pressure used only for the lumping strategy. This part needs more clarity.
AR: Sorry for the ambiguity. For each strategy, the search for potential reduction is conducted following the reverse lists of reaction/species. For example, with removing reactions, GENOA attempts to remove the reaction from the end to the beginning of the reaction list. When applied to the jumping strategy, GENOA
tries to jump the species that has the highest generation and then move down to the species that has the lowest generation.

In lumping, we consider that a condensable species should be first grouped with another compound that has a similar volatility. Thus, the saturation vapor pressure is used to determine the most appropriate lumpable species. The saturation vapor pressure of species is not affected by other strategies and is therefore not used as a criterion for them.

We have added the explanation to the paper:

Section 2.1.4, line 185:

Moreover, the searches for viable reductions via removing are conducted in reverse order of the reaction/species list, beginning with the species/reactions of the highest generation which means that GENOA attempts to remove reactions from the bottom of the list and moves to the previous reactions. The same reverse sequence is followed for other strategies, where the jumping strategy, for instance, GENOA tries to jump the species that has the highest generation and then move down to the species that has the lowest generation. Among all reduction strategies, only lumping alters the saturation vapor pressure of condensable species. Therefore, a rank of saturation vapor pressure for condensable compounds is used exclusively for lumping to determine the most appropriate lumpable species.

RC: Table 5: Are the two conditions ADD1 and ADD2 high or low NOx regimes?

AR: ADD1 is under low NO\textsubscript{x} regime and ADD2 is under high NO\textsubscript{x} regimes. We have added columns in Table 1 (Table 5 in the paper), i.e., the average SOA concentration and NO reactive ratio with RO\textsubscript{2} (R_{RO\textsubscript{2}-NO}), which more clearly indicates the chemical regimes of conditions. If R_{RO\textsubscript{2}-NO} is high, the conditions are in the high NO\textsubscript{x} regime. Otherwise, the conditions are in the low NO\textsubscript{x} regime. For clarity, the digits in Table 5 of the paper have also been changed for temperature and relative humidity.

RC: Figure 2: How do I interpret this? Is the top bar representing 0H and the bottom one representing 12h? Please add this to the figure caption, since 0 h and 12 h are not represented by empty and hashed lines.

AR: Thank you for pointing out the error in Figure 2. The hash line for the 12-h condition was not shown. The figure (Figure 2 in the paper) has been revised to Fig. 3.

RC: L233: I wonder what are the conditions which cause such low O3 and NO3 concentrations?

AR: This condition is located in the northern part of Italy, within the Alpine arch, close to the metropolitan city of Milan. The concentrations of NO transported from polluted areas consume O\textsubscript{3} and NO\textsubscript{3} and produce NO\textsubscript{2}, which explains the low concentrations of O\textsubscript{3} and NO\textsubscript{3} in this area.

We have added the more information about this "ADD2" training condition to the paper:
Table 1: Geographic and meteorological conditions of the training dataset

<table>
<thead>
<tr>
<th>Condition Name</th>
<th>Lat</th>
<th>Lon</th>
<th>Time</th>
<th>TEMP</th>
<th>RH</th>
<th>R_NO\textsuperscript{b}</th>
<th>SOA\textsuperscript{c}</th>
</tr>
</thead>
<tbody>
<tr>
<td>OH NO</td>
<td>36.0</td>
<td>15.4</td>
<td>Jul.</td>
<td>299</td>
<td>679</td>
<td>60</td>
<td>4.1</td>
</tr>
<tr>
<td>OH HO\textsubscript{2}</td>
<td>32.0</td>
<td>-9.4</td>
<td>Jul.</td>
<td>295</td>
<td>296</td>
<td>20</td>
<td>6.1</td>
</tr>
<tr>
<td>NO\textsubscript{3} NO</td>
<td>40.25</td>
<td>-3.4</td>
<td>Jul.</td>
<td>302</td>
<td>2302</td>
<td>69</td>
<td>4.4</td>
</tr>
<tr>
<td>NO\textsubscript{3} HO\textsubscript{2}</td>
<td>32.0</td>
<td>36.6</td>
<td>Aug.</td>
<td>302</td>
<td>2302</td>
<td>29</td>
<td>5.7</td>
</tr>
<tr>
<td>O\textsubscript{3} NO</td>
<td>69.0</td>
<td>33.8</td>
<td>Jan.</td>
<td>260</td>
<td>7</td>
<td>99</td>
<td>5.2</td>
</tr>
<tr>
<td>O\textsubscript{3} HO\textsubscript{2}</td>
<td>68.0</td>
<td>18.2</td>
<td>Dec.</td>
<td>265</td>
<td>5</td>
<td>25</td>
<td>4.6</td>
</tr>
<tr>
<td>ADD1</td>
<td>41.5</td>
<td>-14.2</td>
<td>Dec.</td>
<td>288</td>
<td>6</td>
<td>20</td>
<td>5.5</td>
</tr>
<tr>
<td>ADD2</td>
<td>45.75</td>
<td>9.0</td>
<td>Dec.</td>
<td>279</td>
<td>1</td>
<td>100</td>
<td>4.4</td>
</tr>
</tbody>
</table>

\textsuperscript{a} from left to right: name, latitude, longitude, time period, average temperature, average relatively humidity, daily average NO reacting ratio, simulated total SOA concentration of the training conditions.

\textsuperscript{b} the daily average NO reacting ratio is calculated out of the RO\textsubscript{2} reactivity of NO, HO\textsubscript{2}, NO\textsubscript{3}, and RO\textsubscript{2}. Conditions with high R_NO ratio are considered as in high NO\textsubscript{x} regime.

\textsuperscript{c} the initial concentration of BCARY is 5 \( \mu g/m^3 \).

Figure 3: A bar plot showing the occupancy of seven reacting ratios in BCARY initiation reactions and RO\textsubscript{2} reactions, under the training conditions at midnight (0 h, top bar) and noon (12 h, bottom bar with slash). From left to right, six ratios are presented on each bar in the following order: R_O\textsubscript{2}, R_OH, R_NO\textsubscript{3}, R_RO\textsubscript{2}−NO, R_RO\textsubscript{2}−HO\textsubscript{2}, R_RO\textsubscript{2}−NO\textsubscript{3}, and R_RO\textsubscript{2}−RO\textsubscript{2} (No display if ratio is zero).
Section 2.2, line 232:

One specific exception is the additional condition ADD2, in which is located in the northern part of Italy, within the Alpine arch, close to the metropolitan city of Milan. This condition is in extremely high-NO\textsubscript{x} regime, as high concentrations of NO are transported from polluted areas. These high NO concentrations consume O\textsubscript{3} and NO\textsubscript{3}, causing low concentrations of O\textsubscript{3} and NO\textsubscript{3}. At night, ADD2 has a high R\textsubscript{OH} of 95% at midnight is not due to an abundance of OH, but rather to extremely low concentrations of O\textsubscript{3} (2.9 × 10\textsuperscript{-4} ppb) and NO\textsubscript{3} (1.1 × 10\textsuperscript{-9} ppb) that leads to an absence of nighttime reactivity.

RC: \texttt{L258: Is Kelvin effect not taken into account for gas-to-particle partitioning?}
AR: The Kelvin effect is not taken into account in the simulations, as thermodynamic equilibrium is assumed between gas and particle phases.

RC: \texttt{L262: ”vapor pressure is computed using Mydral and Talkowsky”. This phrase is repeated already in the earlier part of the sentence.}
AR: Rephrased.

AR: Corrected.

RC: Appendix A, L485: There seems to be a contradiction here. The authors say that v1b2 (Nannolal and Jacob and Reid) show the best estimate in comparison with the experiments “As shown in Fig.A1, the SOA distribution simulated with “v1b2” agrees best with the experimental data. Therefore, this method with the vapor pressure computed by Myrdal and Yalkowsky (1997) and the boiling point computed by Joback and Reid (1987) is used in the BCARY reduction”. But in the manuscript why has Myrdal and Talkowsky been selected in place of Nannolal 2008. Why?
AR: Thank you very much for pointing out this error. ”v1” is the method of Nannolal et al. (2008), but in the paper, it was incorrectly referred to as the method of Myrdal and Yalkowsky (1997).

We have corrected the typo:

\textit{Appendix A, line 481:}

Eight methods are provided in UManSysProp, including SIMPOL.1 of Pankow and Asher (2008) (“\textit{sim}”), EVAPORATION of Compernolle et al. (2011) (“\textit{evp}”), and six methods out of the combination of two methods to compute the vapor pressure (“\textit{v0}”: Myrdal and Yalkowsky (1997) and “\textit{v1}”: Nannoolal et al. (2004) and Nannoolal et al. (2008)) and three methods to compute the boiling point (“\textit{b0}”: Nannoolal et al. (2004), “\textit{b1}”: Stein and Brown (1994), and “\textit{b2}”: Joback and Reid (1987)). As shown in Fig. A1, the SOA distribution simulated with “\textit{v1b2}” agrees best with the experimental data. Therefore, this method with the vapor pressure computed by Myrdal and Yalkowsky (1997) and the boiling point computed by Joback and Reid (1987) is used in the BCARY reduction. The results simulated with the final reduced mechanism “Rdc.” is also presented in Fig. A1, which has a great resemblance to the experimental data.

RC: \texttt{L275: I think it would be better to rephrase the sentence defining the FME. Is it so that the simulation error is the larger FME of the two errors I.e the FME of day 1 and the FME of rest of the simulation days? This has}
be more clear in the text.

AR: The following sentences have been rephrased to better explain how we calculate error:

Section 2.4, line 274:

To address the difference of The error of one simulation is defined as the larger of the FME on day one and the FME on days two to five, in order to address the difference in performance of the reduced mechanisms at the early stage of the simulations (first day SOA formation dominates) and at the later stage, the error of one simulation is defined as the larger FME between the FME of the first simulation day 1 and the FME from day 2 to 5, and it is compared (SOA aging dominates). This error is used to evaluate reduction by comparing it to the error tolerance specified in training (Eq. 1). For the evaluation on the training dataset, two errors are estimated compared to the previously verified reduced mechanism with a tolerance denoted \( \epsilon_{\text{pre}} \), and the MCM mechanism with a tolerance denoted \( \epsilon_{\text{ref}} \). The error tolerances are used to restrict both the maximum and the average (half of the tolerance) errors of the training conditions. As for the evaluation on the pre-testing dataset, only one error compared to the MCM mechanism is calculated, with an error tolerance denoted \( \epsilon_{\text{pre-testing}} \). The error tolerances \( \epsilon_{\text{max}} \) and \( \epsilon_{\text{max-testing}} \) are set to the average and maximum errors, respectively.

RC: Eq 1: What does i1 and i2 represent?

AR: i1 and i2 are meaningless and have been removed from Eq.1.

RC: L283: Why does the \( \epsilon_{\text{pre}} \) vary in such a way with respect to \( \epsilon_{\text{ref}} \). More explanation is need here.

AR: \( \epsilon_{\text{ref}} \) and \( \epsilon_{\text{pre}} \) are the criteria we set for evaluating the reduction. The difference between the two criteria is that \( \epsilon_{\text{ref}} \) is compared to the reference mechanism and \( \epsilon_{\text{pre}} \) is compared to the previous validated mechanism. \( \epsilon_{\text{ref}} \) is used to track the performance of the reduction, while \( \epsilon_{\text{pre}} \) is used to avoid large errors introduced by one reduction attempt. So logically, \( \epsilon_{\text{pre}} \) should be less or equal than \( \epsilon_{\text{ref}} \).

In practice, when \( \epsilon_{\text{ref}} \) increases by 1 %, the value of \( \epsilon_{\text{pre}} \) is set from 1 % (minimum value) to the value of \( \epsilon_{\text{ref}} \). By doing this, GENOA first accepts reductions that introduce small errors compared to the previous validated mechanism, and then accepts reductions that introduce larger errors up to \( \epsilon_{\text{ref}} \).

The explanation has been added to the paper:

Section 2.4, line 282:

In order to begin with a conservative BCARY reduction, the initial values of \( \epsilon_{\text{pre}} \) and \( \epsilon_{\text{ref}} \) are both set to 1 %. The values of these error tolerances are then increased to larger values, reflecting the looser criteria used throughout the reducing. \( \epsilon_{\text{ref}} \) is used to track the performance of the reduction, while \( \epsilon_{\text{pre}} \) is used to avoid large errors introduced by one reduction attempt. Therefore, \( \epsilon_{\text{pre}} \) is lower or equal than \( \epsilon_{\text{ref}} \). For every 1 % increase in \( \epsilon_{\text{ref}} \), \( \epsilon_{\text{pre}} \) is stepped up by 1 % from 1 % to the value of \( \epsilon_{\text{ref}} \). By doing this, GENOA first accepts reductions that introduce small errors compared to the previous validated mechanism, and then accepts reductions that introduce larger errors up to \( \epsilon_{\text{ref}} \).

RC: L288: Is this true? As mentioned the <3% avg error is for pre-testing + training dataset. Will the average error still be < 3% for test dataset?

AR: For the BCARY reduction, the pre-testing dataset selected can provide an accurate representation of the average conditions in the testing dataset. Thus, when the average error of the pre-testing condition is less than
3 %, the error of the testing dataset is less than 3 % as well.

Generally, it depends on whether the pre-testing dataset is representative of the testing dataset. As the pre-testing dataset is selected randomly from the testing dataset, it may not initially be a reliable representation of the testing dataset (e.g., the error of pre-testing is much smaller or much larger than the error of testing). It is necessary to modify the pre-testing dataset in such a case. For example, a few conditions with large errors from the testing dataset can be added to the pre-testing dataset to improve the performance of pre-testing if the error of pre-testing is smaller than the error of testing.

We have added the explanation to the paper:

Section 2.2.2, line 240:

Meanwhile, the size of the mechanism has already been significantly reduced, which makes the evaluation of each reduction attempt on the pre-testing dataset less computationally expensive.

In principle, the pre-testing dataset should be able to provide a fairly accurate representation of the testing dataset. However, this may not always be the case, since the pre-testing dataset is selected almost randomly from the testing dataset. Therefore, an adjustment may be required to increase the representativeness of the pre-testing dataset by adding or removing a few conditions.

RC: L303: How does GENOA decide what condensable species to remove? Is it based on species super saturation values?

AR: In the reduction via removing gas-particle partitioning, GENOA tries to remove the partitioning of each condensable regardless of the saturation vapor pressure. The reduction is accepted only if the errors of training/pre-testing conditions are small enough.

In the late-stage reduction, the aerosol-oriented treatments are applied. Since there is strong competition among reduction strategies at the late stage of the reduction process, these treatments are used to reduce species rather than reactions, thereby reducing condensable species.

To avoid confusion, we have rephrased the statements about the aerosol-oriented treatments:

Section 2.5, line 301:

These treatments, which reduce species rather than reactions, are done when the size of the mechanism is below a certain threshold (20 for BCARY reduction). Consequently, the late-stage treatments encourage the reduction via the removing of condensable species, and are referred to as the aerosol-oriented treatments.

RC: L326: Effective partitioning coeff. is temperature dependent. What is the assumed temperature for the classification of SVOCs, LVOCs and ELVOCs? And since the authors use a range of temperatures in their training conditions (268-302 K) does it really make any sense in classifying the SVOCs, LVOCs and ELVOCs at an arbitrary temperature?

AR: The effective partitioning coefficient ($K_p$) is computed at 298 K. We have added the assumed temperature for $K_p$ in the paper.

RC: L332: Are the species in the reduced Rdc. and Khan 2017 mechanism overlapping or identical?

AR: The mechanism of [Khan et al., 2017] preserves information regarding the reaction/species of the first and
second generations of the MCM mechanism. In contrast, the "Rdc." mechanism may preserve information concerning up to the tenth generation of MCM. Thus, the "Rdc." mechanism may provide a more detailed description of SOA formation and aging than the mechanism of [Khan et al., 2017].

**RC:** L341: Condensable species drop to less than 20? It seems that the condensable species is 10% of its original values (10% of 493 48/49 species or thereabout) as shown in Figure 3. Also $\epsilon_{pre}$ is not shown in Figure 3. Also on L 346 it is mentioned as 41 species on the 75th time. So what is the correct number?

**AR:** The number of condensable species indeed dropped to 20 at the 74th step, with a fraction of 5.61% and the initial number of condensable species in MCM BCARY mechanism is 356. As of the beginning of the 75th step (by the end of the 74th step), the number of species is 41, including both condensable and non-condensable species.

Though the numbers in the text have been verified to be accurate, the number of reduction steps might not be clear. According to figure 3 in the paper, the size at the nth step refers to the size at the end of the nth reduction step. However, in the text, we confused it with the size at the beginning of the nth reduction step. Accordingly, the following statements have been revised:

**Section 3.1, line 341:**

- Early stage, from the first to the 74th reduction step, where the number of condensable species drops to 20. By the end of the 74th reduction step, the mechanism is reduced to 68 reactions and 41 species (including 20 condensable species). The early-stage reduction is trained only on the training dataset with the seven pre-described reduction strategies. After $\epsilon_{ref}$ reaching 3%, the list of $B_{ref}$ is changed from [0.05, 0.10, 0.50] to [0.10, 0.50, 1.0].

- Late stage I, from the 75th to the 107th reduction step, where the number of condensables decrease to. By the end of the 107th reduction step, the reduced mechanism consists of 38 reactions and 19 species (including 7 and no more condensable species), and no further reduction can be found within $\epsilon_{ref} \leq 10\%$ and $\epsilon_{pre} \leq 10\%$. On the 75th reduction step, the size of the mechanism drops to 68 reactions and 41 species. From there, the In this stage, the reduction is trained on the pre-testing dataset if the condensable species are removed with lumping, replacing, or jumping. For reduction with other types of reduction strategies, it is first trained on the training dataset and then on the pre-testing datasets. From all reduced mechanisms with seven condensable species, GENOA selected the one with the minimum average errors on the pre-testing dataset (2.44%) to start the next stage.

- Late stage II, from the 108th to the 113rd reduction step. At this stage, the reduction strategy of removing elementary-like reactions is applied to the training. All reductions that reduce the condensables are evaluated exclusively on the pre-testing dataset. From the 108th reduction step to the end of the training, the size of the reduced mechanism was reduced from 37 reactions and 19 species to 23 reactions and 15 species, among which the number of condensable species is reduced from 7 to 6. The average (maximum) error of the final reduced mechanism "Rdc." is 2.65% (17.00%) under the pre-testing dataset compared to MCM.

As $\epsilon_{pre}$ is only compared to the previously validated mechanism, it does not reflect the performance of the mechanism as does $\epsilon_{ref}$. Therefore, we keep $\epsilon_{ref}$ only as an estimation of the performance of the reduction at different stages of the reduction in Figure 3 of the paper.

**RC:** L346-348: Aren't all the reduction strategies trained first with training data sent and then with pretesting data
set? So what is the difference here between, lumping, jumping and replacing compared to other strategies?

AR: In the late stage of reduction, the reduction via removing is evaluated first on the training dataset, and then on the pre-testing dataset, whereas the reduction via lumping, replacing, and jumping is evaluated only on the pre-testing dataset. The reason is that lumping, replacing, and jumping may be more effective in terms of altering the scheme. Compared to removing, they offer more possibilities for reducing species. Therefore, reductions via lumping, replacing, and jumping are evaluated only on the pre-testing dataset. This treatment was tested and adopted for the BCARY reduction. It can be turned off by the user when applied to other reductions.

RC: L351: “evacuated”-> “evaluated”

AR: Corrected.

RC: L358: It should be specified that lumping reduces the condensable species by 35 %.

AR: We have removed the misleading statement:

Section 3.1, line 357:

As expected, the reduction strategy of removing reactions contributes the most to the decrease in the number of reactions (48 %), followed by the strategy of removing species with a contribution of 37 % and lumping with a contribution of 31 %. Meanwhile, both lumping and removing species are significant in the reduction of species, by 35 % and 31 %, respectively.

RC: L366: There is not mC133O in Figure 4.

AR: C133O in the "Rdc." mechanism has the same properties as the one with the same name in the MCM mechanism. Therefore, the name C133O remains unchanged after reduction.

RC: L392: Why is more uncertainty found in regions with low RH and high temperatures? Is it because the training data set does not have enough data to work with in these conditions? Looking at figure 6, it shows that even Russian data points have high uncertainty between 3-6%. This cannot be only due to low RH and high temperature. What could be the other possible reasons for that?

AR: The relatively high error in the regions with low RH and high temperatures indicates that such conditions are not sufficiently represented in the pre-testing dataset. As these conditions are rare to encounter over Europe, they are not included in the pre-testing dataset, such as not increasing the size of the mechanism.

Considering that the average error for pre-testing is 3 %, there are, of course, conditions with errors greater than 3 %, which are placed in the error category between 3 % and 6 %. As we be seen in Fig. 5 of the main paper, the testing errors in July and August (corresponding to the results in Fig. 6 of the paper) are actually very close to 3 %. These conditions are scattered throughout Europe, such as in the Russian area, in northern Europe, as well as in the Mediterranean.

RC: Figure 8: Didn’t the authors claim that PAN is under-represented in GENOA on L 418? But Figure 8 shows higher PAN concentrations for GENOA and lower for MCM.

AR: We apologize for the mix-up in group names in the paper, and there is no doubt that the PAN concentrations are overestimated by the reduced mechanism shown in Figure 8 of the paper.

We have corrected the typo:
In comparison to MCM, only two condensable species containing nitrogen are retained in the "Rdc." mechanism: NBCOOH and C131PAN, leading to an overestimation of the nitrate mass of the underestimation of the organic nitrate group (0.10 µg m⁻³ in MCM and 0.30-0.04 in "Rdc.") and an underestimation of the overestimation of the nitrate mass of the peroxyacetyl nitrate group (0.31-0.10 µg m⁻³ in MCM and 0.04-0.30 in "Rdc.").

RC: L438: It should be explained why due to different volatility species Rdc delays SOA production. Is it due to low LVOC concentrations or high ELVOC concentrations?

AR: Sorry for the misleading. The statement "Rdc delays SOA production" may not accurately describe the general situation in simulations that involve extreme SOA loading and large errors.

Thus, we have removed it from the discussion:

The result indicates that the "Rdc." mechanism may introduce relatively large uncertainty with extreme SOA loading (conditions that were outside the range of conditions used for the construction of the "Rdc." mechanism). Results indicate that the "Rdc." mechanism may delay the SOA formation under large mass loading (larger than 500 µg m⁻³), because of differences of the volatility of the oxidation products which was outside the range of conditions used for the construction of the "Rdc." mechanism and MCM.

References


Authors’ Response to Reviews of
GENerator of reduced Organic Aerosol mechanism (GENOA v1.0): An automatic generation tool of semi-explicit mechanisms
Zhizhao Wang, Florian Couvidat, Karine Sartelet

Reviewer # 3

RC: This paper presented a reduction strategy and developed a software package to reduce the mechanism of SOA formation. The paper is reasonably well written. The method described is innovative and effective. However, there are some information misleading and inappropriate. The current manuscripts needs to be modified before it is accepted for publication.

AR: We would like to thank reviewer # 3 for the positive comments and constructive suggestions, which helped us to substantially improve our manuscript. We have carefully considered all of these comments and revised the manuscript accordingly. Please note that the line and section numbers mentioned in the response correspond to the version of the manuscript before revision.

1. General comments

RC: The authors chose sesquiterpenes as an example to show the methods to reduce SOA formation mechanism.

AR: Sesquiterpene was selected because it is a well-known source of SOA, and its formation mechanism is well documented. Sesquiterpene is also an ideal candidate for model development and demonstration of the reduction methodology, as the oxidation products of sesquiterpene are less volatile and tend to condense more readily than those of lighter molecules such as monoterpene and isoprene.

We have added the reason why we selected sesquiterpene to the main paper:

Section 1, line 72:

The application of GENOA to the MCM degradation scheme of β-caryophyllene (BCARY) (Jenkin et al., 2012) is described in Sect. 3. The β-caryophyllene species is selected because it widely serves as a benchmark for modeling the tropospheric chemistry and SOA formation relevant to is selected for investigation and demonstration of the GENOA algorithm, because it is one of the most abundant and representative sesquiterpenes (SQT) (e.g., Li et al., 2015; Xavier et al., 2019), and its degradation has been evaluated in chamber simulations (Jenkin et al., 2012). The BCARY scheme in the STOCHEM-CRI mechanism (Khan et al., 2017) and Sesquiterpenes are a well-known source of SOAs (Hellén et al., 2020; Tasoglou and Pandis, 2015 and their degradation mechanisms (as BCARY) is well documented in the experiment data of Tasoglou and Pandis (2015) are also used for evaluation
One obvious question is that most of the reactions listed are linear and therefore analytical solutions can be achieved. In such case, the reduction may be of less significance.

The formation of SOA from VOC degradation involves many processes, including gas-phase multi-generation oxidation and gas-to-particle mass transfer. These processes are highly non-linear. For example, the gas-to-particle mass transfer depends on the concentration and composition of both the gas and the particle phases. As the particle composition depends on the environmental conditions and on the formation of other organic compounds (i.e., non-ideality), it is not possible to define an analytical solution. The process of SOA formation from sesquiterpene is, therefore, complex and non-linear.

Moreover, the authors did not show the impact of reduction on the concentration of short-lifetime oxidants, such as OH, HO2 and NO3, which have an important influence on SOA yields. How are the coupling between VOC species and oxidants resolved? Reasonable explanations should be highlighted.

The reduced SOA mechanism focuses solely on reproducing the SOA concentration of those simulated with explicit mechanisms, with fewer species and reactions. Thus, the oxidant concentrations are simulated using implicit gas-phase chemical mechanisms. Current reduced mechanisms do not affect the concentrations of O3 and major radicals (OH, NO3, NO, HO2) produced by the implicit gas-phase mechanism.

In air quality modeling, implicit SOA mechanisms are often based on the surrogate approach (e.g., Odum’s two-product) or the Volatility Basis Set. They are added to implicit gas-phase mechanisms without altering the pathways of oxidants. The same approach is adopted for our semi-explicit SOA mechanisms. Our reduced SOA mechanisms, which include gas-phase chemical mechanisms and aerosol properties for condensible gas-phase species derived from certain SOA precursors, can be added to the implicit gas-phase mechanism in the 3-D model. For example, the 3-D CHIMERE model can simulate SOA using the implicit gas-phase mechanism MELCHIOR2 and SOA mechanism H2O [Couvidat et al., 2018]. With our reduced SOA mechanisms, SOAs in CHIMERE are simulated by MELCHIOR2, the reduced SOA mechanisms generated by GENOA, and H2O for other SOA precursors not covered by our mechanisms.

For the paper, We have added the explanation of how the coupling is generally resolved between VOC species and oxidants in 3-D models:

Section 1, line 36:

Along with To complete implicit gas-phase mechanisms, implicit SOA mechanisms have been developed (Kim et al., 2011), which model the SOA formation specifically without modifying the concentrations of ozone and major radicals. In 3-D modeling, implicit SOA mechanisms or parameterizations are usually added to implicit gas-phase mechanisms, conserving the oxidant chemistry of the implicit gas-phase mechanism.

Implicit SOA mechanisms are often established based on experimental data from smog chamber experiments to represent the formation and evolution of SOA, such as the two-product empirical SOA
We have also added the following statements about our semi-explicit mechanisms:

Section 2, line 80:

The generated semi-explicit mechanisms are designed to preserve the accuracy of explicit mechanisms for SOA formation, while keeping the number of reactions/species low enough to be suitable for large-scale modeling, particularly 3D in 3-D AQMs. The focus of the semi-explicit mechanism is solely on the accurate modelling of SOA. Because ozone, major radicals, and other inorganics are also affected by inorganic and other VOC chemistry, their concentrations are not tracked with the semi-explicit mechanism. Instead, they are simulated using existing implicit gas-phase chemical mechanisms.

Additionally, details about implicit gas-phase mechanisms of the 3-D CHIMERE model and how 3-D CHIMERE results are used for 0-D simulations have been added to the paper:

Section 2.2, line 193:

The version of CHIMERE and its configuration is CHIMERE model and the configuration used for the simulation are described in Lanzafame et al. (2022). The monthly average 3-D CHIMERE simulations were conducted with the implicit gas-phase MELCHIOR2 mechanism (Derognat et al., 2003), which contains 120 reactions and less than 80 lumped species. The MELCHIOR2 mechanism describes the degradation of sesquiterpenes by three oxidant initiated reactions (HUMULE reacts with OH, O3, and NO3, respectively), where the the species HUMULE represents the lumped class of all sesquiterpenes.

The monthly diurnal profiles of hourly meteorological data (e.g., temperature, relative humidity), and hourly concentrations of oxidants and inorganic species were extracted from each location. That information is required in the 0-D simulations with SSH-aerosol (see section 2.4) to reproduce SOA concentrations and compositions under near-realistic conditions. Since the reduced SOA mechanism focuses only on SOA formation, the meteorological data and the concentrations of oxidants, radicals and inorganics are assumed to remain intact during the 0-D SOA simulation. The coordinates and time of each condition are also provided to calculate the solar zenith angle. Because the reduction focuses on the impact on SOA variation, and because no inorganic reactions are considered in the reduced chemical mechanism, the oxidant, radical and inorganic concentrations are fixed as the hourly background, as well as the environmental parameters, are fixed to the diurnal profiles obtained from the CHIMERE data in 0-D SOA simulations. The concentration of HUMULE (denoted C_{SQT} as the CHIMERE surrogate for sesquiterpene (denoted C_{SQT}) is used to estimate the SQT concentration.

Moreover, the GENOA methodology would preserve both SOA and oxidant concentrations in the reduced SOA mechanism if relevant criteria were included in the training process. This may be a potential direction for future development of the GENOA algorithm.

RC: Since GENOA is a semi-explicit mechanism and designed to be used in 3-D models, how can the species in GENOA be matched with the species in 3-D models. One problem that arises is whether GENOA is scale adaptive so that the transport of organic species is well resolved depending on the problem of concern. If not, the package may be useful in optimizing mechanisms for specific urban scenarios and then used in regulatory modeling rather than stated "multi-scale AQM" by the authors.
AR: As we mentioned in the previous comment, semi-explicit SOA mechanisms are added to the implicit mechanisms of the 3-D model. The additional species from semi-explicit SOA mechanisms are then transported by the 3-D model. In the case of sesquiterpene-SOA, only 14 gas-phase species within six condensable species need to be added to the 3-D model, which is a computational scale that is fully acceptable for transport in 3-D modelling. When applied to nested simulation domains (for example over an European domain, a domain over a country and over a city), the user need to use the same semi-explicit SOA mechanisms across all domains, so that the species are the same for transport. In that case, the SOA mechanism has to be trained under all conditions that might be encountered in each domain. As such, GENOA is not "scale adaptive", since the size of the mechanisms does not change with the domain size. Therefore, transport of those additional VOC species is not a problem. To avoid confusion, we have removed the word "multi-scale" from the paper.

RC: It would be more useful if the manuscript can include some information on memory optimization and computational efficiency.

AR: Thanks for the suggestion. We have included the reduced CPU time for the testing process in the paper:

```
Section 3.2.1, line 380:

During the testing procedure, the "Rdc." mechanism is evaluated at 12,159 locations, with two different starting times (0 h and 12 h). The 0-D testing for "Rdc." took approximately 2% of the CPU time consumed by MCM.
```

In addition, we have tested the memory usage of the MCM and RDC mechanisms with the box model SSH-aerosol. As shown in Fig[1], the peak memory consumption is reduced by 96% (from 11.8 MiB to 577.1 KiB) after the reduction. It should be noted that the result of 0-D testing may not be linear to the computational efficiency of 3-D modelling since 3-D simulations include other processes (e.g., transport, deposition). However, the results confirm that the semi-explicit SOA mechanism is conducive to large-scale modeling.

Generally speaking, in 3-D modelling, the most time-consuming process is solving the gas-particle partitioning in SOA modelling. The number of condensable species has a significant impact on CPU performance. As for memory consumption, it varies with the number of species transported especially in the condensed phase, as those are often considered for different particle sizes. Therefore, decreasing the number of compounds that may partition to the condensed phase is a priority for semi-explicit mechanisms. For sesquiterpene, the number of condensables increases only by six, which is acceptable for most large-scale simulations.

### 2. Specific comments

RC: **L26: three-dimensional models -> three-dimensional (3-D) models**

AR: Revised.

RC: **L27: 3D -> 3-D**

AR: Changed.

RC: **L32: carbon-bound -> carbon-bond**

AR: Changed.
Figure 1: Memory maps for MCM (top panel) and "Rdc." (bottom panel) mechanisms simulated with SSH-aerosol under the same conditions. The graphs are generated with massif-visualizer.
Section 1, line 34:

These mechanisms were primarily developed for ozone simulation and reflect only the most significant chemical phenomena occurring in the atmosphere, where VOCs are represented by implicit gas-phase mechanisms. They were developed and validated to simulate the concentrations of oxidants and other conventional air pollutants such as ozone and NO$_2$. In these mechanisms, VOCs have been grouped into a limited number of model species because of computational considerations, and the SOA formation is usually not considered.

Table 1: As the authors stated "using surrogates assigned to molecular structures", listing the information of molecular structure of major species in another table would be better for other researchers to understand the reduction strategy.

The N/C ratio, however, is underestimated by the "Rdc." mechanism by 37% on average (ratio equal to 0.019 in MCM and to 0.012 in "Rdc."). The N/C ratio is consistent with the nitrogen-containing SOA mass distribution discussed previously, as a result of indicating the over reducing organic nitrates in the building process of "Rdc.". A total of three nitrogen-containing organics (NBCO$_2$, NBCOOH, and C131PAN) are preserved in "Rdc.", of which two (NBCO$_2$, NBCOOH) are first-generation products. Therefore, during the first 10 hours, the N/C ratio curve simulated by "Rdc." drops, whereas in MCM it increases as higher-generation nitrates are produced.

References

<table>
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<th>NBCOOH</th>
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Figure 2: UNIFAC functional group decomposition for the condensable species in the "Rdc." mechanism.