



# Molecular representation of benzene and phenol secondary organic aerosols

Aurélien Le Bayon<sup>1</sup>, Aliisa Ojala<sup>2</sup>, Siddharth Iyer<sup>2</sup>, Zhizhao Wang<sup>3</sup>, Victor Lannuque<sup>4</sup>, Florian Couvidat<sup>4</sup>, Raluca Ciuraru<sup>5</sup>, and Karine Sartelet<sup>1</sup>

<sup>1</sup>CEREA, Ecole nationale des ponts et chaussées, EDF R&D, Institut Polytechnique de Paris, IPSL, 77455 Marne-la-vallée, France

<sup>2</sup>Aerosol Physics Laboratory, Tampere University, 33720 Tampere, Finland

<sup>3</sup>College of Engineering Center for Environmental Research and Technology (CE-CERT), University of California, Riverside, CA 92521, USA

<sup>4</sup>Institut National de l'Environnement Industriel et des Risques, 60550 Verneuil-en-Halatte, France

<sup>5</sup>ECOSYS, University Paris-Saclay, INRAE, AgroParisTech, 91120 Palaiseau, France

**Correspondence:** Aurélien Le Bayon (aurelien.le-bayon@enpc.fr), Siddharth Iyer (siddharth.iyer@tuni.fi), and Karine Sartelet (karine.sartelet@enpc.fr)

## Abstract.

Recent experimental and theoretical work has highlighted missing pathways in the oxidation of aromatic compounds, with consequences for the formation of highly oxygenated products relevant to secondary organic aerosol (SOA) formation. In this study we develop an updated, quasi-explicit oxidation mechanism for benzene based on Master Chemical Mechanism (MCM) v3.3.1, extended to represent key multi-generation chemistry. Quantum chemical calculations are used to derive the formation and evolution of geminal-diol bicyclic peroxy radicals and to parameterize the subsequent molecular rearrangements. The mechanism further incorporates autoxidation sequences previously developed for peroxy and alkoxy radicals, successive OH additions and cyclic epoxides formation. This approach enables a more mechanistic description of aromatic oxidation leading to highly oxygenated, low-volatility products. The mechanism is evaluated using box-model simulations against a set of chamber experiments conducted under various conditions. Simulated aerosol concentrations agree well with observations and emphasize the dominant contribution of the newly implemented pathways. The quasi-explicit mechanism is subsequently reduced using the GENerator of reduced Organic Aerosol mechanisms (GENOA), resulting in a semi-explicit mechanism reduced to 1 % of its original number of species while reproducing SOA mass with a mean error of 3.9 % relative to the quasi-explicit scheme. Using the validated reduced mechanism, zero-dimensional simulations under contrasted atmospheric conditions are conducted to estimate the SOA compounds that are formed in the early stages of oxidation and those that dominate the system at later stages.

## 1 Introduction

Volatile organic compounds (VOCs) play an important role in the formation of secondary organic aerosols (SOA). In the atmosphere, these compounds may react with oxidants such as OH, O<sub>3</sub> and NO<sub>3</sub>, leading to the formation of more functionalized



20 organic species (Jacob, 1999; Kroll and Seinfeld, 2008). These newly formed species may have a lower saturation vapor pressure and are more likely to participate to the formation and growth of SOA (Pankow, 1994), which have multiple impacts on health (Mauderly and Chow, 2008; Pope et al., 2002) and environment (Kanakidou et al., 2005).

Aromatic compounds account for a significant proportion of anthropogenic VOCs (Reimann and Lewis, 2007). Benzene, the simplest aromatic compound, is considered a carcinogen and endocrine disruptor (Baan et al., 2009), and it can affect the  
25 circulatory system or lead to the development of chronic diseases (Lan et al., 2004; Bahadar et al., 2014). This compound is emitted by many sources: traffic, cigarette smoke, and industrial solvent use, particularly in sectors such as textiles, pesticides and pharmaceuticals (Weisel, 2010). Phenol is the main product of benzene oxidation by the hydroxyl radical. It may also be primarily emitted by numerous primary sources in the atmosphere, such as biomass burning, traffic exhaust and manure spreading in agriculture (Mohr et al., 2013; Sun et al., 2008). Oxidation of benzene and phenol leads to the formation of  
30 functionalised products that may form SOA, including glyoxal and methylglyoxal (Jang et al., 2002).

In three-dimensional (3D) simulation models, the formation of SOA from a VOC precursor is usually parameterized based on yields derived from atmospheric simulation chamber experiments. Typical approaches include the two-compound Odum approach (Odum et al., 1996), the surrogate approach (Pun et al., 2006; Couvidat et al., 2012), or the volatility basis set (Donahue et al., 2006). While computationally efficient, these parameterizations do not explicitly resolve the molecular composition  
35 of the particle phase, or do so only for a limited number of representative compounds in the case of the surrogate approach (Couvidat et al., 2012; Majdi et al., 2019). To improve the representation of aerosol chemical composition and reduce reliance on empirical fitting, semi-explicit approaches are emerging (Wang et al., 2022; Luttkus et al., 2024; Wieser et al., 2024). These approaches selectively retain the main products and reaction pathways from explicit oxidation mechanisms, along with their subsequent degradation products.

40 Quasi-explicit oxidation schemes provide a comprehensive representation of atmospheric photooxidation products and reactions (Bloss et al., 2005; Aumont et al., 2005), but their complexity limits their use in 3D air-quality models. Hence Wang et al. (2022, 2023) developed an algorithm to reduce quasi-explicit chemical mechanisms to sizes suitable for 3D atmospheric simulations, while conserving the predominant pathways and characteristics of SOA compounds. In this algorithm, called  
45 GENERator of reduced Organic Aerosol mechanism (GENOA), the main oxidation pathways are identified, accounting for radical chemistry, which has been shown to be crucial for monoterpenes (Wang et al., 2024). The SOA chemical composition is accurately represented in terms of functional groups (Wang et al., 2022), making it possible to capture interactions between compounds through non-ideality and partitioning into both organic and aqueous phases (Couvidat and Sartelet, 2015; Kim et al., 2019). Coupled to 3D air-quality models, the reduced mechanisms enable mapping of the molecular composition of SOA on multiple spatial scales, as demonstrated for terpenes at the European scale (Wang et al., 2024) and for toluene from  
50 regional to street-scale applications (Sartelet et al., 2024).

The Master Chemical Mechanism (MCM) (Jenkin et al., 2003; Bloss et al., 2005) is a quasi-explicit mechanism that accurately represents precursor degradation and formation of first-generation oxidation products (Calvert et al., 2002). However, based on the results of experimental studies on benzene oxidation (Nakao et al., 2011; Borrás and Tortajada-Genaro, 2012; Priestley et al., 2021; Cheng et al., 2021; Choi et al., 2024), several compounds and oxidation pathways are not included in



55 this scheme which especially underrepresents the formation of highly oxidized compounds and organonitrates, thus strongly  
impacting SOA formation. In recent years, experimental data analysis led to the identification of new oxidation pathways with  
adjustment of previously existing ones, to better represent the formation of specific compounds, such as nitrocatechol (Harrison  
et al., 2005) and the formation of observed fragmentation products, such as glyoxal and muconaldehyde (Wang et al., 2013) or  
ketene-enols and small carboxylic acids (Xu et al., 2020; Wang et al., 2020). An autoxidation mechanism was also developed  
60 for benzene, allowing for a better representation of several classes of compounds, such as peroxides and organonitrates (Pichel-  
storfer et al., 2024). Pathways related to the rearrangement of bicyclic peroxy radicals (BPR) are still missing, especially for  
geminal diol derived BPRs formed from the oxidation of phenol and catechol.

This work aims to construct a semi-explicit scheme for benzene oxidation that integrates the main SOA formation pathways  
relevant under atmospheric conditions. First, a quasi-explicit oxidation scheme is developed, incorporating recent advances  
65 in the understanding of benzene oxidation. This mechanism, based on the Master Chemical Mechanism and enriched by the  
state-of-the-art known oxidation routes, is presented in section 2. Quantum chemical calculations are then used to investigate  
the role of BPR rearrangements, which may be particularly relevant for geminal diols formed during the oxidation of phenol  
and catechol. Pathways related to BPR rearrangement are determined in section 3. Comparison with experimental data is used  
to validate the quasi-explicit scheme in section 4. The quasi-explicit scheme is reduced using GENOA for 3D atmospheric  
70 air-quality modeling by training over Europe in section 5. The relative influence of the different pathways for benzene and  
phenol SOA formation under atmospheric conditions is studied in section 6.

## 2 Benzene and phenol oxidation mechanism

### 2.1 The MCM near-explicit baseline

The quasi-explicit benzene oxidation mechanism developed in this work is based on the MCM v3.3.1 (Jenkin et al., 2003;  
75 Bloss et al., 2005) to which several oxidation pathways are added. The MCM scheme integrates 135 species and 334 reactions,  
initiated by the benzene oxidation by the hydroxyl radical. Three distinct compounds result from this first step, phenol from  
hydroxyl radical addition on the aromatic cycle, a BPR from the addition of OH and two O<sub>2</sub> and a dicarbonyl epoxide through  
OH and O<sub>2</sub> addition followed by ring opening.

### 2.2 Successive OH addition pathway

80 In addition to the H-abstraction from the aromatic ring, OH can react with phenols via addition to the aromatic ring. In the  
benzene MCM scheme, two successive OH additions are represented, leading to the formation of two phenolic compounds,  
phenol and catechol (dihydroxybenzene). Owing to the electrophilic nature of the hydroxyl group, the kinetics of OH addition  
become increasingly important as the number of OH-substituted carbons on the aromatic ring increases (Atkinson et al., 2006).  
Following the work of Schwantes et al. (2017) on toluene oxidation, further OH addition is considered in the quasi-explicit  
85 scheme, leading to the formation of new phenolic compounds, tri, tetra and pentahydroxybenzene. These compounds further



react in the atmosphere and lead to the formation of BPRs through H abstraction and O<sub>2</sub> addition, alkyl peroxides through RO<sub>2</sub> + NO reactions and quinones formation through bicyclic alkoxy radical degradation. The branching ratios used in the multi-hydroxylation pathway could be assumed to be the same as their methylated equivalent presented in Schwantes et al. (2017) for cresol, i.e., 7 % for H-abstraction and 93 % for addition. However, recent studies indicate a higher branching ratio  
90 for abstraction in the reaction of catechol with OH, Finewax et al. (2018) proposes an abstraction branching ratio of 30 % with an addition rate of 70 %. The kinetic constants are extracted from the MCM when available or calculated using the Structure-Activity Relationship (SAR) method implemented in the generator of explicit mechanisms GECKO-A (Generator for Explicit Chemistry and Kinetics in the Atmosphere) (Jenkin et al., 2018). For aromatic compounds, the kinetic constant is calculated based on the number of substituted carbons and the nature and position of the substituents on the aromatic ring (see example  
95 in Supplement S1). The reactions and compounds resulting from the successive hydroxylation of benzene are integrated into the quasi-explicit mechanism (see Supplement S2).

### 2.3 Peroxy and alkoxy radicals autoxidation pathway

Autoxidation, defined as successive intramolecular reactions followed at each step by O<sub>2</sub> addition reactions, results in the formation of peroxy radicals (RO<sub>2</sub>) with high oxygenated content. Bimolecular reactions involving peroxy radicals may also  
100 lead to the formation of alkoxy radicals. Radical termination in the autoxidation cycle leads to the formation of organonitrates (reaction with NO), hydroperoxides (reaction with HO<sub>2</sub>), carbonyl compounds and dialkyl peroxide dimers (ROOR'). These dimers are compounds of extremely low volatility and result from the accretion reactions between peroxy radicals RO<sub>2</sub>–R'O<sub>2</sub> (Berndt et al., 2018).

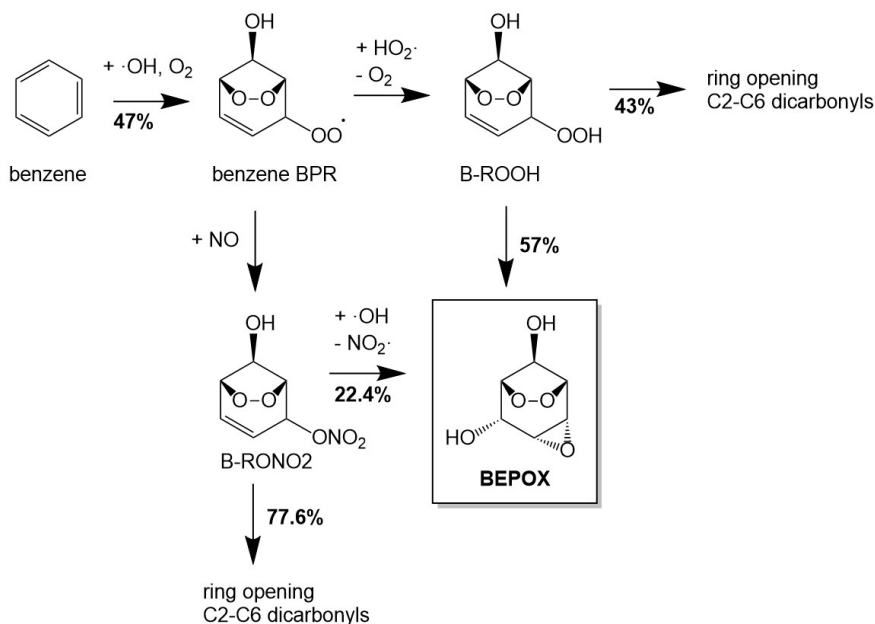
The autoAPRAM framework developed by Pichelstorfer et al. (2024) allows to generate complete autoxidation mechanisms  
105 using specified alkoxy and peroxy radicals as input. Two sets of standard reaction types are implemented in the framework: the first set comprises twelve types of reactions specific to peroxy radicals, taking into account reactions with radicals (NO, HO<sub>2</sub>, RO<sub>2</sub>), possible autoxidation reactions (H-shift and O<sub>2</sub> addition), accretion reactions, alkoxy radical formation from bimolecular reactions and peroxy radical termination (Crouse et al., 2012; Bianchi et al., 2019; Orlando and Tyndall, 2012). The second set defines alkoxy radicals chemistry through three reactions and includes fragmentation, autoxidation and reactions  
110 forming closed shell products (see Supplement S9). These reaction sets are applied to each alkoxy and peroxy radical from the input scheme and each newly formed radical, generating the complete autoxidation mechanism.

Using the autoAPRAM framework with the peroxy and alkoxy radicals from the benzene MCM mechanism as input, the autoxidation mechanism generated in Pichelstorfer et al. (2024) comprises 869 species and 934 reactions. Over 85 % of these reactions stem from the accretion processes, as each peroxy radicals combination must be considered as a reaction. This  
115 mechanism is integrated in the quasi-explicit scheme to take into account the autoxidation processes initiated by the benzene MCM scheme.



## 2.4 Cyclic epoxide formation pathway

Epoxide compounds exhibit significant uptake in the particle phase, thus playing an important role in aerosol formation and growth (Piletic et al., 2013). For isoprene chemistry, epoxide formation is explained as the OH-oxidation of a carbon double bond neighboring a hydroperoxide group (Paulot et al., 2009; Lin et al., 2012). In a similar fashion, Fu et al. (2023) identified the oxidation pathways leading to the formation of cyclic epoxides from toluene. From electronic structure and kinetic calculations, OH oxidation of toluene-derived bicyclic peroxides (T-ROOH) and nitrates (T-RONO<sub>2</sub>) can lead to the formation of the cyclic epoxide, with a 56.1 % and 22.4 % yield respectively. Other competitive reaction pathways are discussed in Fu et al. (2023), leading to the formation of a ring-open epoxide with a 1.44 % yield as well as highly volatile C<sub>2</sub>-C<sub>4</sub> dicarbonyls with a 42.4 % yield for T-ROOH. The oxidation of T-RONO<sub>2</sub> essentially leads to the formation of light dicarbonyls (56.0 %), dicarbonyl retaining the O-O bridge (21.4 %) and the nitrated equivalent of the epoxide formed by T-ROOH (0.2 %). In the absence of further information for benzene, the branching ratios presented in Fu et al. (2023) are used for the OH oxidation of benzene-derived bicyclic peroxide and nitrate. Minor oxidation pathways (<2 % yield) are not considered in the implemented mechanism. The compounds and adapted branching ratios involved in the formation of BEPOX are described in Fig. 1.



**Figure 1.** Formation of the cyclic epoxide BEPOX from B-ROOH and B-RONO<sub>2</sub> oxidation and competing reaction pathways.

## 130 2.5 Adaptation of existing pathways

The oxidation pathways proposed by MCM v3.3.1 for benzene have been updated based on recent experimental results and theoretical calculations. In MCM v3.3.1, the oxidation reaction of benzene by OH leads to the formation of phenol (53 %),



of benzene-BPR (35.2 %) and of an open-chain epoxide (11,8 %). Experimentally, the epoxide is observed with a very low yield (Zaytsev et al., 2019; Xu et al., 2020) and epoxide compounds are rather formed from BPR pathways, as discussed in section 2.4. First generation epoxide formation is therefore removed from the quasi-explicit scheme and the branching ratio leading to the BPR formation is updated to 47 % (Xu et al., 2020).

In the benzene MCM scheme, the bicyclic alkoxy radical formed from benzene BPR is assumed to degrade exclusively via decomposition into glyoxal, butenedial, and furanone. However, a thermokinetic study conducted by Wang et al. (2020) demonstrated that an alternative H-shift pathway can compete with decomposition, leading to the formation of a ketene-enol intermediate. This intermediate rapidly oxidizes to formyl ketene, glyoxal, and formic acid. The same study also investigated butenedial oxidation, and identified the 3-carbon aldehyde acrolein as an additional degradation product. To better represent the bicyclic alkoxy radical degradation, oxidation schemes of ketenol (derived from Wang et al. (2020)) and acrolein (from MCM v3.3.1) are included in the quasi-explicit benzene scheme.

Due to their high level of functionalization and very low saturation vapor pressures, most end products resulting from the newly implemented pathways partition almost entirely into the particulate phase. For compounds whose volatility is high enough to allow their presence in both gas and particulate phases, loss reactions are implemented to prevent their accumulation. Following the approach used in Lannuque et al. (2023), degradation via photolysis and reactions with OH, NO, and NO<sub>3</sub> radicals are considered for these compounds without explicitly accounting for their oxidation products and using the SAR method to estimate their kinetic rate constants.

### 3 Rearrangement of the bicyclic peroxy radicals

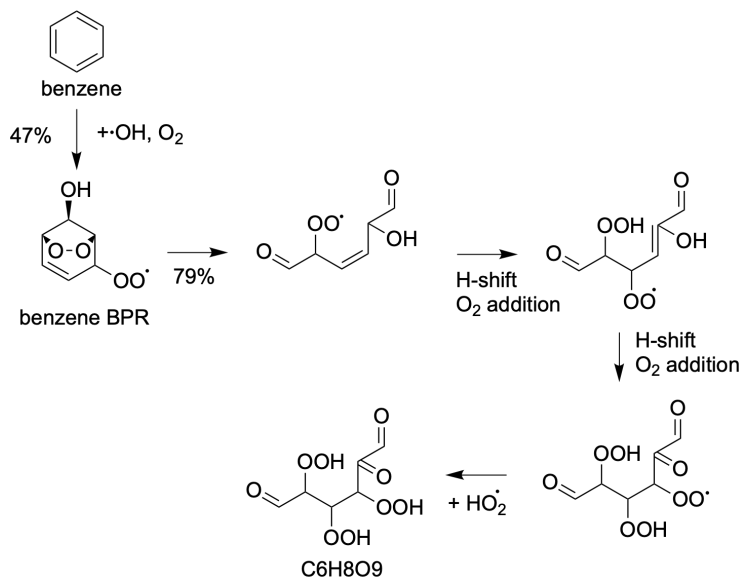
#### 3.1 Benzene BPR

Bicyclic peroxy radicals are key intermediates in the chemistry of aromatic compounds, formed through OH addition on the ring followed by O<sub>2</sub>-additions. In the MCM scheme, benzene BPR forms a bicyclic alkoxy radical mainly through reaction with NO, being able to undergo ring-opening and ultimately forming C<sub>2</sub> and C<sub>4</sub> compounds, such as glyoxal, butenedial and furanone.

Non-cyclic compounds may also be formed from spontaneous rearrangement of BPR, as detailed in Iyer et al. (2023) for toluene. This process leads to the breakage of the peroxide bridge along one of the adjacent C-C bonds, linked to the OH functional group. Following ring-breakage, the ring-opened RO<sub>2</sub> undergoes two consecutive autoxidation cycles. A 1,6 H-shift from the -OH substituted carbon first leads to a resonance stabilized alkyl radical, that then rapidly adds O<sub>2</sub> to form a C<sub>6</sub>H<sub>7</sub>O<sub>7</sub> peroxy radical. This in turn has a 1,6 enolic H-shift, a reaction reported previously to be exceptionally fast (Peeters and Nguyen, 2012; Franzon et al., 2025), forming a C<sub>6</sub>H<sub>7</sub>O<sub>9</sub> peroxy radical. Its hydrogenation by reaction with HO<sub>2</sub> leads to the formation of the main termination product of this cycle, the trihydroperoxide C<sub>6</sub>H<sub>8</sub>O<sub>9</sub>. Due to its low saturation vapor pressure, it is mainly found in the particulate phase and further autoxidation reactions are not represented. The autoxidation sequence previously described and represented in Fig. 2 is implemented in the quasi-explicit oxidation scheme. The kinetic



165 constants for benzene BPR rearrangement and further reactions are based on the thermodynamic calculations presented in Iyer et al. (2023).



**Figure 2.** Simplified autoxidation pathway and termination of benzene BPR rearrangement as represented in the quasi-explicit mechanism.

### 3.2 Geminal diol BPR

In commonly used oxidation schemes, hydroxyl radical addition only occurs on non-substituted ring carbons, leading to the formation of vicinal diols. However, OH addition can also occur at already substituted carbons. This can lead to geminal diol  
170 BPRs that can undergo molecular rearrangement with rate constants that are several orders of magnitude higher compared to non-oxygenated aromatics. (Ojala et al., 2025). As a result, the oxidation of phenol and catechol via the geminal diol pathway can rapidly yield highly oxygenated organic molecules (HOM), including C<sub>6</sub> carboxylic acids and even diacids in the case of  
175 catechol. The yields of HOM via this pathway are a function of the initial OH addition yield, branching ratio to the formation of BPR intermediates, molecular rearrangement rate coefficient of the BPR and subsequent chemistry following molecular rearrangement. Reaction rate constants and resulting product structures are determined using quantum chemical calculations, as is now detailed.

#### 3.2.1 Quantum chemical calculations

The rate coefficients are calculated using multi-conformer transition state theory (MC-TST), with the equation:

$$k = \kappa \frac{k_b T}{h} \frac{\sum_i Q_i e^{-\frac{\Delta E_i}{k_b T}}}{\sum_j Q_j e^{-\frac{\Delta E_j}{k_b T}}} e^{-\frac{E_i - E_j}{k_b T}}$$



180 where  $k_b$  is the Boltzmann constant,  $T$  is the temperature,  $h$  is the Planck constant,  $Q_k$  are the partition functions and  $E_k$  are the energies. The index  $i$  refers to the transition states and  $j$  to the reactant, the summations runs over all relevant conformers and the energy differences in the sum refer to the energy difference in relation to the lowest energy conformer. The  $\kappa$  is the tunneling coefficient calculated with the Eckart tunneling method, which uses the energies of the optimized intrinsic reaction coordinate (IRC) endpoints and the magnitude of the imaginary frequency of the lowest energy transition state conformer.

185 The quantum chemical calculations follow the steps detailed in Møller et al. (2016). Full conformer sampling of all studied molecules is performed with the Spartan '24 program using the MMFF method. Single-point calculations are performed for the conformers at the B3LYP/6-31+G(d) level of theory (henceforth denoted B3LYP) with structures within the cutoff of 5 kcal/mol in relation to the lowest energy conformer being selected for further optimization. If many conformers are present, an additional geometry optimization at the same B3LYP level is used, and conformers outside a 2 kcal/mol cutoff are rejected. Further density

190 functional theory geometry optimization is done at  $\omega$ B97X-D/aug-cc-pVTZ level of theory (denoted  $\omega$ B97 level of theory), with the same level of theory used for the frequency calculation for determining the partition function. Transition states are found by first carrying out constrained optimizations of guess transition state structures, followed by unconstrained transition state optimizations using Gaussian 16 software, both at B3LYP level of theory. Conformer sampling,  $\omega$ B97 optimization and frequency calculations are performed on the lowest energy conformers. Further refining at ROHF-ROCCSD(T)-F12a/VDZ-F12

195 (abbreviated F12) level of theory is done for the lowest energy conformers of all studied molecules, and the IRC endpoints.  $\omega$ B97 level of theory is used for H-shift reactions from OH groups, as the F12-level calculations for this reaction class have previously been shown to lead to artifact solutions (Møller et al., 2019).

### 3.2.2 Endocyclization branching ratios of the geminal diol mechanism

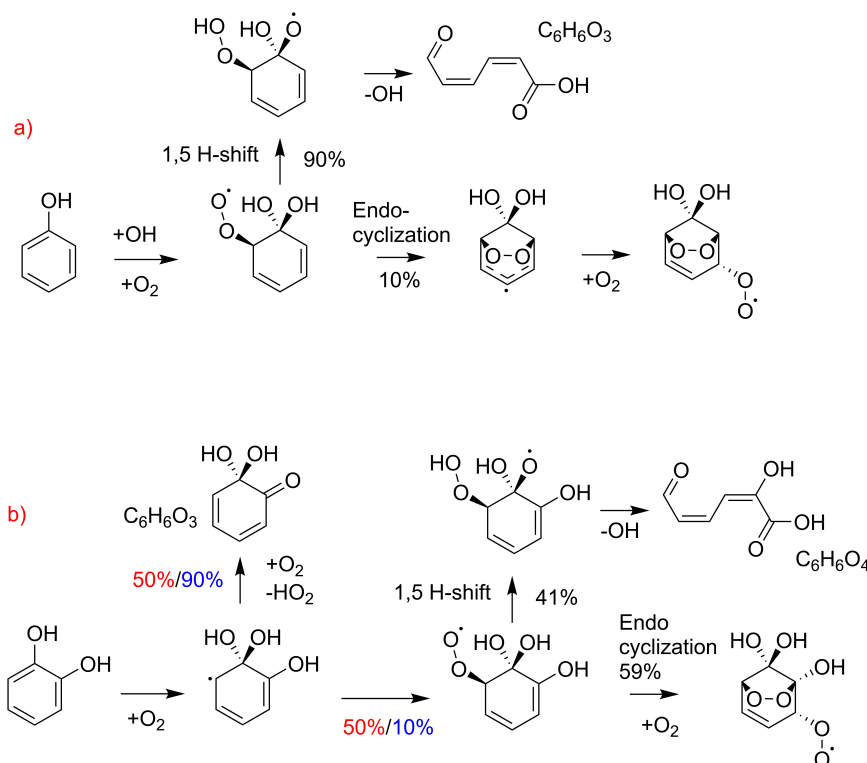
The formation of BPRs requires 5-member oxygen bridge formation over the location of the newly added OH group. Forming

200 the bridge this way is important as the H-O interaction makes the endocyclization reaction competitive. This results in a delocalized alkyl radical, which forms a BPR with the addition of  $O_2$ . Ignoring chirality, one geminal diol BPR structure is possible for phenol due to the symmetry of the system, while two are possible for catechol. The endocyclization reactions and, consequently, the formation of geminal diol BPRs, compete with 1,5 H-shift reactions for phenol and catechol, and an additional  $HO_2$  loss reaction for catechol specifically. This pathway leads to the formation of a closed shell  $C_6H_6O_3$  compound

205 containing geminal diol and carbonyl functionalities. The 1,5 H-shift scheme is illustrated in Fig. 3, with the rates given in Table 1.

**Table 1.** Rate coefficients for the reactions relevant for endocyclization ( $s^{-1}$ )

	Phenol	Catechol
1,5 H-shift	1.52	30.87
Endocyclization	0.17	44.68
Branching ratio	10.0 %	59.1 %



**Figure 3.** Endocyclization. a) for phenol and b) for catechol. Endocyclization has branching ratios of 10 % and 59 % for phenol and catechol respectively.

The endocyclization branching ratio is calculated from equation  $BR_{path} = \frac{k_{path}}{k_{total}}$ . The tunneling factor is subject to uncertainties due to the labile nature of the product alkyl radical, as discussed in Ojala et al. (2025). For this reason, a tunneling factor of 18.0 is used in analogy to para-cresol, yielding an endocyclization branching ratio of 10 %. For catechol, the alkyl radical is stable at the B3LYP level of theory and this is used for the tunneling calculations, yielding a branching ratio of 59 %.

Due to the asymmetry of the molecule, the  $O_2$  addition to catechol can happen to two different locations and the subsequent chemistry is site-specific (see 3). The  $O_2$  addition to the OH-substituted carbon leads to termination via  $HO_2$  loss forming a closed shell carbonyl compound, hindering BPR formation. On the other hand,  $O_2$  addition to the unsubstituted carbon in ortho position relative to the geminal diol carbon lead to endocyclization. The branching between the two  $O_2$  addition sites cannot be accurately determined using quantum chemistry methods. To account for this, two scenarios were considered, corresponding to  $O_2$  addition branching ratios of 10 % and 50 % to the pathway leading to endocyclization. Further details are provided regarding the mechanisms in Table 2 and a sensitivity analysis is conducted in section 4.5.4 regarding the endocyclization branching ratio.

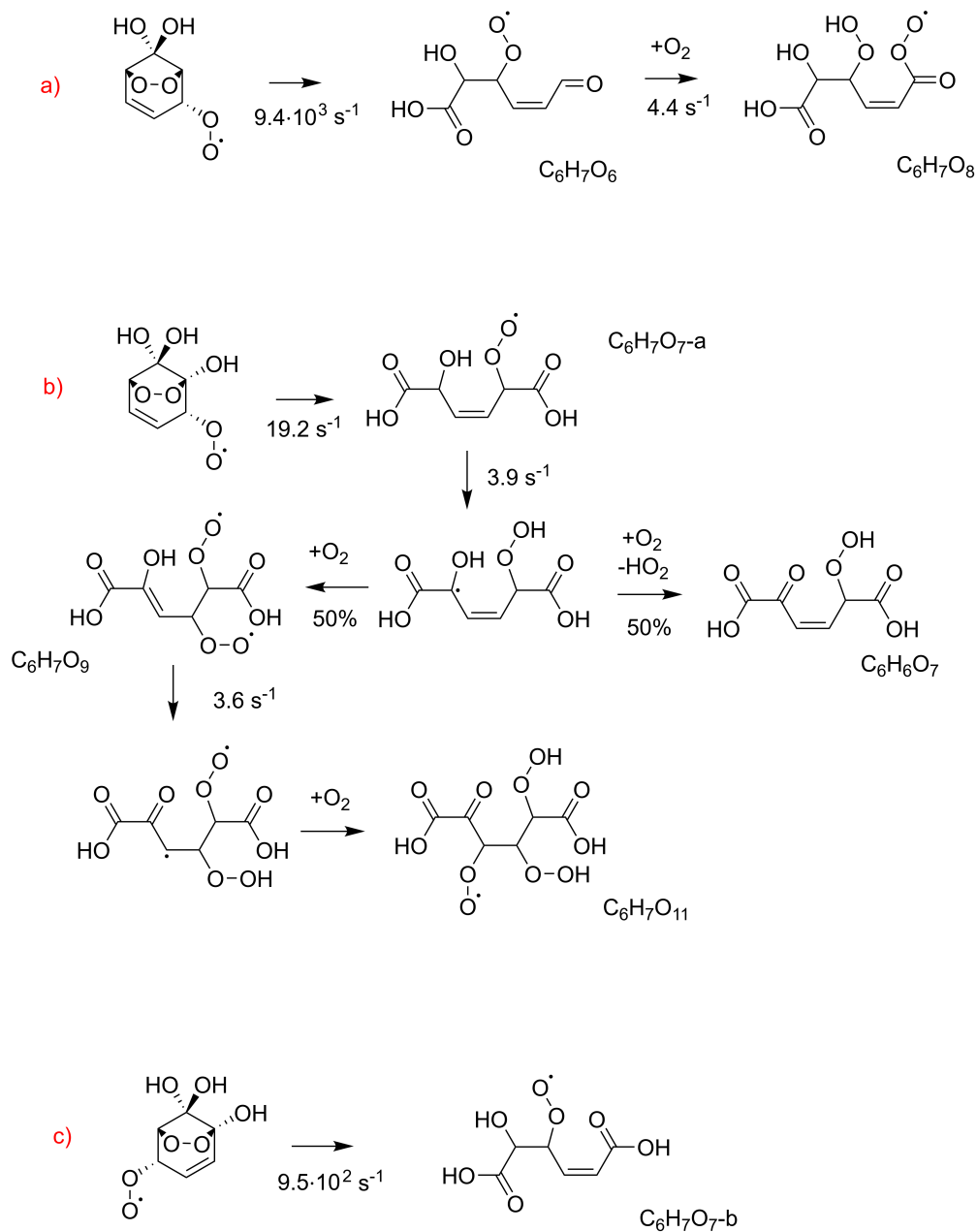


### 3.2.3 Molecular rearrangement

220 Endocyclization is quickly followed by  $O_2$  addition to form a geminal diol BPR, which can undergo a molecular rearrangement reaction as described previously for toluene and other non-oxygenated aromatic hydrocarbons in Iyer et al. (2023). Phenol has only one BPR isomer due to the symmetry of the molecule, whereas catechol has two BPR isomers depending on which carbon the  $O_2$  adds to. Both of these have different molecular rearrangement reaction rate coefficients that lead to two unique ring opened peroxy radicals. The molecular rearrangement reaction and the subsequent chemistry is shown in Fig. 4 for all of these.

225 The molecular rearrangement reaction can proceed in two ways,  $C_1$  or  $C_2$ , depending on which of the two C-C bonds involved in the reaction is broken (Iyer et al., 2023; Ojala et al., 2025). The transition state for the  $C_1$  pathway exhibits an intramolecular interaction between the -OO and -OH groups, reducing the reaction barrier of the rearrangement. In the presence of this interaction, the transition state for phenol tends to constantly optimize to  $C_1$  regardless of the bond constraints used. In the case of catechol, both BPRs undergo molecular rearrangement only along one direction as shown in Fig. 4. We  
230 were unable to find the transition state for the other pathway for either BPR isomer, likely due to the fact that it leads to geminal diol intermediates that are energetically unfavorable.

The  $C_1$  pathway for phenol yields a  $C_6H_7O_6$  peroxy radical that can undergo a 1,6 aldehydic H-shift followed by  $O_2$  addition to form a  $C_6H_7O_8$  peroxy radical (Fig. 4 a). To simplify the mechanism in the model, the two catechol BPR isomers are treated as a single system, both with molecular rearrangement rate coefficients of  $19.2\text{ s}^{-1}$ , leading to an equal 50% yields  
235 of ring opened peroxy radicals  $C_6H_7O_7\text{-a}$  and  $C_6H_7O_7\text{-b}$ . The peroxy radical isomer from  $C_1$  (Fig. 4 c) lacks an obvious labile H-atom and is unlikely to undergo further autoxidation under atmospheric conditions. This is different for the peroxy radical isomer that forms along  $C_2$  (Fig. 4 b), which can undergo two rapid 1,6 H-shift, identical to those shown for benzene BPR in Fig. 4, to produce  $C_6H_7O_{11}$  peroxy radicals. Note that a fraction of the alkyl radical following the first 1,6 H-shift could also terminate to a carbonyl compound by reacting with  $O_2$ . It is difficult to accurately estimate the branching between  
240 radical termination and further autoxidation, so we assume a 50:50 ratio here. Termination of the autoxidation mechanism is considered through reactions with  $HO_2$  and  $NO$ , forming the hydroperoxide or the nitrated equivalent of the formed peroxy radicals, able to partition. The oxidation of intermediate volatile compounds formed during this process (such as the closed-shell species  $C_6H_6O_3$ ) is incorporated into the model by adding its first-generation products to the mechanism using the GECKO-A algorithm (Aumont et al., 2005), allowing for the contribution of these products to SOA formation to be accounted  
245 for in the quasi-explicit mechanism. Based on these findings, geminal diol BPR rearrangement pathways and the associated products are incorporated into the quasi-explicit benzene mechanism.



**Figure 4.** Oxidation schemes of phenol- and catechol-derived geminal diol BPRs. a) Molecular rearrangement of the phenol-derived BPR leads to the formation of compounds with 8 oxygen atoms. b,c) Molecular rearrangement of the two catechol-derived BPRs. One leads to compounds with up to 11 oxygen atoms, while the other is unlikely to oxidize beyond 7 oxygen atoms due to a lack of labile H-atoms.



## 4 Evaluation by comparison to chamber measurements

In order to evaluate the quasi-explicit mechanism developed and its ability to reproduce the formation of SOA, model simulations were compared to experimental results obtained from atmospheric chamber experiments from Choi et al. (2024).  
250 The simulations were performed using the SSH-aerosol v2.0 box model, explicitly representing gas–particle partitioning and multiphase chemical processes (Sartelet et al., 2026).

### 4.1 Presentation of the chemical mechanisms

In the evaluation, several chemical schemes are compared, including the original MCM v3.3.1 mechanism, which serves as the baseline for the developments presented here, and four additional mechanisms designed to isolate the effects of the newly  
255 implemented pathways and associated uncertainties. An overview of the chemical mechanisms considered in this work is provided in Table 2. Mech 1 corresponds to MCM v3.3.1 with revised pathways, addition of multi-hydroxylation, peroxy/alkoxy autoxidation, cyclic epoxides formation and BPR rearrangement, as detailed in section 2. In addition to these pathways, the formation and rearrangement of geminal diol BPRs is taken into account in Mech 2 and Mech 3. The Mech 2 and Mech 3 mechanisms differ in the branching ratios between H-abstraction and OH addition for the OH reaction with catechol, as well  
260 as for subsequent reactions involving more highly hydroxylated species formed after further OH addition. Finally, Mech 4 differs from Mech 2 in the endocyclization branching ratios, as described in section 3.2.2.

**Table 2.** Summary table of the benzene oxidation schemes considered in this study. The *abs.-add.* column describes branching ratios used for OH addition and H abstraction in reactions of phenols with OH. Branching ratio is abbreviated as BR in the header.

Name	Pathways	BR abs.-add.	BR endocycl.	Nb reactions	Nb species
MCM	benzene MCM scheme V3.3.1	-	-	334	135
Mech 1	MCM revised + multi-hydroxylation, autoxidation, epoxides, BPR rearrangement	7 % - 93 %	-	1 494	1 102
Mech 2	Mech 1 + geminal diol BPR rearrangement	7 % - 93 %	50 %	1 766	1 162
Mech 3	Mech 1 + geminal diol BPR rearrangement	30 % - 70 %	50 %	1 766	1 162
Mech 4	Mech 1 + geminal diol BPR rearrangement	7 % - 93 %	10 %	1 766	1 162

### 4.2 Presentation of the experiments

The set of 16 atmospheric chamber experiments described in Choi et al. (2024) covers a broad range of chemical conditions designed to represent aromatic oxidation and SOA formation. The experiments include two precursors: phenol (experiments  
265 1–8) and benzene (experiments 9–16) and explore variability in NO<sub>x</sub> levels, as well as the presence, acidity and concentrations in inorganic seed aerosol. For each precursor, the first two experiments are conducted without seed aerosol, the next two experiments include acidic inorganic seeds and the final four employ neutral seed aerosols. Additional details on NO<sub>x</sub> regimes, seeds nature and meteorological data used are provided in Table 3.



**Table 3.** Experimental conditions of the phenol and benzene simulations, including NO<sub>x</sub> conditions, initial aerosol seeds, and meteorological data.

Phenol exp.	NO <sub>x</sub> regime	Seed type	Meteo. data <sup>1</sup>	Benzene exp.	NO <sub>x</sub> regime	Seed type	Meteo. data <sup>1</sup>
1	high	no seed	detailed	9	low	no seed	detailed
2	low	no seed	range	10	high	no seed	detailed
3	high	acidic	detailed	11	low	acidic	detailed
4	low	acidic	range	12	high	acidic	detailed
5	high	neutral	range	13	low	neutral	detailed
6	high	neutral	range	14	high	neutral	detailed
7	low	neutral	range	15	low	neutral	detailed
8	low	neutral	range	16	low	neutral	detailed

<sup>1</sup> Detailed meteorological data correspond to time-resolved temperature and relative humidity provided by the experimental team (Choi et al., 2024). When only a range minimum and maximum values were available, simulations were performed using mean values.

The experiments are conducted over 8 hours under natural daylight, with radical production driven by photolysis. To ensure measurable oxidation and SOA formation within the chamber timescale, the initial precursor concentrations are substantially higher than ambient levels, particularly for benzene, which reacts relatively slowly under atmospheric conditions.

### 4.3 Model presentation and simulation setup

The simulations were performed using the SSH-aerosol V2.0 box model (Sartelet et al., 2026), a multiphase chemistry–aerosol box model designed to simulate the evolution of gas-phase chemistry and aerosol composition. Aerosol dynamics and multiphase chemical processes are explicitly represented. Semi-volatile organic compounds may partition between the gas, organic and aqueous phases depending on their physicochemical properties and ambient conditions. The quasi-explicit benzene oxidation mechanism is integrated in the model in order to compare simulated SOA formation from benzene and phenol to the chamber experimental results.

Experimental conditions described in Choi et al. (2024) are used to constrain the initial state of the simulations. The simulations are conducted over eight hours with outputs every five minutes. For all benzene-based experiments, as well as experiments 1 and 3 for phenol, meteorological conditions (temperature, humidity) were provided by Choi et al. (2024) and updated in the simulations every five minutes. For the remaining experiments, the temperature and relative humidity are prescribed at each time step using the mean values of the ranges reported in the experiment descriptions. Photolysis rates are calculated based on the location of the chamber and the date of the experiments. An attenuation coefficient is applied to these rates to represent the impact of clouds. This coefficient is adjusted so that the amount of precursor consumed at the end of the simulation corresponds to the one obtained experimentally. Initial precursor concentrations range from 79 ppb to 264 ppb for phenol and from 168 to 325 for benzene. Nitrous acid is introduced as an OH radical source in the benzene experiments only. The initial NO<sub>x</sub> concentration indicated for each experiment is divided between NO and NO<sub>2</sub> to correspond to the initial concentrations

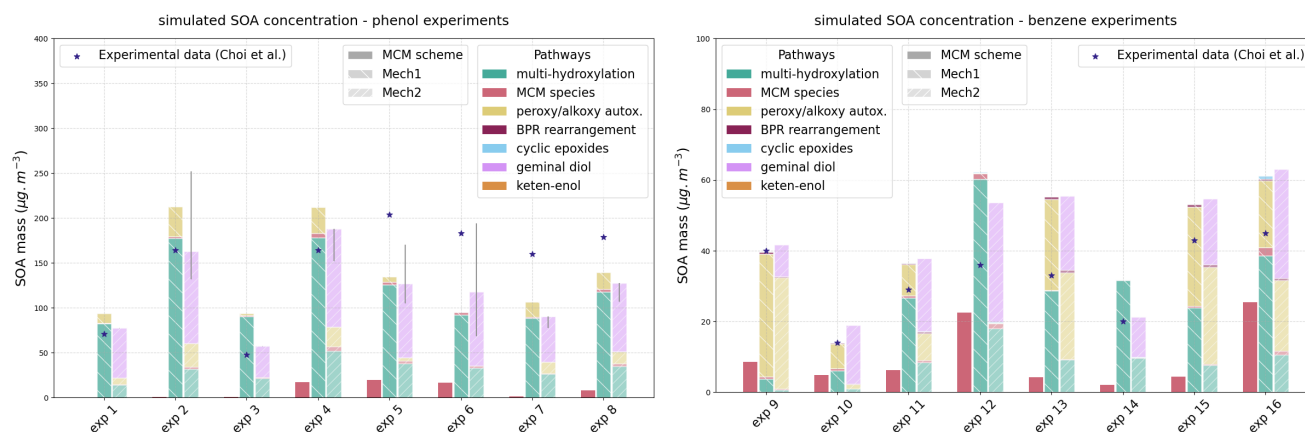


presented in the graphs of Choi et al. (2024). The concentrations of inorganic aerosols introduced at the initial state range from 62 to 2201  $\mu\text{g m}^{-3}$  in the phenol experiments and from 10 to 377  $\mu\text{g m}^{-3}$  in the benzene experiments.

Losses on the simulation chamber walls are included using the parameterizations included in SSH-aerosol (Sartelet et al., 2026). Semi-volatile gas-phase compounds undergo reversible wall uptake controlled by kinetic transfer and wall-gas partitioning. The model parameters are set to  $\frac{S}{V} = 1.654 \text{ m}^{-1}$  for the surface over volume ratio, and to  $k_e = 0.1 \text{ s}^{-1}$  for the eddy diffusivity coefficient (Huang et al., 2018). Particle losses are treated as irreversible first-order processes accounting for diffusion, settling, and turbulence. Wall-loss rates are determined from compound properties and fixed chamber parameters (geometry and eddy diffusivity), assuming a 104  $\text{m}^3$  chamber. To better represent NOx chemistry, a wall loss parametrization of NO2 representing surface adhesion is implemented in the model following Fiorentino et al. (2021). In this parametrization, the heterogeneous loss of NO2 is represented via a first-order surface uptake process, which depends on an uptake coefficient  $\gamma_{\text{NO}_2}$  (assumed to be  $4.5 \times 10^{-6}$  as in Fiorentino et al. (2021)), on the temperature and the ratio surface over volume of the chamber.

#### 4.4 Relative influence of the different pathways

Simulated SOA concentrations obtained for the 16 chamber experiments are shown and compared with measurements in Fig. 5. Results are presented for three chemical schemes: the reference MCM mechanism, the quasi-explicit Mech 1, and the Mech 2 mechanism, which includes geminal diol chemistry. Experimental observations are indicated by stars.



**Figure 5.** Simulated SOA concentrations obtained with the MCM (left bars for each experiment), Mech 1 (middle bars), Mech 2 schemes (right bars), and comparison to experimental concentrations (stars on the graph). The top panel represents experiments using phenol as the precursor, while the bottom panel shows experiments using benzene. The sensitivity to meteorological data is indicated by gray error bars when detailed temperature and relative humidity data is missing

Across all 16 experiments, simulations using the MCM scheme systematically strongly underestimate SOA concentrations. These results are consistent with previous studies reporting the lack of representation of HOMs in aromatics chemistry, and



motivate the development of quasi-explicit schemes in the first instance. The SOA concentrations are much better represented using the Mech 1 and Mech 2 schemes, within a factor 2 of the experiments, and in most cases within 30 %.

310 The addition of successive hydroxylation steps, radical autoxidation and geminal diol pathway introduces low-volatility compounds, greatly increasing SOA concentrations. These pathways are dominating SOA formation for all experiments, autoxidation being more important when NO<sub>x</sub> levels are low (i.e., experiments 9, 13, 15 and 16) and barely represented in high-NO<sub>x</sub> conditions (experiments 12, 14). The rearrangement pathway from benzene BPR has a low impact on SOA composition due to unfavorable kinetic rate and competition with NO and HO<sub>2</sub> reactions.

315 Mech 1 tends to slightly overrepresent SOA formation for benzene experiments at low-NO<sub>x</sub> levels such as in the experiments 13, 15 and 16, in which dimers from accretion reactions in the autoxidation scheme account for a large proportion of the simulated SOA. As observed by (Choi et al., 2024), experiments conducted with acidic seeds (3, 4, 11 and 12) are also overestimated, with the successive hydroxylation pathway dominating SOA formation. The formation of SOA from phenol using neutral seeds (experiments 5 to 8) is strongly underestimated, with simulations yielding SOA concentrations that are up to 1.9 times lower than the experimental values (experiment 6). The gas phase contains numerous compounds of intermediate  
320 volatility, notably resulting from the degradation of products formed by the successive oxidation pathway such as C<sub>6</sub> carbonyls and quinones resulting from the reaction of catechol with OH. Uncertainties in the gas/particle partitioning of those compounds may partly explain the discrepancy. The hydration of light aldehydes in the aqueous phase could lead to the formation of additional SOA when seed concentrations are high. Under high relative humidity, this mechanism may partly explain the differences between simulated and observed SOA concentrations in experiments 5, 6, and 8.

325 The introduction of geminal diol chemistry in Mech 2 significantly modifies SOA formation and composition. For most experiments, the concentrations simulated using Mech 2 better reproduce the amount of SOA obtained experimentally. As with Mech 1, low-NO<sub>x</sub> concentrations tend to favor autoxidation while successive hydroxylation and its nitrogen derivatives are more prevalent when NO<sub>x</sub> concentrations are high. In all experiments, the reaction pathway forming geminal diols contributes significantly to the formation of SOA: geminal diol-derived BPRs rearrangement directly competes with the successive hydroxylation pathway and tends to reduce the amount of SOA formed. The addition of this oxidation pathway appears to improve  
330 the representation of SOA formation in experiments where Mech 1 overestimate SOA, particularly under conditions in which multi-hydroxylation played a significant role, most notably at high NO<sub>x</sub> conditions (e.g. exp 3 and 12).

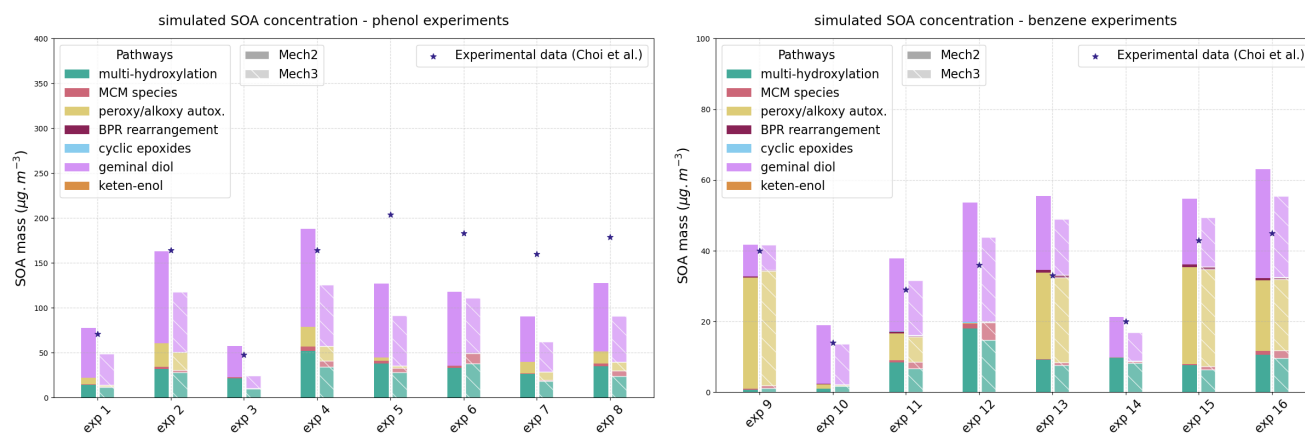
#### 4.5 Sensitivity analysis

335 The simulated concentrations are affected by uncertainties related to model choices and to the representation of environmental conditions. Here, the sensitivity of aromatic oxidation to the competition between OH addition and abstraction pathways is assessed (section 4.5.1), the influence of meteorological conditions (temperature and relative humidity) for experiments where detailed temporal variations are not available (section 4.5.2), the influence of glyoxal gas/particle partitioning (section 4.5.3) and of the endocyclization branching ratio in the geminal diol BPR rearrangement pathway (section 4.5.4). Further studies conducted on the influence of non-ideality and on nitrophenol saturation vapor pressure are presented in Supplement S6 and  
340 S7.



### 4.5.1 Influence of addition/abstraction branching ratios

To quantify the sensitivity of aromatic oxidation to the competition between OH addition and abstraction, the chemical schemes Mech 2 and Mech 3 described in section 4.1 are compared. For each experiment, the left bars represent the scheme with a 7 % abstraction branching ratio for reactions of benzene and phenols with OH whereas the right bars represent the 30 % abstraction  
 345 scheme.



**Figure 6.** SOA concentrations simulated using Mech 2 and Mech 3 and comparison to experimental concentrations (stars on the graph). The left panel represents experiments using phenol as the precursor, while the right panel shows experiments using benzene.

In Mech 3, the increase in the branching ratio in favor of abstraction leads to a decrease in the amount of SOA formed under most of the experimental conditions. Diverting the initial oxidation toward abstraction lowers the yield of these SOA-forming pathways, especially under high-NO<sub>x</sub> conditions where abstraction competes with the formation of low-volatility multi-hydroxylated compounds and organonitrates. In low-NO<sub>x</sub> conditions, SOA formation is mainly driven by autoxidation,  
 350 peroxy and alkoxy radicals formation and accretion reactions, less dependent on multiple OH additions, lowering the impact of the increased abstraction.

### 4.5.2 Uncertainty on meteorology

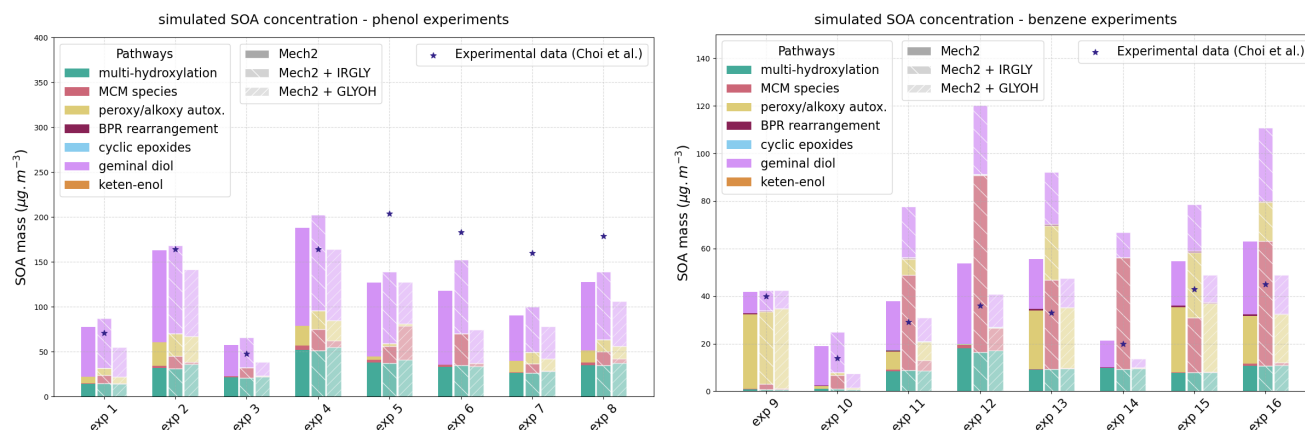
For all benzene-based experiments, as well as experiments 1 and 3 for phenol, the temporal variations of meteorological conditions (temperature, humidity) were provided. However, only mean values were prescribed in the other experiments.  
 355 To assess the impact of these uncertainties on temperature and humidity on the concentrations, the phenol experiments are simulated using the minimum and maximum temperature and humidity observed during the experiments. The error bars shown in Fig. 5 for experiments 2 and 4 to 8 represent the maximum and minimum SOA concentrations obtained using these extreme values. Experiments 2, 5, and 6 show the highest sensitivity to meteorological variability, associated with large variations in relative humidity and temperature over the course of the experiments. This sensitivity likely contributes to the underestimation



360 of SOA concentrations in these cases when average meteorological conditions are assumed. Conversely, experiment 7 shows little sensitivity to variations in temperature and relative humidity due to the dry nature of the seeds and the average relative humidity level being representative of extreme conditions.

### 4.5.3 Influence of glyoxal gas/particle partitioning

365 In SSH-aerosol, glyoxal can partition through its dissolution into the particle aqueous phase, where rapid hydration shifts the gas–particle equilibrium toward the condensed phase. In addition to this reversible partitioning process, the model includes an irreversible parametrized reaction that converts glyoxal into a non-volatile product. The impact of hydration and of the irreversible parametrized reaction on the simulated concentrations of the chamber experiments are illustrated in Fig. 7.



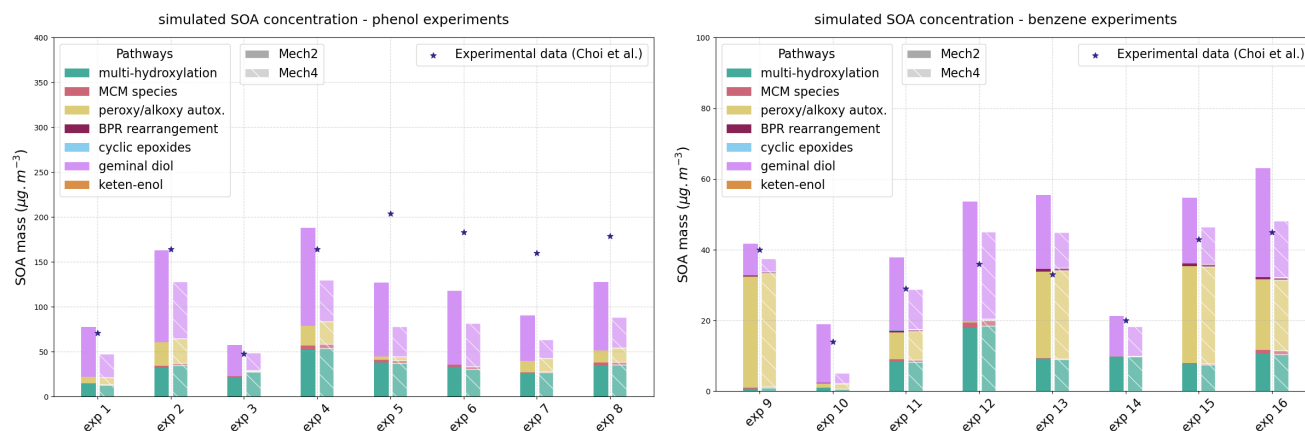
**Figure 7.** SOA concentrations simulated using Mech 2, Mech 2 + IRGLY taking into account the irreversible partitioning of glyoxal and Mech 2 + GLYOH taking into account glyoxal hydration. The left panel represents experiments using phenol as the precursor, while the right panel shows experiments using benzene.

The implementation of the parameterization for irreversible glyoxal partitioning leads to increased SOA formation under all experimental conditions and to an overestimation of concentrations, especially for benzene experiments conducted in the presence of seeds (Experiments 11 and 12). Experiments using phenol as a precursor also show an increase in SOA concentrations, although to a much lesser extent. Most glyoxal formation results from the decomposition of the benzene bicyclic alkoxy radical, whose formation pathway competes with phenol formation.

375 On the contrary, the hydration of glyoxal leads to an overall decrease in SOA formation. The formation of hydrated and dihydrated glyoxal increases its uptake in the particulate phase and reduces the availability of glyoxal in the gas phase. This has the effect of reducing the oxidative capacity of the environment as well as the formation of HOMs via benzene oxidation, since the photolysis of glyoxal in the gas phase leads to the formation of HO<sub>2</sub>. This reaction allows for a more accurate representation of SOA concentrations in cases previously overestimated, particularly experiments 4, 13, 15, and 16.



#### 4.5.4 Influence of the endocyclization branching ratio



**Figure 8.** SOA concentrations simulated using Mech 2 and Mech 4. The left panel represents experiments using phenol as the precursor, while the right panel shows experiments using benzene.

Sensitivity simulations are performed using Mech 4 to assess the impact of the branching ratio leading to endocyclization and to compound C6H6O3 in catechol oxidation. Reducing the endocyclization branching ratio from 50 % to 10 % leads to an important decrease in simulated SOA concentrations across most experimental conditions, as seen in Fig. 8. This reduction is due to a lower yield of HOMs as a larger fraction of the oxydated precursor mass is redirected toward C6H6O3, whose first oxidation products are more volatile. The reduction in SOA mass formed is particularly pronounced in experiments where SOA formation is strongly influenced by geminal diol chemistry, for which the reduced endocyclization branching leads to a substantial underestimation of aerosol mass relative to chamber observations. Overall, the comparison indicates that the 50 % approximation on endocyclization branching ratio provides a consistent representation of SOA mass observed experimentally.

## 5 Reduced scheme for 3D atmospheric modeling

The quasi-explicit oxidation mechanisms developed in this work provide a detailed description of the chemical processes controlling benzene SOA formation but remain computationally too expensive for direct implementation in three-dimensional chemical transport models. It is reduced with the GENERator of reduced Organic Aerosol mechanisms (GENOA), which is an automated framework designed to generate reduced chemical schemes from explicit mechanisms by retaining species and reactions based on their impact on SOA formation. The reduced chemical schemes enable the integration of aerosol-relevant chemistry into air-quality models while maintaining consistency with the detailed quasi-explicit mechanism.



## 5.1 GENOA model presentation

395 The GENOA v2 algorithm (Wang et al., 2022) is designed to reduce the size of oxidation mechanisms while maintaining their ability to reproduce SOA formation across a range of VOC precursors and atmospheric conditions, thereby enabling access to detailed SOA composition in 3D simulations. The reduction process relies on several elementary reduction strategies, including removal of species and/or reactions, jumping of an intermediate species, lumping of two similar compounds, and replacement of one by the other. These operations are applied iteratively to the input quasi-explicit scheme and generate a sequence of  
400 intermediate simplified mechanisms converging toward a final reduced scheme. After each reduction step, the intermediate mechanism is tested using the 0D box model SSH-aerosol under a set of European atmospheric conditions, to ensure its accuracy compared to the quasi-explicit scheme. User-defined errors control the deviation tolerated from the reference scheme, allowing the generation of reduced mechanisms with different levels of complexity depending on the intended application.

As in Wang et al. (2022) and Sartelet et al. (2024), the GENOA reduction framework relies on a hierarchical evaluation  
405 strategy based on three complementary datasets. Throughout the reduction procedure, an eight-condition core dataset is used to evaluate candidate reduced mechanisms, combining six chemically extreme conditions from Wang et al. (2022) and two additional high-NO<sub>x</sub> urban conditions from Sarica et al. (2023). To better reflect typical atmospheric variability and test robustness beyond this core set, the evaluation dataset is then expanded to 152 conditions. Finally, a comprehensive testing dataset of 9,433 European conditions is used for the final assessment of the reduced mechanism. For all datasets, box-model  
410 simulations are performed over five days with different starting times (12 am, 7 am, 12 pm, and 8 pm) to represent both daytime and nighttime chemistry and to capture SOA formation and aging.

## 5.2 Mechanism reduction

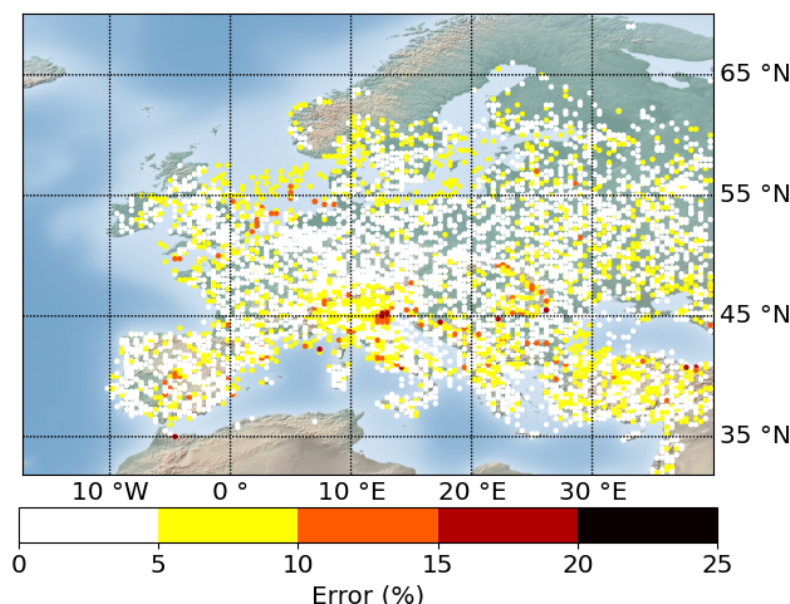
The reduction is carried out in two successive stages. In the first stage, only four reduction strategies are applied in order to simplify the mechanism while largely preserving its original chemical structure and interpretability. During this stage, the  
415 tolerated average error relative to the reference mechanism ( $\epsilon_{\text{mean}}$ ) is gradually increased from 0.5 % to 5 %, while the maximum tolerated error ( $\epsilon_{\text{max}}$ ) increases from 1 % to 30 %. At the end of the first reduction cycle, the mechanism contains 76 reactions and 70 species, including 35 condensable species. It reproduces SOA mass concentrations with an average error of 1.78 % and a maximum error of 13.32 % over the 9433 testing conditions.

In the second reduction cycle, additional lumping and jumping strategies are introduced with the specific goal of minimizing  
420 the number of condensable species. During this cycle,  $\epsilon_{\text{mean}}$  is allowed to increase up to 10 %, while  $\epsilon_{\text{max}}$  remains at 30 %. The second reduction cycle results in a strongly reduced mechanism, retaining only 30 reactions, 28 species, including 12 condensable, which represents 2.6 % of the original reactions and 1.2 % of the original condensable species.

The reduced mechanism resulting from the second reduction cycle reproduces SOA mass concentrations predicted by the Mech 2 mechanism within 3.94 % on average over the 9433 tested conditions, and a maximum error of 22.5 % indicating  
425 that the main processes controlling SOA formation from benzene oxidation are well represented and preserved in the reduced scheme. The spatial distribution of the relative error between the reference mechanism Mech 2 and the reduced mechanism is

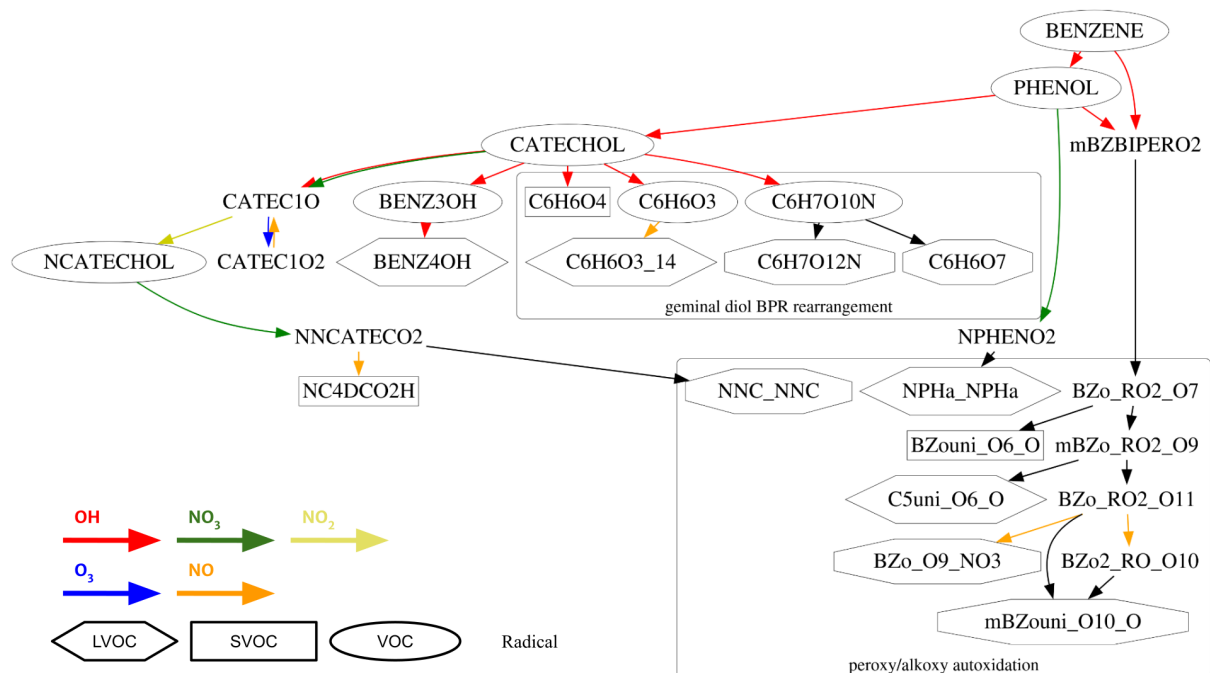


shown in Fig. 9. Over most locations, errors remain below 5 %. Errors exceeding 10 % are confined to a limited number of locations, while errors above 15 % are rare and are mainly found over North Italy.



**Figure 9.** Geographical repartition of the error between the quasi-explicit Mech 2 and the reduced mechanisms.

As illustrated in Fig. 10, the reduced mechanism preserves the main oxidation pathways contributing to SOA formation and  
430 identified in section 4. It contains only 12 condensable species. Regarding the oxidation pathways, the successive hydroxyla-  
tion, the autoxidation of alkoxy and peroxy radicals and the geminal diol BPR rearrangement pathways are retained. Successive  
hydroxylation processes lead to the formation of BENZ3OH and BENZ4OH. In high-NO<sub>x</sub> conditions, organonitrates forma-  
tion is explicitly represented with the formation of nitrocatechol ultimately leading to the highly oxygenated NC4DCO2H  
or to the NNC\_NNC and NPHa\_NPHa dimers from accretion reactions. As developed in section 3.2, catechol oxidation and  
435 geminal diol BPR rearrangement quickly yields condensable compounds such as species C6H6O4, C6H6O7 and C6H7O12N  
of the reduced mechanism. Other species represented on the graph result from autoxidation of benzene and phenol BPRs. Ter-  
mination of the alkoxy and peroxy compounds resulting from the oxidation of phenol BPR is represented through condensable  
compounds such as C5\_uni\_O6\_O, C5\_uni\_O6\_O, BZo\_O9\_NO3, BZouni\_O6\_O and mBZouni\_O10\_O.

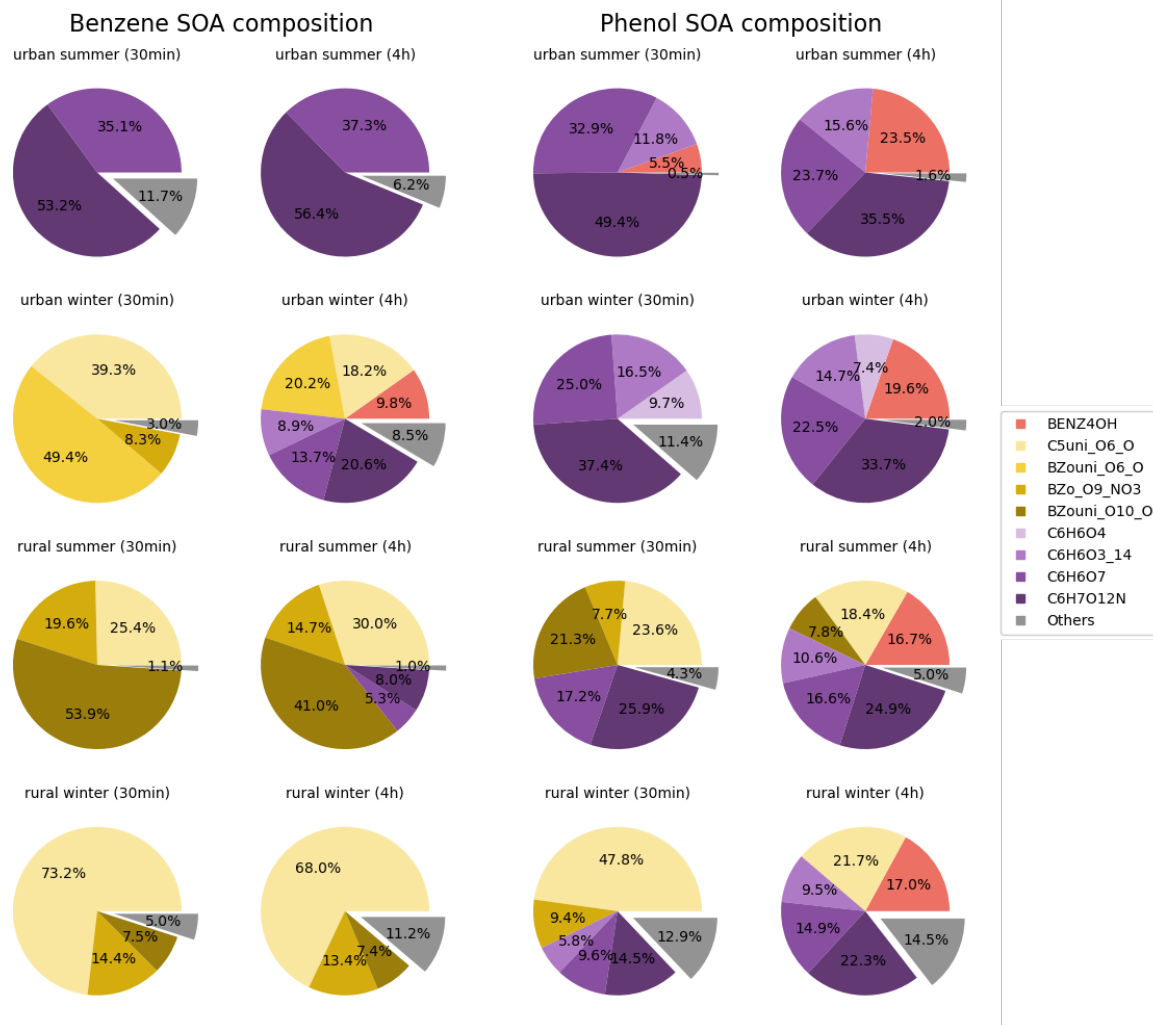


**Figure 10.** Graphical representation of the reduced scheme. Compounds shown with oval frames are VOCs ( $P^{sat} > 10^{-4}$  atm), rectangular frames are SVOCs ( $10^{-4} > P^{sat} > 10^{-9}$  atm), hexagons are LVOCs ( $10^{-9} > P^{sat} > 10^{-13}$  atm) and frameless compounds are radicals. Arrow color indicates the oxidant involved in the reactions.

## 6 Evaluation of the main chemical pathways of benzene and phenol SOA formation

440 Identifying the major oxidation products formed under atmospherically relevant conditions is essential for improving our understanding of SOA formation and evolution. Beyond the nature of the products formed, their temporal evolution provides key insight into the dominant chemical pathways, allowing rapidly formed species associated with early oxidation stages to be distinguished from compounds that accumulate or persist on longer timescales. To address this, we analyze oxidation product distributions using box-model simulations conducted with SSH-aerosol under constant oxidant concentrations representative of either rural or urban conditions. Fig. 11 compares the benzene and phenol SOA composition after short (30 min) and longer  
 445 of either rural or urban conditions. Fig. 11 compares the benzene and phenol SOA composition after short (30 min) and longer (4 h) simulation times in a winter/summer and rural/urban conditions and beginning at 12pm, to assess the compounds that are formed efficiently in the early stages of oxidation and those that dominate the system at later stages.

The simulations show significant contrasts between urban and rural environments and exhibit a strong seasonal dependence. Stronger photochemical activity and higher OH concentrations in summer favor SOA formation via the geminal diol pathway, whereas autoxidation becomes relatively more important in rural environments than in urban settings. For benzene, in rural conditions and in urban winter conditions, SOA formation is initially dominated by autoxidation, leading to efficient formation  
 450 of highly oxygenated products at early oxidation times. In winter, reduced photochemical activity favor RO<sub>2</sub> self-reactions and



**Figure 11.** Benzene and phenol SOA composition after short (30 min) and longer (4 h) simulation times in a winter/summer and rural/urban conditions. Compounds representing under 5 % of the SOA mass are aggregated in the "Others" category.

intramolecular H-shifts, leading to a dominant contribution of autoxidation products to SOA mass, even in urban conditions. In contrast, urban summer environments exhibit an important influence of the geminal diol pathway. Over time, competition emerges between the direct autoxidation pathways of benzene and the contribution of phenol and catechol via autoxidation along the geminal diol pathway, resulting in an important change in SOA composition between the early stage of oxidation and more aged conditions, especially for urban winter conditions. For phenol, autoxidation is less dependent on seasonal effects, being negligible in the urban environment and accounting for more than 50 % at early stages of oxidation in the rural case. As seen previously for benzene, a substantial fraction of the phenol SOA originates from the geminal diol pathway, with this contribution being particularly pronounced under urban conditions and increasing over time.



## 7 Conclusions

In this work, we developed a quasi-explicit oxidation mechanism for the formation of benzene and phenol SOAs. This mechanism builds on MCM v3.3.1 and oxidation pathways that are overly simplified or missing are introduced, most notably geminal diol BPRs formation and rearrangement, successive hydroxylation, and peroxy radicals autoxidation, providing a more realistic  
465 description of how benzene oxidation progresses toward low-volatility products and SOA formation.

Both comparisons to chamber experiments and simulations in atmospheric conditions show that these pathways play a central role in determining both the concentrations and the molecular composition of benzene and phenol-derived SOAs. Under most conditions, the formation of SOA from benzene is dominated by rearrangement of geminal diol BPRs, accounting on average for 50.0 % of the mass across benzene experiments and 62.8 % for phenol. Successive hydroxylation and peroxy  
470 radicals autoxidation, more dependent on NO<sub>x</sub> and OH conditions, account on average for 19.4 % and 28.5 % of benzene SOA formation, respectively. These added pathways allow for a greater precision on modeled SOA concentration, the benzene MCM mechanism reproducing the experimental values with a mean relative error of 83.2 % compared to 28.2 % for the developed mechanism.

To bridge the gap between chemical detail and large-scale modeling, the quasi-explicit mechanism was reduced using the  
475 GENOA algorithm, from 1 772 reactions and 1 160 species to 30 reactions and 28 species. Despite this strong reduction in complexity, the reduced mechanism preserves the three main oxidation pathways controlling SOA formation with an average precision of 3.9 %, making it suitable for implementation in three-dimensional air-quality models while remaining consistent with the underlying chemistry.



480 *Code and data availability.* The code SSH-aerosol v2.0 is available in the github platform (<https://github.com/sshaerosol/ssh-aerosol>, last access: 4th June 2026), and in <https://doi.org/10.5281/zenodo.14196277> (Sartelet et al., 2026)

*Author contributions.* ALB and KS designed the research. ALB, KS, SI and AO contributed to the development of the chemical mechanisms considered in this study. ALB and KS conducted the formal analysis. AO and SI performed and analysed the quantum chemistry calculations. ALB performed the box-model simulations and mechanism reduction, and visualized the data. KS and RC supervised the research. ALB, AO, SI and KS wrote the manuscript draft. ZW, VL, FC and RC reviewed and edited the manuscript.

485 *Competing interests.* The authors declare they have no competing interest.

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490 processes.



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