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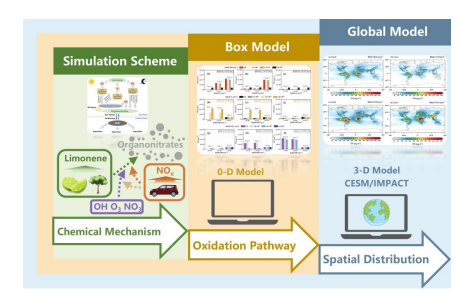


- Competing multiple oxidation pathways shape
- 2 atmospheric limonene-derived organonitrates simulated
- 3 with updated explicit chemical mechanisms
- 4 Qinghao Guo<sup>1</sup>, Haofei Zhang<sup>2</sup>, Bo Long<sup>3</sup>, Lehui Cui<sup>1</sup>, Yiyang Sun<sup>1</sup>, Hao Liu<sup>1</sup>, Yaxin
- 5 Liu<sup>1</sup>, Yunting Xiao<sup>1</sup>, Pingqing Fu<sup>1</sup> and Jialei Zhu<sup>1,\*</sup>
- 6 1 Institute of Surface-Earth System Science, School of Earth System Science, Tianjin University, Tianjin, 300072, China;
- 7 2 Department of Chemistry, University of California, Riverside, California 92521, USA;
- 8 3 College of Materials Science and Engineering, Guizhou Minzu University, Guiyang 550025, China.
- 9 Correspondence to: Jialei Zhu, Email: zhujialei@tju.edu.cn

Abstract. Organonitrates (ON) are key components of secondary organic aerosols (SOA) with potential environmental and climate effects. However, ON formation from limonene, a major monoterpene, and its sensitivity to oxidation pathways remain insufficiently explored due to the absence of models with explicit chemical mechanisms. This study advances the representation of limonene-derived ON formation by incorporating 90 gas-phase reactions and 39 intermediates across three oxidation pathways (O<sub>3</sub>, OH, NO<sub>3</sub>) into both a chemical box model and a global model. Box model sensitivity experiments revealed that competition among major oxidation pathways, coupled with the high yield of limonenederived ON from O3-initiated oxidation, leads to increased limonene-derived ON production when the O3-initiated pathway is enhanced, whereas strengthening the OH- or NO3-initiated pathways reduces ON formation. Compared to the box model, the global simulation exhibits stronger nonlinear responses and great spatiotemporal variability in limonene-derived ON formation across different oxidation pathways. This is primarily driven by the complex distribution of precursors and oxidants, as well as changing in dominate chemical pathways under various meteorological conditions. In the presence of the other two pathways, increasing the O3- or NO3-initiated oxidation pathway reduces the global limonene-derived ON burden by 19.9% and 17.3%, respectively, whereas enhancing the OH-initiated pathway increases it by 44.7%. limonene-derived ON chemistry developed in this study not only enhances the global model's ability to simulate ON formation evaluated through comparison with observations but also demonstrates an approach based on explicit chemical mechanisms that establishes a methodological framework for simulating the chemical formation processes of SOA.







## 1 Introduction

Secondary organic aerosols (SOA) represent a substantial fraction of fine particulate matter and contribute to global public health risk, deterioration of air quality and climate change (Collaborators, 2024; Lelieveld et al., 2015; Tao et al., 2017). Among chemical constituents, organonitrates (ON) are of particular interest owing to their large fraction in SOA (5%-77%) (Farmer et al., 2010; Kiendler-Scharr et al., 2016). The rate of particulate ON formation contributes strongly to the rate of SOA formation at night, which emphasizes the important roles of particulate ON in ambient SOA (Guo et al., 2024). The nitrate group in ON would influence the physical and chemical properties of SOA, such as decreasing saturated vapor pressure of the product molecule (Capouet and Müller, 2006). ON are secondary compounds formed via the oxidation of volatile organic compounds (VOCs) in the presence of nitrogen oxides (NO<sub>x</sub> = NO + NO<sub>2</sub>), substantially influencing NO<sub>x</sub> cycling and formation of ozone (O<sub>3</sub>) and HONO (Perring et al., 2013). In global scale, VOCs are mainly emitted from biogenic sources, while NO<sub>x</sub> are emitted from a wide variety of anthropogenic sources (Ng et al., 2017; Glasius and Goldstein, 2016). Therefore, a thorough investigation of ON is warranted to advance our understanding of interaction between biogenic and anthropogenic emissions.

The chemical formation mechanisms of ON are complex, hampering efforts to simulate and control SOA. In the daytime, hydroxyl radicals (OH) and ozone  $(O_3)$  oxidation of VOCs can produce peroxy





47 radical (RO<sub>2</sub>), which reacts with NO<sub>x</sub> to produce ON (Perring et al., 2013), while the reaction between 48 nitrate radicals (NO<sub>3</sub>) and VOCs dominates the generation of ON in the nighttime (Rollins et al., 2009; 49 Perring et al., 2013; Ng et al., 2017). Furthermore, the coexistence among OH, O<sub>3</sub> and NO<sub>3</sub> has been 50 investigated in VOCs nocturnal oxidation (Brown and Stutz, 2012; Barber et al., 2018; Kwan et al., 2012; 51 Chen et al., 2022). Compared with single oxidant, the introduction of multiple oxidants is evaded for the 52 possible complex reaction mechanisms for VOCs. The regeneration of OH would change the O<sub>3</sub> 53 oxidation process to form SOA (Sato et al., 2013). Chamber experiments show that SOA from NO<sub>3</sub> 54 oxidation of VOCs are affected by oxidation of NO2 by O3 (Ng et al., 2017). Therefore, VOCs are 55 oxidized through the synergistic effects of multiple oxidants, driving the chemical formation of ON. 56 However, ON formation from the VOCs oxidation governed by mixing oxidants has not been fully 57 understood. In particular, the impact of oxidation pathways on the ON formation and spatial distribution 58 are still unclear. 59 As one of typical biogenic volatile organic compounds (BVOCs) (10% of monoterpenes), limonene 60 is mostly emitted from citrus plants and coniferous trees, with a total emission rate of ~11 Tg·yr<sup>-1</sup> 61 (Guenther et al., 2012; Sindelarova et al., 2014). Limonene has unique structure with an endocyclic 62 double bond and an exocyclic double bond, which makes it reactive towards atmospheric oxidants 63 (Surratt et al., 2008). Higher ON (30-72%) and SOA yields (17-40%) through NO<sub>3</sub>-initiated oxidation 64 of limonene than other monoterpenes have been observed in laboratory experiments (Fry et al., 2014; 65 Hallquist et al., 1999; Spittler et al., 2006; Moldanova and Ljungström, 2000; Fry et al., 2011). It has 66 been well demonstrated that limonene + NO<sub>3</sub> is most important pathway to form limonene-derived ON 67 (Kilgour et al., 2024; Ehn et al., 2014; Jokinen et al., 2015; Zhao et al., 2015). Furthermore, recent study 68 found the primary nitrooxy RO2 formed through NO3 addition to limonene occurs at both at endocyclic 69 double bond and the exocyclic double bond. These products could undergo autoxidation, which is fast 70 enough to RO<sub>2</sub> bimolecular reactions (Mayorga et al., 2022). The molecular compositions and formation 71 mechanism of limonene-derived ON have been well investigated through observations and laboratory 72 studies, while their description in models remains not explicit and advanced. 73 The early atmospheric model utilizes empirical yields and empirical coefficients for predicting 74 limonene-derived SOA production in simulation (Yu et al., 2019). Currently, chemical mechanisms are 75 simplified according to analogies with structurally similar compounds in most of regional and global





models due to simplicity and efficiency in calculation (Fisher et al., 2016; Li et al., 2023a). Nevertheless, previous model studies have not included the formation mechanism of limonene-derived ON in detail (Pye et al., 2015; Li et al., 2023a; Zare et al., 2019). Simplified mechanisms make it difficult to understand explicit limonene-derived ON formation process and the influence of interaction between multiple oxidation pathways on ON formation.

Herein, we investigated the impacts of multiple oxidation pathways on limonene-derived ON using both chemical box model and global model, which were developed to include explicit chemical mechanisms for limonene-derived ON formation. The effect of competition among individual oxidation pathways on limonene-derived ON formation were discussed using a chemical box model based on proposed mechanisms. The simulation framework of explicit chemical mechanisms was integrated into global model to evaluate the spatial distributions of limonene-derived ON and contributions of individual oxidation pathways. This study presents a numerical simulation framework for atmospheric chemical processes and aims at enhancing the ability of models to simulate ON and understand the competition effects among atmospheric oxidation pathways on SOA formation, improving atmospheric composition

forecasts and informing interaction between biogenic and anthropogenic emissions.

#### 2 Methodology

## 2.1 Limonene-derived ON formation mechanism

In order to simulate ON via the gas-phase oxidation of limonene, the chemical mechanism used in our model was updated with gas-phase chemical mechanisms of limonene-derived ON based on recent laboratory studies (Mayorga et al., 2022) and Master Chemical Mechanism (MCM, v3.3.1). The explicit chemical mechanism of limonene-derived ON involves three initial oxidation pathways: OH-, O<sub>3</sub>- and NO<sub>3</sub>-initiated oxidation (Fig. 1). The detailed formulas of species could be found in Table S1. The updated explicit formation mechanisms were list in Fig. S1 and Table S2. Compared to the MCM mechanism, the chemical mechanism of limonene-derived ON formation used in this study is developed to include: (1) NO<sub>3</sub> addition at three different carbonsites. The branching ratios of the three C<sub>10</sub>H<sub>16</sub>NO<sub>5</sub>-RO<sub>2</sub> isomers were estimated to 0.63:0.34:0.03 based on previous laboratory studies (Leungsakul et al., 2005; Wang and Wang, 2021). (2) Sequential NO<sub>3</sub> oxidation reactions to form ON for all the products that contain double bonds from OH- and O<sub>3</sub>-initiated oxidation in MCM. The rate constants were set to





be the same as those used in MCM for limonaldehyde. (3) The formation of a ring-opened nitrooxy  $RO_2$  in the presence of  $O_2$  due to bond scission of the two endocyclic nitrooxy  $RO_2$ , and its branching ratio was estimated (Draper et al., 2019; Kurten et al., 2017; Guo et al., 2022). (4) H-shifts of the exocyclic  $C_{10}H_{16}NO_4$ -RO. (5) Bimolecular and unimolecular reactions of the  $C_{10}H_{16}NO_6$ -RO<sub>2</sub> and  $C_{10}H_{16}NO_7$ -RO<sub>2</sub>. The rate constants for the bimolecular reactions are the same as those used in MCM, and autoxidation rate constants are calculated. In our explicit chemical mechanism, more intermediates and chemical processes of limonene-derived ON were distinguished than simplified mechanisms used in previous models.

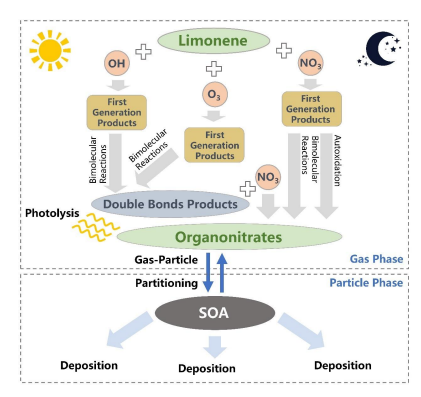


Figure 1. Schematic diagram of the limonene-derived ON formation pathways included in this work.

We assumed highly oxidated ON products ( $C_{10}H_{13}NO_7$ ,  $C_{10}H_{15}NO_4$ ,  $C_{10}H_{15}NO_5$ ,  $C_{10}H_{15}NO_6$ ,  $C_{10}H_{15}NO_7$ ,  $C_{10}H_{15}NO_8$ ,  $C_{10}H_{17}NO_5$ ,  $C_{10}H_{17}NO_6$ ,  $C_{10}H_{17}NO_7$ ) to be semi- to low-volatile which can condense into the particulate phase upon formation. Their vapor pressures are estimated to calculate gasparticle partitioning (Table S3).

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The vapor pressures of the above-mentioned ON species were estimated using the EVAPORATION method (Compernolle et al., 2011). It is one of the widely used group contribution-based models to predict molecular vapor pressures. As the model input, the structures of the ON species were obtained from prior work (Mayorga et al., 2022) and converted to SMILES strings. To validate the method, the predicted vapor pressures were also compared with another commonly used method, SIMPOL 1 (Pankow and Asher, 2008), which requires only the functional groups as inputs. The two methods predict vapor pressures within one order of magnitude in most cases, supporting our estimation.

#### 2.2 Chemical box model

A zero-dimensional (0-D) chemical box model was used to examine the chemical processes of limonenederived ON, investigating the contributions of atmospheric oxidants and oxidant pathways. The chemical mechanism presented in Fig. S1 and Table S2 was applied in this box model. To calculate the total production of limonene-derived ON, processes such as photolysis, dilution, and deposition were ignored for all chemical species in the model. The temperature was set to 298 K in the model. The initial concentrations of limonene and other atmospheric components for all cases were set as shown in Table S5. Limonene at a concentration of 1.0×10<sup>11</sup> molecules cm<sup>-3</sup> was used as the precursor for ON (Guo et al., 2022; Luo et al., 2023). The initial concentration of OH,  $O_3$  and  $NO_3$  spanned  $1.0 \times 10^5$  to  $1.0 \times 10^{19}$ molecules·cm<sup>-3</sup>,  $1.0\times10^{11}$  to  $1.0\times10^{18}$  molecules·cm<sup>-3</sup> and  $1.0\times10^{9}$  to  $1.0\times10^{17}$  molecules·cm<sup>-3</sup>, respectively. The low values represent typical atmospheric concentrations of these species, which are within the range of those reported in previous studies (Shen et al., 2021; Liu et al., 2023; Matsunaga and Ziemann, 2019). The medium to high values represent extreme conditions, in order to investigate the significant impact of oxidants on limonene-derived ON across a broad spectrum of oxidant levels. Chamber experiments were simulated by the box model under ideal situation, which has been specifically design to analyze chemical processes, while simulations under real atmospheric condition were carried using global model in sect. 2.3. We conducted sensitivity tests (Sect. S1 in the supplement) to examine oxidation pathways for formation of limonene-derived ON. Sensitivity tests under single initial oxidation were set. Building upon this foundation, sensitivity tests with multiple oxidation pathways were implemented: (1) introducing secondary oxidant across three concentration gradients under fixed primary oxidant levels, followed by (2) increasing concentration of third oxidant with three concentration gradients. A summary of all cases can be found in Table S4.





#### 2.3 Simulation of global limonene-derived ON

We used the Community Earth System Model (CESM) version 1.2.2.1 coupled with the University of Michigan Integrated Massively Parallel Atmospheric Chemical Transport (IMPACT) aerosol model with a resolution of 1.9° × 2.5° for this study. The IMPACT aerosol module gets the meteorology field from the CESM model at each time step, while changes in the aerosols in IMPACT do not provide feedback to the CESM. The emission of precursors BVOCs are estimated by the Model of Emissions of Gases and Aerosols from Nature inventory (MEGAN) coupled to CESM/IMPACT model. The developed explicit gas phase chemical mechanism same as used in above chemical box model was applied to simulate the formation of limonene-derived ON. The highly oxidated limonene-derived ON considered as semi-volatile species partitioning from gas phase to particle phase contributes to SOA. A base case (Case0) was designed to simulate limonene-derived ON under all three initial oxidation pathways, and six sensitivity experiments were designed for simulating global burden limonene-derived ON under two initial oxidation pathways (Case 1-3) and single initial oxidation pathway (Case 4-6), respectively. Above seven cases were summarized in Supplementary Sect. S2 and Table S6.

#### 3 Results and discussion

#### 3.1 Limonene-derived ON formation through individual initial oxidation pathway.

We employed a chemical box model to simulate limonene-derived ON formed through three initial oxidation pathways, considering various oxidant concentrations (Fig. 2). These simulations were designed to evaluate the effect of increasing oxidant concentrations on the yield of limonene-derived ON from each initial oxidation pathway. In the case with individual OH oxidation pathway, the concentration of limonene-derived ON increases as the initial OH concentration increases (Fig. 2a), following a pattern to that of limonene consumption (Fig. S2a). Initial OH concentration increases from  $1.0 \times 10^5$  to  $1.0 \times 10^{19}$  molecules cm<sup>-3</sup>, resulting in ~20.0-fold increase in the production of limonene-derived ON. At this stage, limonene is not completely consumed by OH, indicating that higher initial OH concentration will increase consumption of limonene to produce more ON.





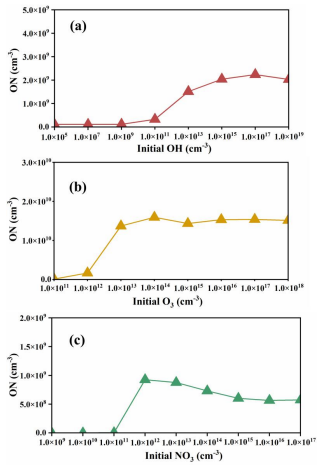


Figure 2. Variations of limonene-derived ON in individual oxidation pathway under different oxidant concentrations. The triangles represent concentration of limonene-derived ON in each experiment. The lines represent the trend of limonene-derived ON. The three datapoint colors represent three initial oxidation pathways (red for OH-initiated oxidation, yellow for  $O_3$ -initiated oxidation, green for  $NO_3$ -initiated oxidation).

In the case with individual  $O_3$  oxidation pathway, the limonene-derived ON increases first and then maintains a relatively stable production with the increase of initial  $O_3$  concentration (Fig. 2b). Limonene is not completely consumed when initial  $O_3$  concentrations below  $1.0 \times 10^{14}$  molecules cm<sup>-3</sup>. Increased consumption of limonene lead to an increase in ON production with increased  $O_3$  concentration.

Different from the cases of the OH- and O<sub>3</sub>-initiated oxidation pathways, limonene-derived ON increases when initial NO<sub>3</sub> concentrations below 1.0×10<sup>12</sup> molecules·cm<sup>-3</sup> could be caused by incompletely consumed limonene (Fig. 2c). The increased consumption of limonene with increase in





concentrations of NO<sub>3</sub> lead to the increased production of ON. However, as initial NO<sub>3</sub> concentrations continued to increase, limonene-derived ON production decrease. Compared to the case with initial NO<sub>3</sub> concentration of  $1.0\times10^{12}$  molecules cm<sup>-3</sup>, reaction of LIMAL and NO<sub>3</sub> become the dominant pathway in the case with initial NO<sub>3</sub> concentration of  $1.0\times10^{17}$  molecules·cm<sup>-3</sup>. Low yield (9.2%) of this pathway results in decreased limonene-derived ON (green box in Fig. S1). The results mean that at low initial oxidant concentration, limonene-derived ON shows a strong dependence on initial oxidant concentration, and the dependence on intermediate reaction rates becomes more important at high initial oxidant concentration.

In addition, average concentration of ON of OH-, O<sub>3</sub>- and NO<sub>3</sub>-initiated oxidation pathways when oxidations are sufficient are calculated separately. The O<sub>3</sub>-initiated oxidation pathway (1.5×10<sup>10</sup> molecules·cm<sup>-3</sup> limonene-derived ON produced) yields more ON than the OH- (2.1×10<sup>9</sup> molecules·cm<sup>-3</sup> limonene-derived ON produced) and NO<sub>3</sub>-initiated (7.1×10<sup>8</sup> molecules·cm<sup>-3</sup> limonene-derived ON produced) oxidation pathways when limonene initial concentration is constant. This indicates that under initial conditions with sufficient oxidation, O<sub>3</sub>-initiated oxidation pathway of limonene has highest yield of ON, which is about 15.0%, while that is low by OH- (2.1%) and NO<sub>3</sub>-initiated (0.7%) oxidation pathway. This difference in the ON yield among various oxidation pathways will be used to explain the contributions of each oxidation pathway to ON concentration in the following discussion.

## ${\bf 3.2~Effects~of~multiple~oxidation~pathways~on~limonene-derived~ON~formation.}$

Compared to the simulation scheme with individual oxidation pathway discussed above, introducing multiple oxidation pathways leads to comprehensive competition among them, which results in a nonlinear response of ON concentration to changes in the initial concentrations of oxidants. Figure 3 shows the dependence of limonene-derived ON on initial concentration of oxidants when include two initial oxidation pathways. The addition of oxidants has various effects on the yield of limonene-derived ON. When the initial concentration of oxidant is low  $(1.0\times10^5 \text{ molecules}\cdot\text{cm}^{-3} \text{ for OH}, 1.0\times10^{11} \text{ molecules}\cdot\text{cm}^{-3} \text{ for O}_3, 1.0\times10^9 \text{ molecules}\cdot\text{cm}^{-3} \text{ for NO}_3)$ , the initial limonene will not be completely consumed. In all case with low concentration of oxidants, adding another oxidant with oxidation pathway will increase consumption of limonene, leading to increase in the limonene-derived ON production. When the initial concentration of oxidants is high, limonene will be nearly or completely consumed. In

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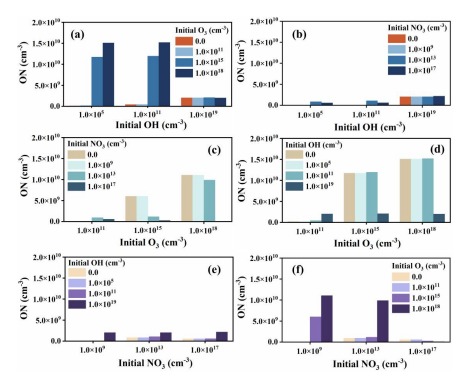
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these cases, the production of limonene-derived ON will be determined by the competition between the two oxidation pathways. The product of limonene-derived ON steadily increased as the initial concentration of O3 increases from 0 to 1.0×1018 molecules cm<sup>-3</sup> when the initial concentration of OH or NO<sub>3</sub> is constant (Fig. 3a, f). According to the chemical mechanism applied in the model, the reaction between limonene and O<sub>3</sub> has higher rate than OH and NO<sub>3</sub> in these cases (Table S7). As a result, in the presence of O<sub>3</sub>, the oxidation of limonene with O<sub>3</sub> proceeds more rapidly than with OH or NO<sub>3</sub>, leading to higher concentration of limonene-derived ON due to the high yield of O3 oxidation pathway as discussed in above section (compare Fig. 2b with Fig. 2a and 2c). In contrast, compared to only including O<sub>3</sub> oxidation pathway, adding oxidation pathways with OH or NO<sub>3</sub> will result in a decrease in limonenederived ON production (Fig. 3c, d), because some limonene that would have reacted with O<sub>3</sub> is instead converted to ON through the OH or NO<sub>3</sub> pathways with lower yield. Therefore, the dominant oxidation pathway and its ON yield determine the impact of the competition between the two oxidation pathways on the final limonene-derived ON production. A similar phenomena observed in laboratory study shows that NO<sub>x</sub> influences γ-terpinene ozonolysis by enhancing NO<sub>3</sub> production at high NO<sub>x</sub> levels, which subsequently leads to NO<sub>3</sub> preferentially consuming γ-terpinene over O<sub>3</sub> (Xu et al., 2020), illustrating the competition between oxidants. The addition of the OH-initiated oxidation pathway results in a small increase in ON production compared to NO<sub>3</sub>-initiated oxidation alone (Fig. 3e), due to the slightly higher yields of limonene-derived ON for OH-initiated oxidation pathway. The ON production would not change much when add the NO3-initiated oxidation pathway compared to the case with OH-initiated oxidation pathway alone (Fig. 3b) because of unchanged the main initial oxidation pathway. These sensitivity experiments suggest that competition of oxidation pathways plays an important role in formation of limonene-derived ON.





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Figure 3. Simulated limonene-derived ON in two initial oxidation pathways under different oxidant conditions, including variation of production of limonene-derived ON with adding (a) initial O3 concentration and (b) initial NO<sub>3</sub> concentration in the three OH levels; variation of limonene-derived ON production with adding (c) initial OH concentration and (d) initial NO<sub>3</sub> concentration in the three O<sub>3</sub> levels; variation of limonene-derived ON production with adding (e) initial OH concentration and (f) initial O3 concentration in the three NO3 levels.

240 Based on the production variations of limonene-derived ON in the cases with one and two initial 241 242 243 244 245

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oxidation pathways discussed above, the comprehensive impact of multiple oxidants on limonenederived ON formation in the cases with multiple initial oxidation pathways are analyzed (Fig. 4). The results can be summarized into three types. The Type 1 is the cases when limonene is not completely consumed (Fig. S4). When two initial oxidant concentration is low (Fig. 4a, d, g) and medium (Fig. 4d, g), the addition of third oxidant increases the production of limonene-derived ON because the addition of the third oxidant increases consumption of limonene. If the oxidant concentration is sufficient to consume up limonene, the production of limonene-derived ON will be determined by the competition between initial oxidation pathways. The Type 2 is the cases with large changes of limonene-derived ON. Under low NO<sub>3</sub> and moderate and high O<sub>3</sub> conditions, the production of limonene-derived ON decreases

with adding OH (Fig. 4a, b), because some limonene that would have reacted with O<sub>3</sub> is instead converted 11

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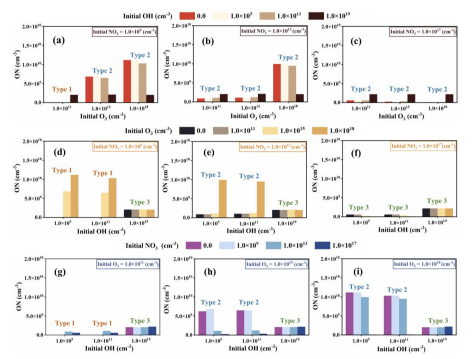
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to ON through the OH pathways with lower yield. The formation of limonene-derived ON shows similar pattern for Type 2 in Figure 4h and i. On the one hand, the yield of limonene-derived ON from NO<sub>3</sub>initiated oxidation is lowest, so the production of limonene-derived ON will decrease when the formation of limonene-derived ON from this pathway becomes the dominant formation route. On the other hand, adding NO<sub>3</sub>-initiated oxidation pathway also consumes NO<sub>3</sub> that would have reacted with the product of the OH- and O3-initiated oxidation, resulting in decrease production of limonene-derived ON. The changes in ON production with constant initial concentration of limonene and various oxidation pathways indicate the interactions of different oxidation process of limonene. In contrast to OH- and NO<sub>3</sub>-initiated oxidation pathway, adding oxidation pathways with O<sub>3</sub> will result in increase in limonenederived ON production (Fig. 4e), due to higher yield of limonene-derived ON from O3-initated oxidation pathway than OH- and NO<sub>3</sub>-initated oxidation pathways. Since the yield of limonene-derived ON of OHinitiated oxidation is higher than NO3-initiated oxidation, the production of limonene-derived ON decreases (Fig. 4c) as the main oxidation pathway changes from NO<sub>3</sub> to OH oxidation (Table S8). Additionally, in some sensitivity experiments (Fig. 4d-i), ON concentration do not change much with the addition of an oxidation pathway (Type 3). This could be explained by minimal competition with the rapid main oxidation pathway. These sensitivity experiments suggest that the limonene-derived ON production in the simulated system are not only controlled by limonene concentration, but also affected by synergic effect of multiple oxidants and oxidation pathways.





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Figure 4. The influence of adding OH-, O<sub>3</sub>- and NO<sub>3</sub>-initiated oxidation pathways on the production of limonenederived ON under different oxidant conditions, including variation of limonene-derived ON production with adding initial OH concentration in the three O<sub>3</sub> levels under (a) low, (b) moderate and (c) high NO<sub>3</sub> levels; variation of limonene-derived ON production with adding initial O<sub>3</sub> concentration in the three OH levels under (d) low, (e) moderate and (f) high NO<sub>3</sub> levels; variation of limonene-derived ON production with adding initial NO<sub>3</sub> concentration in the three OH levels under (d) low, (e) moderate and (f) high O<sub>3</sub> levels. In each panel, the types marked on the columns show the cases when limonene is not completely consumed (type 1) and almost completely consumed (large (type 2) and small (type 3) changes in limonene-derived ON production).

## ${\bf 3.3}\ Contribution\ of\ each\ oxidation\ pathway\ to\ global\ limonene-derived\ ON.$

Global simulation using CESM/IMPACT model was performed to characterize the spatial and temporal distribution of limonene-derived ON and the contributions of each oxidation pathway to global burden. Incorporation of formation of limonene-derived ON reduces underestimation of simulated ON by comparison with observations summarized in the literature (Sect. S3 in the supplement) (Li et al., 2023b). The spatial distribution of limonene-derived ON is shown in Fig. 5a. The simulated global mean limonene-derived ON burden is about  $60.5~\mu g \cdot m^{-2}$ , and the highest burdens (>500  $\mu g \cdot m^{-2}$ ) are predicted over tropical forest regions of central Africa. As the primary precursor of limonene-derived ON, the concentration of limonene dominates the yield of these ON compounds. The seasonal cycle of simulated limonene-derived ON is presented in Fig. S6, which is mainly depend on limonene levels. Global average





limonene-derived ON burden peaks in the summer (69.2 µg·m²) due to highest global average limonene concentration (Fig. S7b), while the large burden of limonene-derived ON in fall is driven by the presence of both high limonene concentration (Fig. S7c) and NO concentration (Fig. S8c) compared to spring and winter. In contrast, the burden of limonene-derived ON is lowest in winter (48.1 µg·m²) because of lowest concentration of limonene (Fig. S7d). Beyond the effects of limonene and NO concentrations, oxidant levels and oxidation pathways also affect the formation mechanisms and production of limonene-derived ON, which may explain the highest burden in regions such as Central Africa, rather than Amazon where limonene concentrations are highest over the world (Fig. S9a). The concentration of oxidants is inherently low in Amazon (Fig. S9b-d) and most oxidants are consumed by reaction with isoprene and some other monoterpene with higher concentrations compared to limonene. Consequently, only a few of oxidants are available to react with limonene, leading to low burden of limonene-derived ON in Amazon despite the highest burden of limonene there. Therefore, high concentrations of limonene-derived ON can only form when both high limonene and oxidant concentrations are present simultaneously.

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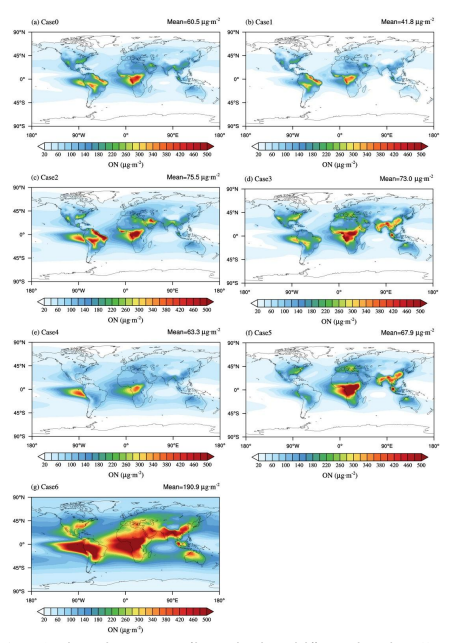


Figure 5. Annual mean column concentration of limonene-derived ON with different simulation schemes. (a) Run with three initial oxidation pathways (Case0), (b) without OH-initiated oxidation pathway (Case1), (c) without O3-initiated oxidation pathway (Case2), (d) without NO3-initiated oxidation pathway (Case3), (e) without O3- and NO3-initiated oxidation pathways (Case4), (f) without OH- and NO3-initiated oxidation pathways (Case5) and (g) without OH- and O3-initiated oxidation pathway (Case6).

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To quantify the contribution of each oxidation pathway to the formation of limonene-derived ON in different regions, we conducted a series of sensitivity experiments (Case1 to 6 introduced in method section) on the oxidation pathways (Fig. 5b-g). Our simulations indicate that increasing O<sub>3</sub> and NO<sub>3</sub>initiated oxidation pathways result in 15.5% and 18.0% increase of global average burden of limonenederived ON, respectively, compared to OH-initiated oxidation pathway alone (Fig. S10a, b). This is primarily because higher yields of limonene-derived ON associated with the O3- and NO3-initiated oxidation pathways compared to OH-initiated oxidation pathways. When compared to O3-initiated oxidation pathway alone (Fig. S10c, d), the addition of OH- or NO<sub>3</sub>-initiated pathways result in increased burden of limonene-derived ON in the limonene-sufficient region (e.g. Amazon), owing to adding a limonene-derived ON formation pathway to consume more limonene. However, in the limonenedeficient yet NO<sub>3</sub>-sufficient regions (e.g. Central Africa, Mediterranean, and middle and low latitude of Asia), increasing the OH- or NO3-initiated oxidation pathways reduces the burden of limonene-derived ON. This occurs because the oxidation of limonene by OH or NO<sub>3</sub> suppresses O<sub>3</sub>-initiated oxidation, which otherwise produces limonene-derived ON with a high yield. Additionally, if limonene undergoes oxidation by NO<sub>3</sub>, the availability of NO<sub>3</sub> for the nitration of OH- and O<sub>3</sub>-initiated oxidation products of limonene will decrease, resulting in a reduction in limonene-derived ON. The addition of OH- and O3initiated oxidation pathways reduces global average burden of limonene-derived ON by 60.5% and 78.1% respectively, compared to the case with NO3-initiated oxidation pathway alone (Fig. S10e, f). This reduction is likely due to insufficient NO<sub>3</sub> oxidation at night to further oxidize intermediates produced from OH- and O3-initiated limonene oxidation during the day, limiting the formation of limonene-derived ON at night. The burden of limonene-derived ON undergoes a noticeable change when an additional oxidation pathway is introduced to the existing two pathways (Fig. S11). Adding OH-initiated oxidation pathway increases the global average burden of limonene-derived ON from 41.8 to 60.5 µg·m<sup>-2</sup>, by 44.7%, while adding O<sub>3</sub>-initiated oxidation pathway decrease that from 75.5 to 60.5 µg·m<sup>-2</sup>, by 19.9% (Fig. S11a, b), which was attributed to the competition between the OH and O<sub>3</sub> oxidation pathways for reactions with limonene. When the O3-initiated oxidation pathway produces the same amount of limonene-derived ON as the OH-initiated pathway, it consumes more NO3. As a result, increasing the O3 oxidation pathway reduces the availability of NO3 for the nitration of intermediate oxidation products, thereby lowering the





total limonene-derived ON yield across all three pathways. In contrast, enhancing the OH oxidation pathway increases the total yield. Moreover, the addition of the NO<sub>3</sub>-initiated oxidation pathway increases burden of limonene-derived ON in the limonene-sufficient region even over 150 μg·m<sup>-2</sup> (Fig. S11c). However, in the region with high NO<sub>3</sub> concentration, the burden of limonene-derived ON decreases (Fig. S11c) because the NO<sub>3</sub>-initiated oxidation pathway yields less limonene-derived ON than the O<sub>3</sub>- and OH-initiated oxidation pathways. These results highlight the different nonlinear responses of limonene-derived ON to multiple oxidation pathways under varying oxidation conditions and precursor concentrations. This difference contributes to the disparity between the global model results and the idealized experimental results from the box model, emphasizing the importance of developing explicit chemical mechanisms in global models for understanding SOA formation processes.

### 4 Conclusion and implications

In this work, the explicit chemical mechanism is developed to simulate formation and spatial distribution of limonene-derived ON using a chemical box model and global model CESM/IMPACT. Under multiple initial oxidation pathways, limonene-derived ON shows non-linear variations with different oxidant conditions, which is controlled by the synergetic effects of multiple oxidants. When limonene is not consumed, adding another oxidant with oxidation pathway will increase limonene-derived ON due to increased consumption of limonene. When limonene is completely consumed, limonene-derived ON production is dominated by competition of oxidation pathways. The production of limonene-derived ON is increased by O<sub>3</sub>-initiated oxidation pathway while decreased by OH and NO<sub>3</sub>-initiated oxidation pathway. This is mainly because limonene oxidated by O<sub>3</sub> produces more ON than OH and NO<sub>3</sub>, resulting from the simulation under individual initial oxidation pathway.

The global model simulation indicates that oxidation process is important for limonene-derived ON formation in addition to limonene concentration. Global limonene-derived ON burden decreases by 19.9% and 17.3% due to O<sub>3</sub>- and NO<sub>3</sub>-initiated oxidation pathway, while OH-initiated oxidation pathway increases global limonene-derived ON burden by 44.7% compared the case only including the other two oxidation pathways. These differences can be attributed to the complex nonlinear response of limonene-derived ON yield to different reaction pathways under varying precursor and oxidant conditions.





The chemical mechanism of ON formation could have an influence on the formation and spatial
distribution of ON. We only include main oxidation process published to date in the model. Although
uncertainties remain in simulating limonene-derived ON due to limited knowledge of its formation
mechanism, this work offers an improvement in the global model's ability to simulate ON and presents
a methodological framework for simulating SOA and their chemical processes. This framework can be
used in the future to improve SOA burden predictions and provide a comprehensive understanding of the
complex interactions between multiple oxidation pathways, which are crucial for SOA formation (Chen
et al., 2022; Zang et al., 2024). Quantitative understanding of these complex interactions in contributing
to SOA formation is essential for understanding the contributions of interaction between anthropogenic
and natural emissions to atmospheric aerosols, and for providing more effective measures to reduce
aerosol pollution.
<b>Data availability.</b> Simulation data are available upon request to the corresponding authors.
Author contributions. QG and JZ designed the study, developed the chemical box model and global
model conducted the simulations, analyzed the data, and wrote the manuscript. HZ and BL provided the
laboratory data. PF, LC, YS, HL, YL and YX contributed to the discussion and revision of the paper.
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