RE: A point-to-point response to reviewers' comments

"Oxygenated products formed from OH-initiated reactions of trimethylbenzene: Autoxidation and accretion" (acp-2020-165) by Yuwei Wang, Archit Mehra, Jordan E. Krechmer, Gan Yang, Xiaoyu Hu, Yiqun Lu, Andrew Lambe, Manjula Canagaratna, Jianmin Chen, Douglas Worsnop, Hugh Coe, Lin Wang

Dear Prof. Dr. Markus Ammann,

We are very grateful to the helpful comments from the reviewers, and have carefully revised our manuscript accordingly. A point-to-point response to the comments, which are repeated in italic, is given below.

We are looking forward to the reviewers' feedback and your decision with the revision.

Best regards,

Lin Wang Fudan University lin_wang@fudan.edu.cn

Reviewer #1

GENERAL COMMENTS

In this manuscript the authors present results of an experimental study of the products of the reaction of trimethylbenzene isomers with OH radicals in the absence and presence of NOx. Reactions are conducted in a flow tube reactor and gas-phase products are analyzed using three types of online mass spectrometry that in combination allow detection of products ranging from very low to very high oxidation state. The authors detect a variety of monomer and dimer products that are not currently incorporated into the Master Chemical Mechanism, which is widely used to model the atmospheric chemistry of organic compounds. The authors propose structures for the compounds (they are preliminary since only elemental formulas and deuterium labeling data are available) and mechanisms by which they could be formed. Fortunately, the authors have kept the paper concise, and not attempted to overinterpret the results or overwhelm readers with data. Although it is difficult to judge the importance of the results because of the lack of quantitative analysis, aromatics are an important class of compounds whose chemistry is not well understood, the work in technically sound, and it should be useful for others to build on. I think the paper should be published in ACP after the following minor comments and have been addressed.

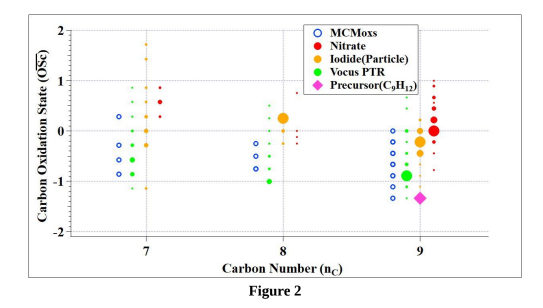
Response. We are very grateful to the positive viewing of our manuscript by Reviewer #1, and have now revised our manuscript accordingly.

SPECIFIC COMMENTS

1. Line 227-228: How can an oxidation product have an oxidation state lower than TMB?

Response. In the original manuscript, these products refer to ions of $C_8H_{15}O^+$, $C_8H_{15}O_2^+$, $C_8H_{17}O^+$, and $C_8H_{17}O_2^+$ detected only by Vocus PTR, all of which have very low signal intensities ranging from 14.67 cps to 78.47 cps. As a comparison, the ion counts for the most abundant C9 products ($C_9H_{11}O^+$) and C8 products ($C_8H_{11}O^+$) detected by Vocus PTR were 13887 cps and 5023 cps, respectively. The concentrations of these ions should be quite low, considering the high sensitivity of Vocus PTR to compounds with a few oxygen atoms (around 8000 cps/ppb to xylene and TMB, and 8500 cps/ppb to methyl ethyl ketone as calibrated with a commercial calibration cylinder). At the same time, there is a documented history of fragmentation during PTR ionization, leading to a leakage of oxygen atom(s) (de Gouw and Warneke, 2007; Gueneron et al., 2015; Karl et al., 2018; Tani, 2013; Yuan et al., 2017). Though oxygenated VOCs are reported to be less significantly fragmented than alkanes and many alkenes (Yuan et al., 2017), a leakage of oxygen due to fragmentation is typically on the order of < 5% for ketones, 15% for alcohols (Karl et al., 2018; Španěl and Smith, 2013). Hence, we postulate that these ions might come from fragmentation of parent compounds in the FIMR (focusing ion-molecule reactor) of Vocus PTR. We have removed this statement in Line 225-228 of the original manuscript and updated Figure 2.

The revised Figure 2 is shown below.



2. Lines 260-262 (and elsewhere): Because of the high sensitivity of the nitrate-CIMS to oxidized products I do not think you can assume that products formed in these experiments did not involve multiple OH reactions. For example, in the work by Krechmer et al., EST (2016) they formed multigeneration products in chamber experiments in which the lights were only on for 10 s.

Response. We agree with this reviewer that multiple OH attacks can occur in our reaction system, which is evidenced by the observation of $C_9H_{16}O_{6-9}$ products, as 16 hydrogen atoms in the molecular formula can be regarded as a characteristic of the second-generation products according to Molteni et al. (2018). In fact, we mentioned the possibility of multiple OH-attacks in multiple places in our manuscript, i.e., statements in Line 238 of the original manuscript and Table 2.

In Line 260-262 of the original manuscript, we intended to state that $C_9H_{12}O_6$ as a multi-generation hydroperoxyl product as predicted by MCM should not have a comparable yield as that formed through the carbonyl termination reaction of a peroxy radical that is involved in autoxidation, since a multigeneration product is not favored at OH exposure as short as one life time of TMB. $C_9H_{14}O_6$ is another product predicted by MCM but we missed it in main text of the original manuscript. $C_9H_{14}O_6$ is unlikely to be formed with a considerable yield through the MCM route for a similar reason.

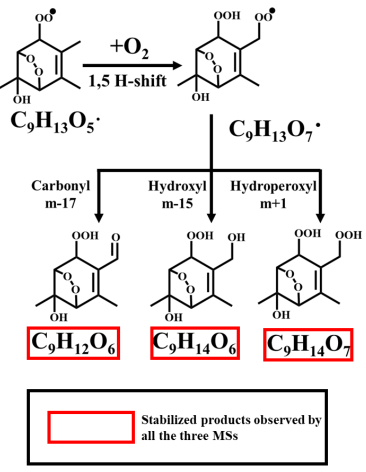
To avoid misunderstanding, we have revised our manuscript, which (Line 268-275) reads " $C_9H_{12}O_6$ is one of the only two signals that have been predicted by MCM,..., which is unlikely to contribute a lot to the observed signal of $C_9H_{12}O_6$ since the concentration of a multi-generation product is not expected to be high at OH exposure as short as one lifetime of TMB. $C_9H_{14}O_6$ is the other one, presumed to be a hydroperoxyl product of a second-generation peroxy radical formed via the epoxy-oxy pathway (MCM name: TM124MUOOH), which is unlikely to be formed through the MCM route with a considerable yield, either."

Please also refer to our response to Comment #3 from Reviewer #2.

3. *Line 325-326: The proposed isomerization of the bicyclic alkoxy radical would never compete with ring-opening pathways. See Vereecken and Peeters, PCCP (2009, 2010) estimation methods for these pathways.*

Response. Thanks for pointing out this issue. After a detailed discussion with Dr. Vereecken, quote, "For this structure, consider that the H-migration is across a trans-substituted alkene: the CH₃-C=C-CHO· carbons are all in one plane. Furthermore, because of the ring structure, the CHO· group cannot rotate the oxygen towards the CH₃ H-atom. Hence, to shift, the H-atom would have to "leap" almost 5 angstroms without being attached to anything, whereas normally it is less than 1.4 angstroms from either or both of the starting-C/ending-O atom", we agree with reviewer and Dr. Vereecken that the 1, 5-H-shift of this alkoxy radical is virtually impossible, and have deleted this pathway in the revised manuscript.

We have updated Scheme 1 as shown below, where the formation of $C_9H_{12}O_6$, $C_9H_{14}O_6$, and $C_9H_{14}O_7$ was proposed with the involvement of an autoxidation step.



Scheme 1

The text in Line 324-327 of the original manuscript have been deleted. We now state in the revised manuscript that (Line355-358)"An autoxidation reaction pathway that can explain the observation of $C_9H_{10}D_2O_7$ in the 1,2,4-(1-methyl-D3)-TMB + OH experiment is currently unavailable, although we speculate that a "peroxy-alkoxy-peroxy" conversion is likely involved during the formation of $C_9H_{12}O_7$ according to the number of oxygen atoms".

4. Line 362: Please provide support (such as references) for the assumption of identical charging efficiencies. Given the high sensitivity of the nitrate-CIMS to oxidation state and chemical structure this sounds like a very poor assumption. I suspect this makes the fractions quoted in sections 3.3 and 3.4 misleading and possibly results is wrong conclusions. The only way I think one can deal with this is to always refer to these as a comparison of fractional signals or some such thing.

Response. We believe that this assumption is the best option when there are no real measurements. Hyttinen et al. (2015) modelled the charging of highly oxygenated products from cyclohexene ozonolysis using nitrate-CIMS, showing a similar charging efficiency of nitrate source for highly oxygenated compounds. Ehn et al. (2014) assumed that the nitrate source has the same sensitivity for all highly oxygenated molecules. We agree with this reviewer that nitrate-CIMS is quite sensitive to the oxidation state of compounds to be measured. However, once a compound is highly oxygenated (i.e., contains 6 or more oxygen atoms) or has at least two hydrogen bond donor functional groups (for example, hydroperoxide, OOH), it can be assumed to be charged at the collision limit (Ehn et al, 2014; Hyttinen et al., 2015). Clearly, HOM monomer and dimer contain more than two hydrogen bond donor functional groups or 6 or more oxygen atoms. Though dimers possess more functional groups that favor the binding with NO₃⁻ than monomers do, the charging efficiency cannot be higher than collision limit.

We have cited the above references to support our assumption, which (Line 379-380) reads "The charging efficiency for C9 and C18 products is assumed to be identical in Nitrate CI-APi-TOF (Ehn et al., 2014; Hyttinen et al., 2015)".

5. *Line 407-409: Can you distinguish peroxynitrates by time profiles, since they should decay by reversible decomposition on short timescales (though they may be reformed)?*

Response. This is a good point. However, during our experimental procedure, we focused more on the establishment of stable signals of key products, which was determined by multiple factors including wall loss, and did not try to tackle the stability of products by terminating the reaction and monitoring the decay of signals. Thus, a comparison of time profiles of various nitrogen-containing products is currently not available.

TECHNICAL COMMENTS

1. Line 289: Should be "summarizes".

Response. We have replaced "summarizes" with "summarizes".

2. *Line 345: Should be "more O atoms"*.**Response**. We have revised our manuscript accordingly.

3. Line 362: Should replace "charge" with "charging" or "ionization". **Response**. We have replaced "charge" with "charging".

4. *Line 457: Delete "a"*. **Response**. We have removed this "a".

5. *Line 459: Should be "linearly dependent"*.**Response**. We have revised the text as "linearly dependent on".

Reviewer #2

The authors describing experimental results of the OH radical initiated oxidation of different trimethylbenzenes carried out in a flow-through apparatus at atmospheric pressure in air. OH has been produced via 254 nm photolysis of ozone in presence of water vapour. Qualitative results of end product analysis are provided from three mass spec techniques. The authors chose relatively high initial reactant concentrations, $(3.5 - 5.2) \times 10^{12}$ molecules cm⁻³, with a reactant conversion of 62.3 % each within the overall residence time of 80 s. Nothing is said regarding the RO₂ radical profiles in the experiments. It can be assumed from the stated reaction parameters that RO₂ levels are substantially higher than atmospheric. Consequently, especially RO₂ self- and cross reactions are favoured, which are of less importance for the RO₂ fate under atmospheric conditions. Thus, it 's not so surprising that a very big fraction of C18 products has been detected. And that 's my main concern: Are the observed product distributions from these experiments relevant for the atmosphere?

Response. Indeed, the initial reactant concentrations are much higher than those under atmospheric conditions. However, high concentrations of VOCs were deliberately chosen in this study. We aim to experimentally observe highly oxygenated products to confirm the possibility of autoxidation, and to propose the detailed autoxidation pathways via the comparison between reactions of un-deuterated and partially deuterated reactants. High concentrations of reactants will certainly help identify the highly oxygenated products that are of low volatility and easy to loss. At the same time, we did not over-interpret our results by hinting that the observed product distributions from the experiments are the same as those in the ambient atmosphere. Our viewpoints in section 3.3, where the characteristics of C18 products are discussed, are (1) to confirm the extensive existence of highly oxygenated RO₂ radicals, in other words, the extensive existence of the autoxidation pathways in the OH-initiated oxidation of TMB as an echo of the last sentence in section 3.1; (2) to provide an evidence on the structural enhancement in accretion product formation. In summary, what we have focused on is the formation mechanism and chemical fates of the RO₂ radicals and HOM products.

To clarify this point, we now state in our revised manuscript (Line 217-220) that "Also note that the concentrations of precursors in our experiments were much higher than the atmospheric ones. These concentrations were deliberately chosen to help identify the highly oxygenated products that are of low volatility and easy to loss in the sampling, but subject to the side effect that the relative significance of different pathways could be altered"

and Line (423-425) that "Again, it should be noted that this result was obtained under the condition of very high concentrations of precursors and thus the relative fractions of products could be different under ambient conditions".

Some other points that should be considered:
1. - Line 137 – 141: Why did the authors take nitrous oxide as precursor for NO and NO₂?

Response. Parts-per-million (ppm) levels of O_3 are required to generate OH radicals, which prevent sustained NO_x (especially NO) mixing ratios at sufficient levels to compete with HO₂ as a sink for RO₂, due to the fast conversion of NO_x to nitric acid (HNO₃) via the reactions of NO+O₃ \rightarrow NO₂+O₂ and NO₂+OH \rightarrow HNO₃. On the other hand, N₂O is a better NO_x precursor specifically in OFR studies for the following reasons, as described in Lambe et al. (2017) and Peng et al. (2018), and recently reviewed in Peng et al. (2020):

- 1) The spatial distribution of NO and NO₂ generated via the N₂O + O(¹D) reaction is more homogenous than what is achieved by simple additions of NO and/or NO₂, because of the continuous production of O(¹D) from the O₃ photolysis inside the reactor.
- 2) Steady-state mixing ratios of NO from $O(^{1}D) + N_{2}O$ reactions are orders of magnitude higher than that from a simple NO injection

2. And again here: What is the NO and NO2 profile in the experiments? One example from modelling should be given in the manuscript in order to allow the readership to get an impression for this.

Response. Thanks for this suggestion. We utilized a photochemical model (PAM_chem_v8) (Lambe et al., 2017; Li et al., 2015; Peng et al., 2015) to investigate the NOx concentrations and NO/NO₂ profiles in the OFR. Unfortunately, the NO/NO₂ concentrations in the model output (tens of ppb) are much larger than our reported values (a few ppb) in our Table 2 of the original manuscript, whereas the comparison of $[O_3]$ and [1,2,4-TMB] between modelled values and measured ones looks fine (Figures R1 and R2).

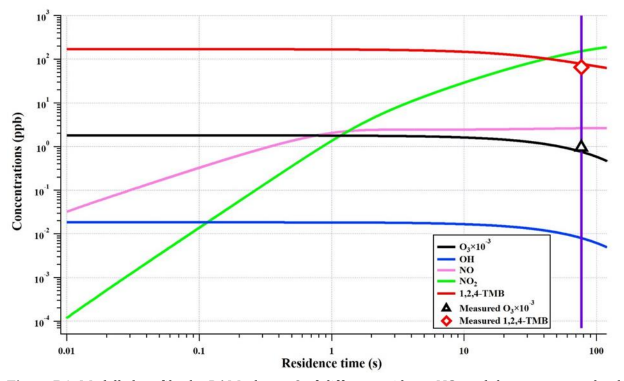


Figure R1. Modelled profiles by PAM_chem_v8 of different oxidants, NO_x and the precursor under the settings of "low NO_x experiment" (initial $[O_3] = 1.8$ ppm, initial [1,2,4-TMB] = 170 ppb, and irradiance of

254 nm Lamps = 2.0×10^{15} ph cm⁻² s). The measured [O₃] and [1,2,4-TMB] at the exit of OFR are shown by a triangle and a diamond, respectively, in the plot. The vertical purple line represents a residence time of 77.3 s."

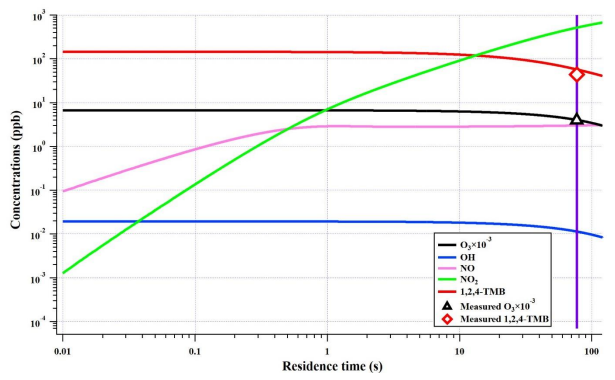


Figure R2. Modelled profiles by PAM_chem_v8 of different oxidants, NO_x and the precursor under the settings of "higher NO_x experiment" (initial $[O_3] = 6.7$ ppm, initial [1,2,4-TMB] = 145 ppb, and irradiance of 254 nm Lamps = 1.28×10^{15} ph cm⁻² s). The measured $[O_3]$ and [1,2,4-TMB] at the exit of OFR are shown by a triangle and a diamond, respectively, in the plot. The vertical purple line represents a residence time of 77.3 s."

To investigate this discrepancy, we recalibrated our NO_x monitor and performed a series of new experiments under experimental conditions similar to experiments #7 and #8 (Table R1), and generally observed a factor of two discrepancy between modelled and measured NO/NO₂ (Figure R3 and R4). The mean ratio of modelled-to-measured [NO] at the exit of OFR were 0.64 ± 0.04 and 0.98 ± 0.01 for Exp. #R1-R4 (low NO_x) and Exp. #R5-R8 (higher NO_x), respectively, whereas those of modelled-to-measured [NO₂] at the exit of OFR were 0.51 ± 0.07 and 0.73 ± 0.01 for Exp. #R1-R4 and Exp. #R5-R8, respectively. Hence, we reached a conclusion that our NO_x monitor malfunctioned during our previous experiments. Since the setting of mass spectrometers have altered significantly and thus the new mass spectrometric results are not directly comparable to those in previous experiment. We decide to keep the previous mass spectrometric results but report the modelled NO_x concentrations, which have no impacts on the conclusions of this study.

Table R1. Summary of experimental conditions for a series of new experiments carried out during the revision. The total flow was set to be 10.4 slpm and $[N_2O]$ was the same as that in the previous experiments.

| # | Precu rsor | Precursor concentratio n (ppb) | Consump tion of precursor (%) | RH(%) | O ₃ concentration (ppb) | Measured NO(ppb)/NO ₂ (p pb) | Modelled NO(ppb)/NO ₂ (ppb) |
|----|---------------|--------------------------------------|--|-----------|--|---|--|
| R1 | 1,2,4- TMB | 182 | 59.2 | 15.2 | 990 | 3.8/336.5 | 2.5/152.1 |
| R2 | 1,2,4- TMB | 191 | 51.7 | 15.2 | 1185 | 2.03/244.9 | 1.6/113.2 |
| R3 | 1,2,4- TMB | 205 | 60.2 | 15 | 868 | 5.38/360.6 | 3.2/185.0 |
| R4 | 1,2,4- TMB | 220 | 41.5 | 14.9 | 1342 | 1.17/178.1 | 1.1/84.1 |
| R5 | 1,2,4- TMB | 141 | 51.3 | 8.7 | 3953 | 3.2/791.8 | 3.1/550.9 |
| R6 | 1,2,4- TMB | 147 | 46.7 | 8.7 | 4671 | 1.8/528.2 | 1.7/365.1 |
| R7 | 1,2,4- TMB | 150 | 52.0 | 8.7 | 3375 | 4.5/1005.5 | 4.5/692.0 |
| R8 | 1,2,4- TMB | 155 | 52.9 | 8.5 | 2946 | 6.1/1079.9 | 6.0/854.1 |

Exp. #R1-R4 correspond to Exp. #7 (low NO_x), and Exp. #R5-R8 correspond to Exp. #8 (high NO_x). In Exp. #R1-R4, the 254 nm lamps were tuned to get different NO/NO₂ levels and so were in Exp. #R5-R8. Reported [NO] and [NO₂] are values at the exit of OFR.

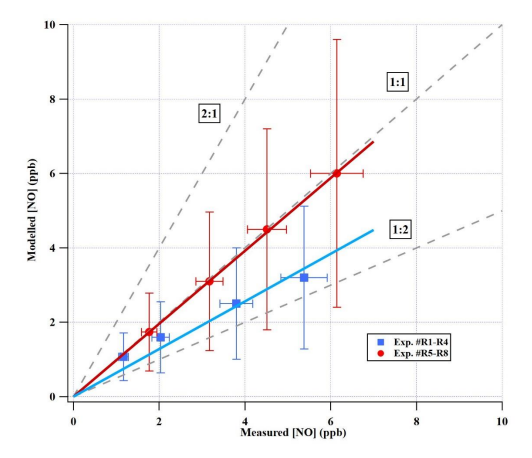


Figure R3. Measured *v.s.* modelled [NO] at a residence time of 77.3 s at the exit of the OFR. Error bars represent either \pm 60% uncertainty in model outputs (Peng et al., 2015) or \pm 10% precision in [NO] measurements by a calibrated NO_x monitor. The mean ratio of modelled-to-measured [NO] at the exit of OFR were 0.64 \pm 0.04 and 0.98 \pm 0.01 for Exp. #R1-R4 and Exp. #R5-R8, respectively.

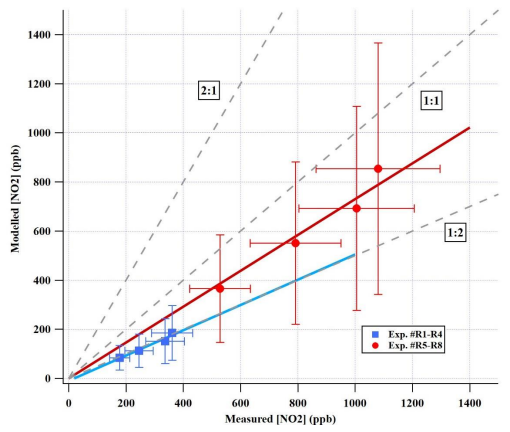


Figure R4. Measured *v.s.* modelled [NO₂] at a residence time of 77.3 s at the exit of the OFR. Error bars represent either \pm 60% uncertainty in model outputs (Peng et al., 2015) or \pm 20% precision in [NO₂] measurements by a calibrated NO_x monitor. The mean ratio of modelled-to-measured [NO₂] at the exit of OFR were 0.51 \pm 0.07 and 0.73 \pm 0.01 for Exp. #R1-R4 and Exp. #R5-R8, respectively.

We have now stated in our revised manuscript (Line 119-121) that "In addition, an ozone monitor (Model 106-M, 2B technologies) was utilized to measure ozone concentration, whereas a set of ...",

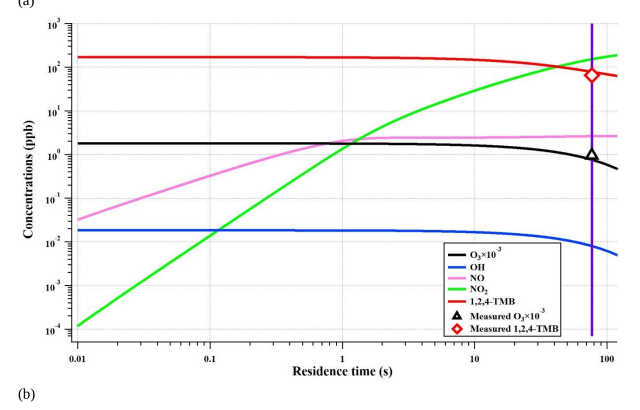
(Line 127) that "approximately 80 seconds (77.3 seconds at 10.4 slpm)",

(Line 143-145) that "A photochemical model (PAM_chem_v8) (Lambe et al., 2017; Li et al., 2015; Peng et al., 2015) was implemented to constrain the NO/NO₂ profiles in the experiments, whose details are presented in Section S1",

(Line 428-432) that "To constrain the NO_x level in the OFR, the profiles of NO/NO₂ were modelled by PAM_chem_v8, as shown in Figure S5. The mathematically-averaged NO_x levels in the low NO_x

experiment (Exp. #7) and higher NO_x experiment (Exp. #8) were 92 ppb (2.5 ppb NO + 89.5 ppb NO₂) and 295.3 ppb (2.9 ppb NO + 292.4 ppb NO₂), respectively. The NO_x/VOC in our experiments is comparable to ambient values in polluted areas. The NO_x/(Δ VOC) was around 0.9 in the low NO_x experiment and 2.9 in the higher NO_x one."

and (Section S1 in the supplement) that "Figure S5 shows the modelled profiles of the major oxidants, NO_x , and the precursor under the settings of Exp. #7 and Exp. #8 in Table 1. In the low NO_x experiment, the modelled $[O_3]$ is 20% lower than the measured value at the exit of OFR whereas the modelled [1,2,4-TMB] is 19% higher than the measured one. In the higher NO_x experiment, the modelled $[O_3]$ is 3% higher than the measured value whereas the modelled [1,2,4-TMB] is 23% higher than the measured one. (a)



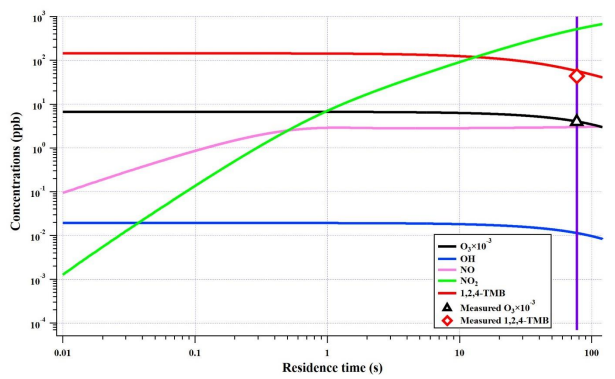


Figure S5. Modelled profiles by PAM_chem_v8 of different oxidants, NO_x and the precursor under the settings of (a) low NO_x experiment (initial $[O_3] = 1.8$ ppm, initial [1,2,4-TMB] = 170 ppb, and irradiance of 254 nm Lamps = 2.0×10^{15} ph cm⁻² s), and (b) higher NO_x experiment (initial $[O_3] = 6.7$ ppm, initial [1,2,4-TMB] = 145 ppb, and irradiance of 254 nm Lamps = 1.28×10^{15} ph cm⁻² s). The measured $[O_3]$ and [1,2,4-TMB] at the exit of OFR are shown by a triangle and a diamond in the plot. The vertical purple line represents a residence time of 77.3 s.

The updated Table 1 is shown below

| # | Precursor | Experimental | Precursor | Consumption | RH | Total | O ₃ |
|---|-----------|--------------|---------------|--------------|------|--------|----------------|
| | | condition | concentration | of precursor | (%) | flow | concentration |
| | | | (ppb) | (%) | | rate | (ppb) |
| | | | | | | (slpm) | |
| 1 | 1,2,4- | OH | 158 | 59.3 | 12.5 | 10 | 712 |
| | TMB | | | | | | |
| 2 | 1,3,5- | OH | 118 | 62.8 | 13.6 | 10 | 845 |
| | TMB | | | | | | |
| 3 | 1,2,3- | OH | 214 | 58.4 | 8.1 | 10 | 1426 |
| | TMB | | | | | | |
| 4 | 1,2,4-(1- | OH | 155 | 62.0 | 11.6 | 10 | 1003 |
| | methyl- | | | | | | |
| | D3)-TMB | | | | | | |
| 5 | 1,2,4-(2- | OH | 169 | 61.8 | 12.5 | 10 | 776 |
| | methyl- | | | | | | |
| | D3)-TMB | | | | | | |

Table 1. Summary of experimental conditions.

| 6 | 1,2,4-(4- | OH | 166 | 62.8 | 11.5 | 10 | 886 |
|---|-----------|------------------------------------|-----|------|------|------|------|
| | methyl- | | | | | | |
| | D3)-TMB | | | | | | |
| 7 | 1,2,4- | Low NO _x (2.5 | 170 | 61.5 | 12.7 | 10.4 | 944 |
| | TMB | ppb NO + 89.5 | | | | | |
| | | ppb NO ₂) ^a | | | | | |
| 8 | 1,2,4- | Higher NO _x | 145 | 69.7 | 9.3 | 10.4 | 3911 |
| | TMB | (2.9 ppb NO + | | | | | |
| | | 292.4 ppb | | | | | |
| | | NO ₂) ^a | | | | | |

^a Modelled mathematically-averaged NO/NO₂ concentrations in the OFR are shown here because of the malfunction of a NO_x monitor. The model underestimates [NO] and [NO₂] by up to a factor of 2, according to separate experiments that are not presented.

Lastly, the text in the Line 403-405 of the original manuscript that "The NO_x levels in the low NO_x experiment (Exp. #7) and higher NO_x experiment (Exp. #8) were 0.8 ppb and 6.5 ppb, respectively. Compared to the ambient values in polluted areas, this NO_x/VOC is low. The NO_x/(Δ VOC) was around 0.8% in the low NOx experiment and 6.4% in higher NO_x one" have been deleted.

3. - Line 197 - 203: From kinetic perspective, secondary chemistry, i.e. OH + product steps, cannot be neglected for a reactant conversion of 62.3 % in this system. What does it mean "one oxidation lifetime"?

Response. The statement "under this condition, the production of the first-generation products is generally favored" does not exclude the possibility of secondary reactions. "One oxidation lifetime" means the concentration of the reactant decreases to 1/e of its initial concentration, i.e., consumption of 63.2% of the initial reactant. To clarify our points, we have now stated in our revised manuscript that (Line 204-208) "...so that the OH exposure in the OFR was close to one oxidation lifetime of TMB, i.e., consumption of (1-1/e) of the initial TMB. Under this condition, the production of the first-generation products is generally favored and the multi-generation products are also present, if the subsequent loss reactions for these products are assumed to proceed in the similar rate."

Please also refer to our response to Comment #2 from Reviewer #1.

4. -Table 1:The authors also used huge ozone concentrations in their runs, $(1.7 - 9.6) \times 10^{13}$ molecules cm⁻³. After first OH attack the trimethylbenzene loses its aromaticity forming a series of unsaturated closed-shell products. What about the possible ozonolysis of these products?

Response. The significance of ozonolysis of these products can be estimated using the concentrations of ozone and OH, and the associated reaction rates with unsaturated closed-shell products. The decay of a VOC by reactions with OH/O₃ is defined as,

$$\frac{d[VOC]}{dt} = -k_{VOC+oxidant}[Oxidant][VOC]$$

where $k_{VOC+oxidant}$ is the reaction rates of VOC with OH/O₃. As a result, the lifetime of a VOC due to its reactions with OH/O₃ is defined as,

$$t_{OH} = \frac{1}{k_{VOC+oxidant} [\text{Oxidant}]}$$

Since the concentration of TMB can be measured online via Vocus PTR, we can determine the averaged OH radical concentration in the oxidation flow reactor according to the decay of TMB, whereas the concertation of O_3 can be measured with an O_3 box.

Now let's take the 1,3,5-TMB + OH experiment for example, where $[OH] = 2.2 \times 10^8$ molecules cm⁻³, and $[O_3] = 2.11 \times 10^{13}$ molecules cm⁻³. No systematic research has been performed on the reaction rate of ozone with the first-generation oxidation products of trimethylbenzene. However, the reaction rates for ozonolysis of VOCs typically range from 10^{-16} to 10^{-18} cm³ molecules⁻¹ s⁻¹. A distinct feature of the aromatic oxidation is the faster oxidation rates of the first-generation products as compared to the parent molecule (Garmash et al., 2020). The pi-electron structure of the aromatic ring makes the parent molecule less susceptible towards OH oxidation compared to most organic molecules. It is thus assumed that the reaction rate between OH and the typical bicyclic first-generation product is ~ 1.5×10^{-10} molecules cm⁻³ (MCM v3.3.1, available at: http://mcm.leeds.ac.uk/MCM). Therefore, we can determine the lifetimes of unsaturated closed-shell products against OH and ozone, respectively.

$$t_{OH} = 30.3 \text{ s}$$

 $t_{O_3} = 473.9 \sim 47393.3 \text{ s}$

Hence, ozonolysis is not expected to be a significant reaction route in our system, which is consistent with a former study (Molteni et al., 2018). Also, Berndt et al. (2018) show that the formation of ROOR' accretion products from TMB is a pure $RO_2+R'O_2$ gas-phase reaction without any hidden effects exerted by ozonolysis.

5. - Schemes: It should be clarified what the authors mean with "stabilized products".

Response. Here, the term "stabilized products" refers to "non-radical products", which shows up for a lot of times in the manuscript. On the other hand, several researchers used "closed-shell products", which we prefer not to use. We have revised our manuscript and added a clarification at the position where this term shows up for the first time (Line 96 - 97):" Subsequent reactions of the intermediates will lead to the formation of stabilized products (or non-radical products),"

6. - Figure 5: A C_{18} product fraction of more than 50% is very surprising for me. This finding should be discussed in respect of rates of the competing steps R5 - R8.

Response. In fact, our study is not the first one that observed a dimer product fraction of more than 50%, as mentioned in Line 377-379 from the previous version of manuscript. It is likely a result from the high

precursor concentrations and should not be implied to the ambient as discussed in our response to the general comment from Reviewer #2.

We have now stated in our manuscript that (Line 393-417) "In the 1,3,5-TMB oxidation experiments (Exp. #2), where the highest C18 dimer fraction was observed, the mole fraction of the C18 dimers is likely determined by the competition of reactions R5, R6, R7, and R8, which can be mathematically expressed as

$$f_{C18} = \frac{0.5 \times k_{R8}[\text{RO}_2]}{k_{R5,R6}[\text{RO}_2] + k_{R7}[\text{HO}_2] + 0.5 \times k_{R8}[\text{RO}_2]}$$
(1)

where $k_{R5,R6}$ stands for the reaction rates for R5 and R6, assumed to be around 8.8 ×10⁻¹³ cm³ molecule⁻¹ s⁻¹ by MCM, k_{R7} is the reaction rate for R7, set at a typical value of 2 ×10⁻¹¹ cm³ molecule⁻¹ s⁻¹ (Berndt et al., 2018b; Bianchi et al., 2019), and k_{R8} is the reaction rate of R8 for BPRs generated by 1,3,5-TMB, which has recently been measured to be as fast as 10⁻¹⁰ cm³ molecule⁻¹ s⁻¹ (Berndt et al., 2018b).

Since the concentration of HO₂ in the OFR was not measured, we utilized a kinetic reaction model (PAM_chem_v8) to characterize the concentration profiles of oxidants in the OFR, which include OH, O₃, HO₂, and H₂O₂. A detailed description of this model is given in section S1 of the supplement and the modelled profiles of oxidants and precursors are shown in Figure S4. According to the model, the steady-state concentration of HO₂ in the Exp. #2 was around 18 ppt (~ 4.5×10^8 molecules cm⁻³). On the other hand, it is difficult to evaluate the effective concentration of the RO₂ radicals in the system, because RO₂ with low oxidation states will not form HOMs via reactions R5-R8. Therefore, we estimated the concentration of RO₂ in Eq. (1) to be close to that of BPRs in the OFR. According to MCM v3.3.1, the branching ratio for the peroxide-bicyclic pathway in the OH oxidation of 1,3,5-TMB is 79%, so that the concentration of BPRs was roughly estimated to be 58.5 ppb (~ 1.5×10^{12} molecules cm⁻³, 79% of the reacted 1,3,5-TMB). Hence, the fraction of C18 dimer is estimated to be around 98%. Clearly, this estimation itself comes with a large uncertainty, and the estimated fraction can only be regarded as an indication of explainable high yields of C18 dimers instead of a rigorous number.

In fact, under our experimental conditions, the C18 dimer fraction in the 1,3,5-TMB experiments was around 86.5%, which is much higher than the dimer fraction of 42.6%-56.5% re-calculated using the measured C9 and C18 signals by Tsiligiannis et al. (2019), 43.3%-52.4% modelled by Tsiligiannis et al. (2019), and 39% reported by Molteni et al. (2018)....",

and in Section S1 of the supplement that "PAM_chem_v8 is a model developed in conjunction with the PAM, which includes the chemistry of photolysis of oxygen, water vapor, and other trace gases by the primary wavelengths in mercury lamps (254 nm and 185 nm) (Lambe et al., 2017; Li et al., 2015; Peng et al., 2015). Simplified VOC and RO₂ chemistry are also included, but the first-generation stabilized products and the second-generation organic radical products do not react further in the model.

Fig S4 shows the modelled concentration profiles of different oxidants and 1,3,5-TMB with an irradiance of 1.64×10^{15} ph cm⁻² s by 254 nm lamps. The initial concentrations of O₃ ([O₃]) (1.2 ppm) and 1,3,5-TMB ([1,3,5-TMB]) were measured before turning on the 254 nm lamps. [O₃] and [1,3,5-TMB] at an 80 s residence time were also measured. The modelled [O₃] is 20% lower than the measured value whereas modelled [1,3,5-TMB] is very close (4% higher) to the measured one, which shows the reliability of this model.

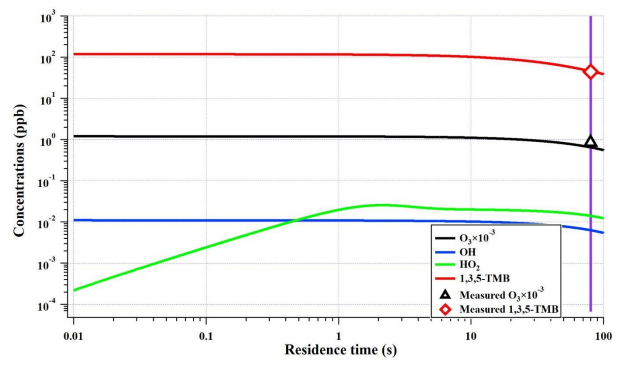


Figure S4. Concentration profiles of different oxidants and 1,3,5-TMB outputted by PAM_chem_v8 under the settings of Exp. #2. Initial [O₃] and [1,3,5-TMB] are 1.2 ppm and 118 ppb, respectively, which were used as input of the model. The measured [O₃] and [1,3,5-TMB] at the exit of OFR are shown by a triangle and a diamond, respectively. Input of irradiance of 254 nm lamps, I_{254} , is 1.64×10^{15} ph cm⁻² s, which was measured with a photodiode in the OFR. The vertical purple line represents a residence time of 80 s"

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1 Oxygenated products formed from OH-initiated reactions of

2 trimethylbenzene: Autoxidation and accretion

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- 15 Abstract. Gas-phase oxidation pathways and products of anthropogenic volatile organic compounds 16 (VOCs), mainly aromatics, are the subject of intensive research with attention paid to their contributions to 17 secondary organic aerosol (SOA) formation and potentially, new particle formation (NPF) in the urban 18 atmosphere. In this study, a series of OH-initiated oxidation experiments of trimethylbenzene (TMB, C_9H_{12}) 19 including 1,2,4-TMB, 1,3,5-TMB, 1,2,3-TMB, and 1,2,4-(methyl-D3)-TMBs (C₉H₉D₃) were investigated 20 in an oxidation flow reactor (OFR), in the absence and presence of NO_x . Products were measured using a 21 suite of state-of-the-art instruments, i.e., a nitrate-based chemical ionization - atmospheric pressure interface time-of-flight mass spectrometer (Nitrate CI-APi-TOF), an iodide-adduct chemical ionization -22 23 time-of-flight mass spectrometer (Iodide CI-TOF) equipped with a Filter Inlet for Gases and AEROsols 24 (FIGAERO), and a Vocus proton-transfer-reaction mass spectrometer (Vocus PTR). A large number of C9 25 products with 1-11 oxygen atoms and C18 products presumably formed from dimerization of C9 peroxy 26 radicals were observed, hinting the extensive existence of autoxidation and accretion reaction pathways in 27 the OH-initiated oxidation reactions of TMBs. Oxidation products of 1,2,4-(methyl-D3)-TMBs with 28 deuterium atoms in different methyl substituents were then used as a molecular basis to propose potential 29 autoxidation reaction pathways. Accretion of C9 peroxy radicals is the most significant for aromatics with meta-substituents and the least for aromatics with ortho-substituents, if the number and size of substituted 30 31 groups are identical. The presence of NO_x would suppress the formation of C18 highly oxygenated 32 molecules (HOMs) and enhance the formation of organonitrates, and even dinitrate organic compounds. 33 Our results show that the oxidation products of TMB are much more diverse and could be more oxygenated than the current mechanisms predict. 34
- 35

36 **1 Introduction**

Oxidation products of volatile organic compounds (VOCs) contribute significantly to the formation of
secondary organic aerosols (SOAs) (Ng et al., 2010; Zhang et al., 2007), which raises a globally ubiquitous
health and environmental concern (Hallquist et al., 2009). There have been numerous studies that aim to
construct detailed VOC oxidation mechanisms to advance our understanding on VOC degradation, SOA
formation, and ozone formation (Atkinson, 1986; Atkinson and Arey, 2003; Atkinson and Carter, 1984;
Kroll and Seinfeld, 2008; Ziemann and Atkinson, 2012). Based on the hypothesis that the products and

kinetics of many unstudied chemical reactions can be proposed by analogy to known reactions of similar
chemical species (Ziemann and Atkinson, 2012) and/or predicted by the structure-activity relationships

- 45 (Kwok and Atkinson, 1995), the Master Chemical Mechanism (MCM) is developed as a nearly explicit
- 46 chemical mechanism, describing the degradation of numerous VOCs (Bloss et al., 2005; Jenkin et al., 2003;
- 47 Saunders et al., 2003). Due to the high complexity of VOC oxidation processes, it is not surprising that
- 48 mechanisms leading to the formation of previously unidentified species are still missing.

49 The formation of highly oxygenated organic molecules (HOMs) through the autoxidation pathway during VOC oxidation is such an example. HOMs refer to organic compounds typically containing six or 50 more oxygen atoms that are formed in the gas phase (Bianchi et al., 2019). Autoxidation is a chemical 51 52 process where an alkyl peroxy radical (RO_2) undergoes an intramolecular hydrogen shift followed by 53 addition of a molecular oxygen, resulting in a more oxygenated RO₂ radical (Crounse et al., 2013; Ehn et al., 2014). It is an effectively repetitive uni-molecular reaction as the more oxidized RO₂ will serve as a 54 55 parent RO_2 in the next autoxidation reaction, leading to the rapid formation of HOMs in very short time scales (Bianchi et al., 2019; Jørgensen et al., 2016). 56

57 Owing to recent developments in the analytical techniques such as nitrate-anion chemical ionization 58 mass spectrometry (nitrate CIMS), our knowledge on the autoxidation pathway during the oxidation of biogenic volatile organic compounds (BVOCs) has been significantly improved. Certain systems, such as 59 the oxidation of monoterpenes, have been studied extensively, of which ozonolysis has been confirmed as 60 an important source for HOMs (Ehn et al., 2014; Jokinen et al., 2014). The OH-initiated oxidation is also a 61 62 considerable HOM formation source for monoterpenes and isoprene (Krechmer et al., 2015), albeit at lower 63 yields for monoterpenes containing an endocyclic double bond (Jokinen et al., 2014, 2015; Rissanen et al., 64 2015). Detailed mechanisms of monoterpene-derived HOM formation reactions, initiated by ozone or OH, 65 were investigated through theoretical calculations (Berndt et al., 2016), or by analogy to reactions of similar 66 chemical species, i.e., cyclohexene (Rissanen et al., 2014). A couple of studies performed H/D isotope 67 exchange experiments, which can probe the number of hydrogen atoms other than that in C-H, strongly 68 supporting the proposal of autoxidation mechanisms (Ehn et al., 2014; Rissanen et al., 2014). Research on 69 other BVOCs, i.e., isoprene and sesquiterpenes (Crounse et al., 2013; Richters et al., 2016; Teng et al., 70 2017), and on other oxidants, i.e., NO_3 and chlorine (Nah et al., 2016; Wang et al., 2019), indicate the 71 widespread existence of autoxidation pathways in the oxidation of BVOCs. The products formed from 72 autoxidation of biogenic precursors have been proven to play a vital role in atmospheric new particle formation (NPF) because of their low volatility (Ehn et al., 2014; Stolzenburg et al., 2018; Tröstl et al., 73 74 2016).

75 On the other hand, studies on autoxidation of anthropogenic VOCs are rather sparse. Wang et al. (2017) 76 theoretically and experimentally showed the autoxidation route of alkylbenzenes to form HOMs in the gas phase. Identities and yields of HOM products from different aromatics were systematically measured and 77 78 the determined molar HOM yields were in the range of 0.1 % to 2.5 %, which are similar to the molar HOM 79 yields of OH-initiated reactions of BVOCs (Jokinen et al., 2015; Molteni et al., 2018). Currently, aromatics-80 derived HOMs are believed to be formed via many reaction pathways, including accretion, bicyclic intermediate reactions, and multi-generation OH reactions (Berndt et al., 2018b; Garmash et al., 2019; 81 82 Zaytsev et al., 2019). The unimolecular isomerization and autoxidation reactions of aromatic peroxy

radicals have been shown to be fast enough to compete with other bimolecular reactions even under NOconcentrations as high as in urban environment (Tsiligiannis et al., 2019).

85 Trimethylbenzene (TMB) including isomers of 1,3,5-TMB, 1,2,3-TMB, and 1,2,4-TMB is one of the 86 most common anthropogenic VOCs in urban areas. OH-initiated oxidation of TMB is its dominant chemical 87 loss in the atmosphere (Atkinson and Arey, 2003), which proceeds either via H atom abstraction from the 88 methyl substituents or via addition of OH radical onto the aromatic ring (Ziemann and Atkinson, 2012). 89 The H atom abstraction channel is minor in the OH-induced oxidation reactions of TMB, forming dimethyl-90 benzaldehyde. The major channels of OH addition consist of peroxide-bicyclic pathway, phenolic pathway, and epoxy-oxy pathway (Bloss et al., 2005; Calvert et al., 2002; Jenkin et al., 2003). The three TMB isomers 91 92 have different branching ratios for these pathways resulting from the substitution-, site-, and stereo-93 specificity, however specific branching ratios are still in debate. Among these pathways, the peroxide-94 bicyclic pathway has the highest branching ratio and can form bicyclic peroxy radicals (BPRs), which are 95 important intermediates that contribute significantly to the formation of HOMs (Wang et al., 2017). 96 Subsequent reactions of the intermediates will lead to the formation of stabilized products (or non-radical 97 products). On the other hand, the details of the autoxidation mechanisms for anthropogenic precursors 98 remain elusive. Direct measurements of individual H-shift rates, the detailed structure of HOMs, and a 99 robust quantification of HOM yields are still lacking. The detailed kinetics for termination reactions of 100 different RO₂ are also ambiguous. Consequently, it is hard to comprehensively judge the TMB oxidation 101 reaction pathways and products under different atmospheric conditions, and to evaluate the contribution of 102 TMB oxidation to atmospheric NPF and SOA formation.

103 In this study, we studied the OH-initiated oxidation of 1,3,5-TMB, 1,2,3-TMB, and 1,2,4-TMB with 104 a focus on autoxidation and accretion products, via the concurrent usage of a Vocus proton-transfer-reaction 105 time-of-flight mass spectrometry (Vocus PTR), an iodide-adduct chemical ionization - time-of-flight mass 106 spectrometer equipped with a Filter Inlet for Gases and AEROsols (FIGAERO Iodide CI-TOF), and a 107 nitrate-based chemical ionization - atmospheric pressure interface time-of-flight mass spectrometer (Nitrate 108 CI-APi-TOF). Oxidation of 1,2,4-(methyl-D3)-TMBs was investigated to elucidate the detailed 109 autoxidation reaction pathway. The influence of NO_x concentration on product distribution was also 110 investigated.

111

112 **2 Methods**

113 As shown in Figure 1, oxidation experiments of TMB were conducted in a Potential Aerosol Mass 114 (PAM) oxidation flow reactor (OFR, Aerodyne Research, Inc.). A self-prepared VOC cylinder was used to 115 provide a constant source of gaseous TMB as a reactant. O₃/OH was produced in-situ in the PAM and the 116 relative humidity (RH) was regulated by the PAM setup, which will be introduced in details later. A Vocus 117 PTR (Krechmer et al., 2018), a FIGEARO Iodide CI-TOF (Lee et al., 2014; Lopez-Hilfiker et al., 2014), 118 and a Nitrate CI-APi-TOF (Ehn et al., 2014; Eisele and Tanner, 1993) were deployed to detect gaseous 119 products as well as particulate ones. In addition, an ozone monitor (Model 106-M, 2B technologies) and a 120 NO_{*} monitor (Model 42i TLE; Thermo Fisher Scientific) were was utilized to measure trace gas 121 concentrations ozone concentration, whereas a set of Scanning Mobility Particle Sizer (SMPS, consisting of one TSI Model 3080 Long DMA and one TSI Model 3776 Condensation Particle Counter) was employed 122 123 to measure the number size distribution of submicron aerosol particles.

124 **OFR.** In this study, the sum of all the flows in the PAM, including a zero air flow, an ozone (O_3) flow, 125 a TMB/N₂ flow, and a N₂O/N₂ flow depending upon experimental conditions, was kept at either 10 or 10.4 slpm (standard litres per minute, standard to 0 °C, 1 atm), resulting in calculated mean residence times of 126 127 approximately 80 seconds (77.3 seconds at 10.4 slpm). Zero air was generated by a zero gas generator (Sabio Model 1001 Zero Gas Source). A fraction of the zero air was passed through a Nafion humidifier 128 129 (Perma Pure Model FC100-80-6MSS) filled with ultrapure water to achieve the desired RH in the OFR. 130 Ozone was generated by passing 800 sccm (standard cubic centimetre, standard to 0 °C, 1 atm) of zero air through a separate ozone chamber and input into the OFR. In order to create a low HO₂/RO₂ ratio 131 environment to promote the carbonyl and hydroxyl channels to terminate RO_2 radicals, the OFR was 132 operated with only the 254 nm lights on (Lambe et al., 2019), which is referred to as OFR254 mode in 133 134 previous studies (Peng et al., 2015). In OFR254 mode, the primary oxidant production reactions in the OFR 135 are:

136 $O_3 + hv(254 nm) \to O_2 + O(^1D)$

137
$$O(^{1}D) + H_{2}O \to 2OH$$
 (R2)

In some experiments, N₂O (99.999%, Air Liquide) was added at the OFR inlet, corresponding to mixing ratios of 3.4% of the total gas flow rates, which produced NO_x via the following reactions (Lambe et al., 2017):

(*R*1)

 141
 $N_2 O + O(^1D) \rightarrow 2NO$ (R3)

 142
 $NO + O_3 \rightarrow NO_2 + O_2$ (R4)

<u>A photochemical model (PAM_chem_v8)</u> (Lambe et al., 2017; Li et al., 2015; Peng et al., 2015) was
 <u>implemented to constrain the NO/NO₂ profiles in the experiments, whose details are presented in Section</u>
 <u>S1.</u>

Before each experiment, the PAM OFR was purged with zero air under the OFR254 operation mode
until the signals of acetic acid and other common VOC oxidation products decreased to background levels
of the Vocus PTR and CI-TOF that are described below.

149 **Vocus PTR.** The newly developed Vocus PTR has a high sensitivity to a wide range of VOCs and 150 oxygenated volatile organic compounds (OVOCs) (Krechmer et al., 2018; Li et al., 2019; Riva et al., 2019). Its mass resolving power $(m/\Delta m = \sim 12000 \text{ at } 200 \text{ Th}, 1 \text{ Th} = 1 u/e$, where e is the elementary charge and 151 *u* is the atomic mass unit) allows to simultaneously monitor many isobaric species, and even to distinguish 152 153 the very minor mass discrepancy (0.001548 u) between one deuterium atom and two hydrogen atoms. The 154 instrument background together with a quantitative calibration by injection of standards was measured between every two experiments to minimize potential inaccuracies. In our study, the pressure of the 155 focusing ion-molecule reactor (FIMR) was actively maintained at 1.5 mbar resulting in an E/N of the FIMR 156 at 110 Td (1 Td = 1×10^{-17} V cm²), which was generally a moderate operating condition leading to relatively 157 158 little fragmentation of compounds of interest (Gueneron et al., 2015; Yuan et al., 2017).

FIGAERO-Iodide CI-TOF. The Iodide-adduct CI-TOF is able to determine elemental compositions of a suite of atmospheric oxygenated organic species (D'Ambro et al., 2017; Lee et al., 2014; Lopez-Hilfiker et al., 2016). It has increasing sensitivities toward more polar and acidic VOCs (Lee et al., 2014). The mass resolution of the Iodide CI-TOF was tuned to be around 3000. The reagent ion (I⁻) was produced from permeated CH_3I vapor in N₂ by a radioactive source of Am-241 (0.1 mCi). The pressure in the ion-molecule 164 reactor (IMR) was regulated at 100 mbar, whereas the small segmented quadrupole (SSO) pressure was set 165 to be around 2 mbar. The FIGAERO inlet manifold enables the Iodide CI-TOF to measure both gas and particle compositions at a molecular level (Lopez-Hilfiker et al., 2014). In our study, aerosols were collected 166 167 onto a PTFE filter (5µm, Millipore) at 0.96 slpm for 20 min, while the gases were measured simultaneously 168 via a separate dedicated port. Then, a thermal desorption cycle was started 2 minutes after the FIGAERO 169 filter was aligned to a heating tube, through which a heated ultra-high purity nitrogen flow was passed and heated according to a pre-programmed temperature ramp. The ultra-high purity nitrogen was initially held 170 at 25 °C for 2 min, and then heated at a rate of 10 °C min⁻¹ to 200 °C, which was maintained for the 171 remainder of the temperature ramp (50 min in total). 172

173 **Nitrate CI-APi-TOF.** The Nitrate CI-APi-TOF has been increasingly used for the measurement of 174 low volatility organic compounds (LVOC) and extremely low volatility organic compounds (ELVOCs) 175 (Ehn et al., 2014; Hyttinen et al., 2015; Jokinen et al., 2014), which mostly have a high O:C ratio. The 176 resolving power of the Nitrate CI-APi-TOF was up to around 8000 in our study. The selectivity of nitrate 177 ions keeps the spectrum clean from the more abundant, less oxidized compounds in our experiments. Most 178 of the detected species were observed exclusively as adducts with NO₃⁻, a very minor fraction of which 179 contain odd hydrogen numbers and are hence postulated to be radicals but not presented in this manuscript.

180 The concurrent use of three mass spectrometers (MSs) with different reagent ions allows us to obtain 181 a comprehensive picture of the oxidation products of TMB with OH radicals. The detection suitability of 182 these three instruments for oxidation products with various levels of oxidation has been discussed a lot in 183 previous studies (Isaacman-VanWertz et al., 2017; Krechmer et al., 2018; Riva et al., 2019). Generally, 184 Vocus PTR displays selectivity for less oxidized compounds; Iodide CI-TOF favors more oxygenated 185 species; and Nitrate CI-APi-TOF shows the highest efficiency for the most oxidized compounds. Dimer products of TMB oxidation are expected to be detected by Nitrate CI-APi-TOF as clusters with NO₃⁻, which 186 187 is due to the potential hydrogen bond donor functional groups in these molecules, inferred from the 188 abundant oxygen and hydrogen atoms in the formulas. These products should not be detected by Vocus 189 PTR. One explanation is that these molecules are likely to be fragile and therefore have fragmented owing to the protonation or the strong electric field in the FIMR of Vocus PTR. Alternatively, these products 190 191 might not go through the PEEK tube inlet of Vocus PTR. At the same time, the sample inlet for Iodide CI-192 TOF in our experiments is not desirable for the detection of dimer products.

193 To ensure that the reported signal is truly from the sample flow instead of internal background or contamination, subtraction of the mass spectra for the OFR background from the samples has been 194 195 performed for each instrument. In addition, since this study is mostly concerned with identification of oxidation products from OH-initiated reactions of TMBs and elucidation of the potential autoxidation 196 197 pathway, Nitrate CI-APi-TOF and Iodide CI-TOF were hence not calibrated and only the arbitrary signals with MS transmission correction (Heinritzi et al., 2016; Krechmer et al., 2018) were compared within the 198 199 same instrument. It should then be noted that the relative signal intensities are biased among the MSs 200 because of their ionization methods and transmission efficiency.

In each experiment, the Vocus PTR was used to confirm the establishment of stable precursor gas concentrations, and then the pair of 254 nm Hg lamps were turned on to generate the OH radicals and reaction products were analyzed by the MSs. The input RH in the OFR was kept at a low level and the voltage of the Hg lamps was slightly tuned in every experiment, so that the OH exposure in the OFR was close to one oxidation lifetime of TMB (Kurylo and Orkin, 2003), i.e., consumption of $\frac{62.3\%(1-1/e)}{62.3\%(1-1/e)}$ of the initial TMB. Under this condition, the production of the first-generation products is generally favored favoured and the multi-generation products are also present, if the subsequent loss reactions for these products are assumed to proceed in the same similar rate.

209 Table 1 summarizes all the experiments that were performed. Studied were 1,3,5-TMB (\geq 99.0%, 210 Aladdin), 1,2,3-TMB (Analytical standard, Aladdin), 1,2,4-TMB (≥ 99.5%, Aladdin), 1,2,4-(1-methyl-D3)-TMB (≥ 95%, Qingdao Tenglong Weibo Technology Co., Ltd., China), 1,2,4-(2-methyl-D3)-TMB (≥ 211 95%, Qingdao Tenglong Weibo Technology Co., Ltd., China), and 1,2,4-(4-methyl-D3)-TMB (\geq 95%, 212 Qingdao Tenglong Weibo Technology Co., Ltd., China). The structure of these partially deuterated TMBs 213 214 can be found in Figure S1. Note that ozone reactions were not taken into account in this study, because 215 ozone reacts with aromatics at negligible rates, and its reaction rate with oxidation products containing C=C double bonds is much slower compared with that of OH (Jenkin et al., 1997, 2003; Molteni et al., 2018; 216 217 Saunders et al., 2003). Also note that the concentrations of precursors in our experiments were much higher 218 than the atmospheric ones. These concentrations were deliberately chosen to help identify the highly 219 oxygenated products that are of low volatility and easy to loss in the sampling, but subject to the side effect 220 that the relative significance of different pathways could be altered.

221

3 Results and discussion

223 **3.1 Characteristics of C9 products**

Figure 2 presents an overview of C7, C8 and C9 products in a carbon oxidation state (\overline{OS}_{c}) -carbon 224 225 number (n_c) space as observed by three MSs and also those predicted by MCM v3.3.1. Carbon oxidation 226 state is a quantity that increases with the level of oxidation, which reveals the chemical aging of atmospheric 227 organics (Kroll et al., 2011). It is evident that more species were detected by the three MSs, and although 228 there were clear differences between products detected from different MSs, results indicate missing 229 oxidation pathways in the current versions of the MCM (MCM v3.3.1, available at: 230 http://mcm.leeds.ac.uk/MCM). Oxygen-containing C9 products were formed by adding functional groups 231 to the carbon skeleton, whereas C7 and C8 products resulted from carbon-carbon scission of the original 232 carbon skeleton together with functionalization. A large proportion of C7-C9 products were more oxidized 233 than those predicted by MCM, hinting the existence of highly efficient oxidation pathways. At the same 234 time, some of the C7 and C8 products were characterized with unexpected low \overline{OS}_{r} , of which a few were 235 even less oxidised than the precursor. The observation of these products is another indication for the 236 existence of missing pathways in the current oxidation mechanisms. 237 Recent studies have emphasized on the importance of the peroxide-bicyclic pathway in producing

highly oxygenated compounds in the oxidation of alkylbenzenes (Wang et al., 2017; Zaytsev et al., 2019), which leads to the formation of ring-retaining products. Therefore, here we further investigated C9 products of TMB oxidation detected by the three MSs (Figure 3). $C_9H_{10}O_{1-6}$, $C_9H_{12}O_{1-7}$, and $C_9H_{14}O_{4-6}$ contributed to the most of the signal intensities in Vocus PTR (Figure 3a). Compounds with fewer hydrogen atoms than TMB in Vocus PTR might be formed from hydrogen abstraction reactions. Iodide CI-TOF detected products with five to seven oxygen atoms (Figures 3b & 3c), which is narrower compared with Vocus PTR and Nitrate CI-APi-TOF. Molecules with 18 hydrogen atoms were detected only in Iodide CI-TOF, which 245 is an unexpected high number. These molecules, low in signal intensities in both gas and particle phases, 246 might be formed from multiple OH attacks since each OH attack can only add two hydrogens in maximum 247 onto the parent molecule. The species with the highest signal intensities measured in the gas phase appeared 248 to be $C_9H_{12}O_4$, $C_9H_{12}O_6$, $C_9H_{14}O_5$, and $C_9H_{14}O_6$ in the 1,2,4-TMB + OH experiment, $C_9H_{14}O_5$ and $C_9H_{14}O_6$ in the 1,3,5-TMB + OH experiment, and $C_{9}H_{12}O_{6}$ and $C_{9}H_{12}O_{7}$ in the 1,2,3-TMB + OH experiment (Figure 249 250 3b). Compared with the gas phase, more oxidized particulate products tended to contribute a larger 251 proportion of signal in FIGAERO-Iodide-CI-APi-TOF (Figure 3c). Nevertheless, the gas phase products 252 are emphasized in the current study, which can be detected by and compared among the three instruments. 253 Nitrate CI-APi-TOF detected C9 products containing 12-16 hydrogen atoms and 5-11 oxygen atoms (Figure 254 3d).

255 RO₂ radicals can react in the absence of NO, to form termination products including carbonyls, 256 alcohols, and hydroperoxides via the following reactions (Mentel et al., 2015).

- 257
- 258

259

| $RO_2 + R'O_2 \rightarrow R_HC = O + R' - OH + O_2$ | (<i>R</i> 5) |
|---|---------------|
| $RO_2 + R'O_2 \rightarrow ROH + R'_HC = O + O_2$ | (<i>R</i> 6) |
| $RO_2 + HO_2 \rightarrow ROOH + O_2$ | (<i>R</i> 7) |

260 Here we present a criteria method based on the work of Mentel et al. (2015). For a parent peroxy radical 261 with a molecular mass of m, its termination ought to lead to the formation of a carbonyl, an alcohol, and a hydroperoxyl, which have a molecular mass of m-17, m-15, and m+1, respectively. Since elemental 262 263 formulas as determined by the high-resolution MS do not contain information regarding functional groups 264 or the structure of a molecule, the identified mass spectral signals could be counted as either one of the 265 three categories. Listed in Table 2 are detected stabilized oxidation products in categories of carbonyl, 266 alcohol, and hydroperoxyl, which hints the potential existence of the corresponding peroxy radicals. These 267 stabilized products all contain six or more oxygen atoms, which meet the definition of HOMs (Bianchi et 268 al., 2019). $C_9H_{12}O_6$ is one of the only two signals that have been predicted by MCMthe only signal that has 269 been predicted by MCM, assumed to be a hydroperoxyl product from a ring-opening peroxy radical that 270 goes through multiple OH attack reactions (MCM name: C7MOCOCO3H), which is unlikely to contribute 271 a lot to the observed signal of $C_9H_{12}O_6$ since the concentration of a multi-generation product is not expected 272 to be high at OH exposure as short as one lifetime of TMBwhich is unlikely to occur under our experimental 273 conditions. $\underline{C_9H_{14}O_6}$ is the other one, presumed to be a hydroperoxyl product of a second-generation peroxy radical formed via epoxy-oxy pathway (MCM name: TM124MUOOH), which is unlikely to be formed 274 275 through the MCM route with a considerable yield, either. Four pairs of peroxy radicals, i.e., $C_9H_{13}O_7$ and 276 $C_9H_{13}O_{9^{\bullet}}$, $C_9H_{13}O_{8^{\bullet}}$ and $C_9H_{13}O_{10^{\bullet}}$, $C_9H_{15}O_{7^{\bullet}}$ and $C_9H_{15}O_{9^{\bullet}}$, and $C_9H_{15}O_{8^{\bullet}}$ and $C_9H_{15}O_{10^{\bullet}}$, can be selected 277 from the eight potential peroxy radicals in Table 2. The molecular formulas for the peroxy radicals within 278 each pair differ by $2 \times O$, which is a first evidence for the autoxidation pathway.

279

3.2 Autoxidation mechanisms of 1,2,4-TMB 280

281 The autoxidation pathways were then further elucidated by experiments with isotopically labelled 282 precursors, 1,2,4-(1-methyl-D3)-TMB, 1,2,4-(2-methyl-D3)-TMB, and 1,2,4-(4-methyl-D3)-TMB, whose 283 structure is shown in Figure S1.

284 If an intramolecular hydrogen shift happens during autoxidation with the abstracted hydrogen coming 285 from a methyl group, molecular oxygen will rapidly attach to this carbon-centred radical to form a new alkyl peroxy radical (Bianchi et al., 2019 and reference herein). One potential fate of this R-CH₂OO· radical 286 287 is to lose one of the two remaining hydrogen atoms, forming a carbonyl according to Reaction R5. Thus, one of the three original hydrogen atoms in the methyl group will leave this molecule after an autoxidation 288 289 step (Ehn et al., 2014; Mentel et al., 2015; Molteni et al., 2018; Otkjær et al., 2018; Rissanen et al., 2014; 290 Wang et al., 2017). In the case of a deuterium abstraction from a methyl-D3 group during the autoxidation, 291 an oxidation product with two deuterium atoms ($C_xH_yD_2O_z$) will then be formed, which is presumably a 292 carbonyl. Although an alcohol or a hydroperoxyl could also be formed from a peroxy radical, it is not 293 suitable to utilize the presence of alcohol and hydroperoxyl products as a criteria to judge the existence of 294 autoxidation. The hydroxyl channel of deuterated peroxy radicals can lead to the formation of alcohol products with either 3 or 4 deuterium atoms, depending on the nature of the other reacting RO₂. The slow 295 296 unimolecular reaction rate of deuterated methyl group corresponds to little formation of the products with 297 4 deuterium atoms, whereas our MSs cannot differentiate 3 deuterium atoms either from a molecule with 298 autoxidation and hydroxyl termination or from an untouched methyl-D3 group. On the other hand, the 299 hydroperoxyl channel would lead to the formation of hydroperoxyl products with 3 deuterium atoms, too. 300 Therefore, only the carbonyl channel products of a peroxy radical was used to suggest the potential 301 autoxidation that has occurred.

302 Table 3 summaries summarizes two-deuterium-containing C9 (C9HyD2Oz) products that were detected 303 by Vocus PTR and Nitrate CI-APi-TOF in different isotope labelling experiments: $C_9H_{10}D_2O_6$ in the 1,2,4-(1-methyl-D3)-TMB + OH experiment by Vocus PTR and Nitrate CI-APi-TOF; $C_9H_{10}D_2O_7$ in the 1,2,4-304 305 (1-methyl-D3)-TMB + OH experiment by Vocus PTR; and C₉H₁₂D₂O₈ in the 1,2,4-(4-methyl-D3)-TMB + OH experiment by Nitrate CI-APi-TOF. C₉H₁₀O₇D₂ (234.0703 Th) was expected to be detected by Nitrate 306 307 CI-APi-TOF, but unfortunately an undefined peak (located at 295.9827 Th) covered the position where 308 $C_9H_{10}O_7D_2 \cdot NO_3^-$ (296.0592 Th) was supposed to been identified. $C_9H_{12}D_2O_8$ (252.0814 Th) was not 309 detected by Vocus PTR, likely owing to either its low proton affinity or its partitioning onto the inlet of Vocus PTR, given its high O:C ratio and hence low volatility. However, Nitrate CI-APi-TOF was able to 310 311 detect this very sticky compound, because the nitrate source is constructed with concentric sample and 312 sheath flows that minimize the diffusive losses of samples to the source wall. These results indicate that an 313 intramolecular deuterium-migration happened on the 1-methyl-D3 substituent of the $C_9H_{10}D_3O_4$ and $C_9H_{10}D_3O_5$ radicals, and the 4-methyl-D3 substituent of the $C_9H_{12}D_3O_7$ radical, respectively, then one 314 oxygen was added to the resulting alkyl radicals, and the new peroxy radical reacted to form $C_9H_{10}D_2O_6$, 315 316 $C_9H_{10}D_2O_7$, and $C_9H_{12}D_2O_9$, respectively.

These three compounds ($C_9H_{10}D_2O_6$, $C_9H_{10}D_2O_7$, and $C_9H_{12}D_2O_9$) did not possess high signal intensities, because the deuterium transfer reactions are typically significantly slower for D (²H) nuclei than hydrogen transfer reactions for H (¹H) (Bianchi et al., 2019; Wang et al., 2017). There might be other twodeuterium-containing C9 products in these experiments. However, since many of these signals were at the instrument detection limits or even lower, the nonideal experimental conditions prevent us from confirming

322 more such compounds.

Based on the observed signals of two-deuterium-containing C9 products and structures that have been
 previously proven to favor H-shift reactions (Otkjær et al., 2018), two plausible formation pathways for the
 observed products are proposed.

326 The first one starts with a BPR of $C_9H_{13}O_5^{\bullet}$ as shown in Scheme 1, which is the first BPR formed from 327 C_9H_{12} via the peroxide-bicyclic pathway. The structure of this particular $C_9H_{13}O_5$ is different from what is 328 proposed in MCM v3.3.1, but the position for the initial OH attack, i.e., the 4th carbon on the ring, is feasible owing to the attraction of a substituted group on its para-position (Li and Wang, 2014), and the subsequent 329 addition of O₂ after the initial OH attack along with bicyclization occurs on the same relative position as 330 previous studies have suggested (Bloss et al., 2005; Jenkin et al., 2003). The resulting BPR of 331 $C_9H_{13}O_{5}$, undergoes a hydrogen shift, during which the abstracted hydrogen comes from the methyl 332 333 terminal of an allylic group. This hydrogen is much easier to be abstracted, compared to those in a normal 334 methyl group that are unlikely to go through a hydrogen shift with a peroxy radical (Otkjær et al., 2018). 335 The new BPR of $C_9H_{13}O_7$ then reacts via R5, R6, and R7 to form $C_9H_{12}O_6$, $C_9H_{14}O_6$, and $C_9H_{14}O_7$, respectively. This pathway is suggested by the observation of $C_9H_{10}D_2O_6$ in the 1,2,4-(1-methyl-D3)-TMB 336 337 + OH experiment. On the other hand, $C_9H_{13}O_5$ can alternatively self react or react with a HO₂ radical to 338 form an alkoxy intermediate, which goes through isomerization and addition of an oxygen to form a BPR 339 of C₉H₁₃O₈•. The stabilized products from C₉H₁₃O₈• include C₉H₁₂O₇, C₉H₁₄O₇, and C₉H₁₄O₈. This pathway 340 is suggested by the observation of $C_9H_{10}D_2O_7$ in the 1,2,4 (1 methyl-D3) TMB + OH experiment.

341 It's noted that in all the three isotope experiments, we also detected products of $C_9H_9D_3O_6$ and 342 $C_9H_9D_3O_7$ with much higher signal intensities, indicating the existence of other autoxidation pathways. 343 Thus, it deserves a repeated emphasis here that we only point out feasible pathways that are supported by 344 our isotope experiments in this work, but do not rule out other possibilities.

345 The second pathway is described in scheme 2. This pathway starts from a BPR of $C_9H_{13}O_5$ that is 346 formed by the initial OH attack and subsequent reactions. MCM v3.3.1 includes a BPR with the same 347 structure but does not contain the subsequent reactions. The BPR of $C_9H_{13}O_5$ can be terminated via R5, 348 forming a stabilized hydroxyl product of $C_9H_{14}O_4$, which is subject to a second OH attack and a following addition of O₂, resulting in a new peroxy radical of C₉H₁₅O₇. There are no systematic investigations on the 349 350 effect of a peroxide-bicyclic substitution on the 1,5 H-shift rate constant. However, our data indicate a 351 hydrogen shift can occur on the 4-methyl group, based on which the structure of $C_9H_{15}O_{9^{\bullet}}$ is proposed. The 352 new BPR of C₉H₁₅O₉• is then terminated via R5, R6, and R7, forming stabilized products C₉H₁₄O₈, C₉H₁₆O₈, 353 and $C_9H_{16}O_9$, respectively. This pathway is suggested by the observation of $C_9H_{12}D_2O_8$ in the 1,2,4-(4-354 methyl-D3)-TMB + OH experiment, though other pathways could result in products with the same formula. 355 An autoxidation reaction pathway that can explain the observation of $C_9H_{10}D_2O_7$ in the 1,2,4-(1-356 methyl-D3)-TMB + OH experiment is currently unavailable, although we speculate that a "peroxy-alkoxy-357 peroxy" conversion is likely involved during the formation of $C_9H_{12}O_7$ according to the number of oxygen atoms.

358 359

360 3.3 Characteristics of C18 HOMs

Products with 18 carbon atoms were observed in our experiments by Nitrate CI-APi-TOF, all containing 24-30 hydrogen atoms and 8 or more <u>oxygen</u> atoms ($C_{18}H_{24/26/28/30}O_{>8}$) (Figure 4). C18 products with 26 or 28 hydrogen atoms contributed the most of the signal intensities while those generated by 1,3,5TMB were the most abundant. Recent studies revealed that long-neglected organic peroxide dimer (ROOR')
formation reactions might be an important source of gas-phase dimer compounds, through which two
peroxy radicals form accretion products consisting of the carbon backbone of both reactants (Berndt et al.,
2018a, 2018b; Zhao et al., 2018).

$$RO_2 + R'O_2 \rightarrow ROOR' + O_2 \tag{R8}$$

This reaction has been proved to be another important loss process for RO₂ radicals formed via autoxidation. On account of their extraordinarily low vapor pressure, HOM dimers contribute more significantly to the formation and growth of atmospheric new particles than HOM monomers.

Our C18 oxidation products have similar ion formulas to the dimer products in recent 1,3,5-TMB oxidation experiments (Molteni et al, 2018; Tsiligiannis et al., 2019). In our experiments, the formation of $C_{18}H_{26}O_{8-15}$, $C_{18}H_{28}O_{9-15}$, and $C_{18}H_{30}O_{12-15}$ can be explained by reactions of two $C_{9}H_{13}O_{x}$, one $C_{9}H_{13}O_{x}$ and one $C_{9}H_{15}O_{x}$, and two $C_{9}H_{15}O_{x}$ respectively. $C_{18}H_{24}O_{8-13}$ with low signal intensities were detected by Nitrate CI-APi-TOF, hinting that H-abstraction reactions have occurred leading to a lower hydrogen atom in the product than in the precursor.

378 Figure 5 summarizes the relative contribution of C9 and C18 products formed from TMB oxidation as 379 detected by Nitrate CI-APi-TOF. The charge charging efficiency for C9 and C18 products is assumed to be 380 identical in Nitrate CI-APi-TOF (Ehn et al., 2014; Hyttinen et al., 2015). Hence, the measured relative 381 abundances of the oxidation products, with corrections of the transmission function in the MS, can faithfully 382 represent the product distribution in the experiments. In the Exp. #1-3, the dimers ($C_{18}H_{26}O_{8-15}$) formed 383 from two $C_9H_{13}O_x$ along those ($C_{18}H_{28}O_{9-15}$) from one $C_9H_{13}O_x$ and one $C_9H_{15}O_x$ contributed the most 384 intensity, whereas the most intensive C9 products $(C_9H_{14}O_{5-11})$ could be the alcohol or hydroperoxyl products of $C_9H_{13}O_{x^*}$, or the carbonyl products of $C_9H_{15}O_{x^*}$ (Table S1). 1,2,3-TMB produced the most C9 385 products, 1,2,4-TMB the second, and 1,3,5-TMB the least. An opposite trend was observed for C18 386 products. Therefore, the reduction of C9 products was likely due to the dimer formation. Here, we define 387 388 the C18 fraction as the ratio of the signal intensities of C18 products to the sum of those of C9 and C18 389 products in Nitrate CI-APi-TOF, and the C9 fraction in a similar way. According to our results, the dimer 390 fraction was the highest for aromatics with meta-substituents and the least for aromatics with ortho-391 substituents, if the number and size of substituted groups are identical, while the monomer fraction had an 392 opposite tendency. This can be explained by the stereoselectivity of accretion formation reactions.

In the 1,3,5-TMB oxidation experiments (Exp. #2), where the highest C18 dimer fraction was

395 R6, R7, and R8, which can be mathematically expressed as
$$0.5 \times k$$
 [R0,]

368

$$f_{C18} = \frac{0.5 \times k_{R8}[\text{RO}_2]}{k_{R5,R6}[\text{RO}_2] + k_{R7}[\text{HO}_2] + 0.5 \times k_{R8}[\text{RO}_2]}$$
(1)

397 where $k_{R5,R6}$ stands for the reaction rates for R5 and R6, assumed to be around 8.8×10^{-13} cm³ molecule⁻¹ 398 s⁻¹ by MCM, k_{R7} is the reaction rate for R7, set at a typical value of 2×10^{-11} cm³ molecule⁻¹ s⁻¹ (Berndt et 399 al., 2018b; Bianchi et al., 2019), and k_{R8} is the reaction rate of R8 for BPRs generated by 1,3,5-TMB, which 400 has recently been measured to be as fast as 10^{-10} cm³ molecule⁻¹ s⁻¹ (Berndt et al., 2018b). 401 Since the concentration of HO₂ in the OFR was not measured, we utilized a kinetic reaction model

402 (PAM_chem_v8) to characterize the concentration profiles of oxidants in the OFR, which include OH, O₃,

 HO_2 , and H_2O_2 . A detailed description of this model is given in Section S1 of the supplement and the 403 404 modelled profiles of oxidants and precursors are shown in Figure S4. According to the model, the steadystate concentration of HO₂ in the Exp. #2 was around 18 ppt (~ 4.5×10^8 molecules cm⁻³). On the other 405 406 hand, it is difficult to evaluate the effective concentration of the RO_2 radicals in the system, because RO_2 407 with low oxidation states will not form HOMs via reactions R5-R8. Therefore, we estimated the concentration of RO₂ in Eq. (1) to be close to that of BPRs in the OFR. According to MCM v3.3.1, the 408 409 branching ratio for the peroxide-bicyclic pathway in the OH oxidation of 1,3,5-TMB is 79%, so that the 410 concentration of BPRs was roughly estimated to be 58.5 ppb (~ 1.5×10^{12} molecules cm⁻³, 79% of the reacted 1,3,5-TMB). Hence, the fraction of C18 dimer is estimated to be around 98%. Clearly, this 411 412 estimation itself comes with a large uncertainty, and the estimated fraction can only be regarded as an 413 indication of explainable high yields of C18 dimers instead of a rigorous number.

414 In fact, under our experimental conditions, Under our experimental conditions, the C18 dimer fraction in the 1,3,5-TMB experiments was around 86.5%, which is much higher than the dimer fraction of 42.6%-415 56.5% re-calculated using the measured C9 and C18 signals by Tsiligiannis et al. (2019), 43.3%-52.4% 416 417 modelled by Tsiligiannis et al. (2019), and 39% reported by Molteni et al. (2018). The lack of a m/z-418 transmission correction in the former two studies could partially explain the discrepancy (Molteni et al., 419 2018; Tsiligiannis et al., 2019). On the other hand, this observation could also be due to the much higher RO₂ concentrations in our experiments. The amount of reacted 1,3,5-TMB in our experiment is around 74.1 420 ppb (~ 1.8×10^{12} molecules cm⁻³), whereas in the experiments of Tsiligiannis et al. (2019) and Molteni et 421 al. (2018), the numbers are 26 ppb (~ 6.5×10^{11} molecules cm⁻³) and 22.3 ppb (~ 5.6×10^{11} molecules cm⁻³) 422 423 ³), respectively. Again, it should be reminded that this result was obtained under the condition of very high 424 concentrations of precursors and thus the relative fractions of products could be different under the ambient 425 conditions.

426

427 **3.4 Influence of NO**_x

428 To constrain the NO_x level in the OFR, the profiles of NO/NO₂ were modelled by PAM_chem_v8, as 429 shown in Figure S5. The mathematically-averaged NO_x levels in the low NO_x experiment (Exp. #7) and 430 higher NO_x experiment (Exp. #8) were 92 ppb (2.5 ppb NO + 89.5 ppb NO₂) and 295.3 ppb (2.9 ppb NO + 431 292.4 ppb NO₂), respectively. The NO_x/VOC in our experiments is comparable to ambient values in 432 polluted areas. The NOx/(Δ VOC) was around 0.9 in the low NO_x experiment and 2.9 in the higher NO_x one. 433 Figure 6 describes the distribution of C9 products detected by Nitrate CI-APi-TOF in the absence of NO_x (Exp. #1), a low NO_x experiment (Exp. #7), and a higher NO_x experiment (Exp. #8), respectively. 434 Once NO_x was added, the formation of C9 non-nitrogen products declined down to around 20% of those in 435 436 Exp. #1. The production of C9 non-nitrogen products did not decrease much between low NO_x experiment and higher NO_x experiment, indicating a nonlinear effect of NO_x on the production of C9 non-nitrogen 437 products. Dinitrates $(C_9H_xN_2O_y)$ increased with the NO_x concentration, but C9 organonitrates (ONs, 438 C₉H_xNO_y) slightly reduced in the higher NO_x experiment compared to that in the low one, which indicates 439 440 a complex competition between $RO_2 + RO_2$ and $RO_2 + NO_x$. 441 The observation of C9 products containing 1-2 nitrogen atoms and C18 products with one nitrogen

441 The observation of C9 products containing 1-2 introgen atoms and C18 products with one introgen 442 atom is similar to the results for 1,3,5-TMB oxidation experiments in the presence of NO_x reported by 443 Tsiligiannis et al. (2019). NO_x can perturb the fate of peroxy radicals by the following reactions (Orlando 444 and Tyndall, 2012; Rissanen, 2018):

445 $RO_2 + NO \rightarrow RONO_2$ (R9)446 $RO_2 + NO_2 \rightarrow RO + NO_2$ (R10)447 $RO_2 + NO_2 \rightarrow RO_2NO_2$ (R11)

Competing with the other RO₂ reactions, NO_x can dramatically reduce the formation of C9 non-nitrogen
products. The NO_x levels in the low NO_x experiment (Exp. #7) and higher NO_x experiment (Exp. #8) were
0.8 ppb and 6.5 ppb, respectively. Compared to the ambient values in polluted areas, this NO_x/VOC is low.
The NO_x/(ΔVOC) was around 0.8% in the low NOx experiment and 6.4% in higher NO_x one.

452 Most organonitrates observed in our study were characterized with 13 hydrogen atoms, as detected by Nitrate CI-APi-TOF (Figure S2). All of them contained more than 6 oxygen atoms, with molecular formulas 453 corresponding to bicyclic organonitrates formed from termination reactions of $C_9H_{13}O_x$ with NO or NO₂ 454 455 (i.e., pathway R9 and R11, respectively). The dinitrates were dominated by species with 14 hydrogen atoms 456 (Figure S3). As suggested by Tsiligiannis et al. (2019), an OH radical could attack a nitrated compound 457 that is formed from NO_x termination of a peroxy radical, then an oxygen atom is added (similarly to the reactions from $C_9H_{14}O_7$ to $C_9H_{15}O_7$ in scheme 2), and then the newly formed peroxy radical that have 458 459 already contained one nitrogen will be terminated by NO or NO2 again. Therefore, most of the detected 460 dinitrates were also formed from $C_9H_{13}O_x$.

461 Figure 7a describes the relative intensities of C18 HOMs in Exp. #7, and Exp. #8 as detected by Nitrate 462 CI-APi-TOF, in comparison with their relative intensities in Exp. #1. The relative intensities of most of the 463 C18 HOMs decreased with the NO_x/(Δ VOC), while a few of the C18 HOMs including C₁₈H₂₄O₁₃, C₁₈H₂₆O₁₃, $C_{18}H_{26}O_{14}$, $C_{18}H_{28}O_{12}$ increased slightly in the higher NO_x experiment, potentially from a combined effect 464 of NO_x and OH. The injection of NO_x can compete with the other RO₂ reactions, and thus it consumes 465 466 peroxy radicals that would otherwise go through accretion reactions, which explains the decrease of most 467 C18 HOMs. On the other hand, the introduction of NO_x can increase the oxidation capacity in the OFR, as it does in the ambient environment, leading to the slight enhancement for the few C18 HOMs. After the 468 addition of NO_x , all of the C18 HOMs decreased by more than six times compared with those in no NO_x 469 470 experiments, indicating that the dimers were more strongly influenced than monomers, which is in 471 agreement with a previous study (Tsiligiannis et al., 2019).

The C18 ONs with 25 or 27 hydrogen atoms were detected in the NO_x experiments (Figure 7b). Other 472 473 C18 products containing nitrogen atoms were not detected. The $C_{18}H_{25}NO_x$ might be formed from reactions between a $C_9H_{12}NO_x$ radical and a $C_9H_{13}O_x$ radical, or between a $C_9H_{14}NO_x$ and a $C_9H_{11}O_x$ radical, all 474 of which existed in the system. The C₁₈H₂₇NO_x is most likely to be formed from reactions between a 475 476 $C_9H_{14}NO_x$ radical and a $C_9H_{13}O_x$ radical, which were the most abundant C9 radicals. All the C18 ONs 477 decreased with the increase of $NO_x/(\Delta VOC)$, which is reasonable. Introduction of NO_x into the system 478 triggered reactions between C9 peroxy radicals and NO_x, which consequently reduced the formation of 479 accretion products like C18 ONs.

480

481 **4 Conclusions**

482 The identities and distribution of oxidation products formed from OH-initiated reactions of three 483 TMBs were obtained with a suite of state-of-the-art chemical ionization mass spectrometers. Our recent 484 study shows that the ring-retaining products are more oxygenated and quite a lot of carbon-carbon scission 485 products are missed in the current model, indicating that the degradation products of aromatics are much more diverse than what is available in MCM (Mehra et al., 2020). Because of its important contribution to 486 487 the nucleation and SOA formation in urban areas, the ring-retaining products of TMB deserve a more 488 detailed characterization. Here we have built on that work by showing the formation pathways of ringretaining highly oxygenated products and through identification of accretion products. 489

490 With the assistance of three 1,2,4-(methyl-D3)-TMB experiments we have demonstrated that the rapid 491 formation of HOMs is attributable to the autoxidation pathway during the TMB oxidation. Several plausible 492 autoxidation pathways for OH-initiated reactions of 1,2,4-TMB were proposed, emphasizing on the ring-493 retaining pathways of aromatics, especially the bicyclic-peroxide channel, which is followed by 494 autoxidation that is not shown in the current models, such as MCM. Oxidation of aromatic VOCs was 495 shown in our study to produce HOM dimers, which might be underestimated or even completely ignored 496 in previous studies which utilize techniques not capable of detecting dimers. The structural enhancement 497 for accretion product formation via the $RO_2 + R'O_2$ reaction has been observed, of which the meta-498 substituents was shown to be strongest and ortho-substituents the weakest, though the detailed 499 stereoselectivity for aromatics remains unclear now.

500 In the presence of NO_x whose reaction with RO₂· can compete with RO₂· + RO₂· or RO₂· + 501 HO₂· reactions, ONs and dinitrates will be generated via reactions of NO_x with BPRs in 1,2,4-TMB 502 oxidation system, and dimer products with one nitrogen will be formed via the subsequent reactions. This 503 is consistent with a recent ambient observations in the polluted environment, where ONs, dinitrates, and 504 nitrogen-containing dimers presumably formed from BVOCs and alkylbenzenes were detected (Brean et 505 al., 2019). The formation of ONs and dinitrates from TMB is not linearly dependent on the NO_x 506 concentration, which excludes the possibility of extrapolating our laboratory results to ambient conditions. 507 Nevertheless, the changes of HOM compositions in the presence of NO_x , especially the accretion products, 508 could have an effect on NPF and SOA formation. Previous work has showed that the ring-retaining product 509 formation at NO_x environment tends to be more important for TMB than other single substituted C9 510 aromatics, i.e., isopropylbenzene and propylbenzene, which emphasized the significance of TMB ring-511 retaining oxidation in the urban environment (Mehra et al., 2020). Further research is needed to acquire a 512 quantitative understanding of the role of NO_x in HOM formation.

Clearly, these multifunctional gas phase products appear at different stages of the oxidation chain.
These mass spectra can be used as ideal "fingerprints" of TMB oxidation in the ambient gas phase
measurement to elucidate atmospheric oxidation conditions.

- 516
- 517 *Data availability*. Data related to this article will be available from a persistent repository and upon request518 from corresponding authors.
- 519
- 520 *Supplement*. The supplement related to this article is available online.
- 521

- 522 Author contributions. LW, and YW designed the experiments. YW, GY, XH, and YL carried out the
- 523 instrument deployment and operation. AM, JK, and AL provided technical support. YW analyzed the data.
- 524 YW, LW, and JK wrote the paper. All co-authors discussed the results and commented on the manuscript.
- 525
- 526 *Competing interests.* The authors declare that they have no conflict of interest.
- 527
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| # | Precursor | Experimental | Precursor | Consumption | RH | Total | O ₃ |
|------|--------------|--|------------------------|--------------------|----------|-----------|----------------|
| | | condition | concentrati | of precursor | (%) | flow | concentration |
| | | | on (ppb) | (%) | | rate | (ppb) |
| | | | | | | (slpm) | |
| 1 | 1,2,4- | OH | 158 | 59.3 | 12.5 | 10 | 712 |
| | TMB | | | | | | |
| 2 | 1,3,5- | OH | 118 | 62.8 | 13.6 | 10 | 845 |
| | TMB | | | | | | |
| 3 | 1,2,3- | OH | 214 | 58.4 | 8.1 | 10 | 1426 |
| | TMB | | | | | | |
| 4 | 1,2,4-(1- | OH | 155 | 62.0 | 11.6 | 10 | 1003 |
| | methyl- | | | | | | |
| | D3)-TMB | | | | | | |
| 5 | 1,2,4-(2- | OH | 169 | 61.8 | 12.5 | 10 | 776 |
| | methyl- | | | | | | |
| | D3)-TMB | | | | | | |
| 6 | 1,2,4-(4- | OH | 166 | 62.8 | 11.5 | 10 | 886 |
| | methyl- | | | | | | |
| | D3)-TMB | | | | | | |
| 7 | 1,2,4- | Low NO_x (2.5) | 170 | 61.5 | 12.7 | 10.4 | 944 |
| | TMB | <u>ppb NO + 89.5</u> | | | | | |
| | | <u>ppb NO₂)</u> | | | | | |
| | | ^a (0.8ppb NO _*) | | | | | |
| 8 | 1,2,4- | Higher NO _x | 145 | 69.7 | 9.3 | 10.4 | 3911 |
| | TMB | (6.5ppb NO _*) | | | | | |
| | | <u>(2.9 ppb NO +</u> | | | | | |
| | | <u>292.4 ppb</u> | | | | | |
| | | <u>NO₂) ^a</u> | | | | | |
| /lod | elled mathem | natically-averaged | NO/NO ₂ con | centrations in the | e OFR ar | e shown h | ere because of |

^a Modelled mathematically-averaged NO/NO₂ concentrations in the OFR are shown here because of the malfunction of a NOx monitor. The model underestimates [NO] and [NO₂] by up to a factor of 2, according to separate experiments that are not presented.

Table 2. Oxidation products of 1,2,4-TMB in categories of carbonyl, hydroxyl, and hydroperoxyl according to their molecular mass, as well as the potential peroxy radicals. Numbers in the parenthesis denote the relative intensity detected by Nitrate CI-APi-TOF in the OH-initiated oxidation of 1,2,4-TMB when that of the largest HOM signal ($C_9H_{16}O_8$) is arbitrarily set to be 100%. The relative intensity has been corrected with the relative transmission efficiency of Nitrate CI-APi-TOF.

| The potential peroxy radical <i>m</i> | Carbonyl <i>m</i> -17 | Hydroxyl m-15 | Hydroperoxyl m+1 | |
|---|--|--|--|--|
| $C_9H_{13}O_7$ • | $\begin{array}{c} C_9 H_{12} O_6 \\ (9.2 \ \%) \end{array}^{a,b,c,d}$ | $\begin{array}{c} C_9 H_{14} O_6 {}^{a,b,c,d} \\ (20.3 \ \%) \end{array}$ | $C_9H_{14}O_7^{b,c,d}$ (50.4 %) | |
| $C_9H_{13}O_8$ • | C ₉ H ₁₂ O7 ^{b,c,d} (54.4 %) | $\begin{array}{c}C_{9}H_{14}O_{7}^{\ b,c,d}\\(50.4\ \%)\end{array}$ | $\begin{array}{c} C_9 H_{14} O_8{}^{c,d} \\ (51.6 \ \%) \end{array}$ | |
| C ₉ H ₁₃ O ₉ • | C ₉ H ₁₂ O ₈ ^d (17.3 %) | $\begin{array}{c} C_9 H_{14} O_8{}^{c,d} \\ (51.6 \ \%) \end{array}$ | C ₉ H ₁₄ O ₉ ^d (29.1 %) | |
| $C_{9}H_{13}O_{10}$. | C ₉ H ₁₂ O ₉ ^d (14.9 %) | C9H14O9 ^d (29.1 %) | $\begin{array}{c} C_9 H_{14} O_{10}{}^d \\ (19.8 \ \%) \end{array}$ | |
| C ₉ H ₁₅ O ₇ • | $C_9H_{14}O_6^{a,b,c,d}$ (20.3 %) | C ₉ H ₁₆ O ₆ ^{b,c,d} (2.3 %) | $\begin{array}{c} C_9 H_{16} O_7{}^{b,c,d} \\ (23.5 \ \%) \end{array}$ | |
| $C_9H_{15}O_8$ • | C ₉ H ₁₄ O ₇ ^{b,c,d} (50.4 %) | $\begin{array}{c} C_9 H_{16} O_7{}^{b,c,d} \\ (23.5 \ \%) \end{array}$ | $C_9H_{16}O_8^{c,d}$ (100 %) | |
| $C_9H_{15}O_9$ • | $\begin{array}{c} C_9 H_{14} O_8{}^{c,d} \\ (51.6 \ \%) \end{array}$ | $\begin{array}{c} C_9 H_{16} O_8{}^{c,d} \\ (100 \ \%) \end{array}$ | $C_9H_{16}O_9^d$ (40.5 %) | |
| $C_9H_{15}O_{10}$. | $C_9H_{14}O_9^d$ (29.1 %) | $\begin{array}{c} C_9 H_{16} O_9{}^d \\ (40.5 \ \%) \end{array}$ | $C_9H_{16}O_{10}{}^d$ (7.1 %) | |

^a These compounds are listed in the MCM mechanism of 1,2,4-TMB where they are formed by multiple OH oxidation steps.

^b These compounds were detected by Vocus PTR.

^c These compounds were detected by Iodide CI-TOF in both gas and particle phase.

^d These compounds were detected by Nitrate CI-APi-TOF.

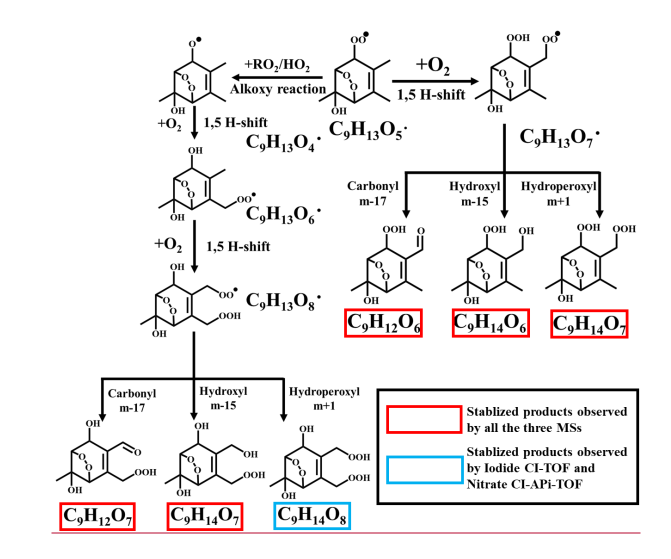
Table 3. Partially deuterated C9 products observed by Vocus PTR and/or Nitrate CI-APi-TOF. "V" and "N" denote observation by Vocus PTR and Nitrate CI-APi-TOF, respectively, whereas "-" means that the product was not observed by any instrument.

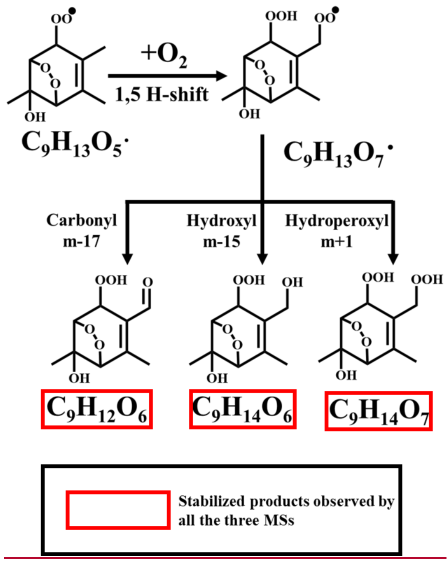
| | 1,2,4-(1-methyl-D3)- TMB | 1,2,4-(2-methyl-D3)- TMB | 1,2,4-(4-methyl-D3)- TMB |
|-------------------|-----------------------------|-----------------------------|-----------------------------|
| $C_9H_{10}D_2O_6$ | V, N | - | - |
| $C_9H_{10}D_2O_7$ | V | - | - |
| $C_9H_{12}D_2O_8$ | - | - | Ν |

Scheme Captions

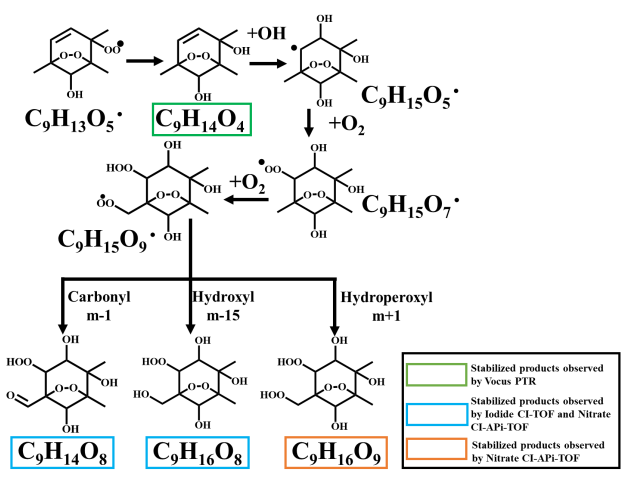
Scheme 1. A proposed autoxidation reaction scheme involving a bicyclic peroxy radical of $C_9H_{13}O_5$.

Scheme 2. A proposed autoxidation reaction scheme involving a bicyclic peroxy radical of $C_9H_{13}O_5$. Note that the reaction has been terminated with the formation of $C_9H_{14}O_4$ and re-initiated by a second OH attack.





Scheme 1



Scheme 2

Figure Captions

Figure 1. Schematics of experimental setup.

Figure 2. Comparison of C7-C9 products observed in the OH-initiated oxidation of 1,2,4-TMB (Exp. #1 in Table 1) with those listed in the MCM mechanism (Bloss et al., 2005). Filled red, orange, and green circles denote observation by Nitrate CI-APi-TOF, Iodide CI-TOF, and Vocus PTR, respectively, whereas open blue circles represent MCM species. The radius of filled circles are proportional to the signals of the compounds in each instrument. The signal of the most abundant product for each instrument is arbitrarily set to be 100%, but note that the arbitrary signals are not comparable among instruments. Symbols have been offset horizontally to avoid overlap.

Figure 3. Distribution of C9 products formed from OH-initiated reactions of TMBs (Exp. #1- 3 in Table 1) by (a) Vocus PTR, (b) Iodide CI-TOF for the gas phase, (c) Iodide CI-TOF for particle phase, and (d) Nitrate CI-APi-TOF. The yield of the most abundant product for each instrument is arbitrarily set to be 100%, but note that the arbitrary yields are not comparable among instruments. Also note that signal of Vocus PTR was processed in a logarithmic way before calculating the arbitrary yield.

Figure 4. (a) Distribution of $C_{18}H_{24}O_{8-13}$ and $C_{18}H_{26}O_{8-15}$ products formed from TMB oxidation experiments (Exp. #1-3 in Table 1), as measured by Nitrate CI-APi-TOF; (b) Distribution of $C_{18}H_{28}O_{9-15}$ and $C_{18}H_{30}O_{12-15}$ formed from TMB oxidation experiments (Exp. #1-3 in Table 1), as measure by Nitrate CI-APi-TOF; and (c) The total signal of C18 products formed from TMB oxidation experiments (Exp. #1-3 in Table 1), as measure by Nitrate CI-APi-TOF; and (c) The total signal of C18 products formed from TMB oxidation experiments (Exp. #1-3 in Table 1), as measure by Nitrate CI-APi-TOF; and (c) The total signal of C18 products formed from TMB oxidation experiments (Exp. #1-3 in Table 1), as measure by Nitrate CI-APi-TOF.

Figure 5. Relative contribution of C9 and C18 products formed from TMB oxidation experiments, as measured by Nitrate CI-APi-TOF. The relative intensity has been corrected with the relative transmission efficiency.

Figure 6. Comparison of C9 products detected by Nitrate CI-APi-TOF with zero, one or two nitrogen atoms formed from 1,2,4-TMB oxidation with different NO_x settings.

Figure 7. (a) Comparison of C18 HOMs formed from 1,2,4-TMB oxidation with different NO_x settings; and (b) Distribution of C18 organonitrates fomed from 1,2,4-TMB oxidation.

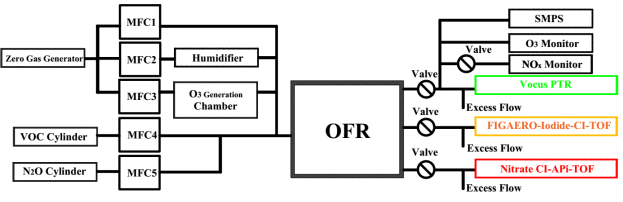


Figure 1

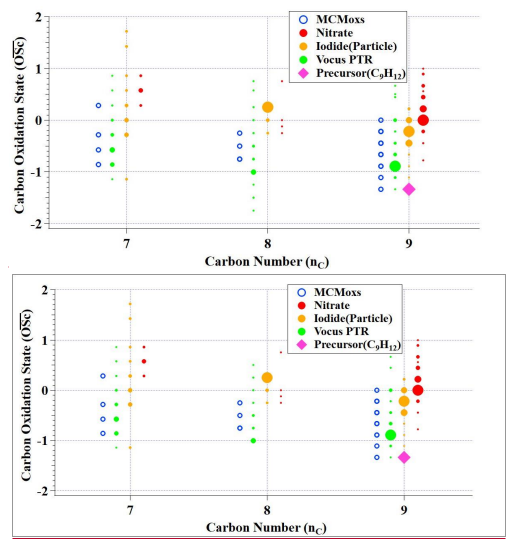


Figure 2

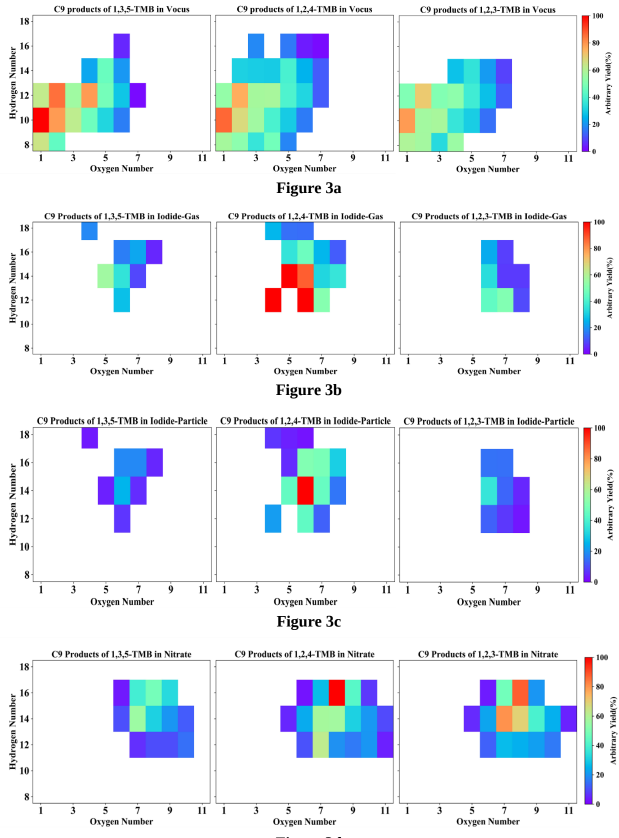
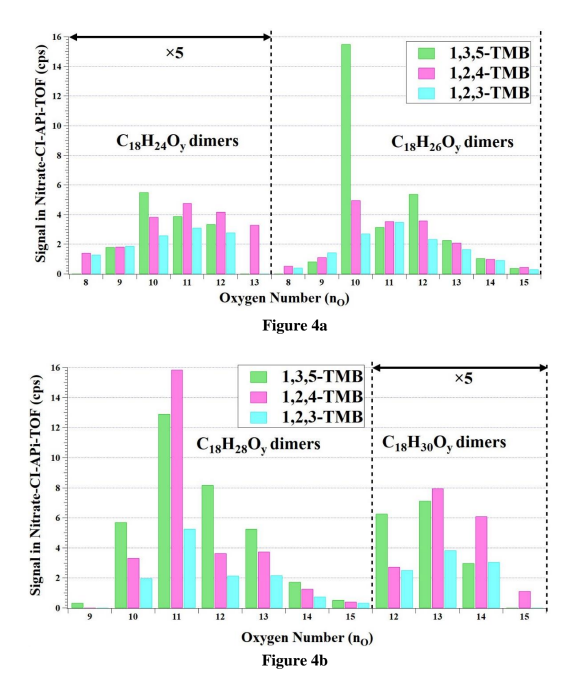


Figure 3d



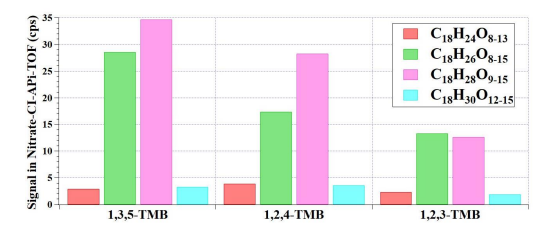


Figure 4c

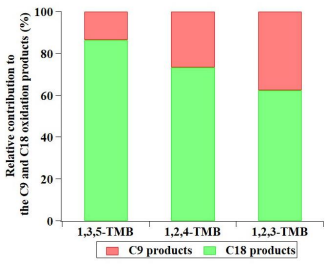


Figure 5

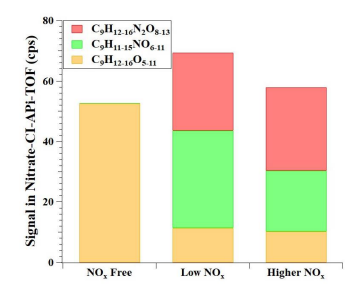
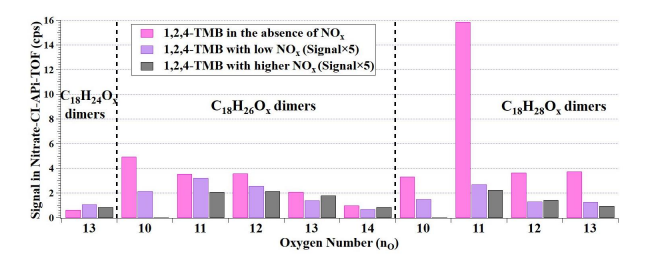


Figure 6





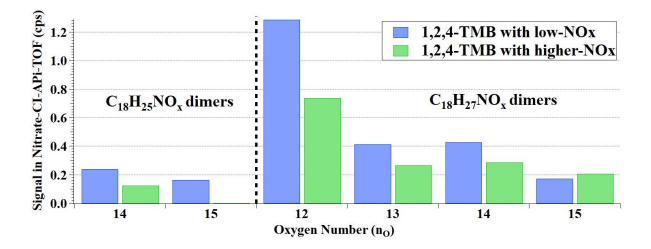


Figure 7b