# 1 Effect of NO<sub>x</sub> on 1,3,5-trimethylbenzene (TMB) oxidation product distribution and particle 2 formation

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### 10 Abstract

11 Secondary organic aerosol (SOA) represents a significant fraction of the tropospheric aerosol and its precursors are volatile organic compounds (VOC). Anthropogenic VOCs (AVOC) dominate the 12 VOC budget in many urban areas with 1,3,5-trimethylbenzene (TMB) being among the most reactive 13 14 aromatic AVOCs. TMB formed highly oxygenated organic molecules (HOM) in NO<sub>x</sub> free environment, which could contribute to new particle formation (NPF) depending on oxidation conditions were 15 elevated OH oxidation enhanced particle formation. The experiments were performed in an oxidation 16 17 flow reactor, the Go:PAM unit, under controlled OH oxidation conditions. By addition of NO<sub>x</sub> to the system we investigated the effect of NO<sub>x</sub> on particle formation and on the product distribution. We show 18 that the formation of HOM and especially HOM accretion products, strongly varies with NO<sub>x</sub> conditions. 19 20 We observe a suppression of HOM and particle formation with increasing  $NO_x/\Delta TMB$  and an increase in the formation of organonitrates (ON) mostly at the expense of HOM accretion products. We propose 21 reaction mechanisms/pathways that explain the formation and observed product distributions with 22 23 respect to oxidation conditions. We hypothesize that, based on our findings from TMB oxidation studies, aromatic AVOCs may not contribute significantly to NPF under typical NOx /AVOC conditions found 24

25 in urban atmospheres.

# 26 **1 Introduction**

Volatile organic compounds (VOC) are ubiquitous in the atmosphere and major precursors for 27 secondary organic aerosol (SOA). SOA represents a dominant fraction of the tropospheric aerosol 28 29 (Hallquist et al., 2009;Shrivastava et al., 2017;Gentner et al., 2017) and affects climate 30 (Intergovernmental Panel on Climate, 2014) and health (WHO, 2016). Consequently research interest 31 in SOA formation and properties is ranging from remote atmospheres (Ehn et al., 2012;Ehn et al., 32 2014;Kristensen et al., 2016) to densely populated and polluted environments (Chan and Yao, 2008;Hu 33 et al., 2015;Guo et al., 2014;Hallquist et al., 2016). Following the study by Ehn et al. (Ehn et al., 2012), highly oxygenated organic molecules (HOM) with low volatilities, formed from the oxidation of 34 biogenic volatile organic compounds (BVOCs), have attracted much research interest (Crounse et al., 35 2013;Ehn et al., 2014;Jokinen et al., 2014;Jokinen et al., 2015;Mentel et al., 2015;Yan et al., 36 2016;Berndt et al., 2016;Bianchi et al., 2019). These compounds have been shown to contribute to new 37 38 particle formation (NPF) and to SOA growth (Ehn et al., 2014;Bianchi et al., 2016;Kirkby et al., 39 2016;Trostl et al., 2016;McFiggans et al., 2019), making them an important factor in the formation of 40 atmospheric SOA. These oxidation products can be either described as HOM based on their high oxygen 41 number (O > 6) (Bianchi et al., 2019) or as extremely low volatile organic compounds (ELVOC) based on their volatility (Donahue et al., 2012; Trostl et al., 2016). In this study, we will refer to oxidation 42 products as HOM, because not all of the measured compounds may fulfill the criteria for ELVOC (Trostl 43

et al., 2016;Kurtén et al., 2016). Gas phase autoxidation of alkylperoxy radicals (RO<sub>2</sub>) has been proposed 44 45 as the formation mechanism for HOM (Crounse et al., 2013;Ehn et al., 2014;Jokinen et al., 2014). After the initial reaction of an oxidant with the VOC and subsequent addition of  $O_2$  to the alkylradical (R), the 46 47 produced RO<sub>2</sub> isomerizes via intra molecular H abstraction (H-shift). During this process a hydroperoxide group and a new R is formed. Additional O2 addition and H-shift sequences can introduce 48 49 large amounts of oxygen to the molecule and subsequently lower the vapour pressure. The chemistry of 50 aromatic compounds is somewhat different compared to other VOCs as they can lose their aromaticity 51 during the initial OH attack while they can retain the ring structure. Moreover, reaction products are 52 more reactive than the parent compound. The produced  $RO_2$  form an oxygen bridge, a bicyclic and 53 potentially a tricyclic alkylradical (Molteni et al., 2018; Wang et al., 2017) before further oxidation 54 processes open the ring structure. For 1,3,5-trimethylbenzene (TMB), emitted from combustion sources in the urban environment, Molteni et al. (2018) proposed a generalized reaction scheme for HOM 55 formation after OH addition. According to their scheme, first generation alkylperoxy radicals with the 56 general formula of C<sub>9</sub>H<sub>13</sub>O<sub>5-11</sub> were formed from the initial OH attack and subsequent H shift and O<sub>2</sub> 57 58 addition sequences. A postulated second OH attack would result in propagating peroxy radical chains 59 yielding radicals with the general formulas C<sub>9</sub>H<sub>15</sub>O<sub>7-11</sub>.

Generally, the termination reaction of RO<sub>2</sub> (with a general m/z = x) with HO<sub>2</sub>, leads to the 60 formation of hydro peroxides (m/z = x + 1) while termination reactions with other RO<sub>2</sub> can lead to the 61 formation of a carbonyl (m/z = x - 17), a hydroxy group (m/z = x - 15) or dimers (m/z = 2x - 32) (Mentel 62 et al., 2015; Jokinen et al., 2014; Rissanen et al., 2014). The propagation reaction of RO<sub>2</sub> with another 63  $RO_2$  or NO also results in the formation of RO (m/z = x-16). The RO can undergo internal H shift leading 64 65 to the formation of a hydroxy group and subsequent a new peroxy radical which can continue the autoxidation sequences. The alkoxy step shifts of the observed m/z by 16 leads to overlap of different 66 termination product sequences. During extensive oxidation the first generation products are subject to 67 secondary chemistry increasing the numbers of products. Molteni et al., 2018) found 68 69 several closed shell monomer products with the general formula C<sub>9</sub>H<sub>12-16</sub>O<sub>5-11</sub> from the oxidation of TMB with OH. The formation of dimers with different number of H atoms in their study was explained 70 by reactions of two first generation RO<sub>2</sub> radicals (C<sub>18</sub>H<sub>26</sub>O<sub>5-10</sub>), one first and one second generation RO<sub>2</sub> 71  $(C_{18}H_{28}O_{9-12})$  or two second generation RO<sub>2</sub> resulting in the dimer  $C_{18}H_{30}O_{11}$ . Although dimers have in 72 73 general lower O:C ratios than monomers, they are expected to be less volatile due to higher molecular 74 weight and more functional groups making them candidates to participate in nucleation processes 75 (Kirkby et al., 2016).

Organonitrates (ON) are formed as soon as sufficient  $NO_x$  is present in the atmosphere. ON are highly important for the reactive nitrogen budget wherein the formation of highly functionalized organic nitrates can contribute significantly to secondary organic aerosol (Lee et al., 2016;Bianchi et al., 2017). In this study we refer to compounds that only consist of H, C and O as HOM monomers or HOM dimers and to N containing compounds as ONs.

81 NO<sub>x</sub> influences the oxidation of organics directly by changing oxidant levels (reducing or 82 increasing OH, depending on the NO<sub>x</sub> regime) and indirectly by influencing RO<sub>2</sub> chemistry. In high NO<sub>x</sub> 83 environment such as urban areas the reaction of NO with RO<sub>2</sub> radicals can compete with the autoxidation 84 mechanism (reaction 1 and 2) and thus potentially inhibit HOM while favouring some ON formation 85 (reaction 1 with a yield of up to 0.3).

$$RO_2 + NO \to RONO_2 \tag{1}$$

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$$\rightarrow RO + NO_2 \tag{2}$$

Although ON may have high O:C they differ from HOM as they contain at least one nitrogen atom.
While HOM formation from BVOCs has been intensively studied, only few studies have been conducted
focusing on the formation of HOM from anthropogenic aromatic volatile organic compounds (AVOCs)

91 (Molteni et al., 2018; Wang et al., 2017). However, these studies indicate that AVOCs have a strong

- potential to form HOM under  $NO_x$  free conditions and proposed that they may play a crucial role in NPF and particle growth of SOA in urban areas. Regarding SOA yields a number of smog chamber studies
- and particle growth of SOA in urban areas. Regarding SOA yields a number of smog chamber studies
   have been conducted in order to investigate the oxidation of TMB with OH radical under different NOx
- and aerosol seed conditions (Paulsen et al., 2005;Rodigast et al., 2017;Wyche et al., 2009;Huang et al.,
- 96 2015;Sato et al., 2012). They reported that under higher NO<sub>x</sub>:VOC conditions, SOA yield is reduced
- 2010, such that 1012). They reported that and thigher  $100_x$  (000 contained, 0011 yield is reduced 97 compared to medium or lower NO<sub>x</sub>:VOC conditions. The NO<sub>x</sub>:VOC, in almost all studies, was < 1, apart
- from one experiment (Wyche et al., 2009), with  $NO_x$ :VOC = 1.9 in which the SOA yield was 0.29,
- 99 compared to yields up to 7.47 under lower NOx conditions.

In this study we investigate the oxidation of TMB in a laminar flow reactor, while different  $NO_x$ and OH conditions were applied. A nitrate chemical ionization atmospheric pressure interface time of flight mass spectrometer (CI-APi-TOF-MS) (Junninen et al., 2010;Jokinen et al., 2012) was used to monitor the oxidation product distribution. We show the formation of HOM and nitrate containing compounds with and without  $NO_x$  added to the reaction system. Possible mechanisms leading to the formation of ON and suppression of particle formation are discussed.

# 106 **2 Materials and Methods**

The measured HOM were generated using the laminar flow Gothenburg Potential Aerosol Mass 107 reactor (Go:PAM), initially described by Watne et al. (Watne et al., 2018). The Go:PAM is a 100 cm 108 long, 9.6 cm wide quartz glass cylinder which is irradiated over a length of 84 cm by two 30 W Phillips 109 TUV lamps (254 nm); a schematic is shown in Figure 1. The OH radicals are produced inside Go:PAM 110 by photolysing O<sub>3</sub> in the presence of water vapour. The O<sub>3</sub> is generated outside Go:PAM by photolyzing 111 pure O<sub>2</sub> (UVP Pen-Ray® Mercury Lamps, 185 nm) and distributed in 3 L min<sup>-1</sup> particle free and 112 humidified air (Milli - Q) over the reactor cross section. The VOC was introduced through a 113 gravimetrically characterized diffusion source (see Figure S2) centrally at the top of the reactor with a 114 flow of 8 L min<sup>-1</sup> while NO was introduced via a NO gas cylinder. Flows were adjusted for a median 115 residence time of 34 s in Go:PAM. A funnel shaped device is subsampling the centre part of the laminar 116 117 flow to minimize wall effects on the sample. A condensation particle counter (CPC, 3775 TSI) was used 118 to measure the number particle concentration in the sample flow. O<sub>3</sub> was monitored by a model 202 monitor (2B Technologies), relative humidity by a Vaisala HMP60 probe and  $NO_x$  by a model 42i 119 monitor (Thermo Scientific) over the course of the experiments. The OH exposure, over the residence 120 time in the reactor, for NO<sub>x</sub> free conditions without added TMB was measured using SO<sub>2</sub> titration 121 (Teledyne T100) as described by Kang et al. (Kang et al., 2007). Gas phase oxidation products were 122 measured with an Atmospheric Pressure interface High Resolution Time of Flight Mass Spectrometer 123 (APi-TOF-MS, Aerodyne Research Inc. & Tofwerk AG) (Junninen et al., 2010; Jokinen et al., 2012) in 124 connection with a A70 CI-inlet (Airmodus Ltd) (Eisele and Tanner, 1993). The CI inlet is a laminar 125 flow inlet operated with a sheath flow of 20 L min<sup>-1</sup> containing NO<sub>3</sub><sup>-1</sup> ions which are generated by 126 ionizing HNO<sub>3</sub> using an <sup>241</sup>Am foil upstream in the inlet design. The sample stream from Go:PAM is 127 introduced in the centre of the sheath flow at a rate of 8 L min<sup>-1</sup>. The NO<sub>3</sub><sup>-</sup> ions are electrostatically 128 pushed into the sample flow and form stable adducts with sample molecules as described by Ehn et al. 129 (2012). The reaction time of oxidation products and NO<sub>3</sub><sup>-</sup> is a few hundred ms before being subsampled 130 into the TOF-MS at 0.8 L min<sup>-1</sup> by a critical orifice. Differential pumping decreased the pressure from 131 103 mbar in the CI source to 10<sup>-6</sup> mbar in the TOF extraction region where HOM are detected as 132 negatively charged clusters with NO<sub>3</sub><sup>-</sup>. 133



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Figure 1: Schematics of the experimental setup with Go:PAM chamber connected to CI- APi-TOF-MS.

137 A kinetic box model was used to simulate the chemistry in the Go:PAM reactor. The core of the 138 model were first described by Watne et al. (Watne et al., 2018). The model consists of 32 species and 68 reactions now including TMB chemistry partly from the MCM v3.3.1 (Jenkin et al., 2003) as well as 139 proposed mechanisms and rate coefficients for NO<sub>2</sub> chemistry (Atkinson et al., 1992;Finlayson-Pitts, 140 141 1999) and highly oxygenated compounds (Ehn et al., 2014; Berndt et al., 2018; Zhao et al., 2018) (see SI Table S2). The photon flux used in the simulations was tuned to match measured decay of O<sub>3</sub> while 142 143 an OH sink was added to match the observed OH exposure in the background experiment, i.e. without 144 the addition of TMB. The model was run for all experiments with and without NO<sub>x</sub>. Primarily, the 145 modelled output on OH exposure for each experiments was used to interpret the results and for 146 calculating the consumed TMB. However, broadly the modelling output was also used to understand 147 the effects of changing experimental conditions on monomer, dimer and organonitrate (ON) production. 148 The experiments without NO<sub>x</sub> were named 1-4 denoting the increase in OH exposure and experiments with NO<sub>x</sub> were denoted according to their NO<sub>x</sub>/ $\Delta$ TMB and high (H) and low (L) OH exposure as seen 149 150 in Table 1.

151 **3 Results and Discussions** 

Table 1 summarises eight experiments where TMB has been oxidised by various amounts of OH using the Go:PAM unit. Generally, a high OH production, induced either by increased light exposure (two lamps) or elevated ozone concentration, resulted in new particles (e.g. exp 3 and 4) while addition of NO<sub>x</sub> reduced or suppressed the particle formation. The results from the kinetic model show that the amount of reacted TMB ranges from 5 - 30 ppb, depending on OH exposure (SI Figure S1). An overview of the oxidation product distribution measured with the CI- APi-TOF-MS for different conditions is

158	shown in Figure 3 and 4. The compounds were detected as nitrate clusters at $m/z = \text{mass}_{\text{compound}} + 62$ .
159	The spectra in Figure 3 and 4 show significant ion signals from oxygenated hydrocarbons retaining the
160	9 carbons from the original TMB with either even H numbers (closed shell) or odd H numbers (open
161	shell) with limited amount of products from fragmentation, i.e. ions with less than nine C. C9 compounds
162	with an O/C ratio of 6/9 or higher were classified as HOM monomers with the general formula
163	$C_9H_{12-16}O_{6-11}$ in the mass range $280 - 360 \text{ m/z}$ . Oxygenated hydrocarbons found in the range $460 - 560$
164	m/z containing 18 C were classified as dimers with chemical formulas C <sub>18</sub> H <sub>24-30</sub> O <sub>10-16</sub> . The monomer
165	with the highest intensity detected was $C_9H_{14}O_7 m/z$ 296. The highest intensities among the dimers were
166	$C_{18}H_{26}O_{10}$ , $C_{18}H_{28}O_{11}$ and $C_{18}H_{28}O_{12}$ at $m/z$ 464, $m/z$ 482 and $m/z$ 498, respectively. In addition to HOM
167	monomers and dimers, nitrogen containing compounds were found as C9 compounds with one or two N
168	or C18 compounds with one N. The nitrogen containing compounds were of the general formulas
169	C <sub>9</sub> H <sub>12-18</sub> NO <sub>6-13</sub> , C <sub>9</sub> H <sub>12-18</sub> N <sub>2</sub> O <sub>8-15</sub> and C <sub>18</sub> H <sub>18-24</sub> NO <sub>6-10</sub> . The dominating ON were C <sub>9</sub> H <sub>13</sub> NO <sub>8</sub> at <i>m/z</i> 325,
170	$C_9H_{15}NO_{10}$ at $m/z$ 359 and $C_9H_{14}N_2O_{10}$ at $m/z$ 372 respectively. In the experiments where NO <sub>x</sub> was
171	added, the formation of ON compounds was increasing with the NOx concentration. In parallel, the
172	levels of HOM monomers and dimers were reduced with NOx concentration, where dimers were
173	stronger affected than monomers. Even if the fragmentation products were limited some fragmentation
174	leading to less than 9 carbons could be observed. The most prominent fragments were assigned
175	molecular formulas C <sub>4</sub> H <sub>7</sub> NO <sub>7</sub> at $m/z$ 243, C <sub>4</sub> H <sub>6</sub> O <sub>12</sub> at $m/z$ 246, C <sub>5</sub> H <sub>6</sub> O <sub>12</sub> at $m/z$ 258 and C <sub>6</sub> H <sub>9</sub> NO <sub>7</sub> at $m/z$
176	269. Some compounds with C numbers of 15 and 17 were detected in the range $270 - 560 \text{ m/z}$ but their
177	contribution to the total signal was negligible (Figure 3).
170	

179Table 1: Experimental conditions for experiments with 30 ppb TMB. Ozone and initial NOx180concentration at time 0 are given in ppb and explicitly modelled OH exposure in molecules s cm<sup>-</sup>

<sup>3</sup>. TMB reacted (Δ TMB) in ppb after a reaction time of 34 s and particle number concentration

182 given in # cm<sup>-3</sup> after reaching steady state in Go:PAM. RH in all experiments was 38%.

#	[O3]0	[NO <sub>x</sub> ]0	OH exposure	Δ TMB	NO <sub>x</sub> /ΔTMB	Particle number	Contribution of top 10 species (%)
1	~19	5	$3.5 \times 10^{9}$	5.4	0.9	-	28.6
2	~19	5	$7.1 \times 10^9$	9.9	0.5	-	29.8
3	~100	3	$3.8 \times 10^{10}$	26	0.1	$60 \pm 14$	38.6
4	~100	3	2.1×10 <sup>11</sup>	30	0.1	$1610\pm217$	35.9
NOx9	~9	82	6.3×10 <sup>9</sup>	9	9.1	-	52.4
NOx3 <sub>L</sub>	~12	38	7.9×10 <sup>9</sup>	11	3.5	-	42.3
NOx3 <sub>H</sub>	~100	79	$3.1 \times 10^{10}$	25	3.2	-	34.2
NOx1	~100	35	9.1×10 <sup>10</sup>	30	1.2	$170 \pm 50$	30.5

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184 The relative contribution of the different compound classes to the total assigned signal are shown in 185 Figure 2. It is apparent that HOM monomers and dimers dominate in the experiments with low  $NO_x$ . The contribution of the monomers to the total oxidation product signal ranges from 20.7- 42.1% and 186 dimers make 6.8 - 43.3 % of the total, depending on experimental conditions. Dimer contributions are 187 highest at high OH exposure (exp 3 and 4 with estimated OH exposure of  $3.8 \times 10^{10}$  and  $2.1 \times 10^{11}$ 188 molecules s cm<sup>-3</sup>, respectively) and decrease with increasing NO<sub>x</sub>. Nitrated compounds dominated the 189 spectra with contributions up to  $\sim$ 75% in the experiments with highest amount of NO<sub>x</sub>. Surprisingly, 190 some nitrated compounds were also found in the experiments without added NO<sub>x</sub> which may stem from 191 background NO contamination (~3-5ppb). 192



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Figure 2: A) Overview of different compound groups to the total explained signal. Top panel illustrates the influence of a decrease of OH exposure (exp 4 - exp 1) and further decrease after adding NO<sub>x</sub> in exp NOx3L – exp NOx9. Dimers show a larger relative reduction than monomers with increasing NO<sub>x</sub>/VOC. Bottom panel shows the influence of increased NO<sub>x</sub>/VOC on the product distribution. Experiments 4, 3 and NOx1 resulted in particle formation. B) Modelled product distribution shown as lumped categories of nitrated compounds, HOM monomers and dimers and their relative contributions

201 Recently, Molteni et al. (2018) assigned 17 compounds making up 80% of the total detected signal for HOM oxidation products from the reaction of TMB with OH. Their compound with the highest fraction 202 of the signal (24.2%) was the dimer C<sub>18</sub>H<sub>26</sub>O<sub>8</sub>. This compound was not detected in our study. We did 203 neither detected deprotonated compounds in the mass range 270 - 560 m/z nor HOM monomers with 204 205 17 H nor compounds with O/C<0.55 which were found by Molteni et al. (2018). However, we do find 206 10 of the previously reported 17 monomers and dimers in our spectra. The oxidation product 207 distributions in our experiments are in general term more diverse, i.e. we found more compounds with smaller yields compared to Molteni et al. (2018). In our experiments the highest 20 compounds together 208 explain 46 - 63 % of the total signal with the individual highest oxidation products contributing only 209 between 4.4 and 16%. These differences can be the result of different experimental conditions and set 210 up. In our study the residence time is almost double compared to Molteni et al., leading to the formation 211 of more oxidized compounds, especially more oxidized dimers, which have been reported in this study. 212 In addition we produce OH radicals through irradiation at 254 nm in the full length of the flow reactor 213 214 enhancing the effects of secondary chemistry. Despite these differences there is a general agreement on the conclusions for the NO<sub>x</sub> free conditions with the Molteni et al. study where one rapidly form HOM 215

216 of very low volatility, that can initiate NPF.



Figure 3: Mass spectra of all experiments without NO<sub>x</sub>: 1, 2, 3, and 4. Panel A shows experiments 3 (light blue) and 4 (blue) with OH exposure of 2.87×10<sup>11</sup> and 3.47×10<sup>11</sup> molecules s cm<sup>-3</sup> respectively.Panel B shows experiment 1 (light blue) and 2 (blue) with of 3.47×10<sup>10</sup> and  $7.62 \times 10^{10}$  molecules s cm<sup>-3</sup> respectively. The signal at m/z 426 is associated with the used mass calibrant PFHA. Note the ten times lower Normalised Signal scale in B.

- 217 The amount of TMB reacted after 34 s in the experiments ranges from 30 ppb (almost all) in experiment
- 4 to 5 ppb in experiments 1 and NOx9 (comp. Table 1 and Figure S1). The signal intensities of the HOM 218
- 219 monomers are increasing with increasing OH exposure. Dimer compounds also increase with increasing OH and reach their highest levels in exp 3, resulting in the highest ratio of dimer to monomer (see Figure
- 220
- 2). Faster conversion of TMB will result in higher initial  $RO_2$  levels enabling faster  $RO_2 + RO_2$  (self-) 221
- reaction. The concentration profile of  $RO_2$  in exp 1 and 2 is lower and more evenly spread out over the 222
- length of Go:PAM (Figure S1) and the influence of the RO<sub>2</sub> + RO<sub>2</sub> reaction will be less in these 223 224 experiments compared to exp 3 and 4. Enhanced OH exposure does not only affect the monomer/dimer
- 225 ratio but also the total amount of compounds measured. Observed compounds were 10 times higher in
- 226 the high OH exposure exp 3, 4, compared to exp 1, 2. This was also valid for the experiments with added 227  $NO_x$  (NOx1 and  $NOx3_H$  vs  $NOx3_L$  and NOx9). Under high OH exposure (exp 4) the major HOM monomer is  $C_9H_{14}O_7$  with a contribution of 5.5 % to the total explained signal, followed by  $C_9H_{16}O_9$ 228 229 and C<sub>9</sub>H<sub>14</sub>O<sub>8</sub>, contributing 4.6 and 4.5 % respectively.
- 230 At the lowest OH exposure (exp 1) the major signals comprise the monomer  $C_9H_{12}O_{10}$  and the dimers C<sub>18</sub>H<sub>26</sub>O<sub>10</sub> and C<sub>18</sub>H<sub>28</sub>O<sub>11</sub>, contributing 4.4, 4.0 and 3.4 % to the total. These dimer compounds, 231 C<sub>18</sub>H<sub>26</sub>O<sub>10</sub> and C<sub>18</sub>H<sub>28</sub>O<sub>11</sub>, are the highest signals in exp 2 and 3. Two open shell species were found 232 233 among the larger signals in exp 1 and 2 (see Figure 3): C<sub>9</sub>H<sub>15</sub>O<sub>8</sub> and C<sub>9</sub>H<sub>15</sub>O<sub>7</sub>.
- 234 The number of H atoms is a characteristic for HOM monomers and dimers. An overview of different oxidation product generations is given in Table 2. We observe HOM monomers with 12-16 H 235

and all compounds with an even number of H are closed shell products. Compounds with an uneven 236 237 number of H are open shell molecules (radicals). Following the proposed termination scheme by Mentel et al. (2015) compounds with 12 H can be identified as first generation monomers (terminated from 238  $C_9H_{13}O_x$  radicals, reaction 3) and compounds with 16 H as second generation monomers (terminated 239 from C<sub>9</sub>H<sub>15</sub>O<sub>x</sub> radicals, reaction 4). C<sub>9</sub>H<sub>14</sub>O<sub>x</sub> can be either first or second generation products and 240 241 originate from either  $C_9H_{13}O_x$  or  $C_9H_{15}O_x$  (reactions 5 and 6).

- $C_9H_{13}O_x + RO_2 \rightarrow C_9H_{12}O_{x-1} + ROH + O_2$ (3)242
- 243  $C_9H_{15}O_x + RO_2 \rightarrow C_9H_{16}O_{x-1} + R'CHO + O_2$ (4)
- $C_9H_{13}O_{x+1} + RO_2 \rightarrow C_9H_{14}O_x + R'CHO + O_2$ (5) 244

245 
$$C_9 H_{15} O_{x+1} + RO_2 \to C_9 H_{14} O_x + ROH + O_2$$
 (6)

246 The RO<sub>2</sub> radicals C<sub>9</sub>H<sub>13,15</sub>O<sub>x</sub> can form dimers likely via the reaction

$$HOM - RO_2 + HOM - RO_2 \rightarrow HOM - DIMER + O_2 \tag{7}$$

Such reaction is possible between any HOM-RO<sub>2</sub>, first or second generation (and other C<sub>9</sub> peroxy 248 249 radicals with sufficient high abundance). Two first generation HOM-RO<sub>2</sub> will result in C<sub>18</sub>H<sub>26</sub>O<sub>x</sub> dimers 250 (reaction 8) while one first and one second generation HOM-RO<sub>2</sub> produce a  $C_{18}H_{28}O_x$  (reaction 9) and dimerization of two second generation HOM-RO2 will result in dimers of the formula C18H30Ox (reaction 251 10). 252

253 
$$C_9H_{13}O_x + C_9H_{13}O_x \to C_{18}H_{26}O_{x-2} + O_2$$
 (8)

247

$$C_9 H_{13} O_x + C_9 H_{15} O_x \to C_{18} H_{28} O_{x-2} + O_2$$
(9)

$$C_9 H_{15} O_x + C_9 H_{15} O_x \to C_{18} H_{30} O_{x-2} + O_2 \tag{10}$$

A closer examination of the contribution of different HOM generations in Table 2 shows that 256 257 first generation monomers with 12 H are showing the highest contribution in experiments with low OH exposure (exp 1, 2) while monomers with 14 H gain importance with higher OH exposure. The second 258 generation monomers with 16 H dominate in experiments with the highest OH exposure (exp 3, 4). The 259 dimer population with 28 H has a larger fraction of the total signal at higher OH exposures compared to 260 261 dimers with 26 H. Dimer population with 30 H is generally lower than other dimers but has the highest 262 fraction in exp 4. The overall dimer fraction of up to 43% in this study (Figure 2) is similar to the dimer fraction of 40% reported by Molteni et al. (2018). However, the relative contributions of monomer and 263 dimer generations differ. We find higher contributions of  $H_{12}$  monomers (up to 11%) and a higher 264 contribution of H<sub>28</sub> dimer (up to 12%) under our experimental conditions. The contribution of H<sub>14</sub> 265 monomers and  $H_{26}$  dimers is significantly less compared to Molteni et al. (2018). 266

Increasing OH exposure promotes second OH attacks on oxidation products leading to the 267 observed reduction of first generation products ( $C_9H_{12}O_x$ ) as well as increase of second generation 268 products (C<sub>9</sub>H<sub>14</sub>O<sub>x</sub> and C<sub>9</sub>H<sub>16</sub>O<sub>x</sub>). The C<sub>9</sub>H<sub>14</sub>O<sub>x</sub> products have mainly characteristics of second 269 270 generation products, as their contribution is enhanced in the experiments with higher OH exposures 271 (Table S1), in which there is an enhanced possibility for secondary chemistry initiated by reaction of OH with the first generation products. The increased oxidation degree can also explain the formation of 272 dimers (C<sub>18</sub>H<sub>28</sub>O<sub>x</sub>) from first and second generation RO<sub>2</sub> (C<sub>9</sub>H<sub>13</sub>O<sub>x</sub> and C<sub>9</sub>H<sub>15</sub>O<sub>x</sub>) at higher OH exposure 273 and the increase in second generation dimers. Open shell species are observed as first generation  $RO_2$ 274  $(C_9H_{13}O_x)$  and have a higher contribution lower at OH exposures. Second generation RO<sub>2</sub>  $(C_9H_{15}O_x)$ 275 276 have the highest contribution in exp 3. At the highest OH exposures in exp 4, the contribution of  $C_9H_{15}O_x$ radicals, one of the top ten contributors to the signal (Table S1), is reduced, while the contribution of 277 278 the second generation products  $(C_9H_{14}O_x \text{ and } C_9H_{16}O_x)$  and dimers increased.

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Compound family	1	2	3	4	NOx9	NOx3 <sub>L</sub>	NOx3 <sub>H</sub>	NOx1
$C_9H_{12}O_x$	11.3	7.8	3.5	5.4	8.5	9.3	5.9	5.4
$C_9H_{13}O_x$	5.7	4.5	3.3	2.1	6.0	7.0	5.6	4.2
$C_9H_{14}O_x$	8.3	10.8	9.9	17.4	4.3	6.6	8.0	13.0
$C_9H_{15}O_x$	5.6	7.2	9.4	4.1	2.4	3.3	4.6	4.7
$C_9H_{16}O_x$	4.8	8.0	7.7	14.5	1.5	2.5	4.8	10.3
$C_{18}H_{26}O_x$	8.5	9.1	8.5	9.3	0.8	2.0	2.7	6.0
$C_{18}H_{28}O_x$	7.1	9.9	9.3	11.0	0.4	1.2	2.6	6.9
$C_{18}H_{30}O_x$	0.7	1.0	2.3	2.6	0.4	0.4	0.9	1.7
$C_9H_{13}NO_x$	6.2	4.7	0.6	0.6	26.8	17.1	10.1	4.8
C <sub>9</sub> H <sub>15</sub> NO <sub>x</sub>	6.4	5.6	0.7	0.9	3.7	10.3	14.5	8.5
$C_9H_{14}N_2O_x$	2.1	1.7	1.0	1.1	18.7	11.8	7.5	1.9

**Table 2: Contribution of oxidation product families to the total signal between 270 – 560** *m/z* 

In this study the contribution of  $C_{18}H_{28}O_x$  shows that both first and second generation HOM-RO<sub>2</sub> were dimerising. The kinetics of dimer formation, if produced from RO<sub>2</sub> self-reaction, depends on the square of [HOM-RO<sub>2</sub>] and their relative importance will increase with the RO<sub>2</sub> concentration. Increased local RO<sub>2</sub> concentrations (in the first part of Go:PAM) would explain the increase of dimers with increasing OH exposure.

Substantial particle formation was observed in exp 4, under the highest OH exposure. Although 287 288 the amount of reacted TMB in exp 3 and 4 is similar (26 and 30 ppb respectively), significant particle 289 formation was not observed under the conditions of exp 3. The rate at which new particle formation (nucleation) occurs is related to the chemical composition and concentration of the nucleating species 290 (McGraw and Zhang, 2008). After reaching the critical nucleus, particle growth becomes spontaneous 291 in the presence of condensable vapour. Apparently, the local concentration of nucleating species or 292 condensable vapour was not high enough in exp 3 to yield large numbers of particles, compared to exp 293 294 4. According to recent studies (Ehn et al., 2014;Trostl et al., 2016;Mohr et al., 2017;McFiggans et al., 2019) dimers play an important role in new particle formation. Mohr et al. (2017) found decreased levels 295 296 of gas phase dimers in ambient air during NPF events, which is in line with our observations of lower 297 dimer levels in the presence of particles in exp 4, compared to exp 3. A large enough concentration of 298 low volatility dimers obviously helps forming critical nucleii that then grow by condensation. Note that the newly formed particles will provide an additional sink for dimers and thus reduce their presence as 299 observable gas phase products at the end of the flow reactor. 300

### 302 Influence of NO<sub>x</sub>

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303 In the experiments NOx1, NOx3<sub>H</sub>, NOx3<sub>L</sub> and NOx9, the NO<sub>x</sub> levels were increased (Table 1). As already has been described NO<sub>x</sub> was introduced to the Go:PAM as NO. After the addition of ozone, 304 305 the ozone concentration decreases from 100 ppb to  $\sim$ 80 ppb at the experiment with lower NO<sub>x</sub> levels 306 and to  $\sim$ 50 ppb at the experiment with higher NO<sub>x</sub> levels, as it reacts with NO producing NO<sub>2</sub>. For both 307 high and low NO<sub>x</sub> conditions there is NO left after the initial reaction with ozone (see grey areas of Figure S3). The presence of NO<sub>x</sub> gave nitrogen containing C<sub>9</sub> compounds with one or two N atoms and 308 C<sub>18</sub> compounds with only one N atom, in addition to HOM monomers and HOM dimers. These 309 compounds are expected to be nitrates or peroxy nitrates, as it is highly unlikely to form nitro-aromatic 310 compounds from TMB (Sato et al., 2012). N-containing compounds were the dominating species in 311 experiments with NO<sub>x</sub> (see Figure 2), except for the experiment NOx1 with the smallest amount of NO<sub>x</sub> 312

added and a high OH exposure (NOx1). The amount of nitrated compounds increased with the amount of added NO<sub>x</sub> at the expense of HOM monomers and HOM dimers as illustrated by the arrows in Figure 2. The effect of the added NO<sub>x</sub> is attenuated at higher OH exposure. E.g., the nitrated monomers are reduced from 35.0 and 38.9% at low OH exposure down to 17.7 and 30.2% at high OH exposure. Under elevated NO<sub>x</sub> conditions nitrated species were found among the 10 compounds with the highest contribution to the respective total signal, (see the top-ten lists Table S1 and Figure 4).

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Figure 4: Comparison of mass spectra of HOM and nitrates.Panel A shows experiment 4 (blue), NOx1 (orange) and experiment NOx3<sub>H</sub> (brown). Panel B shows experiments 2 (blue), NOx3<sub>L</sub>

### 323 (orange) and NOx9 (brown).

For the two experiments with low OH exposure where the nitrated species dominated, seven out 324 of the top ten species were nitrated with the highest signal arising from the di-nitrated compound 325  $(C_9H_{14}N_2O_{10})$ . In these two experiments the top-ten compounds contributed the most to the observed 326 signal (42.3 and 52.4%); most likely owing to the high fraction of nitrated species acting as radical chain 327 termination products. Under highest NOx conditions, the high OH exposure experiment (NOx $3_{\rm H}$ ) has 328 six nitrated compounds in the top-ten list with the di-nitrated  $C_9H_{14}N_2O_{10}$  on the second rank. In 329 experiment NOx1 only one nitrated species (C<sub>9</sub>H<sub>15</sub>NO<sub>10</sub>) is found in the top-ten list despite the elevated 330 NOx conditions. 331

Dimer formation is drastically reduced in the presence of elevated  $NO_x$ . The  $NO_x$  effect on NPF was clear and high  $NO_x/\Delta$ TMB supresses NPF from TMB oxidation, a trend that was also observed by Wildt et al. (2014) and Lehtipalo et al. (2018) for NPF from monoterpene oxidation. Whenever the products were dominated by ON (NOx3<sub>L</sub>, NOx9 and NOx3<sub>H</sub>) particle formation was not observed. Reduced particle formation was observed in NOx1 compared to exp 1 which had a higher contribution of HOM. Owing to the importance of dimers for NPF, as reported by Lehtipalo et al. (2018), we suggest

- that the reaction of  $RO_2$  + NO resulting in ON is responsible for the observed reduced particle formation,
- because it competes with the dimer formation from  $RO_2 + RO_2$ , This reaction is also reducing the
- contribution of HOM monomers and dimers with increasing  $NO_x/\Delta TMB$  and decreasing OH exposure
- from a total of 61% in exp NOx1 (OH exposure= $9.1 \times 10^{10}$  molecules s cm<sup>-3</sup>) to 27.5% in NOx9 (OH exposure= $6.3 \times 10^9$  molecules s cm<sup>-3</sup>). The reaction RO<sub>2</sub> + NO reduce the amount of HOM monomers
- and dimers in competing with autoxidation and termination by  $RO_2+RO_2$  or  $RO_2+HO_2$ . The yield of ON
- from NO+RO<sub>2</sub> might be significant (e.g. up to 0.3) based on the measurable increase of ON and the
- 345 decrease of NO<sub>x</sub> in the system (Figure S3).
- The contribution of HOM monomer generations follows the general trend observed in 346 347 experiments without NO<sub>x</sub>. The first generation HOM have higher contribution at low OH exposures in 348 NOx3<sub>L</sub> and NOx9 and the contribution of second generation HOM is higher the higher the OH exposure is. Similarly, the contribution of C<sub>9</sub>H<sub>14</sub>O<sub>x</sub> is increasing with increasing OH exposure. Analogously to the 349 families of HOM, different families of nitrates can be defined. Table 2 gives an overview of the nitrate 350 families and how they contribute to the total signal in the experiments. At lowest OH exposures we find 351 352 the highest contributions of first generation nitrates ( $C_9H_{13}NO_x$ ) as well as di-nitrates ( $C_9H_{14}N_2O_x$ ). The contribution of second generation nitrates (C<sub>9</sub>H<sub>15</sub>NO<sub>x</sub>) is increasing with increasing OH exposure and is 353 354 highest in NOx3<sub>H.</sub>
- For the formation of observed first generation nitrates  $C_9H_{13}NO_{6-12}$  ( $C_9H_{13}NO_x$  in Table 2) we propose the reaction of a first generation (HOM-)RO<sub>2</sub> with NO following the pathway:
- 357  $C_9H_{13}O_{5-11} + NO \to C_9H_{13}NO_{6-12}$  (11)
- The precursor  $C_9H_{13}O_5$  is formed after one autoxidation step and its termination reaction with NO results in  $C_9H_{13}NO_6$  which has only minor contribution.  $C_9H_{13}NO_{7\&8}$ , with higher contribution in exp NOx3<sub>L</sub> and NOx9 can be formed from the radicals  $C_9H_{13}O_6$  and  $C_9H_{13}O_7$  respectively. The even oxygen number in  $C_9H_{13}O_6$  indicates that the compound should have undergone a transformation to RO (via reaction with RO<sub>2</sub> or NO) and subsequent H-shift and further O<sub>2</sub> addition (Vereecken and Peeters, 2010;Mentel et al., 2015). A proposed detailed reaction mechanism is depicted in the Figure 5.
- Second generation nitrates (C<sub>9</sub>H<sub>15</sub>NO<sub>8-12</sub>) can be formed after an additional OH attack (and introduction 364 365 of an additional H) on a first generation (HOM) monomer, which explains the increase of these compounds with increasing OH exposure. The termination of the RO<sub>2</sub> radical chain with NO (reaction 366 12) will then lead to the formation of the second generation nitrate. The formation of  $RO_2$  precursor 367 species with 7-8 O numbers, i.e. C<sub>9</sub>H<sub>15</sub>O<sub>7-8</sub> likely stem from compounds terminated earlier in the radical 368 chain process ( $C_{9}H_{14}O_{4-5}$ ), which do not fall in the typical HOM class ( $O:C \ge 6:9$ ). The reaction of a first 369 generation nitrate with OH, followed by autoxidation could also possibly produce C<sub>9</sub>H<sub>15</sub>NO<sub>8-12</sub> by 370 371 terminating via peroxy (reaction 13) or hydroperoxy pathway (reaction 14).
- 372  $C_9H_{15}O_{7-11} + NO \rightarrow C_9H_{15}NO_{8-12}$  (12)
- 373  $C_9H_{14}NO_{9-13} + RO_2 \rightarrow C_9H_{15}NO_{8-12} + Carbonyl products + O_2$  (13)

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$$C_9H_{14}NO_{8-12} + HO_2 \rightarrow C_9H_{15}NO_{8-12} + O_2$$
 (14)

For the formation of the most abundant di-nitrates of the formula  $C_9H_{14}N_2O_{8-12}$  (reaction 15), OH has to attack a nitrated compound  $C_9H_{13}NO_{6-10}$  and the RO<sub>2</sub> radical chain has to be terminated with NO (Figure 5).

- 378  $C_9H_{14}NO_{7-11} + NO \rightarrow C_9H_{14}N_2O_{8-12}$  (15)
- The precursor species  $C_9H_{14}NO_7$  would be in this case formed from the OH attack on the smallest possible nitrate  $C_9H_{13}NO_4$  (formed after one autoxidation step and NO termination).

- 381 It should be noted that the formation of peroxy nitrates via the reaction  $RO_2 + NO_2 \rightarrow RO_2NO_2$  with
- alkyl and acyl-RO<sub>2</sub> (PAN-like) cannot be ruled out as a potential formation mechanisms of nitrates.

The volatility of the produced ON seems to be too high (compared to dimers) to initiate NPF at the

384 present concentrations as seen from the reduction in particle formation potential from exp 4 (HOM

dominated, particle formation) to NOx1 (reduction in HOM and increase in ON, reduced particle

386 formation) and NOx $3_{\rm H}$  (ON dominated, no particle formation).



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In Figure 2B, the calculated HOM dimer contribution are the sum of medium (produced from
 HOM-RO<sub>2</sub>+RO<sub>2</sub>), and highly oxidised dimers (produced from HOM-RO<sub>2</sub> + HOM-RO<sub>2</sub>) while the ON

Figure 5: The proposed radical reaction mechanism for the formation of some of the mono- and di-nitrates from TMB mentioned in this study. Two separate mechanisms were suggested for the species with the formula  $C_9H_{13}NO_8$ , which formation pathways are based on (a) (Wang et al., 2017) and (b) (Molteni et al., 2018).

<sup>392</sup> Figure 2B shows the results for the evolution of HOM monomer, HOM Dimers and organic 393 nitrates (ON) as calculated using the kinetic model described in detailed in SI. The model used has a 394 very simplified scheme for TMB oxidation and subsequent RO<sub>2</sub> chemistry. We implemented a few rate coefficients suggested in the literature in order to demonstrate how those compare with our experimental 395 results. Rate coefficients for RO<sub>2</sub> termination reactions from MCM in combination with rate coefficients 396 397 of dimer formation by Berndt et al. (2018) (case 1 and 2 in SI Table S2) led to an overestimation of dimer compounds. Better representation of the observations were achieved by applying a) the rate 398 coefficients proposed by (Zhao et al., 2018) for dimer formation (2.0×10<sup>-12</sup> cm<sup>3</sup>molecules<sup>-1</sup> s<sup>-1</sup>) and 399  $1.0 \times 10^{-12}$  cm<sup>3</sup> molecules<sup>-1</sup> s<sup>-1</sup> for termination and RO formation with branching ratios of 0.4 and 0.6, 400 respectively; b) the rate coefficient  $(1.0 \times 10^{-11} \text{ cm}^3 \text{ molecules}^{-1} \text{ s}^{-1})$  from Berndt et al. (2018) for HOMRO<sub>2</sub> 401 + NO with a branching ratio of 0.3 for ON formation (reaction 56 in Table S2). 402

405 includes both organic nitrates (produced from HOM-RO<sub>2</sub>+NO) and peroxy nitrates (produced from HOM-RO<sub>2</sub>+NO<sub>2</sub>). The increased production of monomers calculated for exp 1 and 2 is in agreement 406 with experimental results, where we observed a slightly larger contribution from monomers compared 407 to dimers. Calculated concentrations for dimers are similar in exp 3 and higher in exp 4, compared to 408 monomers. Secondary chemistry (reaction of OH with the products and possible formation of a second 409 410 generation of HOM or nitrates) was not taken into consideration in the model. For the experiment with 411 NO<sub>x</sub> (Figure 2B) the modeled product distribution follows the general trend of monomer, dimer and 412 nitrate that we observe in the exp NOx3<sub>L</sub> and NOx9 with a general higher nitrate production compared 413 to monomers and dimers. For the NO<sub>x</sub> experiments with high initial ozone, the model can reproduce the 414 higher HOM monomer and dimer levels in NOx1 but slightly overestimates the contribution of dimers. 415 Particle formation was observed in NOx1 which might explain the overestimation owing to missing condensation sink in the model. Modelling NOx3<sub>H</sub> gives an overestimation of monomers and dimers. 416 In NOx3<sub>H</sub> modelled dimers start forming after  $\sim$ 15 sec. Almost all NO is converted to ON or NO<sub>2</sub> at this 417 point and the reaction HOM-RO2+NO does not produce additional ON and the modelled levels of ON 418 419 reach a plateau while contribution of HOM dimers can increase. For exp NOx1 and NOx3<sub>H</sub> the model is in better agreement with the measurements if only the HOM - dimer formation (from HOM-420 RO<sub>2</sub>+HOM-RO<sub>2</sub>) is taken into account, i.e. highly oxidized dimers excluding medium oxidized dimer 421 (from HOM-RO<sub>2</sub> + RO<sub>2</sub>). Generally, the simplified model was able to support our analysis of the TMB 422

chemistry as described above. Furthermore, it gave some support to the recently suggested mechanisms
 included in case 3 (e.g. (Zhao et al., 2018))

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#### 4 Atmospheric implication and Conclusion

427 We have measured the formation of HOM monomers and dimers from OH initiated oxidation 428 of TMB. The experiments with highest OH exposures lead to particle formation when  $NO_x$  was not 429 added. With increasing OH exposure and increased likelihood of a second OH attack, we observe a 430 higher contribution from second generation oxidation products and dimers in general. The latter is 431 attributed to the increased RO<sub>2</sub> concentrations from the increased/fast TMB consumption by OH. The 432 observed products in this study match what would be expected as termination products from previously 433 proposed reaction mechanisms for HOM formation.

The addition of NO<sub>x</sub> to simulate urban condition leads to the formation of ON in addition to 434 435 HOM and a reduction in particle formation potential. We observe that the formation of ON is increasing with increasing  $NO_x/\Delta TMB$ , mostly at the expense of dimers. The presence of ONs, formed at the 436 expense of dimers, can explain the decreased tendency for particle formation. We therefore suggest that 437 the reaction HOM-RO<sub>2</sub>+NO competes with HOM-RO<sub>2</sub> self-reaction yielding primarily a reduction in 438 dimer formation, which is responsible for the reduction in particle formation. The experimental designed 439 using the Go:PAM with concentrations of  $HO_x$  (and  $RO_x$ ) higher than ambient would attenuate the 440 influence of added NO<sub>x</sub>. This will further emphasis the implication of our findings and most likely the 441 442 NO<sub>x</sub> effect would be even more important in the urban atmosphere.

443 According to studies by (Molteni et al., 2018) and (Wang et al., 2017) HOM formation from 444 AVOCs was observed and consequently AVOC-HOM were suggested as potential contributors to 445 observed NPF in urban atmospheres. In our study, under  $NO_x$  free conditions, we found several of 446 previous identified HOM even if we did not fully agree with the identity and relative importance of all 447 HOM. However, the oxidation in polluted environments will happen under elevated  $NO_x$  levels and, as 448 has been shown here this can lead to formation of ON instead of HOM and subsequently a reduction in 449 NPF potential. We conclude that for interpretation of NPF from aromatics in urban areas care should be

- 450 taken and the OH exposure,  $NO_x$  levels and  $RO_2$  concentrations need to be considered in details since
- 451 they will largely determine if the HOM-RO<sub>2</sub>+NO can compete with reactions yielding HOM, and
- 452 especially HOM dimers.
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- 455 *Data availability*: The data set is available upon request by contacting Mattias Hallquist 456 (hallq@chem.gu.se).
- 457 *Competing Interests*: The authors declare that they have no conflict of interest.
- 458 *Author contribution*: J.H., E.T. T.M. and M.H. designed the experiments. J.H and E.T performed the 459 experiments and data analysis. E.T. performed the modelling. C.M.S. designed the chemical 460 mechanisms. E.T., J.H. and M.H. wrote the paper. All authors commented on the paper and were 461 involved in the scientific interpretation and discussion.
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