

Molecular composition and volatility of multi-generation products formed from isoprene oxidation by nitrate radical

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Abstract

Isoprene oxidation by nitrate radical (NO_3) is a potentially important source of secondary organic aerosol (SOA). It is suggested that the second or later-generation products are the more substantial contributors to SOA. However, there are few studies investigating the multi-generation chemistry of isoprene- NO_3 reaction, and information about the volatility of different isoprene nitrates, which is essential to evaluate their potential to form SOA and determine their atmospheric fate, is rare. In this work, we studied the reaction between isoprene and NO_3 in the SAPHIR chamber (Jülich) under near atmospheric conditions. Various oxidation products were measured by a high-resolution time-of-flight chemical ionization mass spectrometer using Br^- as the reagent ion. Most of the products detected are organic nitrates, and they are grouped into monomers (C_4 - and C_5 -products), and dimers (C_{10} -products) with 1–3 nitrate groups according to their chemical composition. Most of the observed products match expected termination products observed in previous studies, but some compounds such as monomers and dimers with three nitrogen atoms were rarely reported in the literature as gas-phase products from isoprene oxidation by NO_3 . Possible formation mechanisms for these compounds are proposed. The multi-generation chemistry of isoprene and NO_3 is characterized by taking advantages of the time behavior of different products. In addition, the vapor pressures of diverse isoprene nitrates are calculated by different parametrization methods. An estimation of the vapor pressure is also derived from their condensation behavior. According to our results, isoprene monomers belong to intermediate volatility or semi-volatile organic compounds and thus have little effect on SOA formation. In contrast, the dimers are expected to have low or

43 extremely low volatility, indicating that they are potentially substantial contributors to SOA. However, the
44 monomers constitute 80% of the total explained signals on average, while the dimers contribute less than 2%,
45 suggesting that the contribution of isoprene NO₃ oxidation to SOA by condensation should be low under
46 atmospheric conditions. We expect a SOA mass yield of about 5 % from the wall loss and dilution corrected
47 mass concentrations, assuming that all of the isoprene dimers in the low- or extremely low-volatility organic
48 compound (LVOC or ELVOC) range will condense completely.

49 1. Introduction

50 Atmospheric submicron aerosols have an adverse effect on air quality, human health and climate (Jimenez et al.,
51 2009; Pöschl, 2005). Secondary organic aerosol (SOA), which is formed from oxidation of volatile organic
52 compounds (VOC) followed by gas-to-particle partitioning, comprise a large fraction (20-90%) of the
53 submicron aerosol mass (Jimenez et al., 2009; Zhang et al., 2007). It is confirmed that a significant proportion of
54 SOA arises from biogenic VOC (BVOC) oxidation (Hallquist et al., 2009; Spracklen et al., 2011).

55 Isoprene is globally the most abundant non-methane volatile organic compound originating from
56 vegetation, with emissions estimated to be 440-660 Tg yr⁻¹(Guenther et al., 2012). Due to its high abundance, as
57 well as its high reactivity with atmospheric oxidants, isoprene plays a significant role in tropospheric chemistry,
58 and its chemistry affects the global aerosol burden and distribution (Carlton et al., 2009; Fry et al., 2018; Ng et
59 al., 2008, 2017; Surratt et al., 2010), although its SOA yield is much lower than those of monoterpenes and
60 sesquiterpenes (Friedman and Farmer, 2018; Kim et al., 2015; Marais et al., 2016; , McFiggans, et al. 2019;
61 Mutzel et al., 2016; Ng et al., 2007, 2008; Surratt et al., 2010; Thornton et al., 2020). Recent model simulations
62 suggested the isoprene-derived SOA production is 56.7 Tg C yr⁻¹, contributing up to 41% of global SOA
63 (Stadtler et al., 2018). Observations in southeastern United States suggested that isoprene-derived SOA makes
64 up 17- 48% of total organic aerosol (Hu et al., 2015; Kim et al., 2015; Marais et al., 2016). As a consequence, it
65 is essential to fully characterize the potential of isoprene to form condensable products and its contribution to
66 SOA formation (Carlton et al., 2009).

67 Although the majority of isoprene emissions is emitted by plants and is light-dependent, isoprene emitted
68 in the day can persist in the boundary layer after sunset, and its mixing ratio can remain as high as several ppb
69 (Brown et al., 2009; Starn et al., 1998; Stroud et al., 2002; Warneke et al., 2004). During the daytime, isoprene
70 is primarily oxidized by the hydroxyl radical (OH) and somewhat by ozone (O₃), but its main oxidizers shift to
71 nitrate radical (NO₃) and O₃ in the nighttime (Wennberg et al., 2018). Due to the higher reactivity of NO₃ with
72 isoprene ($k_{NO_3} = 6.5 \times 10^{-13} \text{ cm}^3 \text{ molecules}^{-1}\text{s}^{-1}$ and $k_{O_3} = 1.28 \times 10^{-17} \text{ cm}^3 \text{ molecules}^{-1}\text{s}^{-1}$ at 298 K, respectively,
73 IUPAC), a considerable fraction of the residual isoprene would be oxidized by NO₃ at night, and therefore
74 nocturnal nitrate radical chemistry is typically thought to be of significant importance for isoprene, especially in
75 regions where sufficient nitrogen oxides are available (Brown et al., 2009; Fry et al., 2018; Ng et al., 2017;
76 Wennberg et al., 2018). Although reaction with NO₃ only represents ~ 5-6% of isoprene loss, it accounts for a
77 large proportion of organic nitrates derived from isoprene oxidation (~ 40-50%) (Wennberg et al., 2018).
78 Therefore, reaction of isoprene with NO₃ is a potential source of SOA. In addition, it is found from both
79 laboratory and chamber experiments that the SOA yield of isoprene from NO₃ oxidation is higher than that from
80 OH or O₃ oxidation, which is typically less than 5% (Carlton et al., 2009; Dommen et al., 2009; Kleindienst et
81 al., 2007; Kroll et al., 2006). For example, Ng et al. (2008) concluded the isoprene SOA yield from NO₃ was in
82 the range of 4.3% to 23.8%, depending on RO₂ fate (higher SOA yield when the experiments were dominated
83 by RO₂+RO₂ rather than RO₂+NO₃ reaction). Rollins et al. (2009) also observed a high SOA yield from
84 isoprene (14%) when both of its double bonds were oxidized by NO₃. In an aircraft study in the southeastern
85 United States, Fry et al. (2018) derived an isoprene-NO₃ SOA yield as large as 27% on average under high NO_x
86 conditions, although their mass yield estimation was indirect, and based on a molar yield determination of 9 ±
87 5%. In light of the relatively high SOA yield from NO₃ oxidation, even though only a minor fraction of isoprene

88 is oxidized by NO₃, the SOA formed at nighttime would still probably be comparable to that produced at
89 daytime (Brown et al., 2009; Fry et al., 2018).

90 However, isoprene-NO₃ chemistry (Wennberg et al. 2018, Vereecken et al. 2021) has received less
91 attention than the extensively studied OH- or O₃-initiated oxidation (Barber et al., 2018; Novelli et al., 2020;
92 Peeters et al., 2014; Wang et al., 2018; Wennberg et al., 2018; Whalley et al., 2012). It has been recognized that
93 later-generation oxidation of isoprene by NO₃ makes more significant contribution to SOA formation (Carlton et
94 al., 2009; Fry et al., 2018; Rollins et al., 2009). Nevertheless, although the importance of multi-generation NO₃
95 oxidation of isoprene to SOA formation has been recognized, few studies extended the investigation beyond the
96 first-generation oxidation, and details of isoprene-NO₃ multi-generation chemistry are still lacking.

97 Organic compounds, especially highly oxygenated organic molecules (HOM) that have low or extremely
98 low volatility, contribute significantly to SOA formation by condensation, or even form new particles (Bianchi
99 et al., 2019; Ehn et al., 2014; Kirkby et al., 2016, Tröstl et al., 2016). Previous studies have confirmed that low-
100 volatility products from isoprene-NO₃ reaction are the major precursors to SOA (Ng et al., 2008; Rollins et al.,
101 2009; Schwantes et al., 2019). Here the low-volatility compounds refer to gas phase products that allow
102 fractions to exist in particle-phase, and may include the groups of organic compounds with intermediate
103 volatility (IVOC, $300 < C^* < 3 \times 10^6 \mu\text{g m}^{-3}$), semi-volatility (SVOC, $0.3 < C^* < 300 \mu\text{g m}^{-3}$), low volatility (LVOC,
104 $3 \times 10^{-5} < C^* < 0.3 \mu\text{g m}^{-3}$) and extremely low volatility (ELVOC, $C^* < 3 \times 10^{-5} \mu\text{g m}^{-3}$) as proposed by Donahue et al.
105 (2012). In general, SVOC, LVOC and ELVOC can contribute to the SOA formation (Jimenez et al., 2009). In
106 order to evaluate the potential of oxygenated products to form SOA, information about their vapor pressures is
107 essential. However, due to the high degree of functionalization, low or extremely low volatility, as well as
108 uncertainties in quantification and molecular structures, it is challenging to determine the exact vapor pressure
109 of highly oxidized products. Detailed information on the volatilities of different generation products is lacking,
110 which impedes the assessment of their contribution to SOA formation.

111 In this work, we present the results of chamber experiments on isoprene oxidation by NO₃ under near
112 atmospheric conditions, where NO₃ was produced in situ by O₃ reaction with NO₂. Subsequent characteristics of
113 multi-generation chemistry of isoprene with NO₃ are investigated. By examining the time evolution of various
114 gas-phase products, we propose possible reaction mechanisms that help to get the possible functionalization of
115 the products. Saturation vapor pressures of the major gas-phase products observed by HR-ToF-CIMS are
116 predicted by using different parameterization methods that are widely-used or state-of-the-art in literature. In
117 addition, we estimate the vapor pressure derived from equilibrium partitioning coefficient according to the
118 condensation behavior of different products in experiments with and without seed aerosols. Based on these
119 results, the volatility of the major oxidation products stemming from isoprene-NO₃ reaction and their potential
120 to form SOA are evaluated.

121 **2. Experimental and methods**

122 **2.1 Atmospheric simulation chamber SAPHIR**

123 All the data presented here were measured in the atmospheric simulation chamber SAPHIR (Simulation of
124 Atmospheric PHotochemical In a large Reaction Chamber) at Forschungszentrum Jülich, Germany, which is
125 designed to investigate the oxidation processes of both biogenic and anthropogenic trace gases and formation of

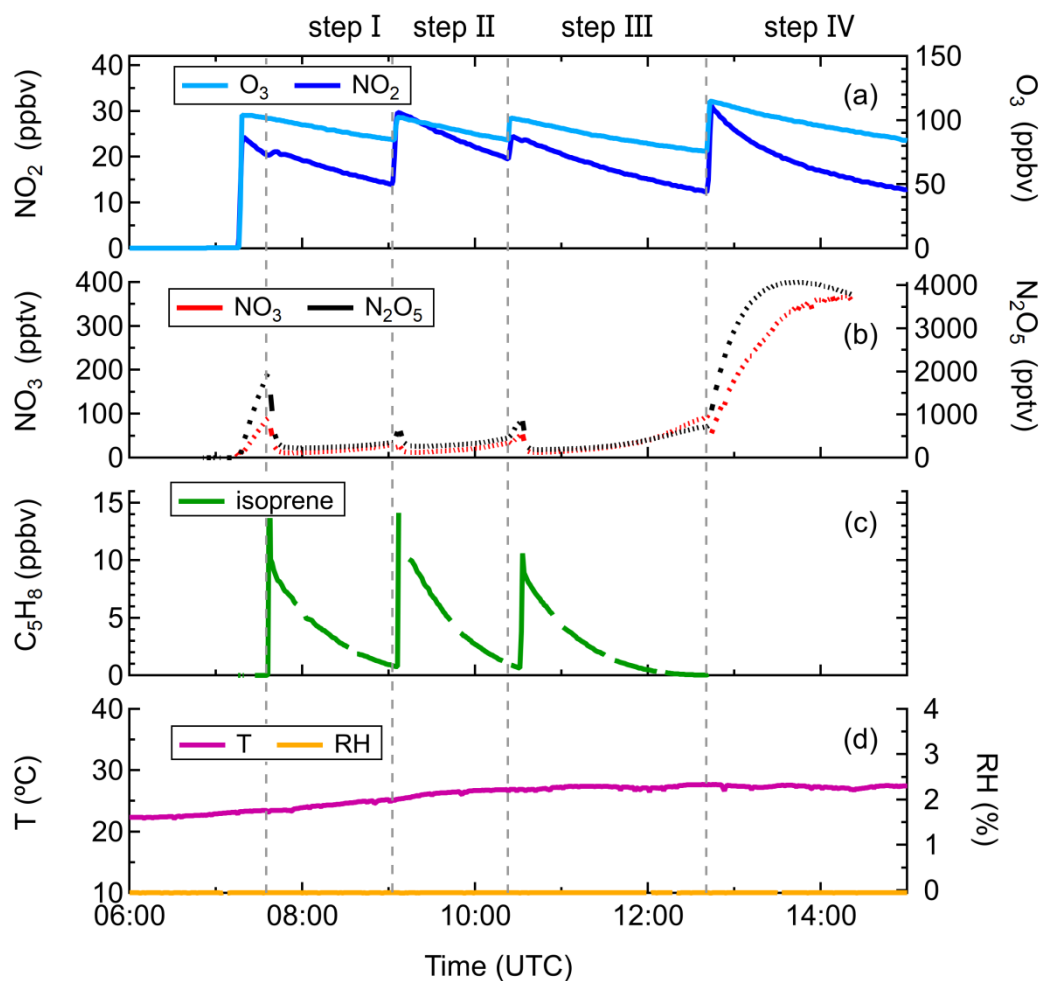
126 secondary particles and pollutants under near atmospheric conditions. The SAPHIR chamber is a double-walled
127 Teflon (FEP) cylinder with a volume of 270 m³ (5 meters in diameter and 18 meters in length). The large
128 volume-to-surface ratio (1 m) allows experiments to be conducted under natural conditions and reduces
129 interference from the chamber walls. The chamber is equipped with a shutter system which can be opened to
130 admit sunlight for photochemical experiments or closed to mimic nighttime conditions. There are two fans
131 inside the chamber to ensure good mixing of trace gases (within 2 minutes). The chamber is filled with synthetic
132 air made from mixing of ultrapure nitrogen and oxygen (Linde, purity $\geq 99.99990\%$) and is slightly over-
133 pressured (~ 35 Pa) to prevent intrusion of outside air into the chamber. Due to small leakage (~ 7 m³ h⁻¹) and
134 gas consumption by instrument sampling, a replenishment flow is provided by a flow control, which leads to a
135 dilution rate of 4%–7% per hour. A more detailed description of the chamber set-up and its characterization can
136 be found elsewhere (Rohrer et al., 2005).

137 **2.2 Experiment description**

138 A series of experiments investigating the oxidation of isoprene by NO₃ were conducted in the SAPHIR chamber
139 in August 2018 (ISOPNO₃ campaign) under different chemical conditions. In this work, we primarily focus on
140 an experiment conducted on 08 August 2018 that examined the fast oxidation of isoprene by NO₃ (up to ~ 130
141 pptv) without seed aerosols. The experiment was performed under dry (RH < 5%) and dark condition, and
142 employed injections of O₃ and NO₂ as source of NO₃, where O₃ was generated by a silent discharge ozoniser
143 (O3onia), and high-purity NO₂ was introduced from a gas bottle (Linde, purity >99%).

144 Before the experiment, the chamber was flushed overnight with a total amount of ~ 1800 m³ synthetic air to
145 minimize any remaining contamination. At the beginning of the experiment, the chamber air was slightly
146 humidified (RH < 0.1%) by flushing water vapor from boiling Milli-Q[®] water into the chamber. Thereafter, O₃
147 and NO₂ were added to the chamber in succession, and their concentrations in the chamber after injection were
148 approximately 100 and 25 ppbv, respectively, as shown in Fig. 1. After that, ~ 10 ppbv of isoprene was injected
149 using a GC syringe, initiating the reaction with NO₃. The period between the first and second injection is
150 defined as “step I”, as so on for the other three periods. The second injection was done when isoprene from the
151 first injection was almost completely consumed, to reach concentrations of O₃, NO₂, and isoprene in the
152 chamber of ~ 100 , 30, and 10 ppbv, respectively. About 1.5 hours later, the chemistry was further accelerated by
153 a third injection of precursors, and accordingly the concentrations of O₃, NO₂, and isoprene reached ~ 100 , 25,
154 and 10 ppbv, respectively. Two hours later, the fourth addition was made and the concentrations of O₃ and NO₂
155 increased to approximately 115 ppbv and 30 ppbv, respectively, aiming to promote further oxidation of early
156 generation products. In total the system was kept running for about 7.5 h. Calculation from measurements of
157 isoprene, O₃, OH, NO₃ and dilution indicates that NO₃ contributed for more than 90% of the chemical losses of
158 isoprene, as shown in Fig. S1, with reaction with O₃ being a minor pathway in our system. The reaction of
159 isoprene with OH was not considered as OH concentration was below the detection limit of the instrument in
160 this study (Fig. S2). Thus, losses due to reaction with OH could not be quantified from the measurement, but
161 have been determined to contribute about 10% of the isoprene losses according to a recently published
162 modelling work based on the same campaign, with the contribution of the NO₃ reaction accounting for up to 80%
163 accordingly (Vereecken et al., 2021).

164 A complementary experiment was conducted on 14 August 2018 under similar conditions but with seed
 165 aerosols. Approximately $60 \mu\text{g m}^{-3}$ of ammonium sulfate aerosol was added at the beginning of the experiment.
 166 Thereafter, approximate 100 and 20 ppbv of O_3 and NO_2 were introduced to the chamber to produce NO_3 ,
 167 followed by addition of ~ 10 ppbv of isoprene in about 30 minutes later (see Fig. S3). Another 6 ppbv of NO_2
 168 and 10 ppbv of isoprene were added about one hour later to accelerate the reaction. At the last injection, only O_3
 169 (~ 50 ppbv) and NO_2 (~ 7 ppbv) were added, similar as for the experiment without seeds. The experiment lasted
 170 for about 8 h. The results were used to investigate the condensation behavior of various gas-phase products from
 171 isoprene oxidation, aiming to estimate equilibrium partitioning coefficients and vapor pressures.



172
 173 **Figure 1: Measurements of (A) O_3 and NO_2 , (B) NO_3 and N_2O_5 , (C) isoprene and (D) temperature and relative**
 174 **humidity in the chamber during the experiment on 08 August, 2018.**

175 2.3 Instrumentation

176 A high-resolution time-of-flight chemical ionization mass spectrometer (HR-ToF-CIMS, Aerodyne Research
 177 Inc., hereafter CIMS) was used to continuously measure the gas-phase products from isoprene oxidation by NO_3 .
 178 The ToF-MS was operated in ‘V’ mode with the mass resolution power between 3000–4000 Th/Th. In order to
 179 reduce the losses of HO_2 radicals and HOM on the tubing, a customized inlet (Albrecht et al., 2019) was directly
 180 connected to the chamber. The CIMS was operated in negative ion mode using Br^- as the reagent ion, which is
 181 selective to polar species such as acids, hydroxy or nitrooxy carbonyls, as well as HO_2 radicals (Albrecht et al.,
 182 2019; Ng et al., 2008; Rissanen et al., 2019; Riva et al., 2019).

183 Bromide ions were generated by passing a mixture of 10 standard cubic centimeters per minute of 0.4%
184 CF₃Br in nitrogen and 2 standard liter per minute nitrogen through a 370 MBq ²¹⁰Po source (Type P-2021-5000,
185 NDR Static Control LLC, USA), resulting in ~10⁵ ion counts per second (Albrecht et al., 2019). In our system,
186 on average, about 190 ions were identified for each mass spectrum on average, most of which were detected as
187 adducts with Br⁻, while some acidic compounds (~ 7% of the total) like nitric acid (HNO₃), glycolic acid
188 (C₂H₄O₃), and malonic acid (C₃H₄O₄) were also detected as deprotonated ions. In addition, there were some ions
189 (~ 10% of the total) identified as adducts with NO₃⁻. The isotope distribution of ⁷⁹Br and ⁸¹Br is approximately
190 1:1, therefore two signals appear at $m/z = MW+79$ and $m/z = MW+81$ with MW being the molecular mass of the
191 molecule that is detected as cluster with Br⁻. In this work, we will use Thomson (Th) as the unit for mass-to-
192 charge (m/z), and the m/z of molecules discussed in following include the mass contribution from Br⁻ (m/z 79) if
193 there is no other annotation.

194 In order to have an indicator of the CIMS performance, perfluoropentanoic acid (PFPA, C₅F₉HO₂) was
195 used as an internal standard. For m/z calibration, five isolated peaks were used, including Br⁻ (m/z 79), H₂OBr⁻
196 (m/z 97), HNO₃Br⁻ (m/z 142), C₅F₉O₂⁻ (m/z 263), and C₅F₉HO₂Br⁻ (m/z 343), covering the mass range of
197 dominant products. The averaged accuracies of all five calibrated masses were below 5 ppm over the whole
198 measurement period. However, due to the low signal intensity, the PFPA cluster (C₁₀F₁₈O₄H⁻, m/z 527) was not
199 suited for mass calibration, and there were no other suitable masses with sufficient intensity and high accuracy
200 that could be used to calibrate the higher mass range. Therefore, peak fitting in the mass range between 300 to
201 500+ Th might have higher uncertainties. The CIMS was optimized to gain a maximum signal of [HO₂*Br]⁻
202 isotopes, which are weakly bounded clusters. This was achieved by adjusting step by step the electrostatic field
203 in the transfer stage to minimize fragmentation. During the campaign, the settings of CIMS were kept
204 unchanged to keep a similar performance. However, the signal of reagent ion Br⁻ decreased by about 65% (from
205 ~ 100, 000 to 34, 000 counts s⁻¹) over the campaign duration of four weeks. In order to minimize the effect of
206 drift in performance, we used the normalized (by the sum of the total ion counts) signals for analysis. The
207 sensitivity for total carbon was calculated by determining the slope of wall-loss corrected total carbon signals
208 detected by CIMS (only the identified peaks were considered) versus isoprene consumed. As illustrated in Fig.
209 S4a, the CIMS sensitivities were roughly identical in two experiments (0.026 ± 0.002 norm. count s⁻¹ ppbv⁻¹ on
210 08 August, and 0.022 ± 0.001 norm. count s⁻¹ ppbv⁻¹ on 14 August), indicating that different experimental
211 conditions over two days had an insignificant impact on CIMS sensitivity for total carbon and thus the data from
212 these days are comparable. In addition, an inter-comparison of measurements by Br⁻ CIMS and I⁻ CIMS were
213 made. As shown in Fig. S4b, the measurements of C₃H₆N₂O₈ from the two instruments are well linearly
214 correlated with each other at the early oxidation stages. However, the correlation coefficients of measurements
215 from two instruments deviated from experiment to experiment. This is probably related to different experimental
216 conditions, which might lead to different chemical processes and thus formation of isomers. Since CIMS with
217 different reagent ions might have different sensitivities to isomers, and may be selective for different
218 compounds, the correlation coefficients of measurements from Br⁻ and I⁻ CIMS may differ from day to day.
219 Moreover, the Br⁻ CIMS was not tuned during the campaign while the I⁻ CIMS was optimized from time to time.
220 In general, the performance of Br⁻ CIMS was stable and the data taken by it are reliable.

221 The mass spectra data were processed using the software “Tofware” embedded in Igor as provided by
222 Aerodyne Research Inc. (<https://www.tofwerk.com/software/tofware/?cn-reloaded=1>). Peaks detected in the

223 mass spectra could be isolated and identified according to their exact mass, and molecular formulas and the
224 corresponding intensities were obtained by high-resolution peak fitting. Due to a lack of authentic standards for
225 the products, it is difficult to quantitatively determine their individual absolute concentrations, but we have
226 calculated the bulk sensitivity for organonitrates by using the sum of organic nitrate signals from Br CIMS
227 divided by measurements of the total alkyl nitrates from a thermal dissociation-cavity ring-down spectrometer
228 during the experiment . The estimated bulk sensitivities for organonitrates are 0.016 ± 0.001 and 0.022 ± 0.001
229 norm. count s^{-1} ppbv $^{-1}$ on 08 August and 14 August, respectively, as shown in Fig. S4c, comparable to the
230 sensitivity for total carbon, but smaller than the sensitivity for salicylic acid determined by an independent
231 calibration (163 norm. count μg^{-1} on average as shown in Fig. S4d, equal to 0.07 norm. count s^{-1} ppbv $^{-1}$). The
232 bulk sensitivity for organonitrates enables estimation of the absolute concentrations of products assuming that
233 they have identical sensitivity. In this study we use the normalized signals instead of absolute concentrations for
234 analysis. This is sufficient here because our analysis focuses on the time evolution of signals and the relative
235 changes of intensities, so the absolute concentrations are not necessarily needed. The sensitivity derived above is
236 only used to convert the signals of dimers to concentrations in order to estimate the SOA yield.

237 Isoprene was measured by a Vocus proton transfer reaction time-of-flight mass spectrometer (Aerodyne
238 Research Inc., hereafter Vocus), which has a higher mass resolving power (nominal 10000 Th/Th) and less inlet
239 wall losses and sampling delays compared to traditional PTR-MS (Krechmer et al., 2018). The mixing ratio of
240 O_3 was monitored by an UV absorption instrument, and that of NO_2 was monitored by a chemiluminescence
241 instrument and a custom-built cavity ring-down spectrometer (CRDS). The concentrations of NO_3 and N_2O_5
242 were detected by two custom-built CRDS instruments (Dubé et al., 2006; Sobanski et al., 2016). In addition,
243 temperature and pressure inside the chamber were monitored by an ultra-sonic anemometer and a pressure
244 sensor, respectively. The relative humidity was primarily detected as water mixing ratio by a Picarro CRDS
245 instrument (Crosson, 2008).

246 The particle number concentrations and their size distributions were measured by a condensation particle
247 counter (TSI 3783, hereafter CPC) and a scan mobility particle sizer (TSI 3081 electrostatic classifier combined
248 with TSI 3025 CPC, hereafter SMPS). The aerosol chemical composition was identified by a high-resolution
249 time of flight aerosol mass spectrometer (HR-ToF-AMS, Aerodyne Research Inc., hereafter AMS). The
250 ionization efficiency of AMS was determined by using the monodisperse aerosol generated from NH_4NO_3 and
251 $(NH_4)_2SO_4$ solutions. The collection efficiency (CE) could be estimated based on the particle mass concentration
252 yielded from AMS and that derived from SMPS. In this study, the average CE value of 0.5 is used for correction.

253 **2.4 Methods to estimate saturation vapor pressure**

254 The pure liquid saturation vapor pressure is a thermodynamic metric relevant for the partitioning equilibrium of
255 organic molecules, which determines their propensity to form SOA (Compernelle et al., 2011; O'Meara et al.,
256 2014; Pankow and Asher, 2008). Due to their complex functionalities and low or extremely low volatility, it is
257 challenging to determine the vapor pressures of highly oxidized molecules. As a result, theoretical and
258 semiempirical methods are usually used for vapor pressure estimation. Commonly used semiempirical methods
259 include composition-activity (CA), group-contribution (GC), and structure-activity (SA) methods. The CA
260 methods are the easiest to use, as they only require information on molecular composition for estimation. They
261 are widely applied in context of the two-dimensional volatility basis set (2D-VBS) (Donahue et al., 2011). For

GC methods, the exact functional groups are required to calculate the saturation vapor pressure. The SIMPOL.1 (Pankow and Asher, 2008), the parameterization as described by Nannoolal et al. (2008), and EVAPORATION (Compernelle et al., 2011) are three widely used GC methods. Structure-activity methods can provide more accurate estimates with sophisticated treatments of intramolecular interactions like intramolecular hydrogen-bonding (Bilde et al., 2015). However, detailed molecular properties such as boiling point and evaporation enthalpy are required for estimation, which are generally obtained by complex and time-consuming quantum chemical calculations. Therefore, SA methods are not applied for vapor pressure estimation in this study.

Saturation concentration (C_i^* , mass based) is related to saturation vapor pressure and can be calculated following Eq. (1) (Donahue et al., 2006). The $\log_{10}(C_i^*)$ is a metric used in the 2D-VBS method to evaluate the volatility of organic molecules.

$$C_i^* = \frac{M_i 10^6 \zeta_i p_i^\circ}{RT} \quad (1)$$

where R ($8.206 \times 10^{-5} \text{ m}^3 \text{ atm K}^{-1} \text{ mol}^{-1}$) is the gas constant, T (K) is the temperature, M_i (g mol^{-1}) is the molecular weight of compound i , ζ_i is the activity coefficient of compound i and here is assumed to be 1 (Donahue et al., 2006), p_i° (atm) is the pure liquid saturation vapor pressure at temperature T (298 K).

In this study, different CA methods are applied to calculate the saturation vapor pressures of various oxidation products from isoprene reaction with NO_3 . These include parameterizations that were constrained by chamber measurements as proposed by Donahue et al. (2011), Mohr et al. (2019), and Peräkylä et al. (2019). All of these three parameterization methods have included the effect of the presence of nitrate groups on vapor-pressure estimation. Further we test the GC methods proposed by Nannoolal et al. (2008), Pankow and Asher (2008, SIMPOL.1), and Compernelle et al. (2011, EVAPORATION). All the methods used in this study are summarized in Table 1. The calculations of EVAPORATION and the Nannool method were done via the online molecular and multiphase property prediction facility UManSysProp (http://umansysprop.seaes.manchester.ac.uk/tool/vapour_pressure). For the latter the boiling point parameterization method needs to be predefined, and that from Nannoolal et al. (2004) was adopted as recommended by O'Meara et al. (2014). The information about molecular structures needed for the calculation is inferred from mechanistic information, which is described in detail in Sect. 2.5.

Table 1: Summary of estimation methods of saturation vapor pressure used in this study

Estimation method	Methodology	Input information			Reference
		molecular formula	functional groups	others	
Donahue et al.	CA ^a	√			Donahue et al., 2011
Mohr et al.	ICA ^b	√			Mohr et al., 2019
Peräkylä et al.	ICA ^b	√			Peräkylä et al., 2020
Nannoolal et al.	GC ^c	√	√	√ ^d	Nannoolal et al., 2008
SIMPOL.1	GC	√	√		Pankow and Asher, 2008
EVAPORATION	GC	√	√		Compernelle

This study

EXP^c

289 ^a abbreviation of composition-activity method; ^b abbreviation of improved composition-activity method, which
 290 modified the parameterization based on chamber measurements to better fit HOMs; ^c abbreviation of group-
 291 contribution method; ^d boiling point parameterization method is also required to be defined; ^e abbreviation of
 292 experimental method.

293 In addition, we take advantage of the measurements in this study to calculate the gas-particle equilibrium
 294 partitioning coefficient (K) by comparing experiments with and without seed aerosols. The partitioning
 295 coefficient K can be converted to saturation concentration C* by Eq. (2).

$$296 \quad K_i = \frac{C_{i,p}}{C_{i,g} \times C_{OA}} = \frac{1}{C_i^*} \quad (2)$$

297 where $C_{i,g}$ and $C_{i,p}$ are the gas- and particle-phase concentrations ($\mu\text{g m}^{-3}$) of species i , respectively, and C_{OA} is
 298 the organic aerosol concentration ($\mu\text{g m}^{-3}$). In this study, $C_{i,g}$ is signal of species i from CIMS in the experiment
 299 with seeds, and $C_{i,p}$ is the difference of signals between experiment without and with seeds (under the same
 300 isoprene consumption condition). The C_{OA} in the experiment with seeds is in a range of 1-4 $\mu\text{g m}^{-3}$.

301 2.5 Pathways to the multifunctional oxidation products

302 2.5.1 Basic peroxy and alkoxy radical chemistry

303 As mentioned before, information about molecular structures (at least functional groups) is required to calculate
 304 vapor pressures by using GC methods. Although the high-resolution ToF-CIMS allows for determining
 305 chemical composition of the detected ions, it is unable to provide information about molecular structures, so that
 306 the constitutional or configurational isomers with the same mass cannot be distinguished without additional
 307 information. Fortunately, knowledge of detailed chemical formation mechanisms can help inferring the
 308 molecular structure information. However, the development of a comprehensive, multi-generational kinetic
 309 mechanism for NO₃-initiated oxidation of isoprene is outside the scope of the current paper. Instead, in order to
 310 link the observed mass peaks to representative molecular structures, we developed a framework tracing the
 311 chemical oxidation mechanisms by taking well-known oxidation steps to predict the most likely isomeric forms
 312 of the functionalized products formed in the isoprene oxidation. For this purpose, we rely on the extensive
 313 literature on isoprene, alkylperoxy radical, and alkoxy radical chemistry (Atkinson, 2007; Atkinson and Arey,
 314 2003; Bianchi et al., 2019; Crouse et al., 2013; Ehn et al., 2014; Jenkin et al., 2015; Jenkin et al., 2019; Kwan
 315 et al., 2012; Mentel et al., 2015; Ng et al., 2008; Noveli et al., 2021; Orlando et al., 2003; Orlando and Tyndall,
 316 2012; Rollins et al., 2009; Schwantes et al., 2015; Vereecken and Francisco, 2012; Vereecken and Peeters, 2009,
 317 2010; Vereecken et al., 2021; Wennberg et al., 2018; Ziemann and Atkinson, 2012). This framework is depicted
 318 in the supporting information and will be discussed in more detail in Sect. 2.5.2 and Sect. 2.5.3. They are based
 319 on the following main reactivity trends.

320 Generally, RO₂ radicals can react with other RO₂ and HO₂ radicals. There are three major channels for the
 321 reaction between two RO₂ radicals, leading to alkoxy radicals (RO) (Reaction R1a), as well as termination
 322 products like alcohols, aldehydes or ketones (Reaction R1b) and accretion products (Reaction R1c). These
 323 reactions should take place with the first-generation peroxy radicals, as well as with the higher generation RO₂

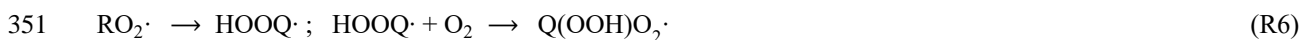
324 radicals formed in the later oxidation steps. Hydroperoxides can be formed from the reaction of RO₂ with HO₂
 325 radicals (Reaction R2a). This reaction can also yield alkoxy radicals (Reaction R2b).



331 In the presence of NO_x, RO₂ radicals can also react with NO and NO₂, leading to the formation of alkoxy
 332 radicals (R3a), organic nitrates (R3b), and peroxy nitrates (R4) (including peroxyacyl nitrates, PANs, if R =
 333 R'C(O)-). The channel that results in RO radicals is the major pathway for the reaction of RO₂ radicals with NO
 334 (Ziemann and Atkinson, 2012). However, reactions of RO₂ radicals with NO (Reaction R3a and R3b) can be
 335 neglected in this study due to the high O₃ concentration, which results in rapid conversion of NO to NO₂. The
 336 peroxy nitrates formed from the reaction of RO₂ with NO₂ will undergo rapid thermal decomposition under our
 337 experimental conditions, with exception of PANs. The reaction of RO₂ with NO₃ radicals mainly forms NO₂ and
 338 alkoxy radicals (Reaction R5), which will continue the radical chains (Reaction R7).



343 In addition to bimolecular reactions, intramolecular rearrangement (H-migration) is a competitive reaction
 344 pathway for RO₂ radicals. RO₂ radicals can undergo H-migration to form a hydroperoxy functionality (-OOH)
 345 and a radical site that can subsequently recombine with an O₂ molecule, leading to the formation of a new, more
 346 oxidized substituted RO₂ (Reaction R6). This process is the so-called "autoxidation" path and has been
 347 confirmed as a significantly important way for SOA formation (Crouse et al., 2013; Ehn et al., 2014; Mentel et
 348 al., 2015; Praske et al., 2018; Rissanen et al., 2014). The rates of RO₂ H-migration are strongly dependent on the
 349 structure of RO₂ radicals, and the most likely routes can be derived based on the structure-activity relationship
 350 proposed by Vereecken and Nozière (2020).



352 The RO radicals formed in in the reaction of RO₂ + RO₂ typically have three accessible pathways,
 353 including isomerization by H-migration (Reaction R7a), fragmentation (Reaction R7b) and less important here,
 354 reaction with O₂ (Reaction R7c). Like H-migration in RO₂, rearrangement by H-shift in RO radicals leads to the
 355 formation of more oxidized RO₂ radicals. Fragmentation leads to smaller carbon chains, and this becomes more
 356 important for alkoxy radicals with a higher number of (oxygen-bearing) substituents (Vereecken and Peeters,
 357 2009, 2010).



361 In addition to the above general reaction pathways, we include a number of other reactions in the
362 framework, such as fragmentation of peroxy radicals, epoxidation of β -OOH alkyl radicals, and unimolecular
363 termination of nitrooxy or hydroperoxyl peroxy radicals. Details can be found in the supporting information.

364 2.5.2 Formation of first-generation products

365 Here “first-generation products” refers to the closed-shell compounds from the first attack of NO_3 at the
366 isoprene double bonds, while “second-generation products” follow an addition of NO_3 to the remaining double
367 bond (or any other oxidation reaction) of a first-generation product. Addition of a NO_3 radical to one of isoprene
368 double bonds and subsequent addition of O_2 to the resulting (delocalized) radical sites leads to the formation of
369 nitrooxy alkylperoxy radicals (INO_2 , $\text{C}_5\text{H}_8\text{NO}_3$). Since isoprene contains two double bonds, NO_3 can attack any
370 of the four positions on the conjugated carbon bonds, resulting in eight possible INO_2 isomers (including six
371 constitutional and two conformational isomers), as shown in Scheme S1. However, both theoretical and
372 experimental studies suggest that the addition occurs preferably at the primary and terminal carbons, wherein C1
373 addition seems to be preferred over C4 addition (Schwantes et al., 2015; Suh et al., 2001; Wennberg et al., 2018).
374 As the GC methods have limited or no ability to distinguish between positional isomers (Kurten et al., 2016), we
375 take exemplarily the products following the C1 addition for the vapor pressure analysis in this study.

376 The initial peroxy radicals ($\text{C}_5\text{H}_8\text{NO}_3$) can undergo rearrangement by H shift from C–H bonds with
377 subsequent O_2 addition, yielding new –OOH functionalized peroxy radicals (Reaction R6). Repeating this
378 process can lead to the formation of a series of peroxy radicals and termination products with stepwise
379 increasing number of oxygen atoms by 2, as shown in the conceptual scheme Scheme S2. This is the RO_2
380 autoxidation channel and the molecular formula of peroxy radicals formed via consecutive O_2 additions can be
381 represented as $\text{C}_5\text{H}_8\text{NO}_{(3+2n)}$ ($n \geq 1$, number of autoxidation steps). The autoxidation chain can be terminated
382 when the H-shift occurs at a carbon with an –OOH or – ONO_2 group attached, leading to carbonyl formation
383 with OH or NO_2 loss (Anglada et al., 2016; Bianchi et al., 2019; Vereecken, 2008; Vereecken et al., 2004). The
384 closed-shell products formed in these termination steps have the general molecular formula $\text{C}_5\text{H}_7\text{NO}_{(5+2n-1)}$ (OH
385 loss channel) or $\text{C}_5\text{H}_8\text{O}_{(3+2n-2)}$ (NO_2 loss channel).

386 The $\text{C}_5\text{H}_8\text{NO}_{(3+2n)}$ peroxy radicals can also react with HO_2 radicals to form –OOH functionalized
387 termination products with the general molecular formula $\text{C}_5\text{H}_9\text{NO}_{(3+2n)}$ (Reaction R2a), or yielding the alkoxy
388 radicals $\text{C}_5\text{H}_8\text{NO}_{(3+2n-1)}$ (Reaction R2b). In addition, the $\text{C}_5\text{H}_8\text{NO}_{(3+2n)}$ peroxy radicals can react with other RO_2
389 radicals (Reaction R1a-R1c). The reaction R1a leads to the formation of alkoxy radicals ($\text{C}_5\text{H}_8\text{NO}_{(3+2n-1)}$) while
390 R1b forms closed-shell products either with a carbonyl group ($\text{C}_5\text{H}_7\text{NO}_{(3+2n-1)}$) or a hydroxyl group
391 ($\text{C}_5\text{H}_9\text{NO}_{(5+2n-1)}$). Alternatively, dimers can be formed following Reaction R1c, which have then two – ONO_2
392 groups and at least 8 oxygen atoms depending on the formula of RO_2 radicals involved, as shown in Table S1.

393 The alkoxy radicals from reactions R1a and R2b can undergo unimolecular rearrangement by H shift with
394 subsequent O_2 addition, similar to the RO_2 radicals, forming new RO_2 radicals with a –OH group (Reaction
395 R7a). As mentioned above, when the H-shift occurs at a carbon with an –OOH or – ONO_2 group attached, the
396 resulting intermediates tend to lose an OH group or NO_2 (Bianchi et al., 2019), yielding the closed-shell
397 carbonyl products with general formulas $\text{C}_5\text{H}_7\text{NO}_{(5+2n-2)}$ or $\text{C}_5\text{H}_8\text{O}_{(3+2n-3)}$ respectively, as shown in the conceptual
398 scheme Scheme S3. The newly-formed RO_2 radicals from alkoxy H-shift channel can follow the peroxy
399 pathways (Reaction R1-R6) like other RO_2 radicals, leading to a diversity of compounds like hydroperoxides

400 (Reaction R2a, $C_5H_9NO_{(3+2n+1)}$), alcohols (Reaction R1b, $C_5H_9NO_{(3+2n)}$), aldehydes (Reaction R1b, $C_5H_7NO_{(3+2n)}$)
401 as well as accretion products (Reaction R1c, $C_{10}H_{16}N_2O_x$), as depicted in Scheme S3. Alternatively, they can
402 also yield alkoxy radicals again following reactions R1a and R2b and continue so on. Furthermore, the alkoxy
403 radicals can break apart into two fragments according to Reaction R7b.

404 In general, the alkoxy reaction pathways diversify the parity of the oxygen number of the products from the
405 reaction of isoprene with NO_3 , and the compounds formed via these reactions generally have one less or one
406 more oxygen atom compared to those formed from straight peroxy reaction pathways. With help of the
407 mechanistic framework described above, we can infer the functionality of first-generation products. This is
408 exemplified in Scheme S5 and S6 for the major first-generation C_5 products. In addition, the reaction pathways
409 and their corresponding structures of the first-generation C_{10} dimers ($C_{10}H_{16}N_2O_x$) are summarized in Scheme
410 S13.

411 **2.5.3 Formation of second-generation products**

412 Nitrate radicals can oxidize the first-generation products once again at the double bond remaining ($k_{NO_3}(298K) \sim$
413 $3\text{-}11 \times 10^{14} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, Wennberg et al., 2018). This leads eventually to “second-generation” products that
414 contain at least two nitrogen atoms. Addition of NO_3 radical to the remaining double bond of the first-generation
415 products results in the formation of dinitrooxy peroxy radicals. We assume that dinitrooxy peroxy radicals can
416 undergo unimolecular and bimolecular reactions (Reaction R1–R6) in analogy to nitrooxy peroxy radicals,
417 which lead to secondary products containing two or more nitrogen atoms, as summarized in the conceptual
418 scheme Scheme S4.

419 The reaction of first-generation nitrooxy peroxy radicals with NO_2 can also yield 2N-compounds (Reaction
420 R4), however these 2N-compounds ought to be under first-generation products by definition. Such species are
421 not discussed in detail here but will be covered to catch the diversity of the functionalities for the vapor pressure
422 estimation. With the help of this secondary reaction framework, we can propose functional groups for the major
423 second-generation products. Scheme S8 – S10 depict the detailed (possible) reaction pathways that lead to the
424 formation of detected C_5 dinitrates, as well as their possible structures. Furthermore, the proposed formation
425 mechanism and their structures for C_5 trinitrates are shown in Scheme S12, while those for the second-
426 generation C_{10} dimers ($C_{10}H_{17}N_3O_x$ and $C_{10}H_{18}N_4O_x$) are depicted in Scheme S13.

427 **2.5.4 Formation of fragmentation products**

428 In addition to the multigenerational C_5 and C_{10} products, fragmentation products can be formed from the
429 reaction of isoprene with NO_3 . As mentioned above, the alkoxy radicals can undergo C–C bond scission,
430 producing a carbonyl compound and an alkyl fragment (Reaction R7b). As shown in Scheme S7, when the
431 secondary nitrooxy alkoxy radicals from the further oxidation of C_5 carbonyl compounds ($C_5H_8O_2$ and $C_5H_8O_3$
432 here) undergo unimolecular decomposition, C_4 carbonyl products ($C_4H_7NO_5$ and $C_4H_7NO_6$, respectively) are
433 formed as well as formyl radicals. Since the bond fission can occur at different positions, the generation of more
434 reactive C_2 and C_3 carbonyl compounds are possible. In addition, the C_4 carbonyl compounds are possibly
435 generated through peroxy radical arrangement by 1,4 H-shift and subsequent acyl radical bond scission reactions
436 (see Scheme S7). The C_4 dinitrates can be formed following similar chemistry, as depicted in Scheme S11.

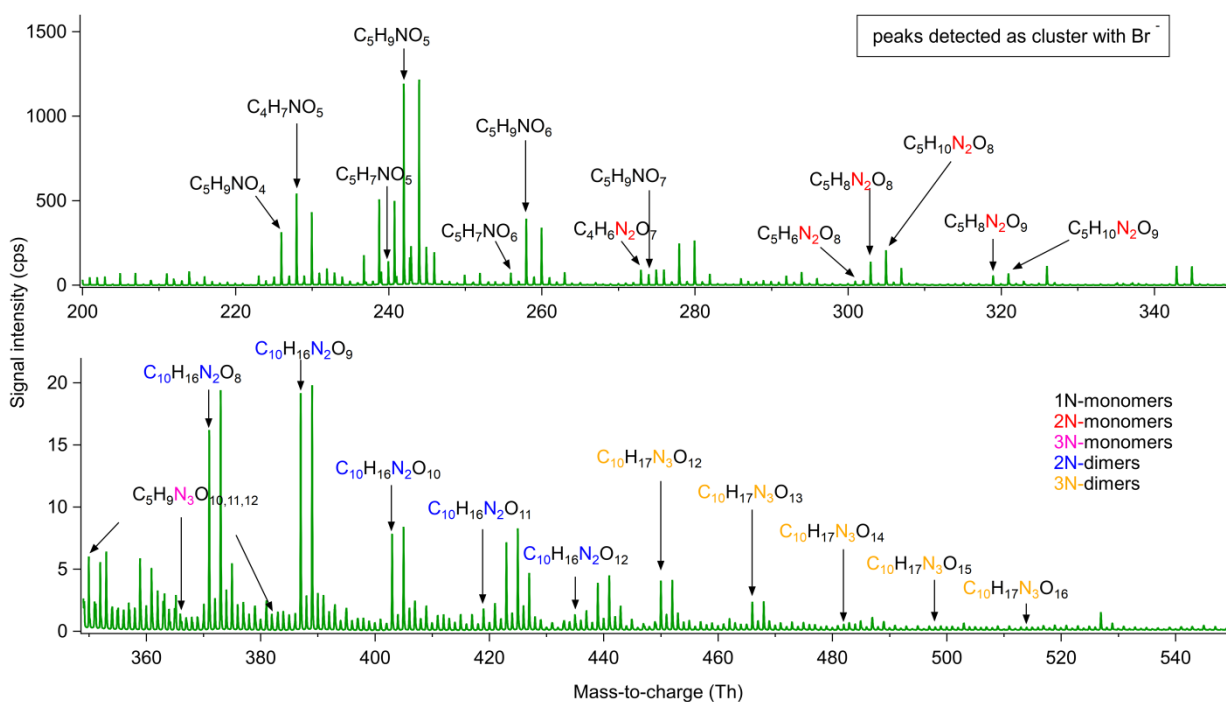
437 2.5.5 Candidate structures for vapor pressure estimation

438 Among all gas-phase products detected by CIMS, we selected 32 major representative organonitrates formed
439 from isoprene oxidation by NO_3 radicals. Their structures are rationalized by the corresponding molecular
440 formulas and proposed formation mechanisms in the reaction framework. Table S2 summarizes all the
441 exemplified structures used for vapor pressure estimation. The functional groups covered in the selected
442 structures include nitrate, hydroxyl, ketone, aldehyde, carboxylic acid, peroxide, hydroperoxide, hydroperoxy
443 acid, peroxyxynitrate, peroxyacyl nitrate and epoxide. The structural information allows calculation of the
444 saturation vapor pressure by GC methods.

445 3. Results and discussion

446 3.1 Chemical composition of oxidation products

447 Figure 2 illustrates the average mass spectra of the whole experiment measured by Br⁻CIMS for isoprene- NO_3
448 reaction. Chemical sum formulas were attributed to most of the detected ions. The gas-phase products were
449 separated into two major groups according to their chemical composition, including monomers comprising C_5
450 compounds and dimers containing C_{10} compounds. There were also products from decomposition reactions with
451 $\text{C}_{<5}$, which were merged into monomers. The monomers and dimers were further classified into five subgroups
452 as follows. Monomers consisting of compounds with one nitrogen atom (hereafter 1N-monomers) and two or
453 three N atoms (2N- or 3N-monomers) mainly accumulate in m/z 220–280 Th, m/z 300–340 Th and 350–390 Th,
454 respectively, while dimers containing compounds with two N atoms (2N-dimers) and three N atoms (3N-dimers)
455 appear in m/z 370–440 Th and 450–520 Th, respectively. As shown in Fig. 2, the signal intensities decrease
456 from 1N-monomers, 2N-monomers, 2N-dimers to 3N-monomers and 3N-dimers. Many of the compounds
457 detected in this work were also observed in previous isoprene- NO_3 systems (Kwan et al., 2012; Ng et al., 2008;
458 Schwantes et al., 2015). In this work, only closed-shell products are considered for analysis.



459

460 **Figure 2: Averaged mass spectra for isoprene-NO₃ experiment on 8 August, 2018. Molecular formulas were**
461 **determined according to the accurate mass data provided by HR-ToF-CIMS.**

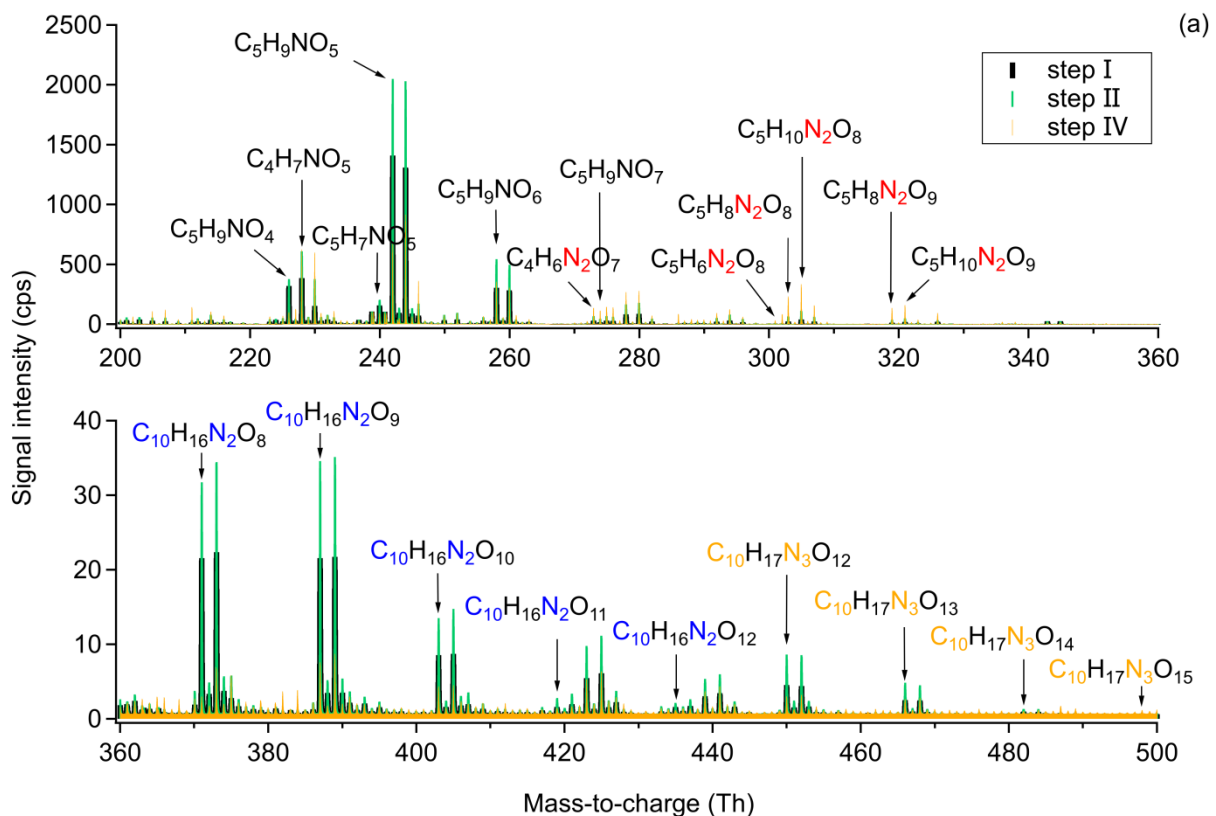
462 The 1N-monomer C₅H₉NO₅ at *m/z* 242 is the dominant product formed from the NO₃-induced isoprene
463 oxidation in our experiment, followed by the 1N-decomposition product C₄H₇NO₅ at *m/z* 228. In addition to
464 C₅H₉NO₅, several analogues with molecular formulas C₅H₇NO₄₋₇ and C₅H₉NO₄ are in relatively high abundance.
465 C₅H_{8,10}N₂O_{8,9} and C₅H₉N₃O₁₀₋₁₂ are the major 2N- and 3N-monomers. Their signal intensities are one to two
466 orders of magnitude lower than those of 1N-monomers. According to the chemical composition, the 1N-
467 monomers are likely to be the first-generation products from NO₃ oxidation of isoprene, while the 2N- and 3N-
468 monomers probably arise from the further oxidation of 1N-monomers by NO₃, which therefore should be
469 second- or later-generation products. As mentioned before, the reaction of nitrooxy alkylperoxy radicals with
470 NO₂ can lead to the formation of peroxy nitrates (for the special case peroxyacyl nitrates, PAN-like) containing
471 two N atoms. The peroxy nitrates will decompose rapidly under experimental conditions, whereas the PAN-like
472 compounds are more stable (with lifetimes ranging from minutes to weeks at 298K and ambient temperature).
473 Such C₅ PAN-like compounds are isomers of aforementioned 2N-monomers, but ought to be first-generation
474 products. In addition to C₅-2N-monomers, we observe some C₄-2N-monomers with relatively high intensity,
475 such as C₄H₆N₂O₇ at *m/z* 273 and C₄H₈N₂O₈ at *m/z* 291. It is proposed that such C₄ dinitrates originate from the
476 further oxidation of C₅ carbonyl compounds followed by unimolecular decomposition (Schwantes et al., 2015;
477 Wennberg et al., 2018), as shown in Scheme S11.

478 2N-Dimers are C₁₀ compounds with 8-12 oxygen atoms (C₁₀H₁₆N₂O₈₋₁₂), and their signal intensities are
479 relatively low compared to that of monomers, approximately three orders of magnitude lower. They might be
480 ROOR products from the self or cross reaction of two nitrooxy peroxy radicals (Berndt et al., 2018). 3N-Dimers
481 are molecules consisting of 12–16 oxygen atoms (C₁₀H₁₇N₃O₁₂₋₁₆). They are probably formed from further
482 oxidation of 2N-dimers or from the cross reaction of a nitrooxy peroxy radical with a dinitrooxy peroxy radicals.

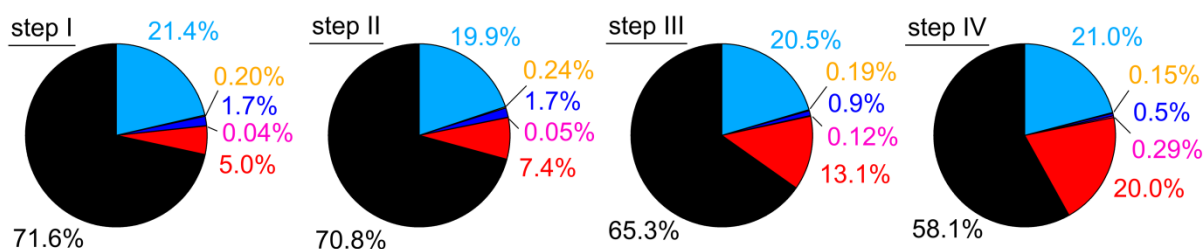
483 **3.2 Multi-generation chemistry**

484 **3.2.1 Molecular composition for each step**

485 As mentioned in Sect. 2.2, there were four injections during the experiment on 8 August (denoted as step I, II,
486 III, IV in Fig. 3), wherein in the first three injections all components, O₃, NO₂, and isoprene, were added, while
487 in the last step only O₃ and NO₂ were injected to promote the further oxidation of early-generation products. The
488 extended oxidation time with reinjection of oxidants provides the opportunity to investigate the multi-generation
489 oxidation chemistry of isoprene-NO₃ system. The mass spectra show only slow changes in the concentrations
490 during the last period of each step, indicating weak chemical evolution. Therefore, we use integrated mass
491 spectra over the last 10 minutes of each step for further analysis. Due to the similarity of the integrated mass
492 spectra for step II and step III, the latter is omitted in Fig. 3.



1N-, 2N-, 3N-monomers, 2N-, 3N-dimers, others



493

494 **Figure 3: Comparison of the chemical composition of each oxidation step. (A) Averaged mass spectra for step I, II,**
 495 **and IV, with the omitted spectrum of step III being very similar to that of step II. (B) Relative contribution of**
 496 **different chemical groups for each oxidation step. Only organic products were counted for analysis. ‘Others’ refers to**
 497 **CHO compounds without containing nitrogen atoms (e.g., C₅H₈O₂ and C₅H₈O₃).**

498 As shown in Fig. 3a, large amounts of 1N-monomers were formed from NO₃ oxidation of isoprene in step I,
 499 wherein C₅H₉NO₅, C₅H₉NO₆, and C₄H₇NO₅ are the most abundant compounds in signal. The 2N-monomers,
 500 which are expected from further oxidation of 1N-monomers, are much less compared to 1N-monomers,
 501 accounting for 5.0% of the total organic signals, with the 3N-monomers even less (0.04%). The low
 502 contributions of second-generation products probably results from the relatively high concentration of isoprene
 503 in step I, reducing the possibility for further oxidation of first-generation products. These results indicate that the
 504 system is dominated by first-generation chemistry at the early stage and therefore the oxidation state of products
 505 is low. In addition to monomers, some 2N- and 3N-dimers are observed. They contribute 1.7% and 0.2%,
 506 respectively, to the total organic signals, as shown in Fig. 3b. The low signal intensity of dimers probably results
 507 from their small yield under our experimental conditions. In this case their contribution to SOA formation might

508 be small. However, a part of the dimers condense onto chamber wall due to their low volatility, so only a
509 smaller portion exists in the gas phase (compare Table S3 and Fig. S5).

510 In step II, the secondary chemistry was accelerated by further addition of O₃ and NO₂, but the primary
511 chemistry was also maintained by isoprene injection. As a result, more 1N-monomers (e.g. C₅H₉NO_{4,5,6}) were
512 formed compared to step I, as well as dimers (e.g., C₁₀H₁₆N₂O_{8,9,10} and C₁₀H₁₇N₃O_{12,13}), as shown in Fig. 3a. The
513 signals of 2N-monomers almost double in this period compared to those in step I, and their relative contribution
514 increase from 5.0% to 7.4%. This is attributed to the further oxidation of first-generation products formed in
515 step I. The relative contributions of different chemical groups exhibited in Fig. 3b clearly show that, although
516 NO₃ produced from the second addition of NO₂ and O₃ still primarily reacted with newly-injected isoprene,
517 reaction of NO₃ with the first-generation oxidation products retaining a double bond was inevitable, leading to
518 more second-generation 2N- or 3N-products compared to step I. The visibly increasing fraction of 2N-
519 monomers indicates that the second-generation chemistry started to play a more important role than that in the
520 early stage. In step III, the chemical process proceeded similarly, and thus is not further discussed here.

521 Due to the favorable conditions for further oxidation, the signals of 1N-monomers (such as C₅H₉NO₄,
522 C₅H₉NO₅, and C₅H₉NO₆), as well as 2N- and 3N-dimers, dropped dramatically in step IV, with their relative
523 contributions decreasing to 58.1%, 0.5%, and 0.15%, respectively. The decrease in signals of dimers is primarily
524 ascribed to lack of isoprene, as there were less peroxy radicals under this condition, and hence less dimers were
525 formed. In addition, their condensation on the wall and dilution also contributed to the decreasing signals.
526 Furthermore, dimers with 2 or 3 nitrogen atoms possess at least one double bond in their molecular structures
527 and can thus be further oxidized under high NO₃ condition to form 4N- or 5N-dimers. However, only few 4N-
528 dimers and no 5N-dimers were detected by CIMS, suggesting that the 4N- and 5N-dimers were either not
529 formed, or if present, with lower absolute concentrations below the detection limit (approximately 5×10⁷ and
530 5×10⁵ molecules cm⁻³ for salicylic acid and acetic acid, for an integration time of 60 s). condensed on the wall
531 due to their low volatilities. In contrast, 2N- and 3N-monomers increase significantly, with their relative
532 contributions ascending to 20.0% and 0.29%, respectively. This indicates that 2N- and 3N-monomers might be
533 second- or later-generation products that are formed from the further oxidation of first-generation products.
534 Additionally, unlike the C₅ monomers, the signal of C₄H₇NO₅ increased in step IV, indicating that there is a new
535 formation pathway for C₄H₇NO₅ under excess NO₃ condition. No double bond can remain in such products, as
536 otherwise they would be oxidized and their signal should decay instead.

537 In summary, above findings confirm that multi-generation chemistry happened during the NO₃-initiated
538 isoprene oxidation, and that the later generation oxidation was promoted by “excess” NO₃ radicals.

539 3.2.2 Carbon oxidation state ($\overline{\text{OS}}_{\text{C}}$)

540 The oxidation state of carbon ($\overline{\text{OS}}_{\text{C}}$) is defined as the charge a carbon atom takes with assumption that it loses
541 completely all electrons in bonds to more electronegative atoms and vice versa (Kroll et al., 2011). This quantity
542 is a metric for the degree of oxidation and will increase with oxidation. Moreover, $\overline{\text{OS}}_{\text{C}}$ together with carbon
543 number can be used to constrain the composition of organic mixtures and provide insights into their evolutions.
544 The carbon oxidation state of a species is determined by the relative abundances and oxidation states of non-
545 carbon atoms in the compound. Since we observed nitrate groups in the products, $\overline{\text{OS}}_{\text{C}}$ is defined by Eq. (3). In
546 this study, the group-averaged $\overline{\text{OS}}_{\text{C}}$ is the signal-weighted mean average carbon oxidation state of compounds

547 with the same carbon number, and the bulk-averaged \overline{OS}_C is the signal-weighted mean average carbon oxidation
548 state of all detected compounds in the system.

$$549 \quad \overline{OS}_C = \frac{2 \times n_O - n_H - 5 \times n_N}{n_C} \quad (3)$$

550 wherein, n_O , n_H , and n_N are the number of the respective atoms in the molecular formula.

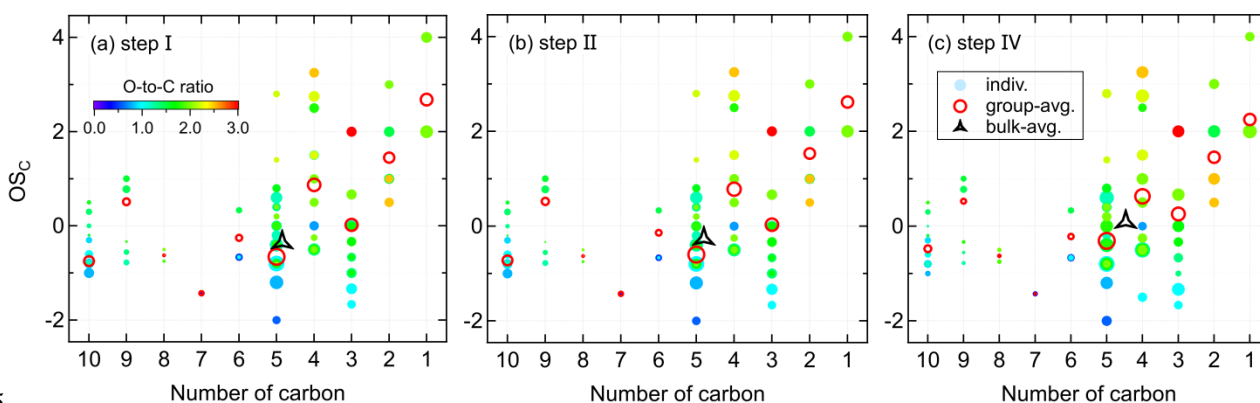
551 Figure 4 shows the distribution of gas-phase products from the isoprene-NO₃ system in the oxidation state
552 versus carbon number (OS_C vs n_C) space. The bulk-averaged \overline{OS}_C is -0.35 in step I, wherein the smaller
553 molecules ($C_{\leq 4}$) have higher oxidation states than the larger molecules. The group-averaged oxidation state of
554 C₅ compounds is relatively low ($\overline{OS}_{C=5} = -0.66$), indicating that both of the oxidation and autoxidation degree of
555 isoprene are quite low during this period. This is consistent with the conclusion made previously from mass
556 spectra results that at the early stage isoprene-NO₃ oxidation was dominated by first-generation chemistry.

557 The system \overline{OS}_C increases to -0.26 in step II, confirming that first-generation products were further
558 oxidized after the second injection. During this step, the \overline{OS}_C of most compound groups increase only weakly,
559 except for that of the C₅ compounds. The group-averaged \overline{OS}_C of C₅ compounds increases to -0.60 in step II,
560 which is the major contributor to the increase of \overline{OS}_C of the whole system. The increase of \overline{OS}_C of C₅
561 compounds is largely attributed to the formation of 2N-monomers expected from further oxidation of existing
562 1N-products formed in step I. This is confirmed by the detectable increase of 2N- and 3N-monomers in the mass
563 spectra and their higher relative contributions to total signals (see Fig. 3). In addition to C₅ compounds, the \overline{OS}_C
564 of C₃ and C₆ products increase significantly in step II.

565 In step IV, the secondary oxidation was largely accelerated by reinjection of O₃ and NO₂, and hence the
566 system oxidation degree increases, with the bulk-averaged \overline{OS}_C growing substantially to 0.09. Similarly, the
567 significant increase of system \overline{OS}_C is mainly attributed to the C₅ compounds, with their group-averaged \overline{OS}_C
568 increasing to -0.31. In addition, the \overline{OS}_C of C₁₀ compounds increased evidently despite their decreasing signals,
569 suggesting C₁₀ dimers were further oxidized as well in step IV. It is worth noting that the average carbon
570 number decreases step by step with increasing \overline{OS}_C . This is the case because fewer C₁₀ products, but more
571 fragments were formed with the reaction proceeding, as shown in Fig.4 by the decreasing peak areas of larger
572 molecules but converse trend for smaller molecules. One conceivable explanation for the decreasing dimers but
573 increasing fragments with the increasing \overline{OS}_C is that, with more highly oxidized RO₂ formed under high NO₃
574 condition, the prevailing fate of RO₂ changes from dimerization to forming alkoxy radicals, which would
575 undergo unimolecular decomposition rapidly, especially when there is a neighboring oxygen-containing
576 functional group (Molteni et al., 2019).

577 In the oxidation system, the increase in \overline{OS}_C is attributed to the formation of bonds between carbon and
578 oxygen as well as other electronegative atoms, and/ or the breaking of bonds between carbon and hydrogen and
579 other electropositive atoms (Kroll et al., 2011). The -ONO₂ group has an oxidation state of -1, which means that
580 addition of a -ONO₂ group to isoprene will increase its OS_C by 0.2. According to our estimates, the values of
581 system \overline{OS}_C increased by 1.25 (step I), 0.09 (step II), and 0.35 (step IV), indicating that the increases in \overline{OS}_C are
582 not only due to addition of -ONO₂ group(s) but also to other oxygen-containing functionalities. In addition to
583 functionalization, it is possible that other reactions such as fragmentation and oligomerization which can
584 increase or reduce the oxidation state were involved during the reaction.

585 As mentioned above, the average carbon oxidation state of a mixture of molecules largely depends on its
 586 chemical composition. Therefore, for different oxidation systems, their \overline{OS}_C may differ due to different
 587 precursors and oxidation conditions. In our study, the \overline{OS}_C of NO_3 -initiated isoprene oxidation system increased
 588 from -0.35 to 0.09 with further oxidation. For OH- and O_3 -initiated systems, the average oxidation state of
 589 laboratory-generated isoprene SOA are reported to range from -1.3 to -0.2, as listed in Table S4. It seems that
 590 the SOA generated from chloride-initiated oxidation of isoprene is more oxidized compared to other isoprene
 591 oxidation systems, for which the \overline{OS}_C can be as high as +1.8 according to limited studies (Wang and Ruiz, 2017).
 592 With regard to ambient measurements, the calculated \overline{OS}_C values of organic aerosol and aerosol fractions fell
 593 into a wider range between -2 to +2, depending on the site position and the corresponding oxidation
 594 environment of that site (Table S4).



595
 596 **Figure 4: Distribution of gas-phase products from isoprene oxidation by NO_3 in the carbon oxidation state (OS_C)**
 597 **versus carbon number (nc) space. Markers are colored by oxygen-to-carbon molar ratio and sized by the logarithm**
 598 **of peak areas. The group-averaged and bulk-averaged \overline{OS}_C are signal-weighted mean average carbon oxidation state**
 599 **of compounds with the same carbon number and of all detected compounds, respectively.**

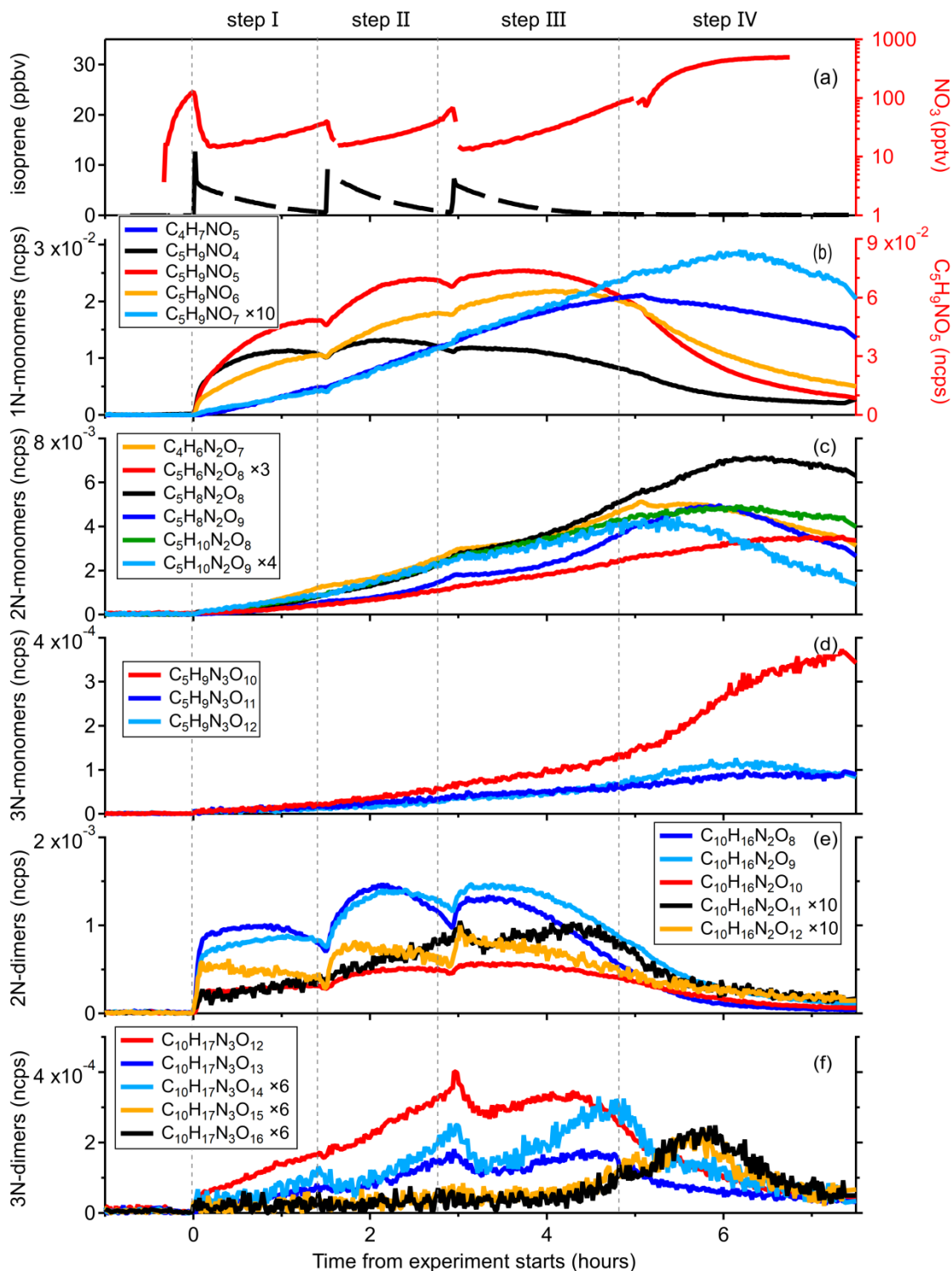
600 In summary, isoprene and its products undergo further oxidation by NO_3 , leading to an increase in degree
 601 of oxidation of products as the reaction proceeds. The increasing bulk \overline{OS}_C is largely governed by the highly
 602 oxidized C_5 compounds. In addition, more fragments but fewer dimers are formed as the \overline{OS}_C increases, which
 603 can be probably explained by the change of RO_2 fate from prevailing dimerization to fragmentation through the
 604 alkoxy radical channel.

605 3.2.3 Characteristics of different-generation products

606 (1) 1N-monomers

607 To illustrate the multi-generation chemistry involved in the isoprene- NO_3 reaction system, Fig. 5 shows the time
 608 evolution of the major gas-phase products. The signal of the most abundant compounds, $\text{C}_5\text{H}_9\text{NO}_5$, increases
 609 rapidly as soon as the reaction was initiated, reaching a maximum when its chemical production rate matches its
 610 loss rate (including chemical destruction, wall loss, dilution, etc.), and decreases slowly thereafter. Its time
 611 behavior in the first three steps is similar. In step IV, however, the injection of O_3 and NO_2 resulted in a strong
 612 decay of $\text{C}_5\text{H}_9\text{NO}_5$, owing to the occurrence of further oxidation by NO_3 . The time behavior suggests that
 613 $\text{C}_5\text{H}_9\text{NO}_5$ signal is dominated by first-generation oxidation products, and the same conclusion can be made for
 614 $\text{C}_5\text{H}_9\text{NO}_4$ and $\text{C}_5\text{H}_9\text{NO}_6$. According to the mechanistic framework developed above, the $\text{C}_5\text{H}_9\text{NO}_4$, $\text{C}_5\text{H}_9\text{NO}_5$,
 615 and $\text{C}_5\text{H}_9\text{NO}_6$ compounds most likely correspond to hydroxyl nitrates, nitrooxy hydroperoxides, and hydroxy

616 hydroperoxy nitrates, respectively, but other constitutional isomers are possible. They were already observed in
 617 previous studies and were proposed to form through reactions of INO_2 radicals with RO_2 , HO_2 , and
 618 unimolecular rearrangement, as shown in Scheme S5 (Ng et al., 2008; Kwan et al., 2012; Schwantes et al., 2015;
 619 Wennberg et al., 2018).



620
 621 **Figure 5: Time evolution of selected gas-phase compounds measured during the isoprene - NO_3 experiment on 08**
 622 **August, 2018. (a) Time series of O_3 , NO_2 , NO_3 and isoprene. (b)–(f) Time evolution of major 1N-monomers ($\text{C}_5\text{H}_9\text{NO}_4$ -**
 623 **7 and $\text{C}_4\text{H}_7\text{NO}_5$), 2N-monomers ($\text{C}_4\text{H}_6\text{N}_2\text{O}_7$, $\text{C}_5\text{H}_6\text{N}_2\text{O}_8$, and $\text{C}_5\text{H}_{8,10}\text{N}_2\text{O}_{8,9}$), 3N-monomers ($\text{C}_5\text{H}_9\text{N}_3\text{O}_{10-12}$), 2N-dimers**
 624 **($\text{C}_{10}\text{H}_{16}\text{N}_2\text{O}_{8-12}$), and 3N-dimers ($\text{C}_{10}\text{H}_{17}\text{N}_3\text{O}_{12-16}$).**

625 As shown in Fig. 5b, the temporal evolution of $C_5H_9NO_7$ (m/z 274) is different to $C_5H_9NO_{4-6}$ compounds,
626 suggesting that it has a completely different formation pathway. Specifically, the formation rate of $C_5H_9NO_7$ is
627 initially much slower than that of $C_5H_9NO_{4-6}$ but accelerates to become comparable to them later as the
628 experiment proceeds, i.e. when a multitude of first-generation products are accumulated. This implies that
629 $C_5H_9NO_7$ is produced from the further oxidation of first-generation products, and its signal is dominated by
630 second-generation products. Based on its molecular composition, $C_5H_9NO_7$ could be the dihydroperoxy nitrate
631 as shown in Scheme S5, but its formation through the reaction of HO_2 with nitrooxy hydroperoxy radical from
632 INO_2 autoxidation suggests it should be first-generation products, not in accordance with the time behavior we
633 actually observe. Consequently, we can conclude that it is not the major formation pathway that contributed to
634 $C_5H_9NO_7$ observed in this study. As shown in Scheme S7, the first-generation C_5 hydroxy carbonyl ($C_5H_8O_2$,
635 m/z 179) can be further oxidized by NO_3 and the resulting alkyl radical would rapidly recombine with O_2 ,
636 producing a new peroxy radical, which then reacts with HO_2 radicals to form $C_5H_9NO_7$ (hydroxy hydroperoxy
637 carbonyl nitrate). Similarly, the C_5 hydroperoxy carbonyl ($C_5H_8O_3$, m/z 195) can also lead to the formation of
638 such $C_5H_9NO_7$ (isomer of that formed through $C_5H_8O_2$ channel) through further oxidation (see Scheme S7).
639 According to above two mechanisms, $C_5H_9NO_7$ formed following such reaction pathways should be second-
640 generation products, better consistent with its time behavior.

641 Considering its similar time behavior to $C_5H_9NO_7$, the observed $C_4H_7NO_5$ (m/z 228) signal is likewise
642 thought to be dominated by second-generation products. Schwantes et al. (2015) proposed such a C_4 product
643 based on OH-initiated chemistry, but as the OH concentration in our system was below the detection limit
644 during the experiment (see Fig. S2), this formation pathway cannot apply in our situation. Instead, we suggest
645 that $C_4H_7NO_5$ is formed through the unimolecular decomposition of the C_5 alkoxy or acyl radicals, which result
646 from further oxidation of the C_5 hydroxy carbonyl ($C_5H_8O_2$, m/z 179), as shown in Scheme S7. It should be
647 pointed out here that there may be reaction pathways forming $C_4H_7NO_5$ as first-generation products that are not
648 considered here, whereas it is no doubt that the second-generation chemistry played a dominant role in $C_4H_7NO_5$
649 formation according to its time evolution measured by CIMS.

650 Although $C_4H_7NO_5$ and $C_5H_9NO_7$ show similar time behaviors in the first three steps, it seems that they
651 followed fairly different reaction pathways when the concentration of NO_3 in the chamber increased
652 dramatically in step IV. As shown in Fig. 5b, the signal of $C_4H_7NO_5$ drops immediately after the injection of O_3
653 and NO_2 , while that of $C_5H_9NO_7$ continues to increase, although its formation rate becomes slightly lower with
654 increasing NO_3 concentration. The decay of $C_4H_7NO_5$ signal can be explained by more chemical destruction or
655 less production under high NO_3 condition, wherein the latter seems more sensible in terms of its structure (no
656 double bond remaining). As shown in Scheme S7, the second-generation $C_4H_7NO_5$ and $C_5H_9NO_7$ compounds
657 share the same precursor in the $C_5H_8O_2$ channel. Consequently, the production of $C_5H_9NO_7$ through this
658 pathway would be interrupted immediately after the injection of O_3 and NO_2 like $C_4H_7NO_5$. In reality, its signal
659 might decay even faster due to the larger reaction rate of RO_2 H-shift (leading to the formation of $C_4H_7NO_5$)
660 than that of RO_2 reacting with HO_2 (leading to the formation of $C_5H_9NO_7$). As presented by Vereecken and
661 Nozière (2020), the rate coefficient of aldehydic H-shift is $\geq 0.5 \text{ s}^{-1}$ (298 K), while the pseudo first order rate
662 coefficient of RO_2 reacting with HO_2 is $\sim 10^{-3} \text{ s}^{-1}$ (k (298 K) = $5 \times 10^{-12} \text{ cm}^3 \text{ molecules}^{-1}\text{s}^{-1}$ (Atkinson, 2007), and
663 $[HO_2] \sim 4 \times 10^8 \text{ molecules cm}^{-3}$), about two orders of magnitude smaller. This result implies that the increasing
664 $C_5H_9NO_7$ observed is contributed to by other formation pathways. As mentioned before, $C_5H_9NO_7$ can also be

665 produced by $C_5H_8O_3$ oxidation. We find that the signal of $C_4H_7NO_6$ (m/z 244), which results from $C_5H_8O_3$
666 oxidation as well, remains increasing after the injection of O_3 and NO_2 . This tentatively confirms that the
667 production of $C_5H_9NO_7$ in step IV is mainly from $C_5H_8O_3$ oxidation channel. More experimental or theoretical
668 studies are needed to provide insights into these differences.

669 (2) 2N- and 3N-monomers

670 As shown in Fig. 5c, 2N-monomers formed much slower than 1N-monomers in the early stage, but their
671 formation rates were accelerated in step II and step III, probably due to the accumulation of first-generation
672 products. According to our mechanistic framework, 2N-monomers are second-generation products resulting
673 from the further oxidation of 1N-monomers by NO_3 , which is consistent with their time behaviors detected by
674 CIMS.

675 Like $C_4H_7NO_5$ and $C_5H_9NO_7$, different 2N-monomers have similar behavior in the first three steps, but they
676 are obviously different in step IV when the concentration of NO_3 increased drastically in the chamber. For
677 instance, the signals of $C_5H_8N_2O_8$, $C_5H_8N_2O_9$ and $C_5H_{10}N_2O_8$ continue to increase after the injection of O_3 and
678 NO_2 , while that of $C_5H_{10}N_2O_9$ drops immediately. This is related to their detailed formation mechanisms which
679 are outside the scope of this study. Furthermore, $C_5H_8N_2O_9$ and $C_5H_{10}N_2O_9$ decay a little bit faster than
680 $C_5H_8N_2O_8$ and $C_5H_{10}N_2O_8$, which might be related to their volatility and will be further discussed in next section.

681 Different from other 2N-monomers, the signals of $C_5H_6N_2O_8$ (m/z 301) increases continuously under high
682 NO_3 condition, although its net formation rate is almost zero at the end of step IV. The characteristics of
683 $C_5H_6N_2O_8$ under high NO_3 condition reflects its different formation pathways from other dinitrates, and without
684 having a comprehensive knowledge of its chemical mechanism, we are unable to tell what exactly leads to the
685 differences. In the Master Chemical Mechanism (MCM v3.3.1), $C_5H_6N_2O_8$ is proposed to be a PAN-like
686 compound stemming from the C_5 nitrooxy carbonyl ($C_5H_7NO_4$)
687 (<http://mcm.leeds.ac.uk/MCM/browse.htm?species=NC4CHO>). Such $C_5H_6N_2O_8$ compound would react with
688 NO_3 radicals due to the remaining double bond, and hence this cannot be the predominant formation pathway of
689 the $C_5H_6N_2O_8$ observed in this study. Based on the formation mechanism of dinitrooxyepoxides ($C_5H_8N_2O_7$)
690 proposed by Kwan et al. (2012), we suggest that $C_5H_6N_2O_8$ can also be a dinitrooxyepoxide resulting from
691 cyclization of specific hydroperoxy alkyl radicals, as shown in Scheme S10. Alternatively, the C_5 hydroxy
692 nitrate ($C_5H_9NO_4$) can be oxidized by NO_3 and then react with NO_3 radicals again, forming $C_5H_6N_2O_8$ with two
693 aldehyde groups ultimately (see Scheme S10). According to the proposed mechanisms above, $C_5H_6N_2O_8$ formed
694 through the first two pathways are second-generation products, while those from the third channel are third-
695 generation products, in accordance with its time behavior measured by CIMS.

696 In addition to C_5 -2N-monomers, we observe some C_4 dinitrates such as $C_4H_6N_2O_7$ (m/z 273) and $C_4H_8N_2O_8$
697 (m/z 291), and the signal intensity of $C_4H_6N_2O_7$ is comparable to the major C_5 -2N-monomers. C_4 dinitrates have
698 rarely been mentioned in previous isoprene- NO_3 studies. As shown in Fig. 5c, $C_4H_6N_2O_7$ has similar time
699 behavior to C_5 -2N-monomers, and hence is thought to be second-generation products. Wennberg et al. (2018)
700 proposed that such a C_4 dinitrate was generated from OH-initiated further oxidation of $C_5H_7NO_4$. However, this
701 is not applicable here due to a lack of OH radicals in our system. Instead, we propose that the $C_4H_6N_2O_7$
702 observed in this study is dinitrooxy carbonyl compound resulting from NO_3 oxidation of $C_5H_7NO_4$ with
703 subsequent unimolecular decomposition (see Scheme S11 for details).

704 As shown in Fig. 5d, 3N-monomers are generated more slowly than 1N-monomers, but their signals grow
705 gradually as the experiment proceeds, with a significant increase especially for $C_5H_9N_3O_{10}$ in the last step.
706 Furthermore, we can see from Fig. 5c and Fig. 5d that the signals of C_5 trinitrates in step IV appear
707 anticorrelated to that of $C_5H_{10}N_2O_8$ and $C_5H_{10}N_2O_9$. The gas-phase 3N-monomers have rarely been reported in
708 previous literature. Ng et al. (2008) observed $C_5H_9N_3O_{10}$ compound in the particle-phase and assumed that it
709 was produced from NO_3 oxidation of the C_5 hydroxy nitrate ($C_5H_9NO_4$). Similarly, $C_5H_9N_3O_{11}$ and $C_5H_9N_3O_{12}$
710 can be formed through NO_3 reacting with dinitrooxy peroxy radicals, which result from corresponding first-
711 generation nitrooxy compounds (C_5 hydroperoxy nitrate, $C_5H_9NO_5$ or C_5 hydroxy hydroperoxy nitrate, $C_5H_9NO_6$)
712 oxidation by NO_3 radicals, as shown in Scheme S12. 3N-Monomers formed following such pathways are
713 second-generation products by definition. Regarding the rising signals of 3N-monomers in step IV, one
714 explanation is that although the reaction of dinitrooxy peroxy radicals with NO_3 is not an oxidation process,
715 their formation can be significantly facilitated by increasing NO_3 concentration. It is also possible that 3N-
716 monomers are formed through H-abstraction of 2N-monomers. NO_3 radicals can abstract the hydrogen of
717 dihydroxy dinitrate ($C_5H_{10}N_2O_8$) or hydroxyl hydroperoxy dinitrate ($C_5H_{10}N_2O_9$) from the carbon with an $-OH$,
718 $-OOH$ or $-ONO_2$ group attached, leading to alkyl radicals that can subsequently recombine with O_2 and then
719 react with NO_2 or NO_3 , yielding trinitrates or peroxytrinitrates containing three nitrogen atoms. 3N-Monomers
720 stemming from such reactions ought to be third-generation products. However, we should point out that 3N-
721 monomers formed following H-abstraction pathway are less likely because abstracting hydrogen from the
722 hydroxyl, hydroperoxy or nitrooxy carbon would lead to fragmentation at most cases (Bianchi et al., 2019).

723 In addition, it is interesting to note that the signal of $C_5H_9N_3O_{10}$ increases continuously throughout step IV,
724 whereas that of $C_5H_9N_3O_{11}$ and $C_5H_9N_3O_{12}$ drop after a short period of growth. Meanwhile, the production of
725 $C_5H_9N_3O_{10}$ is facilitated by the increasing NO_3 concentration compared to that of $C_5H_9N_3O_{12}$ and $C_5H_9N_3O_{11}$.
726 Currently, we cannot explain what exactly causes these differences, but we suspect that there may be different
727 chemical pathways forming different 3N-monomers that are not covered here and may also be related to their
728 different physical properties, such as vapor pressures.

729 (3) 2N- and 3N-dimers

730 As shown in Fig. 5e, 2N-dimers (except for $C_{10}H_{16}N_2O_{11}$) display very similar time behavior to 1N-monomer,
731 which form rapidly after each injection, indicating that the signals of 2N-dimers are dominated by first-
732 generation products like most 1N-monomers. It is noted that the time behavior of $C_{10}H_{16}N_2O_{11}$ (m/z 419) is
733 completely different from that of other 2N-dimers. As illustrated in Fig. 5e, the production rate of $C_{10}H_{16}N_2O_{11}$
734 is initially much slower compared to other dimers. Besides, its signal increases monotonically in the first two
735 oxidation stages, whereas that of the others always increase first, approaching the maximum as its chemical
736 production competes against the losses, and decrease gradually thereafter. The special time behavior of
737 $C_{10}H_{16}N_2O_{11}$ suggests that it has a different formation pathway from other 2N-dimers, and its signal is most
738 likely dominated by secondary products. In addition, we find that the signal of $C_{10}H_{16}N_2O_{12}$ always starts to
739 decay earlier than that of $C_{10}H_{16}N_2O_8$ and $C_{10}H_{16}N_2O_9$. If we assume that their production rates have the same
740 order of magnitude (confirming by their formation rates after each injection), then it can be concluded that
741 $C_{10}H_{16}N_2O_{12}$ had additional chemical destruction, or its volatility is much lower than $C_{10}H_{16}N_2O_8$ and
742 $C_{10}H_{16}N_2O_9$ and hence has more rapid lost on the wall. It seems the second hypothesis is more likely when

743 comparing its signal with and without dilution and wall-loss corrections (see Fig. S5). More detailed discussion
744 about volatilities of different isoprene organonitrates will be provided in the next section.

745 It is proposed that dimers (ROOR') are likely formed through the self- or cross-reaction of two peroxy
746 radicals (Berndt et al. 2018). Consequently, the generation number of dimers depends only on how the involved
747 peroxy radicals are formed. Table S1 summarizes the possible permutation scheme of 2N-dimers from RO₂ +
748 RO' reactions, and their structural information can be found in Scheme S13. For example, self-reaction of two
749 C₅ nitrooxy peroxy radicals (C₅H₈NO₅) leads to the formation of C₁₀H₁₆N₂O₈ compound, while recombination
750 of two C₅ nitrooxy hydroxyl peroxy radicals (C₅H₈NO₆) or a C₅ nitrooxy peroxy radical (C₅H₈NO₅) with a C₅
751 nitrooxy hydroperoxy peroxy radical (C₅H₈NO₇) results in C₁₀H₁₆N₂O₁₀ compound. According to their time
752 behavior, 2N-dimers (except for C₁₀H₁₆N₂O₁₁) are thought to be first-generation products, and from this fact we
753 can infer that the peroxy radicals contributing to dimer formation are dominated by first-generation
754 intermediates. With regard to C₁₀H₁₆N₂O₁₁, we conclude that it is most likely a secondary product considering
755 its typical second-generation behavior. In other words, at least one of the two C₅ nitrooxy peroxy radicals
756 involved in formation of C₁₀H₁₆N₂O₁₁ must be a secondary intermediate. As listed in Table S1, C₁₀H₁₆N₂O₁₁ can
757 be formed through C₅H₈NO₆ + C₅H₈NO₇ or C₅H₈NO₆ + C₅H₈NO₇ reactions, wherein C₅H₈NO₇ and C₅H₈NO₈
758 would be secondary peroxy radicals if they are formed through NO₃ further oxidation of the C₅ hydroxy
759 carbonyl compounds (C₅H₈O₂ or C₅H₈O₃), as shown in Scheme S7. In addition, it is possible that C₁₀H₁₆N₂O₁₁ is
760 formed from a C₅ hydroxy peroxy radical C₅H₉O₃ reacting with a C₅ dinitrooxy hydroxy carbonyl peroxy radical
761 C₅H₇N₂O₁₀ (from C₅H₇NO₅ oxidation by NO₃), as we observe high abundant C₅H₁₀O₃ during the experiment,
762 although C₅H₁₀O₃ is assumed to be the major product of the OH-initiated chemistry.

763 Apart from 2N-dimers, we observe detectable signals at *m/z* 450, 466, 482, 498 and 514, which are
764 identified as 3N-dimers with molecular formulas C₁₀H₁₇N₃O₁₂₋₁₆. C₁₀H₁₇N₃O₁₂ and C₁₀H₁₇N₃O₁₃ were detected
765 in the particle-phase in previous study, suggesting that they have low volatility and can contribute to SOA
766 formation (Ng et al., 2008). As shown in Fig. 5f, 3N-dimers form much slower than 2N-dimers, but their
767 productions are accelerated as the experiment proceeds. This is similar to the characteristics of second-
768 generation 2N- and 3N-monomers to some degree, suggesting that the signals of 3N-dimers we observed are
769 most likely dominated by secondary or even later-generation compounds.

770 It is worth noting that C₁₀H₁₇N₃O₁₂₋₁₄ and C₁₀H₁₇N₃O_{15,16} have two completely different types of time
771 behavior. The signals of C₁₀H₁₇N₃O₁₂, C₁₀H₁₇N₃O₁₃ and C₁₀H₁₇N₃O₁₄ more or less increase in the first three
772 oxidation steps and start to decline in the late of step III with increasing NO₃ concentration. As depicted in
773 Scheme S13, 3N-dimers can result from further oxidation of 2N-dimers or the cross-reaction of a first-
774 generation nitrooxy peroxy radical with a secondary dinitrooxy peroxy radical. Accordingly, such 3N-dimers are
775 thought to be second-generation products, and they would further react with NO₃ due to the remaining double
776 bond in their molecular structure, leading to severe chemical destruction of these compounds under high NO₃
777 condition. This is consistent with the time behavior of C₁₀H₁₇N₃O₁₂, C₁₀H₁₇N₃O₁₃ and C₁₀H₁₇N₃O₁₄. In contrast,
778 C₁₀H₁₇N₃O₁₅ and C₁₀H₁₇N₃O₁₆ are formed even more slowly, and their production in the first four hours is close
779 to zero. However, their signals start to climb in the late of step III, during which that of C₁₀H₁₇N₃O₁₂,
780 C₁₀H₁₇N₃O₁₃ and C₁₀H₁₇N₃O₁₄ decline. This suggests that C₁₀H₁₇N₃O₁₅ and C₁₀H₁₇N₃O₁₆ formed under high NO₃
781 condition probably result from further reactions of C₁₀H₁₇N₃O₁₂₋₁₄. However, this assumption is highly uncertain

782 and more experimental and theoretical studies are needed to substantiate it. In terms of their time behavior,
783 $C_{10}H_{17}N_3O_{15}$ and $C_{10}H_{17}N_3O_{16}$ are thought to be third- or even later-generation products.

784 3.3 Volatility distribution of isoprene nitrates

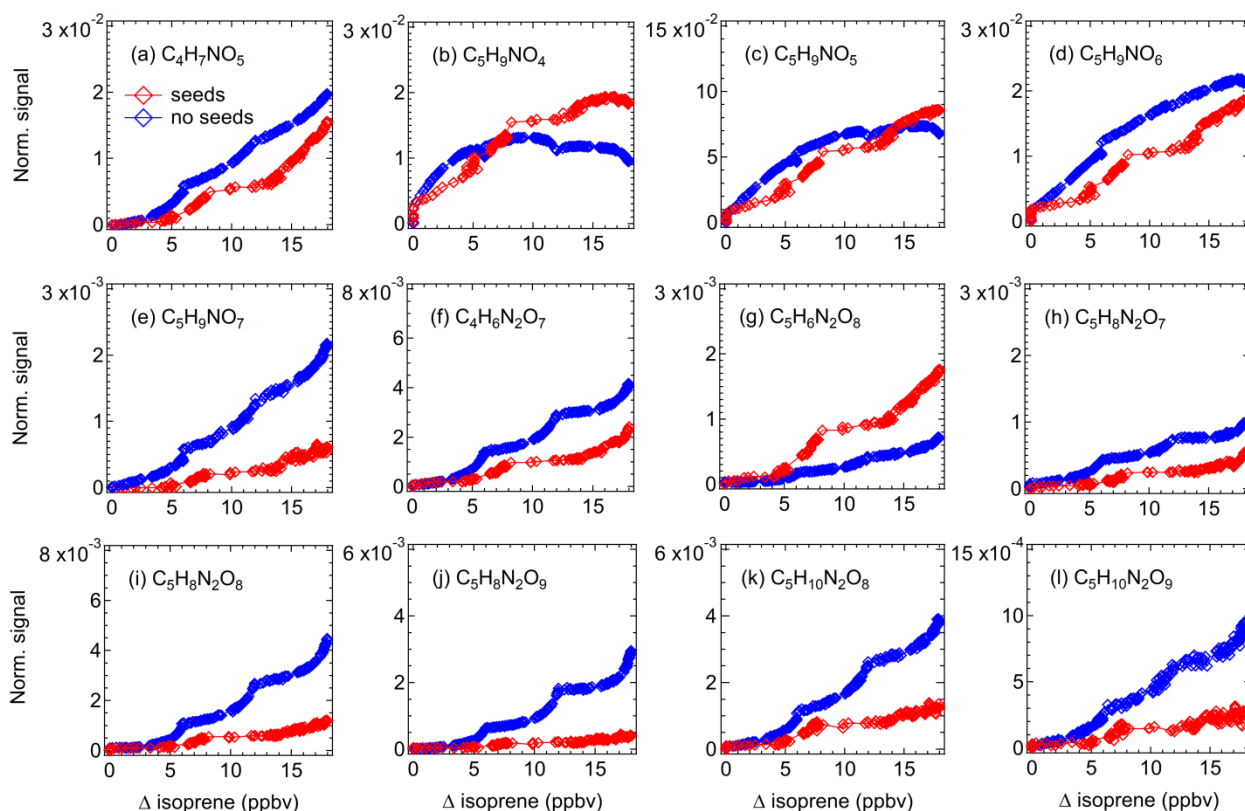
785 3.3.1 C^* estimated by experimental methods

786 Detailed information about the volatility of organic molecules is essential to evaluate their potential to form
787 SOA. In order to investigate the potential contribution of various isoprene oxidation products to SOA formation,
788 we use our (limited) experimental data to estimate the vapor pressure of different isoprene organonitrates on the
789 basis of their condensation behavior. Figure 6 shows how the signals of gas-phase products change in
790 experiments with and without seed aerosols (ammonium sulfate). Please note that while the two experiments
791 were conducted under similar conditions, the procedures could not be kept fully identical as aerosol seeding
792 required specific measures and the oxidation chemistry might be slightly altered (e.g., due to initiation of
793 heterogeneous reactions).

794 As shown in Fig. 6, the signals of most of the selected compounds decline when there are seed aerosols in
795 the chamber, indicating that part of the condensable vapors is partitioned to the particle-phase due to the
796 introduction of condensation sinks. The decrease in signal differs for different products, mostly depending on
797 their vapor pressures. As expected, the lower volatility of a compound the higher the fraction that condenses.
798 For instance, the signal of $C_5H_9NO_7$ decreases by more than 70% in experiment with seed aerosols, compared to
799 less than 40% on average for other less-oxidized 1N-monomers. In some cases (e.g., $C_5H_9NO_4$ and $C_5H_9NO_5$)
800 however, the product signals in experiment with seed aerosols are higher than that without seeds after the
801 consumed isoprene exceeding a certain level. In addition, the signal of $C_5H_6N_2O_8$ in the experiment with seeds
802 is always higher compared to that without seeds. One explanation for this phenomenon is the effect of
803 heterogeneous reactions. It is likely that some condensed compound (denoted as A) can react on the particle
804 surface to form new products with the molecular composition of compound B, or alternatively forming a
805 precursor of B. When they evaporate back to the gas phase, it can result in an increase in signal of compound B.
806 That's why a higher signal was observed for such compounds in experiment with seeds than that without seeds,
807 as observed for $C_5H_6N_2O_8$ in this case.

808 Based on the observed condensation behavior of different products, we can derive their vapor pressures
809 from the gas-particle equilibrium partitioning coefficients by Eq. (2). As depicted in Fig. 7, the saturation
810 concentrations of different organonitrates show a decreasing tendency from 1N-, 2N-monomer and 3N-
811 monomers to 2N- and 3N-dimers, suggesting that dimers have a higher propensity of condensation and
812 contribute to SOA formation. This is partly related to their molecular weight, as larger molecules generally have
813 lower vapor pressures. However, it cannot explain all the features of the volatility distribution. For example,
814 $C_5H_9NO_6$ (corresponding to No.8 in Fig.7) has higher mass than $C_5H_9NO_5$ (corresponding to No.7 in Fig.7) but
815 is predicted to have higher vapor pressure. In general, chemical composition and functionalities have significant
816 effects on vapor pressure. For instance, the 2D-VBS composition-activity relationship suggests that each carbon
817 and oxygen decrease C^* by 0.475 and 1.75 decades, respectively (Donahue et al., 2011). Different functional
818 groups also have very different effect on volatility. For example, each hydroxyl group ($-OH$) or hydroperoxy
819 group ($-OOH$) typically reduces the volatility by 2.4 to 2.5 decades, while the less polar carbonyl group ($=O$)
820 reduces the volatility by 1 decade (Pankow and Asher 2008, Donahue et al., 2011). The nitrooxy group ($-ONO_2$)

821 has a similar reductive effect on vapor pressure, which typically reduces C^* by 2.5 orders of magnitude (Pankow
 822 and Asher, 2008). Here, the irregularly high vapor pressure of $C_5H_9NO_6$ is most likely attributed to the
 823 functional groups it contains. As listed in Table S2, $C_5H_9NO_6$ is proposed to be nitrooxy hydroxy hydroperoxyl
 824 compound, which consists of two highly polar functional groups $-OH$ and $-OOH$, contributing to formation of
 825 intramolecular H-bonding that can significantly increase the vapor pressure (Bilde et al., 2015; Kurten et al.,
 826 2016), while $C_5H_9NO_5$ only contains a $-OOH$ group and hence cannot form intramolecular H-bonding. This
 827 explanation is also valid for $C_5H_9N_3O_{10}$ and $C_5H_9N_3O_{12}$. In summary, these findings underline that the
 828 constitutional and configurational information of a molecule is critical for vapor-pressure estimation.



829
 830 **Figure 6: Time evolution of selected major gas-phase products during experiments with (red) and without (blue) seed**
 831 **aerosols (ammonium sulfate). Signals have been corrected for dilution.**

832 3.3.2 C^* estimated by different parametrization methods

833 For comparison, we also adopt different parameterization methods to estimate the saturation vapor pressures of
 834 isoprene oxidation products based on their molecular composition and the proposed structures, with the results
 835 depicted in Fig. 7. In general, the saturation concentrations calculated by different parameterization methods
 836 show a similar volatility distribution to that calculated by experimental method, with C^* of 1N-, 2N- and 3N-
 837 monomers, 2N- and 3N-dimers decreasing in turn. However, different parameterization methods lead to the
 838 predicted vapor concentrations with a variability of several orders of magnitude for the same compound, and the
 839 discrepancies become larger and larger with more complicated molecules. In addition, C^* of structural isomers
 840 calculated by the same method could span several decades.

841 As shown in Fig. 7, the Donahue et al. parameterization mostly provides lower C^* compared to the three
 842 GC methods, with a maximum discrepancy up to 12 orders of magnitude for dimers. With regard to smaller and
 843 less oxidized 1N-monomers, predicted C^* values from different methods are in relatively good agreement with

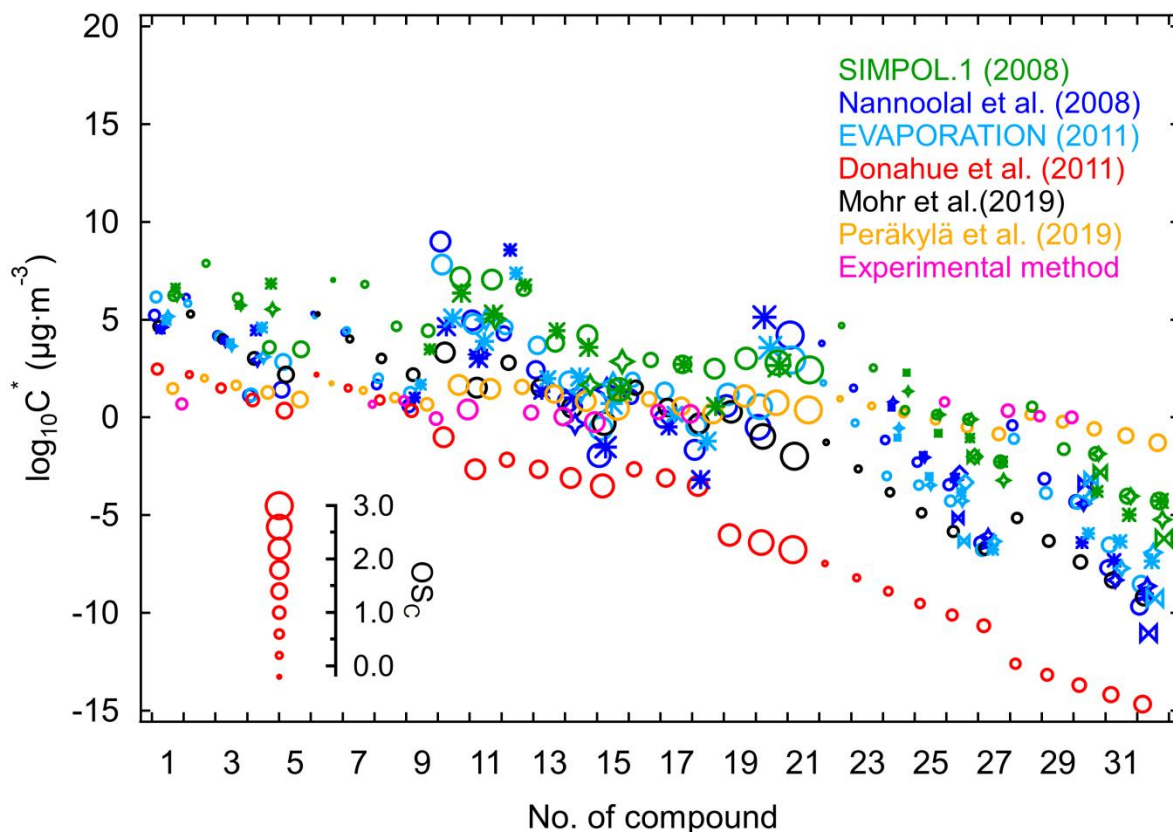
844 each other, whereas the disagreement increases to 11 orders of magnitude for 2N- and 3N-monomers. This is
845 mainly the case because the organic molecules were regarded as a mixture of =O and –OH functional groups in
846 the Donahue et al. parameterization, and their relative abundance was assumed to be 1:1 (Donahue et al., 2011).
847 In consequence, the –OOH functional group in peroxides is treated as two –OH groups when adapting the
848 method proposed by Donahue et al. (2011). However, it is demonstrated that the extra oxygen in peroxy
849 moieties has little contribution to reduce vapor pressure (Pankow and Asher et al., 2008), hence treating –OOH
850 equivalent to two –OH functional groups would underestimate the vapor pressures of hydroperoxyl compounds.
851 Furthermore, organic compounds consisting of multiple polar functional groups (such as hydroperoxy, peroxy
852 acid, and peroxide functional groups) tend to form intramolecular H-bonding, which would increase the vapor
853 pressure (Bilde et al., 2015; Kurten et al., 2016). All these issues contribute to an underestimation of the vapor
854 pressures of multifunctional products when using the Donahue et al. parameterization. Mohr et al. (2019)
855 improved the parameterization for vapor-pressure estimation by taking the presence of –OOH functional groups
856 in HOM explicitly into consideration and revising the parameters to reduce the effect of –OOH on depressing C^* .
857 Consequently, the Mohr et al. parameterization effectively reduces the discrepancy between its estimates and
858 those predicted by the GC methods, with the differences within 6 orders of magnitude. Nevertheless, there is a
859 slight tendency to underestimate the vapor pressures of 3N-monomers and dimers. The Peräkylä et al.
860 parameterization method, which was derived from measurements of the condensation behavior of HOM
861 produced from α -pinene ozonolysis, predicts similar C^* to Donahue et al. method for 1N-monomers, but higher
862 C^* for 2N- and 3N-monomers like the Mohr et al. method. As for dimers, especially for the 3N-dimers
863 containing more multifunctional groups, the Peräkylä et al. method even predicts higher C^* than the GC
864 methods in most cases.

865 Three GC methods predict similar saturation vapor pressures for different isoprene nitrates in this work,
866 with the differences within 5 orders of magnitudes. Generally, the SIMPOL.1 method always provides higher C^*
867 compared to another two methods, and the disagreement between methods becomes larger for molecules
868 containing multifunctional groups. For instance, the vapor-pressure discrepancy between SIMPOL.1 and
869 another two GC methods are both 2 orders of magnitude for $C_5H_9NO_{4,5}$ and $C_{10}H_{17}N_3O_{12-14}$, but it increased up
870 to 4 and 5 orders of magnitude, respectively, for $C_5H_9NO_{6,7}$ and $C_{10}H_{17}N_3O_{15,16}$.

871 It is worth noting that the Nannoolal et al. method is able to distinguish between positional isomers (e.g.,
872 the estimated C^* for two $C_5H_{10}N_2O_9$ isomers are 0.858 and 0.333 $\mu\text{g m}^{-3}$, respectively), whereas such capacity of
873 EVAPORATION method is limited (e.g., it is able to distinguish between the position isomers of $C_5H_{10}N_2O_9$,
874 but it predicts identical C^* for $C_{10}H_{16}N_2O_{11}$ isomers). In this respect, the SIMPOL.1 method cannot distinguish
875 between positional isomers at all. Moreover, SIMPOL.1 method predicts smaller differences between functional
876 group isomers for 1N-monomers and 3N-dimers compared to the Nannoolal et al. method and the
877 EVAPORATION, but there is no such regular pattern for 2N-monomers and 2N-dimers.

878 By comparing the results calculated by experimental method with those by different parameterization
879 methods, we can see that the GC methods predict high saturation concentrations for 1N-monomers than the
880 experimental method, while the Donahue et al. and Peräkylä et al. method provide similar C^* values. With
881 regard to 2N-monomers, the GC methods predict higher vapor pressures compared to the experimental method,
882 but the discrepancy decreases with decreasing saturation concentration. The disagreement of C^* for 2N-
883 monomers estimated by experimental method and the Mohr et al. or Peräkylä et al. method are within 2 orders

884 of magnitude. In terms of low-volatility dimers, however, the vapor pressures calculated by the experimental
885 method were 1–3 orders of magnitude larger than that predicted by the parameterization methods except for the
886 Peräkylä et al. method. The Peräkylä et al. method provides the most similar predictions to the experimental
887 method for isoprene oxidation products in the full volatility range, with the disagreement within 1 order of
888 magnitude.

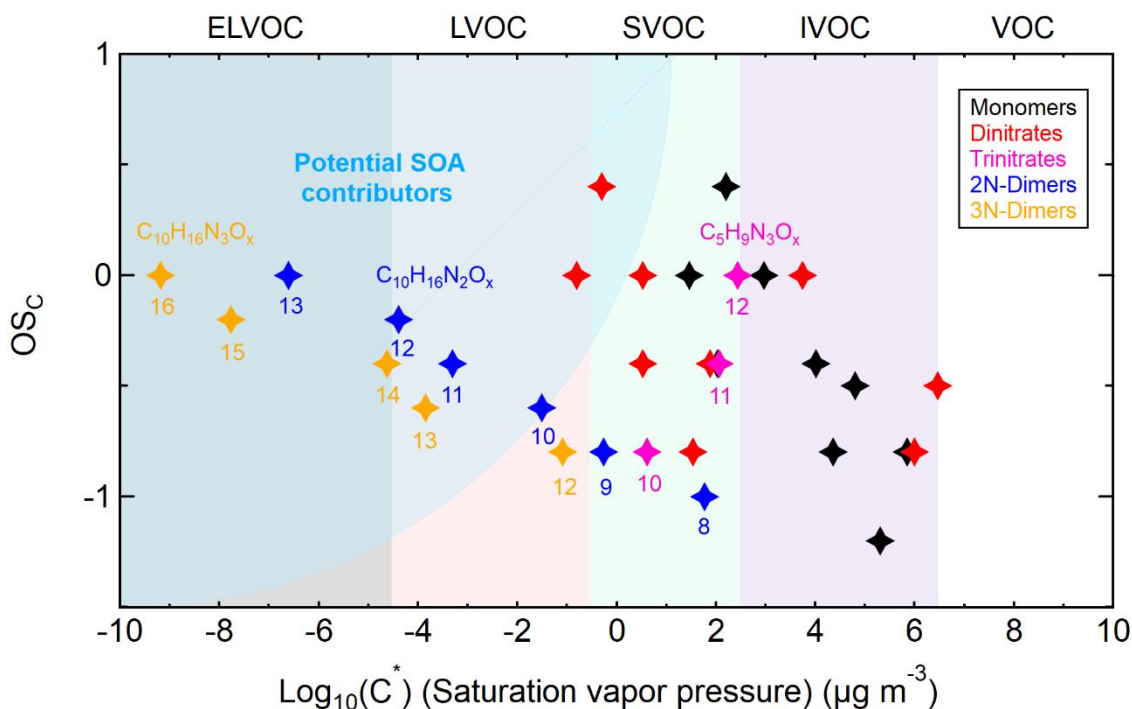


889
890 **Figure 7:** Saturation concentrations (in $\mu\text{g m}^{-3}$, at 298.15 K) of isoprene organonitrates estimated by using
891 experimental and parameterization methods. The numbers correspond with the compound numbers of given in Table
892 S2 (No. 1–9, 10–18, 19–21, 22–27, and 28–32 corresponding to 1N-monomers, 2N-monomers, 3N-monomer, 2N-
893 dimers and 3N-dimers, respectively). Marker shapes indicate different isomers, with their size scaled by carbon
894 oxidation state (*OSC*).

895 In general, the vapor pressures estimated experimentally in this study are very close to that calculated by
896 Peräkylä et al. method for which the estimation parameters were also derived experimentally. The discrepancy
897 between the experimental and the GC methods spans several orders of magnitude depending on different
898 compounds, with the GC methods predicting higher C^* for less-functionalized 1N-monomers, approximate C^*
899 for 2N-monomers, but lower C^* for highly functionalized dimers. It is difficult to tell which method is more
900 reliable without any measured saturation vapor pressure data on such multifunctional organic nitrates. However,
901 considering the fact that the existing GC methods tend to underestimate saturation vapor pressures of the highly
902 functionalized organic molecules due to their limited capability to deal with intramolecular interactions (e.g. the
903 intramolecular hydrogen bonding formed among polar functional groups), and the well consistent results of two
904 experimentally derived methods, we suggest that the experimental method might be a good choice to determine
905 the volatility of highly oxidized compounds accurately.

906 **3.3.3 Volatility distribution of isoprene nitrates and expected SOA yields.**

907 Although the vapor pressures calculated by different methods show a variability of several orders of magnitude,
 908 the predicted volatility distributions of different organic groups are consistent. To eliminate the discrepancy
 909 caused by methods and get an average trend of the volatility distribution of various isoprene nitrates, we use the
 910 median value of C^* calculated by different methods as the estimator of the vapor pressure for each nitrate
 911 compound.
 912



913
 914 **Figure 8: Volatility distribution of different organonitrates formed from NO_3 -initiated isoprene oxidation. The**
 915 **volatility classes are indicated along the top with corresponding colors in the plot. The position of potential SOA**
 916 **contributors is determined depending on the exact functionalities of molecules adapted from Bianchi et al. (2019).**

917 The average carbon oxidation state is plotted against $\text{Log}_{10}(C^*)$ in Fig. 8 to describe the volatility
 918 distribution of organic nitrates formed from isoprene oxidation by NO_3 . Generally, the volatility of measured
 919 gas-phase products spans a wide range from IVOC to ELVOC, wherein all of the 1N-monomers fall in the
 920 IVOC or SVOC range, suggesting that 1N-monomers have low potential to form SOA by simple condensation
 921 as long as the organic aerosol load is less than $200 \mu\text{g m}^{-3}$. The addition of a second or third $-\text{NO}_3$ functional
 922 group decreases C^* of most 2N- and 3N-monomers by 2-3 decades compared with 1N-monomers, and most of
 923 them belong to SVOC. They will start to condense in significant fractions if the organic aerosol load is in a
 924 range of $1\text{-}10 \mu\text{g m}^{-3}$, which means 2N- and 3N-monomers with $\text{OS}_c > -0.8$ may contribute to SOA formation
 925 under atmospheric conditions. With regard to dimers, all 3N-dimers and 2N-dimers (except for $\text{C}_{10}\text{H}_{16}\text{N}_{2}\text{O}_{8,9}$)
 926 are in LVOC or even ELVOC range, indicating isoprene dimers had high propensity to form SOA even at
 927 organic aerosol loads $\ll 1 \mu\text{g/m}^3$. However, we would like to emphasize here that the signals of 2N- and 3N-
 928 dimers only account for less than 2% on average of the total assigned signals, as shown in Fig. S6. This suggests
 929 that the SOA yield of isoprene from NO_3 oxidation by condensation should be low under atmospheric
 930 conditions.

931 The fate of RO₂ determines the product distribution directly and hence could substantially affect SOA
932 yields and aerosol physicochemical properties (Boyd et al., 2015; Fry et al., 2018; Ng et al., 2008; Schwantes et
933 al., 2015; Ziemann and Atkinson, 2012). Consequently, it would be helpful to provide SOA yields together with
934 the fate of RO₂. In our experiment, reactions with HO₂ and NO₃ the dominant loss channels for the initially
935 formed RO₂ from isoprene oxidation by NO₃, contributing for ~ 53% and ~ 30% of overall RO₂ loss; RO₂ + RO₂
936 reactions contributed a minor fraction (~ 13%) followed by unimolecular reactions with a contribution of ~ 5%,
937 according to modelling results (Brownwood et al., 2021). More details about the modelling and the results can
938 be found elsewhere (Brownwood et al., 2021; Vereecken et al., 2021).

939 In polluted urban regions, the fate of RO₂ is typically dominated by RO₂ + NO₃, while in the more pristine
940 environment, the RO₂ + HO₂ reaction will dominate RO₂ fate (Bianchi et al., 2019; Boyd et al., 2015; Brown
941 and Stutz, 2012). RO₂ + HO₂ was more important in the chamber than that in ambient and enhanced RO₂ + HO₂
942 would potentially lead to less dimer formation by RO₂ + RO₂ reactions and hence reducing SOA yields.
943 However, a recent work from Brownwood et al. (2021) based on the same campaign as this study pointed out
944 that the bulk aerosol composition and SOA yields were largely independent of RO₂ fate. Similarly, Boyd et al.
945 (2015) found for β-pinene-NO₃ system that RO₂ fate (“RO₂ + NO₃ dominant” vs “RO₂ + HO₂ dominant”) had
946 only few effects on SOA formation. Therefore, the SOA yield estimated in this study is expected to be
947 comparable to that in the atmosphere.

948 Assuming that the dimers in the LVOC or ELVOC range will condense onto particles, we estimated a SOA
949 mass yield for condensation of isoprene organic nitrates of about 5 % ± 2 %. This value is based on an averaged
950 bulk organonitrate sensitivity of 0.019 norm. count s⁻¹ ppbv⁻¹ and has been corrected for wall loss and dilution
951 (see Fig. S7, with uncorrected SOA mass yield of about 2 %). The estimated SOA mass yield is within the range
952 of those reported in the literature, but at the lower end (4.3% to 23.8% depending on RO₂ fate, Ng et al., 2008;
953 0.7% for first generation oxidation and 14% after oxidation of both double bonds, Rollins et al., 2009; 27% on
954 average for ambient measurements, Fry et al., 2018). The SOA yield will probably become somewhat higher if
955 taking the contribution of SVOCs (including C₁₀H₁₆N₂O₈, C₁₀H₁₆N₂O₉ and some other monomers, as shown in
956 Fig. 8) into consideration. Our finding is commensurable with the SOA yield for isoprene organic nitrates of 2-6%
957 derived from HR-AMS measurements in the same campaign (Brownwood et al., 2021).

958 In addition, Br⁻ adduct ionization CIMS is selective for HO₂ and less oxidized organic compounds
959 (Albrecht et al., 2019; Rissanen et al., 2019), so it is reasonable to assume that there were more highly oxidized
960 products that were not detected by Br⁻ CIMS. This assumption is confirmed by measurements with a NO₃⁻ CIMS
961 performed in another isoprene-NO₃ experiment in SAPHIR (Zhao et al., 2021). Zhao et al. (2021) observed a
962 higher fraction of dimers and more highly oxidized monomers and dimers, as well as trimers (C₁₅ compounds).
963 As a consequence, the SOA yields derived from NO₃⁻ CIMS measurements is slightly higher.

964 From these points of view our yield is more a lower limit. However, even if we assume an error of a factor
965 of 2, the SOA yield of isoprene organic nitrates by condensation is more likely in a range of about 10% or less
966 than in the higher range of 20-30% published in the literature. Of course, by our method we cannot cover any
967 liquid phase processes that would lead to additional SOA beyond the condensation of the target compounds.

968 **4. Conclusions and implication**

969 In this work, a gas-phase experiment conducted in the SAPHIR chamber under near atmospheric conditions in
970 the dark was analyzed to primarily investigate the multi-generation chemistry of isoprene-NO₃ system. The
971 characteristics of a diversity of isoprene nitrates were measured by the CIMS using Br⁻ as the reagent ion.
972 Isoprene 1N-, 2N-, and 3N-monomers and 2N- and 3N-dimers have different time behaviors, indicating the
973 occurrence of multi-generation oxidation during this process. Based on their specific time behaviors as well as
974 the general knowledge of isoprene and radical chemistry, the possible formation mechanisms of these
975 compounds are proposed.

976 In order to evaluate the potential contribution of various isoprene nitrates to SOA formation, different
977 composition-activity and group-contribution methods were used to estimate their saturation vapor pressures. We
978 also calculated the vapor pressures of isoprene oxidation products based on the gas-particle equilibrium
979 coefficients derived from condensation measurements. The vapor pressures estimated by different methods
980 spans several orders of magnitude, and the discrepancies increase as the compounds become highly
981 functionalized. It shows that existing composition-activity methods (especially the Donahue et al. method)
982 seriously underestimate the saturation vapor pressure of multifunctional low-volatility molecules compared to
983 the experimental methods. The group-contribution methods seem to have a better performance than the CA
984 methods on this aspect, but they still have a tendency to underestimate the vapor pressures of multifunctional
985 molecules. We suggest that experimental methods is a good choice to estimate the volatility of highly oxidized
986 compounds accurately.

987 According to our results, 1N-monomers and most 2N and 3N-nitrates fall in the IVOC or SVOC range.
988 Therefore, they have, with a few exceptions, low potential to form SOA at atmospheric organic aerosol loads. In
989 contrast, 2N- and 3N-dimers are estimated to have low or extremely low volatility, indicating that they are
990 significant contributors to SOA formation, although dimers constitute less than 2% of the total explained signals.
991 In this study, no new particle formation events were observed. Assuming that the dimers in the LVOC or
992 ELVOC range will condense onto particles completely, we estimate a SOA mass yield of about 5 % ± 2 %,
993 which is a lower limit if one takes a possible contribution of the minor dimer products as well as SVOC species
994 into consideration. Both the volatility distribution and calculated SOA yields indicate that isoprene dimers
995 formed from NO₃ oxidation are the major contributors to SOA formation.

996 **Data availability**

997 All data given in figures can be displayed in tables or in digital form. This includes the data given in the
998 Supplement. Please send all requests for data to t.mentel@fz-juelich.de and r.wu@fz-juelich.de.

999 **Author contributions**

1000 HF, JNC, JLF, SSB, AW, and AKS designed the study. Instrument deployment and data analysis were carried
1001 out by RW, ET, SK, SRA, LH, AN, HF, RT, TH, PTMC, JS, FB, BB, JAT. RW, LV, ET, DZ, JAT, MH, TFM
1002 interpreted the compiled data set. RW, TFM, LV wrote the manuscript. All co-authors discussed the results and
1003 commented on the manuscript.

1004 **Competing interests**

1005 The authors declare that they have no conflict of interest.

1006 **Acknowledgements**

1007 This work has received funding from the European Research Council (ERC) and European Commission (EC)
1008 under the European Union's Horizon 2020 research and innovation program (SARLEP grant agreement No.
1009 681529, Eurochamp 2020 grant agreement No. 730997 and FORCeS, grant No.821205). R.Wu gratefully
1010 acknowledges the fellowship from Helmholtz-OCPC (Office of China Postdoc Council) Postdoc Program for
1011 research support. M. Hallquist, E. Tsiligiannis and Th. F. Mentel acknowledge support by the Swedish Research
1012 Council (grant numbers 2014-05332 and 2018-04430) and FORMAS (grant numbers 942-2015-1537 and xxx-
1013 2020-yyyy).

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