# **OBSERVATIONS OF HIGHLY OXIDISED** 1 **MOLECULES AND PARTICLE NUCLEATION** 2 **IN THE ATMOSPHERE OF BEIJING** 3 4 James Brean<sup>1</sup>, Roy M. Harrison<sup>1\*†</sup>, Zongbo Shi<sup>1</sup> 5 David C.S. Beddows<sup>1</sup>, W. Joe F. Acton<sup>2</sup> 6 C. Nicholas Hewitt<sup>2</sup>, Freya A. Squires<sup>3</sup> and James Lee<sup>3</sup> 7 8 9 <sup>1</sup>Division of Environmental Health and Risk Management, 10 School of Geography, Earth and Environmental Sciences 11 **University of Birmingham** 12 **Edgbaston, Birmingham B15 2TT** 13 **United Kingdom** 14 15 <sup>2</sup>Lancaster Environment Centre 16 Lancaster University, Lancaster LA1 4YQ 17 **United Kingdom** 18 19 <sup>3</sup>National Centre for Atmospheric Science 20 **Wolfson Atmospheric Chemistry Laboratory** 21 **University of York, York YO10 5DD** 22 **United Kingdom** 23 24

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# 25 ABSTRACT

Particle nucleation is one of the main sources of atmospheric particulate matter by number, with new 26 particles having great relevance for human health and climate. Highly oxidised multifunctional 27 28 organic molecules (HOMs) have been recently identified as key constituents in the growth, and, sometimes, in initial formation of new particles. While there have been many studies of HOMs in 29 atmospheric chambers, flow tubes and clean environments, analyses of data from polluted 30 environments are scarce. Here, measurements of HOMs and particle size distributions down to small 31 molecular clusters are presented alongside volatile organic compounds (VOC) and trace gas data from 32 a campaign in June 2017, in Beijing. Many gas phase HOMs have been characterised and their 33 temporal trends and behaviours analysed in the context of new particle formation. The HOMs 34 identified have a comparable degree of oxidation to those seen in other, cleaner, environments, likely 35 due to an interplay between the higher temperatures facilitating rapid hydrogen abstractions and the 36 higher concentrations of NO<sub>x</sub> and other RO<sub>2</sub><sup>-</sup> terminators ending the autoxidation sequence more 37 rapidly. Our data indicate that alkylbenzenes, monoterpenes, and isoprene are important precursor 38 VOCs for HOMs in Beijing. Many of the C<sub>5</sub> and C<sub>10</sub> compounds derived from isoprene and 39 monoterpenes have a slightly greater degree of average oxidation state of carbon compared to those 40 from other precursors. Most HOMs except for large dimers have daytime peak concentrations, 41 42 indicating the importance of OH<sup>-</sup> chemistry in the formation of HOMs, as O<sub>3</sub> tends to be lower on days with higher HOM concentrations ; similarly, VOC concentrations are lower on the days with 43 higher HOM concentrations. The daytime peaks of HOMs coincide with the growth of freshly formed 44 new particles, and their initial formation coincides with the peak in sulfuric acid vapours, suggesting 45 that the nucleation process is sulfuric acid-dependent, with HOMs contributing to subsequent particle 46 47 growth.

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## 50 1. INTRODUCTION

Atmospheric particle nucleation, or the formation of solid or liquid particles from vapour phase 51 precursors is one of the dominant sources of global aerosol by number, with primary emissions 52 53 typically dominating the mass loadings (Tomasi et al., 2016). New particle formation (NPF) or the secondary formation of fresh particles is a two-step process comprising of initial homogeneous 54 nucleation of thermodynamically stable clusters and their subsequent growth. The rate of growth 55 needs be fast enough to out-compete the loss of these particles by coagulation and condensation 56 processes in order for the new particles to grow, and hence NPF is a function of the competition 57 between source and sink (Gong et al., 2010). New particle formation has been shown to occur 58 across a wide range of environments (Kulmala et al., 2005). The high particle load in urban 59 environments was thought to suppress new particle formation until measurements in the early 2000s 60 (McMurry et al., 2000; Shi et al., 2001; Alam et al., 2003), with frequent occurrences observed even 61 in the most polluted urban centres. NPF events in Beijing occur on about 40% of days annually, 62 with the highest rates in the spring (Wu et al., 2007, 2008; Wang et al., 2016). Chu et al. (2019) 63 review many studies of NPF which have taken place in China and highlight the need for long-term 64 observations and mechanistic studies. 65

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NPF can lead to production of cloud condensation nuclei (CCN) (Wiedensohler et al., 2009; Yu and 67 Luo, 2009; Yue et al., 2011; Kerminen et al., 2012) which influences the radiative atmospheric 68 forcing (Penner et al., 2011). A high particle count, such as that caused by nucleation events, has 69 been shown to precede haze events in environments such as Beijing (Guo et al., 2014). These events 70 are detrimental to health and quality of life. The sub-100 nm fraction of particles to which new 71 72 particle formation contributes to is often referred to as the ultrafine fraction. Ultrafine particles (UFPs) pose risks to human health due to their high number concentration. UFPs exhibit gas-like 73 behaviour and enter all parts of the lung before penetrating into the bloodstream (Miller et al., 74 2017). They can initiate inflammation via oxidative stress responses, progressing conditions such as 75

atherosclerosis and initiating cardiovascular responses such as hypertension through to myocardial
infarction (Delfino et al., 2005; Brook et al., 2010).

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79 Highly oxidised multifunctional molecules (HOMs), organic molecules with O:C ratios >0.6, are the result of atmospheric autoxidation and have recently been subject to much investigation, in part 80 81 because the extremely low volatilities arising from their high O:C ratios favour their condensation into the particulate phase. HOMs are most well characterised as the product of oxidation of the 82 biogenic monoterpenoid compound  $\alpha$ -pinene (Riccobono et al., 2014; Tröstl et al., 2016; Bianchi et 83 al., 2017). Although globally, biogenic volatile organic compound (BVOC) concentrations far 84 exceed anthropogenic volatile organic compound (AVOC) concentrations, in the urban environment 85 the anthropogenic fraction is far more significant. Formation of HOMs from aromatic compounds 86 has been demonstrated in laboratory studies and these have been hypothesised to be large drivers of 87 NPF in urban environments (Wang et al., 2017; Molteni et al., 2018; Qi et al., 2018). The formation 88 of HOMs through autoxidation processes begins with the reaction of VOCs with OH, O<sub>3</sub> or NO<sub>3</sub>; 89 formation of a peroxy radical (RO<sub>2</sub>) is followed by rapid O<sub>2</sub> additions and intra-molecular hydrogen 90 abstractions (Jokinen et al., 2014; Rissanen et al., 2014; Kurtén et al., 2015). Furthermore, 91 generation of oligomers from stabilised Criegee intermediates arising from short chain alkenes has 92 93 been hypothesised as a contributor of Extremely Low Volatility Organic Compounds (ELVOCs) and Low Volatility Organic Compounds (LVOCs) (Zhao et al., 2015). The low volatilities of these 94 molecules arise from their numerous oxygen-containing functionalities, and this allows them to 95 make a significant contribution to early stage particle growth where other species cannot due to the 96 Kelvin effect (Tröstl et al., 2016), although the contribution of HOMs to the initial molecular 97 98 clusters is still debated (Kurtén et al., 2016; Elm et al., 2017; Myllys et al., 2017).

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Recent technological advances have facilitated insights into the very first steps of nucleation which
 were previously unseen, with mass spectrometric techniques such as the Atmospheric Pressure

Interface Time of Flight Mass Spectrometer (APi-ToF) and its chemical ionisation counterpart (CI-102 APi-ToF) allowing for high mass and time resolution measurements of low volatility compounds 103 and molecular clusters. Diethylene glycol based particle counters, such as the Particle Size 104 105 Magnifier (PSM) allow for measurements of particle size distributions down to the smallest molecular clusters nearing 1 nm. Recent chamber studies have elucidated the contribution of 106 individual species to particle nucleation, ammonia and amines greatly enhancing the rate of sulfuric 107 acid nucleation (Kirkby et al., 2011; Almeida et al., 2013). In these studies, HOMs have been 108 identified, formed through autoxidation mechanisms (Schobesberger et al., 2013; Riccobono et al., 109 2014; Ehn et al., 2014). These are key to early particle growth (Tröstl et al., 2016) and can nucleate 110 111 even in the absence of sulfuric acid in chambers (Kirkby et al., 2016) and in the free troposphere (Rose et al., 2018). In this paper, we report the results of HOM and particle size measurements 112 during a summer campaign in Beijing, China, 113

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115 **2. DATA AND METHODS** 

#### 116 **2.1.** Sampling Site

Sampling was performed as part of the Air Pollution and Human Health in a Developing Megacity 117 (APHH-Beijing) campaign, a large international collaborative project examining emissions, 118 processes and health effects of air pollution. For a comprehensive overview of the programme, see 119 Shi et al. (2019). All sampling was conducted across a one month period at the Institute for 120 Atmospheric Physics (IAP), Chinese Academy of Sciences, Beijing (39°58.53'N, 116°22.69'E). 121 The sampling was conducted from a shipping container, with sampling inlets 1-2 metres above 122 ground level, the nearest road being 30 metres away. Meteorological parameters (wind speed, wind 123 direction, relative humidity (RH) and temperature) were measured at the IAP meteorological tower, 124 20 metres away from the sampling site, 30 metres from the nearest road at a height of 120 metres. 125 Data was continuously taken from the CI-APi-ToF during a two week period, but due to data losses 126

only five days of data is presented here. Particle size distribution measurements were taken during a
33 day period from 24/05/2017 – 26/06/2017.

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# 2.2 Chemical Ionisation Atmospheric Pressure Interface Time of Flight Mass Spectrometry

The Aerodyne Nitrate Chemical Ionisation Atmospheric Pressure Interface Time of Flight Mass 132 Spectrometer (CI-APi-ToF) was used to make measurements of neutral oxidised organic 133 compounds, sulfuric acid and their molecular clusters at high time resolution with high resolving 134 power. The ionization system charges molecules by adduct formation, such as in the case of organic 135 compounds with two or more hydrogen bond donor groups (Hyttinen et al., 2015), or proton 136 transfer in the case of strong acids like sulfuric acid. Hydroxyl or hydroperoxyl functionalities are 137 both common hydrogen bond donating groups, with hydroperoxyl being the more efficient 138 hydrogen bond donor (Møller et al., 2017). This instrument has been explained in great detail 139 elsewhere (Junninen et al., 2010; Jokinen et al., 2012), but briefly the front end consists of a 140 141 chemical ionisation system where a 10 LPM sample flow is drawn in through the 1 metre length 1" OD stainless steel tubing opening. A secondary flow was run parallel and concentric to this sample 142 flow, rendering the reaction chamber effectively wall-less. A 3 SCCM flow of a carrier gas (N<sub>2</sub>) is 143 passed over a reservoir of liquid HNO<sub>3</sub>, entraining vapour which is subsequently ionised to NO<sub>3</sub><sup>-</sup> via 144 an X-ray source. This flow is then guided into the sample flow. . The nitrate ions will then charge 145 molecules either by clustering or proton transfer. The mixed flows travelling at 10 LPM enter the 146 critical orifice at the front end of the instrument at 0.8 LPM and are guided through a series of 147 differentially pumped chambers before reaching the ToF analyser. Two of these chambers contain 148 149 quadrupoles which can be used to select greater sensitivity for certain mass ranges, and the voltages across each individual chamber can be tuned to maximise sensitivity and resolution for ions of 150 interest. Mass spectra are taken at a frequency of 20 kHz but are recorded at a rate of 1 Hz. All data 151 analysis was carried out in the Tofware package in Igor Pro 6 (Tofwerk AG, Switzerland). A seven 152

153	point mass calibration was performed for every minute of data, and all data was normalised to
154	signal at 62, 80 and 125 $m/Q$ to account for fluctuations in ion signal, these masses representing
155	NO <sub>3</sub> <sup>-</sup> , H <sub>2</sub> ONO <sub>3</sub> <sup>-</sup> and HNO <sub>3</sub> NO <sub>3</sub> <sup>-</sup> respectively. The resultant normalised counts have been multiplied
156	by $10^5$ so magnitudes are similar to the original count rates. Typical values for calibration
157	coefficients range from $10^9$ - $10^{10}$ molecules cm <sup>-3</sup> from these normalised data (Kürten et al., 2012),
158	producing peak sulfuric acid concentrations in the range of 10 <sup>6</sup> molecules cm <sup>-3</sup> . From the very
159	limited periods with simultaneous data for SO <sub>2</sub> , OH radical and condensation sink, it was possible
160	to calculate $H_2SO_4$ concentrations of $10^3$ to $10^5$ molec cm <sup>-3</sup> , in which range the calibration constant
161	was $7.0 \pm 1.6 \text{ x } 10^8 \text{ cm}^{-3}$ which fits well with that expected for this concentration range (Kürten et
162	al., 2012). The nitrate-water cluster is included as the presence of many nitrate-water clusters of the
163	general formula $(H_2O)_x(HNO_3)_yNO_3^-$ were found, where $x = (1, 2, 3 20)$ and $y = (0, 1)$ . No
164	sensitivity calibration was performed for these measurements, and so all values are reported in
165	normalised signal intensity. Due to the high resolving power of the CI-APi-ToF system (mass
166	resolution of 3500 m/dm and mass accuracy of 20 ppm at 288 m/Q), multiple peaks can be fit at the
167	same unit mass and their molecular formulae assigned. These peaks follow the general formula
168	$C_xH_yO_zN_w$ where $x = 2-20$ , $y = 2-32$ , $z = 4-16$ and $w = 0-2$ , spanning from small organic acids like
169	oxalic and malonic acid through to large dimers of oxidised monoterpene RO2 radicals such as
170	$C_{20}H_{31}O_9N$ . Beyond 500 <i>m/Q</i> , peak fitting and assignment of compositions becomes problematic as
171	signal decreases, mass accuracy decreases, and the total number of chemical compositions
172	increases, so peaks above the $C_{20}$ region have not been assigned, and a number of peaks have been
173	unassigned due to this uncertainty (Cubison and Jimenez, 2015). As proton transfer mostly happens
174	with acids, and nearly all HOM molecules will be charged by adduct formation it is possible to infer
175	the uncharged formula; therefore all HOMs from here onwards will be listed as their uncharged
176	form.

## 179 2.3. Size Distribution Measurements

Two Scanning Mobility Particle Sizer (SMPS) instruments measured particle size distributions at 180 15 minute time resolution, one LongSMPS (TSI 3080 EC, 3082 Long DMA, 3775 CPC, TSI, USA) 181 182 and one NanoSMPS (3082 EC, 3082 Nano DMA, 3776 CPC, TSI, USA) measuring the ranges 14-615 nm and 4-65 nm respectively. A Particle Size Magnifier (A10, Airmodus, FN) linked to a CPC 183 (3775, TSI, USA) measured the sub-3 nm size fraction. The PSM was run in stepping mode, 184 operating at four different saturator flows to vary the lowest size cut-off of particles that it will grow 185 (this cut-off is technically a point of 50% detection efficiency) of <1.30, 1.36, 1.67 and 2.01 nm. 186 The instrument switched between saturator flows per 2.5 minutes, giving a sub-2.01 nm size 187 distribution every 10 minutes. The data was treated with a moving average filter to account for 188 jumps in total particle count, and due to the similar behaviour of the two upper and two lower size 189 cuts, these have been averaged to two size cuts at 1.30 and 1.84 nm. 190

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#### 192 **2.4.** Calculations

193 The condensation sink (CS) was calculated from the size distribution data as follows:

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$$CS = 4\pi D \sum_{d'p} \beta_{m,d'p} d'_p N_{d'p}$$
(1)

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where D is the diffusion coefficient of the diffusing vapour (assumed sulfuric acid),  $\beta_m$  is a transition regime correction (Kulmala et al., 2012), d'<sub>p</sub> is particle diameter, and N<sub>d'p</sub> is the number of particles at diameter d'<sub>p</sub>.

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## 200 **2.5.** Other Measurements

Measurements of the classical air pollutants were measured on the same site, and have been reported in the campaign overview paper (Shi et al., 2019). SO<sub>2</sub> was measured using a 43i SO<sub>2</sub> analyser (ThermoFisher Scientific, USA), O<sub>3</sub> with a 49i O<sub>3</sub> analyser (ThermoFisher Scientific, USA) and NO<sub>x</sub> with a 42i-TL Trace NO<sub>x</sub> analyser (ThermoFisher Scientific, USA), and a T500U

CAPS NO<sub>2</sub> analyser (Teledyne API, USA). VOC mixing ratios were measured using a Proton
 Transfer Reaction-Time of Flight-Mass Spectrometer (PTR-ToF 2000, Ionicon, Austria).

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## 208 **3. RESULTS AND DISCUSSION**

# 209 **3.1.** Characteristics of Sampling Period

A total of five days of CI-API-ToF data were collected successfully, from 2017/06/21 midday 210 through 2017/06/26 midday. New particle formation events were observed on 24<sup>th</sup> June in the late 211 afternoon and 25th June at midday. Some nighttime formation of molecular clusters was seen 212 earlier in the campaign, as were several peaks to the 1.5 - 100 nm size range, likely from pollutant 213 plumes containing freshly nucleating condensable materials. The trace gases, O<sub>3</sub>, SO<sub>2</sub>, NO and NO<sub>2</sub> 214 are plotted in the Figure S1.  $O_3$  shows mid-afternoon peaks, around ~120 ppb on the first two days 215 of the campaign, and 50-70 ppb for the latter days. SO<sub>2</sub> shows a large peak, reaching 4 ppb on 22/06 216 but <1 ppb for the rest of campaign. NO shows strong mid-morning rush hour related peaks, 217 declining towards midday due to being rapidly consumed by O<sub>3</sub>. NO<sub>2</sub> shows large traffic related 218 peaks. The sulfuric acid signal across this period as measured by NO<sub>3</sub><sup>-</sup>CI-APi-ToF showed strong 219 midday peaks, with signal highest on 24/06/2017 and 25/06/2017. The meteorological data are 220 shown in Figure S2 alongside condensation sink (CS). The conditions were generally warm and 221 humid, with temperature reaching its maximum on 25/06/2017, with a peak hourly temperature of 222 31°C. High temperatures were seen on 21/06 and 24/06 also, of 30°C and 26°C respectively. 223

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## 225 **3.2.** Gas Phase HOM Chemistry

# 226 **3.2.1. Bulk chemical properties**

For the peaks that have had chemical formulae assigned, oxidation state of carbon, or  $OS_c$ , can be used to describe their bulk oxidation chemistry.  $OS_c$  is defined as (Kroll et al., 2011)

$$OSc = (2 \times 0:C) - H:C \quad (2)$$

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This does not account for the presence of nitrate ester groups, which has been accounted for 232 233 previously by subtracting five times the N:C ratio (Massoli et al., 2018), under the assumption that all nitrogen containing functionality is in the form of nitrate ester (RONO<sub>2</sub>) groups. In Beijing, 234 multiple sources of nitrate-containing organic compounds are seen, in the forms of amines, nitriles 235 and heterocycles. The variation of oxidation state with carbon number  $(C_n)$  without correction for 236 nitrate esters is plotted in Figure 1. The average oxidation state of carbon in this dataset tends to 237 decrease with an increase to  $C_n$ , highest where  $C_n = 5$ , attributable both to high O:C and peak area 238 for the peak assigned to  $C_5H_{10}N_2O_8$  at m/Q 288.  $C_n = 5$  also shows the greatest distribution of 239 oxidation states, likely due to the high ambient concentration of isoprene and therefore its many 240 241 oxidation products being of high enough signal for many well resolved peaks to be seen in this dataset. It is worth noting that some of the ions plotted here may not form through peroxy radical 242 autoxidation, such as C<sub>5</sub>H<sub>10</sub>N<sub>2</sub>O<sub>8</sub>, which may be a second-generation oxidation product of isoprene 243 under high NO<sub>x</sub> (Lee et al., 2016).  $C_n$ =10 and 15 also see a small increase to average oxidation 244 number compared to their neighbours. The lower oxidation state of the larger products is likely a 245 246 function of two things. First and foremost, any autoxidation mechanism must undergo more steps in order for a larger molecule to reach an equivalent O:C ratio with a smaller one, and the equivalent 247 O:C ratio is ultimately less likely to be reached before the radical is terminated (Massoli et al., 248 2018). Secondly, the lower vapour pressures of these larger products will lead to their partitioning 249 into the condensed phase more readily than the smaller, thus they are more rapidly lost (Mutzel et 250 al., 2015). 251

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The degrees of OSc observed here are similar to those seen in other environments such as during the SOAS campaign in 2013 in southern United States, characterised by low NO/NO<sub>2</sub> and high temperatures, where campaign averages of 0.3 ppb, 0.4-0.5 ppb, and 25°C respectively were

measured, although an additional parameter to account for nitrogen containing VOCs is included in 256 the calculation (Massoli et al., 2018). The OS<sub>c</sub> observed in Beijing is also higher than that seen in 257 the boreal forest environment of Hyytiälä, despite extremely low NO<sub>x</sub> concentrations, likely due to 258 259 low temperature conditions dominating in those conditions (Schobesberger et al., 2013). These relatively similar degrees of oxidation to those seen in other, cleaner, environments are likely due to 260 an interplay between the higher temperatures facilitating rapid hydrogen abstractions (Crounse et 261 al., 2013; Quéléver et al., 2018) and the higher concentrations of NO<sub>x</sub>, HO<sub>2</sub>, and other RO<sub>2</sub> 262 molecules terminating the autoxidation sequence more efficiently (Praske et al., 2018, Rissanen, 263 2018, Garmash et al., 2019). 264

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A mass defect plot is shown in Figure 2, which shows nominal mass plotted against mass defect for 266 all peaks in this dataset. Mass defect is defined as the ion mass minus integer mass. This is shown 267 for two separate daytime periods, one where nucleation was not occurring and HOM concentrations 268 are lower (10:30 – 12:00 23/06/2017) and one where nucleation was occurring under high HOM 269 270 concentrations  $(10:30 - 12:00\ 25/06/2017)$ . The band of lower mass defect is characterised by a number of large peaks with high signal, for example, at m/O 436 the ion (C<sub>2</sub>H<sub>7</sub>N)<sub>2</sub>(H<sub>2</sub>SO<sub>4</sub>)<sub>2</sub>HSO<sub>4</sub><sup>-</sup>. 271 The upper component of the mass defect is dominated by organic compounds, the upper end of 272 more positive mass defect is occupied by molecules with more <sup>1</sup>H (mass defect 7.825 mDa) and <sup>14</sup>N 273 (mass defect 3.074 mDa). The end of less positive mass defect has lower <sup>1</sup>H and more <sup>16</sup>O (mass 274 defect -5.085 mDa); alternatively put, the mass defect reflects the variation in  $OS_c$ . The organic 275 components with more positive mass defects will be more volatile than their lower mass defect 276 counterparts as they will contain fewer oxygen functionalities (Tröstl et al., 2016, Stolzenburg et al., 277 278 2018). These higher volatility products may still contribute to larger size particle growth. The more negative mass defect components will be those of greater O:C and therefore lower volatility, 279 LVOCs, and the yet larger and more oxidised components, ELVOCs (Tröstl et al., 2016). During 280 the nucleation period, the signal intensity for the species in the upper band of more negative mass 281

defect have the most marked increase in concentration, with significantly less difference >500 m/Q. This region 200-400 m/Q will contain most of the C<sub>5</sub>+ monomer HOMs seen in this dataset.

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#### 285 **3.2.2. Diurnal trends of HOMs**

Temporal trends of HOMs in the urban atmosphere can reveal their sources and behaviour in the 286 atmosphere. Most of the HOM species peak in the daytime. These species all follow a similar 287 diurnal trend, as shown in Figure 3. Both the concentrations of O<sub>3</sub> and OH<sup>-</sup> are high during the 288 summer period in Beijing (although the nitrate chemical ionisation technique is not sensitive to all 289 OH oxidation products (Berndt et al., 2015)). Figure S1 shows the time series of concentrations of 290 NO which is considered a dominant peroxy radical terminator of particular importance in the 291 polluted urban environment (Khan et al., 2015). Radicals such as HO<sub>2</sub><sup>-</sup> and RO<sub>2</sub><sup>-</sup> also typically peak 292 during daytime. The HOM components peaking in the daytime are presumed to be the oxidation 293 products of a mixture of anthropogenic and biogenic components, such as alkylbenzenes, 294 monoterpenes and isoprene. The oxidation of monoterpenes, specifically the monoterpene  $\alpha$ -pinene, 295 has been the subject of extensive study recently, with the O<sub>3</sub>-initiated autoxidation sequence being 296 the best characterised (Ehn et al., 2014; Jokinen et al., 2014; Kurtén et al., 2015; Kirkby et al., 297 2016); ozonolysis of  $\alpha$ -pinene opens the ring structure and produces a RO<sub>2</sub> radical (Kirkby et al., 298 2016). In the case of aromatics, OH addition to the ring and the subsequently formed bicyclic 299 peroxy radical is the basis for the autooxidation of compounds such as xylenes and 300 trimethylbenzenes (Molteni et al., 2018; Wu et al., 2017). 301

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The identified compounds have been roughly separated into several categories, each of these plotted in Figure 3. Figure 3ashows the separation of components into non-nitrogen containing HOMs, and nitrogen containing HOMs, or organonitrates (ONs). The ON signal is much higher than that of the HOM, attributable in part to a few ions of high signal, such as the isoprene organonitrate  $C_5H_{10}N_2O_8$ . A few similar structural formulae are seen ( $C_5H_{10}N_2O_6$ ,  $C_5H_{11}NO_6$ ,  $C_5H_{11}NO_7$ , etc),

some of which have been identified as important gas phase oxidation products of isoprene under 308 high NO<sub>x</sub> conditions (Xiong et al., 2015), and their contribution to SOA has been explored 309 previously (Lee et al., 2016). A high nitrophenol signal is also seen,  $C_6H_5NO_3$ . The signal for HOM 310 311 compounds is less dominated by a few large ions. The prevalence of ON compounds points towards the important role of NO<sub>x</sub> as a peroxy radical terminator, with the probability for the RO<sub>2</sub> + NO<sub>x</sub> 312 reaction to produce nitrate ester compounds increasing with the size of the RO<sub>2</sub> molecule (Atkinson 313 et al., 1982). The NO<sub>x</sub> concentrations in urban Beijing are approximately a factor of 10 higher than 314 seen at the Hyytiälä station in Finland as reported by Yan et al. (2016), and hence it is expected to 315 be a more significant peroxy radical terminator. 316

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Despite the very large fluxes of anthropogenic organic pollutants in Beijing, biogenic emissions are 318 still an important source of reactive VOCs in the city, with abundant isoprene oxidation products 319 observed (see above), as well as monoterpene monomers ( $C_{10}H_{16}O_9$ ,  $C_{10}H_{15}O_9N$ ) and some dimer 320 products ( $C_{20}H_{30}O_{11}$ ,  $C_{20}H_{31}O_{11}N$ ). The time series of the signals of all  $C_5$ ,  $C_{10}$  and  $C_{20}$  molecules is 321 plotted in Figure 3b, with C<sub>5</sub> species assumed to be isoprene dominated, C<sub>10</sub> and C<sub>20</sub> assumed to be 322 monoterpene dominated. Signals for isoprene oxidation products are higher, with abundant 323 isoprene nitrate and dinitrate products.  $C_{10}$  products show similar behaviour, with, for example, 324 several  $C_{10}H_{15}O_xN x = 5-9$  compounds seen. The  $C_{20}$  signal intensities are low, and follow the 325 general formula  $C_{20}H_xO_yN_z$ , where x = 26-32, y = 7-11 and z = 0-2; in Figure 3 the signal for  $C_{20}$ 326 compounds has been multiplied by a factor of 50 for visibility. The low signals reflect the lack of 327 RO<sub>2</sub> cross reactions necessary for the production of these accretion products. 328

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Other identified peaks are plotted in Figure 3c. The  $C_2$ - $C_4$  components are summed together, these being small organic acids such as malonic acid and oxalic acid, as well as products such as  $C_4H_7O_6N$ . Malonic acid is the most prominent here, seen both as an  $NO_3^-$  adduct ( $C_3H_4O_4NO_3^-$ ) and a proton transfer product ( $C_3H_3O_4^-$ ) at a ratio of around 2:3. The  $C_6$ - $C_9$  components are assumed to

be dominated by oxidation products of alkylbenzenes such as C<sub>8</sub>H<sub>12</sub>O<sub>5</sub>, although fragments of other 334 compounds, i.e., monoterpenes, can also occupy this region (Isaacman-Vanwertz et al., 2018). It is 335 assumed the majority of the signal for these peaks come from alkylbenzenes. This assumption is 336 337 supported by the relative signal intensity ratios of the oxygen numbers of monomer  $C_8H_{12}O_n$ compounds being similar to those seen for xylene oxidation products in previous work (Molteni et 338 al., 2018). The largest fraction,  $C_{11}$  through  $C_{18}$  includes the larger compounds, oxidation products 339 of larger aromatics, or products of the cross reaction of smaller RO<sub>2</sub> radicals. Here they are grouped 340 without more sophisticated disaggregation as they all follow much the same time series, species 341 such as  $C_{11}H_{11}O_8N$  following the same temporal trends as  $C_{15}H_{16}O_9$  and  $C_{16}H_{24}O_{12}$ . 342

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Nearly all ions with the exception of the larger compounds attributed to the cross reaction of  $C_{10}$ 344 monomers follow similar temporal patterns, with the majority of peaks occurring in the daytime. 345 This reflects the importance of the concentration of atmospheric oxidants. Some selected oxidation 346 products are plotted against their precursor VOCs in Figure 4. The concentration of isoprene is 347 plotted against the signal of a nitrate HOM product, C<sub>5</sub>H<sub>9</sub>NO<sub>6</sub> (Xiong et al., 2015; Lee et al., 2016), 348 while monoterpenes are plotted against C<sub>10</sub>H<sub>16</sub>O<sub>9</sub> (Ehn et al., 2014; Berndt et al., 2016; Yan et al., 349 2016; Kirkby et al., 2016; Massoli et al., 2018), and C<sub>2</sub>-benzenes against C<sub>8</sub>H<sub>12</sub>O<sub>6</sub> (Molteni et al., 350 2018; Wang et al., 2017). The first half of the time series shows little correlation between the VOC 351 species and the resultant oxidation products, while isoprene, monoterpenes and C2-benzenes follow 352 their usual diurnal cycles, isoprene having the most distinct with a strong midday peak. The latter 353 two days, however, show similar and coinciding peaks in both the VOCs and HOMs - HOMs show 354 afternoon peaks on both days, and an initial shelf on the final half day. The C<sub>5</sub>H<sub>9</sub>NO<sub>6</sub> peak follows 355 356 some of the peaks of the isoprene, but not all (e.g., morning shelf of isoprene on 24/06). Concentrations of isoprene do not seem to determine directly the signal of HOM, as the day with 357 the lowest isoprene of all is the day with highest C<sub>5</sub>H<sub>9</sub>NO<sub>6</sub>. The C<sub>10</sub>H<sub>16</sub>O<sub>9</sub> trace has coincidental 358

peaks with the monoterpene trace also, including two 4-hour separated simultaneous peaks on

25/06. The peaks in the concentrations of C<sub>2</sub>-benzenes are nearly synchronous with the peaks in 360 C<sub>8</sub>H<sub>12</sub>O<sub>6</sub>, for which the dataexhibit a strong mid afternoon peak likely due to the lack of an efficient 361 ozonolysis reaction pathway; the main oxidant of C<sub>2</sub>-benzenes is the OH radical. Trends of both C<sub>3</sub> 362 363 benzenes and their HOMs are much the same as C<sub>2</sub> benzenes as discussed above, pointing to similar sources and oxidation chemistries. . The concentration of precursor VOC is likely a driving force in 364 the identity and quantity of various HOM products, but not the sole determinant, as while there are 365 simultaneous peaks of VOCs and HOMs, both the condensation sink and oxidant concentrations 366 also influence HOM product signals. 367

368

The first half of campaign measurements is marked by an episode of low HOM signals. A diurnal cycle still exists but it is weak. The radiation intensity was significantly lower on these prior days than it was on the 24th. No data is available for the final period of measurement. Ozone is higher on the prior measurement days with lower HOM signals (see Figure S1). It is therefore plausible that light intensity, and therefore OH concentration is one of the main drivers of HOM concentrations in Beijing.

375

The C<sub>20</sub> compounds plotted in Figure 3b show no strong diurnal sequence, contrasting with other 376 HOMs. We can presume that all  $C_{20}$  compounds identified are the result of the reaction of two 377 monoterpenoid  $C_{10} RO_2$  radicals, a reasonable assumption as all identified  $C_{20}$  species follow the 378 general formula outlined for these reactions ( $C_{20}H_{28-32}O_{6-16}$ ). The formation of  $C_{20}$  dimers is 379 dependent upon two processes, initial oxidation of monoterpenes, and RO2-RO2 termination. Initial 380 oxidation is contingent upon oxidant concentration, which is highest in the daytime, and  $RO_2$ -RO<sub>2</sub>. 381 382 termination is contingent upon the probability of the molecular collision between the RO<sub>2</sub>. molecules occurring before other radical termination (i.e., RO<sub>2</sub>-NO<sub>x</sub>, or RO<sub>2</sub>-HO<sub>2</sub>). There is likely 383 a strong diurnal sequence in the dominant RO2 termination mechanisms across the day period, and 384 the combination of the two factors discussed above results in there being no strong diurnal trend in 385

these molecules. A lower oxidant concentration at night results in less  $RO_2$  molecules, but less NO and  $HO_2$  results in a greater chance for those  $RO_2$  molecules to dimerise (Rissanen, 2018, Garmash et al., 2019). As the levels of  $NO_x$  in Beijing fall, the peroxy radical termination reactions will be less probable compared to continued autoxidation (Praske et al., 2018), and it is expected that more oxidised HOM products will be seen with lower volatilities and therefore a greater potential contribution to earlier stage particle formation and growth.

392

# 393 **3.3.** New Particle Formation

Nearly all the signal intensity in the CI-APi-ToF instrument arises from molecules charged by NO<sub>3</sub><sup>-</sup>, 394 therefore plotting the unit mass resolution data (the data gained by integrating over the entire area at 395 each m/Q integer) against time describes simply the evolution of oxidised organic molecules, acids 396 and their molecular clusters both with each other and stabilising amine species. This is done in 397 Figure 5. As the signal intensity varies by factors of 10 from mass to mass, each value has been 398 normalised so they have maxima at 1. This has been done separately for two days for clarity, as the 399 signal intensity also varies from day to day. PSM data for these two days is plotted in Figure 5 also, 400 with both total particle count >1.30 nm in black and the number difference between the lower and 401 upper size cuts (1.30 and 1.84 nm) in blue, which shows the number of particles between these 402 sizes. The relationship between mass and electrical mobility diameter can be defined thus (Tammet, 403 1995), 404

405

$$d_e = (\frac{6m}{\pi \rho})^{\frac{1}{3}} + d_g \qquad (3)$$

406

where  $d_e$  is the electrical mobility diameter of the cluster or particle, m is the mass of the cluster or particle expressed in kg,  $\rho$  is the density and  $d_g$  is the effective gas diameter, determined to be 0.3 nm for smaller particles (Larriba et al., 2011). We can use this to draw a comparison between the PSM and CI-APi-ToF measurements. If a density of 1.2 g cm<sup>-3</sup> is assumed, then once molecular clusters reach the >400 *m/Q* range, they will be seen in the lowest size cut of the PSM, or >700 *m/Q*  412 if a density of 2.0 g cm<sup>-3</sup> is assumed. A full table of densities is provided in the Supplementary
413 Information.

414

415 A burst in the signal seen by the CI-APi-TOF occurs first in the late morning in the top panel of Figure 5, and this is at the same time as peaks begin to rise in the identified HOMs (see Figure 3). 416 Here, the PSM is not available due to an instrumental fault until 16:00; however, at that point, an 417 elevation to particle count and a large elevation to cluster count can be seen. Moving into the 418 evening period, the mass contour shows peaks to larger masses >400 m/Q. This is likely dimerised 419 compounds and products of NO<sub>3</sub> chemistry with little contribution to newly forming particles, but 420 still sensitive to chemical ionisation by  $NO_3^{-}$ . Many of these peaks cannot be assigned due to 421 uncertainties in the structural formula assignment for higher mass peaks, as the number of possible 422 dimerised compounds is many, being the combination of most possible RO<sub>2</sub> radicals. Graphically, 423 these are over-represented in Figure 5 due to the normalisation, their signals (especially >500 m/Q) 424 are much lower than the signals <400 m/Q. 425

426

The second day plotted in the lower panel of Figure 5 (25/06/2017) shows a strong afternoon peak 427 to the HOMs (for most HOMs, stronger than that on the day prior). Particle formation is shown in 428 the PSM data. A strong midday peak to particle number is seen with two distinct peaks to cluster 429 count. These two peaks are not coincidental with the two peaks to HOM signal (i.e., nitrogen-430 containing HOMs in Figure 3a peaking at 11:00 and 16:00). Sulfuric acid, however, does peak 431 synchronously with the particle number count. Sulfuric acid is plotted across the contour plot in 432 Figure 6, where PSM data is also shown in the bottom panel. The peak to CI-APi-TOF mass signal, 433 visible in Figure 5 occurs at around 12:00/13:00, peaks in the PSM cluster count occur at 10:00 and 434 13:00. Peaks in mass up to 550 m/Q are seen in the CI-APi-ToF at 13:00. Assuming the density of 435 these species is  $\leq 1.6$  g cm<sup>-3</sup> then these will be suitably sized to be grown in the PSM saturator. 436 These newly formed particles then go on to grow and contribute significantly to the larger particle 437

438 count (Figure S3). As initial particle formation coincides with sulfuric acid signal peaks and before
439 HOM signals peak, it can be assumed on these days, the HOM contribution to the initial particle
440 formation is modest.

441

There is recent strong evidence to suggest that the driving force of the earliest stages of particle 442 formation in urban Shanghai is from sulfuric acid and C<sub>2</sub>-amines (Yao et al., 2018), and the 443 coincidental peaks of sulfuric acid with new particles as seen in Figure 6 suggest a similar 444 behaviour. Dimethylamine (DMA) can efficiently stabilise the sulfuric acid clusters (Almeida et al., 445 2013). Here, few larger sulfuric acid-DMA clusters were visible in the dataset, as seen in the work 446 by Yao et al., 2018, although five sulfuric acid-dimethylamine (SA-DMA) ions were observed, the 447 others were likely too low in signal to be confidently resolved from their neighbouring peaks; 448 however, clusters of up to 4 sulfuric acid ions and 3 dimethylamine molecules were seen, with 449 similar diurnal trends to sulfuric acid. The scarcity of SA-DMA clusters is likely due to 450 instrumental conditions, rather than their absence in the atmosphere. The nitrate chemical ionisation 451 system tends to evaporate amine compounds upon charging, and as specific voltage-tuning setups 452 can lend themselves towards preservation or breakage of molecular clusters, the signal for larger 453 sulfuric acid clusters was also very weak. The formation of HOM-sulfuric acid clusters is unlikely 454 under atmospheric conditions (Elm et al., 2017) and few of these were observed. Signals of HOMs 455 seem to coincide with later particle growth; it can be expected that HOM molecules make a more 456 significant contribution to particle growth than to early particle formation, with the largest and most 457 oxidised being involved in early growth, and the smaller and less oxidised contributing to later 458 growth as the necessary vapour pressure properties become less demanding. 459

460

## 461 **4. CONCLUSIONS**

The average degree of HOM oxidation in Beijing is comparable with that seen in otherenvironments. Rapid intramolecular hydrogen shifts during autoxidation due to the higher

temperatures are probably offset by the frequent termination reactions due to high  $NO_x$ 

465 concentrations.  $OS_c$  values seem to be marginally higher for biogenic species.

466

The temporal trend of nearly every HOM shows afternoon or evening maxima. Both O<sub>3</sub> and OH have high daytime concentrations and these likely drive the initial oxidation steps. The species arising from alkylbenzene precursors show sharper afternoon peaks, probably since their oxidation is OH dominated. Many of the rest of the peaks, coming from largely BVOC precursors show broader daytime peaks, being influenced by O<sub>3</sub> also. There seems to be no direct link between VOC concentrations and HOM signals, with days of lower precursor VOC sometimes having higher HOM signals and vice versa.

474

Initial particle formation coincides with peak sulfuric acid signals, while the growth of the particles 475 correlates more closely with the signals of HOMs. This is very similar to behaviour observed in a 476 study of NPF in Shanghai which was attributed to sulfuric acid-dimethylamine-water nucleation 477 with condensing organic species contributing to particle growth (Yao et al., 2018), and this is 478 further backed up by numerous SA-DMA clusters present in this dataset. The freshly formed 479 particles grow and contribute significantly to total particle loading. This is visible when the unit 480 mass CI-APi-ToF data is plotted as a contour plot, and further to this is visible in the PSM data, 481 with bursts to both total number count >1.30 nm and the number of molecular clusters between 1.30 482 and 1.84 nm. As NO<sub>x</sub> levels fall in Beijing due to traffic emission control measures being enforced 483 it is likely that autoxidation will become increasingly significant in the new particle formation 484 processes. The number of molecules detected by the  $NO_3^{-3}$  CIMS is undoubtedly many more than 485 486 have had formulae assigned here, but to identify more requires a more sophisticated data deconvolution. 487

488

489

# 490 DATA ACCESSIBILITY

- 491 Data supporting this publication are openly available from the UBIRA eData repository at
   492 https://doi.org/10.25500/edata.bham.00000304
- 493

#### 494 AUTHOR CONTRIBUTIONS

The study was conceived and planned by RMH and ZS. DCSB and JB set up and operated the main instrumental measurements, and JB prepared the first draft of the paper and responded to comments from RMH and ZS. CNH and WJA contributed the hydrocarbon data and provided comments on the draft manuscript, and ES and JL contributed the gas phase pollutant data.

499

#### 500 **COMPETING INTERESTS**

501 The authors have no conflict of interests.

502

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# 892 **FIGURE LEGENDS:**

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- Figure 1 Oxidation state of carbon calculated as two times the oxygen to carbon ratio minus the hydrogen to carbon ratio against carbon number for (colored) individual ions and (blue circles) signal weighted average for each carbon number. Area and colour are both proportional to the peak area for each ion
- **Figure 2** Mass defect plot of fitted mass spectral peaks between 100-600mass units on (a) 10:30 899 - 12:00 23/06/2017, a non nucleation day, and (b) 10:30 -12:00 25/06/2017, a 900 nucleation day. Mass defect can be defined as the mass - integer mass. The size of point 901 is proportional to the signal intensity. As <sup>1</sup>H has a positive mass defect (1.007276 Da), 902 the upward trend along the horizontal indicates increasing carbon chain length, and 903 differences at similar masses are due to increasing oxygen functionality, clustering with 904 species such as sulfuric acid (negative mass defect) and ammonia (positive mass 905 defect), as <sup>16</sup>O and <sup>32</sup>S have negative mass defects (15.9949 and 31.9721 Da 906 respectively), while <sup>14</sup>N has a positive mass defect at 14.0031 Da. 907
- Figure 3 Summed time series of the concentrations of (A) all non-nitrogen containing HOMs and all organonitrates identified, (B) C5, C10 and C20 components, assumed to be dominated by isoprene, monoterpene monomer and monoterpene dimers, signal for C20 multiplied 50 times to fit scale, and (C) summed C6 C9 components, and summed C11 C18 components, assumed to be dominated by alkylbenzenes and other larger components respectively.
- Figure 4 Time series for the whole sampling campaign for the concentrations of (left axis) VOCs as measured by PTR-ToF and (right axis) a selected HOM product associated with that precursor.
- 920Figure 5Normalised unit mass NO3- CI-APi-ToF signal intensity on 24/06/2017 (A) and92125/06/2017 (B). Each individual unit mass was normalised to a maximum of 1. Each922period is normalised separately so the individual signal maxima on each day are visible.923The graph is plotted between 200-600 mass units, with every 10 mass units averaged for924simplicity. On the secondary axis is plotted PSM data, both total particle count >1.30925nm (black trace) and total clusters between 1.30 and 1.84 nm (blue trace). Data is926plotted at 1 hour time resolution.
- 927928Figure 6SMPS + PSM contour plot for two nucleation days on 24/06/2017 and 25/06/2017. Data929in bottom panel is from the PSM instrument, top panel from NanoSMPS, units in colour930bar are  $log_{10}$  (dN/logD<sub>p</sub>) for N in cm<sup>-3</sup>. Points signify normalised sulfuric acid931concentration (right axis) as measured by CI-APi-ToF.



Figure 1. Oxidation state of carbon plotted against carbon number for (colored) individual ions and
(blue circles) signal weighted average for each carbon number. Area and colour are both proportional
to the peak area for each ion.



942 Figure 2. Mass defect plot of fitted mass spectral peaks between 100-600mass units on (a) 10:30 -943 12:00 23/06/2017, a non nucleation day, and (b) 10:30 -12:00 25/06/2017, a nucleation day. Mass defect can be defined as the mass - integer mass. The size of point is proportional to the signal 944 intensity. As <sup>1</sup>H has a positive mass defect (1.007276 Da), the upward trend along the horizontal 945 indicates increasing carbon chain length, and differences at similar masses are due to increasing 946 oxygen functionality, clustering with species such as sulfuric acid (negative mass defect) and 947 ammonia (positive mass defect), as <sup>16</sup>O and <sup>32</sup>S have negative mass defects (15.9949 and 31.9721 Da 948 respectively), while <sup>14</sup>N has a positive mass defect at 14.0031 Da. 949

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Figure 3. Summed time series of the concentrations of (A) all non-nitrogen containing HOMs and all organonitrates identified, (B) C5, C10 and C20 components, assumed to be dominated by isoprene, monoterpene monomer and monoterpene dimers, signal for C20 multiplied 50 times to fit scale, and (C) summed C6 - C9 components, and summed C11 - C18 components, assumed to be dominated by alkylbenzenes and other larger components respectively.



Figure 4. Time series for the whole sampling campaign for the concentrations of (left axis) VOCs as
 measured by PTR-ToF and (right axis) a selected HOM product associated with that precursor.



Figure 5. Normalised unit mass NO<sub>3</sub><sup>-</sup> CI-APi-ToF signal intensity on 24/06/2017 (A) and 25/06/2017
(B). Each individual unit mass was normalised to a maximum of 1. Each period is normalised separately so the individual signal maxima on each day are visible. The graph is plotted between 200-600 mass units, with every 10 mass units averaged for simplicity. On the secondary axis is plotted PSM data, both total particle count >1.30 nm (black trace) and total clusters between 1.30 and 1.84 nm (blue trace). Data is plotted at 1 hour time resolution.



Figure 6. SMPS + PSM contour plot for two nucleation days on 24/06/2017 and 25/06/2017. Data in bottom panel is from the PSM instrument, top panel from NanoSMPS, units in colour bar are  $log_{10}$ (dN/logD<sub>p</sub>) for N in cm<sup>-3</sup>. Points signify normalised sulfuric acid concentration (right axis) as measured by CI-APi-ToF.