

<https://helda.helsinki.fi>

Contribution of Atmospheric Oxygenated Organic Compounds to Particle Growth in an Urban Environment

Qiao, Xiaohui

2021-10-19

Qiao , X , Yan , C , Li , X , Guo , Y , Yin , R , Deng , C , Li , C , Nie , W , Wang , M , Cai , R , Huang , D , Wang , Z , Yao , L , Worsnop , D R , Bianchi , F , Liu , Y , Donahue , N M , Kulmala , M & Jiang , J 2021 , ' Contribution of Atmospheric Oxygenated Organic Compounds to Particle Growth in an Urban Environment ' , Environmental Science & Technology , vol. 55 , no. 20 , pp. 13646-13656 . <https://doi.org/10.1021/acs.est.1c02095>

<http://hdl.handle.net/10138/341154>

<https://doi.org/10.1021/acs.est.1c02095>

submittedVersion

Downloaded from Helda, University of Helsinki institutional repository.

This is an electronic reprint of the original article.

This reprint may differ from the original in pagination and typographic detail.

Please cite the original version.

This document is confidential and is proprietary to the American Chemical Society and its authors. Do not copy or disclose without written permission. If you have received this item in error, notify the sender and delete all copies.

**Contribution of atmospheric oxygenated organic compounds
to particle growth in an urban environment**

Journal:	<i>Environmental Science & Technology</i>
Manuscript ID	es-2021-02095k.R2
Manuscript Type:	Article
Date Submitted by the Author:	14-Aug-2021
Complete List of Authors:	<p>Qiao, Xiaohui; Tsinghua University, School of Environment Yan, Chao; University of Helsinki, Institute for Atmospheric and Earth System Research INAR / Physics Li, Xiaoxiao; Tsinghua University, School of Environment Guo, Yishuo; Beijing University of Chemical Technology, Beijing Advanced Innovation center for Soft Matter Science and Engineering Yin, Rujing; Tsinghua University Deng, Chenjuan; Tsinghua University, School of Environment Li, Chang; Beijing University of Chemical Technology, Beijing Advanced Innovation center for Soft Matter Science and Engineering Nie, Wei; Nanjing University, School of Atmospheric Sciences Wang, MingYi; Carnegie Mellon University, Chemistry Cai, Runlong; University of Helsinki, Faculty of Science Huang, Dan Dan; Shanghai Academy of Environmental Sciences, CBME Wang, Zhe; The Hong Kong University of Science and Technology, Division of Environment and Sustainability Yao, Lei; University of Helsinki, Institute for Atmospheric and Earth System Research INAR / Physics Worsnop, Douglas; Aerodyne Research Inc, Bianchi, Federico; University of Helsinki, Institute for Atmospheric and Earth System Research INAR / Physics Liu, Yongchun; Beijing University of Chemical Technology, Beijing Advanced Innovation center for Soft Matter Science and Engineering Donahue, Neil M.; Carnegie Mellon University, Chemistry Kulmala, Markku; Helsingin Yliopisto, Dept. of Physical Sciences Jiang, Jingkun; Tsinghua University, School of Environment</p>

SCHOLARONE™
Manuscripts

1 Contribution of atmospheric oxygenated organic compounds to
2 particle growth in an urban environment

3

4 *Xiaohui Qiao*¹, *Chao Yan*^{2,3,*}, *Xiaoxiao Li*¹, *YiShuo Guo*³, *Rujing Yin*¹, *Chenjuan Deng*¹, *Chang*
5 *Li*³, *Wei Nie*⁴, *Mingyi Wang*⁵, *Runlong Cai*^{2,3}, *Dandan Huang*⁷, *Zhe Wang*⁸, *Lei Yao*², *Douglas R.*
6 *Worsnop*^{2,9}, *Federico Bianchi*^{2,3}, *Yongchun Liu*³, *Neil M. Donahue*^{5,6}, *Markku Kulmala*^{2,3},
7 *Jingkun Jiang*^{1,*}

8 ¹ State Key Joint Laboratory of Environment Simulation and Pollution Control, School of
9 Environment, Tsinghua University, 100084 Beijing

10 ² Institute for Atmospheric and Earth System Research/Physics, Faculty of Science, University of
11 Helsinki, 00014 Helsinki, Finland

12 ³ Aerosol and Haze Laboratory, Beijing Advanced Innovation Center for Soft Matter Science and
13 Engineering, Beijing University of Chemical Technology, 100029 Beijing, China

14 ⁴ Joint International research Laboratory of Atmospheric and Earth System Research, School of
15 Atmospheric Sciences, Nanjing University, Nanjing, China

16 ⁵ Center for Atmospheric Particle Studies, Carnegie Mellon University, Pittsburgh, PA, USA

17 ⁶ Department of Chemistry, Carnegie Mellon University, Pittsburgh, PA, USA

18 ⁷ State Environmental Protection Key Laboratory of Formation and Prevention of Urban Air
19 Pollution Complex, Shanghai Academy of Environmental Sciences, Shanghai, China

20 ⁸ Division of Environment and Sustainability, The Hong Kong University of Science and
21 Technology, Hong Kong SAR, China

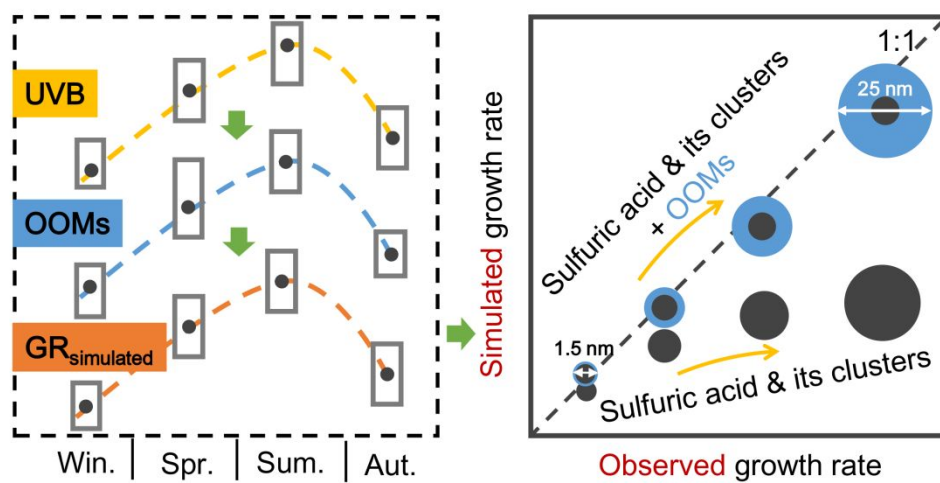
22 ⁹ Aerodyne Research Inc., Billerica, Massachusetts 01821, USA

23

24

25

26 Abstract Graphic:



27

28

29 **ABSTRACT.** Gas-phase oxygenated organic molecules (OOMs) can contribute substantially to
30 the growth of newly formed particles. However, the characteristics of OOMs and their
31 contributions to particle growth rate are not well understood in urban area, which has complex
32 anthropogenic emissions and atmospheric conditions. We performed long-term measurement of
33 gas-phase OOMs in urban Beijing during 2018-2019 using nitrate-based chemical ionization mass
34 spectrometry. OOM concentrations showed clear seasonal variations, with the highest in summer
35 and the lowest in winter. Correspondingly, calculated particle growth rates due to OOM
36 condensation were highest in summer, followed by spring, autumn, and winter. One prominent
37 feature of OOMs in this urban environment was a high fraction (~75%) of nitrogen-containing
38 OOMs. These nitrogen-containing OOMs contributed only 50%-60% of the total growth rate led
39 by OOM condensation, owing to their slightly higher volatility than non-nitrate OOMs. By
40 comparing the calculated condensation growth rates and the observed particle growth rates, we
41 showed that sulfuric acid and its clusters are the main contributors to the growth of sub-3 nm
42 particles, with OOMs significantly promoting the growth of 3-25 nm particles. In winter Beijing,
43 there are missing contributors to the growth of particles above 3 nm, which remain to be further
44 investigated.

45 **KEYWORDS.** Oxygenated Organic Molecules, Seasonal Variation, Particle Growth, Nitrogen-
46 containing OOMs, Urban Environment

47 **SYNOPSIS.** Atmospheric oxygenated organic molecules with clear seasonal variations contribute
48 significantly to the growth of newly formed nanoparticles in urban Beijing.

49

50 **1 Introduction**

51 Atmospheric new particle formation (NPF) happens ubiquitously in various environments, and
52 particles growing past a size of around 50 nm can be activated as the cloud condensation nuclei
53 (CCN), which influence cloud formation and climate.¹⁻³ NPF in polluted environments, despite the
54 severe suppression by high aerosol loadings, has been shown to produce a large number of
55 nanoparticles; these particles, if they survive scavenging by large existing particles, can grow and
56 act as CCN.⁴⁻⁶ Therefore, as a crucial determining factor for particle survival, particle growth rate
57 can substantially influence the CCN budget.⁷

58 Condensable vapors, such as sulfuric acid, are important contributors to the growth of particles in
59 various environments due to their low volatility.⁸⁻¹⁰ Observations in Atlanta indicated that
60 condensation of sulfuric acid and the subsequent equilibration with ammonia could substantially
61 account for nanoparticle growth.¹¹ Also, it has been illustrated recently that condensation of H₂SO₄
62 and H₂SO₄-amine clusters contribute significantly to the growth of sub-3 nm particles in urban
63 Beijing; however, sulfuric acid and its clusters alone cannot explain the growth of particles above
64 3 nm.¹⁰ This is still far from “climate-relevant” size. There are still some missing contributors to
65 particle growth, especially beyond the earliest stages.

66 Gas-phase oxygenated organic molecules (OOMs) are important for particle growth and even
67 largely explain the particle growth rate in forested areas owing to their low volatility.¹²⁻¹⁴ OOMs
68 can be produced from the oxidation of volatile organic compounds (VOCs), including
69 anthropogenic volatile organic compounds (AVOCs, e.g., benzene and toluene) and biogenic
70 volatile organic compounds (BVOCs, e.g., isoprene, monoterpenes, and sesquiterpenes).¹⁵⁻¹⁸
71 Chamber experiments have shown that the oxidation products of monoterpenes alone can trigger
72 particle formation and drive the following growth.¹⁹⁻²² Consistently, ambient observations in a
73 boreal forest in southern Finland (Hyytiälä) have shown that BVOC-derived OOMs are sufficient
74 to explain particle growth over the size range of 3-50 nm.¹³

75 In contrast to remote forests, AVOCs are usually the dominant VOC species in urban environments,
76 and NO_x levels are usually much higher. Thus the characteristics of OOMs in polluted urban
77 environments might be substantially different from those in forested areas. For BVOCs, auto-
78 oxidation of RO₂ radicals is an important pathway of OOM formation.^{14, 18, 23} Each auto-oxidation
79 step results in an addition of two oxygen atoms, which greatly increases the oxygen number of
80 OOMs.^{16, 24, 25} For AVOCs, multi-generation oxidation is also an important pathway in OOMs
81 formation.²⁶ This is mainly because the abundant NO_x in urban areas can suppress the auto-
82 oxidation and terminate the chain propagation,²⁷ but also because the oxidation lifetime of later-
83 generation AVOC products can be much lower than the precursor AVOCs.²⁸ As a result, OOMs
84 generated under high NO_x conditions may have relatively higher volatilities than those in low NO_x
85 conditions.^{26, 27, 29} In wintertime Beijing, Yan et al.³⁰ demonstrated that OOMs are important for
86 particle growth to CCN size. In another study in summertime Beijing, Brean et al.³¹ showed that
87 elevated OOM concentrations and particle growth coincided, implying an important contribution

88 of OOMs to particle growth in urban areas. However, neither study had a quantitative evaluation
89 of the contribution of OOMs to particle growth, and the contribution in different seasons has not
90 been examined.

91 We conducted measurements of OOMs in urban Beijing during Jan. 2018 – Aug. 2019 covering
92 four seasons. We investigated the overall characteristics of OOMs in urban Beijing in comparison
93 to those in other atmospheric environments. We further evaluated the contribution of OOMs to
94 particle growth over a size range of 1.5-25 nm and explored the dominant factors of its seasonal
95 variation. Finally, we compared the calculated particle growth rates explained by sulfuric acid
96 (include its clusters) and non-/nitrated OOMs to the observed values.

97 **2 Methods**

98 **2.1 Measurements**

99 The field measurements were conducted at an urban site located inside the west campus of Beijing
100 University of Chemical Technology. The station is ~500 meters west to the Third Ring Road and
101 is surrounded by residential and commercial areas. A detailed description about this site can be
102 found in previous studies.^{10, 32} The measurements were between Jan. 2018 and Aug. 2019, covering
103 all the four seasons and including 27 NPF events with reliable OOMs and particle growth rate data
104 in total. The exact duration of the four seasons are Mar. 3 – Apr. 7 2018 for spring with 5 NPF
105 cases, Oct. 16 – Nov. 31 for autumn with 5 NPF cases, Jan. 25 – Feb. 6 and Dec. 10 – Dec. 24
106 2018 for winter with 11 NPF cases, and Jun. 15 – Aug. 31 2019 for summer with 6 NPF cases,
107 respectively.

108 OOMs and H₂SO₄ were measured with nitrate-based chemical ionization atmospheric pressure
109 interface time-of-flight mass spectrometers (CI-APi-ToF, Aerodyne Research Inc.).^{33, 34} A high-
110 resolution time-of-flight mass spectrometer with the m/z resolution of ~4000 Th/Th (CI-APi-
111 HToF) was used in 2018, and a long time-of-flight mass spectrometer with the m/z resolution of
112 ~10000 Th/Th (CI-APi-LToF) was used in 2019. Both instruments were calibrated for H₂SO₄
113 sensitivity and m/z-dependent transmission efficiency following previous studies.^{20, 35} Details of
114 the sampling configurations are similar with those previously described.¹⁰ In addition, H₂SO₄
115 clusters in this study refer to clusters containing up to four H₂SO₄ molecules and some base

116 molecules as stabilizing agents. The concentration of these clusters was estimated using the
117 simplified kinetic model described in Cai et al.³⁶

118 Particle number concentrations and size distributions ranging from 1 nm to 10 μm were measured
119 by a diethylene glycol scanning mobility particle spectrometer (DEG-SMPS; 1-7.5 nm)^{37, 38}
120 together with a particle size distribution system (PSD; 3-10 μm).³⁹ A core sampling method was
121 used in the DEG-SMPS system to achieve an aerosol sampling efficiency close to 100%.⁴⁰ A
122 miniature cylindrical differential mobility analyzer (mini-cyDMA) was equipped for sub-10 nm
123 aerosol classification and a modified DEG-UCPC with two-stage temperature control chamber
124 was used for as the front equipment for particle counting.⁴¹ Meteorological data, including
125 temperature, ambient pressure, wind speed, and wind direction, were measured with a local
126 weather station (Vaisala, AWS310).

127 2.2 OOMs quantification

128 OOM identification was performed with the tofTools package developed by Junninen et al.⁴² The
129 low background noises of the CI-APi-ToF are automatically removed in software. After obtaining
130 the signals of assigned peaks, the concentrations of OOMs are calculated using Eq. 1.^{20, 35}

$$[\text{OOM}_i] = C \times T_i \times \ln \left(1 + \frac{[\text{OOM}_i \cdot \text{NO}_3^-]}{\sum_{i=0}^2 [(\text{HNO}_3)_i \cdot \text{NO}_3^-]} \right) \quad \text{Eq. (1)}$$

131 Here, OOM signals in counts per second $[\text{OOM}_i \cdot \text{NO}_3^-]$ are normalized to the sum of reagent ion
132 signals, including NO_3^- , $(\text{HNO}_3)\text{NO}_3^-$, and $(\text{HNO}_3)_2\text{NO}_3^-$. T_i is the mass-dependent transmission
133 efficiency of the instrument, which was determined by adding different perfluorinated acid vapors
134 in sufficient amounts to deplete NO_3^- .³⁵ C is the calibration coefficient obtained for H_2SO_4 using
135 a homemade H_2SO_4 generator.⁴³ We conducted four calibrations during the observation period.
136 The calibration coefficients are 4.79×10^9 and $4.5 \times 10^9 \text{ cm}^{-3} \cdot \text{cps}^{-1}$ for CI-APi-HToF in 2018, 6.07
137 $\times 10^9$ and $6.47 \times 10^9 \text{ cm}^{-3} \cdot \text{cps}^{-1}$ for CI-APi-LToF in 2019. Here, we assume OOMs and H_2SO_4
138 have the same collision and binding efficiency with NO_3^- .¹⁸ The charging efficiency for some less
139 functionalized OOMs is likely lower than that for H_2SO_4 , and therefore, the total OOM
140 concentration measured by nitrate-based CI-APi-TOF should be regarded as a lower limit.

141 2.3 OOMs volatility estimation

142 Due to the lack of molecular structure information, OOM volatility was estimated using volatility
 143 parameterization methods based on the number of carbon, oxygen, and nitrogen atoms of the
 144 identified molecule.⁴⁴ As suggested by a recent study,⁴⁵ two volatility parameterization methods
 145 that treat OOMs with different functional groups were adopted in this study. One is for OOMs that
 146 mainly contain hydroperoxide groups, such as products via auto-oxidation:¹³

$$\log_{10} C_{300K}^* = (25 - n_C)b_C - (n_O - 3n_N)b_O - 2\frac{(n_O - 3n_N)n_C}{(n_C + n_O - 3n_N)}b_{CO} - n_Nb_N \quad \text{Eq. (2)}$$

147 where $b_C = 0.475$, $b_O = 0.2$, $b_{CO} = 0.9$, and $b_N = 2.5$, and n_C , n_O , and n_N are the number of carbon,
 148 oxygen, and nitrogen atoms in the compounds, respectively. The other is for OOMs that mainly
 149 contain hydroxyl and carboxylic groups, such as products via multi-generation oxidation:⁴⁵

$$\log_{10} C_{300K}^* = (25 - n_C)b_C - (n_O - 2n_N)b_O \quad \text{Eq. (3)}$$

150 where $b_C = 0.475$, and $b_O = 2.3$. A nitrate group reduces the similar extent of vapor pressure like
 151 a hydroxyl, and is replaced by hydroxyl for simplicity.⁴⁵ In terms of the oxidation pathway and the
 152 resulting functional groups, oxidation products of isoprene are also suitable for Eq. 3.²⁴ An
 153 empirical dividing line of $H/C = -0.2 \cdot O/C + 1.5$ is adopted in this study to separate OOMs with
 154 different kinds of main functional groups corresponding to the above two parameterization
 155 methods. The volatility of OOM above the line is estimated by Eq. 2, and otherwise by Eq. 3. More
 156 details about the volatility estimation method are in the supplemental information.

157 OOMs measured by nitrate CI-APi-ToF are partly extremely low or low volatility organic
 158 compounds in Beijing (Fig. S6). The over-saturated concentration of an OOM determines whether
 159 it can condense onto particulate matter or not. Due to the Kelvin effect, lower volatility is required
 160 for OOMs to condense onto smaller particles. According to the measured OOMs concentration in
 161 Beijing, OOMs with $\log_{10}(c^*) \geq 1$ could hardly condense on particles smaller than 25 nm (Fig.
 162 S16). Therefore, we define OOMs with $\log_{10}(c^*) \leq 0$ as condensable OOMs, most of which are
 163 extremely low-volatility organic compounds (ELVOC, c^* in 10^{-8} - 10^{-4} $\mu\text{g}\cdot\text{m}^{-3}$) and low-volatility
 164 organic compounds (LVOC, c^* in 10^{-4} - 10^{-1} $\mu\text{g}\cdot\text{m}^{-3}$).⁴⁴

165 2.4 Vapor condensation growth rate simulation

166 The condensation growth rate (GR), defined as $GR = dd_p/dt$, represents the change in particle
 167 size per unit time. The particle growth rate contributed by OOM condensation for a given
 168 diameter was calculated as Eq. 4,^{19, 46} and GR for a particle size bin is calculated by averaging
 169 the GR of all the particles in this bin.

$$\frac{dd_p}{dt} = \left(\frac{d_p + d_i}{d_p}\right)^2 \frac{\bar{c}_{i,p}}{2\rho} \alpha_{i,p} B_{i,p} [C_i^v - a_{i,p} C_i^0] \quad \text{Eq. (4)}$$

170 where d_p , d_i are the diameter of the particle and the condensable vapor, respectively; $\bar{c}_{i,p}$ is the
 171 mean relative thermal velocity between a condensable vapor molecular and a particle; ρ is the
 172 density of the particles; $\alpha_{i,p}$ is the accommodation coefficient; $B_{i,p}$ is the Kinetic oriented
 173 transition-regime diffusion correction factor. The detailed calculations and assumptions of the
 174 above parameters are in SI.

175 The square bracket term in Eq. 4 represents the driving force of OOM condensation, i.e. the over-
 176 saturated concentration of the condensable vapor. C_i^v and C_i^0 are the actual vapor concentration and
 177 saturation vapor concentration, respectively. $a_{i,p}$ is the particulate-phase activity, and $a_{i,p} = X_{i,p} \gamma_{i,p}$
 178 $K_{i,p}$, where $\gamma_{i,p}$ is the mass-based activity coefficient in the organic condensed phase, which is
 179 assumed to be 1. $K_{i,p}$ is the Kelvin coefficient, which is estimated by $K_{i,p} = 10^{d_{pK}/d_p}$. Assuming a
 180 particle surface tension of $0.034 \text{ N} \cdot \text{m}^{-1}$,^{47, 48} and a mean molar weight of $250 \text{ g} \cdot \text{mol}^{-1}$ at 300 K,
 181 the corresponding Kelvin diameter (d_{pK}) is 5 nm. $X_{i,p}$ is the mass fraction of OOM_i at the particle
 182 size of d_p . In this study, $X_{i,p}$ changes along with the growth of a new particle, and it was estimated
 183 by simulating the cumulative mass of OOM_i in the particulate phase based on the mass flux of
 184 OOM_i condensation.

185 The condensation growth rates contributed by H_2SO_4 and $(\text{H}_2\text{SO}_4)_n(\text{DMA})_n$ are calculated
 186 following the method reported by Stolzenburg et al.,⁹ which includes the effect of van der Waals
 187 forces in vapour condensation. The details of the calculation are supplemented in the SI.

188 It should be noted that the theoretical GR in Eq. 4 has a corresponding relationship with particle
 189 sizes (d_p) and time (t). For a particle size bin, e.g. 1.5-3 nm, the period used to calculate the

190 theoretical $GR_{1.5-3}$ is consistent with that of observed $GR_{1.5-3}$ with log-normal distribution function
191 method.⁴⁹ The time windows used for GR calculations are the exact periods when NPF events
192 occur and develop. The criteria to identified NPF events and the calculation of observed GR is
193 consistent with a previous study.¹⁰

194 3 Results and discussion

195 3.1 OOMs observed in urban Beijing

196 An overall of ~1100 organic formulas were identified within the m/z range of 200-450 Th for each
197 season. The observed OOMs species in four seasons are generally similar in terms of the total
198 number of identified molecules and the proportion of non-nitrogen OOM molecules (Fig. S1). The
199 mass-defect plot of the averaged OOMs in spring is shown in Fig. 1 as an example.

200 Several groups of homologues were observed in both non-nitrogen OOMs and nitrogen-containing
201 OOMs. Points lined on the direction of $+CH_2$ are homologues, differed by an integral number of
202 $-CH_2-$ group. The multiple series of homologous oxidation products are likely a result of the co-
203 existed homologues precursors in urban areas, e.g. benzene, toluene, and xylene.⁵⁰

204 =====

205 Place Figure 1 Here

206 =====

207 Another important feature of OOMs in Beijing is the higher proportion of nitrogen-containing
208 OOMs in comparison to that of forested area (Figs. S2 & S3). In our measurements, ~30% of the
209 identified molecules are non-nitrogen OOMs and ~70% are nitrogen-containing OOMs. Among
210 them, 35-52% of the total concentration of nitrogen-containing OOMs are OOMs with more than
211 one nitrogen atom. The termination of auto-oxidation chain propagation by NO_x results in carbonyl
212 (RO_2+NO reaction) or nitrate functional group (RO_2+NO_x reaction).¹⁴ Multi-generation oxidation
213 of RO_2+NO_x may explain the formation of OOMs with more than one nitrogen atom. Nitrate
214 radical (NO_3) oxidation may also play a role. In addition to the high proportion of nitrogen-

215 containing OOMs, strong auto-oxidation processes with more than two consecutive oxygen
216 additions are rarely observed; this is consistent with NO_x reactions shortening RO_2 lifetimes in the
217 urban environment.

218 Although the OOM speciation in each season is quite similar, the total concentration of OOMs in
219 Beijing shows clear seasonal variations while falling within a similar range to concentrations
220 observed elsewhere. As shown in Fig. 2, the seasonal median concentration of total OOMs in
221 Beijing is the highest in summer ($4.0 \times 10^8 \text{ cm}^{-3}$), followed by spring ($3.3 \times 10^8 \text{ cm}^{-3}$), autumn (1.1
222 $\times 10^8 \text{ cm}^{-3}$), and winter ($3.2 \times 10^7 \text{ cm}^{-3}$). Similarly, the concentration of condensable OOMs in
223 urban Beijing, accounting for 17-35% of total OOM concentration, also has the same seasonal
224 variations. The high total OOM concentration in summer is most likely related to the enhanced
225 photochemistry due to stronger solar radiation and higher temperature (Fig. S8). From a global
226 perspective, OOMs measured by nitrate CI-APi-ToF range from 10^7 to 10^9 cm^{-3} . It should be noted
227 that, nitrated phenols can contribute to 14-78% of the total OOM concentration in urban Beijing,
228 which is higher than their contribution of $\sim 16\%$ in those remote forest areas.⁵¹ However, as nitrated
229 phenols are too volatile to have significant contribution to particle growth, they are excluded in
230 the following discussion. The information of identified nitrated phenol peaks is supplemented in
231 the SI.

232 =====

233 Place Figure 2 Here

234 =====

235 The concentrations of both non-nitrogen OOMs and nitrogen-containing OOMs increase along
236 with solar radiation (UVB) in the morning, which also coincides with particle growth processes.
237 Such patterns are consistent during all four seasons (Fig. S4). The averaged diurnal variations in
238 summer are shown in Fig. 3 as an example. Sulfuric acid starts to rise at around 5 am (local time)
239 and reaches a peak at around 10 am. Both non-nitrogen OOMs and nitrogen-containing OOMs
240 start to rise at 6 am; nitrogen-containing OOMs reach a peak at 11 am and non-nitrate OOMs

241 reached the peak one hour later. In daytime, OOMs increase along with the increase of solar
242 radiation due to photochemistry, and decrease more slowly than sulfuric acid. This slower decrease
243 of OOMs can be attributed to the continuing formation via ozone oxidation and the lower removal
244 efficiency via condensation onto particles compared to sulfuric acid; it will be discussed in Section
245 3.2. In addition, the secondary peak of nitrogen-containing OOMs at around 10 pm may be due to
246 the onset of NO_3 chemistry at nighttime.

247

=====

248

Place Figure 3 Here

249

=====

250 3.2 Particle growth contributed by OOMs condensation

251 The growth rates of 1.5-25 nm particles driven by condensation of OOMs are approximately 0.25-
252 3.4 $\text{nm}\cdot\text{h}^{-1}$, and show clear seasonal variations. As shown in Fig. 4, the condensation rates of
253 OOMs, i.e., the sum of non-nitrate and nitrate OOMs, are the highest in summer, ranging from
254 1.7-3.4 $\text{nm}\cdot\text{h}^{-1}$, followed by 0.7-2.0 $\text{nm}\cdot\text{h}^{-1}$ of spring, 0.96-1.48 $\text{nm}\cdot\text{h}^{-1}$ of autumn, and 0.25-0.3
255 $\text{nm}\cdot\text{h}^{-1}$ of winter. For particles in the same size range, the growth rates contributed by OOMs in
256 summer can be 6.5-13 times of those in winter. In addition, the contribution of nitrogen-containing
257 OOMs to particle growth is larger than that of non-nitrate HOMs in all size ranges and all four
258 seasons, with calculated growth rates of 0.2-2.6 $\text{nm}\cdot\text{h}^{-1}$ and 0.05-0.9 $\text{nm}\cdot\text{h}^{-1}$, respectively.
259 Moreover, particle growth contributed by sulfuric acid and its clusters, nitrogen-containing OOMs,
260 and non-nitrate OOMs have different size dependencies. With increasing particle diameter, the
261 contribution by OOMs shows a flat or increasing trend, while the contribution by sulfuric acid
262 considerably decreases. Taking summer as an example, from 1.5-3 nm to 15-25 nm, the
263 contribution of sulfuric acid and its clusters decreased from 2.95 to 1.18 $\text{nm}\cdot\text{h}^{-1}$, nitrogen-
264 containing OOMs increased from 1.03 to 2.76 $\text{nm}\cdot\text{h}^{-1}$, and non-nitrogen OOMs increased from
265 0.78 to 1.0 $\text{nm}\cdot\text{h}^{-1}$.

266

=====

267

Place Figure 4 Here

268

=====

269 The difference of condensation growth rates during four seasons is primarily due to the strong
270 seasonal variation of the condensable OOM concentration. As shown in Figs. 5a & b, the
271 condensation growth rate contributed by OOMs correlated well (Pearson correlation coefficient of
272 0.95-0.99 in linear scale) with the condensable OOM concentration for each particle size range
273 (Fig. S10). This is because that the volatility distributions of both non-nitrogen OOMs and
274 nitrogen-containing OOMs are relatively stable in four seasons (Fig. S11). Therefore, for any given
275 particle size, the driving force of particle growth is almost solely determined by the gas-phase
276 concentration of condensable OOMs. As shown in Fig. 6, the concentrations of both non-nitrogen
277 OOMs and nitrogen-containing OOMs vary by almost one order of magnitude, leading to similar
278 extent of variations in particle growth rate.

279

=====

280

Place Figure 5 Here

281

=====

282

=====

283

Place Figure 6 Here

284

=====

285 In addition to vapor concentration, the effectiveness of vapor condensation also influences the
286 contribution of a certain gaseous species to particle growth. Here, we use the term "growth
287 promotion efficiency (GPE)", defined as the resulted growth rate per unit concentration (e.g., $\text{nm}\cdot\text{h}^{-1}$
288 $1/\text{cm}^{-3}$). From this perspective, sulfuric acid and its clusters are the most efficient condensable
289 vapors, followed by non-nitrate OOMs and nitrogen-containing OOMs (Fig. 5c). The GPE is

290 essentially determined by the bulk volatility and the averaged molar mass of different species.⁵²
291 Sulfuric acid, although with a small molar mass, can hardly evaporate, so that it has the highest
292 efficiency. For the case of OOMs, non-nitrogen OOMs and nitrogen-containing OOMs have
293 similar molar mass, but the bulk volatility of non-nitrate OOMs is lower than that of nitrogen-
294 containing OOMs (Figs. S12-13). Therefore, the GPE of non-nitrate OOMs is higher than that of
295 nitrogen-containing OOMs, and with the same vapor concentration, non-nitrate OOMs lead to
296 larger growth rates than nitrogen-containing OOMs do.

297 Taking both the vapor concentration and GPE into consideration, we could explain the difference
298 of condensation growth rate between non-nitrogen OOMs and nitrogen-containing OOMs. The
299 total concentrations of condensable nitrogen-containing OOMs are 2.67-3.37 times of non-
300 nitrogen OOMs (Fig. 6), but the lower GPE of nitrogen-containing OOMs (Fig. 5c) partially offsets
301 the higher concentration. As a whole, the growth rates contributed by nitrogen-containing OOMs
302 are 1-2.33 times of that by non-nitrogen OOMs.

303 As also shown in Fig.5c, the GPE of different species varies differently across particle sizes. For
304 sulfuric acid and its clusters, their GPE decrease with increased particle sizes. This is because
305 condensation of same amount of sulfuric acid causes smaller increases in diameters for large
306 particles than for small particles. Similar effect is expected for OOMs, however, OOM
307 condensation is also strongly influenced by the Kelvin effect. More OOMs can condense as
308 particles grow larger and the Kelvin effect diminishes, which compensates the size effect and leads
309 to an overall weak size dependence.

310 **4 Atmospheric implications**

311 This study reports the seasonal characteristics of OOMs in an urban environment and supplements
312 for the global scheme¹⁴, which helps to better understand OOMs formation in different
313 environments. We found an atmospheric relevance between solar radiation and OOMs
314 concentration in a seasonal scale: with the increase of solar radiation intensity (from winter to
315 summer), the concentration of day-time OOMs rises rapidly first and then increases slowly (Fig.
316 S8a). The strength of solar radiation is correlated with OH radical concentration and atmospheric
317 temperature.^{53, 54} Since the concentration of precursors in winter is not significantly lower than
318 other seasons (Fig. S9a) and its loss through condensation sink is not higher than other seasons

319 (Fig. S9c), the lower oxidant concentrations (e.g. OH radical) and lower temperature may
320 substantially limit the formation of OOMs in wintertime. Compared with the seasonal variations
321 of VOC precursors and condensation sink, photochemical strength seems to play a more important
322 role in the seasonal variation of OOM concentrations based on its better consistency (Fig. S9).
323 This correlation may also exist in other mid-latitude urban and forested environments that with
324 distinct seasonal changes.

325 In urban Beijing, sulfuric acid and its clusters can explain a significant fraction of the observed
326 growth of 1.5-3 nm particles¹⁰, and OOMs have a significant contribution to the growth of 3-25
327 nm particles (Fig. 7). The higher contribution of sulfuric acid and its clusters in 1.5-3 nm particles
328 can be explained by its high GPE at this size range (Fig. 5c). In addition, the concentration of
329 sulfuric acid rises early and steeply during the time window when the initial growth of new
330 particles occurs (Fig. 3). For larger (3-25 nm) particles, despite the lower GPE of OOMs than
331 sulfuric acid, OOMs start to dominate particle growth, due to the much higher vapor concentration
332 and the increasingly weaker Kelvin effect. Overall, the contributions by sulfuric acid and OOMs
333 can explain a major fraction of the observed growth rate in all size ranges of 1.5-25 nm.

334 It was reported that NPF events in Beijing can contribute to the surface area concentration and
335 mass concentration of atmospheric aerosols (mainly governed by particles larger than 100 nm),
336 especially under conditions with a fast growth of newly formed particles.^{5, 55} Although this study
337 focused on the growth of nanoparticles, the condensation of OOMs is expected to contribute
338 significantly to the growth of large particles as well.

339 However, winter is an exception. A clear discrepancy remains between simulated and observed
340 growth rate in 3-25 nm particles, which may be an indication of some missing contributors. One
341 plausible cause is the underestimation of condensable OOM concentration at low temperatures.
342 Nitrate CI-APi-ToF is less sensitive to moderately oxidized molecules,⁵⁶ while with the decrease
343 of temperature, some moderately oxidized organics may become condensable and contribute to
344 particle growth.⁵⁷ Meanwhile, particle-phase processes could promote particle growth through
345 lowering the volatility of particles-phase organics,⁵⁸ including condensed phase reactions and
346 oligomerization.⁵⁹ Another possible reasons is the missing contribution by HNO₃ and NH₃ at low
347 temperatures, as suggested by a recent chamber study.⁶⁰

348

=====

349

Place Figure 7 Here

350

=====

351 In some cases, the simulated particle growth rates are higher than observed values (Fig. 7). This
352 may arise from two sources of uncertainties. One is the uncertainty in growth rate simulation,
353 which mainly comes from the estimation of OOMs volatility (Fig. S5). The other is from the
354 calculation of observed growth rate. As illustrated by Deng et al.,¹⁰ growth rates calculated using
355 the appearance time method and the log-normal distribution method can sometimes differ by a
356 factor of 2. We investigated the results of appearance time growth rate and corrected the influence
357 by particle coagulation⁶¹ (Figs. S14-15). The ratios of simulated growth rate over observed growth
358 rate are lower when particle growth rates are calculated with the appearance time method. However,
359 our findings on the important role of OOMs in explaining the growth rate of 3-25 nm particles
360 remain valid. In addition, the sum of OOMs, sulfuric acid and its clusters is still far from explaining
361 the observed growth rate (appearance time method) in winter.

362

363 **Supporting Information**

364 It includes the detailed information on the calculation method of simulated growth rate;
365 comparison of observed oxygenated organic molecules in all of the four seasons using Venn plot
366 (Figure S1); the proportion of carbon-resolved non-nitrogen OOMs and nitrogen-containing
367 OOMs in Beijing and other atmospheric environment (Figures S2 & S3); the diurnal variation of
368 sulfuric acid, non-nitrogen OOMs and nitrogen-containing OOMs in all of the four seasons (Figure
369 S4); the comparison of two volatility parameterization methods (Figure S5); the volatility
370 distribution of observed oxygenated organic molecules (OOMs) in Beijing for four seasons (Figure
371 S6); the relations between H to C ratio and O to C ratio of known OOMs peaklists from laboratory

372 studies (Figure S7); the relations of condensable oxygenated organic molecules with temperature
373 and solar radiation (Figure S8); seasonal variations of volatile organic compounds, temperature
374 and solar radiation, and condensation sink (CS) in Beijing (Figure S9); relations of size-resolved
375 simulated growth rate and condensing vapor concentration (Figure S10); the relations of driving
376 force and OOMs concentration (Figure S11); the growth efficiency of different kinds of
377 condensing vapors (Figure S12); the volatility distribution of non-nitrogen OOMs and nitrogen-
378 containing OOMs (Figure S13); the relations of simulated GR and observed GR (appearance time
379 method) for different particle size bins (Figures S14 & S15); the condensation growth rate
380 contributed by OOMs with different volatilities for 1.5-25 nm particles (Figure S16); six examples
381 for peak identification in tofTool (Figure S17); concentration of total OOMs in new particle
382 formation (NPF) days and non-NPF days. (Figure S18); and the identified nitrated phenol peaks
383 during the measurement (Table S1).

384

385 AUTHOR INFORMATION

386 **Corresponding Author**

387 * **Jingkun Jiang** – State Key Joint Laboratory of Environment Simulation and Pollution Control,
388 School of Environment, Tsinghua University, 100084 Beijing, China; Email:
389 jjangjk@tsinghua.edu.cn

390 * **Chao Yan** – Institute for Atmospheric and Earth System Research Physics, Faculty of Science,
391 University of Helsinki, 00014 Helsinki, Finland; Aerosol and Haze Laboratory, Beijing Advanced
392 Innovation Center for Soft Matter Science and Engineering, Beijing University of Chemical
393 Technology, 100029 Beijing, China; Email: chao.yan@helsinki.fi

394 **Notes**

395 The authors declare no competing financial interest.

396 **Acknowledgement**

397 Financial support from the National Science Foundation of China (92044301, 21876094, and
398 41730106) and Samsung PM_{2.5} SRP are acknowledged.

399

400 **Reference**

- 401 1. Kulmala, M.; Vehkamäki, H.; Petäjä, T.; Dal Maso, M.; Lauri, A.; Kerminen, V. M.;
402 Birmili, W.; McMurry, P. H., Formation and growth rates of ultrafine atmospheric particles: A
403 review of observations. *Journal of Aerosol Science* **2004**, *35*, 143-176.
- 404 2. Kerminen, V. M.; Lihavainen, H.; Komppula, M.; Viisanen, Y.; Kulmala, M., Direct
405 observational evidence linking atmospheric aerosol formation and cloud droplet activation.
406 *Geophysical Research Letters* **2005**, *32*, 1-4.
- 407 3. Merikanto, J.; Spracklen, D. V.; Mann, G. W.; Pickering, S. J.; Carslaw, K. S., Impact of
408 nucleation on global CCN. *Atmospheric Chemistry and Physics* **2009**, *9*, 8601-8616.
- 409 4. McMurry, P. H.; Friedlander, S. K., New particle formation in the presence of an aerosol.
410 *Atmospheric Environment* **1979**, *13*, (12), 1635-1651.
- 411 5. Guo, S.; Hu, M.; Zamora, M. L.; Peng, J.; Shang, D.; Zheng, J.; Du, Z.; Wu, Z.; Shao,
412 M.; Zeng, L.; Molina, M. J.; Zhang, R., Elucidating severe urban haze formation in China.
413 *Proceedings of the National Academy of Sciences of the United States of America* **2014**, *111*,
414 17373-17378.
- 415 6. Cai, R. L.; Yang, D. S.; Fu, Y. Y.; Wang, X.; Li, X. X.; Ma, Y.; Hao, J. M.; Zheng, J.;
416 Jiang, J. K., Aerosol surface area concentration: a governing factor in new particle formation in
417 Beijing. *Atmospheric Chemistry and Physics* **2017**, *17*, (20), 12327-12340.
- 418 7. Kulmala, M.; Kerminen, V. M.; Petäjä, T.; Ding, A. J.; Wang, L., Atmospheric gas-to-
419 particle conversion: Why NPF events are observed in megacities? *Faraday Discussions* **2017**,
420 *200*, 271-288.

- 421 8. McMurry, P. H.; Fink, M.; Sakurai, H.; Stolzenburg, M. R.; Mauldin, I. L.; Smith, J.;
422 Eisele, F.; Moore, K.; Sjostedt, S.; Tanner, D.; Huey, L. G.; Nowak, J. B.; Edgerton, E.; Voisin,
423 D., A criterion for new particle formation in the sulfur-rich Atlanta atmosphere. *Journal of*
424 *Geophysical Research Atmospheres* **2005**, *110*, 1-10.
- 425 9. Stolzenburg, D.; Simon, M.; Ranjithkumar, A.; Kürten, A.; Lehtipalo, K.; Gordon, H.;
426 Ehrhart, S.; Finkenzeller, H.; Pichelstorfer, L.; Nieminen, T.; He, X.-C.; Brilke, S.; Xiao, M.;
427 Amorim, A.; Baalbaki, R.; Baccharini, A.; Beck, L.; Bräkling, S.; Caudillo Murillo, L.; Chen, D.;
428 Chu, B.; Dada, L.; Dias, A.; Dommen, J.; Duplissy, J.; El Haddad, I.; Fischer, L.; Gonzalez
429 Carracedo, L.; Heinritzi, M.; Kim, C.; Koenig, T. K.; Kong, W.; Lamkaddam, H.; Lee, C. P.;
430 Leiminger, M.; Li, Z.; Makhmutov, V.; Manninen, H. E.; Marie, G.; Marten, R.; Müller, T.; Nie,
431 W.; Partoll, E.; Petäjä, T.; Pfeifer, J.; Philippov, M.; Rissanen, M. P.; Rörup, B.; Schobesberger,
432 S.; Schuchmann, S.; Shen, J.; Sipilä, M.; Steiner, G.; Stozhkov, Y.; Tauber, C.; Tham, Y. J.;
433 Tomé, A.; Vazquez-Pufleau, M.; Wagner, A. C.; Wang, M.; Wang, Y.; Weber, S. K.; Wimmer,
434 D.; Wlasits, P. J.; Wu, Y.; Ye, Q.; Zauner-Wieczorek, M.; Baltensperger, U.; Carslaw, K. S.;
435 Curtius, J.; Donahue, N. M.; Flagan, R. C.; Hansel, A.; Kulmala, M.; Lelieveld, J.; Volkamer, R.;
436 Kirkby, J.; Winkler, P. M., Enhanced growth rate of atmospheric particles from sulfuric acid.
437 *Atmospheric Chemistry and Physics* **2020**, *20*, (12), 7359-7372.
- 438 10. Deng, C.; Fu, Y.; Dada, L.; Yan, C.; Cai, R.; Yang, D.; Zhou, Y.; Yin, R.; Lu, Y.; Li, X.;
439 Qiao, X.; Fan, X.; Nie, W.; Kontkanen, J.; Kangasluoma, J.; Chu, B.; Ding, A.; Kerminen, V.
440 M.; Paasonen, P.; Worsnop, D. R.; Bianchi, F.; Liu, Y.; Zheng, J.; Wang, L.; Kulmala, M.; Jiang,
441 J., Seasonal Characteristics of New Particle Formation and Growth in Urban Beijing.
442 *Environmental Science & Technology* **2020**, *54*, (14), 8547-8557.
- 443 11. Stolzenburg, M. R.; McMurry, P. H.; Sakurai, H.; Smith, J. N.; Mauldin, R. L.; Eisele, F.
444 L.; Clement, C. F., Growth rates of freshly nucleated atmospheric particles in Atlanta. *Journal of*
445 *Geophysical Research Atmospheres* **2005**, *110*, 1-10.
- 446 12. Riipinen, I.; Yli-Juuti, T.; Pierce, J. R.; Petäjä, T.; Worsnop, D. R.; Kulmala, M.;
447 Donahue, N. M., The contribution of organics to atmospheric nanoparticle growth. *Nature*
448 *Geoscience* **2012**, *5*, 453-458.
- 449 13. Mohr, C.; Thornton, J. A.; Heitto, A.; Lopez-Hilfiker, F. D.; Lutz, A.; Riipinen, I.; Hong,
450 J.; Donahue, N. M.; Hallquist, M.; Petäjä, T.; Kulmala, M.; Yli-Juuti, T., Molecular
451 identification of organic vapors driving atmospheric nanoparticle growth. *Nature*
452 *Communications* **2019**, *10*, 1-7.

- 453 14. Bianchi, F.; Kurten, T.; Riva, M.; Mohr, C.; Rissanen, M. P.; Roldin, P.; Berndt, T.;
454 Crounse, J. D.; Wennberg, P. O.; Mentel, T. F.; Wildt, J.; Junninen, H.; Jokinen, T.; Kulmala,
455 M.; Worsnop, D. R.; Thornton, J. A.; Donahue, N.; Kjaergaard, H. G.; Ehn, M., Highly
456 Oxygenated Organic Molecules (HOM) from Gas-Phase Autoxidation Involving Peroxy
457 Radicals: A Key Contributor to Atmospheric Aerosol. *Chemical Reviews* **2019**, *119*, (6), 3472-
458 3509.
- 459 15. Molteni, U.; Bianchi, F.; Klein, F.; El Haddad, I.; Frege, C.; Rossi, M. J.; Dommen, J.;
460 Baltensperger, U., Formation of highly oxygenated organic molecules from aromatic
461 compounds. *Atmospheric Chemistry and Physics* **2018**, *18*, 1909-1921.
- 462 16. Jokinen, T.; Kausiala, O.; Garmash, O.; Peräkylä, O.; Junninen, H.; Schobesberger, S.;
463 Yan, C.; Sipilä, M.; Rissanen, M. P., Production of highly oxidized organic compounds from
464 ozonolysis of β -caryophyllene: Laboratory and field measurements. *Boreal Environment*
465 *Research* **2016**, *21*, 262-273.
- 466 17. Lee, A.; Goldstein, A. H.; Kroll, J. H.; Ng, N. L.; Varutbangkul, V.; Flagan, R. C.;
467 Seinfeld, J. H., Gas-phase products and secondary aerosol yields from the photooxidation of 16
468 different terpenes. *Journal of Geophysical Research Atmospheres* **2006**, *111*, 1-25.
- 469 18. Ehn, M.; Thornton, J. A.; Kleist, E.; Sipilä, M.; Junninen, H.; Pullinen, I.; Springer, M.;
470 Rubach, F.; Tillmann, R.; Lee, B.; Lopez-Hilfiker, F.; Andres, S.; Acir, I. H.; Rissanen, M.;
471 Jokinen, T.; Schobesberger, S.; Kangasluoma, J.; Kontkanen, J.; Nieminen, T.; Kurten, T.;
472 Nielsen, L. B.; Jorgensen, S.; Kjaergaard, H. G.; Canagaratna, M.; Maso, M. D.; Berndt, T.;
473 Petaja, T.; Wahner, A.; Kerminen, V. M.; Kulmala, M.; Worsnop, D. R.; Wildt, J.; Mentel, T. F.,
474 A large source of low-volatility secondary organic aerosol. *Nature* **2014**, *506*, 476-479.
- 475 19. Tröstl, J.; Chuang, W. K.; Gordon, H.; Heinritzi, M.; Yan, C.; Molteni, U.; Ahlm, L.;
476 Frege, C.; Bianchi, F.; Wagner, R.; Simon, M.; Lehtipalo, K.; Williamson, C.; Craven, J. S.;
477 Duplissy, J.; Adamov, A.; Almeida, J.; Bernhammer, A. K.; Breitenlechner, M.; Brilke, S.; Dias,
478 A.; Ehrhart, S.; Flagan, R. C.; Franchin, A.; Fuchs, C.; Guida, R.; Gysel, M.; Hansel, A.; Hoyle,
479 C. R.; Jokinen, T.; Junninen, H.; Kangasluoma, J.; Keskinen, H.; Kim, J.; Krapf, M.; Kurten, A.;
480 Laaksonen, A.; Lawler, M.; Leiminger, M.; Mathot, S.; Mohler, O.; Nieminen, T.; Onnela, A.;
481 Petaja, T.; Piel, F. M.; Miettinen, P.; Rissanen, M. P.; Rondo, L.; Sarnela, N.; Schobesberger, S.;
482 Sengupta, K.; Sipilä, M.; Smith, J. N.; Steiner, G.; Tome, A.; Virtanen, A.; Wagner, A. C.;
483 Weingartner, E.; Wimmer, D.; Winkler, P. M.; Ye, P.; Carslaw, K. S.; Curtius, J.; Dommen, J.;
484 Kirkby, J.; Kulmala, M.; Riipinen, I.; Worsnop, D. R.; Donahue, N. M.; Baltensperger, U., The

- 485 role of low-volatility organic compounds in initial particle growth in the atmosphere. *Nature*
486 **2016**, *533*, (7604), 527-31.
- 487 20. Kirkby, J.; Duplissy, J.; Sengupta, K.; Frege, C.; Gordon, H.; Williamson, C.; Heinritzi,
488 M.; Simon, M.; Yan, C.; Almeida, J.; Trostl, J.; Nieminen, T.; Ortega, I. K.; Wagner, R.;
489 Adamov, A.; Amorim, A.; Bernhammer, A. K.; Bianchi, F.; Breitenlechner, M.; Brilke, S.; Chen,
490 X.; Craven, J.; Dias, A.; Ehrhart, S.; Flagan, R. C.; Franchin, A.; Fuchs, C.; Guida, R.; Hakala,
491 J.; Hoyle, C. R.; Jokinen, T.; Junninen, H.; Kangasluoma, J.; Kim, J.; Krapf, M.; Kurten, A.;
492 Laaksonen, A.; Lehtipalo, K.; Makhmutov, V.; Mathot, S.; Molteni, U.; Onnela, A.; Perakyla,
493 O.; Piel, F.; Petaja, T.; Praplan, A. P.; Pringle, K.; Rap, A.; Richards, N. A.; Riipinen, I.;
494 Rissanen, M. P.; Rondo, L.; Sarnela, N.; Schobesberger, S.; Scott, C. E.; Seinfeld, J. H.; Sipila,
495 M.; Steiner, G.; Stozhkov, Y.; Stratmann, F.; Tome, A.; Virtanen, A.; Vogel, A. L.; Wagner, A.
496 C.; Wagner, P. E.; Weingartner, E.; Wimmer, D.; Winkler, P. M.; Ye, P.; Zhang, X.; Hansel, A.;
497 Dommen, J.; Donahue, N. M.; Worsnop, D. R.; Baltensperger, U.; Kulmala, M.; Carslaw, K. S.;
498 Curtius, J., Ion-induced nucleation of pure biogenic particles. *Nature* **2016**, *533*, 521-526.
- 499 21. Schobesberger, S.; Junninen, H.; Bianchi, F.; Lonn, G.; Ehn, M.; Lehtipalo, K.; Dommen,
500 J.; Ehrhart, S.; Ortega, I. K.; Franchin, A.; Nieminen, T.; Riccobono, F.; Hutterli, M.; Duplissy,
501 J.; Almeida, J.; Amorim, A.; Breitenlechner, M.; Downard, A. J.; Dunne, E. M.; Flagan, R. C.;
502 Kajos, M.; Keskinen, H.; Kirkby, J.; Kupc, A.; Kurten, A.; Kurten, T.; Laaksonen, A.; Mathot,
503 S.; Onnela, A.; Praplan, A. P.; Rondo, L.; Santos, F. D.; Schallhart, S.; Schnitzhofer, R.; Sipila,
504 M.; Tome, A.; Tsagkogeorgas, G.; Vehkamaki, H.; Wimmer, D.; Baltensperger, U.; Carslaw, K.
505 S.; Curtius, J.; Hansel, A.; Petaja, T.; Kulmala, M.; Donahue, N. M.; Worsnop, D. R., Molecular
506 understanding of atmospheric particle formation from sulfuric acid and large oxidized organic
507 molecules. *Proceedings of the National Academy of Sciences of the United States of America*
508 **2013**, *110*, (43), 17223-8.
- 509 22. Zhao, J.; Ortega, J.; Chen, M.; McMurry, P. H.; Smith, J. N., Dependence of particle
510 nucleation and growth on high molecular weight gas phase products during ozonolysis of α -
511 pinene. *Atmospheric Chemistry and Physics* **2013**, *13*, 7631-7644.
- 512 23. Crounse, J. D.; Nielsen, L. B.; Jørgensen, S.; Kjaergaard, H. G.; Wennberg, P. O.,
513 Autoxidation of organic compounds in the atmosphere. *Journal of Physical Chemistry Letters*
514 **2013**, *4*, 3513-3520.

- 515 24. Wang, S.; Riva, M.; Yan, C.; Ehn, M.; Wang, L., Primary Formation of Highly Oxidized
516 Multifunctional Products in the OH-Initiated Oxidation of Isoprene: A Combined Theoretical
517 and Experimental Study. *Environmental Science and Technology* **2018**, *52*, 12255-12264.
- 518 25. Krechmer, J. E.; Coggon, M. M.; Massoli, P.; Nguyen, T. B.; Crouse, J. D.; Hu, W.;
519 Day, D. A.; Tyndall, G. S.; Henze, D. K.; Rivera-Rios, J. C.; Nowak, J. B.; Kimmel, J. R.;
520 Mauldin, R. L., 3rd; Stark, H.; Jayne, J. T.; Sipila, M.; Junninen, H.; Clair, J. M.; Zhang, X.;
521 Feiner, P. A.; Zhang, L.; Miller, D. O.; Brune, W. H.; Keutsch, F. N.; Wennberg, P. O.; Seinfeld,
522 J. H.; Worsnop, D. R.; Jimenez, J. L.; Canagaratna, M. R., Formation of Low Volatility Organic
523 Compounds and Secondary Organic Aerosol from Isoprene Hydroxyhydroperoxide Low-NO
524 Oxidation. *Environmental Science & Technology* **2015**, *49*, (17), 10330-9.
- 525 26. Garmash, O.; Rissanen, M. P.; Pullinen, I.; Schmitt, S.; Kausiala, O.; Tillmann, R.; Zhao,
526 D.; Percival, C.; Bannan, T. J.; Priestley, M.; Hallquist, A. M.; Kleist, E.; Kiendler-Scharr, A.;
527 Hallquist, M.; Berndt, T.; McFiggans, G.; Wildt, J.; Mentel, T. F.; Ehn, M., Multi-generation OH
528 oxidation as a source for highly oxygenated organic molecules from aromatics. *Atmospheric*
529 *Chemistry and Physics* **2020**, *20*, 515-537.
- 530 27. Rissanen, M. P., NO₂ Suppression of Autoxidation-Inhibition of Gas-Phase Highly
531 Oxidized Dimer Product Formation. *ACS Earth and Space Chemistry* **2018**, *2*, 1211-1219.
- 532 28. Xiao, M.; Hoyle, C.; Dada, L.; Stolzenburg, D.; Kürten, A.; Wang, M.; Lamkaddam, H.;
533 Garmash, O.; Mentler, B.; Molteni, U.; Baccharini, A.; Simon, M.; He, X.-C.; Lehtipalo, K.;
534 Ahonen, L.; Baalbaki, R.; Bauer, P.; Beck, L.; Bell, D.; Bianchi, F.; Brilke, S.; Chen, D.; Chiu,
535 R.; Dias, A.; Duplissy, J.; Finkenzeller, H.; Gordon, H.; Hofbauer, V.; Kim, C.; Koenig, T.;
536 Lampilahti, J.; Lee, C. P.; Li, Z.; Mai, H.; Makhmutov, V.; Manninen, H.; Marten, R.; Mathot,
537 S.; Mauldin, R.; Nie, W.; Onnela, A.; Partoll, E.; Petäjä, T.; Pfeifer, J.; Pospisilova, V.;
538 Quéléver, L.; Rissanen, M.; Schobesberger, S.; Schuchmann, S.; Stozhkov, Y.; Tauber, C.;
539 Tham, Y. J.; Tomé, A.; Vazquez-Pufleau, M.; Wagner, A.; Wanger, R.; Wang, Y.; Weitz, L.;
540 Wimmer, D.; Wu, Y.; Yan, C.; Ye, P.; Ye, Q.; Zha, Q.; Zhou, X.; Amorim, A.; Carslaw, K.;
541 Curtius, J.; Hansel, A.; Volkamer, R.; Winkler, P.; Flagan, R.; Kulmala, M.; Worsnop, D.;
542 Kirkby, J.; Donahue, N.; Baltensperger, U.; El Haddad, I.; Dommen, J., The driving factors of
543 new particle formation and growth in the polluted boundary layer. *Atmospheric Chemistry and*
544 *Physics Discussions* **2021**, <https://doi.org/10.5194/acp-2020-1323>.
- 545 29. Yan, C.; Nie, W.; Vogel, A. L.; Dada, L.; Lehtipalo, K.; Stolzenburg, D.; Wagner, R.;
546 Rissanen, M. P.; Xiao, M.; Ahonen, L.; Fischer, L.; Rose, C.; Bianchi, F.; Gordon, H.; Simon,

- 547 M.; Heinritzi, M.; Garmash, O.; Roldin, P.; Dias, A.; Ye, P.; Hofbauer, V.; Amorim, A.; Bauer,
548 P. S.; Bergen, A.; Bernhammer, A. K.; Breitenlechner, M.; Brilke, S.; Buchholz, A.; Mazon, S.
549 B.; Canagaratna, M. R.; Chen, X.; Ding, A.; Dommen, J.; Draper, D. C.; Duplissy, J.; Frege, C.;
550 Heyn, C.; Guida, R.; Hakala, J.; Heikkinen, L.; Hoyle, C. R.; Jokinen, T.; Kangasluoma, J.;
551 Kirkby, J.; Kontkanen, J.; Kurten, A.; Lawler, M. J.; Mai, H.; Mathot, S.; Mauldin, R. L., 3rd;
552 Molteni, U.; Nichman, L.; Nieminen, T.; Nowak, J.; Ojdanic, A.; Onnela, A.; Pajunoja, A.;
553 Petaja, T.; Piel, F.; Quelever, L. L. J.; Sarnela, N.; Schallhart, S.; Sengupta, K.; Sipila, M.; Tome,
554 A.; Trostl, J.; Vaisanen, O.; Wagner, A. C.; Ylisirnio, A.; Zha, Q.; Baltensperger, U.; Carslaw,
555 K. S.; Curtius, J.; Flagan, R. C.; Hansel, A.; Riipinen, I.; Smith, J. N.; Virtanen, A.; Winkler, P.
556 M.; Donahue, N. M.; Kerminen, V. M.; Kulmala, M.; Ehn, M.; Worsnop, D. R., Size-dependent
557 influence of NO_x on the growth rates of organic aerosol particles. *Science Advances* **2020**, *6*,
558 (22), 4945.
- 559 30. Yan, C.; Yin, R. J.; Lu, Y. Q.; Dada, L. N.; Yang, D. S.; Fu, Y. Y.; Kontkanen, J.; Deng,
560 C. J.; Garmash, O.; Ruan, J. X.; Baalbaki, R.; Schervish, M.; Cai, R. L.; Bloss, M.; Chan, T.;
561 Chen, T. Z.; Chen, Q.; Chen, X. M.; Chen, Y.; Chu, B. W.; Dallenbach, K.; Foreback, B.; He, X.
562 C.; Heikkinen, L.; Jokinen, T.; Junninen, H.; Kangasluoma, J.; Kokkonen, T.; Kurppa, M.;
563 Lehtipalo, K.; Li, H. Y.; Li, H.; Li, X. X.; Liu, Y. L.; Ma, Q. X.; Paasonen, P.; Rantala, P.;
564 Pileci, R. E.; Rusanen, A.; Sarnela, N.; Simonen, P.; Wang, S. X.; Wang, W. G.; Wang, Y. H.;
565 Xue, M.; Yang, G.; Yao, L.; Zhou, Y.; Kujansuu, J.; Petaja, T.; Nie, W.; Ma, Y.; Ge, M. F.; He,
566 H.; Donahue, N. M.; Worsnop, D. R.; Kerminen, V. M.; Wang, L.; Liu, Y. C.; Zheng, J.; Kulma,
567 M.; Jiang, J. K.; Bianchi, F., The Synergistic Role of Sulfuric Acid, Bases, and Oxidized
568 Organics Governing New-Particle Formation in Beijing. *Geophysical Research Letters* **2021**, *48*,
569 (7), 91944.
- 570 31. Brean, J.; Harrison, R. M.; Shi, Z. B.; Beddows, D. C. S.; Acton, W. J. F.; Hewitt, C. N.;
571 Squires, F. A.; Lee, J., Observations of highly oxidized molecules and particle nucleation in the
572 atmosphere of Beijing. *Atmospheric Chemistry and Physics* **2019**, *19*, (23), 14933-14947.
- 573 32. Liu, Y.; Yan, C.; Feng, Z.; Zheng, F.; Fan, X.; Zhang, Y.; Li, C.; Zhou, Y.; Lin, Z.; Guo,
574 Y.; Zhang, Y.; Ma, L.; Zhou, W.; Liu, Z.; Dada, L.; Daellenbach, K.; Kontkanen, J.; Cai, R.;
575 Chan, T.; Kulmala, M., Continuous and comprehensive atmospheric observations in Beijing: a
576 station to understand the complex urban atmospheric environment. *Big Earth Data* **2020**, *4*, 295-
577 321.

- 578 33. Jokinen, T.; Sipilä, M.; Junninen, H.; Ehn, M.; Lönn, G.; Hakala, J.; Petäjä, T.; Mauldin,
579 R. L.; Kulmala, M.; Worsnop, D. R., Atmospheric sulphuric acid and neutral cluster
580 measurements using CI-API-TOF. *Atmospheric Chemistry and Physics* **2012**, *12*, 4117-4125.
- 581 34. Bertram, T. H.; Kimmel, J. R.; Crisp, T. A.; Ryder, O. S.; Yatavelli, R. L. N.; Thornton,
582 J. A.; Cubison, M. J.; Gonin, M.; Worsnop, D. R., A field-deployable, chemical ionization time-
583 of-flight mass spectrometer. *Atmospheric Measurement Techniques* **2011**, *4*, 1471-1479.
- 584 35. Heinritzi, M.; Simon, M.; Steiner, G.; Wagner, A. C.; Kürten, A.; Hansel, A.; Curtius, J.,
585 Characterization of the mass-dependent transmission efficiency of a CIMS. *Atmospheric*
586 *Measurement Techniques* **2016**, *9*, 1449-1460.
- 587 36. Cai, R.; Yan, C.; Yang, D.; Yin, R.; Lu, Y.; Deng, C.; Fu, Y.; Ruan, J.; Li, X.;
588 Kontkanen, J.; Zhang, Q.; Kangasluoma, J.; Ma, Y.; Hao, J.; Worsnop, D. R.; Bianchi, F.;
589 Paasonen, P.; Kerminen, V.-M.; Liu, Y.; Wang, L.; Zheng, J.; Kulmala, M.; Jiang, J., Sulfuric
590 acid-amine nucleation in urban Beijing. *Atmospheric Chemistry and Physics* **2021**, *21*, (4),
591 2457-2468.
- 592 37. Jiang, J.; Chen, M.; Kuang, C.; Attoui, M.; McMurry, P. H., Electrical Mobility
593 Spectrometer Using a Diethylene Glycol Condensation Particle Counter for Measurement of
594 Aerosol Size Distributions Down to 1 nm. *Aerosol Science and Technology* **2011**, *45*, (4), 510-
595 521.
- 596 38. Cai, R.; Chen, D.-R.; Hao, J.; Jiang, J., A miniature cylindrical differential mobility
597 analyzer for sub-3 nm particle sizing. *Journal of Aerosol Science* **2017**, *106*, 111-119.
- 598 39. Liu, J.; Jiang, J.; Zhang, Q.; Deng, J.; Hao, J., A spectrometer for measuring particle size
599 distributions in the range of 3 nm to 10 μm . *Frontiers of Environmental Science & Engineering*
600 **2016**, *10*, (1), 63-72.
- 601 40. Fu, Y.; Xue, M.; Cai, R.; Kangasluoma, J.; Jiang, J., Theoretical and experimental
602 analysis of the core sampling method: Reducing diffusional losses in aerosol sampling line.
603 *Aerosol Science and Technology* **2019**, *53*, 793-801.
- 604 41. Cai, R.; Chen, D. R.; Hao, J.; Jiang, J., A miniature cylindrical differential mobility
605 analyzer for sub-3 nm particle sizing. *Journal of Aerosol Science* **2017**, *106*, 111-119.
- 606 42. Junninen, H.; Ehn, M.; Petäjä; Luosujärvi, L.; Kotiaho, T.; Kostianen, R.; Rohner, U.;
607 Gonin, M.; Fuhrer, K.; Kulmala, M.; Worsnop, D. R., A high-resolution mass spectrometer to
608 measure atmospheric ion composition. *Atmospheric Measurement Techniques* **2010**, *3*, 1039-
609 1053.

- 610 43. Kurten, A.; Rondo, L.; Ehrhart, S.; Curtius, J., Calibration of a chemical ionization mass
611 spectrometer for the measurement of gaseous sulfuric acid. *Journal of Physical Chemistry A*
612 **2012**, *116*, 6375-6386.
- 613 44. Donahue, N. M.; Kroll, J. H.; Pandis, S. N.; Robinson, A. L., A two-dimensional
614 volatility basis set-Part 2: Diagnostics of organic-aerosol evolution. *Atmospheric Chemistry and*
615 *Physics* **2012**, *12*, 615-634.
- 616 45. Wang, M.; Chen, D.; Xiao, M.; Ye, Q.; Stolzenburg, D.; Hofbauer, V.; Ye, P.; Vogel, A.
617 L.; Mauldin, R. L., 3rd; Amorim, A.; Baccharini, A.; Baumgartner, B.; Brilke, S.; Dada, L.; Dias,
618 A.; Duplissy, J.; Finkenzeller, H.; Garmash, O.; He, X. C.; Hoyle, C. R.; Kim, C.; Kvashnin, A.;
619 Lehtipalo, K.; Fischer, L.; Molteni, U.; Petaja, T.; Pospisilova, V.; Quelever, L. L. J.; Rissanen,
620 M.; Simon, M.; Tauber, C.; Tome, A.; Wagner, A. C.; Weitz, L.; Volkamer, R.; Winkler, P. M.;
621 Kirkby, J.; Worsnop, D. R.; Kulmala, M.; Baltensperger, U.; Dommen, J.; El-Haddad, I.;
622 Donahue, N. M., Photo-oxidation of Aromatic Hydrocarbons Produces Low-Volatility Organic
623 Compounds. *Environmental Science & Technology* **2020**, *54*, (13), 7911-7921.
- 624 46. Seinfeld, J. H.; Pandis, S. N., *Atmospheric chemistry and physics: from air pollution to*
625 *climate change*. 3rd ed.; John Wiley & Sons, Inc.: Hoboken, New Jersey, 2016.
- 626 47. Fenelonov, V. B.; Kodenov, G. G.; Kostrovsky, V. G., On the dependence of surface
627 tension of liquids on the curvature of the liquid-vapor interface. *Journal of Physical Chemistry B*
628 **2001**, *105*, 1050-1055.
- 629 48. Yli-Juuti, T.; Barsanti, K.; Hildebrandt Ruiz, L.; Kieloaho, A. J.; Makkonen, U.; Petäjä,
630 T.; Ruuskanen, T.; Kulmala, M.; Riipinen, I., Model for acid-base chemistry in nanoparticle
631 growth (MABNAG). *Atmospheric Chemistry and Physics* **2013**, *13*, 12507-12524.
- 632 49. Kulmala, M.; Petäjä, T.; Nieminen, T.; Sipilä, M.; Manninen, H. E.; Lehtipalo, K.; Dal
633 Maso, M.; Aalto, P. P.; Junninen, H.; Paasonen, P.; Riipinen, I.; Lehtinen, K. E. J.; Laaksonen,
634 A.; Kerminen, V. M., Measurement of the nucleation of atmospheric aerosol particles. *Nature*
635 *Protocols* **2012**, *7*, 1651-1667.
- 636 50. Sheng, J.; Zhao, D.; Ding, D.; Li, X.; Huang, M.; Gao, Y.; Quan, J.; Zhang, Q.,
637 Characterizing the level, photochemical reactivity, emission, and source contribution of the
638 volatile organic compounds based on PTR-TOF-MS during winter haze period in Beijing, China.
639 *Atmospheric Research* **2018**, *212*, 54-63.
- 640 51. Yan, C.; Nie, W.; Äijälä, M.; Rissanen, M. P.; Canagaratna, M. R.; Massoli, P.; Junninen,
641 H.; Jokinen, T.; Sarnela, N.; Häme, S. A. K.; Schobesberger, S.; Canonaco, F.; Yao, L.; Prévôt,

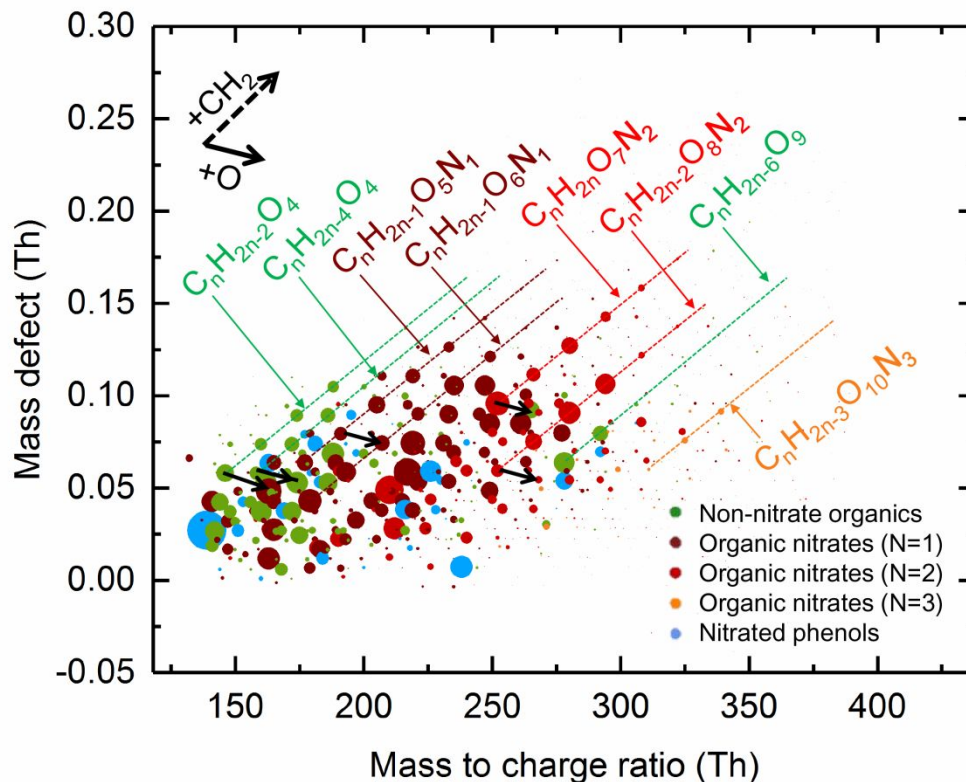
- 642 A. S. H.; Petäjä, T.; Kulmala, M.; Sipilä, M.; Worsnop, D. R.; Ehn, M., Source characterization
643 of highly oxidized multifunctional compounds in a boreal forest environment using positive
644 matrix factorization. *Atmospheric Chemistry and Physics* **2016**, *16*, 12715-12731.
- 645 52. Nieminen, T.; Lehtinen, K. E. J.; Kulmala, M., Sub-10 nm particle growth by vapor
646 condensation-effects of vapor molecule size and particle thermal speed. *Atmospheric Chemistry
647 and Physics* **2010**, *10*, 9773-9779.
- 648 53. Rohrer, F.; Berresheim, H., Strong correlation between levels of tropospheric hydroxyl
649 radicals and solar ultraviolet radiation. *Nature* **2006**, *442*, 184-187.
- 650 54. Tan, Z.; Lu, K.; Hofzumahaus, A.; Fuchs, H.; Bohn, B.; Holland, F.; Liu, Y.; Rohrer, F.;
651 Shao, M.; Sun, K.; Wu, Y.; Zeng, L.; Zhang, Y.; Zou, Q.; Kiendler-Scharr, A.; Wahner, A.;
652 Zhang, Y., Experimental budgets of OH, HO₂ and RO₂ radicals and implications for ozone
653 formation in the Pearl River Delta in China 2014. *Atmospheric Chemistry and Physics* **2019**, *19*,
654 7129-7150.
- 655 55. Kulmala, M.; Dada, L.; Daellenbach, K. R.; Yan, C.; Stolzenburg, D.; Kontkanen, J.;
656 Ezhova, E.; Hakala, S.; Tuovinen, S.; Kokkonen, T. V.; Kurppa, M.; Cai, R.; Zhou, Y.; Yin, R.;
657 Baalbaki, R.; Chan, T.; Chu, B.; Deng, C.; Fu, Y.; Ge, M.; He, H.; Heikkinen, L.; Junninen, H.;
658 Liu, Y.; Lu, Y.; Nie, W.; Rusanen, A.; Vakkari, V.; Wang, Y.; Yang, G.; Yao, L.; Zheng, J.;
659 Kujansuu, J.; Kangasluoma, J.; Petaja, T.; Paasonen, P.; Jarvi, L.; Worsnop, D.; Ding, A.; Liu,
660 Y.; Wang, L.; Jiang, J.; Bianchi, F.; Kerminen, V. M., Is reducing new particle formation a
661 plausible solution to mitigate particulate air pollution in Beijing and other Chinese megacities?
662 *Faraday Discussions* **2021**, *226*, 334-347.
- 663 56. Riva, M.; Rantala, P.; Krechmer, J. E.; Peräkylä, O.; Zhang, Y.; Heikkinen, L.; Garmash,
664 O.; Yan, C.; Kulmala, M.; Worsnop, D.; Ehn, M., Evaluating the performance of five different
665 chemical ionization techniques for detecting gaseous oxygenated organic species. *Atmospheric
666 Measurement Techniques* **2019**, *12*, 2403-2421.
- 667 57. Ye, Q.; Wang, M.; Hofbauer, V.; Stolzenburg, D.; Chen, D.; Schervish, M.; Vogel, A.;
668 Mauldin, R. L.; Baalbaki, R.; Brilke, S.; Dada, L.; Dias, A.; Duplissy, J.; El Haddad, I.;
669 Finkenzeller, H.; Fischer, L.; He, X.; Kim, C.; Kürten, A.; Lamkaddam, H.; Lee, C. P.;
670 Lehtipalo, K.; Leiminger, M.; Manninen, H. E.; Marten, R.; Mentler, B.; Partoll, E.; Petäjä, T.;
671 Rissanen, M.; Schobesberger, S.; Schuchmann, S.; Simon, M.; Tham, Y. J.; Vazquez-Pufleau,
672 M.; Wagner, A. C.; Wang, Y.; Wu, Y.; Xiao, M.; Baltensperger, U.; Curtius, J.; Flagan, R.;
673 Kirkby, J.; Kulmala, M.; Volkamer, R.; Winkler, P. M.; Worsnop, D.; Donahue, N. M.,

- 674 Molecular Composition and Volatility of Nucleated Particles from α -Pinene Oxidation between -
675 50 °c and +25 °c. *Environmental Science and Technology* **2019**, *53*, 12357-12365.
- 676 58. Olenius, T.; Yli-Juuti, T.; Elm, J.; Kontkanen, J.; Riipinen, I., Chapter 11 - New Particle
677 Formation and Growth: Creating a New Atmospheric Phase Interface. In *Physical Chemistry of*
678 *Gas-Liquid Interfaces*, Faust, J. A.; House, J. E., Eds. Elsevier: 2018; pp 315-352.
- 679 59. Hallquist, M.; Wenger, J. C.; Baltensperger, U.; Rudich, Y.; Simpson, D.; Claeys, M.;
680 Dommen, J.; Donahue, N. M.; George, C.; Goldstein, A. H.; Hamilton, J. F.; Herrmann, H.;
681 Hoffmann, T.; Iinuma, Y.; Jang, M.; Jenkin, M. E.; Jimenez, J. L.; Kiendler-Scharr, A.;
682 Maenhaut, W.; McFiggans, G.; Mentel, T. F.; Monod, A.; Prevot, A. S. H.; Seinfeld, J. H.;
683 Surratt, J. D.; Szmigielski, R.; Wildt, J., The formation, properties and impact of secondary
684 organic aerosol: current and emerging issues. *Atmospheric Chemistry and Physics* **2009**, *9*, (14),
685 5155-5236.
- 686 60. Wang, M.; Kong, W.; Marten, R.; He, X. C.; Chen, D.; Pfeifer, J.; Heitto, A.; Kontkanen,
687 J.; Dada, L.; Kurten, A.; Yli-Juuti, T.; Manninen, H. E.; Amanatidis, S.; Amorim, A.; Baalbaki,
688 R.; Baccarini, A.; Bell, D. M.; Bertozzi, B.; Brakling, S.; Brilke, S.; Murillo, L. C.; Chiu, R.;
689 Chu, B.; De Menezes, L. P.; Duplissy, J.; Finkenzeller, H.; Carracedo, L. G.; Granzin, M.;
690 Guida, R.; Hansel, A.; Hofbauer, V.; Krechmer, J.; Lehtipalo, K.; Lamkaddam, H.; Lampimaki,
691 M.; Lee, C. P.; Makhmutov, V.; Marie, G.; Mathot, S.; Mauldin, R. L.; Mentler, B.; Muller, T.;
692 Onnela, A.; Partoll, E.; Petaja, T.; Philippov, M.; Pospisilova, V.; Ranjithkumar, A.; Rissanen,
693 M.; Rorup, B.; Scholz, W.; Shen, J.; Simon, M.; Sipila, M.; Steiner, G.; Stolzenburg, D.; Tham,
694 Y. J.; Tome, A.; Wagner, A. C.; Wang, D. S.; Wang, Y.; Weber, S. K.; Winkler, P. M.; Wlasits,
695 P. J.; Wu, Y.; Xiao, M.; Ye, Q.; Zauner-Wieczorek, M.; Zhou, X.; Volkamer, R.; Riipinen, I.;
696 Dommen, J.; Curtius, J.; Baltensperger, U.; Kulmala, M.; Worsnop, D. R.; Kirkby, J.; Seinfeld, J.
697 H.; El-Haddad, I.; Flagan, R. C.; Donahue, N. M., Rapid growth of new atmospheric particles by
698 nitric acid and ammonia condensation. *Nature* **2020**, *581*, (7807), 184-189.
- 699 61. Cai, R.; Li, C.; He, X.-C.; Deng, C.; Lu, Y.; Yin, R.; Yan, C.; Wang, L.; Jiang, J.;
700 Kulmala, M.; Kangasluoma, J., Impacts of coagulation on the appearance time method for new
701 particle growth rate evaluation and their corrections. *Atmospheric Chemistry and Physics* **2021**,
702 *21*, (3), 2287-2304.
- 703 62. Zha, Q.; Yan, C.; Junninen, H.; Riva, M.; Sarnela, N.; Aalto, J.; Quéléver, L.; Schallhart,
704 S.; Dada, L.; Heikkinen, L.; Peräkylä, O.; Zou, J.; Rose, C.; Wang, Y.; Mammarella, I.; Katul,
705 G.; Vesala, T.; Worsnop, D. R.; Kulmala, M.; Petäjä, T.; Bianchi, F.; Ehn, M., Vertical

- 706 characterization of highly oxygenated molecules (HOMs) below and above a boreal forest
707 canopy. *Atmospheric Chemistry and Physics* **2018**, *18*, 17437-17450.
- 708 63. Kürten, A.; Bergen, A.; Heinritzi, M.; Leiminger, M.; Lorenz, V.; Piel, F.; Simon, M.;
709 Sitals, R.; Wagner, A. C.; Curtius, J., Observation of new particle formation and measurement of
710 sulfuric acid, ammonia, amines and highly oxidized organic molecules at a rural site in central
711 Germany. *Atmospheric Chemistry and Physics* **2016**, *16*, 12793-12813.
- 712 64. Mutzel, A.; Poulain, L.; Berndt, T.; Iinuma, Y.; Rodigast, M.; Böge, O.; Richters, S.;
713 Spindler, G.; Sipilä, M.; Jokinen, T.; Kulmala, M.; Herrmann, H., Highly Oxidized
714 Multifunctional Organic Compounds Observed in Tropospheric Particles: A Field and
715 Laboratory Study. *Environmental Science and Technology* **2015**, *49*, 7754-7761.
- 716 65. Sarnela, N.; Jokinen, T.; Nieminen, T.; Lehtipalo, K.; Junninen, H.; Kangasluoma, J.;
717 Hakala, J.; Taipale, R.; Schobesberger, S.; Sipilä, M.; Larnimaa, K.; Westerholm, H.; Heijari, J.;
718 Kerminen, V. M.; Petäjä, T.; Kulmala, M., Sulphuric acid and aerosol particle production in the
719 vicinity of an oil refinery. *Atmospheric Environment* **2015**, *119*, 156-166.
- 720

721 **Figures.**

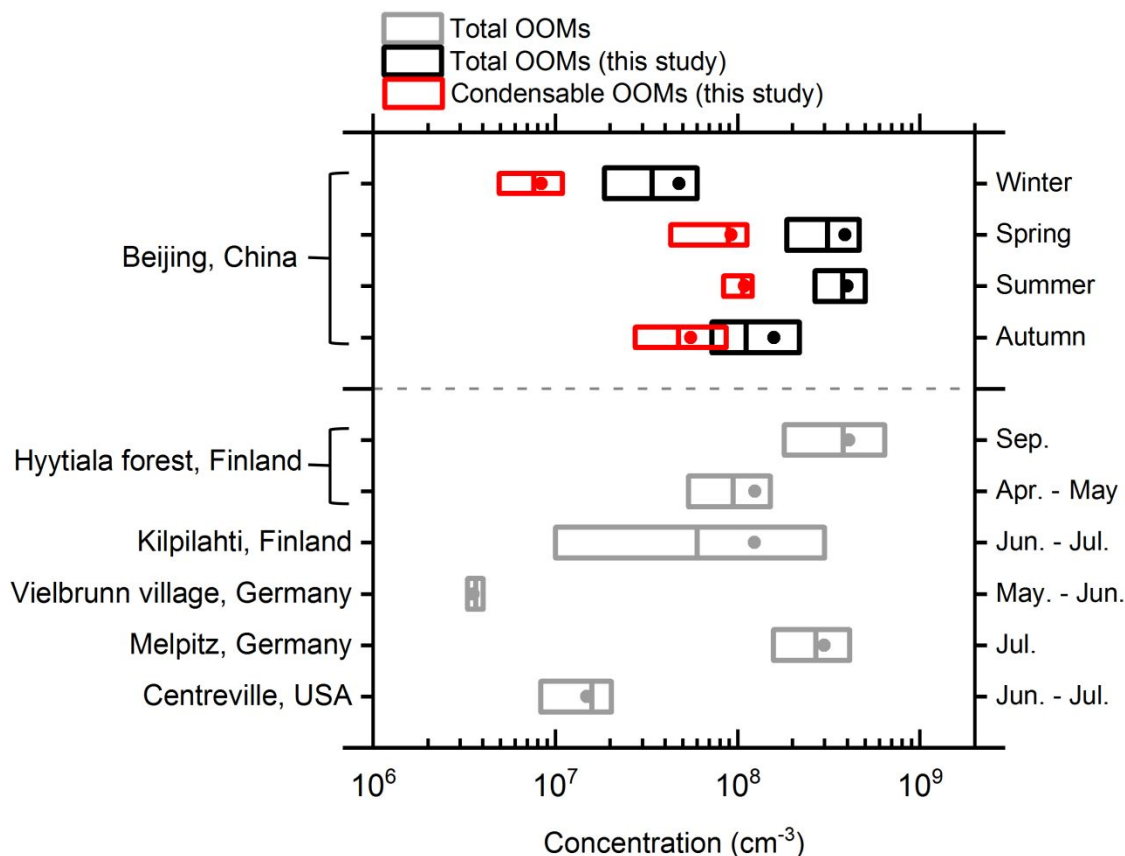
722



723

724 **Figure 1.** Mass-defect plot of observed oxygenated organic molecules (OOMs) in Beijing of
 725 spring. The detected OOMs are marked in the mass-defect plot by their exact mass and mass defect
 726 (exact mass subtracted by its unit mass). The differences in molecular composition are shown at
 727 regular spacing in the mass-defect plot. Adjacent points on the two direction lines are differed by
 728 a $-\text{CH}_2-$ group or oxygen atom in composition. The general chemical formulas are annotated by
 729 the lines. OOMs with different numbers of nitrogen atoms are shown in different colors. Nitrated
 730 phenols are classified separately. The size of the circle corresponds to the concentration of OOMs.

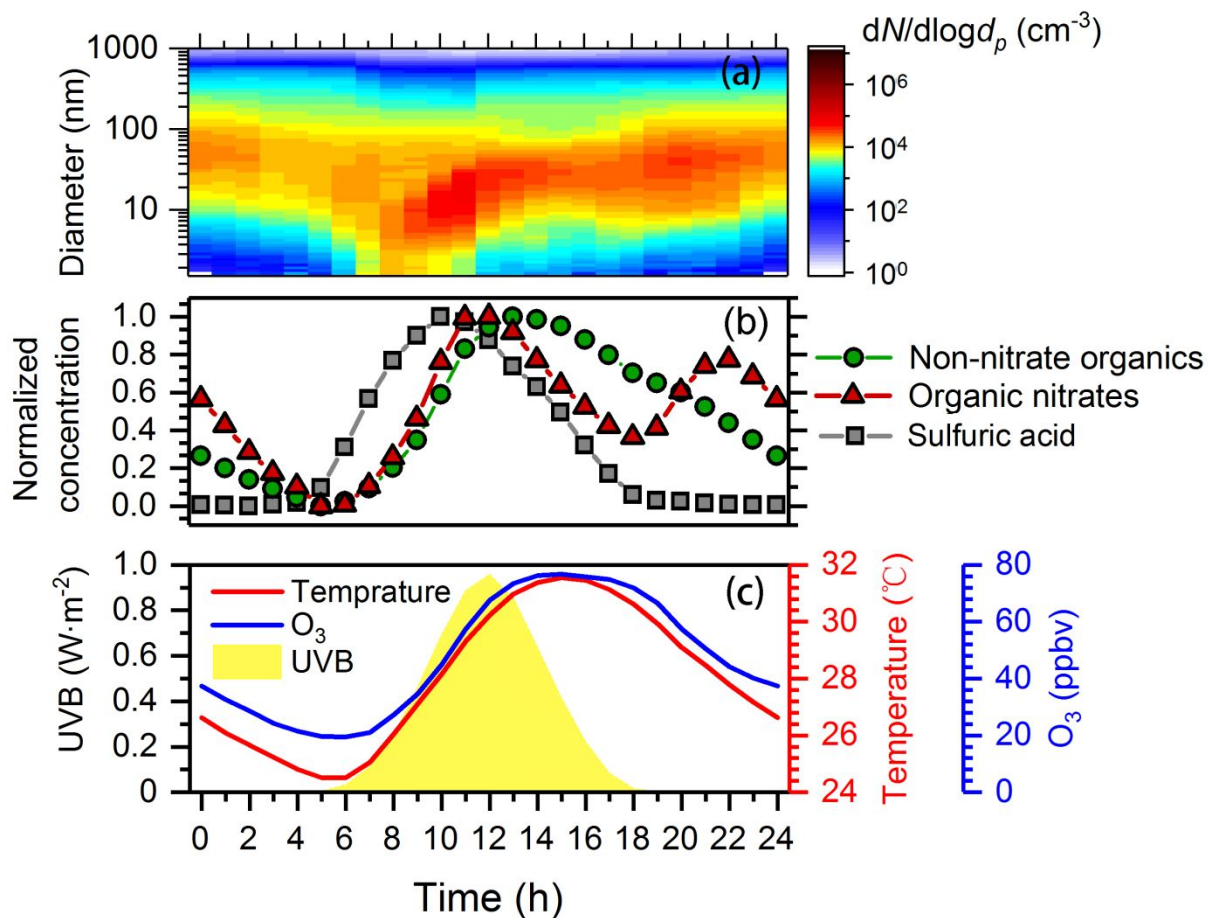
731



732

733 **Figure 2.** Comparison of atmospheric oxygenated organic molecules (OOMs) measured by nitrate
 734 CI-APi-ToF at different locations worldwide. Boxes in black and red are results of total OOMs
 735 and condensable OOMs of urban Beijing during the whole observation periods in this study. Boxes
 736 in gray are previous results around the world, including Yan et al.⁵¹ and Zha et al.⁶² in Hyytiälä
 737 forest, Kürten et al.⁶³ in Vielbrunn village, Mutzel et al.⁶⁴ in Kilpilahti, Sarnela et al.⁶⁵ in
 738 Centreville, and Krechmer et al.²⁵ in Melpitz. Among them, Vielbrunn and Centreville only report
 739 the main oxidation products of monoterpene and isoprene respectively. The boundaries of boxes
 740 on x-axis correspond to the 25th and 75th values. Lines and points in boxes are median and mean
 741 concentration respectively.

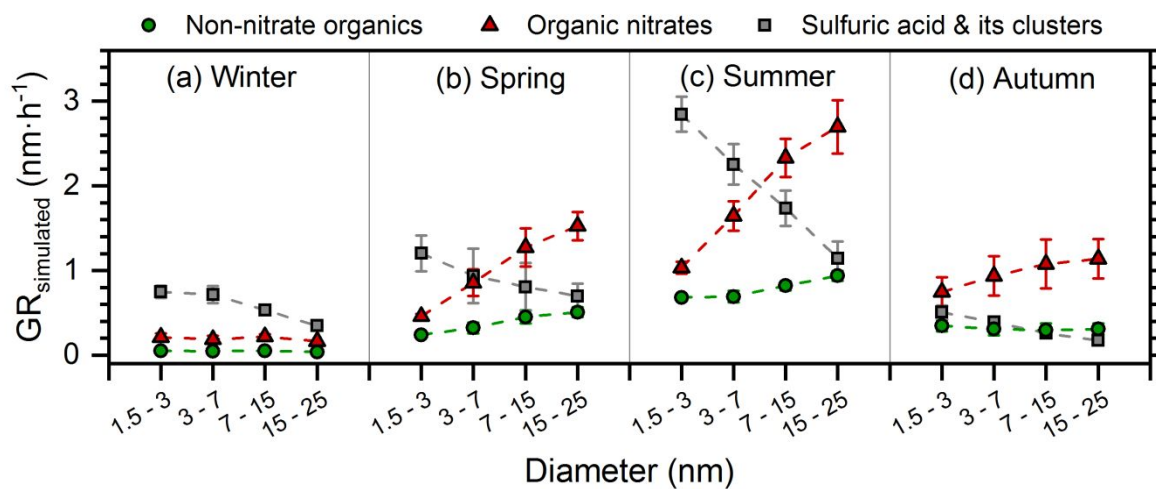
742



743

744 **Figure 3.** Diurnal variation of observed oxygenated organic compounds and other measured
745 parameters in summer. Note that the particle size distribution shown in the top panel is the
746 averaged result of new particle formation days.

747

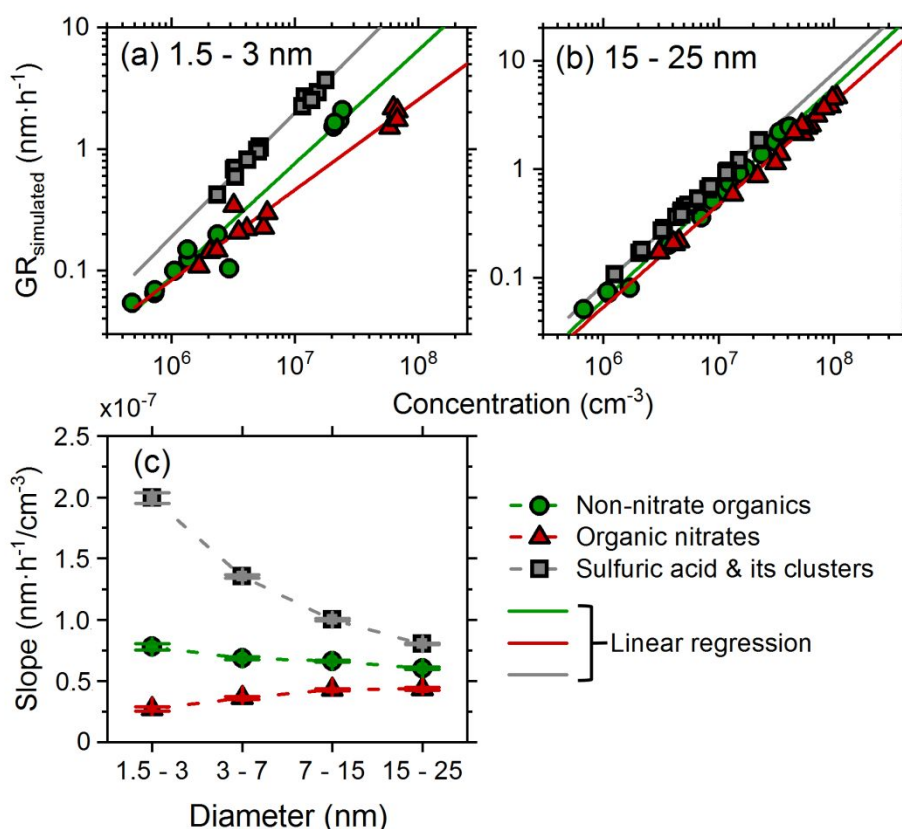


748

749 **Figure 4.** Simulated growth rates contributed by sulfuric acid and its clusters, non-nitrogen OOMs,
 750 and nitrogen-containing OOMs. Markers are the averaged results. Error bars are the standard
 751 deviations of the simulated results. There are 11, 5, 6 and 5 new particles cases for winter, spring,
 752 summer and autumn respectively. The particle growth period of each case (generally 7 am-12 pm)
 753 is used for simulation.

754

755



756

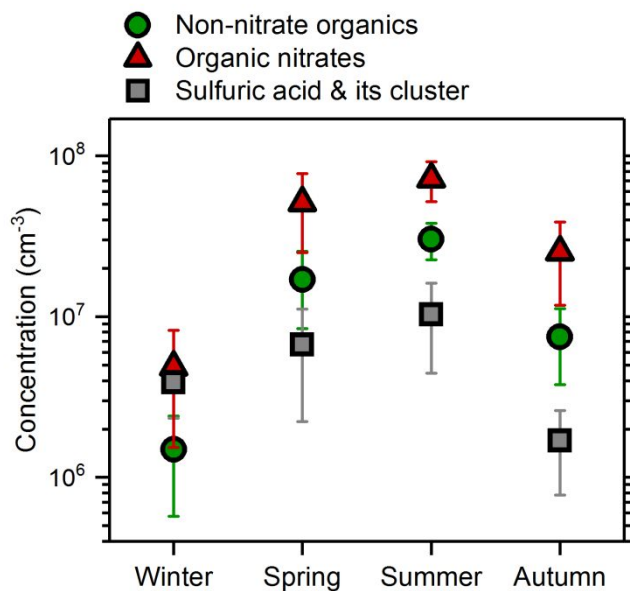
757 **Figure 5.** Growth promotion efficiency of different condensing vapors in condensation growth
 758 rate. (a)-(b) Vapor concentration and simulated condensation growth rate for different particle size
 759 bins, including sulfuric acid and its clusters, non-nitrogen OOMs, and nitrogen-containing OOMs.
 760 Solid lines are the linear regressions for different vapors. (c) Slopes of linear regressions
 761 ($GR_{\text{simulated}}$ v.s. concentration) for different particle size bins, including 1-3 nm, 3-7 nm, 7-15 nm,
 762 and 15-25 nm. The green circle represents non-nitrogen OOMs, the red triangle represents
 763 nitrogen-containing OOMs, and the gray square represents sulfuric acid and its clusters. The caps
 764 of markers in panel (c) are error bars for linear regression uncertainties, which range in 7.2×10^{-10}
 765 - $4.3 \times 10^{-9} \text{ nm}\cdot\text{h}^{-1}/\text{cm}^{-3}$.

766

767

768

769

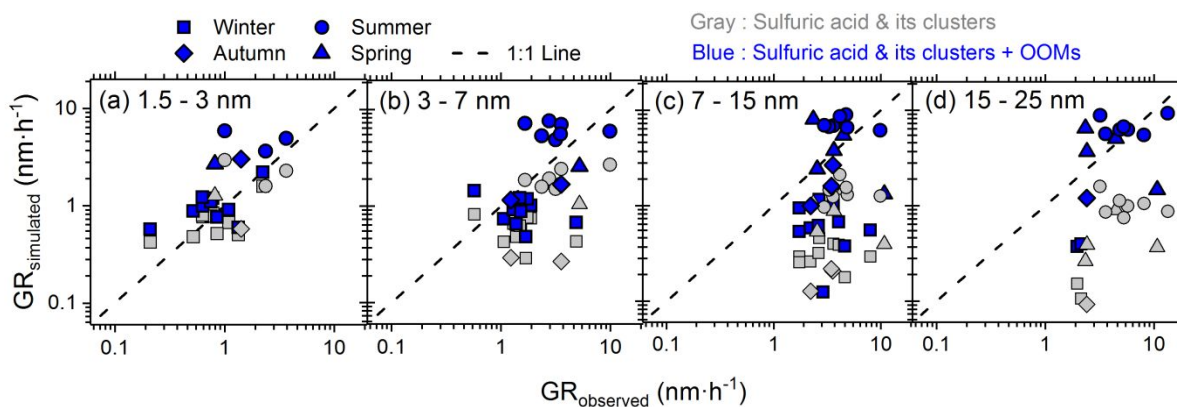


770

771 **Figure 6.** Concentrations of condensable oxygenated organic molecules (OOMs) in four seasons.
772 Markers are the averaged results. Error bars are the standard deviations. The green circle represents
773 non-nitrogen OOMs, the red triangle represents nitrogen-containing OOMs, and the gray square
774 represents sulfuric acid and its clusters.

775

776



777

778 **Figure 7.** The relations of simulated growth rate and observed growth rate for different particle
779 size bins. Results of different seasons are shown in different shapes of markers. The x-axis is the
780 observed growth rate and the y-axis is the simulated growth rate. The simulated growth rate of
781 sulfuric acid and its clusters are shown in gray, and the blue ones are sulfuric acid and its clusters
782 plus OOMs. The dashed lines are 1:1 lines. Note the observed growth rates are calculated using
783 mode fitting method.