1	Measurement Report: Effects of anthropogenic emissions and
2	environmental factors on biogenic secondary organic aerosol
3	(BSOA) formation in a coastal city of Southeastern China
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5 6	Youwei Hong ^{a,b,c,d*} , Xinbei Xu ^{a,b,c} , Dan Liao ^e , Taotao Liu ^{a,b,c} ,Xiaoting Ji ^{a,b,c} , Ke Xu ^{a,b,d} , Chunyang Liao ^f , Ting Wang ^g , Chunshui Lin ^g , Jinsheng Chen ^{a,b,c*}
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8 9	^a Center for Excellence in Regional Atmospheric Environment, Institute of Urban Environment, Chinese Academy of Sciences, Xiamen, 361021, China
10 11	^b Key Lab of Urban Environment and Health, Institute of Urban Environment, Chinese Academy of Sciences, Xiamen, 361021, China
12	^c University of Chinese Academy of Sciences, Beijing, 100049, China
13	^d School of Life Sciences, Hebei University, Baoding, 071000, China
14	^e College of Environment and Public Health, Xiamen Huaxia University, Xiamen 361024, China
15 16	^f State Key Laboratory of Environmental Chemistry and Ecotoxicology, Research Center for Eco- Environmental Sciences, Chinese Academy of Sciences, Beijing 100085, China
17	^g Institute of Earth Environment, Chinese Academy of Sciences, Xi'an, 710061, China
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19	*Corresponding author E-mail: Jinsheng Chen (jschen@iue.ac.cn); Youwei Hong
20	(<u>ywhong@iue.ac.cn</u>)
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34 Abstract:

To better understand the formation of biogenic secondary organic aerosol (BSOA), 35 aerosol samples with a 4 h time resolution were collected during summer and 36 wintertime in the southeast of China, along with on-line measurements of trace gases, 37 aerosol chemical compositions, and meteorological parameters. The samples were 38 analyzed by gas chromatography-mass spectrometry for PM_{2.5}-bound SOA tracers, 39 including isoprene (SOA_I), α/β -pinene (SOA_M), β -caryophyllene (SOA_C), and toluene 40 (ASOA). The average concentrations of total SOA tracers in winter and summer were 41 38.8 and 111.9 ng m⁻³, respectively, with the predominance of SOA_M (70.1% and 42 45.8%), followed by SOA_I (14.0% and 45.6%), ASOA (11.0% and 6.2%) and SOA_C 43 (4.9% and 2.3%). Compared to those in winter, the majority of BSOA tracers in summer 44 showed significant positive correlations with Ox (O_3+NO_2) (r = 0.443 \sim 0.808), HONO 45 $(r = 0.299 \sim 0.601)$, ultraviolet (UV) $(r = 0.382 \sim 0.588)$ and temperature (T) $(r = 0.299 \sim 0.601)$ 46 $0.529 \sim 0.852$), indicating the influence of photochemical oxidation under relatively 47 clean conditions. However, in winter, BSOA tracers were significantly correlated with 48 $PM_{2.5}$ (r = 0.407 \sim 0.867), NO_3^- (r = 0.416 \sim 0.884), SO_4^{2-} (r = 0.419 \sim 0.813), and NH_3 49 $(r = 0.440 \sim 0.757)$, attributed to the contributions of anthropogenic emissions. Major 50 51 BSOA tracers in both seasons was linearly correlated with aerosol acidity (pH) (r =0.421 \sim 0.752), liquid water content (LWC) (r = 0.403 \sim 0.876) and SO₄²⁻ (r = 0.419 \sim 52 0.813). The results indicated that acid-catalyzed reactive uptake onto sulfate aerosol 53 particles enhanced the formation of BSOA. In summer, the clean air mass originated 54 from the ocean, and chlorine depletion was observed. We also found that concentrations 55 of the total SOA tracers was correlated with HCl ($R^2 = 0.545$) and chlorine ions (r = 56 57 $0.280 \sim 0.639$) in PM_{2.5}, reflecting the contribution of Cl-initiated VOC oxidations to 58 the formation of SOA. In winter, the northeast dominant wind direction brought 59 continental polluted air mass to the monitoring site, affecting the transformation of BSOA tracers. This implied that anthropogenic emissions, atmospheric oxidation 60 capacity and halogen chemistry have significant effects on the formation of BSOA in 61 62 the southeast coastal area.

Keywords: SOA tracers; biogenic volatile organic compounds; anthropogenic
pollutants; atmospheric oxidation capacity; coastal area

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66 **1. Introduction**

67 Secondary organic aerosol (SOA) has attracted widespread scientific research concerns, due to its potential impacts on climate change, human health and air quality 68 (Shrivastava et al., 2017; Reid et al., 2018; Zhu et al., 2019; Wang et al., 2021b). 69 70 Understanding the formation of SOA and assessing its relevance for environmental 71 effects become an integral part of aerosol chemistry (Charan et al., 2019; Xiao et al., 2020; Palmer et al., 2022). However, due to its complex precursors and atmospheric 72 physical or chemical processes, SOA prediction by air quality models remains highly 73 uncertain (McFiggans et al., 2019). Therefore, it is necessary to better explore missed 74 75 SOA sources and unknown SOA formation mechanisms.

SOA is produced by the conversion of biogenic and anthropogenic volatile organic 76 77 compounds (BVOCs and AVOCs) through complex homogeneous and heterogeneous 78 reactions (Charan et al., 2019; Xiao et al., 2020; Mahilang et al., 2021). BVOCs are the 79 main precursors of SOA on a global scale, while AVOCs are the predominant 80 contributor to SOA in urban areas (Hallquist et al., 2009; Wang et al., 2021a). Recently, 81 laboratory, field and modeling studies have shown that anthropogenic emissions greatly affect the formation of BSOA (Hoyle et al., 2011; Shrivastava et al., 2019; Zhang et al., 82 83 2019b; Zhang et al., 2019c; Mahilang et al., 2021; Xu et al., 2021). Anthropogenic air pollutants, such as NOx, SO₂, NH₃ and aerosols, could influence the conversion of 84 85 BVOCs to the particulate phase and the production of nitrogen and sulfur compounds 86 (Wang et al., 2020). NOx is one of the important drivers of SOA formation and yields 87 during both daytime and nighttime through alternating the fate of peroxy radicals (RO₂ \cdot) (Sarrafzadeh et al., 2016; Newland et al., 2021). While ·OH dominates the 88 photochemical oxidation of BVOC during daylight hours, and NO₃ becomes one of the 89 90 main oxidants for biogenic SOA and organic nitrate formation at night. SO₂ also plays

an important role in changing SOA formation from BVOC photooxidation and 91 92 ozonolysis through sulfuric acid formation and acid-catalyzed heterogeneous reactions (Zhao et al., 2018; Zhang et al., 2019b; Xu et al., 2021). In addition, NH₃ and amines 93 can affect the SOA yields and composition through both gas-phase and heterogeneous 94 reactions, by reacting with sulfuric or nitric acid to generate secondary inorganic 95 aerosols (SIA) (Ma et al., 2018; Liu et al., 2021; Lv et al., 2022). However, due to 96 97 complex precursors and atmospheric processes, the combined effects of anthropogenic 98 emissions and meteorological factors on the formation of SOA are not fully understood.

99 The coastal area of southeastern China is under the East Asian monsoon control, which cause an obvious alternation of polluted and clean air masses from continental 100 and ocean area, respectively (Wu et al., 2019; Hong et al., 2021). Also, the local 101 102 geographical environment, including relatively high humidity, dense vegetation and 103 strong atmospheric oxidation capacity, provides a good chance to study the sources and formation mechanisms of SOA. In our previous studies, ground-based observations in 104 a mountainous forest area of this region showed that BSOA tracers were the largest 105 106 contributor to SOA, and the aerosols were highly oxidized (Hong et al., 2019). However, with the development of rapid urbanization, anthropogenic emissions will be of great 107 significance on SOA formation (Liu et al., 2020). Halogen radicals (chlorine, bromine, 108 109 iodine) have an important role in tropospheric oxidants chemistry and OA formation (Wang et al., 2021c). Therefore, it is necessary to investigate the sources and formation 110 mechanisms of SOA in coastal urban areas, and so as to provide a scientific basis for 111 112 the estimation of regional SOA budgets and PM_{2.5} pollution control.

In this study, a continuous PM_{2.5} sampling campaign with a 4 h time resolution was conducted in a coastal city of southeastern China during the winter and summertime period. Seasonal, diurnal variations and SOC contributions of SOA tracers were analyzed. Atmospheric process identified by SOA tracers in different seasons were further analyzed. Finally, the combined effects of anthropogenic emissions and major environmental factors on promoting SOA formation was discussed.

119 **2. Materials and methods**

120 2.1 Sample collection

121 The sampling was performed at the Institute of Urban Environment, Chinese Academy of Sciences (118.06° E, 24.61° N), which is located in a suburban area of 122 123 Xiamen, a coastal city of southeastern China. Detailed information of the air monitoring 124 supersite was described in our previous study (Hong et al., 2021). Briefly, time-resolved 125 (00:00-08:00, 08:00-12:00, 12:00-16:00, 16:00-20:00, 20:00-24:00 CST - China Standard Time) PM_{2.5} samples were collected on the rooftop of the station (about 70m 126 above the ground). The sampling was carried out by using a high volume $(1.05 \text{ m}^3 \text{ min}^{-1})$ 127 sampler (TH-1000C, Wuhan Tianhong, China) with a PM_{2.5} inlet from 10 to 18 January, 128 and from 5 to 14 July 2020. All samples were collected onto pre-baked (450 °C, 6 h) 129 130 quartz fiber filters. Field blank samples were also collected. The sample filters were separately sealed in aluminum foil and stored in a freezer (-20 °C) prior to analysis. 131

132 2.2 SOA tracers analysis by GC/MS

133 The isoprene-derived SOA (SOA_I) tracers included 2 methyltetrols (MTLs: 2methylthreitol (MTL1) and 2-methylerythritol (MTL2)), C5-alkene triols (cis-2-134 methyl-1,3,4-trihydroxy-1-butene, trans-2-methyl-1,3,4-trihydroxy-1-butene, and 3-135 methyl-2,3,4-trihydroxy-1-butene) and 2-methylglyceric 136 acid (MGA). The monoterpene-derived SOA (SOA_M) tracers were composed of pinic acid (PA), pinonic 137 acid (PNA), 3-hydroxyglutaric acid (HGA), 3-methyl-1,2,3-butanetricarboxylic acid 138 (MBTCA), 3-hydro-4,4-dimethyglutaric acid (HDMGA), and 3-acetylglutaric acid 139 (AGA). The β -caryophyllene-derived SOA (SOA_C) tracer was β -caryophyllenic acid 140 141 (CA), the toluene-derived SOA (SOA_A) tracer was 2,3-Dihydroxy-4-oxopentanoic acid (DHOPA) and levoglucosan (LEV) as a tracer of biomass burning. Due to the lack of 142 authentic standards, surrogate standards (including erythritol, malic acid, PA and 143 citramalic acid) were used to compensate for unavoidable assay variance of SOA_I, 144 SOA_M, SOA_C and SOA_A tracer in each sample during the pretreatment process, 145 respectively (Fu et al., 2009). However, inherent low volatility of isoprene SOA tracers 146

could cause the uncertainties of using the GC/MS method, and low-volatility oligomers
might break down into monomers, such as C5-alkene triols and 2-methyltetrols (LopezHilfiker et al., 2016; Hu et al., 2016). Therefore, quantifying the abundance of certain
SOA tracers remained a lot of uncertainties.

The analytical procedure of fifteen SOA tracers was published in our previous 151 studies (Hong et al., 2019; Liu et al., 2020). Briefly, the filter samples were 152 ultrasonically extracted with a mixture of dichloromethane and methanol (2:1, v/v) for 153 154 10 min three times. The mixed extracts were filtered with a PTFE filter (0.22 μ m), and dried with high purity N_2 (99.99%), and then derivatized with 60 μ L of 155 N,O-bis-(trimethylsilyl) trifluoroacetamide (BSTFA) with 1% trimethylsilyl chloride 156 and 10 µL of pyridine at 70 °C for 3 h. At last, 140 µL of internal standard solution (¹³ 157 C n-alkane solution, 1.507 ng μ L⁻¹) was added into the samples. Then, relative response 158 factors (RRFs) of surrogate and internal standard were calculated to quantify the 159 targeted organic tracers in each sample. Details of SOA tracer's calculated 160 concentrations based on RRFs were presented in our previous studies (Hong et al., 2019; 161 162 Liu et al., 2020).

Fifteen SOA tracers were determined by GC-MSD (7890A/5975C, Agilent 163 Technologies, Inc., USA) with a DB-5 MS silica capillary column (i.d. 30×0.25 mm, 164 0.25 µm film thickness). 1 µL sample was injected with splitless mode and high purity 165 helium (99.999%) was used as carrier gas at a stable flow of 1.0 mL/min. The GC 166 temperature was initiated at 100 °C (held for 1 min) and then to 300 °C at 5 °C min⁻¹, 167 and kept at 300 °C for 10 min. The operation mode is electron ionization (EI) mode of 168 70 ev. The method detection limits (MDLs) for erythritol and PNA were 0.01 and 0.02 169 ng m⁻³, respectively. The recoveries of erythritol, PNA, malic acid, PA and citramalic 170 acid were 67±2%, 73±1%, 75±1%, 88±7% and 82±8%, respectively. SOA tracers were 171 not detected in the field blank samples. 172

173 *2.3 Observations in the air monitoring supersite*

Water-soluble inorganic ions (WSII) in $PM_{2.5}$ (Cl⁻, SO_4^{2-} , NO_3^{-} , Na^+ , K^+ , NH_4^+ , Mg²⁺, and Ca²⁺) and gas pollutants (HCl, HONO, HNO₃, NH₃) were hourly measured using a monitoring device for aerosols and gases in ambient Air (MARGA 2080;

- 177 Metrohm Applikon B.V.; Delft, Netherlands). Internal calibration was carried out using
- LiBr standard solutions. The detection limit of Cl^{-} , SO_4^{2-} , NO_3^{-} , Na^+ , K^+ , NH_4^+ , Mg^{2+} ,
- and Ca^{2+} were 0.01, 0.04, 0.05, 0.05, 0.09, 0.05, 0.06 and 0.09 µg m⁻³, respectively.

Hourly mass concentrations of PM_{2.5} and PM₁₀ were measured by using a tapered 180 element oscillating microbalance (TEOM1405, Thermo Scientific Corp., MA, USA). 181 NO₂, SO₂ and O₃ were monitored using continuous gas analyzers (TEI 42i, 43i, and 182 183 49i, Thermo Scientific Corp., MA, USA). Ambient meteorological parameters including relative humidity (RH), temperature (T), wind speed (WS), and wind 184 direction (WD) were obtained by an ultrasonic atmospherium (150WX, Airmar, the 185 USA). Photolysis frequencies were determined using a photolysis spectrometer (PFS-186 100, Focused Photonics Inc., Hangzhou, China), including the photolysis rate constants 187 J (O¹D), J (HCHO M), J (HCHO R), J (NO₂), J (H₂O₂), J (HONO), J (NO₃ M) and 188 J (NO₃ R), and the spectral band ranged from 270 to 790 nm. Boundary layer height 189 (BLH) based on ERA-5 reanalysis dataset was downloaded from the following link 190 191 https://www.ecmwf.int/en/forecasts/datasets/reanalysis-datasets/era5.

192 2.4 Estimation of SOC using a tracer-based method

The fraction of SOC formed by the oxidation of monoterpene, isoprene, β -193 caryophyllene and toluene was estimated using a tracer-based method (Kleindienst et 194 al., 2007; Hong et al., 2019). It is defined as $[SOC] = \sum i[tri]/f_{SOC}$, where [SOC]195 represents the mass concentration of SOC (μ gC m⁻³) and $\sum i$ [tri] means the sum of the 196 concentration of individual SOA tracer ($\mu g m^{-3}$). The carbon mass fractions (f_{SOC}) of 197 monoterpene, isoprene, β -caryophyllene and toluene were $0.231 \pm 0.111, 0.155 \pm 0.039$, 198 199 0.023 ± 0.005 and 0.008 ± 0.003 , respectively, based on smog-chamber experimental data (Kleindienst et al., 2007). 200

201 2.5 Aerosol acidity and OH calculation

The forward mode of ISORROPIA II thermodynamic model was used to calculate the aerosol acidity (pH) (Fountoukis and Nenes, 2007). ISORROPIA II can calculate liquid water content (LWC), based on total SO_4^{2-} , NO_3^{-} (gas HNO₃ plus particle NO_3^{-}),

205 Cl^- , ammonia (gas NH₃ plus particle NH₄⁺), non-volatile cations (Na⁺, K⁺, Ca²⁺, Mg²⁺),

and meteorological factors (RH and T) (Rumsey et al., 2014; Guo et al., 2016). The pH

value from ISORROPIA II was calculated using the following equation:

$$pH = - lg \left(\frac{1000 \times H^+}{LWC}\right)$$

209 where H^+ is the hydronium ion concentration loading for an air sample ($\mu g/m^3$).

The OH concentration ([OH]) was estimated using the NO₂ and HONO concentrations and the photolysis rate constants (J) of NO₂, O₃, and HONO, according to the following improved empirical formula (Wen et al., 2019).

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$$[OH] = 4.1 \times 10^9 \times \frac{J(O^1D)^{0.83} \times J(NO_2)^{0.19} \times (140 \times NO_2 + 1) + HONO \times J(HONO)}{0.41 \times NO_2^2 + 1.7 \times NO_2 + 1 + NO \times k_{NO+OH} + HONO \times k_{HONO+OH}}$$

214 2.6 Statistical analysis

Correlation analysis by SPSS 22.0 software (IBM, Armonk, NY, USA) was used to study the relationship among SOA tracers, meteorological parameters and criteria air pollutants. One-way analysis of variance (ANOVA) was adopted to examine the variations of different factors.

219 2.7. Backward trajectory analysis

Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) was used to analyze the impacts of air masses on Xiamen during different seasons. 72 h backward trajectories were calculated every hour at a height of 500 m. The meteorological data with a resolution of 1° longitude \times 1° latitude was obtained from the NCEP/GDAS. Cluster analysis was adopted using the total spatial variance (TSV).

225 **3 R**

3 Results and discussion

226 *3.1. Overview of air pollutants*

The concentrations of criteria air pollutants, including SO₂, CO, NO₂, O₃, PM_{2.5} and PM₁₀, and meteorological parameters during wintertime and summertime were shown in Fig.1. The concentrations of PM_{2.5} in winter ranged from 14.9 to 75.3 μ g m⁻³ with an average of 42.1 μ g m⁻³, which was much higher than that (the average of 18.4

 μ g m⁻³) in summer, ranging from 12.8 to 46.4 μ g m⁻³. The concentrations of CO, NO₂ 231 and PM_{10} showed similar seasonal trends to the pattern of $PM_{2.5}$. In contrast, O_3 had the 232 highest concentration in summer, which was attributed to the formation of 233 photochemical reaction under strong UV radiation and the weak titration of nitrogen 234 oxides. Meanwhile, the concentrations of SO₂ ($8.37\pm0.79 \ \mu g \ m^{-3}$) in summer was also 235 higher than that $(2.63 \pm 1.95 \ \mu g \ m^{-3})$ in winter, mainly attributed to the influence of coal 236 combustion and ship emissions. The monitoring site was located approximately 15 km 237 238 away from Xiamen port area and a coal-fired power plant (4×300 kW) in the south. Southerly winds were prevailed in summer, which might cause the relative high 239 concentration of SO₂ in the monitoring site. 240



241



243 during the sampling period

244 *3.2 Temporal variations of SOA tracers* and estimated *SOC*

245 Temporal variations of individual SOA tracer are shown in Fig.S1. The average

concentrations of total SOA tracers in winter and summer were 37.3 and 111.3 ng m⁻³,

respectively. The predominance of SOA_M (26.6 ng m⁻³), followed by ASOA (4.60 ng 247 m⁻³), SOA_I (4.35 ng m⁻³) and SOA_C (1.76 ng m⁻³) was observed in winter while SOA_I 248 (54.4 ng m^{-3}) and SOA_M (47.8 ng m^{-3}) in summer were the main contributors to total 249 SOA tracers, followed by ASOA (6.64 ng m^{-3}) and SOA_C (2.45 ng m^{-3}). In summer, 250 BSOA tracers showed much higher concentrations in the daytime (149.3 ng m^{-3}) than 251 in the nighttime (60.1 ng m^{-3}), while inverse results were observed in winter (30.4 ng 252 m^{-3} and 35.0 ng m^{-3} in the daytime and nighttime, respectively). As shown in Table S2, 253 in summer, SOA_I in the daytime ranged from 21.3 to 293.2 ng m⁻³ (average of 254 82.0 ± 66.2 ng m⁻³) and the concentrations of SOA_I ranging from 6.81 to 110.1 ng m⁻³ 255 (average of 26.8 ± 24.6 ng m⁻³) were observed in the nighttime. However, in winter, the 256 concentrations of isoprene SOA tracers in the daytime ranging from 1.36 to 11.1 ng m⁻³ 257 (average of 3.79 ± 2.37 ng m⁻³) were lower than those (average of 4.91 ± 3.75 ng m⁻³) in 258 the nighttime. As shown in Fig. 2, diurnal variations of SOAM, SOAI, CPA and DHOPA 259 tracers in summer showed high levels in the afternoon (12:00-16:00 CST), due to the 260 impacts of beneficial photochemical oxidation conditions caused by high temperature 261 and strong UV radiation. The related SOA tracers were consisted with the emissions of 262 their precursors including biogenic and anthropogenic VOCs, similar to our previous 263 studies (Hong et al., 2019; Liu et al., 2020). However, the SOA tracers in winter showed 264 the lowest concentrations in the morning (8:00–12:00 CST), related with the favorable 265 dispersion conditions caused by the increasing planetary boundary layer height (BLH) 266 (Fig.1). Levoglucosan (LEV), a typical tracer of biomass burning, similar seasonal and 267 diurnal trend to other tracers was observed. However, LEV may not be as stable in the 268 atmosphere, especially under high relative humidity conditions (Hoffmann et al., 2010). 269 270 In this study, maybe, it's hard to reflect the real concentration of LEV. A correlation of CPA with LEV was carried out (Fig.S2), just to discuss the impacts of biomass burning 271 on the distribution of CPA tracers through local or long-range transport. Totally, high 272 concentrations of BSOA tracers was found in the daytime and in summer, indicating 273 the effects of temperature on biogenic VOCs emissions and their photochemical 274 275 oxidations. And the concentrations of BSOA tracers in winter increased in the nighttime,



276 due to the changing of nocturnal boundary layer.

Figure 2. Diurnal variation of individual SOA tracer during the wintertime (A,
C, and E) and summertime (B, D, and F)

As shown in Fig.S2a, b, SOA tracers-based SOC in winter and summer was estimated. The concentrations of SOC in summer was higher than that in winter, attributed to the increase of flourishing vegetation emissions and photochemical reactions under high temperature and strong solar radiation conditions. For individual SOA tracer, the

concentrations of monoterpene-derived SOC was comparable to the toluene-derived 286 SOC, which were higher than isoprene-derived SOC and β -caryophyllene-derived SOC. 287 An obvious trend of diurnal variations of isoprene-derived SOC in summer was 288 observed, which was consistent with the diurnal pattern of isoprene concentration 289 (Fig.S3). However, no similar trend was found in winter, attributed to the influence of 290 291 low temperature on inhibiting the emissions of isoprene from various kinds of plants. In addition, the toluene, monoterpene, isoprene and β -caryophyllene-derived SOC in 292 293 summer accounted for 40.0%, 39.2%, 15.7% and 5.1% of the total SOC, respectively 294 (Fig.S2c, d). However, in winter, the percentages of toluene, monoterpene, isoprene and β -caryophyllene-derived SOC were 47.2%, 42.1%, 3.2% and 7.6%, respectively. 295 The percentages of isoprene-derived SOC estimated from different precursors varied 296 297 significantly among the seasons. High temperature enhanced the emissions of isoprene, and strong solar radiation favored the formation of isoprene SOA tracers, contributing 298 to the highest isoprene-derived SOC percentage in summer (Ding et al., 2014). And the 299 highest percentages of toluene-derived SOC (47.2%) in winter were related with 300 301 anthropogenic emissions and adverse diffusion conditions.

302 3.3 Atmospheric process indication of BSOA tracers

As shown in Fig.3, percentages of different types of SOA tracers in winter and 303 summer were calculated. In summer, the monoterpene, isoprene, toluene and β -304 305 caryophyllene SOA tracers accounted for 45.8%, 45.6%, 6.2% and 2.3% of the total SOA tracers, respectively. However, in winter, the percentages of monoterpene, 306 isoprene, toluene and β -caryophyllene SOA tracers were 70.1%, 14.0%, 11.0% and 307 4.9%, respectively. The percentage of SOA_I tracers decreased sharply, due to the 308 309 impacts of temperature on isoprene emissions, which was consisted with our previous 310 findings (Hong et al., 2019). Meanwhile, the concentrations of SOA_M tracers were the largest in both seasons, due to a large amount of monoterpene emissions from the 311 related plant species. Xiamen, an international garden city, located in coastal area of 312 southeastern China. Monoterpene, such as α/β -pinene, is mostly emitted by coniferous 313 plant and most flowers and fruits, while isoprene originates from broad-leaved trees 314



and deciduous plants (Ding et al., 2014; Shrivastava et al., 2017; Yang et al., 2021).

Figure 3. Percentages of isoprene, monoterpene, β-caryophyllene and toluene
SOA tracers in winter (a) and summer (b)

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The first (PA and PNA) and later generation (HGA, AGA, HDMGA and MBTCA) 319 products were used to evaluate the aging degree of SOA_M (Ding et al., 2014; Hong et 320 al., 2019). In this study, HGA (32.2%) was the major component of α/β -pinene tracers, 321 followed by PA (30.5%), PNA (21.8%), HDMGA (7.3%), MBTCA (6.8%), and AGA 322 (1.5%). The percentage of PA and PNA were much higher than those in mountainous 323 background areas (PA: 9% and PNA: 3%)(Hong et al., 2019), suggesting the 324 325 contribution of preliminary products to SOA in urban areas. As shown in Fig.3, the percentages of PA and PNA in winter (21.8% and 14.2%) were higher than those in 326 summer (14.2% and 10.7%). Reacted with atmospheric oxidants including O₃ and OH, 327 PA and PNA were transformed into MBTCA (Offenberg et al., 2007). This is the reason 328 why the proportions of PA and PNA had a significant decreasing trend from winter to 329 summer. The ratio of MBTCA/(PA+PNA) in summer and winter were 0.16±0.09 and 330 0.12±0.07, respectively, which also proved the impacts of atmospheric oxidation 331 capacity on the aging degree of SOA_M. In addition, the ratio of HGA/MBTCA could 332 333 be used to distinguish the contribution of α -pinene or β -pinene to the SOA_M formation 334 (Jaoui et al., 2005; Ding et al., 2014). Low ratio of HGA/MBTCA (~1.0) showed that α -pinene was the major precursor for SOA_M (Lewandowski et al., 2013). The ratio of 335

HGA/MBTCA with an average of 5.78 in Xiamen was high, suggesting the contribution
of β-pinene to SOA_M.

338 As shown in Fig.3, MTLs and C5 alkene triols were the main components of the total SOA_I, with an average percentage of 68.0±14.9%, indicating a low-NOx 339 environment (Ding et al., 2014; Liu et al., 2020). In summer, the percentages of MTLs 340 and C5 alkene triols to the total SOA tracers in summer (21.8% and 14.2%) were 341 obviously higher than those in winter (4.2% and 4.3%). This was consisted with the 342 fact that the concentrations of NO₂ (14.8 \pm 7.46 µg m⁻³) in summer was significantly 343 lower than that $(32.7\pm32.6 \,\mu g \,m^{-3})$ in winter. Previous studies found that MTLs and C5 344 alkene triols were formed by the OH and HO₂ radicals via the HO₂ channel under low-345 NOx conditions (Surratt et al., 2010). C5 alkene triols are mainly produced by acid 346 catalyzed reaction of Isoprene Epoxydiols (IEPOX) in the gas phase, while MTLs are 347 formed by ring opening products of IEPOX (Surratt et al., 2007; Surratt et al., 2010). 348 And the ozonolysis of isoprene was also an important pathway for MTLs in the 349 350 presence of acid sulfate aerosols (Riva et al., 2016).

351 CPA, the typical tracer of sesquiterpenes, is formed by the photooxidation of β caryophyllene (Jaoui et al., 2007). As shown in Fig.3, CPA in winter and summer 352 accounted for 5.0% and 2.3% of the total SOA tracers, respectively. This is because 353 that the percentage of SOA_I has significant increase in summer. And the concentrations 354 of CPA ($2.5\pm2.0 \text{ ng m}^{-3}$) in summer were higher than that ($1.7\pm0.8 \text{ ng m}^{-3}$) in winter, 355 probably attributed to the emissions of β -caryophyllene driven by temperature and solar 356 radiation. The CPA has a good correlation with DHOPA in summer (Fig.S4), 357 suggesting the influence of photochemical oxidation (Liu et al., 2020). However, the 358 359 CPA were not correlated with LEV in both seasons, reflecting the limited contribution 360 of biomass burning (Zhang et al., 2019c).

361 *3.4 Impacts of aerosol acidity on BSOA formation*

Aerosol acidity (pH) was an important factor on SOA formation (Surratt et al.,
2007; Offenberg et al., 2009; Zhang et al., 2019b; Zhang et al., 2019d). Time series of

aerosol pH calculated by ISORROPIA II is shown in Fig.4. The PM_{2.5} in Xiamen was

moderately acidic with daily pH range from 3.68 to 4.67. The highest aerosol pH was
observed in winter, and the lowest pH in summer. This is with similar seasonal trend,
closing to the Yangtze River Delta (YRD) region, but obviously lower levels than those
in NCP cities of China (Zhou et al., 2021). In general, the aerosol pH in Chinese cities
were higher than those in US and European.





Figure 4. Diurnal variations of aerosol acidity (pH) during the wintertime (a)
and summertime (b) period (The boxes with error bars represent the 10th, 25th,
75th, and 90th percentiles)

374 A declining trend pH during the daytime was observed (Fig. 4), which was related to the changes of chemical compositions and environmental conditions. The aerosol pH 375 levels (~3 to 6) was related with a shift from sulfate- to nitrate-dominated aerosols (Guo 376 377 et al., 2017). According to the multiphase buffer theory, the peak buffer pH (pKa*) 378 regulated the aerosol pH, and temperature could obviously cause the variation of aerosol pH (Zheng et al., 2020). To further discuss the impacts of aerosol acidity on 379 380 BSOA formation in coastal city, we analyzed the relationship between BSOA tracers and seed particles with different pH and liquid water content (LWC) (Fig. S5 and Table 381 382 1).

Season	SOA tracer	Ηd	LWC	ONOH	PM _{2.5}	CI-	NO ₃ -	$SO4^{2-}$	$\rm NH_3$	SO_2	NO_2	Ox	Т	RH	UV
	C5	.584**	.701**	.534**	**069.	.569**	.710**	.663**	.705**	0.308	.353*	0.203	.361*	0.140	0.200
	MTLs	.590**	.705**	.431*	.665**	.639**	.707**	.651**	.757**	0.185	0.229	0.098	.353*	0.295	-0.068
	MGA	.390*	.707**	0.261	.668**	0.081	.758**	.572**	0.284	0.172	0.123	.374*	.377*	-0.019	0.238
	PA	.432*	.403**	.463**	.407**	.481*	.416*	.488*	.440*	.446*	0.241	-0.193	.319*	-0.205	0.145
WINTER	PNA	.489**	.579**	0.311	.459**	.516**	.573**	.533**	.543**	0.08	0.071	-0.101	0.121	.337*	-0.122
(n=39)	HGA	.443*	.829**	.352*	.834**	.600**	.847**	.754**	.641**	0.275	0.299	.451**	.451**	0.043	0.210
	MBTCA	.433*	.678**	.447**	.670**	.435*	.733**	.589**	.710**	.327*	0.253	.492**	.552**	-0.158	0.317
	HDMGA	.421*	.876**	.401*	.867**	.631**	.884**	.813**	.643**	.335*	.321*	.526**	.485**	-0.049	0.327
	AGA	.570**	.575**	.370*	.488**	.577**	.566**	.544**	.731**	0.126	0.181	0.019	0.279	0.298	-0.122
	CPA	0.212	.462**	-0.068	.452**	.483**	.437*	.419*	0.255	-0.15	-0.170	0.016	0.079	0.200	-0.144
	C5	495**	.425**	0.160	.622**	340*	0.268	.625**	.436**	0.254	0.025	.649**	.573**	529**	0.247
	MTLs	551**	0.131	0.055	0.272	439**	0.131	.428**	.304*	0.089	-0.278	.550**	$.610^{**}$	594**	0.263
	MGA	540**	0.029	0.116	0.132	403**	0.066	.472**	0.270	0.096	410**	.443**	.633**	668**	.382*
	PA	633**	.483**	.601**	.461**	-0.135	.541**	.502**	.405*	0.037	0.238	.456**	.626**	558**	.400*
SUMMER	PNA	664**	.616**	.387**	.812**	389**	.450**	.784**	.503**	0.269	.294*	.769**	.718**	631**	.404*
(n=50)	HGA	607**	.612**	.299*	.836**	384**	.447**	.770**	.539**	.316*	0.272	.808**	.670**	599**	0.322
	MBTCA	752**	.415**	0.237	.577**	382**	.359*	.636**	.501**	0.201	-0.052	.712**	.852**	816**	.588**
	HDMGA	525**	.618**	.299*	.833**	342*	.408**	.768**	.488**	.358*	.365**	.746**	.574**	500**	0.240
	AGA	684**	.592**	.447**	.766**	334*	.479**	.735**	.435**	0.244	0.271	.694**	.720**	634**	.477**
	CPA	552**	.625**	.441**	.780**	280*	.453**	.763**	.307*	.299*	.503**	.611**	.529**	458**	0.305
*/**Correlatio	n coefficients wi	th an asteris.	k indicate st	tatistically si	ignificant re	lationships :	at $a = 0.05$,	and two ast	erisks mean	ı significant	at $a = 0.01$.				

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387 In Table 1, the BSOA tracers was linearly correlated with aerosol acidity (pH) and SO₄²⁻. In summer, BSOA tracers in the particle phase were found to increase with 388 increasing acidity, which was attributed to the presence of acid catalyzed aerosols. For 389 example, isoprene SOA tracers is mainly formed through acid-catalyzed reactive uptake 390 of isoprene-derived epoxydiols (IEPOX) onto sulfate aerosol particles. In our previous 391 studies, we have reported that high concentration of MTLs was related with sulfate, 392 which could significantly promote the formation of isoprene-SOA tracers (Liu et al., 393 2020). Other studies also found that sulfate could increase the BSOA production by 394 395 promoting acid-catalyzed ring-opening reactions (Xu et al., 2015). In contrast, positive correlations between BSOA tracers and aerosol pH in winter were observed, indicating 396 that the formation of BSOA was predominantly enhanced by other factors, except for 397 the aerosol acidity. The aerosol pH in winter was higher than those in summer, probably 398 due to the influence of nitrate-dominated aerosols. Also, the aged aerosols through long-399 range transport might result in the increase of BSOA tracers and aerosol pH. 400

In addition, positive correlation between BSOA tracers and LWC was observed 401 (Table 1), probably attributed to the effects of the LWC on determining the peak buffer 402 pH (pKa*). Zheng et al. (2020) reported that the buffering effect of ammonia suppresses 403 404 the contribution of different chemical compositions in aerosol particles, making LWC the primary determinant of aerosol pH. Other studies have demonstrated that the uptake 405 406 coefficient of first-generation oxidation products, especially for carbonyl compounds, might depend on RH (Luo et al., 2019). Meanwhile, high LWC could reduce the aerosol 407 408 particle viscosity, which was benefit to the generation of the reactive intermediate such as IEPOX, or other oxidation products of VOC into aqueous-phase of aerosol particles, 409 410 thereby promoting the formation of BSOA (Zhang et al., 2019b; Zhang et al., 2019d).

411 3.5 Impacts of chlorine on BSOA formation

412 Halogen radicals (Cl, Br and I) originated from sea salt aerosol (SSA) have an important role in tropospheric oxidants chemistry (Wang et al., 2021c). In this study, 413 chlorine depletion was frequently observed in summer (Fig.5b), indicating that HCl can 414 be formed through acid displacement of sea salt aerosol Cl⁻ by H₂SO₄ and HNO₃ 415 produced from anthropogenic emissions of SO₂ and NOx. Moreover, concentrations of 416 the total SOA tracers were positively correlated with HCl (Fig.5a), suggesting the 417 enhancement of SOA precursors transformation. Previous studies have found that Cl-418 initiated VOC oxidations could contribute to the formation of SOA (Wang and Ruiz, 419

421



422 Figure 5. Correlations of total SOA tracers and HCl (a) and chlorine depletion
423 (b) in different seasons

Under ammonia-rich conditions, HCl partitioned into the aqueous particulate 424 phase mostly took place, and chlorine ions could affect aqueous oxidation of secondary 425 organic compounds (Xu et al., 2021). As shown in Table 1, most of SOA tracers in 426 winter were correlated with the concentrations of chlorine ions in PM_{2.5}, while inverse 427 results were observed in summer. In winter, the dominant wind direction is northeast 428 429 (Fig.6), and chlorine mainly come from continental polluted air mass, such as industrial 430 and combustion emissions. So, anthropogenic pollutants through long-range transport might cause the enhancement of SOA tracer concentrations at the monitoring site. 431 432 However, in summer, negative correlations of BSOA tracers and chlorine ions in PM_{2.5} was found, probably due to the influence of chlorine depletion. As shown in Fig. 6, the 433 434 dominant wind direction is southerly, and chlorine mainly originated from the spray of 435 sea salt.





Figure 6. Backward trajectories analyses during the winter (a) and summertime(b)

439 *3.6. Enhanced formation of BSOA by anthropogenic emissions*

Recent studies had indicated that anthropogenic emissions might affect SOA formation through multiple chemical processes, based on laboratory studies and field observations (Kari et al., 2019; Shrivastava et al., 2019; Zhang et al., 2019c; Cheng et al., 2021; Xu et al., 2021). In this study, we conducted the correlation analysis of individual SOA tracers and Ox (= O_3 +NO₂), HONO, OH, SO₂, NH₃, PM_{2.5}, sulfate, nitrate, as well as meteorological parameters (including T, RH and UV) (Table 1).

Most of SOA tracers have a significant positive correlation with NH₃, suggesting 446 an enhancement effect on the formation of SOA (Table 1). NH₃ could affect the SOA 447 yields through both gas-phase and heterogeneous reactions (Na et al., 2007; Ma et al., 448 2018; Hao et al., 2020). Gas-phase reaction between NH_3 and organic acids (such as 449 PA and PNA) produced ammonium salts in the particle phase, which contributed to the 450 increased SOA formation. However, not all gas-phase organic acids (e.g., MGA and 451 452 pyruvic acid) could demonstrate gas-to-particle conversion (Na et al., 2007). When SOA formation had ceased, the addition of excessive NH₃ would result in the rapid 453 454 decomposition of the main SOA species, due to the nucleophilic attack of NH₃ (Ma et al., 2018). 455





457 Figure 7. Relationships of SOA tracers and HONO (a) and its estimated OH (b)

As an indicator of atmospheric oxidation capacity, the tropospheric odd oxygen 458 Ox (O_3+NO_2) was calculated. As shown in Table 1, the majority of SOA tracers in 459 summer showed significant positive correlations with Ox (R>0.5, P<0.001). However, 460 in winter, a part of SOA_M tracers (e.g. HGA, MBTCA and HDMGA) were found to be 461 significantly correlated with Ox. In addition, HONO and OH radicals, another critical 462 463 indicator of atmospheric oxidation capacity, was also discussed. In this study, the concentration of OH radicals calculated from HONO in summer was higher than those 464 in winter. In summer, the SOA_I tracers was correlated with OH radicals (Fig.7b), 465 466 consisted with previous findings that OH radicals could promote the formation of SOA (Sarrafzadeh et al., 2016; Liu et al., 2019; Song et al., 2019; Zhang et al., 2019a). Due 467 468 to its photolysis to produce OH radicals during the daytime, HONO could facilitate SOA formation. In winter, the concentrations of SOA_I, SOA_M and ASOA tracers were 469 470 correlated with HONO (Fig.7a). These results indicated high concentrations of HONO 471 and sufficient ultraviolet radiation could enhance the photochemical reactions of VOCs. 472 Which was consisted with our previous results on the formation of peroxyacetyl nitrate (PAN) (Hu et al., 2020). As for T and UV, it exhibited significantly positive correlations 473 with the related SOA tracers, especially in summer. These results suggested that SOA 474 tracers were produced from the photo-oxidation of VOC precursors (Cheng et al., 2021). 475 In addition, the SOA tracers were significantly positive correlated with PM_{2.5} and 476 its components including NO₃- and SO₄²⁻. In coastal cities of southeastern China, with 477

the development of rapid urbanization, air pollution caused by motor vehicles and
industrial emissions is becoming more frequent in winter (Wu et al., 2020). The Xiamen
port is one of the top 10 ports in China, resulting the impacts of ship emissions and port

481 activities on ambient air quality (Xu et al., 2018), and the numbers of motor vehicles increased sharply in recent years. We also found that the 90th percentile of maximum 482 daily average 8h (MDA8) O₃ concentrations in Xiamen was significantly increased 483 from 2015 to 2020 (Fig. S6). During the past several years, the elevated secondary 484 inorganic components, including NO₃-, SO₄²⁻ and NH₄⁺, accounted for 40-50% of the 485 total PM_{2.5}, and OM ranged from 30% to 40% (Wu et al., 2019; Hong et al., 2021). 486 These results also implied the effects of anthropogenic emissions and enhanced 487 atmospheric oxidation capacity on secondary formation of aerosol particles under 488 489 atmospheric stagnant conditions.

490 **Conclusions**

Pollution characteristics and source identification of BSOA tracers during the 491 summer and winter in coastal areas of southeastern China were investigated. The 492 average concentration of total BSOA tracers in summer was higher than that in winter, 493 with the predominance of SOA_M, followed by SOA_I and SOA_C. The BSOA tracers in 494 495 summer were predominantly produced by the influence of photochemical oxidation under relatively clean conditions. However, in winter, the formation of BSOA tracers 496 were attributed to the impacts of anthropogenic emissions and atmospheric stagnant 497 498 conditions. In addition, the results also indicated that acid-catalyzed reactive uptake onto sulfate aerosol particles enhanced the formation of BSOA in both seasons. We 499 further found that Cl-initiated VOC oxidations has potentially accelerated the 500 transformation of BSOA precursors through sea salt aerosol originated from the ocean 501 502 in summer and anthropogenic emissions in winter. This study demonstrated that the combined effects of anthropogenic pollutants and atmospheric oxidation capacity on 503 504 the formation of BSOA in coastal area.

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506 *Data Availability.* The data set related to this work can be accessed via 507 https://doi.org/10.5281/zenodo.6376025 (Hong, 2022). The details are also available 508 upon request from the corresponding author (ywhong@iue.ac.cn).

509

510 Authorship Contribution Statement. Youwei Hong and Xinbei Xu contributed equally 511 to this work. Youwei Hong designed and wrote the manuscript. Xinbei Xu collected the 512 data, contributed to the data analysis. Dan Liao, Taotao Liu, Xiaoting Ji and Ke Xu 513 performed modeling analyses and data analysis. Jinsheng Chen supported funding of observation and research. Chunyang Liao, Ting Wang and Chunshui Lin contributed torevise the manuscript.

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Competing interests. The authors declare that they have no conflict of interest.

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537 **Reference**

- Charan, S. M., Huang, Y., and Seinfeld, J. H.: Computational Simulation of Secondary
 Organic Aerosol Formation in Laboratory Chambers, Chem. Rev., 119, 11912-11944,
 10.1021/acs.chemrev.9b00358, 2019.
- 541 Cheng, Y., Ma, Y., and Hu, D.: Tracer-based source apportioning of atmospheric organic
 542 carbon and the influence of anthropogenic emissions on secondary organic aerosol
 543 formation in Hong Kong, Atmos. Chem. Phys., 21, 10589-10608, 10.5194/acp-21544 10589-2021, 2021.
- 545 Dhulipala, S. V., Bhandari, S., and Hildebrandt Ruiz, L.: Formation of oxidized organic
 546 compounds from Cl-initiated oxidation of toluene, Atmospheric Environment, 199, 265547 273, 10.1016/j.atmosenv.2018.11.002, 2019.
- 548 Ding, X., He, Q.-F., Shen, R.-Q., Yu, Q.-Q., and Wang, X.-M.: Spatial distributions of
 549 secondary organic aerosols from isoprene, monoterpenes, beta-caryophyllene, and
 550 aromatics over China during summer, Journal of Geophysical Research-Atmospheres,
 551 119, 11877-11891, 10.1002/2014jd021748, 2014.

- Fountoukis, C., and Nenes, A.: ISORROPIA II: a computationally efficient thermodynamic equilibrium model for K^+ -Ca²⁺-Mg²⁺-NH₄⁺ -Na⁺-SO₄²⁻ -NO₃⁻-Cl⁻-H₂O aerosols, Atmos. Chem. Phys., 7, 4639-4659, 10.5194/acp-7-4639-2007, 2007.
- Fu, P., Kawamura, K., Chen, J., and Barrie, L. A.: Isoprene, Monoterpene, and Sesquiterpene
 Oxidation Products in the High Arctic Aerosols during Late Winter to Early Summer,
 Environmental Science & Technology, 43, 4022-4028, 10.1021/es803669a, 2009.
- Guo, H., Sullivan, A. P., Campuzano-Jost, P., Schroder, J. C., Lopez-Hilfiker, F. D.,
 Dibb, J. E., Jimenez, J. L., Thornton, J. A., Brown, S. S., Nenes, A., and Weber,
 R. J.: Fine particle pH and the partitioning of nitric acid during winter in the
 northeastern United States, Journal of Geophysical Research: Atmospheres, 121,
 10,355-310,376, https://doi.org/10.1002/2016JD025311, 2016.
- Guo, H., Weber, R. J., and Nenes, A.: High levels of ammonia do not raise fine particle pH
 sufficiently to yield nitrogen oxide-dominated sulfate production, Scientific Reports, 7,
 12109, 10.1038/s41598-017-11704-0, 2017.
- Hallquist, M., Wenger, J. C., Baltensperger, U., Rudich, Y., Simpson, D., Claeys, M.,
 Dommen, J., Donahue, N. M., George, C., Goldstein, A. H., Hamilton, J. F., Herrmann,
 H., Hoffmann, T., Iinuma, Y., Jang, M., Jenkin, M. E., Jimenez, J. L., Kiendler-Scharr,
 A., Maenhaut, W., McFiggans, G., Mentel, T. F., Monod, A., Prevot, A. S. H., Seinfeld,
 J. H., Surratt, J. D., Szmigielski, R., and Wildt, J.: The formation, properties and impact
 of secondary organic aerosol: current and emerging issues, Atmospheric Chemistry and
 Physics, 9, 5155-5236, 10.5194/acp-9-5155-2009, 2009.
- Hao, L., Kari, E., Leskinen, A., Worsnop, D. R., and Virtanen, A.: Direct contribution of
 ammonia to α-pinene secondary organic aerosol formation, Atmos. Chem. Phys., 20,
 14393-14405, 10.5194/acp-20-14393-2020, 2020.
- Hoffmann, D., Tilgner, A., Iinuma, Y., and Herrmann, H.: Atmospheric Stability of
 Levoglucosan: A Detailed Laboratory and Modeling Study, Environmental Science &
 Technology, 44, 694-699, 10.1021/es902476f, 2010.
- Hong, Y., Xu, X., Liao, D., Zheng, R., Ji, X., Chen, Y., Xu, L., Li, M., Wang, H., Xiao, H.,
 Choi, S.-D., and Chen, J.: Source apportionment of PM2.5 and sulfate formation during
 the COVID-19 lockdown in a coastal city of southeast China, Environmental Pollution,
 286, 117577, https://doi.org/10.1016/j.envpol.2021.117577, 2021.
- Hong, youwei. (2022). Dataset for ACP by Hong et al., 2022 [Data set]. Zenodo.
 https://doi.org/10.5281/zenodo.6376025
- Hong, Z., Zhang, H., Zhang, Y., Xu, L., Liu, T., Xiao, H., Hong, Y., Chen, J., Li, M., Deng,
 J., Wu, X., Hu, B., and Chen, X.: Secondary organic aerosol of PM2.5 in a mountainous
 forest area in southeastern China: Molecular compositions and tracers implication,
 Science of the Total Environment, 653, 496-503, 10.1016/j.scitotenv.2018.10.370, 2019.
- Hoyle, C. R., Boy, M., Donahue, N. M., Fry, J. L., Glasius, M., Guenther, A., Hallar, A. G.,
 Hartz, K. H., Petters, M. D., Petaja, T., Rosenoern, T., and Sullivan, A. P.: A review of
 the anthropogenic influence on biogenic secondary organic aerosol, Atmospheric
 Chemistry and Physics, 11, 321-343, 10.5194/acp-11-321-2011, 2011.
- Hu, B., Liu, T., Hong, Y., Xu, L., Li, M., Wu, X., Wang, H., Chen, J., and Chen, J.:
 Characteristics of peroxyacetyl nitrate (PAN) in a coastal city of southeastern China:
 Photochemical mechanism and pollution process, Science of the Total Environment,
 719, 10.1016/j.scitotenv.2020.137493, 2020.
- Hu, W., Palm, B. B., Day, D. A., Campuzano-Jost, P., Krechmer, J. E., Peng, Z., de Sa, S. S.,
 Martin, S. T., Alexander, M. L., Baumann, K., Hacker, L., Kiendler-Scharr, A., Koss, A.
 R., de Gouw, J. A., Goldstein, A. H., Seco, R., Sjostedt, S. J., Park, J.-H., Guenther, A.
 B., Kim, S., Canonaco, F., Prevot, A. S. H., Brune, W. H., and Jimenez, J. L.: Volatility
 and lifetime against OH heterogeneous reaction of ambient isoprene-epoxydiols-derived

603 secondary organic aerosol (IEPOX-SOA), Atmospheric Chemistry and Physics, 16, 604 11563-11580, 10.5194/acp-16-11563-2016, 2016. Jaoui, M., Kleindienst, T. E., Lewandowski, M., Offenberg, J. H., and Edney, E. O.: 605 606 Identification and quantification of aerosol polar oxygenated compounds bearing carboxylic or hydroxyl groups, 2. Organic tracer compounds from monoterpenes. 607 Environmental Science & Technology, 39, 5661-5673, 10.1021/es048111b, 2005. 608 Jaoui, M., Lewandowski, M., Kleindienst, T. E., Offenberg, J. H., and Edney, E. O.: β-609 caryophyllinic acid: An atmospheric tracer forβ-caryophyllene secondary organic 610 611 aerosol, Geophysical Research Letters, 34, 10.1029/2006gl028827, 2007. Kari, E., Hao, L. Q., Ylisirnio, A., Buchholz, A., Leskinen, A., Yli-Pirila, P., Nuutinen, I., 612 Kuuspalo, K., Jokiniemi, J., Faiola, C. L., Schobesberger, S., and Virtanen, A.: Potential 613 614 dual effect of anthropogenic emissions on the formation of biogenic secondary organic 615 aerosol (BSOA), Atmospheric Chemistry and Physics, 19, 15651-15671, 10.5194/acp-19-15651-2019, 2019. 616 Kleindienst, T. E., Jaoui, M., Lewandowski, M., Offenberg, J. H., Lewis, C. W., Bhave, P. V., 617 and Edney, E. O.: Estimates of the contributions of biogenic and anthropogenic 618 hydrocarbons to secondary organic aerosol at a southeastern US location, Atmospheric 619 Environment, 41, 8288-8300, 10.1016/j.atmosenv.2007.06.045, 2007. 620 621 Lewandowski, M., Piletic, I. R., Kleindienst, T. E., Offenberg, J. H., Beaver, M. R., Jaoui, M., 622 Docherty, K. S., and Edney, E. O.: Secondary organic aerosol characterisation at field 623 sites across the United States during the spring-summer period, International Journal of Environmental Analytical Chemistry, 93, 1084-1103, 10.1080/03067319.2013.803545, 624 625 2013. Liu, S., Tsona, N. T., Zhang, Q., Jia, L., Xu, Y., and Du, L.: Influence of relative humidity on 626 627 cyclohexene SOA formation from OH photooxidation, Chemosphere, 231, 478-486, 10.1016/j.chemosphere.2019.05.131, 2019. 628 629 Liu, S., Huang, D., Wang, Y., Zhang, S., Wu, C., Du, W., and Wang, G.: Synergetic effect of 630 NH₃ and NOx on the production and optical absorption of secondary organic aerosol 631 formation from toluene photooxidation, Atmos. Chem. Phys. Discuss., 2021, 1-38, 10.5194/acp-2021-560, 2021. 632 633 Liu, T., Hu, B., Xu, X., Hong, Y., Zhang, Y., Wu, X., Xu, L., Li, M., Chen, Y., Chen, X., and Chen, J.: Characteristics of PM2.5-bound secondary organic aerosol tracers in a coastal 634 635 city in Southeastern China: Seasonal patterns and pollution identification, Atmospheric Environment, 237, 10.1016/j.atmosenv.2020.117710, 2020. 636 Lopez-Hilfiker, F. D., Mohr, C., D'Ambro, E. L., Lutz, A., Riedel, T. P., Gaston, C. J., Iyer, 637 S., Zhang, Z., Gold, A., Surratt, J. D., Lee, B. H., Kurten, T., Hu, W. W., Jimenez, J., 638 Hallquist, M., and Thornton, J. A.: Molecular Composition and Volatility of Organic 639 Aerosol in the Southeastern US: Implications for IEPDX Derived SOA, Environmental 640 641 Science & Technology, 50, 2200-2209, 10.1021/acs.est.5b04769, 2016. Lowes, S., Jersey, J., Shoup, R., Garofolo, F., Savoie, N., Mortz, E., Needham, S., Caturla, M. 642 C., Steffen, R., Sheldon, C., Hayes, R., Samuels, T., Di Donato, L., Kamerud, J., 643 Michael, S., Lin, Z. P., Hillier, J., Moussallie, M., Teixeira, L. D., Rocci, M., Buonarati, 644 M., Truog, J., Hussain, S., Lundberg, R., Breau, A., Zhang, T. Y., Jonker, J., Berger, N., 645 646 Gagnon-Carignan, S., Nehls, C., Nicholson, R., Hilhorst, M., Karnik, S., de Boer, T., Houghton, R., Smith, K., Cojocaru, L., Allen, M., Harter, T., Fatmi, S., Sayyarpour, F., 647 Vija, J., Malone, M., and Heller, D.: Recommendations on: internal standard criteria, 648 stability, incurred sample reanalysis and recent 483s by the Global CRO Council for 649 Bioanalysis, Bioanalysis, 3, 1323-1332, 10.4155/Bio.11.135, 2011. 650 Luo, H., Jia, L., Wan, Q., An, T., and Wang, Y.: Role of liquid water in the formation of O-3 651 and SOA particles from 1,2,3-trimethylbenzene, Atmospheric Environment, 217, 652 10.1016/j.atmosenv.2019.116955, 2019. 653 Lv, S., Wang, F., Wu, C., Chen, Y., Liu, S., Zhang, S., Li, D., Du, W., Zhang, F., Wang, H., 654 655 Huang, C., Fu, Q., Duan, Y., and Wang, G.: Gas-to-Aerosol Phase Partitioning of

656 Atmospheric Water-Soluble Organic Compounds at a Rural Site in China: An Enhancing Effect of NH3 on SOA Formation, Environmental Science & Technology, 657 10.1021/acs.est.1c06855, 2022. 658 Ma, Q., Lin, X. X., Yang, C. G., Long, B., Gai, Y. B., and Zhang, W. J.: The influences of 659 ammonia on aerosol formation in the ozonolysis of styrene: roles of Criegee intermediate 660 reactions, Roy Soc Open Sci, 5, ARTN 17217110.1098/rsos.172171, 2018. 661 Mahilang, M., Deb, M. K., and Pervez, S.: Biogenic secondary organic aerosols: A review on 662 formation mechanism, analytical challenges and environmental impacts, Chemosphere, 663 664 262, 10.1016/j.chemosphere.2020.127771, 2021. McFiggans, G., Mentel, T. F., Wildt, J., Pullinen, I., Kang, S., Kleist, E., Schmitt, S., 665 Springer, M., Tillmann, R., Wu, C., Zhao, D. F., Hallquist, M., Faxon, C., Le Breton, M., 666 667 Hallquist, A. M., Simpson, D., Bergstrom, R., Jenkin, M. E., Ehn, M., Thornton, J. A., Alfarra, M. R., Bannan, T. J., Percival, C. J., Priestley, M., Topping, D., and Kiendler-668 Scharr, A.: Secondary organic aerosol reduced by mixture of atmospheric vapours, 669 670 Nature, 565, 587-593, 10.1038/s41586-018-0871-y, 2019. Na, K., Song, C., Switzer, C., and Cocker, D. R.: Effect of Ammonia on Secondary Organic 671 Aerosol Formation from α -Pinene Ozonolysis in Dry and Humid Conditions, 672 Environmental Science & Technology, 41, 6096-6102, 10.1021/es061956y, 2007. 673 674 Newland, M. J., Bryant, D. J., Dunmore, R. E., Bannan, T. J., Acton, W. J. F., Langford, B., 675 Hopkins, J. R., Squires, F. A., Dixon, W., Drysdale, W. S., Ivatt, P. D., Evans, M. J., Edwards, P. M., Whalley, L. K., Heard, D. E., Slater, E. J., Woodward-Massey, R., Ye, 676 C., Mehra, A., Worrall, S. D., Bacak, A., Coe, H., Percival, C. J., Hewitt, C. N., Lee, J. 677 678 D., Cui, T., Surratt, J. D., Wang, X., Lewis, A. C., Rickard, A. R., and Hamilton, J. F.: Low-NO atmospheric oxidation pathways in a polluted megacity, Atmos. Chem. Phys., 679 680 21, 1613-1625, 10.5194/acp-21-1613-2021, 2021. Offenberg, J. H., Lewis, C. W., Lewandowski, M., Jaoui, M., Kleindienst, T. E., and Edney, 681 682 E. O.: Contributions of toluene and alpha-pinene to SOA formed in an irradiated 683 toluene/alpha-pinene/NOx/air mixture: Comparison of results using C-14 content and 684 SOA organic tracer methods, Environmental Science & Technology, 41, 3972-3976, 10.1021/es070089+, 2007. 685 686 Offenberg, J. H., Lewandowski, M., Edney, E. O., Kleindienst, T. E., and Jaoui, M.: Influence of Aerosol Acidity on the Formation of Secondary Organic Aerosol from Biogenic 687 688 Precursor Hydrocarbons, Environmental Science & Technology, 43, 7742-7747, 10.1021/es901538e, 2009. 689 690 Palmer, P. I., Marvin, M. R., Siddans, R., Kerridge, B. J., and Moore, D. P.: Nocturnal survival of isoprene linked to formation of upper tropospheric organic aerosol, Science, 691 692 375, 562-566, doi:10.1126/science.abg4506, 2022. 693 Reid, J. P., Bertram, A. K., Topping, D. O., Laskin, A., Martin, S. T., Petters, M. D., Pope, F. 694 D., and Rovelli, G.: The viscosity of atmospherically relevant organic particles, Nature Communications, 9, 10.1038/s41467-018-03027-z, 2018. 695 696 Riva, M., Budisulistiorini, S. H., Zhang, Z., Gold, A., and Surratt, J. D.: Chemical characterization of secondary organic aerosol constituents from isoprene ozonolysis in 697 the presence of acidic aerosol, Atmospheric Environment, 130, 5-13, 698 699 10.1016/j.atmosenv.2015.06.027, 2016. Rumsey, I. C., Cowen, K. A., Walker, J. T., Kelly, T. J., Hanft, E. A., Mishoe, K., 700 Rogers, C., Proost, R., Beachley, G. M., Lear, G., Frelink, T., and Otjes, R. P.: 701 An assessment of the performance of the Monitor for AeRosols and GAses in 702 ambient air (MARGA): a semi-continuous method for soluble compounds, 703 Atmos. Chem. Phys., 14, 5639-5658, 10.5194/acp-14-5639-2014, 2014. 704 Sarrafzadeh, M., Wildt, J., Pullinen, I., Springer, M., Kleist, E., Tillmann, R., Schmitt, S. H., 705 Wu, C., Mentel, T. F., Zhao, D., Hastie, D. R., and Kiendler-Scharr, A.: Impact of NOx 706 707 and OH on secondary organic aerosol formation from beta-pinene photooxidation,

708	Atmospheric Chemistry and Physics, 16, 11237-11248, 10.5194/acp-16-11237-2016,
709	2016.
710	Shrivastava, M., Cappa, C. D., Fan, J., Goldstein, A. H., Guenther, A. B., Jimenez, J. L.,
711	Kuang, C., Laskin, A., Martin, S. T., Ng, N. L., Petaja, T., Pierce, J. R., Rasch, P. J.,
712	Roldin, P., Seinfeld, J. H., Shilling, J., Smith, J. N., Thornton, J. A., Volkamer, R.,
713	Wang, J., Worsnop, D. R., Zaveri, R. A., Zelenyuk, A., and Zhang, Q.: Recent advances
714	in understanding secondary organic aerosol: Implications for global climate forcing,
715	Reviews of Geophysics, 55, 509-559, 10.1002/2016rg000540, 2017.
716	Shrivastava, M., Andreae, M. O., Artaxo, P., Barbosa, H. M. J., Berg, L. K., Brito, J., Ching,
717	J., Easter, R. C., Fan, J., Fast, J. D., Feng, Z., Fuentes, J. D., Glasius, M., Goldstein, A.
718	H., Alves, E. G., Gomes, H., Gu, D., Guenther, A., Jathar, S. H., Kim, S., Liu, Y., Lou,
719	S., Martin, S. T., McNeill, V. F., Medeiros, A., de Sa, S. S., Shilling, J. E., Springston, S.
720	R., Souza, R. A. F., Thornton, J. A., Isaacman-VanWertz, G., Yee, L. D., Ynoue, R.,
721	Zaveri, R. A., Zelenyuk, A., and Zhao, C.: Urban pollution greatly enhances formation of
722	natural aerosols over the Amazon rainforest, Nature Communications, 10,
723	10.1038/s41467-019-08909-4, 2019.
724	Song, M., Zhang, C., Wu, H., Mu, Y., Ma, Z., Zhang, Y., Liu, J., and Li, X.: The influence of
725	OH concentration on SOA formation from isoprene photooxidation, Science of the Total
726	Environment, 650, 951-957, 10.1016/j.scitotenv.2018.09.084, 2019.
727	Surratt, J. D., Lewandowski, M., Offenberg, J. H., Jaoui, M., Kleindienst, T. E., Edney, E. O.,
728	and Seinfeld, J. H.: Effect of acidity on secondary organic aerosol formation from
729	isoprene, Environmental Science & Technology, 41, 5363-5369, 10.1021/es0704176,
730	2007.
731	Surratt, J. D., Chan, A. W. H., Eddingsaas, N. C., Chan, M., Loza, C. L., Kwan, A. J., Hersey,
732	S. P., Flagan, R. C., Wennberg, P. O., and Seinfeld, J. H.: Reactive intermediates
733	revealed in secondary organic aerosol formation from isoprene, Proceedings of the
734	National Academy of Sciences of the United States of America, 107, 6640-6645,
735	10.1073/pnas.0911114107, 2010.
736	Wang, D. S., and Ruiz, L. H.: Secondary organic aerosol from chlorine-initiated oxidation of
737	isoprene, Atmos. Chem. Phys., 17, 13491-13508, 10.5194/acp-17-13491-2017, 2017.
738	Wang, J., Ye, J., Zhang, Q., Zhao, J., Wu, Y., Li, J., Liu, D., Li, W., Zhang, Y., Wu, C., Xie,
739	C., Qin, Y., Lei, Y., Huang, X., Guo, J., Liu, P., Fu, P., Li, Y., Lee, H. C., Choi, H.,
740	Zhang, J., Liao, H., Chen, M., Sun, Y., Ge, X., Martin, S. T., and Jacob, D. J.: Aqueous
741	production of secondary organic aerosol from fossil-fuel emissions in winter Beijing
742	haze, Proc Natl Acad Sci U S A, 118, 10.1073/pnas.2022179118, 2021a.
743	Wang, J., Ye, J., Zhang, Q., Zhao, J., Wu, Y., Li, J., Liu, D., Li, W., Zhang, Y., Wu, C., Xie,
744	C., Qin, Y., Lei, Y., Huang, X., Guo, J., Liu, P., Fu, P., Li, Y., Lee, H. C., Choi, H.,
745	Zhang, J., Liao, H., Chen, M., Sun, Y., Ge, X., Martin, S. T., and Jacob, D. J.: Aqueous
746	production of secondary organic aerosol from fossil-fuel emissions in winter Beijing
747	haze, Proceedings of the National Academy of Sciences of the United States of America,
748	118, 10.1073/pnas.2022179118, 2021b.
749	Wang, S., Du, L., Tsona, N. T., Jiang, X., You, B., Xu, L., Yang, Z., and Wang, W.: Effect of
750	NOx and SO2 on the photooxidation of methylglyoxal: Implications in secondary
751	aerosol formation, J Environ Sci (China), 92, 151-162, 10.1016/j.jes.2020.02.011, 2020.
752	Wang, X., Jacob, D. J., Downs, W., Zhai, S., Zhu, L., Shah, V., Holmes, C. D., Sherwen, T.,
753	Alexander, B., Evans, M. J., Eastham, S. D., Neuman, J. A., Veres, P. R., Koenig, T. K.,
754	Volkamer, R., Huey, L. G., Bannan, T. J., Percival, C. J., Lee, B. H., and Thornton, J. A.:
755	Global tropospheric halogen (Cl, Br, I) chemistry and its impact on oxidants, Atmos.
756	Chem. Phys., 21, 13973-13996, 10.5194/acp-21-13973-2021, 2021c.
757	Wen, L., Chen, T., Zheng, P., Wu, L., Wang, X., Mellouki, A., Xue, L., and Wang, W.:
758	Nitrous acid in marine boundary layer over eastern Bohai Sea, China: Characteristics,
759	sources, and implications, Sci. Total Environ., 10.1016/j.scitotenv.2019.03.225, 2019.

- Wu, X., Xu, L. L., Hong, Y. W., Chen, J. F., Qiu, Y. Q., Hu, B. Y., Hong, Z. Y., Zhang, Y.
 R., Liu, T. T., Chen, Y. T., Bian, Y. H., Zhao, G. Q., Chen, J. S., and Li, M. R.: The air
 pollution governed by subtropical high in a coastal city in Southeast China: Formation
 processes and influencing mechanisms, Science of the Total Environment, 692, 11351145, 10.1016/j.scitotenv.2019.07.341, 2019.
- Wu, X., Li, M., Chen, J., Wang, H., Xu, L., Hong, Y., Zhao, G., Hu, B., Zhang, Y., Dan, Y.,
 and Yu, S.: The characteristics of air pollution induced by the quasi-stationary front:
 Formation processes and influencing factors, Science of the Total Environment, 707,
 10.1016/j.scitotenv.2019.136194, 2020.
- Xiao, Y., Wu, Z., Guo, S., He, L., Huang, X., and Hu, M.: Formation mechanism of
 secondary organic aerosol in aerosol liquid water: A review, Chinese Science Bulletin,
 65, 3118-3133, 2020.
- Xu, L., Du, L., Tsona, N. T., and Ge, M. F.: Anthropogenic Effects on Biogenic Secondary
 Organic Aerosol Formation, Advances in Atmospheric Sciences, 38, 1053-1084,
 10.1007/s00376-020-0284-3, 2021.
- Xu, L., Guo, H. Y., Boyd, C. M., Klein, M., Bougiatioti, A., Cerully, K. M., Hite, J. R.,
 Isaacman-VanWertz, G., Kreisberg, N. M., Knote, C., Olson, K., Koss, A., Goldstein, A.
 H., Hering, S. V., de Gouw, J., Baumann, K., Lee, S. H., Nenes, A., Weber, R. J., and
 Ng, N. L.: Effects of anthropogenic emissions on aerosol formation from isoprene and
 monoterpenes in the southeastern United States, Proceedings of the National Academy of
 Sciences of the United States of America, 112, 37-42, 10.1073/pnas.1417609112, 2015.
- Yang, W., Cao, J., Wu, Y., Kong, F., and Li, L.: Review on plant terpenoid emissions
 worldwide and in China, The Science of the total environment, 787, 147454-147454, 10.1016/j.scitotenv.2021.147454, 2021.
- Zhang, J., An, J., Qu, Y., Liu, X., and Chen, Y.: Impacts of potential HONO sources on the
 concentrations of oxidants and secondary organic aerosols in the Beijing-Tianjin-Hebei
 region of China, Science of the Total Environment, 647, 836-852,
 10.1016/j.scitotenv.2018.08.030, 2019a.
- Zhang, P., Chen, T., Liu, J., Liu, C., Ma, J., Ma, Q., Chu, B., and He, H.: Impacts of SO2,
 Relative Humidity, and Seed Acidity on Secondary Organic Aerosol Formation in the
 Ozonolysis of Butyl Vinyl Ether, Environmental Science & Technology, 53, 8845-8853,
 10.1021/acs.est.9b02702, 2019b.
- Zhang, Y.-Q., Chen, D.-H., Ding, X., Li, J., Zhang, T., Wang, J.-Q., Cheng, Q., Jiang, H.,
 Song, W., Ou, Y.-B., Ye, P.-L., Zhang, G., and Wang, X.-M.: Impact of anthropogenic
 emissions on biogenic secondary organic aerosol: observation in the Pearl River Delta,
 southern China, Atmospheric Chemistry and Physics, 19, 14403-14415, 10.5194/acp-1914403-2019, 2019c.
- Zhang, Y., Chen, Y., Lei, Z., Olson, N. E., Riva, M., Koss, A. R., Zhang, Z., Gold, A., Jayne,
 J. T., Worsnop, D. R., Onasch, T. B., Kroll, J. H., Turpin, B. J., Ault, A. P., and Surratt,
 J. D.: Joint Impacts of Acidity and Viscosity on the Formation of Secondary Organic
 Aerosol from Isoprene Epoxydiols (IEPDX) in Phase Separated Particles, Acs Earth and
 Space Chemistry, 3, 2646-2658, 10.1021/acsearthspacechem.9b00209, 2019d.
- Zhao, D. F., Schmitt, S. H., Wang, M. J., Acir, I. H., Tillmann, R., Tan, Z. F., Novelli, A.,
 Fuchs, H., Pullinen, I., Wegener, R., Rohrer, F., Wildt, J., Kiendler-Scharr, A., Wahner,
 A., and Mentel, T. F.: Effects of NOx and SO2 on the secondary organic aerosol
 formation from photooxidation of alpha-pinene and limonene, Atmospheric Chemistry
 and Physics, 18, 1611-1628, 10.5194/acp-18-1611-2018, 2018.
- Zheng, G., Su, H., Wang, S., Andreae, M. O., Poschl, U., and Cheng, Y.: Multiphase buffer
 theory explains contrasts in atmospheric aerosol acidity, Science, 369, 1374-+,
 10.1126/science.aba3719, 2020.
- Zhou, M., Zheng, G., Wang, H., Qiao, L., Zhu, S., Huang, D., An, J., Lou, S., Tao, S., Wang,
 Q., Yan, R., Ma, Y., Chen, C., Cheng, Y., Su, H., and Huang, C.: Long-term trends and

- drivers of aerosol pH in eastern China, Atmos. Chem. Phys. Discuss., 2021, 1-21,
- 813 10.5194/acp-2021-455, 2021.
- Zhu, J., Penner, J. E., Yu, F., Sillman, S., Andreae, M. O., and Coe, H.: Decrease in radiative
 forcing by organic aerosol nucleation, climate, and land use change, Nature
- 816 Communications, 10, 10.1038/s41467-019-08407-7, 2019.
- 817
- 818

819