

1 **Significant role of biomass burning in heavy haze**
2 **formation in Nanjing, a megacity in China: Molecular-**
3 **level insights from intensive PM_{2.5} sampling on winter**
4 **hazy days**

5 Mingjie Kang^{1,2}, Mengying Bao^{1,2,3}, Wenhui Song^{1,2}, Aduburexiati Abulimiti^{1,2},
6 **Changliu Wu^{1,2}**, Fang Cao^{1,2}, Sönke Szidat⁴, Yanlin Zhang^{1,2}

7 ¹ School of Ecology and Applied Meteorology, Nanjing University of Information Science and
8 Technology. Nanjing 210044, China.

9 ² Atmospheric Environment Center, Joint Laboratory for International Cooperation on Climate and
10 Environmental Change, Ministry of Education, Nanjing University of Information Science and
11 Technology. Nanjing 210044, China.

12 ³ Huzhou Meteorological Administration, Huzhou, 313000, China

13 ⁴ Department of Chemistry, Biochemistry and Pharmaceutical Sciences & Oeschger Centre for Climate
14 Change Research, University of Bern, Bern, 3012, Switzerland

15
16 Correspondence to: Yanlin Zhang (zhangyanlin@nuist.edu.cn; dryanlinzhang@outlook.com)

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18 **Abstract.** Reports on molecular-level characterization of primary and secondary constituents
19 in PM_{2.5} at high time resolution are limited to date, especially during haze events. The study
20 explored molecular composition and source contributions of PM_{2.5} with comprehensive
21 analytical methods by conducting intensive sampling at roughly 2-hour intervals during hazy
22 days in winter. Results show that organic matters were the predominant species, followed by
23 NO₃⁻. Biomass burning (BB) was the biggest contributor to organic carbon (OC), both in
24 concentration and in proportion. Radiocarbon analysis of carbonaceous fractions reflects that
25 fossil fuels dominate water-soluble organic carbon (WSOC) (61-82%) likely resulting from
26 increased coal combustion for residential cooking and heating and the coal-fired industry in
27 cold times. Interestingly, the contribution of non-fossils instead of fossil fuels to WSOC
28 enhanced with aggravating haze pollution, coinciding with significantly intensified BB during
29 that time. Other non-fossil sources, including fungal spores and plant debris, showed a larger
30 contribution to OC in light haze episodes. For secondary sources, naphthalene-derived
31 secondary organic carbon (SOC) contributed more to OC in PM_{2.5} (0.27-2.46%) compared to
32 biogenic emissions (0.05-1.10%), suggesting fossil fuels may dominate SOC formation during
33 urban haze events. SOC declined with rising haze pollution and presented high levels on days
34 with high temperature and low relative humidity due to elevated photooxidation. Additionally,
35 BB can raise secondary formation as well as the emissions of other sources, as demonstrated
36 by the significant relationships between BB tracers and many other source tracers. These
37 findings illustrate that BB likely plays a significant role in the heavy winter haze.

38

39 1. Introduction

40 The air quality of China has improved a lot over the past decade due to extensive
41 implementation of emission controls across the country. However, such progress was
42 unexpectedly shattered by severe air pollution happening during COVID-19 lockdown when
43 anthropogenic emissions significantly decreased (Huang et al., 2020b; Le et al., 2020; Wang et
44 al., 2020). This underscores the ongoing challenge of controlling PM_{2.5} pollution, especially
45 during cold seasons in megacities. Additionally, the emergence of ozone (O₃) pollution in many
46 urban areas complicates the situation. Rising O₃ levels, associated with increased atmospheric
47 oxidation capacity (Kang et al., 2021), create more complex air pollution scenarios due to
48 intricate secondary aerosol formations and the combined effects of PM_{2.5} and O₃.

49 PM_{2.5} exerts influences on air visibility, regional/global radiation balance, hydrological cycle
50 (Kaufman et al., 2002), and human and ecosystem health (Alexeeff et al., 2023; Chen et al.,
51 2022; Pope et al., 2004; Wang et al., 2022). In response scientists have carried out a series of
52 studies to analyze aerosol components and emission sources (Cheng et al., 2016; Huang et al.,
53 2014, 2020b, a; Jimenez et al., 2009; Kang et al., 2016, 2018a, b, 2019; Li et al., 2016a; Liu et
54 al., 2014; Sun et al., 2014; Wang et al., 2006; Yang et al., 2024; Zhang et al., 2012, 2018). These
55 studies revealed that PM_{2.5} pollution is formed through mixed interaction of primary and
56 secondary sources, including anthropogenic and biogenic origins. Primary sources mainly
57 contain plant emissions, fungal spores, soil dust, fossil fuel combustion, and biomass burning
58 (BB) (Anon, 2002; Fu et al., 2012; Kang et al., 2018b, a; Morris et al., 2011; Pöschl et al., 2010;
59 Simoneit, 2002; Zhang et al., 2015, 2016) while secondary sources primarily involve
60 homogeneous and heterogeneous reactions of biogenic and anthropogenic precursors (e.g., NO_x,
61 NH₃, SO₂, and VOCs) (Fu et al., 2010; Huang et al., 2014). Many PM_{2.5} species carry origin
62 information and thus can serve as tracers to determine specific sources.

63 For example, saccharides (i.e., anhydrosugars, sugars, and sugar alcohols) are important water-
64 soluble organic constituents of aerosols (Simoneit et al., 2004b; Sindelarova et al., 2014), which
65 can be cloud condensation nucleus and ice nuclei thus influencing Earth's climate and water
66 supply (Kaufman et al., 2002). Among them, levoglucosan is widely used as a typical BB tracer
67 (Elias et al., 2001; Li et al., 2021b; Liu et al., 2013). BB has a substantial impact on the

68 secondary organic aerosols (SOA) budget and climate change (Chen et al., 2017b; Zhang et al.,
69 2024). For example, substituted phenols from lignin combustion, which serve as BB tracers as
70 well, undergo aqueous phase oxidation with photooxidants to form SOA, significantly
71 influencing the evolution of organic aerosols (Zhang et al., 2024). However, the contribution of
72 BB emissions to SOA formation is not yet well understood and is consequently not accurately
73 represented in regional and global atmospheric chemistry models. Sugar alcohols like arabitol
74 and mannitol can be utilized to assess the contribution of airborne fungal spores to carbonaceous
75 aerosols (Bauer et al., 2008a, b; Fu et al., 2012, 2016). Other primary sugars (e.g., glucose) are
76 useful markers for plant pollen, fruits, and detritus (Fu et al., 2016; Puxbaum and Tenze-Kunit,
77 2003).

78 Secondary organic aerosols (SOA) are also a significant fraction, produced by the reactions of
79 oxidants (e.g., OH) with biogenic/anthropogenic VOCs (Claeys et al., 2004; Hallquist et al.,
80 2009; Huang et al., 2014; Mozaffar et al., 2020). Biogenic VOCs, such as isoprene,
81 monoterpenes, and sesquiterpenes, play a vital role in global SOA formation and atmospheric
82 processes (Claeys et al., 2004; Griffin et al., 1999; Guenther et al., 2006; Pöschl et al., 2010;
83 Sindelarova et al., 2014; Zhang et al., 2007), while anthropogenic VOCs (e.g., aromatic
84 hydrocarbons) tend to be more important in populated cities and nearby areas where coal
85 combustion, transportation, solvent use and biofuel/biomass burning contribute significantly
86 (Chen et al., 2017b; Ding et al., 2017; Srivastava et al., 2022). Despite its high importance and
87 wide existence, comprehensive characterization of SOA at the molecular level is difficult
88 because of complex and non-linear reactions and variable meteorological conditions. The lack
89 of molecular-level composition, abundance, and formation mechanisms of SOA at high time
90 resolution introduces inevitable uncertainties in modeling and forecasting air pollutants (Zhang
91 et al., 2022, 2023). Correctly simulating SOA with chemical transport models therefore can
92 become more challenging.

93 Other than the aforementioned organic species in PM_{2.5}, secondary inorganic aerosols (SIA, the
94 sum of sulfate (SO₄²⁻), nitrate (NO₃⁻), and ammonium (NH₄⁺)) equally account for a substantial
95 proportion of fine aerosols, especially on heavy pollution days (Fu et al., 2012; Huang et al.,
96 2014; Lu et al., 2019; Yan et al., 2023). Nitrate and sulfate in PM_{2.5} are mostly formed by

97 secondary formation and are accordingly expected to have significant regional influences once
98 they are emitted, particularly in winter. A recent study reported that nitrate comprised the largest
99 fraction of PM_{2.5} in China during severe haze, and NO_x emission reduction was regarded as an
100 effective measure to combat air pollution (Yan et al., 2023). Nevertheless, this conclusion was
101 challenged by the sustained severe haze during COVID-19 lockdown while NO_x emissions
102 substantially declined (Le et al., 2020), suggesting the complexity of PM_{2.5} pollution and callout
103 of more research work.

104 Although previous studies over past decades provide valuable information about aerosol
105 components, the molecular-level compositions and concentrations of fine particles still have
106 not been well understood due to their high spatial and temporal variability, especially at sub-
107 daily (hourly) levels. One reason is that aerosol properties can be modified at any time during
108 the transport through dry or wet deposition, in-cloud processes, and atmospheric chemical
109 reactions. Intensive aerosol sampling with high time resolution is then necessary for better
110 quantifying the PM_{2.5} components and source contributions. Former researches mostly focused
111 on analyzing the differences between hazy and clean days while very few reported variations
112 among different hazy days on sub-daily (e.g., hourly) basis in part due to the difficulty in too
113 frequent aerosol samplings. However, these molecular-level data at high time resolution are
114 useful and necessary for exploring the key factors controlling haze formation, which is
115 important for setting up regulatory standards in response to rapid changes in aerosol
116 composition and concentrations through time and place. Furthermore, the impacts of aerosol
117 particles with different properties (e.g., chemical composition) on climate (Kanakidou et al.,
118 2005; Kawana et al., 2022) remain unclear, and molecular-level PM_{2.5} components at hourly
119 intervals would greatly help better understand such issues.

120 Herein, we systematically unraveled hourly variation in molecular-level PM_{2.5} components
121 during haze events in Nanjing, a major city of the Yangtze River Delta with concentrated heavy
122 industry and population. Concentrations of major organic and inorganic components such as
123 BB tracers, sugar and sugar alcohols, oxidation products (e.g., biogenic SOA tracers and
124 aromatic acids), and water-soluble ions were measured and compared across three different
125 haze pollution levels. Contributions of primary sources to organic carbon (OC) in PM_{2.5} samples

126 were estimated including BB, fungal spores, and plant debris. Contributions of secondary OC
127 formed by biogenic and anthropogenic VOCs to total OC were also calculated. ¹⁴C
128 measurement were performed on water-soluble organic carbon (WSOC) to accurately quantify
129 the contribution of fossil fuel sources. The molecular-level results of PM_{2.5} components and
130 source contributions at high time resolution will help understand the haze formation and
131 evolution in megacities.

132 **2. Materials and methods**

133 **2.1 Sampling**

134 The sampling site was located on the rooftop of a building at the Nanjing University of
135 Information Science and Technology in Nanjing, China (32.2°N, 118.72°E). A total of 23 PM_{2.5}
136 samples were collected onto Prebaked quartz fiber filters (Pallflex) at a roughly 2-hour interval
137 from 31 December 2017 to 2 January 2018. High-volume air sampler (KC-1000, Qingdao
138 Laoshan Electric Inc., China) was used at a flow rate of 1.05 m³ min⁻¹. The field blank was also
139 collected with pump off during sampling. All the samples were stored in darkness at -20°C for
140 later analysis. In this study, the whole sampling period was divided into three episodes
141 according to PM_{2.5} levels, i.e., > 200, 100–200, and <100 μg m⁻³, representing a haze pollution
142 process from heavily polluted days to moderately polluted days.

143 **2.2 Measurements of organic molecules**

144 Sugar compounds, including anhydrosugars, sugar alcohols, and sugars, were measured using
145 ion chromatography (Dionex ICS-5000+, ThermoFisher Scientific, USA) after being extracted
146 with ultra-pure water (Milli-Q Reference, America). Standard curves establishment and blank
147 correction were conducted during the analysis. Other organic compounds, including biogenic
148 SOA tracers (isoprene, sesquiterpene, and monoterpene), diacids, and other main organic
149 molecules appeared in the present study were determined by gas chromatography/mass
150 spectrometry (Agilent Technologies; Santa Clara, CA). The average recoveries ranged from 70%
151 to 110% and repeatability experiments showed that the deviation was less than 15%. All the
152 data were corrected with field blanks. More details about measurements can be found in
153 previous studies (Bao et al., 2023). The total mass concentrations of SOC produced by isoprene

154 (2-methylglyceric acid and 2-methyltetrols were used), α/β -pinene, and β -caryophyllene were
155 estimated using the tracer-based method by Kleindienst et al. (2007). The BB derived OC and
156 fungal-spore derived OC were calculated using the methods in earlier reports (Bauer et al.,
157 2008a; Fu et al., 2014).

158 **2.3 Measurements of OC, EC, WSOC, and inorganic ions**

159 The elemental and organic carbon content were detected using a Sunset Lab EC/OC Analyzer
160 with the Interagency Monitoring of Protected Visual Environments (IMPROVE) 7-step
161 program heating method. This approach has been proved to be more accurate for EC and OC
162 measurement (Wu et al., 2020). Details about determination of water-soluble OC (WSOC) can
163 be found elsewhere (Bao et al., 2022). The water-soluble ions were measured by ion
164 chromatography (IC), and more detailed information is provided elsewhere (Bao et al., 2023).
165 The detected inorganic ions are listed in Table 1.

166 **2.4 ^{14}C analysis of the carbonaceous fractions**

167 The ^{14}C of WSOC was determined by extracting WSOC using deionized water and then
168 collecting the extracted solution for ^{14}C measurement using chemical wet oxidation of the water
169 extraction eluate (Song et al., 2022). The ^{14}C results are expressed as the fractions of measured
170 carbon, which is calculated as below ($F^{14}\text{C}$):

$$171 \quad F^{14}\text{C} = \frac{(^{14}\text{C}/^{12}\text{C})_{\text{sample}}}{(^{14}\text{C}/^{12}\text{C})_{1950}} \quad (1)$$

172 Where $(^{14}\text{C}/^{12}\text{C})_{1950}$ is the reference isotopic ratio in 1950. Then, these $F^{14}\text{C}$ values were
173 corrected by dividing by the reference value ($f_{nf,ref}$) to remove potential impacts of nuclear
174 bomb tests in the 1950s and 1960s, in order to obtain the non-fossil fractions of WSOC. More
175 details can be found in papers by Song et al. (2022) and Zhang et al. (2017).

$$176 \quad f_{nf} = F^{14}\text{C} / f_{nf,ref} \quad (2)$$

177 **2.5 Backward trajectories below 500 m above ground level**

178 Since regional transport also imposes influences on $\text{PM}_{2.5}$ levels (Chang et al., 2019; Chen et
179 al., 2017a), the Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model was
180 employed to compute backward trajectories of air masses arriving at the sampling site to

181 estimate the impacts of air pollution transport on haze formation (available at
182 <https://www.ready.noaa.gov/hypub-bin/trajtype.pl?runtype=archive>). MODIS active
183 fire/hotspot products were utilized to evaluate the impact of open biomass burning
184 during the entire sampling period. Based on the backward trajectory analysis, the air masses
185 throughout the sampling period were significantly influenced by biomass burning, as illustrated
186 in Fig. S1. By comparison, the third episode showed a greater influx of clean ocean air
187 masses (Fig. S1c).

188 3. Results and discussion

189 3.1 Inorganic ions

190 Table 1 lists the concentrations of identified inorganic ions, in which Cl^- , NO_3^- , SO_4^{2-} ,
191 and NH_4^+ are the major inorganic components during the entire sampling period. The
192 contribution of SIA to total $\text{PM}_{2.5}$ far exceeded that of organic matters (OM) during all
193 haze episodes, suggesting SIA contributes greatly to the occurrence of heavy haze. As
194 illustrated in Figure 1, NO_3^- was found to be the second dominant species (20.1–25.6%)
195 in $\text{PM}_{2.5}$ next to organic matters (OM), particularly in the heaviest haze event, consistent with
196 the findings in a megacity of Canada (Rivellini et al., 2024). However, these percentages are
197 greater than those in other megacities reported by Huang et al. (2014) (7.1–13.6%).
198 Such discrepancy may be caused by the spatial-temporal variations in energy mix and
199 meteorological parameters over years. The predominance of NO_3^- in SIA (30–52%) is
200 in agreement with the study about nitrate aerosols over another megacity in China (~
201 43%) (Fan et al., 2020). The major sources of NO_3^- include vehicles, coal combustion, natural
202 gas burning and biomass burning (Fan et al., 2023; Lin et al., 2024; Zhang et al., 2014a). The
203 rising NO_3^- relative to SO_4^{2-} may be associated with the decline in SO_2 and the rise in
204 NH_3 emissions in recent years, which allows more HNO_3 to condense into particulate
205 NO_3^- (Shah et al., 2024), as indicated by the significant relationship between NO_3^- and
206 NH_4^+ ($r = 0.98$, $p < 0.01$). Higher concentration ($56.0 \pm 4.4 \mu\text{g m}^{-3}$) and contribution (~
207 25.6%) of NO_3^- appeared in the highest- $\text{PM}_{2.5}$ episode. This is probably related to the
208 high relative humidity (RH) in this period (Fig. S2), which usually comes with high
209 aerosol liquid water content (Bian et al., 2014) and accordingly leads to more

210 heterogeneous reactions of nitrate formation (Lin et al., 2020). On the other hand, the
211 relatively colder temperatures in heavy haze episode favor the partitioning of HNO₃
212 from the gas phase to the particle phase. NO₃⁻ was also significantly correlated with
213 non-sea-salt SO₄²⁻ (nss-SO₄²⁻, **calculated by subtracting sea-salt sulfate from the total**
214 **sulfate using the typical sulfate-to-sodium mass ratio of 0.252 in seawater (Yang et al.,**
215 **2015))** ($r = 0.92, p < 0.01$), **suggesting they may share similar sources or formation**
216 **pathways** (Zhang et al., 2014a). Actually, under polluted conditions with high RH,
217 reactive nitrogen chemistry in aerosol water is a source of SO₄²⁻, where NO_x is not only
218 a precursor of nitrate but also an important oxidant for sulfate formation (Cheng et al.,
219 2016). Therefore, NO_x emission reductions have great potential in effectively reducing
220 atmospheric sulfate, nitrate, and even O₃ pollution simultaneously (Kang et al., 2021;
221 Shah et al., 2024). Interestingly, these three SIA components (**NO₃⁻, SO₄²⁻, and NH₄⁺**)
222 were observed to be strongly correlated with BB tracers (e.g., levoglucosan and
223 mannosan), with $p < 0.01$ and r in the range of 0.63–0.80, indicating BB was able to
224 promote the secondary production of SIA significantly. Given that the precursors of
225 NO₃⁻ and SO₄²⁻, i.e., NO_x and SO₂, are mainly contributed by fossil fuel combustion
226 activities (e.g., transportation and industrial emissions) in urban areas, the above
227 relationships thus suggest that BB may contribute greatly to the secondary
228 transformation of fossil-fuel-derived precursors.

229 **3.2 OC, EC, WSOC, and ¹⁴C of WSOC**

230 Similarly, the abundance of EC, OC, WSOC, and WISOC decreased with decreasing PM_{2.5}
231 levels (Table 1), in line with growing wind speeds. Compared with other episodes, the first
232 episode with PM_{2.5} > 200 μg m⁻³ had relatively high RH, low temperature, and low wind speed
233 (Fig. S2), demonstrating adverse meteorological conditions boost haze formation. As displayed
234 in Table 1, the mass concentrations of OC and EC were in the range of 8.74–41.1 and 1.26–
235 3.08 μg m⁻³, respectively. Such OC values are similar to those previously reported in PM_{2.5}
236 aerosols over Nanjing while EC levels are lower (Li et al., 2015, 2016b), reflecting the reduction
237 of primary emissions as a result of tightened emission controls over past years. OC and EC are
238 significantly correlated ($r = 0.87, p < 0.01$, Fig. S3), suggesting they may share common sources,

239 such as BB, vehicle exhaust, and fossil fuel combustion (Ji et al., 2019). OC/EC ratios showed
240 an increasing trend with rising PM_{2.5} levels (from an average of 8.7 to 13.3) (Table 1 and Fig.
241 2), close to those in regions dominated by BB (Boreddy et al., 2018; Zhang et al., 2014b). It
242 was reported that BB tended to emit relatively high fractions of OC rather than EC (Andreae
243 and Merlet, 2001), thus the high OC/EC ratios in this study illustrate substantial contributions
244 from BB, particularly during heavy haze events. In addition, high OC/EC ratios observed in
245 this study (> 2.0–2.2) indicate the presence of secondary organic aerosol (Li et al., 2016b). This
246 may be partially attributed to BB, which is a significant source of oxidants (Chang et al., 2024)
247 and an important contributor to SOA formation (Li et al., 2024; Lim et al., 2019; Yee et al.,
248 2013).

249 OC can be divided into water-soluble organic carbon (WSOC), which is often composed of BB-
250 derived and aged OC, and water-insoluble organic carbon (WISOC), normally representing
251 primary OC (Zhang et al., 2014b). As shown in Fig. 2, WISOC concentration (4.55–25.8 µg m⁻³)
252 is on average higher than WSOC, becoming the major portion of OC. WSOC ranged from
253 3.84 to 18.1 µg m⁻³ with higher values occurring in the most PM_{2.5} polluted episode (14.3 ± 2.62
254 µg m⁻³), comparable to the numbers previously reported in winter (14.0 µg m⁻³) (Li et al., 2018).
255 The ratios of WSOC/OC were relatively higher in more polluted periods (PM_{2.5} > 100 µg m⁻³)
256 with an average of 0.40 ± 0.06 and 0.43 ± 0.03, respectively (Table 1). It was reported that
257 higher WSOC/OC ratios (> 0.4) indicate the significant contribution of secondary organic
258 aerosol and aged aerosols (Boreddy et al., 2018; Ram et al., 2010). Considering the high RH in
259 the most polluted episode, the aqueous-phase oxidations of anthropogenic and/or biogenic
260 VOCs may be partially responsible for more WSOC formation during this period (Youn et al.,
261 2013). In comparison, the lower WSOC/OC ratios (0.35 ± 0.17) in the third episode (PM_{2.5} <
262 100 µg m⁻³) likely suggest rising primary emissions containing large amounts of water-insoluble
263 organics (e.g., lipid compounds), as indicated by greater WISOC/OC ratios during this period
264 (0.65 ± 0.17). In addition to secondary formation, WSOC was also found to be significantly
265 correlated with levoglucosan ($r = 0.74, p < 0.01$), indicating BB was an important contributor
266 to WSOC. Soluble organic gases derived from BB, such as phenols, can react with oxidants in
267 the aqueous phase to form SOA in aerosol liquid water and clouds, significantly contributing to

268 SOA formation. Moreover, this aqueous SOA formation greatly increases as relative humidity
269 (RH) increases (Zhang et al., 2024). Given the high relative humidity during the most polluted
270 periods, aqueous SOA production from BB-derived organic gases mostly likely play a crucial
271 role in heavy haze formation. Aqueous SOA generation from BB emissions was also confirmed
272 by many other studies (Gilardoni et al., 2016; Li et al., 2021a, 2014; Xiao et al., 2022),
273 highlighting the importance of BB emissions in atmospheric oxidation processes. This is also
274 supported by a more recent report that intermediate VOCs emitted by BB make a considerable
275 contribution to SOA (Li et al., 2024), reflecting the significant role of BB in the secondary
276 formation of atmospheric organic aerosols.

277 To track the variation trend of fossil and non-fossil contribution to carbonaceous aerosols during
278 the full course of haze development, the ^{14}C measurement was applied here to quantify fossil
279 and non-fossil sources of WSOC. As presented in Table 1 and Fig. 3, the non-fossil fraction of
280 WSOC was in the range of 18–39% (mean 26%), exhibiting fossil fuel sources were the
281 dominant contributor to WSOC on hazy days (61–82%, 74%) (Fig. S4). Such high fossil
282 contributions were previously observed in another megacity of Beijing during haze events in
283 winter ($\sim 61\%$) (Zhang et al., 2017) and in spring ($\sim 54\%$) (Liu et al., 2016), and these
284 differences in ^{14}C levels of WSOC could result from different origins and formation processes
285 of oxygenated OC in different places and seasons. The high proportion of fossil fuels observed
286 in this study can be attributable to extensive coal combustion for residential cooking and heating
287 on cold days, and industrial activities and traffic emissions in the vicinity of the sampling sites
288 could also contribute. Despite the predominance of fossil fuel sources, it is interesting to note
289 that the contribution of non-fossils, rather than fossil fuels, increased with increasing haze
290 pollution, suggesting non-fossil sources play a key role in the formation of heavy haze.
291 Similarly, the non-fossil fraction of organic aerosols in northern India was found to higher
292 during the more polluted cold period compared to the warm season (Bhattu et al., 2024).
293 Furthermore, the highest percentages of non-fossil sources occurred in the haziest period ($31 \pm$
294 6%) were coincident with the highest BB contributions during this period, which was also
295 evidenced by the correlations between non-fossil WSOC and BB markers (e.g., syringic acid, r
296 $= 0.68$, $p < 0.01$), indicating BB was a significant non-fossil source of WSOC and was likely

297 to be the important driver of heavy winter haze, despite the large amount of fossil fuel
298 contribution at the site. This is further supported by previous reports that emphasized the
299 contribution of aqueous-phase photochemical oxidation of BB organic gases to haze pollution
300 (Xiao et al., 2022; Zhang et al., 2024). This aqueous-phase SOA formation could contribute
301 more than the conventional semi-volatile SOA formation pathways, especially under polluted
302 conditions with high relative humidity (Zhang et al., 2024). Additionally, BB-chlorine
303 emissions could enhance oxidation capacity and further promote secondary aerosol formation
304 (Chang et al., 2024).

305 **3.3 Carbonaceous components**

306 Figure 4 displays the average concentrations of carbonaceous species in PM_{2.5} during three air
307 pollution episodes. Saturated diacids (within 1.66–14.6 $\mu\text{g m}^{-3}$) were the dominant
308 carbonaceous components of PM_{2.5}, followed by sugars and sugar alcohols (278–4936 ng m^{-3})
309 as well as anhydrosugars (79.4–801 ng m^{-3}). Higher anhydrosugar concentrations in the first
310 episode suggest greater BB impacts during heavy haze events. In contrast, the elevated levels
311 of sugars and sugar alcohols in the last two episodes are likely due to increased wind speeds,
312 which enhanced the resuspension of biogenic detritus and soil microbes rich in these substances.
313 Biogenic SOA tracers were minor species during winter haze and showed higher levels in the
314 second episode, probably due to enhanced photooxidation under elevated temperatures and low
315 RH. Similarly, unsaturated aliphatic diacids and aromatic acids presented the same trend as
316 biogenic SOA. Lignin and resin acids, alternative tracers for BB, demonstrated higher
317 concentrations in heavy haze events, as did anhydrosugars, further demonstrating the important
318 role of BB in heavy haze. The individual organic species identified in this study are discussed
319 below and in the Supporting Information document.

320 **3.3.1 Biomass burning tracers (anhydrosugars and lignin/resin acids)**

321 Levoglucosan is a specific indicator of BB and is generated from the thermal degradation of
322 cellulose (Simoneit, 2002). The largest levoglucosan concentration was in the highest-PM_{2.5}
323 episode (average: $471 \pm 122 \text{ ng m}^{-3}$), highlighting the remarkable contributions of BB to severe
324 haze formation (Fig. S5). These Figures are higher than those reported in winter in Beijing

325 (average: 361 ng m⁻³) (Li et al., 2018), and significantly higher than in the marine aerosols
326 (average: 7.3 ng m⁻³) (Kang et al., 2018a). Mannosan and galactosan, isomers of levoglucosan,
327 are main tracers for hemicellulose pyrolysis (Simoneit, 2002). Throughout the sampling period,
328 their concentrations were much lower than those of levoglucosan (Fig. S5 and S6). The
329 significant correlation between mannosan and levoglucosan ($r = 0.78$, $p < 0.01$) is indicative of
330 similar origins at this site.

331 The ratios of levoglucosan to potassium (L/K^+) can serve as an indicator to distinguish burning
332 from different biomasses (Urban et al., 2012). Similar to levoglucosan, K^+ is a BB tracer as
333 well, but there is no significant correlation between K^+ and levoglucosan in this study. This is
334 because in urban areas airborne potassium can also be emitted from other important sources,
335 such as meat cooking, refuse incineration, and resuspension of surface soil and fertilizers
336 (Simoneit, 2002; Urban et al., 2012). On average, the L/K^+ ratios for three episodes were 0.51
337 ± 0.19 , 0.20 ± 0.07 , and 0.44 ± 0.33 , respectively. The lower ratios observed in the second
338 episode might be triggered by the increased wind speeds, which favor the resuspension of
339 **surface soil and fertilizers containing abundant potassium into the air** (Urban et al., 2012). The
340 enhanced chemical degradation of levoglucosan under relatively high temperatures and low RH
341 may also contribute to lower L/K^+ ratios (Li et al., 2021b). In general, the L/K^+ values in this
342 study (0.06–1.04) agree well with those reported for crop and wood burning (Cheng et al., 2013;
343 Urban et al., 2012), implying a mixed biofuel combustion, as indicated by the isomeric ratios
344 of anhydrosugars (Fig. S8).

345 Levoglucosan to OC (L/OC) and to EC (L/EC) ratios have long been used to assess the
346 contribution of BB to aerosol abundance and possible degradation of levoglucosan (Mochida
347 et al., 2010; Sullivan et al., 2008; Zhang et al., 2008). L/OC and L/EC ratios in this study are
348 similar to those values in December in Beijing (Li et al., 2018) but higher than those in marine
349 aerosols in winter (Zhu et al., 2015a). Relatively higher L/OC and L/EC ratios were observed
350 in heavy haze events (Fig. S7), again proving the greater contribution of BB to heavy haze. The
351 overall decreasing L/OC and L/EC ratios with declined $PM_{2.5}$ level might stem from reduced
352 BB activities as well as levoglucosan degradation.

353 Lignin and resin acids are also reported in the smoke aerosols from BB, which can be used as
354 BB markers as well (Simoneit, 2002). In this study, the total lignin and resin acids are found in
355 much lower amounts than anhydrosugars (Fig. 4). A total of three lignin products (i.e., 4-
356 hydroxybenzoic acid, vanillic acid, and syringic acid) and one resin product (dehydroabietic
357 acid) were measured, with higher concentrations occurring in highest-PM_{2.5} episode ($46.5 \pm$
358 38.0 ng m^{-3}), further demonstrating significant BB influence on heavy haze. These values are
359 comparable to those in wintertime aerosols over Beijing (47.5 ng m^{-3}) (Li et al., 2018).
360 Specifically, **syringic acid was found to be the most abundant species among lignin and resin**
361 **acids** during heavy haze events ($\sim 28.0 \text{ ng m}^{-3}$) while dehydroabietic acid dominated in
362 moderate and light haze episodes (~ 14.4 and 17.0 ng m^{-3} , respectively). Dehydroabietic acid
363 and vanillic acid are typical tracers emitted from burning of conifer (softwood fuel), while
364 syringic acid was found enriched in hard wood smoke (Simoneit, 2002). Therefore, the
365 relatively high levels of dehydroabietic acid and syringic acid observed in the highest-PM_{2.5}
366 episode together exhibit greater contributions of mixed wood burning on cold days, during
367 which plentiful firewood were burned for residential cooking and heating in nearby suburbs. 4-
368 hydroxybenzoic acid (4-HBA) is one major molecular tracer identified in the pyrolysis of non-
369 woody vegetation including grass and crop residue, with concentrations in the range of 0.05–
370 9.32 ng m^{-3} . A significant correlation between 4-HBA and vanillic acid was found ($r = 0.86$, p
371 < 0.01), indicating similar sources such as mixed biofuel burnings.

372 **3.3.2 Primary sugars and sugar alcohols**

373 Primary sugars identified in this study mainly include trehalose and glucose with concentrations
374 ranging from $86.5\text{--}3023 \text{ ng m}^{-3}$ and $49.3\text{--}551 \text{ ng m}^{-3}$, respectively. Trehalose is the most
375 abundant saccharide in soils especially in the fine mode (PM_{2.5}) (Jia and Fraser, 2011) and can
376 be used as a potential tracer for resuspension of surface soil and unpaved road dust (Fu et al.,
377 2012). This is supported by the similar change trend of trehalose and nss-Ca²⁺ in the present
378 study, since nss-Ca²⁺ is an indicator for soil dust, particularly in winter and spring (Virkkula et
379 al., 2006). Generally, trehalose showed higher concentrations in the second episode with an
380 average of $1057 \pm 1112 \text{ ng m}^{-3}$, which might be linked to the meteorological parameters like
381 increased wind speeds relative to the other two episodes, enabling more trehalose in surface

382 soil to transport into the air. Glucose is also rich in biologically active soils and was proposed
383 to be a marker for fugitive dust from cultivated land (Rogge et al., 2007). In addition, glucose
384 is abundant in plant tissues as well, such as pollen, fruits, developing leaves, and plant detritus
385 (Graham et al., 2003). Both glucose and trehalose presented higher levels in moderate haze
386 events, indicating enhanced primary biogenic sources during that time due to the rising
387 temperature and wind speeds (Zhu et al., 2015b).

388 Sugar alcohols detected in this study consisted of arabitol, mannitol, and glycerol with
389 concentrations in the range of 4.59–48.2 ng m⁻³, 0.47–24.4 ng m⁻³, and 119–4749 ng m⁻³,
390 respectively. Glycerol was obviously the most abundant sugar alcohols, consistent with
391 previous studies (Kang et al., 2018b; Li et al., 2018; Ren et al., 2020). The levels of glycerol
392 went up when PM_{2.5} concentration declined, with the highest levels present in the lowest-PM_{2.5}
393 episode (~ 2348 ng m⁻³). Such a trend may be explained by the rising local temperature during
394 moderate and light haze events, as lower ambient temperatures can reduce microbial activities
395 like fungal spore release. Conversely, higher concentrations of arabitol and mannitol turned out
396 to exist in the highest-PM_{2.5} episode (> 200 µg m⁻³), when BB greatly intensified. In addition
397 to being emitted directly from natural sources like microbial activities and plant tissues, all
398 these saccharides can be emitted significantly by thermal stripping during BB (Simoneit et al.,
399 2004b). Also, BB can enhance emissions and long-range transport of some non-combusted
400 organic compounds (Medeiros et al., 2006). It was reported that sugar alcohols were associated
401 with airborne detritus from mature leaves and would be more prevalent during the period of
402 leaf senescence (Graham et al., 2003; Medeiros et al., 2006), thus high levels of arabitol and
403 mannitol can be expected in strongly BB-impacted aerosols in winter. This is further supported
404 by the correlations between arabitol/mannitol and levoglucosan ($r = 0.39, p = 0.06$ and $r = 0.40,$
405 $p = 0.06$, respectively). The above results indicate BB may have a greater effect on arabitol and
406 mannitol than on glycerol, suggesting their main sources in the region were different.

407 3.3.3 Biogenic SOA tracers

408 The total levels of biogenic SOA tracers were in the range of 1.80–34.7 ng m⁻³, with higher
409 concentrations in the second episode (averaging 15.8 ng m⁻³) as shown in Fig. 4. Isoprene-
410 derived SOA tracers contributed more to the total biogenic SOA than monoterpene and

411 sesquiterpene combined (Fig. S12). The averaged ratios of isoprene to monoterpene oxidation
412 products for three episodes were 1.16 ± 0.53 , 1.44 ± 0.71 , and 2.16 ± 0.94 , respectively. Such
413 values were lower than those reported in mountain aerosols, Central East China (about 4.9–6.7)
414 (Fu et al., 2010), where large isoprene fluxes and high levels of atmospheric radicals such as
415 OH exist.

416 Isoprene emitted from terrestrial vegetation is the predominant biogenic source of hydrocarbon
417 in the atmosphere though emission of monoterpenes is quite universal among plants (Sharkey
418 et al., 2008). Isoprene has reactive double bonds and hence can be readily oxidized by radicals
419 (e.g., OH) as a source of tropospheric O₃ and SOA (Chameides et al., 1988; Claeys et al., 2004;
420 Lin et al., 2013a). A total of six isoprene-SOA tracers were detected in these samples, including
421 three C5-alkene triols, two 2-methyltetrols, and 2-methylglyceric acid (Table 1 and Fig. S10-
422 S12). All of them showed higher levels in the second episode with average concentrations of
423 $8.58 \pm 2.52 \text{ ng m}^{-3}$ for total isoprene-SOA, $2.20 \pm 0.56 \text{ ng m}^{-3}$ for C5-alkene triols, 3.81 ± 1.20
424 ng m^{-3} for 2-methyltetrols (2-MTs), $2.56 \pm 0.96 \text{ ng m}^{-3}$ for 2-methylglyceric acid (2-MGA),
425 respectively. By comparing the temporal variations of meteorological factors and biogenic SOA
426 concentrations (Fig. S2 and S10), it is not hard to find that the peak concentrations basically
427 appeared under relatively high temperature and low RH conditions, in agreement with results
428 in central China (Li et al., 2013). The similar variation patterns among isoprene SOA tracers
429 suggest they may share common sources and be formed via similar pathways, as indicated by
430 the significant correlations between C5-alkene triols and 2-MTs/2-MGA ($r = 0.89\text{--}0.90$, $p <$
431 0.01). 2-Methyltetrols were the dominant isoprene products ($0.20\text{--}8.71 \text{ ng m}^{-3}$), in line with
432 previous studies (Kang et al., 2018a; Li et al., 2018). Both 2-methyltetrols and C5-alkene triols
433 are produced from the photooxidation of isoprene under low-NO_x (NO_x = NO+NO₂) conditions
434 (Surratt et al., 2006, 2010) while 2-MGA is formed under high-NO_x conditions (Lin et al., 2013b;
435 Surratt et al., 2006). The concentration ratios of C5-alkene triols to 2-methyltetrols did not
436 exhibit significant changes except in the most polluted events (Fig. S13), which imply that their
437 reaction processes may be different during heavy haze compared to moderate and light haze
438 episodes. The answer may lie in the chemical structure of these two species, as C5-alkene triols
439 have a double bond which is prone to be oxidized easily, thus the dropping ratios of C5-alkene

440 triols to 2-methyltetrols therefore probably reflect photochemical aging of organic aerosols over
441 time.

442 Oxidation products of monoterpene include 3-hydroxyglutaric acid (3-HGA), pinonic acid,
443 and pinic acid. The concentrations of total monoterpene-derived SOA were in the range of 1.17–
444 13.5 ng m⁻³, with higher levels occurring in second episode which probably results from the
445 enhanced photooxidation reactions due to increased temperature and declined RH. A clear
446 correlation was found between 3-HGA and pinonic acid ($r = 0.79$, $p < 0.01$), implying similar
447 sources and formation pathways. Pinic acid is a minor compound in monoterpene-derived SOA
448 (0.04–1.81 ng m⁻³), with abundances less than those of 3HGA (0.42–6.60 ng m⁻³) and pinonic
449 acid (0.05–6.91 ng m⁻³) (Fig. S11). Pinic acid correlated with lignin and resin acids such as
450 vanillic acid and 4HBA ($r = 0.69$ – 0.76 , $p < 0.01$), suggesting BB can significantly promote its
451 secondary formation. This is because BB is not only a significant source of air pollutants but
452 also of oxidants (Chang et al., 2024), which enhances oxidation capacity and further promotes
453 photochemistry and SOA formation. However, pinic acid did not exhibit the highest
454 concentration during the heavy haze period with the greatest BB contribution. This may be due
455 to pinic acid undergoing further reactions at high relative humidity, forming highly oxidized
456 polar compounds through the addition of a molecule of water and the opening of the
457 dimethylcyclobutane ring (Claeys et al., 2007).

458 β -caryophyllinic acid is an ozonolysis or photooxidation product of β -caryophyllene (Jaoui et
459 al., 2007), a major species of sesquiterpenes emitted from plants (Duhl et al., 2008). On the
460 whole, there are no pronounced differences in concentrations of β -caryophyllinic acid among
461 the three episodes with the exception of a slightly higher average of 0.29 ng m⁻³ in the lowest-
462 PM_{2.5} event ($< 100 \mu\text{g m}^{-3}$).

463 3.3.4 Aromatic acids

464 Three aromatic acids containing two phthalic acids (phthalic acid and isophthalic acid) and
465 benzoic acid were determined in these aerosols. Relatively higher total abundances of aromatic
466 acids occurred in high-PM_{2.5} episodes ($> 100 \mu\text{g m}^{-3}$) with a concentration range of 8.3–45.1 ng
467 m⁻³. Phthalic acid (Ph) and isophthalic acid (iPh) were the major aromatic acids, with

468 concentrations in the range of 1.45–13.0 ng m⁻³ and 0.98–21.2 ng m⁻³, respectively. The
469 secondary photochemical reactions of polycyclic aromatic hydrocarbons (PAHs) such as
470 naphthalene are possibly the main sources of Ph, which has been proposed as a naphthalene-
471 derived SOA tracer (Fine et al., 2004; Ren et al., 2020). Vehicle exhausts are important sources
472 of naphthalene in urban atmosphere, and therefore transportation emissions were likely to be
473 responsible for the Ph over this site. By comparison, benzoic acid was a minor species in
474 aromatic acids (0.47–11.4 ng m⁻³). It can be directly emitted from vehicle exhaust and
475 secondarily produced through photochemical reactions of aromatic hydrocarbons from traffic
476 emissions such as toluene (Ho et al., 2015; Li et al., 2022; Rogge et al., 1993; Suh et al., 2003).
477 The relationships among Ph, iPh, and benzoic acid ($r = 0.64\text{--}0.79$, $p < 0.01$) suggest they share
478 common sources, such as fossil fuels.

479 3.3.5 Hydroxy-/polyacids

480 Polyacids are reported to be secondary photooxidation products of atmospheric organic
481 precursors (Fu et al., 2008; Kawamura and Sakaguchi, 1999). A total of three hydroxy-
482 /polyacids were measured, including glyceric acid, malic acid, and tartaric acid. The slightly
483 higher content of hydroxy-/polyacids in second episode may be due to enhanced photooxidation
484 reactions under increased temperature and low RH. Malic acid (0.77–6.60 ng m⁻³) is the major
485 compound in hydroxy carboxylic acids, followed by glyceric acid (0.22–6.56 ng m⁻³), while
486 tartaric acid is relatively minor. The above result is consistent with an early report over the
487 polluted East Asia/Pacific region (Simoneit et al., 2004a). In current study, glyceric acid was
488 significantly correlated with tartaric acid ($r = 0.81$, $p < 0.01$), implying similar sources and/or
489 formation pathways. Moreover, glyceric and tartaric acid were found to be significantly
490 correlated with isoprene ($r = 0.71\text{--}0.93$, $p < 0.01$) and monoterpene ($r = 0.65\text{--}0.77$, $p < 0.01$)
491 SOA tracers (e.g., 2-methyltetrols, C5-alkene triols, pinic, and pinonic) while malic acid was
492 positively correlated with glucose ($r = 0.65$, $p < 0.01$). These significant relationships suggest
493 that hydroxy-acids may be secondary oxidation products of biogenic VOCs and sugars
494 (Simoneit et al., 2004a). **There were also pronounced correlations between glyceric acid and**
495 **aromatic acids such as iPh and benzoic acid ($r = 0.63\text{--}0.71$, $p < 0.01$), implying that they may**
496 **undergo similar atmospheric processing pathways.** In addition, glyceric and tartaric acids were

497 significantly correlated with 4HBA and vanillic acid ($r = 0.58\text{--}0.81$, $p < 0.01$), indicating BB
498 contribute to the secondary production of hydroxy-acids.

499 **3.3.6 Dicarboxylic acids**

500 Dicarboxylic acids are predominantly present as air particles rather than in the gas phase due to
501 their low vapor pressures (Limbeck et al., 2001; Saxena and Hildemann, 1996). They contain
502 two carboxyl groups and are the major constituents of water-soluble organics in aerosols
503 (Saxena and Hildemann, 1996), as proved by the significant correlation between WSOC and
504 dicarboxylic acids in this study ($r = 0.74\text{--}0.87$, $p < 0.01$). In addition to being directly released
505 into the air from incomplete combustion of fossil fuels, meat cooking, and biomass burning,
506 they can be also formed by secondary photochemical reactions (Mochida et al., 2003). For
507 instance, isoprene and unsaturated fatty acids are proposed to be sources of dicarboxylic acids
508 in the open ocean (Bikkina et al., 2014). Totally, four saturated dicarboxylic acids (i.e., oxalic,
509 malonic, succinic, and glutaric acid) and two unsaturated dicarboxylic acids (maleic and
510 fumaric acid) were included here. The levels of unsaturated-dicarboxylic acids ($2.48\text{--}69.5$ ng
511 m^{-3}) were far less than those of saturated diacids ($1.66\text{--}14.6$ $\mu\text{g m}^{-3}$). Similar to biogenic SOA,
512 dicarboxylic acids showed higher concentrations in the episode with relatively high temperature
513 and low RH (Fig. 4), which are beneficial for the photochemical oxidation of organic precursors.

514 Malonic acid (C3, $1.48\text{--}14.3$ $\mu\text{g m}^{-3}$) was the most abundant species among measured
515 dicarboxylic acids, followed by oxalic acid (C2, $0.09\text{--}0.74$ $\mu\text{g m}^{-3}$). C2 and C4 (succinic acid)
516 levels are comparable to those reported in $\text{PM}_{2.5}$ aerosols from megacities such as Beijing (Ho
517 et al., 2010) and Guangzhou (Liu et al., 2021). It was deduced that C2 and C3 diacids are likely
518 produced by the oxidation of C4 and other longer chain diacids, whereas those longer-chain
519 diacids (C5–C10) are formed by oxidation of unsaturated fatty acids (Kawamura and Gagosian,
520 1987; Kawamura and Sakaguchi, 1999). This conclusion is supported by the significant
521 correlations between C2 and C4 ($r = 0.86$, $p < 0.01$), C2 and C5 (glutaric acid) ($r = 0.77$, $p <$
522 0.01) and C4 and C5 ($r = 0.60$, $p < 0.01$) in the present study. In comparison with other diacids
523 identified in this study, the relatively higher levels of C2 and C3 may partially result from
524 considerable photodegradation of C4 and C5 in haze events, implying these urban aerosols may
525 have undergone great aging processes. The ratio of C3 to C4 is a useful indicator for elevated

526 photochemical production of dicarboxylic acids in the atmosphere, as C4 is a precursor of C3
527 formation (Kawamura and Ikushima, 1993). In this study, C3 dicarboxylic acid was far more
528 abundant than C4 indicating strong photochemical processes, as also suggested by the high
529 WSOC/OC ratios mentioned earlier. Such findings mean secondary formation is an important
530 pathway of dicarboxylic acids on hazy days in urban Nanjing, apart from primary emissions. It
531 should be noted that C2 and C5 both correlated well with levoglucosan ($r = 0.66\text{--}0.69, p < 0.01$),
532 indicating BB is an alternative source of these diacids and/or can facilitate their oxidation
533 reaction (Kawamura and Bikkina, 2016). Chlorine emissions from BB were found to increase
534 oxidant levels, such as O₃ and OH radicals, largely impacting atmospheric chemistry and
535 oxidation process (Chang et al., 2024).

536 3.4 Contributions of primary and secondary sources to OC

537 To evaluate the contribution of primary (e.g., BB, fungal spores, and plant debris) and
538 secondary sources (e.g., oxidation reactions of PAHs and biogenic VOCs including isoprene,
539 monoterpene, and sesquiterpene) to OC in PM_{2.5}, tracer-based methods were applied here.
540 Details about specific calculation methods and relevant conversion factors can be found in our
541 previous work and other reports (Bauer et al., 2008a; Gelencsér et al., 2007; Holden et al., 2011;
542 Kang et al., 2018a; Kleindienst et al., 2007, 2012; Puxbaum and Tenze-Kunit, 2003).

543 Compared with other primary and secondary sources, BB made an absolutely predominant
544 contribution to aerosol OC throughout the whole sampling period, both in concentration and in
545 proportion (0.72–8.86 $\mu\text{g m}^{-3}$ and 8.29–26.5%). The greatest impact of BB was observed during
546 heavy haze events (mean: $5.79 \pm 1.50 \mu\text{g m}^{-3}$, $16.3 \pm 3.39\%$). This could be attributed to the
547 increased domestic wood/crop combustion for heating and cooking, along with biomass
548 burning in the surrounding area, driven by low temperatures and high relative humidity during
549 this period (Figs. S1-S2). BB-chlorine emissions have been shown to elevate O₃ and OH radical
550 levels, significantly impacting oxidation processes (Chang et al., 2024). In addition, soluble
551 organic gases from BB can dissolve in aerosol/cloud liquid water and subsequently react with
552 aqueous phase oxidants to form SOA, with these reactions increasing with increasing RH
553 (Zhang et al., 2024). Considering the potential atmospheric degradation of levoglucosan, the
554 contribution of BB might be somewhat underestimated and thus the actual BB fraction is likely

555 larger, highlighting the crucial role of BB in haze formation. A higher relative contribution of
556 BB to organic aerosols during the colder period, characterized by elevated PM_{2.5} concentrations,
557 was also recently reported in northern India (Bhattu et al., 2024). Relatively high concentration
558 of fungal-spores-derived OC occurred in the highest-PM_{2.5} episode ($0.44 \pm 0.14 \mu\text{g m}^{-3}$) when
559 BB impacts were significant (Fig. S15), consistent with an earlier study that observed elevated
560 fungal spore tracers on BB-affected days (Yang et al., 2012). This suggests that BB could raise
561 emissions from other sources, such as fungal spores, further exacerbating air pollution.
562 Nonetheless, percentages of fungal spores to OC were on the decline with increasing PM_{2.5}
563 levels with higher fractions displaying in light haze episode ($2.38 \pm 2.26\%$), during which the
564 contribution of BB to OC remained high ($15.9 \pm 7.01\%$). By comparison, concentrations and
565 contributions of OC from plant debris were higher in the second episode ($0.45 \pm 0.21 \mu\text{g m}^{-3}$,
566 $1.99 \pm 1.02\%$), probably on account of increased resuspension of surface soils and road dust
567 resulting from elevated wind speeds and temperatures (Simoneit et al., 2004b). The total
568 abundance of primary OC derived from BB, fungal spores and plant debris ranges from 1.23 to
569 $9.65 \mu\text{g m}^{-3}$ making up 11.3–31.3% of OC, with higher concentrations in the most polluted
570 episode (average: $6.52 \pm 1.62 \mu\text{g m}^{-3}$, $18.4 \pm 3.62\%$). It is noteworthy that despite lower
571 concentrations of total primary OC in light haze episode (PM_{2.5} < 100 $\mu\text{g m}^{-3}$), the contribution
572 of primary OC to aerosol OC was comparable to and even bigger ($19.9 \pm 8.31\%$) than those in
573 heavy and moderate episodes.

574 By comparison, secondary sources (i.e., isoprene, monoterpene, sesquiterpene, and naphthalene)
575 contributed less than primary sources, accounting for only 0.38–3.56% of OC in PM_{2.5}, which
576 probably arose from reduced photolysis during winter due to less intense sunlight. Overall, SOC
577 showed high levels ($0.36 \pm 0.07 \mu\text{g m}^{-3}$) and high contributions ($1.53 \pm 0.37\%$) during periods
578 of high temperatures and low RH, because such weather conditions promote increased
579 photochemical reactions and the production of SOC in the atmosphere. It is notable that
580 naphthalene-derived SOC was the main secondary source of OC, both in concentration (0.04–
581 $0.34 \mu\text{g m}^{-3}$) and in proportion (0.27–2.46%) (Table 2), followed by biogenic isoprene-derived
582 SOC (0.003–0.09 $\mu\text{g m}^{-3}$, 0.01–0.60%), **indicating anthropogenic VOCs make a dominate**
583 **contribution to SOC in these urban aerosols.** Moreover, the total concentrations and fractional

584 contributions of these biogenic SOCs (0.01–0.16 $\mu\text{g m}^{-3}$, 0.05–1.10%) were lower than those
585 from anthropogenic sources, probably due to **significantly reduced biogenic VOC emissions**
586 **and largely increased fossil fuel combustion during cold winter periods**. The abundance and
587 percentage of total primary and secondary OC were 1.54–9.98 $\mu\text{g m}^{-3}$ and 11.9–32.2%,
588 respectively, based on the detected tracers in this study. Such values are comparable to those
589 reported in winter aerosol in Beijing (6.18–38.3%) (Li et al., 2018).

590 **4. Conclusions**

591 **Molecular distributions and high temporal variations of primary and secondary components in**
592 **PM_{2.5} during winter hazy episodes in urban Nanjing were comprehensively characterized**
593 **through intensive sampling. Our results revealed that OM dominated the total PM_{2.5}, followed**
594 **by NO₃⁻. ¹⁴C analysis showed that while fossil fuel sources primarily contributed to WSOC,**
595 **non-fossil sources, notably BB, became more significant as PM_{2.5} pollution intensified. BB**
596 **made a dominant contribution to OC, particularly during severe haze events, likely due to**
597 **aqueous SOA formation from BB-derived organic gases. Other non-fossil sources like fungal**
598 **spores were also elevated by BB, whereas plant debris contributions were higher on lighter hazy**
599 **days with higher wind speeds and temperatures. Overall, these findings highlight the significant**
600 **role of BB in winter haze over Nanjing and underscore the need for further research into the**
601 **molecular-level identification of gaseous species from BB emissions and their role in secondary**
602 **aerosol formation. Additionally, although meteorological parameters have an important**
603 **influence on the development of heavy haze, accurately quantifying their contribution remains**
604 **a challenge for future research.**

605

606 **Data availability.** The dataset for this paper is available upon request from the corresponding
607 author (zhangyanlin@nuist.edu.cn).

608 **Supplement.** Information on Chloride, unsaturated diacids, monocarboxylic acids,
609 Methylglyoxal, Methanesulfonic acid (MSA) in PM_{2.5} were investigated here. HYSPLIT back
610 trajectories initiated over Nanjing (Fig. S1). Time series of meteorological parameters (Fig. S2).
611 Relationship between EC and OC in PM_{2.5} (Fig. S3). Temporal variations of fossil and non-

612 fossil contribution to WSOC (Fig. S4). Temporal variations of biomass burning tracers along
613 with the average concentrations of anhydrosugars and lignin and resin products detected in three
614 episodes (Fig. S5-S6). Temporal variations of ratios of L/M, L/OC, and L/EC, and the average
615 ratios during three episodes (Fig. S7). Comparison of L/M and M/G ratios from literature values
616 and ambient aerosols in this study (Fig. S8). Temporal variations of sugars, sugar alcohols, and
617 biogenic SOA tracers (Fig. S9-S10). Average concentrations of biogenic SOA tracers detected
618 in three episodes (Fig. S11-S12). Temporal variations in the concentration ratios of isoprene
619 oxidation products (Fig. S13). Temporal variations in the biogenic SOC derived from isoprene,
620 monoterpene, and sesquiterpene (Fig. S14). Temporal variations in biomass burning-derived
621 OC, fungal spores-derived OC, and plant debris-derived OC (Fig. S15).

622 **Author contributions.** YLZ designed the research. MYB collected aerosol samples. MYB and
623 WHS performed the laboratory analyses. The paper was written by MJK with editing from all
624 co-authors.

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

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Table 1. Concentrations of PM_{2.5} components in aerosol samples collected in urban Nanjing during polluted episodes.

Species	PM _{2.5} ($\mu\text{g m}^{-3}$)						PM _{2.5} ($\mu\text{g m}^{-3}$)										
	>200						100-200						<100				
	mean	std	min	max	mean	std	min	max	mean	std	min	max	mean	std	min	max	
EC ($\mu\text{g m}^{-3}$)	2.67	0.26	2.27	3.08	2.00	0.08	1.93	2.14	1.73	0.31	1.26	2.24					
OC ($\mu\text{g m}^{-3}$)	35.4	4.78	23.8	41.1	23.7	3.86	18.5	28.7	15.3	6.19	8.74	26.7					
TC ($\mu\text{g m}^{-3}$)	38.1	4.85	26.0	43.4	25.7	3.91	20.5	30.7	17.0	6.39	10.2	28.8					
WSOC ($\mu\text{g m}^{-3}$)	14.3	2.62	8.97	18.1	10.2	1.30	8.11	11.4	6.21	1.90	3.84	8.26					
WISOC ($\mu\text{g m}^{-3}$)	21.1	3.68	14.8	25.8	13.5	2.78	10.4	17.5	9.87	4.64	4.55	19.4					
OC/EC	13.3	2.08	10.5	17.4	11.8	1.74	9.57	14.4	8.70	2.72	6.00	13.2					
WSOC/OC	0.40	0.06	0.31	0.49	0.43	0.03	0.39	0.47	0.35	0.17	nd	0.51					
WISOC/OC	0.60	0.06	0.51	0.69	0.57	0.03	0.53	0.61	0.65	0.17	0.49	1.00					
14C-WSOC	0.31	0.06	0.25	0.39	0.25	0.02	0.23	0.28	0.24	0.04	0.18	0.29					
Inorganic icons ($\mu\text{g m}^{-3}$)																	
F ⁻	0.08	0.03	0.05	0.12	0.16	0.20	0.06	0.52	0.05	0.02	0.02	0.08					
Cl ⁻	7.00	1.66	3.86	10.2	6.51	1.50	4.26	7.86	5.51	2.62	1.88	10.2					
NO ₃ ⁻	56.0	4.39	48.7	62.4	33.9	6.50	24.0	40.1	12.7	4.37	5.75	17.7					

SO ₄ ²⁻	30.9	4.42	26.4	38.8	19.1	3.78	13.2	23.8	10.4	3.95	6.59	19.4
NH ₄ ⁺	28.0	3.20	20.3	30.9	17.1	3.60	10.8	19.7	8.52	2.35	4.97	11.4
PO ₄ ³⁻	0.14	0.02	0.11	0.17	0.07	0.03	0.03	0.12	0.02	0.01	0.01	0.03
Na ⁺	0.73	0.15	0.47	0.98	0.83	0.18	0.59	1.08	0.47	0.16	0.29	0.76
Ca ²⁺	0.73	0.41	0.35	1.58	1.23	0.55	0.76	1.99	0.40	0.16	0.19	0.62
nss-Ca ²⁺	0.70	0.41	0.32	1.55	1.20	0.55	0.73	1.96	0.38	0.16	0.17	0.61
K ⁺	0.98	0.24	0.72	1.52	1.01	0.34	0.62	1.40	0.65	0.48	0.22	1.69
nss-K ⁺	0.95	0.24	0.69	1.49	0.98	0.34	0.60	1.36	0.64	0.48	0.21	1.67
Mg ²⁺	0.69	0.37	0.25	1.18	0.24	0.14	0.10	0.42	0.10	0.07	0.03	0.22
Anhydrosugars (ng m⁻³)												
Levoglucosan (L)	471	122	284	721	185	28.1	142	219	201	121	59.0	395
Galactosan (G)	39.6	19.1	4.84	63.6	73.2	14.8	55.1	94.1	51.0	44.6	6.70	115
Mannosan (M)	45.4	21.2	20.8	81.9	14.8	9.73	4.79	30.3	14.0	8.11	6.63	25.4
L/M	11.5	3.21	5.86	16.5	18.3	12.4	4.67	38.0	22.4	12.7	8.88	38.2
M/G	2.86	4.83	0.41	15.6	0.20	0.13	0.07	0.41	0.66	1.20	nd	3.09
L/K ⁺	0.51	0.19	0.21	0.76	0.20	0.07	0.14	0.29	0.44	0.33	0.06	1.04
Sugar alcohol (ng m⁻³)												

arabitol	30.5	10.3	12.0	44.1	28.8	10.4	16.6	42.1	17.8	13.4	4.59	48.2
mannitol	14.4	6.24	0.47	24.4	14.2	4.12	7.92	18.4	12.9	7.20	2.43	22.0
glycerol	295	151	119	561	1822	1916	376	4062	2348	1334	652	4749
Sugars (ng m⁻³)												
trehalose	851	874	86.5	2970	1057	1112	302	3023	672	521	257	1378
glucose	203	85.1	49.3	377	312	148	193	551	158	56.0	69.8	240
total measured saccharides	1951	896	633	3841	3507	1632	1738	4976	3474	1238	1478	5436
Isoprene SOA tracers (ng m⁻³)												
cis-2-methyl-1,3,4-trihydroxy-1-butene	0.38	0.42	0.02	1.26	0.62	0.17	0.38	0.85	0.36	0.17	0.11	0.68
3-methyl-2,3,4-trihydroxy-1-butene	0.45	0.67	0.03	2.17	0.59	0.24	0.26	0.93	0.64	0.37	0.01	1.07
trans-2-methyl-1,3,4-trihydroxy-1-butene	0.76	0.83	0.03	2.87	0.99	0.53	0.41	1.81	0.74	0.52	0.06	1.55
sum of C5-Alkene triols	1.59	1.83	0.07	6.30	2.20	0.56	1.66	2.91	1.74	0.99	0.18	3.19
2-methylthreitol	0.69	1.16	0.07	3.78	1.52	0.60	0.65	2.26	1.16	0.92	0.03	3.11
2-methylethritol	1.17	1.55	0.10	4.93	2.30	0.69	1.29	2.97	2.10	1.19	0.41	4.30

sum of 2-methyltetros	1.86	2.68	0.20	8.71	3.81	1.20	1.94	4.67	3.26	2.09	0.45	7.41
2-methylglyceric acid	2.05	1.86	0.21	5.93	2.56	0.96	1.13	3.52	1.58	1.09	0.35	3.80
sum of isoprene SOA	5.51	6.23	0.56	20.9	8.58	2.52	4.80	11.1	6.58	4.10	0.97	14.4
Monoterpene SOA tracers (ng m⁻³)												
3HGA	2.45	1.64	0.94	5.52	2.75	2.30	1.02	6.60	0.95	0.39	0.42	1.53
pinonic	1.61	2.15	0.05	6.91	3.41	1.67	1.65	5.64	1.04	0.57	0.38	1.81
pinic	0.32	0.31	0.05	1.06	0.87	0.62	0.24	1.81	0.84	0.69	0.04	1.69
sum of monoterpene SOA	4.38	4.00	1.17	13.5	7.03	3.79	3.22	12.7	2.82	0.90	1.36	4.09
Sesquiterpene SOA tracers (ng m⁻³)												
β -caryophyllinic acid	0.26	0.38	nd	1.03	0.22	0.42	nd	0.97	0.29	0.45	nd	1.33
total measured biogenic SOA tracers	10.2	10.2	1.80	34.7	15.8	5.75	8.14	24.3	9.69	4.92	2.36	18.6
Saturated dicarboxylic acids ($\mu\text{g m}^{-3}$)												
oxalic acid, C2	0.46	0.16	0.23	0.74	0.34	0.11	0.23	0.51	0.18	0.06	0.09	0.30
malonic acid, C3	6.43	2.10	1.51	8.71	10.0	2.41	8.50	14.3	5.96	2.41	1.48	8.14
succinic Acid, C4	0.04	0.02	0.01	0.07	0.03	0.02	0.01	0.06	0.01	0.01	nd	0.02
glutaric acid, C5	0.06	0.02	0.03	0.08	0.04	0.02	0.02	0.06	0.02	0.01	0.01	0.03

sum of saturated diacids	6.99	2.10	1.96	9.19	10.4	2.34	9.12	14.6	6.16	2.41	1.66	8.32
Unsaturated aliphatic diacids (ng m⁻³)												
maleic acid	8.32	5.35	0.86	20.2	21.3	9.11	11.5	33.1	10.79	13.1	1.00	41.9
fumaric acid	11.7	6.84	1.61	27.6	15.5	5.34	8.32	23.2	11.51	8.01	1.70	27.6
M/F	0.71	0.28	0.27	1.26	1.38	0.35	0.80	1.72	0.85	0.44	0.32	1.52
sum of unsaturated aliphatic diacids	20.0	11.8	2.48	47.8	36.8	13.9	21.0	56.3	22.3	20.5	2.70	69.5
Aromatic acids (ng m⁻³)												
phthalic acid (Ph)	8.02	3.05	3.00	12.4	10.5	1.77	8.09	12.8	5.88	3.73	1.45	13.0
isophthalic acid (iPh)	10.1	5.28	0.98	21.2	11.7	6.50	6.75	20.2	5.76	3.32	1.72	11.2
benzoic acid	5.46	2.76	0.47	11.4	5.88	0.52	5.01	6.29	4.47	2.44	1.07	8.41
sum of aromatic acids	23.6	10.2	8.30	45.1	28.1	8.24	21.1	39.3	16.1	8.86	4.25	30.3
Hydroxyl- and polyacids (ng m⁻³)												
glyceric acid	2.20	1.81	0.22	6.56	3.52	1.34	2.00	4.89	2.68	1.48	0.60	5.17
malic acid	3.00	1.45	0.95	5.73	4.32	2.06	1.52	6.60	3.67	1.88	0.77	6.51
tartaric acid	0.45	0.54	0.06	1.89	1.10	0.42	0.49	1.48	1.37	0.83	0.14	2.83
sum of hydroxyl and polyacids	5.66	2.63	1.24	10.4	8.94	3.73	4.01	12.2	7.73	4.14	1.51	14.5

Lignin and resin acids (ng m⁻³)													
4HBA, 4-hydroxybenzoic acid	2.10	2.89	0.36	9.32	2.50	0.86	1.09	3.31	3.40	2.26	0.05	6.02	
vanillic acid	1.12	2.05	0.00	5.96	2.50	0.98	1.23	3.53	4.76	3.36	0.02	8.98	
syringic acid	28.0	40.7	0.23	97.8	0.21	0.20	0.01	0.54	1.18	2.95	0.01	8.47	
dehydroabietic acid	15.3	4.80	4.30	22.7	14.4	7.91	8.22	23.4	17.0	14.0	5.45	40.9	
sum of lignin and resin acids	46.5	38.0	15.8	114	19.7	8.78	10.8	29.5	26.3	15.6	9.58	56.1	
α-Dicarbonyls (ng m⁻³)													
MeGly, methylglyoxal	20.4	29.2	7.47	103	10.1	4.93	6.55	18.7	6.43	3.04	2.12	10.4	
Other species ($\mu\text{g m}^{-3}$)													
MSA ⁻ , methanesulfonic acid	0.09	0.02	0.06	0.12	0.04	0.01	0.02	0.05	0.02	0.01	0.00	0.03	
formic acid	0.18	0.05	0.08	0.25	0.12	0.01	0.11	0.14	0.05	0.02	0.02	0.08	
acetic acid	0.22	0.11	0.07	0.44	0.16	0.07	0.08	0.27	0.05	0.01	0.03	0.06	

Note that: OC=organic carbon; TC=total carbon; WSOC=water-soluble OC; WISOC=water-insoluble OC. nss-Ca²⁺ refers to non-sea-salt Ca²⁺.
nd means not detected. Water-insoluble OC (WISOC) was calculated as the difference between OC and WSOC.

Table 2. Abundance and contributions of OC from primary sources (i.e., biomass burning, fungal spores, and plant debris) and from secondary formation (biogenic and anthropogenic VOCs) to OC in PM_{2.5}.

Abundance ($\mu\text{g m}^{-3}$)	PM _{2.5} concentration ($\mu\text{g m}^{-3}$) >200			100-200			<100					
	mean	std	min	max	mean	std	min	max	mean	std	min	max
BB-OC	5.79	1.50	3.48	8.86	2.27	0.34	1.74	2.69	2.47	1.48	0.72	4.86
Fungal spores-OC	0.44	0.14	0.21	0.62	0.42	0.09	0.32	0.52	0.29	0.18	0.09	0.68
plant debris-OC	0.29	0.12	0.07	0.55	0.45	0.21	0.28	0.80	0.23	0.08	0.10	0.35
sum of POC	6.52	1.62	3.77	9.65	3.14	0.46	2.48	3.67	2.99	1.56	1.23	5.39
Isoprene SOC	0.03	0.03	0.003	0.09	0.04	0.01	0.02	0.05	0.03	0.02	0.01	0.07
Monoterpene SOC	0.02	0.02	0.01	0.06	0.03	0.02	0.01	0.06	0.01	0.004	0.01	0.02
Sesquiterpene SOC	0.01	0.02	0.00	0.04	0.01	0.02	0.00	0.04	0.01	0.02	0.00	0.06
sum of BSOC	0.06	0.05	0.01	0.16	0.08	0.04	0.04	0.15	0.06	0.03	0.01	0.10
Naphthalene SOC	0.21	0.08	0.08	0.32	0.27	0.05	0.21	0.33	0.15	0.10	0.04	0.34
sum of SOC	0.26	0.11	0.09	0.49	0.36	0.07	0.28	0.44	0.21	0.12	0.05	0.41
total	6.79	1.68	3.86	9.98	3.50	0.50	2.76	4.07	3.20	1.57	1.54	5.62
Contribution to OC (%)												

BB-OC	16.3	3.39	10.6	23.6	9.63	0.56	8.96	10.3	15.9	7.01	8.29	26.5
Fungal spores-OC	1.23	0.31	0.74	1.63	1.81	0.47	1.23	2.32	2.38	2.26	0.56	7.50
plant debris-OC	0.83	0.39	0.30	1.74	1.99	1.02	0.98	3.48	1.69	0.75	0.62	2.44
sum of POC	18.4	3.62	12.2	25.7	13.4	1.97	11.3	16.0	19.9	8.31	11.5	31.3
Isoprene SOC	0.07	0.08	0.01	0.25	0.18	0.08	0.07	0.24	0.23	0.18	0.04	0.60
Monoterpene SOC	0.05	0.04	0.02	0.15	0.13	0.08	0.05	0.25	0.09	0.04	0.04	0.15
Sesquiterpene SOC	0.03	0.05	0.00	0.14	0.04	0.08	0.00	0.19	0.13	0.22	0.00	0.66
sum of BSOC	0.15	0.14	0.05	0.43	0.36	0.20	0.14	0.67	0.44	0.34	0.09	1.10
naphthalene SOC	0.59	0.21	0.27	0.88	1.17	0.22	0.88	1.46	1.12	0.84	0.29	2.46
sum of SOC	0.74	0.29	0.38	1.28	1.53	0.37	1.01	1.99	1.57	1.13	0.38	3.56
total	19.1	3.74	12.8	26.6	15.0	2.28	12.6	17.8	21.5	8.29	11.9	32.2

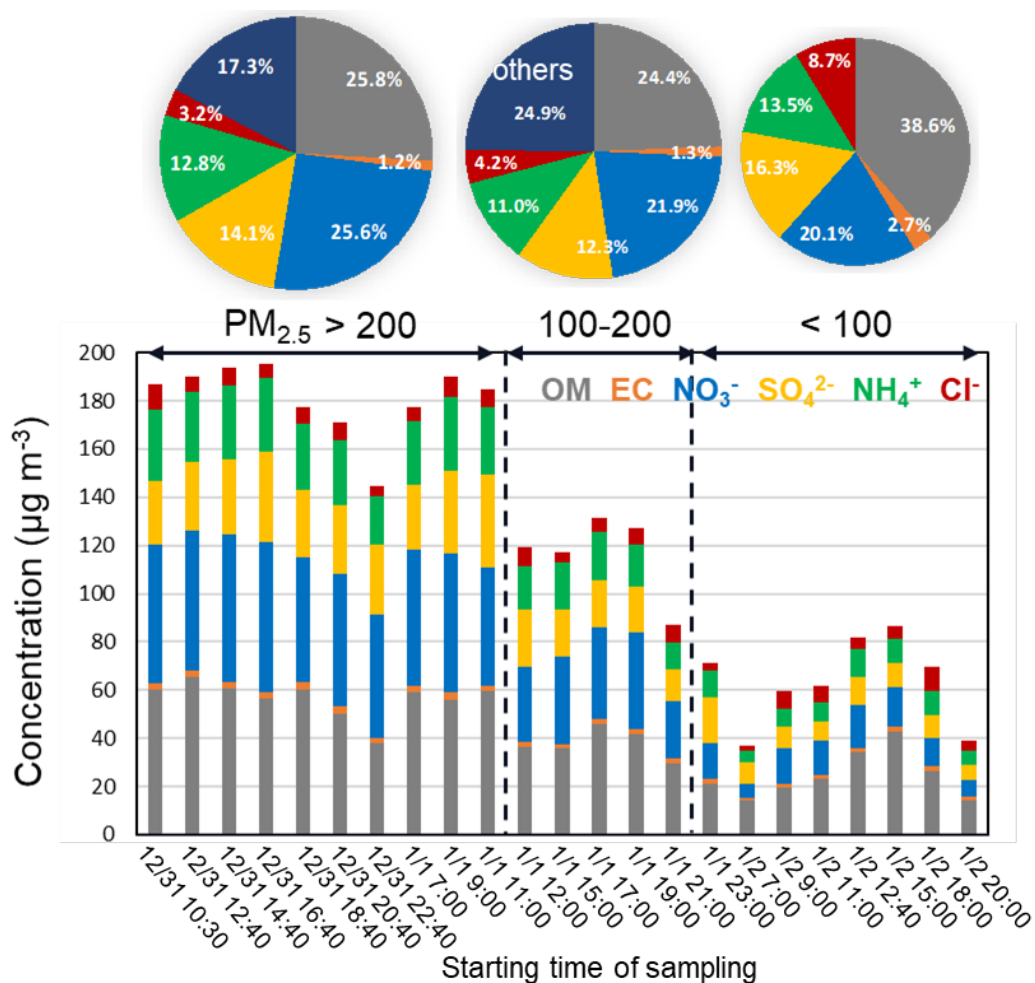


Figure 1. Temporal variations of dominant PM_{2.5} compositions based on different PM_{2.5} levels (i.e., <100, 100-200, and >200 µg m⁻³). The concentrations of organics (OM) were derived from OC concentration by multiplying it by a recommended factor of 1.6 (Turpin et al., 2001). Others represent the fine particles removing the organics, secondary inorganic aerosol (sulfate, nitrate, ammonium) and chloride. The pie charts present the average contribution of major components to PM_{2.5} during three pollution episodes.

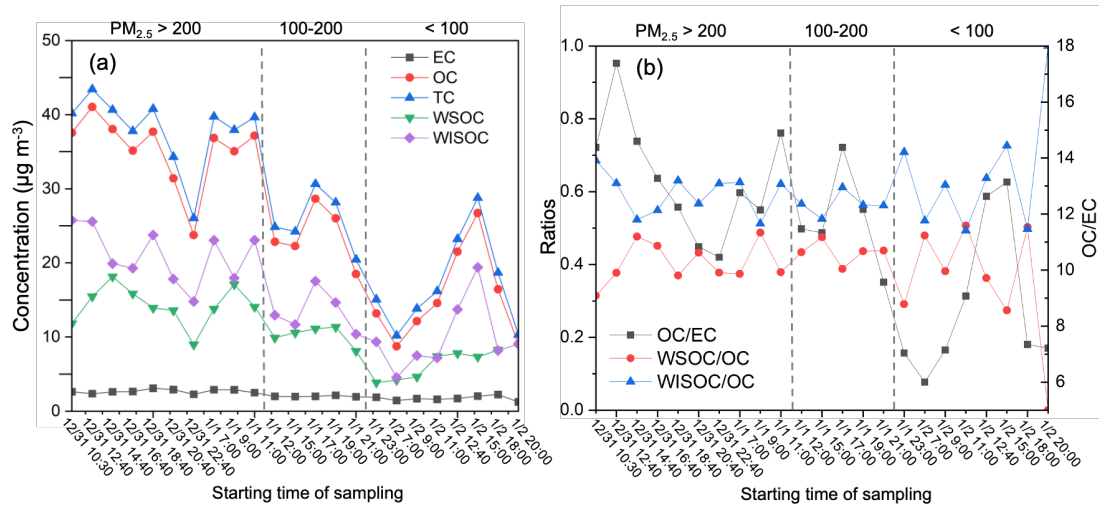


Figure 2. (a) Temporal variations of OC (organic carbon), EC (elemental carbon), WSOC (water-soluble organic carbon), WISOC (water-insoluble organic carbon), total carbon (TC) (units are $\mu\text{g m}^{-3}$), and (b) the ratios of OC/EC, WSOC/OC, and WISOC/OC in PM_{2.5} samples in Nanjing.

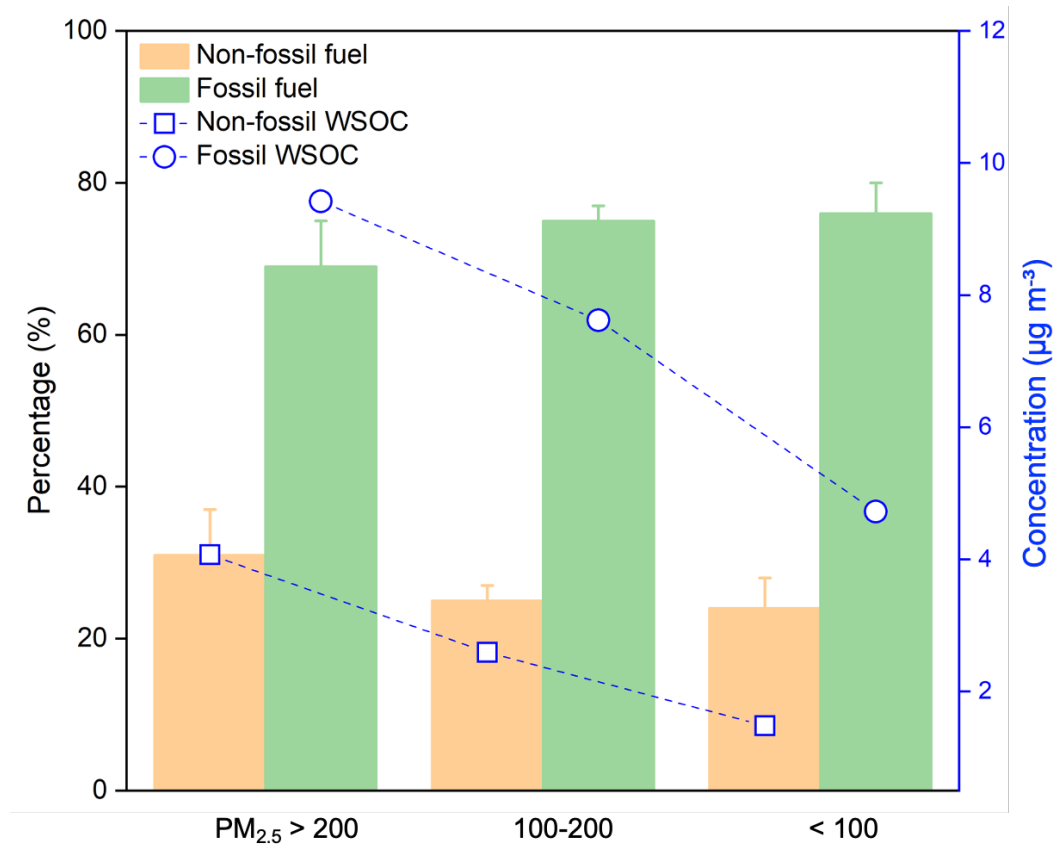


Figure 3. Comparison of fossil and non-fossil source contributions to water-soluble organic carbon (WSOC) in urban PM_{2.5} samples during three haze episodes (i.e., PM_{2.5} > 200, 100-200, and < 100 µg m⁻³).

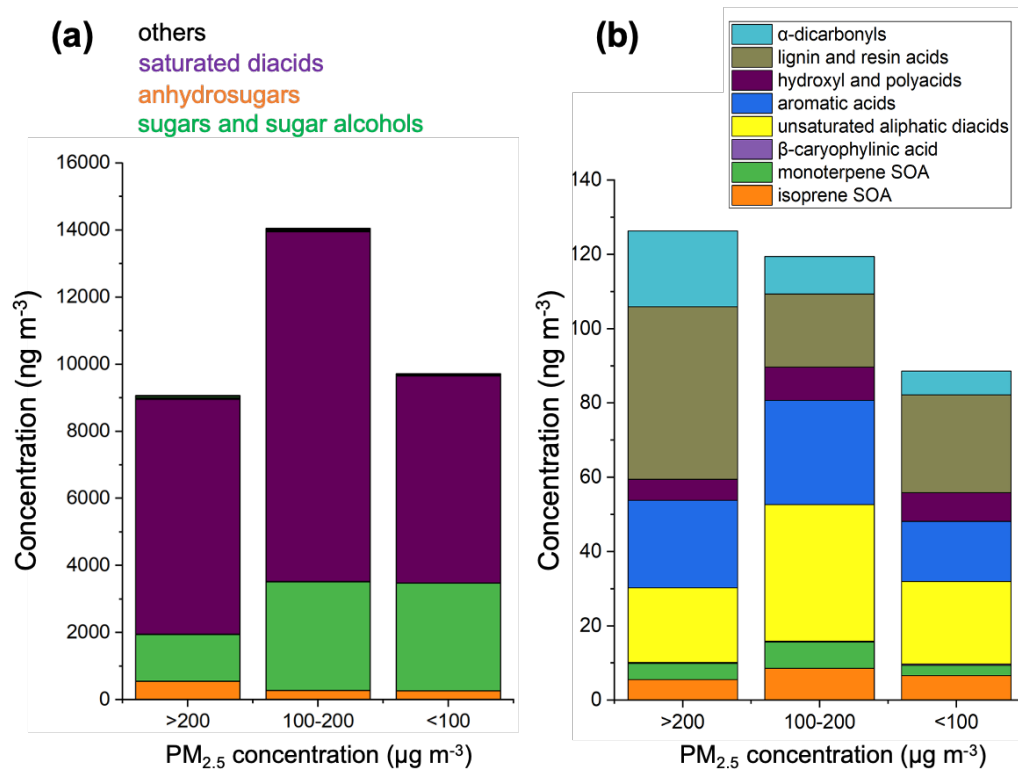


Figure 4. Average concentrations of measured carbonaceous species during three episodes with PM_{2.5} levels in the ranges of > 200, 100-200, and < 100 µg m⁻³, respectively. “others” in (a) denotes the sum of the components presented in (b).

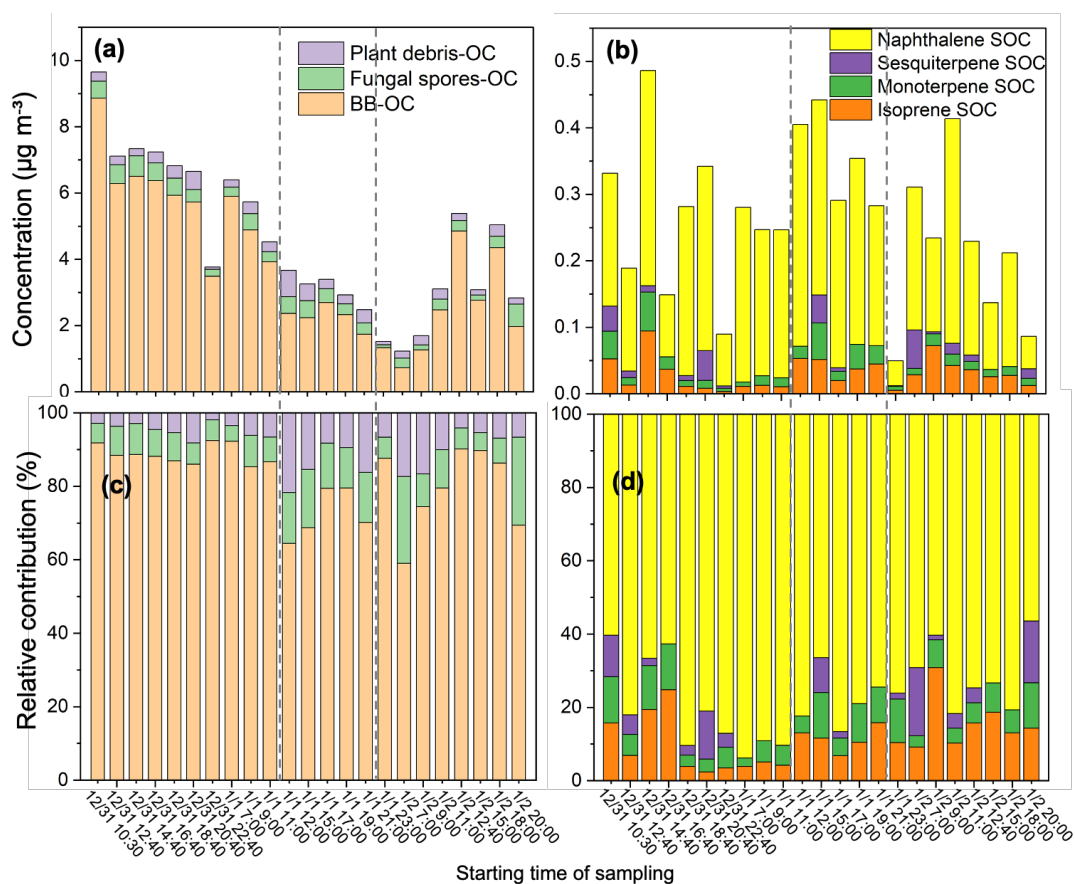


Figure 5. Concentrations of (a) primary organic carbon (OC) derived from biomass burning, fungal spores, and plant debris, and (b) secondary OC generated by isoprene, monoterpene, sesquiterpene, and naphthalene, and relative contribution of these OCs (c and d).

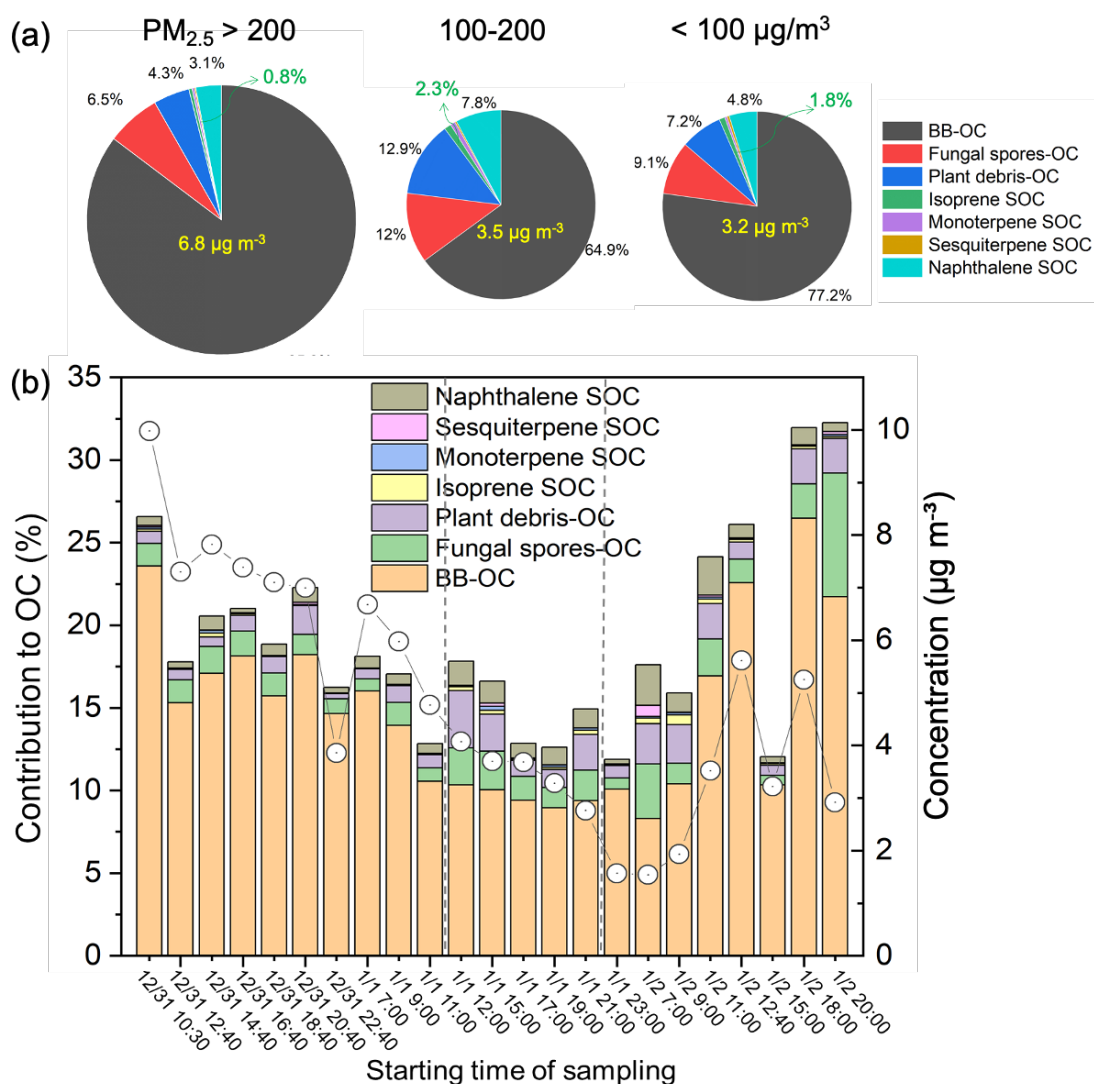


Figure 6. (a) Episode-averaged relative contributions of OC derived from biomass burning, fungal spores, plant debris, isoprene, monoterpene, sesquiterpene, and naphthalene to OC in $PM_{2.5}$ (%). The yellow numbers refer to the total tracer-based OC concentrations attributed to these sources ($\mu\text{g}/\text{m}^3$). Each pie size is proportional to its total tracer-based OC concentration. The green arrows and numbers represent the biogenic SOC fraction contributed by isoprene, monoterpene, and sesquiterpene. (b) Contributions of biomass burning, fungal spores, plant debris, isoprene, monoterpene, sesquiterpene, and naphthalene to OC in $PM_{2.5}$ (%), and OC concentrations attributed to these sources ($\mu\text{g}/\text{m}^3$, white circles).