- 1 Impact of fossil and non-fossil sources on the molecular compositions of water soluble humic-
- 2 like substance in PM_{2.5} at a suburb site of Yangtze River Delta, China
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19 Abstract

20 Atmospheric humic-like substances (HULIS) affect global radiation balance due to their strong light absorption at the ultraviolet wavelength. The potential sources and molecular 21 compositions of water soluble HULIS at a suburb site of Yangtze River Delta from 2017 to 2018 22 were discussed based on the radiocarbon (¹⁴C) analysis combining the Fourier Transform Ion 23 Cyclotron Resonance Mass Spectrometry (FT-ICR MS) technique in this study. The ¹⁴C results 24 showed that the averaged non-fossil source contributions to HULIS were 39 ± 8 % and 36 ± 6 % 25 in summer and winter, respectively, indicating the significant contributions from fossil sources to 26 27 HULIS. The Van Krevelen diagrams obtained from the FT-ICR MS results showed that the proportions of tannins-like and carbohydrates-like groups were higher in summer, suggesting 28 29 significant contribution of HULIS from biogenic secondary organic aerosols (SOA). The higher proportions of condensed aromatic structures in winter suggested increasing anthropogenic 30 emissions. Molecular composition analysis on the CHO, CHON, CHOS, and CHONS subgroups 31

showed the relatively higher intensities of high O-containing macromolecular oligomers in CHO 32 compounds in summer, further indicating stronger biogenic SOA formation in summer. High-33 intensity phenolic substances and flavonoids which were related to biomass burning and polycyclic 34 aromatic hydrocarbons (PAHs) derivatives indicating fossil fuel combustion emissions were found 35 in winter CHO compounds. Besides, two high-intensity CHO compounds containing condensed 36 aromatic ring structures ($C_9H_6O_7$ and $C_{10}H_5O_8$) identified in summer and winter samples were 37 similar to those from off-road engine samples, indicating that traffic emission was one of the 38 important fossil sources of HULIS at the study site. The CHON compounds were mainly composed 39 of nitro compounds or organonitrates with significantly higher intensities in winter, which was 40 associated to biomass burning emission, as well as the enhanced formation of organonitrates due 41 42 to high NO_x in winter. However, the high-intensity CHON molecular formulas in summer were 43 referring to N-heterocyclic aromatic compounds, which were produced from the atmospheric secondary processes involving reduced N species (e.g., ammonium). The S-containing compounds 44 45 were mainly composed of organosulfates (OSs) derived from biogenic precursors, long-chain alkane and aromatic hydrocarbon, illustrating the mixed sources of HULIS. Generally, different 46 47 policies need to be considered for each season due to the different season sources, i.e., biogenic emission in summer and biomass burning in winter for non-fossil source, traffic emission and 48 49 anthropogenic SOA formation in both seasons and additional coal combustion in winter. Measures to control emissions from motor vehicles and industrial processes need to be considered in summer. 50 Additional control measures on coal power plants and biomass burning should be concerned in 51 winter. These findings add to our understanding of the interaction between the sources and the 52 53 molecular compositions of atmospheric HULIS.

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55 **1. Introduction**

Atmospheric humic-like substances (HULIS) have been observed worldwide and can be produced from primary combustion of biomass, fossil fuel, as well as various secondary processes such as photochemical processes of volatile organic compounds (VOCs) and heterogeneous reactions of organic aerosols in the atmosphere (Kuang et al., 2015; Li et al., 2019; Ma et al., 2018; Sun et al., 2021). As important component of brown caron (BrC) aerosols, HULIS species have been widely reported to have a great impact on global radiative budget, contributing to 20-40% of the direct radiative forcing caused by light absorbing aerosols due to its light absorption at the

ultraviolet wavelength (Chung et al., 2012; Zhang et al., 2017; Zhang et al., 2020a; Wang et al., 63 64 2018c). HULIS are a highly complex mixture of polar organic compounds composed of aromatic and hydrophobic aliphatic structures containing carboxyl, carbonyl, and hydroxyl function groups 65 (Zheng et al., 2013; Graber and Rudich, 2006; Zhang et al., 2022b; Zhang et al., 2022c). During 66 the atmospheric secondary oxidation processes, the substitutions of hydrophilic functional groups 67 increased aerosol hygroscopicity (Huo et al., 2021; Jiang et al., 2020). Polycarboxylic acids in 68 HULIS are surface-active and play an important role in the cloud condensation nuclei (CCN) 69 70 activity (Tsui and McNeill, 2018). N-base compounds can promote the generation of atmospheric reactive oxygen species (ROS) which have a great impact on human health (Wang et al., 2017c; 71 De Haan et al., 2018; Song et al., 2022). Identifying the molecular compositions of HULIS is a 72 73 challenge due to complex mixtures contained in HULIS and can help to a better understanding of 74 the processes involving organic compounds in atmosphere (Noziere et al., 2015; Laskin et al., 2018). 75

76 The Fourier-Transform Ion Cyclotron Resonance Mass Spectrometry (FT-ICR MS) coupled with electrospray ionization (ESI) ion source have been widely used in identifying the chemical 77 78 structure of HULIS, providing high mass accuracy and can determine molecular formulas from 79 mixed compounds (Chen et al., 2016; Wang et al., 2019b; Lin et al., 2012a; Jiang et al., 2020). 80 Typical molecular formulas composed of C, H, and O atoms in HULIS were observed being 81 abundant in carboxylic acids, lignin-derived products, and polycyclic aromatic hydrocarbons 82 (PAHs) or their derivatives (Lin et al., 2012a; Sun et al., 2021; Jiang et al., 2020; Huo et al., 2021; Song et al., 2018). In addition, the HULIS formation of N and S containing precursors was also 83 widely detected. The N-containing compounds such as nitroaromatics were important 84 chromophores in HULIS in aged biomass burning organic aerosols (BBOA), as well as in ambient 85 86 aerosols influenced by biomass burning (BB), while reduced N compounds such as N-heterocyclic 87 aromatic compounds were found to be important chromophores in fresh BBOA (Wang et al., 2019b; Song et al., 2022; Jiang et al., 2020; Wang et al., 2017c). Recent laboratory simulation 88 experiments showed that the photooxidation of various anthropogenic VOCs (e.g., naphthalene, 89 benzene, toluene, and ethylbenzene) would be promoted under high NO_x condition, producing 90 91 strongly light absorbing nitroaromatics (Yang et al., 2022; Aiona et al., 2018; Siemens et al., 2022; Xie et al., 2017). Otherwise, nighttime oxidation of biogenic or anthropogenic VOCs, such as 92 benzene/toluene, isoprene (C_5H_8) and monoterpenes ($C_{10}H_{16}$) by NO₃ radicals lead to substantial 93

organonitrates formation, where the VOCs oxidation is strongly affected by NO_x (He et al., 2021;
Shen et al., 2021; Wang et al., 2020; Zheng et al., 2021).

The organosulfates (OSs) and nitrooxy organosulfates (nitrooxy-OSs) have also been found 96 to widely exist in HULIS in different atmospheric environment (Lin et al., 2012b; Lin et al., 2012a; 97 Sun et al., 2021). Field study and laboratory smog chamber experiments have confirmed that OSs 98 and nitrooxy-OSs in the atmosphere mainly come from the O₃, OH, or NO₃ oxidation of biogenic 99 VOCs such as isoprene, α/β -pinene as well as aromatic hydrocarbon in the presence of H₂SO₄/SO₂ 100 (Surratt et al., 2008; Glasius et al., 2021; Yang et al., 2020; Lin et al., 2012b; Huang et al., 2020). 101 Coal combustions were found to be important sources of the aromatic OSs and nitrooxy-OSs in 102 HULIS (Song et al., 2018). Besides, the long-chain alkanes were found to be important precursor 103 of OSs in atmospheric aerosol samples from urban area which was related to vehicle emissions 104 105 (Wang et al., 2019a; Tao et al., 2014).

Nanjing is one of the main cities in the Yangtze River Delta (YRD), which is one of the most 106 developed areas in China. Organic matter can account for 20-40 % of PM_{2.5} in the YRD area due 107 to the impact of complicated sources, especially anthropogenic emissions (Wang et al., 2017a; 108 109 Wang et al., 2016a). Studies have reported that BrC is an important contributor to aerosol light absorption in Nanjing and exhibited obvious seasonal variations, with peaks in wintertime, owing 110 111 to emissions from biomass burning, fossil fuel combustion, and secondary formation (Chen et al., 2018; Cui et al., 2021; Xie et al., 2020; Wang et al., 2018a). Recently, works on the field 112 113 observation of nitrated aromatic compounds (NACs) were conducted to explore the light absorption contributions of NACs to BrC and help to better understand the links between the 114 optical properties and molecular compositions of BrC (Gu et al., 2022; Cao et al., 2023). However, 115 as far as we know, understanding of the sources of atmospheric HULIS at molecular levels was 116 117 still limited. In this work, the molecular compositions of water soluble HULIS isolated from PM_{2.5} 118 samples collected in summertime and wintertime from 2017 to 2018 at Nanjing, China, were investigated combining the FT-ICR MS and radiocarbon (¹⁴C) analysis. We aim to obtain the 119 molecular characteristic differences of water soluble HULIS in summertime and wintertime and 120 to get a better understanding of the influence of different sources on the molecular compositions 121 122 of HULIS.

123 **2. Materials and methods**

124 2.1 Sample collection

The 24 h PM_{2.5} samples were collected on the roof of Wende building, which was about 21 125 m height from the ground at Nanjing University of Information Science and Technology (32.2° N. 126 118.7° E) using a high-volume sampler (KC-1000, Qingdao, China) at a flow rate of 300 L min⁻¹. 127 The study site was located in the northern suburb area of Nanjing, adjacent to G205 State Road 128 and surrounded by an industrial park and residential area. Generally, the study site was affected 129 130 by human activity, industrial emission, and traffic emission. The sample collection was conducted in summer from 12 August 2017 to 26 August 2017 and in winter from 31 December 2017 to 31 131 January 2018. A heavy haze event occurred from 31 December 2017 to 3 January 2018, thus the 132 sample frequency was adjusted to 2 h in daytime and 8 h in nighttime. Field blank filters were 133 performed before and after sample collection for each season. More details about the sample 134 collection can be found in previous research reported by Bao et al. (2022). The air pollutants data 135 136 including PM_{2.5}, SO₂ and NO₂ were provided by China National Environmental Monitoring Centre. Twelve samples were selected for further chemical analysis and the details about the sample 137 138 selection are described in Section 3.1 in this study.

139 2.2 Chemical analysis

140 The solid phase extraction (SPE) cartridge (Oasis HLB, 30 µm, 60 mg/cartridge, Waters, USA) was performed to isolate the water soluble HULIS in this study. Briefly, the prepared water 141 142 extracts passed through the pre-conditioned HLB cartridge firstly, then the retained HULIS on the HLB cartridge were eluted with 2% (v/v) ammonia/methanol and evaporated to dryness under a 143 144 gentle stream of nitrogen gas, then re-dissolved in ultrapure water for the measurement. The carbon fraction in HULIS (HULIS-C) were determinated using a total carbon analyzer (Shimadzu-TOC-145 VCPH, Shimadzu, Japan) with standard deviation of reproducibility test less than 3.5 % and 146 detection limit of 0.14 µg C m⁻³. More details about the HULIS isolation and measurement have 147 148 been described in Bao et al. (2022).

The mass concentrations of the water soluble ions including NO_3^- , NH_4^+ and SO_4^{2-} were measured using an ion chromatography (Dionex ICS-5000+, ThermoFisher Scientific, USA) separated on an AS11 column (4*250 mm, Dionex). Potassium hydrate (KOH) was used as the gradient eluent for anion determination. The levoglucosan concentrations were analyzed using the same ion chromatograph equipped with a CarboPac MA1 analytical column (4*250 mm, Dionex) and an electrochemical detector. Sodium hydroxide (NaOH) was used as the gradient eluent for levoglucosan determination. All data were blank corrected in this study. More details of themethods have been described previously (Liu et al., 2019).

157 2.3 Radiocarbon analysis

For the radiocarbon measurement of the HULIS samples, the organic solvents were firstly 158 evaporated under a gentle flow of ultrapure N₂ for 30-40 minutes in tin cups. After that, the tin 159 cups were wrapped into balls and more than 50 µg of carbon from the HULIS samples was 160 combusted into CO₂ using an elemental analyzer (EA, model vario micro, elemental, Germany), 161 then reduced into graphite targets for ¹⁴C determination at the State Key Laboratory of Organic 162 Geochemistry, Guangzhou Institute of Geochemistry, Guangzhou, China (Jiang et al., 2020). 163 Detailed descriptions of the ¹⁴C data processing can be found in previous study (Mo et al., 2018). 164 Briefly, the ¹⁴C values were expressed as the modern carbon (f_m) fraction after correcting for the 165 δ^{13} C fractionation. The f_m was converted into non-fossil carbon (f_{nf}) fraction with the correction 166 factor of 1.06 ± 0.07 based on the long-term time series of ${}^{14}CO_2$ sampled at the background station 167 in this study (Levin et al., 2013; Levin and Kromer, 2004). ¹⁴C analysis of the oxalic acid standard 168 (IAEA-C7) was conducted in this study (Xu et al., 2021). No field blank correction was performed 169 170 for the carbon isotope analysis since the carbon content in the field blanks was negligible.

171 2.4 High-resolution FT-ICR MS analysis

172 The ultrahigh resolution mass spectra of the HULIS samples were obtained through a SolariX XR FT-ICR MS (Bruker Daltonics, GmbH, Bremen, Germany) equipped with a 9.4 T 173 174 superconducting magnet (Gamry Instruments, Warminster, USA) and a Paracell analyzer cell (Brucker Daltonik GmbH, Bremen, Germany) in the negative ESI mode. The detection mass range 175 176 was set as m/z 150 to 800 and the ion accumulation time was set as 0.65 s. A total of 100 continuous 4M transient data points were superposed to enhance the signal to noise ratio and dynamic range. 177 178 The mass spectrum was externally calibrated with a standard solution of arginine and internal 179 recalibration was performed using typical O_6S_1 chemical species in DataAnalysis ver. 4.4 software (Bruker Daltonics) (Mo et al., 2018; Tang et al., 2020; Jiang et al., 2020). Field blank filters were 180 181 analyzed as same as the samples and all the sample data were blank corrected. More details about the data processing can be found in Text S1 in the supporting information. 182

183 **3. Results and discussion**

184 3.1 General temporal characteristics during the sampling periods

Figure 1 displays the temporal variations of non-fossil contributions to HULIS-C, the mass 185 concentrations of HULIS-C, levoglucosan, NO₃⁻, SO₄²⁻, NH₄⁺, SO₂, NO₂, and PM_{2.5}, as well as the 186 187 relative humidity and temperature during the study periods corresponding to the 12 samples. The 12 samples were named as S1-S6 (summer) and W1-W6 (winter) in chronological order 188 corresponding to the six samples in summer and winter, respectively in this study. The averaged 189 mass concentrations of PM_{2.5} in summer and winter during the selected periods were 21.05 ± 8.05 190 μ g m⁻³ and 445.67 ± 275.00 μ g m⁻³, respectively, indicating the serious pollution level in winter. 191 The daily PM_{2.5} mass concentrations in summer were all below the daily averaged Chinese 192 National Ambient Air Quality Standard (NAAQS) of 35 μ g m⁻³, while the daily PM_{2.5} mass 193 concentrations in winter all exceeded the daily averaged NAAOS of 35 μ g m⁻³, of which the PM_{2.5} 194 mass concentrations of W1-W3 and W6 exceeded 200 µg m⁻³. The averaged mass concentrations 195 of HULIS in summer and winter during the selected periods were 1.83 \pm 0.27 μ g m⁻³ and 4.52 \pm 196 2.29 µg m⁻³, respectively. The averaged HULIS concentration in summer was comparable with 197 those measured in other cities in China, i.e., 1.70 µg m⁻³ in Guangzhou, 1.61 µg m⁻³ in Shanghai 198 and 1.50 µg m⁻³ in Xi'an. Compared with those measured in winter samples in other cities, our 199 result was comparable with those in Xi'an (4.50 μ g m⁻³), a little lower than those in the megacity 200 of Shanghai (5.31 μ g m⁻³) and higher than those in the southern coastal city of Guangzhou (3.6 μ g 201 m⁻³) (Fan et al., 2016; Zhang et al., 2020b; Zhao et al., 2016). 202

As shown in Fig. 1, the mass concentrations of HULIS-C, levoglucosan, water soluble 203 204 secondary inorganic aerosols (SIA), and air pollutants showed similar trends in winter, suggesting the influence of BB and anthropogenic emissions in winter (Wu et al., 2019b). The radiocarbon 205 analysis results showed that the f_{nf} of HULIS-C ranged from 30 % to 50 % with an average 206 contribution of 39 ± 8 % in summer and ranged from 32 % to 48 % with an average contribution 207 208 of 36 ± 6 % in winter, indicating the significant contributions from fossil sources to HULIS at the 209 study site. The 48 h back trajectories (Fig. S1) showed that the study site was affected by the polluted air masses mainly from the northern cities in winter, suggesting the coal combustion 210 contributions to HULIS in winter (Ma et al., 2018; Sun et al., 2021). In addition, significant 211 212 increasing of the levoglucosan and HULIS-C mass concentrations were found from 31 December 213 2017 to 1 January 2018, corresponding to the W1-W3 samples and the maximum of the levoglucosan and HULIS-C mass concentrations were 552.79 ng m⁻³ and 7.40 µg m⁻³, respectively, 214 indicating the BB impact during the periods. In summer, the study site was affected by both 215

regional transport from the nearby cities in the north and west of Nanjing and the Donghai Sea. The anthropogenic emissions from the neighboring cities might cause the anthropogenic SOA formation, i.e., secondary N-containing and S-containing compounds with aromatic structures during the atmospheric transport processes, which was discussed in detail in section 3.4 in this study.

3.2 Mass spectra and molecular formula assignments

Figure S2 and S3 show the negative ion ESI FT-ICR mass spectra of HULIS in summer and 222 223 winter, respectively. The molecular formulas listed are some of the top ten molecular formulas. Thousands of peaks are present in the spectra in the range from m/z 150 to m/z 600 and the most 224 intense ion peaks are those in the range m/z 200-400 in summer and m/z 150-350 in winter. Our 225 results are similar to those found for the ultrahigh resolution mass spectra of water-soluble organic 226 227 compounds in particles produced from BB, coal combustion, vehicle exhaust emissions, as well as in ambient aerosols and cloud water samples, within a reasonable range (Tang et al., 2020; Sun et 228 al., 2021; Song et al., 2018; Song et al., 2019; Bianco et al., 2018). In this study, the assigned 229 molecular formulas were classified into the following four main subgroups based on their 230 231 elemental compositions: CHO (compounds containing only C, H, and O), CHON (compounds containing C, H, O and N), CHOS (compounds containing C, H, O, and S), and CHONS 232 233 (compounds containing C, H, O, N, and S). As shown in Fig. 2, the proportions of the four subgroups accounted for the overall formulas followed as CHO (20 %-27 %), CHON (28 %-43 %), 234 235 CHOS (19 %-26 %), and CHONS (16 %-26 %) in summer, respectively and CHO (15 %-19 %), CHON (30 %-40 %), CHOS (21 %-32 %), and CHONS (20 %-29 %) in winter, respectively. The 236 237 average proportions of the CHO, CHON, CHOS, and CHONS compounds in summer were $22 \pm$ 3%, $36 \pm 5\%$, $22 \pm 3\%$, and $20 \pm 4\%$, respectively. The average proportions of the four subgroups 238 239 in winter were 17 ± 2 %, 32 ± 4 %, 24 ± 3 %, and 27 ± 4 %, respectively. The CHON groups were 240 the major components of molecular formulas, furthermore, the relative intensity of CHON groups increased significantly in winter (Fig. S2 and Fig. S3). Studies have suggested that HULIS emitted 241 242 from biomass burning can produce a high abundance of CHON compounds and S-containing compounds were the dominant component for primary HULIS emitted from coal combustion 243 244 (Zhang et al., 2021; Song et al., 2018). The higher intensity of CHON compounds in winter in this study further indicated the BB contribution. The contributions of S-containing compounds (CHOS 245 and CHONS groups) increased in winter which might be related to the polluted air masses 246

transported from the northern cities with increasing coal combustions emissions in winter (Song et al., 2018). Notably, the relatively higher proportions of CHO and CHON groups in summer were most probably related to the increasing biogenic emissions in summer, resulting in the formation of some high molecular weight oligomers or highly oxidized organonitrates, which was discussed in detail in section 3.4.1 and 3.4.2 in this study.

252 Table S1 and S2 displays the composition characteristics of atmospheric HULIS in the summer and winter samples, including the relative intensity weighted average values of number, 253 molecular weight (MW_w), elemental ratios (O/C_w and H/C_w), double-bond equivalent (DBE_w), 254 255 aromaticity index (AI_w), and DBE/C_w. A total of 14387 and 15731 peaks were detected in the summer and winter samples, respectively. The O/C and H/C ratios are commonly calculated to 256 evaluate the oxidation degree and saturation degree of the compounds, respectively (Ning et al., 257 258 2022). The O/C_w values were in a range of 0.61-0.80 with an average value of 0.71 \pm 0.07 for summer samples and in a range of 0.59-0.67 with an average value of 0.62 \pm 0.03 for winter 259 260 samples, respectively. The higher oxidation degree of summer samples than winter samples indicated stronger secondary HULIS formation in summer. The H/Cw values were in a range of 261 262 1.38-1.46 with an average value of 1.42 ± 0.03 for summer samples and in a range of 1.33-1.41 with an average value of 1.36 ± 0.04 for winter samples, respectively. The O/C_w and H/C_w of each 263 264 molecular subgroup followed a changing trend of CHO < CHON < CHOS < CHONS compounds. Most of the S-containing compounds had a O/C value ≥ 0.7 , suggesting the large amounts of highly 265 266 oxidized OSs in S-containing compounds which contained various functional groups and were mainly from the photochemical oxidation of biogenic or anthropogenic volatile organic 267 268 compounds (VOCs) (Mutzel et al., 2015). The DBE values were calculated to describe the degree of unsaturation of compounds and restricted the assigned molecular formulas with unreasonably 269 270 high or low number of rings or double bonds (Kroll et al., 2011). The related parameter DBE/C 271 was the double-bond equivalent of unit carbon which can reflect the condensed ring structures in the compounds (Jiang et al., 2021). The higher DBEw and DBE/Cw values of CHO and CHON 272 273 compounds were found in this study, indicating the higher unsaturation degree of these two groups.

Considering that double bonds can be formed by heteroatoms especially O atoms, whereas make no contributions to the aromaticity of the compounds, AI_w was calculated to supplement the DBE results (Song et al., 2018; Ning et al., 2019). AI_w can eliminate the contribution of O, N, and S atoms to the C=C double bond density of molecules. The AI_w values of different compounds

groups in HULIS presented the changing trends: AI_w (CHONS) > AI_w (CHON) > AI_w (CHO) > 278 AI_w (CHOS) in summer and AI_w (CHON) > AI_w (CHO) > AI_w (CHONS) > AI_w (CHOS) in winter, 279 respectively. The formulas can be classified into three parts based on AI values proposed by 280 previous studies: aliphatic (AI =0), olefinic ($0 \le AI \le 0.5$) and aromatic (AI >0.5) (Koch and Dittmar, 281 2006). As shown in Fig. S4 and S5, the aliphatic were the main components of S-containing 282 compounds in this study and the olefinic and aromatic were the main components of CHO and 283 CHON compounds. Furthermore, the aromatic proportion of CHO and CHON compounds 284 285 significantly increased in winter, suggesting the increasing anthropogenic emissions in winter.

286 3.3 Comparative analysis using Van Krevelen diagrams

In this study, the Van Krevelen diagrams (Fig. 3) were constructed to display the molecular 287 composition and categorical distribution of the collected samples (Noziere et al., 2015; Patriarca 288 289 et al., 2018; Li et al., 2022). According to the elemental ratios (O/C and H/C ratios) and AI values, seven major compound classes were classified, including lipids-like species, lignins-like species, 290 proteins-like species, tannins-like species, carbohydrates-like species, condensed aromatics 291 structure, and unsaturated hydrocarbons (Table S3). The Van Krevelen diagrams showed similar 292 293 distributions in the 12 samples. The CHO and CHON compounds located in the lower left area 294 and the S-containing compounds located in the upper light area with higher O/C and H/C ratios, 295 indicating a higher degree of oxidation and saturation. The condensed aromatic structure mainly 296 consisted in the CHO and CHON compounds, further suggesting the influence of anthropogenic 297 emissions on the formation of CHO and CHON compounds.

Figure 4 presents the averaged relative contributions of the number of molecular formulas 298 299 from the seven categories in summer and winter samples, respectively. Lignins-like species accounted for the highest proportion of CHO compounds with average contributions of 58 % and 300 301 61 % in summer and winter, respectively, followed by CHON compounds with average 302 contributions of 48 % and 57 % in summer and winter, respectively. Lignins are mainly composed of carboxyl groups, alicyclic rings, aromatic rings, and other O-containing groups. Previous studies 303 304 have reported that lignin was a complex phenolic polymer which usually came from direct biological emissions or combustions of biofuel (Ning et al., 2019; Boreddy et al., 2021; Sun et al., 305 306 2021). Lignins pyrolysis products and other lignins derived molecules have been shown to be 307 oxidized into light absorbing BrC chromophore under certain conditions (Fleming et al., 2020).

Tannins-like species accounted for 21 %, 27 %, 23 %, and 30 % of CHO, CHON, CHOS, and 308 CHONS compounds, respectively in summer which were higher than those in winter with 309 310 contributions of 13 %, 16 %, 16 %, and 23 % to CHO, CHON, CHOS, and CHONS compounds, respectively. Tannins-like species are a series of polyphenolic compounds containing hydroxyls 311 and carboxylic groups which have been widely reported in fogs, cloud water and aerosol samples, 312 313 attributing to highly oxidized organic compounds such as OSs or nitrooxy-OSs produced from the nighttime chemistry between the biogenic VOCs with the NO₃ (Altieri et al., 2009; Bianco et al., 314 315 2018; Ning et al., 2019; Altieri et al., 2008; Shen et al., 2021). Carbohydrates-like species which contain monosaccharide, alditols, and anhydrosugars mainly consisted in CHONS compounds 316 which also had a relative higher proportion of 33 % in summer than that of 29 % in winter (Sun et 317 al., 2021). C₁₀H₁₆NO₇₋₉S, as monoterpene nitrooxy-OSs, showing high relative intensities, were 318 319 typical carbohydrates-like species detected in this study which represented biogenic secondary organic aerosols (SOA) (Ning et al., 2019; Surratt et al., 2008; Wang et al., 2020). Both the higher 320 321 proportions of tannins-like and carbohydrates-like classes in summer indicated stronger biogenic SOA formation in this study. 322

Proteins-like classes mainly consisted in CHOS compounds with average proportions of 29 % and 38 % in summer and winter, respectively. Proteins contain peptide-like structures formed by dehydration with different kinds of amino acids and consist of short chains of amino acid residues (Bianco et al., 2018). These compounds are associated with photochemical oxidation processing in aerosols, thus resulting in the significant formation of OSs from biogenic or anthropogenic precursors in this study (Bigg and Leck, 2008).

329 Higher condensed aromatics were detected in winter with average proportions of 14 % in 330 CHO compounds and 8 % in CHON compounds, respectively which were 2-2.5 times of those in 331 summer. Condensed aromatics are important components of PAHs which were usually emitted 332 from incomplete combustion of fossil fuels (Ma et al., 2020). The increase of the proportion of condensed aromatics in winter indicated the stronger influence of anthropogenic sources on 333 334 HULIS formation. The unsaturated hydrocarbons and lipids-like species showed the lowest molecular number percentage of less than 1 % in this study. Previous studies have shown that the 335 336 lipids-like species were the main components of water insoluble organic compounds in aerosols and could be attributed to monocarboxylic acids (Ning et al., 2022; Wozniak et al., 2008). 337

In summary, both the summer and winter samples were mainly composed of compounds from 338 biogenic origins (lignins-like, tannins-like, proteins-like, and carbohydrates-like species). More 339 340 tannins-like and carbohydrates-like species were detected in summer including large amounts of highly oxidized OSs or nitrooxy-OSs, indicating biogenic SOA formation. More condensed 341 aromatic structures in CHO and CHON compounds were detected in winter, owing to increasing 342 anthropogenic emissions. It is noted that ESI ionization technology is more sensitive for the 343 identification of polar compounds. Therefore, the low polar or nonpolar compounds, such as PAHs 344 or their derivatives from fossil sources, were probably underestimated in this study (Jiang et al., 345 2014; Lin et al., 2018). 346

347 3.4 Molecular composition of HULIS

348 3.4.1 Molecular characteristics of CHO compounds

349 The O/C_w and H/C_w ratios for the CHO compounds were 0.45-0.56 and 1.15-1.30 for the summer samples and 0.42-0.48 and 0.90-1.02 for the winter samples (Table S1 and S2). The 350 351 summer samples showed higher oxidation degree and saturation degree. We firstly plotted the Van Krevelen diagrams of the four molecular subgroups showing relative intensities for all the 12 352 353 samples and similar distributions of the high-intensity compounds were found in the 6 summer samples and the 6 winter samples, respectively. Then we combined all the data in summer and 354 355 winter, respectively. As shown in Fig. 5a and 5d, the CHO compounds in summer with high relative abundance were located at the area within $0.2 \le O/C \le 1.0$ and $1.0 \le H/C \le 1.7$, mainly 356 357 including lignins-like species and tannins-like species which were closely related to biogenic 358 emissions. On the contrary, the condensed aromatics showed high relative abundance in winter, 359 suggesting obviously different sources of HULIS in summer and winter. The DBE values increased with the increasing of the C numbers (Fig. 5b and 5e). The high-intensity CHO 360 361 compounds in HULIS had DBE values between 3-7 with C numbers from 10 to 20 for summer 362 samples. In winter, the high-intensity CHO compounds had DBE values between 7-11 with C numbers from 5 to 15. As mentioned above, the aromatic (AI >0.5) proportion of CHO compounds 363 significantly increased in winter, the higher DBE values in winter further indicated the consists of 364 more highly unsaturated aromatic compounds which reflected the anthropogenic emissions. 365

The CHO compounds were classified according to the number of oxygen atoms to evaluate the oxygen content. As shown in Fig. 5c and 5f, the high-intensity CHO compounds with 6-11 oxygen atom were detected in summer, such as $C_{15}H_{24}O_6$, $C_{15}H_{22}O_{10}$, $C_{18}H_{26}O_8$, and $C_{18}H_{26}O_9$,

these highly oxygenated organic molecules with high molecular weight have also been detected in 369 laboratory α-pinene ozonolysis SOA (Pospisilova et al., 2020). We further classified the CHO 370 371 compounds by different carbon atom numbers. As shown in Fig. S6, the C₁₇-C₂₂ compounds were the main components of the CHO compounds, accounting for more than 50 % of the total number 372 of CHO molecular formulas in both summer and winter seasons. However, the total relative 373 374 intensities of the CHO compounds in summer were significantly higher than those in winter, of which the C₂₃-C₂₆ and C₂₇-C₃₂ compounds were enriched in summer. These high molecular weight 375 compounds were probably oligomers formed from various biogenic precursors, such as isoprene, 376 377 sesquiterpene, and monoterpene (Daellenbach et al., 2019; Berndt et al., 2018). The high intensities of these compounds in summer further indicated the stronger biogenic SOA formation in summer 378 compared with that in winter. 379

High-intensity CHO compounds with 4-9 oxygen atom were detected in winter (Fig. 5c) of 380 which the C₁₄H₁₀O₄ formula with a DBE value of 10 appeared the highest intensity, which was 381 382 probable functional PAHs and have been reported in HULIS from coal combustion smoke particles (Song et al., 2019). As shown in Fig. S2 and S3, the $C_{14}H_{10}O_4$ formula appeared high intensity in 383 all the winter samples, providing the evidence of coal combustion emissions in winter. Some other 384 high-intensity compounds in winter, such as $C_{14}H_8O_4$ and $C_{14}H_8O_5$ both with DBE values of 11, 385 386 and C₁₃H₈O₂, C₁₃H₈O₅, and C₁₃H₈O₆ with DBE values of 10, might refer to hydroxyl substitutions 387 derived from anthracenedione and xanthone, respectively, which have been reported in secondary 388 wood combustion products (Bruns et al., 2015). C₁₅H₁₀O₆, C₁₅H₈O₆, and C₁₆H₁₂O₇ which had 389 DBE values of 11, 12, and 11, respectively, might be flavonoids which had flavone backbone, the 390 key structure of plant pigments, widely existing in plants in nature and could be important sources 391 of BrC chromophores in aged BBOA (Fleming et al., 2020; Lin et al., 2016; Huang et al., 2021). 392 Phenolic substances derived from phenol, guaiacol, and syringol are also widely existed in BBOA, 393 usually from the pyrolysis of lignins in wood, which also play an important role in aqueous-phase SOA formation (Boreddy et al., 2021). For instance, C₁₃H₁₀O₃ and C₁₃H₁₀O₅ are guaiacol 394 derivatives, C₁₅H₁₆O₈ are syringol derivatives and C₁₈H₁₄O₆ and C₁₈H₁₄O₇ are phenol derivatives 395 (Sun et al., 2021). As shown in Fig. S7, the relative intensities of the CHO compounds mentioned 396 397 above produced from BB were found to have similar trends with the mass concentrations of 398 levoglucosan, which were significantly higher in W1-W3 samples, corresponding to the BB period

from 31 December 2017 to 1 January 2018, providing the evidence of BB influence on HULISformation in winter.

401 It is noted that the top compounds $C_9H_6O_7$ and $C_{10}H_6O_8$ were detected both in the summer and winter samples (Fig. S2 and S3), which had DBE values of 7 and 8, respectively, containing 402 abundant condensed aromatic ring structures with high O numbers. Their peaks were also detected 403 404 in the HFO (heavy-fuel-oil)-fueled off-road engine samples reported before, suggesting the traffic emission contributions to HULIS (Cui et al., 2019). This supported the radiocarbon analysis results 405 406 in this study and gave further information that the traffic emissions were important fossil sources 407 in both summer and winter seasons, which was also found in previous research which reported the sources of HULIS based on the positive matrix factorization (PMF) model by Bao et al. (2022). 408

409 3.4.2 Molecular characteristics of CHON compounds

410 The O/C_w of CHON compounds in summer and winter were 0.57-0.71 and 0.52-0.56, respectively, while the H/C_w were 1.20-1.32 and 1.00-1.11, respectively (Table S1 and S2). 411 Compared with the summer CHON compounds, the winter CHON compounds presented 412 significantly higher ion abundance (Fig. 6a and 6d). The most abundant CHON subgroups had 413 414 DBE values of 4-7 and 3-10 in summer and winter, respectively (Fig. 6b and 6e). Similar with the CHO compounds, the higher DBE values of high-intensity CHON compounds in HULIS in winter 415 416 indicated a high prevalence of double bonds or ring structures. According to the N and O number, 417 the CHON compounds were classified into N₁O_x (N₁O₁-N₁O₁₅) and N₂O_x (N₂O₂-N₂O₁₄) subgroups 418 in summer and N₁O_x (N₁O₁-N₁O₁₂) and N₂O_x (N₂O₂-N₂O₁₂) subgroups in winter, respectively (Fig. 6c and 6f). NO₈₋₁₂ and NO₆₋₉ compounds were mostly enriched subgroups in summer and winter, 419 420 respectively. More oxygen-enriched CHON compounds containing O number above 9 were 421 detected in summer, implying the higher oxidation degree for summer samples. In addition, the 422 N_1O_x were both the major compounds represented average of 64 ± 4 % and 61 ± 6 % of the CHON 423 molecular formulas in summer and winter, respectively, indicating the presence of more single nitro/amino substituents in CHON compounds in this study. 424

Among the CHON compounds, 95 ± 1 % and 86 ± 3 % CHON compounds had O/N values ≥ 3 in summer and winter, respectively in this study, indicating these compounds contained large amounts of oxidized nitrogen functional groups such as nitro compounds (-NO₂) and/or organonitrates (-ONO₂) and excess oxygen atoms indicated the existence of other oxygencontaining functional groups (Laskin et al., 2009). The organonitrates formation from NO₃ 430 oxidation of biogenic or anthropogenic VOCs can affect the interactions between anthropogenic 431 and natural emissions (He et al., 2021; Shen et al., 2021; Wang et al., 2020). Organonitrates were 432 found to be important species contributing to SOA formation in the polluted urban environment, 433 which were enhanced under high NO_x level (Zheng et al., 2021). The significant higher relative 434 intensities of CHON compounds in winter indicated that the high NO_x environment in winter 435 promoted the formation of organonitrates and highlighted the importance of orgnonitrates for SOA 436 control in polluted environment.

437 Furthermore, we found that the increase of the relative abundance of CHON compounds in winter was particularly significant in W1-W3 samples (Fig. S2 and S3), corresponding to the BB 438 episode. Phenols produced from the pyrolysis of lignins can react with NO₃ radicals in the 439 440 atmosphere, producing nitrophenols, which have been shown to be important BrC chromophore 441 in BBOA (Wang et al., 2017c; Lin et al., 2016; Cai et al., 2020). It was reported that the gas-phase reactions of NO₃ radicals with phenolic substances took place at least 4 orders of magnitude faster 442 443 than those with aromatic hydrocarbon and even faster in the aqueous phase (Lin et al., 2017). Among the top CHON compounds with high relative abundance in W1-W3 samples, such as 444 445 C₆H₄N₂O₆ and C₇H₆N₂O₆ both with a DBE value of 6, were refer to nitrophenols containing one or two nitrogen-containing functional groups, which have been widely reported in aged BBOA, 446 447 indicating the increasing of the CHON compounds relative intensity in W1-W3 samples were 448 closely related to BB (Lin et al., 2017; Cai et al., 2020; Mohr et al., 2013; Kourtchev et al., 2016; 449 Lin et al., 2016). Some other top CHON compounds in winter samples such as C₉H₄NO₄ and C10H6NO4 with low O/C and H/C ratios most likely indicated the presence of condensed aromatic 450 451 structures in the compounds. The C₉H₄NO₄ compounds were most likely emitted from vehicle 452 emissions which have previously been reported (Cui et al., 2019).

453 It is worth noting that some high-intensity CHON compounds with low O/C and H/C ratios 454 were detected in summer samples in this study (Fig. 6a), which were closely related to aromatic compounds from anthropogenic emissions. The top compounds with molecular formulas of 455 C₈H₅N₂O₂ and C₁₉H₁₁N₂O₄, which had O/N of 1 and 2, respectively, were both reduced N 456 compounds referring to N-heterocyclic compounds. Previously studies have found that the N-457 458 heterocyclic aromatic compounds can be formed through the aldehyde-ammonia reactions (De 459 Haan et al., 2018; Zhang et al., 2022a). This indicated the important role of reduced N species (e.g., 460 ammonium) in the formation of anthropogenic SOA in summer. Our results were consistent with previous study conducted in Xi'an, China which also found formation of reduced N compounds in
light-absorbing aerosols through ammonia involved reactions in summer (Zeng et al., 2021).

463 3.4.3 Molecular characteristics of S-containing compounds (CHOS and CHONS compounds)

The O/C_w of CHOS compounds in summer and winter were 0.60-0.79 and 0.56-0.67, 464 respectively, while the H/Cw were 1.50-1.54 and 1.53-1.72, respectively. The O/Cw of CHONS 465 compounds in summer and winter were 0.82-1.01 and 0.76-0.94, respectively, while the H/C_w 466 were 1.57-1.65 and 1.58-1.66, respectively (Table S1 and S2). As shown in Fig. 7a, 7d, 8a, and 8d, 467 468 the high-intensity S-containing compounds in summer and winter were both located at the area where O/C > 0.5 and H/C > 1.5, respectively. In addition, the relative intensity of S-containing 469 compounds increased with the O/C ratios, suggesting the S-containing compounds were highly 470 oxidized. A small number of high-intensity S-containing compounds with O/C <1.0 and H/C <1.0 471 472 were also found in winter in this study, which might be related to OSs and nitrooxy-OSs produced from the oxidation of aromatic hydrocarbon. The CHOS compounds presenting high relative 473 474 abundance were rich in $O_{6-9}S$ and $O_{5-7}S$ groups in summer and winter, respectively, of which the DBE values were all below 4. The CHONS compounds were rich in O₈₋₁₀S and O₇₋₉S groups in 475 476 summer and winter, respectively, of which the DBE values were all below 6 (Fig. 7b, 7e, 7c, 7f, 8b, 8e, 8c, and 8f). Compared with those of the CHO and CHON compounds, the DBE values of 477 478 S-containing compounds were significantly lower.

479 Among the S-containing compounds, more than 95 % of the CHOS, CHON₁S, and CHON₂S 480 formulas had O/S ratios greater than 4, 7, and 10, respectively, implying these compounds may contain organic sulfate functional groups (-OSO₃) or one or two organic nitrate groups (-ONO₂) 481 482 and these compounds were more likely OSs or nitrooxy-OSs, presenting lower DBE values and 483 higher O/C and H/C ratios (Table S5 and S6) (O'Brien et al., 2014). The high-intensity CHONS 484 compounds observed in this study, such as C₁₀H₁₆NO₇₋₉S, C₁₀H₁₈NO₈₋₉S, C₁₀H₁₈N₂O₁₁S, and 485 $C_9H_{14}NO_{8-9}S$ could be nitrooxy-OSs derived from monoterpenes such as limonene and α -terpinene of which we found the formulas in summer contained more oxygen atoms, indicating the higher 486 oxidation degree of these nitrooxy-OSs in summer (Figure S2 and S3) (Sun et al., 2021; 487 488 Bruggemann et al., 2020; Wang et al., 2020; Wang et al., 2018d).

The CHOS compounds with high intensity abundance, such as typical isoprene epoxydiols (IEPOX) derived OSs with molecular formulas of $C_5H_8O_7S$ and $C_5H_{10}O_7S$ were both detected in the summer and winter samples, of which the relative intensity of $C_5H_8O_7S$ were over 80 % in S1,

S2, S5, and S6 samples, indicating the significant isoprene SOA formation in summer (Kourtchev 492 et al., 2016; Kourtchev et al., 2013). The results were consistent with the PMF results reported by 493 494 Bao et al. (2022). The monoterpenes derived OSs such as C₈H₁₄O₆S, C₈H₁₄O₈S, C₁₀H₁₈O₈, 495 $C_{10}H_{14}O_6$, and $C_{11}H_{16}O_7$ were detected in both summer and winter samples in this study, which could refer to monoterpene-OSs derived from α -pinene, α -terpinene, and limonene (Wang et al., 496 497 2020). Moreover, OSs with high carbon numbers (C \geq 14) such as C₁₄H₂₂O₇S, C₁₄H₂₂O₈S, C14H24O7S, C15H26O7S, C15H24O7S, C15H24O8S, and C16H28O7S were also observed in both 498 499 summer and winter samples. Long-chain alkanes emitted from vehicle emissions might be 500 precursors of these OSs which was consistent with the molecular structures of OSs collected in urban areas affected by traffic emissions such as Shanghai, Los Angeles, and Beijing (Wang et al., 501 2019a; Tao et al., 2014; Wang et al., 2016b). The aromatic OSs such as naphthalene derived OSs 502 503 with molecular formulas of C₁₀H₁₀O₆S, C₁₀H₁₀O₇S, and C₁₀H₁₂O₇S, 2-methylnaphthalene derived OSs with molecular formulas of C₉H₁₂O₆S, C₁₁H₁₂O₇S, and C₁₁H₁₄O₇S, and hydroxybenzene 504 505 derived OSs with molecular formulas of $C_6H_6O_5S$ were also observed in this study (Qi et al., 2021; Riva et al., 2015; Blair et al., 2017). Figure S8 further displays the ternary plot of the relative 506 507 intensities of OSs from biogenic precursors (e.g., isoprene and monoterpenes), long-chain alkanes 508 and aromatic hydrocarbon. As shown in Fig. S8, the biogenic OSs and long-chain alkanes OSs 509 formation were comparable in summer and winter, demonstrating both biogenic and anthropogenic 510 emission contributions to HULIS. The aromatic OSs presented higher relative intensities in winter, 511 further indicating the increasing anthropogenic emissions in winter. The presence of long-chain alkanes derived OSs in both summer and winter seasons provided another evidence that the traffic 512 emission was one of the important fossil sources of HULIS in this study. 513

514 3.5 Comparison with organic compounds in source and atmospheric aerosol samples

515 The O/C and H/C ratios of water soluble HULIS in this study were compared with those of 516 water soluble organic compounds reported in source samples from BB, coal combustions, and vehicle emissions (Tang et al., 2020; Song et al., 2018; Cui et al., 2019; Song et al., 2019), cloud 517 water samples (Bianco et al., 2018; Zhao et al., 2013), rainwater samples (Altieri et al., 2009), fog 518 samples (Brege et al., 2018), as well as aerosol samples collected in Beijing (Jang et al., 2020; Wu 519 520 et al., 2019a; Wang et al., 2018a), Tianjin (Han et al., 2022), Baoding (Sun et al., 2021), Shanghai (Wang et al., 2017b), Guangzhou (Jiang et al., 2021), respectively in China, Mainz (Wang et al., 521 2018b), Cork city (Kourtchev et al., 2014), and Bologna (Brege et al., 2018), respectively in 522

Europe, and Bakersfield (O'Brien et al., 2014) and Virginia (Willoughby et al., 2014), respectively in the United States (Fig. 9). The O/C ratios were obviously higher than those detected in primary BB, coal combustion, and vehicle emission samples. The H/C ratios of the CHO and CHON compounds were comparable with the source samples, indicating the organics in HULIS experienced atmospheric secondary process and the mixed sources of HULIS in this study. The H/C ratios of the S-containing compounds were much higher than those of source samples which could be attributed to the significant organosulfates formation in the atmosphere.

530 The O/C ratios reported in this study were also higher than those reported in aerosol samples in urban area in China, further indicating the serious secondary pollution at Nanjing, China. 531 Among the CHO and CHON compounds, we found that the highest H/C ratio values were observed 532 in the southern city of Guangzhou, followed by those in Nanjing and Shanghai, and the lowest 533 534 values were observed in the northern cities such as Beijing, Tianjin, and Baoding, indicating the higher unsaturation degree of the aerosol samples collected from the northern cities, which were 535 536 also considered as the heavy industrial region in China. The higher H/C ratios of aerosol samples collected in Europe and the United States indicated the less anthropogenic emissions such as 537 538 industrial emissions from those areas.

539 **4. Conclusions**

540 This study focuses on the sources and molecular characteristics differences of water soluble HULIS in summertime and wintertime from 2017 to 2018 at a suburb site of the YRD, China based 541 542 on the radiocarbon analysis and FT-ICR MS measurement with ESI ion source in negative mode. 543 The carbon isotope analysis results highlight the important fossil source contributions to HULIS at the study site. A total of 14387 and 15731 peaks were detected in the summer and winter samples, 544 respectively based on the FT-ICR MS results. The assigned molecular formulas were classified 545 546 into CHO, CHON, CHOS, and CHONS subgroups according to their elemental compositions. The 547 Van Krevelen diagrams showed that more tannins-like and carbohydrates-like species were detected in summer indicating biogenic SOA formation. Whereas more compounds containing 548 condensed aromatic structures were detected in winter which were derived from anthropogenic 549 emissions. The total relative intensity of CHO compounds in summer were significantly higher 550 551 than those in winter, containing lots of macromolecular oligomers derived from biogenic 552 precursors. The high-intensity CHO compounds in winter were mainly aromatic compounds such as phenolic substances and flavonoids which were related to aged BBOA and oxidized PAHs most 553

probably from fossil fuel combustion. On the contrary, the total relative intensity of CHON 554 compounds significantly increased in winter, mainly composed of nitro compounds or 555 organonitrates. The enhanced formation of nitrophenols in winter indicated the BB influence. The 556 increasing organonitrates formation in winter highlighted the secondary N-containing compounds 557 formation via NO₃ radical-initiated oxidation processes. It is worth noting that the top CHON 558 compounds in summer were referring to aromatic reduced N compounds produced from the 559 aldehyde-ammonia reactions. The S-containing compounds were mainly composed of highly 560 oxidized OSs. The monoterpenes derived OSs and long-chain alkanes derived OSs were widely 561 observed in both summer and winter samples, while the aromatic OSs formation were found to be 562 more significant in winter. The presence of long-chain alkanes derived OSs supported the 563 radiocarbon results, indicating that the traffic emission was the important fossil sources at the study 564 565 site. The presence of aromatic secondary N-containing and S-containing compounds provided evidence for the substantial contributions from anthropogenic SOA formation to fossil sources at 566 567 the study site. These results further verified the work reported before by Bao et al. (2022) based on the PMF model which have found the significant anthropogenic SOA and fossil fuel 568 569 combustion contributions to HULIS in urban area in China at molecular level. In addition, strong biogenic emission in summer and BB in winter were found in this study, highlighting the 570 571 importance of different control policies for each season in the future.

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577 **References**

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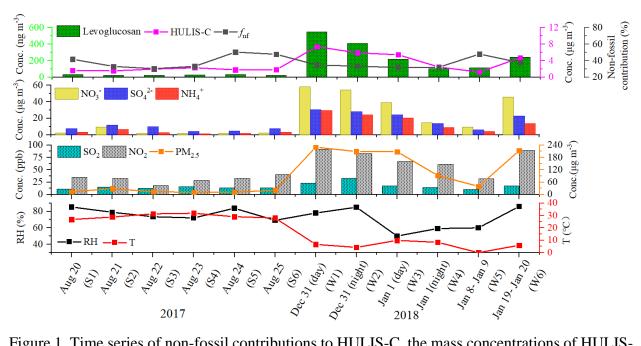
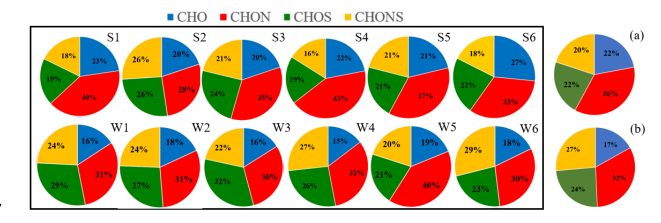


Figure 1. Time series of non-fossil contributions to HULIS-C, the mass concentrations of HULIS-

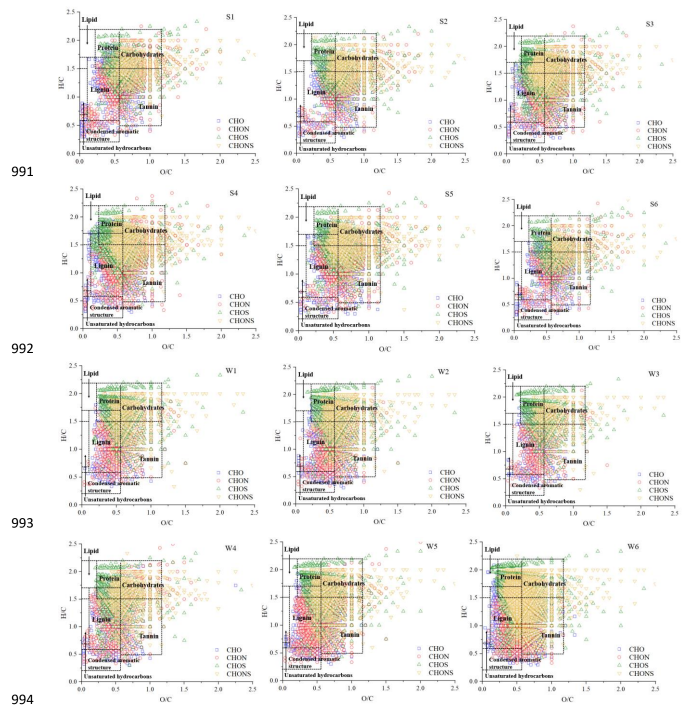
C, Levoglucosan, NO₃⁻, SO₄²⁻, NH₄⁺, SO₂, NO₂, and PM_{2.5}, relative humidity, and temperature during the study periods.



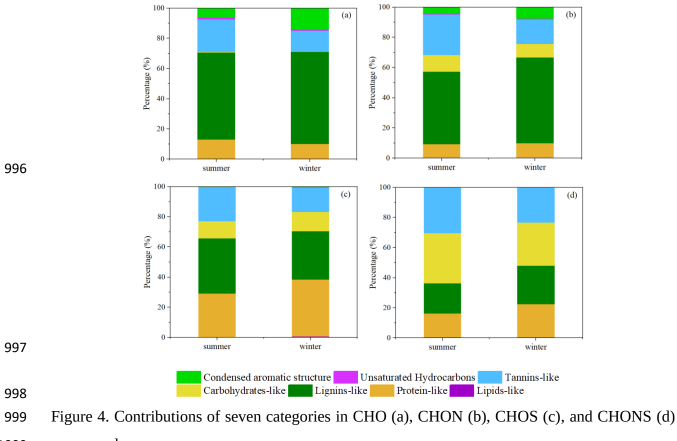
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Figure 2. Pie graph of the number percentages of each elemental formula group for the 12 samples

- 989 plotted in the box and the averaged number percentages of each elemental formula group for the
- 990 summer samples (a) and winter samples (b).



995 Figure 3. Van Krevelen diagrams of the 12 samples.



1000 compounds.

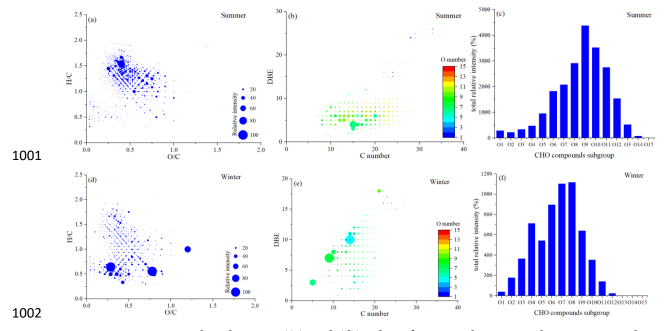


Figure 5. Van Krevelen diagram ((a) and (d)), plot of DBE values vs carbon atom numbers ((b)
and (e)), and the total relative intensity of each subgroup ((c) and (f)) for the CHO compounds in
summer and winter.

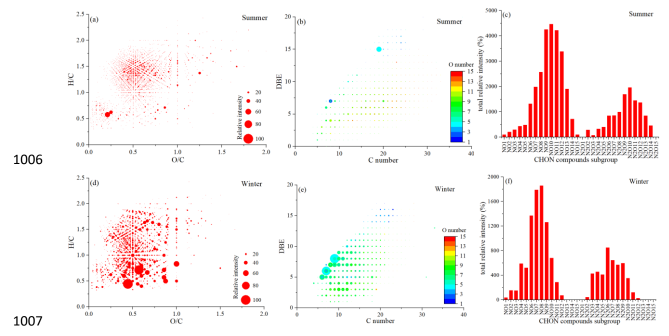


Figure 6. Van Krevelen diagram ((a) and (d)), plot of DBE values vs carbon atom numbers ((b)
and (e)), and the total relative intensity of each subgroup ((c) and (f)) for the CHON compounds
in summer and winter.

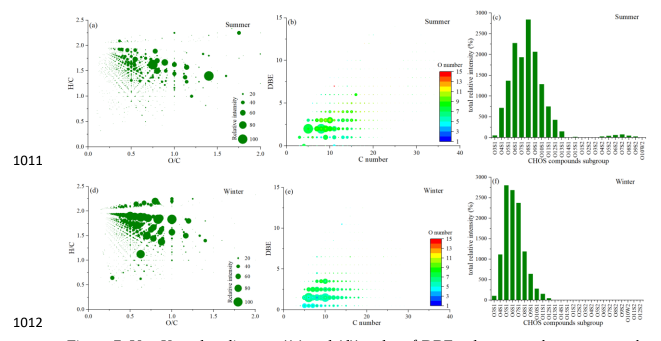


Figure 7. Van Krevelen diagram ((a) and (d)), plot of DBE values vs carbon atom numbers ((b)
and (e)), and the total relative intensity of each subgroup ((c) and (f)) for the CHOS compounds in
summer and winter.

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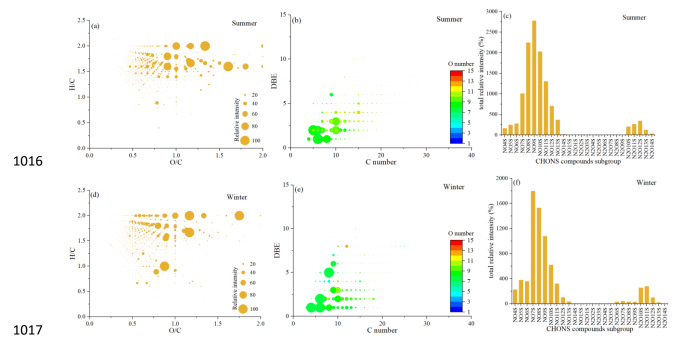


Figure 8. Van Krevelen diagram ((a) and (d)), plot of DBE values vs carbon atom numbers ((b)
and (e)), and the total relative intensity of each subgroup ((c) and (f)) for the CHONS compounds
in summer and winter.

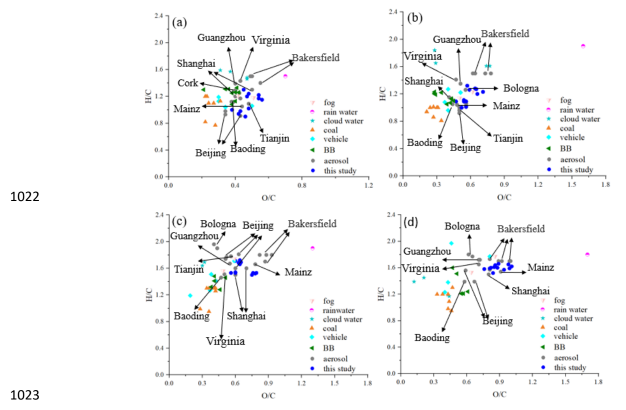


Figure 9. Comparison of O/C and H/C ratios of water soluble organic compounds in different
atmospheric media in CHO (a), CHON (b), CHOS (c), and CHONS (d) compounds.