1 Large contribution of fossil-fuel derived secondary organic

2 carbon to water-soluble organic aerosols in winter haze of China

3 Yan-Lin Zhang^{1,2,3*}, Imad El-Haddad³, Ru-Jin Huang^{3,4*}, Kin-Fai Ho^{4,5}, Jun-Ji Cao^{4*},

4 Yongming Han⁴, Peter Zotter^{3, #}, Carlo Bozzetti³, Kaspar R. Daellenbach³, Jay G. Slowik³, Gary

- 5 Salazar², André S.H. Prévôt^{3*}, Sönke Szidat^{2*}
- 6 ¹Yale-NUIST Center on Atmospheric Environment, Nanjing University of Information Science
- 7 and Technology, 210044 Nanjing, China
- 8 ²Department of Chemistry and Biochemistry & Oeschger Centre for Climate Change Research,
- 9 University of Bern, 3012 Bern, Switzerland
- 10 ³Paul Scherrer Institute (PSI), 5232 Villigen, Switzerland
- ⁴Key Laboratory of Aerosol Chemistry and Physics, Institute of Earth Environment, Chinese
- 12 Academy of Sciences, 710061 Xi'an, China
- ⁵School of Public Health and Primary Care, The Chinese University of Hong Kong, Hong Kong,
 China
- 15 *To whom correspondence should be addressed. E-mail: <u>dryanlinzhang@outlook.com</u> or
- 16 <u>zhangyanlin@nuist.edu.cn</u>(Y.-L.Z.); <u>andre.prevot@psi.ch</u>(A. Prévôt); <u>rujin.huang@ieecas.cn</u>
- 17 (R.-J.H.); jjcao@ieecas.cn (J.J.C.); szidat@dcb.unibe.ch (S.S.).
- 18 Phone: +86 25 5873 1022; fax: +86 25 5873 1193

19 Abstract

20 Water-soluble organic carbon (WSOC) is a large fraction of organic aerosols (OA) globally and 21 has significant impacts on climate and human health. The sources of WSOC remain very 22 uncertain in polluted regions. Here we present a quantitative source apportionment of WSOC isolated from aerosols in China using radiocarbon (¹⁴C) and offline high-resolution time-of-23 24 flight aerosol mass spectrometer measurements. Fossil emissions on average accounted for 32-47% of WSOC. Secondary organic carbon (SOC) dominated both the non-fossil and fossil 25 derived WSOC, highlighting the importance of secondary formation to WSOC in severe winter 26 27 haze episodes. Contributions from fossil emissions to SOC were $61\pm4\%$ and $50\pm9\%$ in 28 Shanghai and Beijing, respectively, significantly larger than those in Guangzhou (36±9%) and 29 Xi'an (26±9%). The most important primary sources were biomass burning emissions, contributing 17-26% of WSOC. The remaining primary sources such as coal combustion, 30 31 cooking and traffic were generally very small but not negligible contributors, as coal 32 combustion contribution could exceed 10%. Taken together with earlier ¹⁴C source 33 apportionment studies in urban, rural, semi-urban, and background regions in Asia, Europe and 34 USA, we demonstrated a dominant contribution of non-fossil emissions (i.e., $75\pm11\%$) to 35 WSOC aerosols in the North Hemisphere; however, the fossil fraction is substantially larger in 36 aerosols from East Asia and the East Asian pollution outflow especially during winter due to 37 increasing coal combustion. Inclusion of our findings can improve a modelling of effects of 38 WSOC aerosols on climate, atmospheric chemistry and public health.

39 1 INTRODUCTION

Water-soluble organic carbon (WSOC) is a large fraction of atmospheric organic 40 aerosols (OA), which contributes approximately 10% to 80% of the total mass of organic carbon 41 42 (OC) in aerosols from urban, rural and remote sites (Zappoli et al., 1999;Weber et al., 43 2007;Ruellan and Cachier, 2001;Wozniak et al., 2012;Mayol-Bracero et al., 2002). Only 10 to 44 20% of total mass of WSOC has been resolved at a molecular level, and it consists of a large 45 variety of chemical species such as mono- and di-carboxylic acids, carbohydrate derivatives, 46 alcohols, aliphatic and aromatic acids and amino acids (Fu et al., 2015; Noziere et al., 2015). 47 Recent studies suggest that the water-soluble fraction of HUmic LIke Substances (HULIS) is a major component of WSOC, which exhibits light-absorbing properties (Limbeck et al., 48 49 2005; Andreae and Gelencser, 2006; Laskin et al., 2015). Therefore, WSOC has significant 50 influences on the Earth's climate either directly by scattering and absorbing radiation or 51 indirectly by altering the hygroscopic properties of aerosols and increasing cloud condensation 52 nuclei (CCN) activity (Asa-Awuku et al., 2011;Cheng et al., 2011;Hecobian et al., 2010).

53 WSOC can be directly emitted as primary particles mainly from biomass burning 54 emissions or produced from secondary organic aerosol (SOA) formation (Sannigrahi et al., 2006;Kondo et al., 2007;Weber et al., 2007;Bozzetti et al., 2017b;Bozzetti et al., 2017a). 55 56 Ambient studies provide evidence that SOA formation through the oxidation of volatile organic compounds (VOCs) and gas-to-particle conversion processes may be a prevalent source of 57 58 WSOC (Kondo et al., 2007; Weber et al., 2007; Miyazaki et al., 2006; Hecobian et al., 2010). WSOC is therefore thought to be a good proxy of secondary organic carbon (SOC) in the 59 absence of biomass burning (Weber et al., 2007). By contrast, water-insoluble OC (WIOC) is 60 thought to be mainly from primary origins with a substantial contribution from fossil fuel 61 62 emissions (Miyazaki et al., 2006;Zhang et al., 2014b).

Due to a large variety of sources and unresolved formation processes of WSOC, their
 relative fossil and non-fossil contributions are still poorly constrained. Radiocarbon (¹⁴C)
 analysis of sub-fractions of organic aerosols such as OC, WIOC and WSOC enable an

66 unambiguous, precise and quantitative determination of their fossil and non-fossil sources 67 (Zhang et al., 2012;Zhang et al., 2014b;Zhang et al., 2014c;Zong et al., 2016;Cao et al., 2017). 68 Meanwhile, the application of aerosol mass spectrometer measurement and positive matrix 69 factorization and multi-linear engine 2 (ME-2) can quantitatively classify organic aerosols into 70 two major types such as hydrocarbon-like OA (HOA) from primary fossil-fuel combustion and 71 oxygenated organic aerosol (OOA) from secondary origin (Zhang et al., 2007; Jimenez et al., 72 2009). Field campaigns with the aerosol mass spectrometer (AMS) have revealed a 73 predominance of OOA in various atmospheric environments, although their sources remain 74 poorly characterized (Zhang et al., 2007; Jimenez et al., 2009). Previous studies found OOA is 75 strongly correlated with WSOC from urban aerosols in Tokyo, Japan, the Pearl River Delta 76 (PRD) in South China and Helsinki, Finland, indicating similar chemical characteristics, sources and formation processes of OOA and WSOC (Kondo et al., 2007;Xiao et al., 77 78 2011; Timonen et al., 2013). Similarly, HOA is mostly water insoluble and the major portion of water insoluble OC (WIOC) can be assigned as HOA (Kondo et al., 2007;Daellenbach et al., 79 80 2016). Therefore, ¹⁴C measurement of WIOC and WSOC aerosols may provide new insights into sources and formation processes of primary and secondary OA, respectively, which also 81 82 will elucidate the origin of HOA and OOA as measured by AMS (Zotter et al., 2014b;Zhang et 83 al., 2017).

In this paper we apply a newly developed method to measure ${}^{14}C$ in WSOC of PM_{2.5} 84 85 (particulate matter with an aerodynamic diameter of small than 2.5 µm) samples collected at 86 four Chinese megacities during an extremely severe haze episode during winter 2013 (Zhang et al., 2015b;Huang et al., 2014). In conjunction with our previous dataset from the same 87 88 campaign, we quantify fossil and non-fossil emissions from primary and secondary sources of 89 WSOC and WIOC. The dataset is also complemented by previous ¹⁴C-based source apportionment studies conducted in urban, rural and remote regions in the North Hemisphere 90 91 to gain an overall picture of the sources of WSOC aerosols.

92 2 MATERIALS AND METHODS

93 2.1 Sampling

94 During January 2013 extremely high concentrations of 24-h PM_{2.5} (i.e. often >100 95 µg/m³) were identified in several large cities in East China (Huang et al., 2014;Zhang et al., 96 2015b). To investigate sources and formation mechanisms of the haze particles, an intensive 97 field campaign was carried out in four large cities, Beijing, Xi'an, Shanghai and Guangzhou, 98 which are representative cities of the Beijing-Tianjin-Hebei region, central-northwest region, 99 Yangtze Delta Region, and Pearl River Delta Region, respectively. The sampling procedures 100 have been previously described in detail elsewhere (Zhang et al., 2015b). Briefly, PM_{2.5} samples 101 were collected on pre-baked (450 °C for 6 hours) quartz filters using high-volume samplers for 102 24 h at a flow rate of ~1.05 m³/min from 5 to 25 January 2013. The sampling sites in each city 103 were located at campuses of universities or at research institutes, at least 100 m away from 104 major emission sources (e.g., roadways, industry and domestic sources). One field blank sample 105 for each site was collected and analyzed. The results reported here were corrected for these field 106 blanks (Zotter et al., 2014a;Cao et al., 2013). All samples were stored at -20 °C before analysis. 107 The $PM_{2.5}$ mass was gravimetrically measured with an analytical microbalance before and after 108 sampling with the same conditions (~12 hour)

109 2.2 OC and EC mass determinations

110 A 1.0 cm² filter punches were used for OC and EC mass determination with a OC/EC 111 analyzer (Model4L) using the EUSAAR_2 protocol (Cavalli et al., 2010). The replicate analysis 112 (n=6) showed an analytical precision with relative standard deviations smaller than 5%, 10%, 113 and 5% for OC, EC and TC, respectively. The field blank of OC was on average 2.0 ± 1.0 114 μ g/cm² (equivalent to ~0.5 μ g/m³), which was used for blank correction for OC. EC data was 115 not corrected for field blank, because such a blank was not detectable.

116 2.3 Offline-AMS measurement and PMF source apportionment

117 The water-soluble extracts from the same samples were analyzed by a high-resolution time of118 flight aerosol mass spectrometer (HR-ToF-AMS) and the resulting mass spectra were used as

an inputs for positive matrix factorization (PMF) for the source apportionment of the WSOC, OC and $PM_{2.5}$. The methodology applied, and the AMS-PMF results obtained are detailed in Huang et al. (2014) and will only be briefly described in the following. Here, only data relative to WSOC are used.

Filter punches (the equivalent of ~4 cm²) were sonicated in 10 mL ultrapure water (18.2 M Ω cm at 25 °C, TOC <3ppb) for 20 min at 30°C. The water extracts were aerosolized and the resulting particles were dried with a silica gel diffusion dryer before analysis by the HR-ToF-AMS. For each measurement ten mass spectra were recorded (AMS V-mode, m/z 12-500), with a collection time for each spectrum of 1 minute.

128 Online AMS measurements provide quantitative mass spectra of submicron non-refractory

129 aerosol species, including organic aerosol and ammonium nitrate and sulfate. However, the

130 offline AMS measurements described herein cannot be directly related to ambient

131 concentrations due to uncertainties in nebulization and AMS lens cut-off. Here, we have

scaled the organic aerosol mass spectra to water soluble organic aerosol concentrations

133 (WSOM), obtained as WSOC times OM/OC ratios. The latter were determined by the high

resolution analysis of the organic aerosol mass spectra, acquired by the AMS.

The quantitative WSOM mass spectra are used together with other aerosol species (listed
below), collectively referred to as 'species' hereafter, as PMF inputs. PMF solves the bilinear
matrix equation:

138
$$X_{ij} = \sum_{k} G_{i,k} F_{k,j} + E_{i,j}$$
 (Eq. 1)

by following a weighted least squares approach. In the equation, *i* represent the time index, *j* a species and *k* the factor number. X_{ij} is the input matrix, $G_{i,k}$ is the matrix of the factor timeseries, $F_{k,j}$ is the matrix of the factor profiles and $E_{i,j}$ the model residual matrix. PMF determines $G_{i,k}$ and $F_{k,j}$ such that the ratio of the Frobenius norm of $E_{i,j}$ over the uncertainty matrix, $s_{i,j}$, used as model input is minimised. 144 The species considered as inputs include the quantitative WSOM mass spectra, organic markers 145 (3 anhydrous sugars, 4 lignin breakdown products, 2 resin acids, 4 hopanes, 19 polycyclic 146 aromatic hydrocarbons and their oxygenated derivatives), EC, and major ions (Cl⁻, NO₃⁻, SO₄²⁻, oxalate, methylsulfonic acid, Na⁺, K⁺, Mg²⁺, Ca²⁺, and NH₄⁺) and residual PM. The latter is the 147 148 difference between total PM2.5 mass and the measured species. It represents our best estimate 149 of the particulate chemical species not measured here, most likely dominated by crustal material. 150 The Source Finder toolkit (SoFi v.4.9) (Canonaco et al., 2013) for IGOR Pro software package 151 (Wavemetrics, Inc., Portland, OR, USA) was used to run the PMF algorithm. The PMF was solved by the Multilinear Engine 2 (ME-2, Paatero, 1999), which allows the constraining of the 152 $F_{k,i}$ elements to vary within a certain range defined by the scalar α ($0 \le \alpha \le 1$), such that the 153 modelled $\mathbf{F}_{k,i}$ equals: 154

155
$$\mathbf{F}'_{k,i} = \mathbf{F}_{k,i} \pm \alpha * \mathbf{F}_{k,i}$$
(Eq. 2)

156 The elements that were constrained in $F_{k,i}$ matrix can be found in Huang et al. (2014). The 157 factors extracted by ME-2 were interpreted to be related to primary emissions from traffic (TR), biomass burning (BB), coal burning (CC), cooking emissions (CI) and dust and from two 158 159 secondary aerosol fractions. The contributions of the water soluble organic aerosol related to 160 these different factors were determined by the multiplying their relative abundance in the factor 161 profiles by the respective factor time-series. The factors WSOM time series were then divided by the respective OM/OC_k calculated from the high-resolution analysis of the factor mass 162 spectral profile to obtain the $WSOC_k$ time series related to each of the factors. The average 163 OM/OCk are: 1.25, 1.39, 1.49, 1.55, 2.25, and 2.4 for TR, CI, BB, CB, SOA, and dust, 164 165 respectively. In the following analysis, the mass of $WSOC_k$ related to coal burning and traffic 166 were assigned to fossil WSOC fraction, while the mass of WSOC_k related to biomass burning 167 and cooking emissions were assigned to non-fossil WSOC fraction (see Sec. 2.5). Meanwhile, 168 the remaining WSOC fractions are assigned to the secondary factors, which can be from both fossil and non-fossil origins, were considered collectively and compared to the unassignedfossil and non-fossil WSOC, to retrieve the origins of this remaining fraction (see Sec. 2.5).

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2.4¹⁴C measurement of WSOC

¹⁴C content of micro-scale WSOC aerosol samples was measured with a newly 173 174 developed method (Zhang et al., 2014c). Briefly, a 16-mm-diameter punch of each filter was 175 extracted using 10 ml ultrapure water with low TOC impurity (less than 5 ppb). The water 176 extracts were recovered in the 20 ml PFA vials and were then pre-frozen at -20 °C more than 5 177 hours before completely dryness in a freeze dryer (Alpha 2-4 LSC, Christ, Germany) for about 178 24 h to 36 h. The residue was re-dissolved in 50 μ l of ultrapure water three times and transferred 179 into 200 μ l tin capsules (Elementar, Germany). The concentrated samples were heated in the oven at 55-60 °C until complete dryness before the ¹⁴C measurements. 180

181 WSOC extracts in tin capsules were then converted to CO₂ by the oxidation of the 182 carbon-containing samples using an Elemental Analyzer (EA, Model Vario Micro, Elementar, Germany) as a combustion unit (up to 1050 °C). The resulting CO₂ was introduced continuously 183 by a versatile gas inlet system into a gas ion source of the accelerator mass spectrometer 184 MICADAS where ¹⁴C of CO₂ was finally measured (Wacker et al., 2013;Salazar et al., 2015). 185 The ¹⁴C content of OC and EC was measured in our previous study (Zhang et al., 2015b). ¹⁴C 186 187 results were expressed as fraction of modern ($f_{\rm M}$), i.e., the fraction of the measured ${}^{14}{\rm C}/{}^{12}{\rm C}$ ratio related to the ¹⁴C/¹²C ratio of the reference year 1950 (Stuiver, 1977). To correct excess ¹⁴C 188 from nuclear bomb tests in the 1950s and 1960s, $f_{\rm M}$ values were converted to the fraction of 189 190 non-fossil ($f_{\rm NF}$) (Zotter et al., 2014a;Zhang et al., 2012):

191 $f_{\rm NF} = f_{\rm M}/f_{\rm M,ref}$ (Eq. 3)

192 $f_{M,ref}$ is a reference value of f_M for non-fossil carbon sources including biogenic and 193 biomass burning emissions, which were estimated as 1.08 ± 0.05 (i.e., $f_{M,ref}=(0.5*1.10+0.5*1.05)$ 194 (see details in (Zhang et al., 2012)) for WSOC samples collected in 2013 according to the 195 contemporary atmospheric $CO_2 f_M$ (Levin et al., 2010) and a tree growth model (Mohn et al., 196 2008).

197 2.5 AMS²-based source apportionment of WSOC

To better understand the origin of WSOC observed at these sites, WSOC sources were apportioned into several major sources by a combination of ¹⁴C and PMF source apportionments (See Figure 1). Here, two "AMS" (i.e., accelerator mass spectrometer and aerosol mass spectrometer), such a combined approach was named as "AMS²-based source apportionment.

203 WSOC concentration from non-fossil (WSOC_{NF}) and fossil (WSOC_F) sources were 204 calculated from:

205
$$WSOC_{NF}=WSOC*f_{NF}(WSOC)$$
 (Eq. 4)

206
$$WSOC_F = WSOC - WSOC_{NF}(Eq. 5)$$

The mass concentration of WSOC was derived from the subtraction of TC mass measured from a water-extracted filter from that measured with an un-treated filter (Zhang et al., 2012):

Based on mass balance, WIOC concentrations from non-fossil (WIOC_{NF}) and fossil
(WIOC_F) sources were calculated from:

- 213 $WIOC_{NF}=OC_{NF}-WSOC_{NF}$ (Eq. 7)
- 214 $WIOC_F=OC_F-WSOC_F$ (Eq.8)

where OC concentrations from non-fossil (OC_{NF}) and fossil (OC_{F}) sources were obtained by mass and ¹⁴C measurement of the OC fraction, which were reported previously (Zhang et al., 2015b).

The non-fossil and fossil-fuel derived WSOC can be apportioned into primary and secondary OC:

220
$$WSOC_{NF}=WSOC_{POC,NF}+WSOC_{SOC,NF}(Eq.9)$$

221
$$WSOC_F = WSOC_{POC,F} + WSOC_{SOC,F} (Eq. 10)$$

WSOC_{POC,NF} can be sub-divided into the following three major primary emissions including
cooking emission (WSOC_{CI}) and biomass burning (WSOC_{BB}).

224
$$WSOC_{POC,NF}=WSOC_{CI}+WSOC_{BB}(Eq.11)$$

Similarly, WSOC_{POC,F} can be sub-divided into the following two major primary emissions
including traffic (WSOC_{TR}) and coal combustion (WSOC_{CB}).

227
$$WSOC_{POC,F}=WSOC_{TR}+WSOC_{CB}(Eq.12)$$

where primary fractions such as $WSOC_{CI}$, $WSOC_{BB}$, $WSOC_{TR}$ and $WSOC_{CB}$ are previously estimated by the off-line AMS-PMF approach (Huang et al., 2014;Daellenbach et al., 2016;Bozzetti et al., 2017a;Bozzetti et al., 2017b).

An uncertainty propagation scheme using a Latin-hypercube sampling (LHS) model was implemented to properly estimate overall uncertainties including measurement uncertainties of the mass determinations of carbon species (i.e., OC, EC, TC, WSOC, WIOC) and ¹⁴C measurement, blank corrections from field blanks, and estimation of $f_{M,ref}$ (Zhang et al., 2015b).

236 **3 RESULTS AND DISCUSSION**

237 **3.1 Overall results**

238 During the haze periods of January 2013, the highest daily average PM_{2.5} concentrations were 239 found in Xi'an (345 μ g/m³) followed by Beijing (158 μ g/m³), Shanghai (90 μ g/m³) and 240 Guangzhou ($68 \mu g/m^3$). These levels were much higher than the China's National ambient Air 241 quality standards (i.e., $35 \,\mu g/m^3$). Indeed, several studies have already reported the chemical 242 composition, source and formation mechanism of PM_{2.5} in many large cities during the haze 243 events of January 2013 in East China. For examples, Huang et al. (2014) revealed that the 244 secondary aerosol formation contributed to 44-71% of OA in Beijing, Xi'an, Shanghai, and 245 Guangzhou during this extremely haze event in China (Huang et al., 2014). By ¹⁴C-based source 246 appointment conducted in the same campaign, Zhang et al. (2015) have reported that 247 carbonaceous aerosol pollution was driven to a large (often dominant) extent by SOA formation 248 from both, fossil and biomass-burning sources (Zhang et al., 2015b). For all four cities, the 24 h average levels of WSOC were significantly correlated with the levels of PM_{2.5} and OC (R=0.99, 249 250 p < 0.01, Figure 2), suggesting that WSOC and OA may have similar sources and formation processes and thus have important implications for OC loadings and associated environmental 251 252 and health effects. However, the sources of WSOC remain poorly constrained. In this study, we measured the ¹⁴C content of WSOC aerosols in six samples (three with the highest, three 253 254 with average PM mass) for each city to report on heavily and moderately polluted days (HPD and MPD, respectively) (Zhang et al., 2015b). The ¹⁴C contents of OC and EC of the same 255 256 samples were reported previously (Zhang et al., 2015b).

257 WSOC on average accounted for 53±8.0% (ranging from 40-65%) of OC including all samples from the four sites, which was consistent with previous estimates . Based on these 258 259 measurements, the concentrations of WSOC from non-fossil sources (WSOC_{NF}) spanned from 1.41 to 45.3 μ g/m³ with a mean of 10.6±12.1 μ g/m³, whereas the corresponding range for 260 261 WSOC from fossil-fuel emissions (WSOC_F) was 0.44 to 20.1 μ g/m³ with a mean of 5.3±4.9 μ g/m³ (Figure 3). Similar to PM_{2.5} levels, the highest concentrations of WSOC_{NF} and WSOC_F 262 263 were observed in Northern China in Xi'an and Beijing (Xi'an>Beijing), followed by the two 264 southern sites Shanghai and Guangzhou. Non-fossil contributions (mean \pm standard deviation) 265 to total WSOC were 53±5%, 75±4%, 48±2% and 68±6% in Beijing, Xi'an, Shanghai, and 266 Guangzhou, respectively. Thus, fossil contributions were notably higher in Beijing and 267 Shanghai than those in Xi'an and Guangzhou. Such a trend was also observed for OC (Zhang 268 et al., 2015b), suggesting relatively high contribution from fossil-fuel emissions to OC and 269 WSOC due to large coal usage. Despite of these fossil emissions, non-fossil sources were 270 considerably important or even dominant contributors for all the studied sites, which may be 271 associated with primary and secondary OA from regional-transported and local biomass 272 burning emissions. As shown in Figure 4, non-fossil WSOC was significantly correlated with 273 levoglucosan, indicating that a large fraction of non-fossil WSOC was indeed from biomass 274 burning emissions. In addition, no significant or only a negative correlation (Figure 4) was 275 found between levoglucosan and fraction of fossil to WSOC, suggesting that fossil-fuel source is very unlikely a major or important contributor of levoglucosan even in the regions (e.g., 276 277 Xi'an and Beijing in this study) where coal combustion is important during the cold period 278 (Zhang et al., 2015a). It should also be noted that formation of SOA derived from biogenic VOCs may also have contributed to WSOC_{NF} in Guangzhou, where temperatures during the 279 sampling period were significantly higher (i.e., 5–18 °C) than those in other cities (i.e., -12 to 280 281 +9 °C)(Bozzetti et al., 2017b). Although both fossil and non-fossil WSOC concentrations were 282 dramatically enhanced during HPD compared to those during MPD, their relative contributions 283 did not change significantly in Beijing and Shanghai whereas a small increasing and decreasing 284 trend in non-fossil fraction was found in Xi'an and Guangzhou, respectively (Figure 3). This 285 suggests that the source pattern of WSOC in Beijing and Shanghai remained similar between 286 HPD and MPD, but the increase in the WSOC concentrations was rather enhanced by additional 287 fossil-fuel and biogenic/biomass burning emissions in Guangzhou and Xi'an, respectively. It 288 should be noted that the meteorological conditions play significant roles on the haze formation 289 in the eastern China during winter 2013, and has already been well documented (Zhang et al., 2014a). However, the details sources of WSOC and WIOC were still unclear. 290

291 **3.2 WSOC versus WIOC**

To compare sources of WSOC and WIOC aerosols, the mass concentrations and ¹⁴C contents 292 of WIOC were also derived based on mass balance. The ¹⁴C-based source apportionment of 293 WIOC and the relationship between f_{NF} (WSOC) and f_{NF} (WIOC) is presented in Figures 5 and 294 295 6a, respectively. It shows that non-fossil contributions to WSOC were larger than those of 296 WIOC for nearly all samples in Beijing, Xi'an and Guangzhou. On average, the majority (60-297 70%) of the fossil OC was water insoluble at these 3 sites (see Figure 6b), indicating that fossil-298 derived OA mostly consisted of hydrophobic components and thus is less water soluble than 299 OA from non-fossil sources. This result is consistent with findings reported elsewhere such as 300 at an urban or rural site in Switzerland (Zhang et al., 2013), a remote site in Hainan Island, 301 South China (Zhang et al., 2014b) and at two rural sites on the east coast of the United States 302 (Wozniak et al., 2012). Meanwhile, the fossil OC in Shanghai, the dominant fraction of OC, 303 was more water soluble (Figure 6b), suggesting an enhanced SOA formation from fossil VOCs 304 from vehicle emissions and/or coal burning for this city. As shown in Figure 6b, non-fossil OA was enriched in water-soluble fractions (i.e., 60%±8%) for all cities, associated with the 305 306 hydrophilic properties of biogenic-derived SOA and biomass-burning derived primary organic 307 aerosol (POA) and SOA, which are composed of a large fraction of polar and highly oxygenated 308 compounds (Mayol-Bracero et al., 2002;Sullivan et al., 2011;Noziere et al., 2015). Thus, non-309 fossil OC has more water-soluble components than fossil ones. It should be noted that relative contributions of WSOC_{NF} and WSOC_F are similar in Beijing and Shanghai, whereas WSOC_{NF} 310 311 is much higher than WSOC_F in Xi'an and Guangzhou. This suggests larger contribution of non-312 fossil sources to WSOC aerosols in Xi'an and Guangzhou than those in Beijing and Shanghai.

313 **3.3 High contribution of secondary formation to WSOC**

WSOC was further apportioned into fossil sources such as coal burning (CB), traffic (TR) and SOC (SOC,F) as well as non-fossil sources such as biomass burning (BB), cooking (CI) and SOC (SOC,NF) using a AMS² based source apportionment (see Sec. 2.5 and Figure 1). SOC dominated WSOC during both the HPD and MPD with a mean contribution of $67\% \pm 9\%$, highlighting the importance of SOC formation to the WSOC aerosols in wintertime pollution 319 events. This is consistent with our previous findings for total PM2.5 mass and bulk carbonaceous 320 aerosols (i.e., total carbon, sum of OC and EC) (Huang et al., 2014; Zhang et al., 2015b). The 321 increase in SOC contribution to WSOC during HPD compared to MPD can be largely due to 322 fossil contribution in Beijing but non-fossil emissions in Xi'an. In Shanghai and Guangzhou, 323 the source pattern of WSOC was not significantly different between MPD and HPD. Fossil 324 contributions to WSOC_{SOC} were 50%±9% in Beijing, 61±4% in Shanghai, associated with SOA 325 from local and transported fossil-fuel derived precursors at these sites (Guo et al., 2014). This 326 contribution drops to 36±9% and 26±9% in Guangzhou and Xi'an, respectively, due to higher 327 biomass-burning contribution to SOC. Despite of the general importance of fossil SOC, 328 formation of non-fossil WSOC_{SOC} becomes especially relevant during HPD especially in Xi'an 329 (Figure 7), which may be explained by competing effects in SOC formation from fossil versus non-fossil precursors. It can be hypothesized for extremely polluted episodes that more 330 331 hydrophilic volatile compounds that were emitted from biomass burning precursors 332 preferentially form SOC compounds via heterogeneous reaction/processing on dust particles compared to highly hydrophobic precursors from fossil sources, a point subjected to future 333 334 laboratory and field experiments. The most important primary sources of WSOC were biomass 335 burning emissions, and their contributions were higher in Xi'an (26%±7%) and Guangzhou 336 $(25\%\pm6\%)$ than those found in Beijing $(17\%\pm6\%)$ and Shanghai $(17\%\pm5\%)$. The remaining 337 primary sources such as coal combustion, cooking and traffic were generally very small 338 contributors of WSOC due to lower water solubility, although coal combustion could exceed 339 10% in Beijing. It should be noted that WSOC was dominated by SOC formation with mean 340 contribution of $61\%\pm10\%$ and $72\%\pm12\%$ (average for all four cities) to non-fossil and fossil-341 fuel derived WSOC, respectively.

342 Summary and implications

Our study demonstrates that non-fossil emissions are generally a dominant contributor of WSOC aerosols during extreme haze events in representative major cities of China, which is in agreement with WSOC source information identified in aerosols with different size fractions 346 (e.g., TSP, PM_{10} and $PM_{2.5}$) observed in the Northern Hemisphere at urban, rural, semi-urban, and background sites in East/South Asia, Europe and USA (Table 1). The ¹⁴C-based source 347 348 apportionment database shows a mean non-fossil fraction of 73±11% across all sites. This 349 overwhelming non-fossil contribution to WSOC is consistently observed throughout the year, 350 which is associated with seasonal-dependent biomass-burning emissions and/or biogenic-351 derived SOC formation. Our study provides evidence that the presence of oxidized OA, which 352 is to a large extent water soluble, in the Northern Hemisphere (Zhang et al., 2007) is mainly 353 derived from biogenic-derived SOA and/or biomass burning sources. The overall importance 354 of non-fossil emissions to the WSOC aerosols results from large contributions of SOC formation from biogenic precursors (e.g., most likely during summer) and relatively high water-355 356 solubility of primary biomass burning particles (e.g., most likely during winter) compared to those emitted from fossil fuel emissions such as coal combustion and vehicle exhaust. Despite 357 358 of the importance of non-fossil sources, a significant fossil fraction is also observed in the WSOC aerosols from polluted regions in East Asia and sites influenced by East Asian 359 continental outflow (Table 1, Figure 8). This fossil contribution is apparently higher than in this 360 361 region than in the USA and Europe, which is due to large industrial and residential coal usage 362 as well as vehicle emissions. From our observation, the increases in the fossil fractions of 363 WSOC were mostly from SOC formation. Since WSOC has hygroscopic properties, our 364 findings suggest that SOC formation from non-fossil emissions have significant implications 365 on aerosol-induced climate effects. In addition, fossil-derived SOC formation may also become 366 important in polluted regions with large amounts of fossil fuel emissions such as in China and 367 other emerging countries. Low combustion efficiencies and consequently high emission factors 368 in most of the combustion processes in China may further be responsible for increased 369 concentrations of fossil precursors which may be oxidized to form water-soluble SOA in the 370 atmosphere and contribute substantially to the WSOC aerosols. The enhanced WSOC levels 371 may be also originate from aging of fossil POA during the long-range transport of aerosols 372 (Kirillova et al., 2014a). It is also interesting to note that fossil contribution during winter in 373 East Asia is generally higher than those in the rest of the year although relatively large fossil

fraction could be occasionally found as well. Such seasonal dependence was not observed in other regions, suggesting the importance of fossil contribution to WSOC due to increasing coal combustions during winter in China. This study provides a more detailed source apportionment of WSOC, which could improve modelling of climate and health effects as well as the understanding of atmospheric chemistry of WSOC in the polluted atmosphere such as China and provide scientific basis for policy decisions on air pollution emissions mitigation.

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636 Figures and Tables

Table 1. Compilation of literature values of relative fossil-fuel contributions (fossil %) to the

638	WSOC aerosols in East/South Asia, USA and Europe.	
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Site	Location	Season	Size	WSOC (µg/m ³)	WSOC/OC	Fossil %	References
East Asia							
Urban	Beijing, China	Winter/2013	PM _{2.5}	19.8	0.49	47	this work
Urban	Xi'an, China	Winter/2013	PM _{2.5}	31.3	0.53	25	this work
Urban	Shanghai, China	Winter/2013	PM _{2.5}	6.5	0.58	52	this work
Urban	Guangzhou, China	Winter/2013	PM _{2.5}	6.6	0.53	32	this work
Urban	Beijing, China	Winter/2014	PM _{2.5}	14.7	0.40	56	(Fang et al., 2017)
Urban	Beijing, China	Winter/2011	PM _{4.3}	15	0.50	55	(Zhang et al., 2014c)
Urban	Beijing, China	Winter/2013	PM _{2.5}	9.3	0.31	54	(Yan et al., 2017)
Urban	Guangzhou, China	Winter/2012/ 2013	PM _{2.5}	4.1	0.38	33	(Liu et al., 2014)
Urban	Guangzhou, China	Winter/2011	PM_{10}	4.5	0.43	28.5	(Zhang et al., 2014c)
Urban	Xi'an, China	Autumn/2009	PM _{2.5}	5.1	0.28	31	(Pavuluri et al., 2013)
Urban	Xi'an, China	Autumn/2010	TSP	8.1	0.28	29	(Pavuluri et al., 2013)
Urban	Wuhan, China	Winter/2013	PM _{2.5}	13.7	0.45	37	(Liu et al., 2016)
Urban	Sapporo, Japan	Summer/Autum n/2010	PM ₃	1	0.43	15	(Pavuluri et al., 2013)
Urban	Sapporo, Japan	Summer/2011	TSP	1.1	0.24	12	(Pavuluri et al., 2013)
Urban	Sapporo, Japan	Spring/2010	TSP	1.1	0.31	11	(Pavuluri et al., 2013)
Urban	Sapporo, Japan	Autumn/2011	TSP	1.8	0.48	18.3	(Pavuluri et al., 2013)
Urban	Sapporo, Japan	Winter/2010	TSP	0.9	0.45	40.2	(Pavuluri et al., 2013)
Background	Jeju Island, Korea	Winter/2014	PM _{2.5}	2.2	0.66	50	(Fang et al., 2017)
Background	Jeju Island, Korea	Spring/2011	PM _{2.5}	2.0		37.5	(Kirillova et al., 2014a)
Background	Jeju Island, Korea	Spring/2011	TSP	3.0		25	(Kirillova et al., 2014a)
Average						33±14	2014a)
South Asia							
Background	Hainan, China	Annual 2005/2006	PM _{2.5}	3.9	0.54	18	(Zhang et al., 2014b)
Background	Hainan, China	Winter 2005/2006	PM _{2.5}	6.2	0.57	14.5	(Zhang et al., 2014b)
Background	Hainan, China	Summer 2005/2006	PM _{2.5}	1.4	0.40	17.7	(Zhang et al., 2014b)
Background	Hanimaadhoo, Maldives	Annual 2008/2009	TSP	0.5		17	(Kirillova et al., 2013)
Background	Sinhagad, India	Annual 2008/2009	TSP	3.0		24	(Kirillova et al., 2013)
Background	Hanimaadhoo, Maldives	Spring/2012	PM _{2.5}	0.6	0.62	14	(Bosch et al., 2014)
Urban	Delhi, India	Winter/2010/ 2011	PM _{2.5}	22.0		21	(Kirillova et al., 2014b)
Average						18±4	

Europe and US	Europe and USA							
Urban	Göteborg, Sweden	Winter/2005	PM _{2.5}	1.1	0.48	23	(Szidat et al., 2009)	
Urban	Göteborg, Sweden	Summer/2006	PM _{2.5}	0.8	0.61	30	(Szidat et al., 2009)	
Rural	Göteborg, Sweden	Winter/2005		1.2	0.53	27	(Szidat et al., 2009)	
Rural/semi- urban	Stockholm, Sweden	Summer/2009	TSP			12	(Kirillova et al., 2010)	
Urban	Zürich, Switzerland	Summer/2002	PM_{10}	2.1	0.54	14	(Szidat et al., 2004)	
Urban	Zürich, Switzerland	Winter/2008	PM_{10}	2.8	0.60	26.8	(Zhang et al., 2013)	
Urban	Moleno, Switzerland	Summer/2006	PM_{10}	5.3	0.67	30	(Zhang et al., 2013)	
Urban	Bern, Switzerland	Winter/2009	PM_{10}		0.39	14	(Zhang et al., 2014c)	
Urban	Atlanta, USA	Summer/2004	PM _{2.5}	2.3	0.59	26.5	(Weber et al., 2007)	
Rural	Millbrook, USA	Annual/2006/20 07	TSP		0.36	12	(Wozniak et al., 2012)	
Rural	Harcum, USA	Annual/2006/20 07	TSP		0.38	14	(Wozniak et al., 2012)	
Regional background	Cesar, Netherlands	Annual/2011/20 12	PM _{2.5}	2.3	0.65	21	(Dusek et al., 2017)	
Average						21±8		

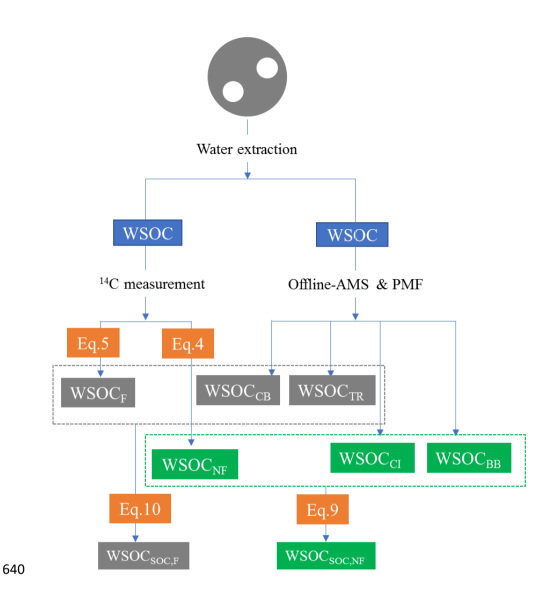


Figure 1. The AMS²-based source apportionment scheme of WSOC aerosols in this study.

- 642 See the main text for the equations (i.e., Eq. 4, 5, 9, 10 in the Sec. 2.5) and the offline-AMS &
- 643 PMF (see the Sec. 2.3).

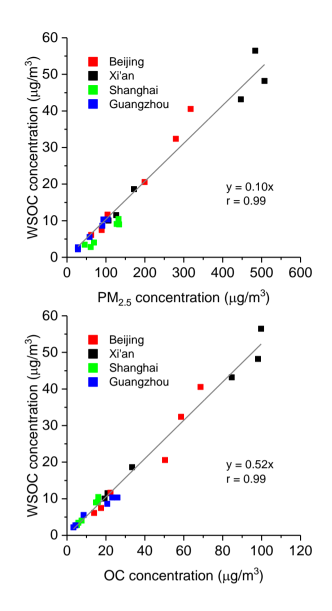
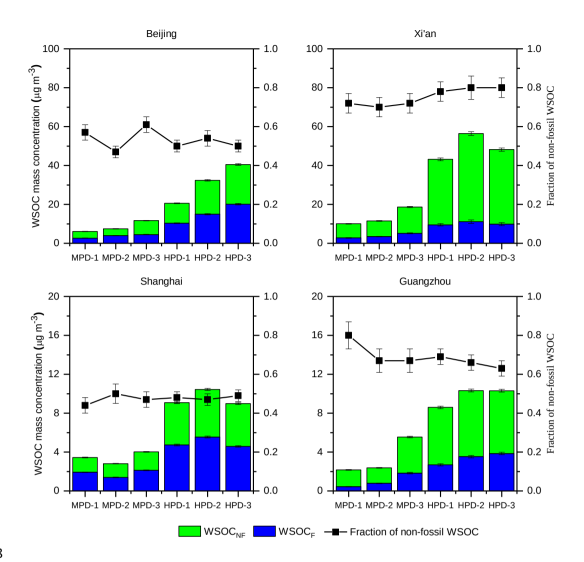


Figure 2. Linear relationships (p<0.01) of WSOC with PM_{2.5} (top) and OC concentrations
(bottom).



649 Figure 3. Mass concentrations ($\mu g/m^3$) of WSOC from non-fossil and fossil-fuel sources

 $(WSOC_{NF} and WSOC_{F}, respectively)$ as well as non-fossil fractions of the WSOC aerosols from

651 Beijing, Xi'an, Shanghai and Guangzhou during moderately polluted days (MPD) and heavily

652 polluted days (HPD). Note the different scaling for different cities.

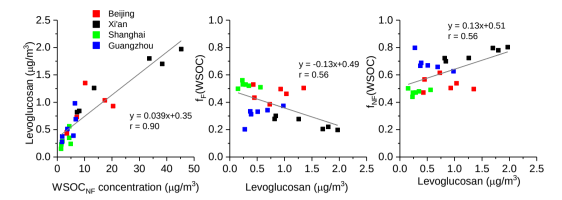


Figure 4. Relationships of non-fossil derived WSOC (WSOC_{NF}) and levoglucosan (left), levoglucosan and fraction of fossil to WSOC (f_F (WSOC)) (middle) and levoglucosan and

fraction of non-fossil to WSOC ($f_{NF}(WSOC)$) (right).

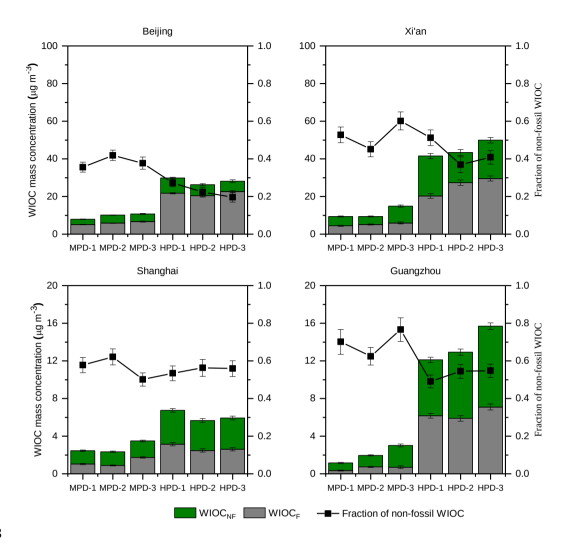
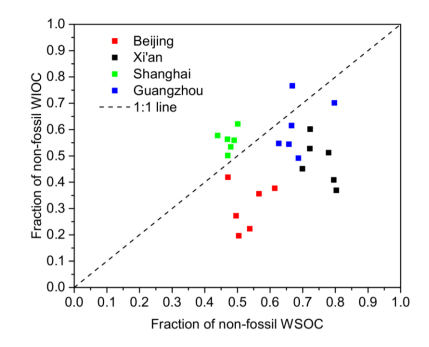


Figure 5. Mass concentrations (μ g/m³) of WIOC from non-fossil and fossil-fuel sources (WIOC _{NF} and WIOC _F, respectively) as well as non-fossil fractions in the WIOC aerosols from Beijing, Xi'an, Shanghai and Guangzhou during moderately polluted days (MPD) and heavily polluted days (HPD). Note the different scaling for different cities.







(a)

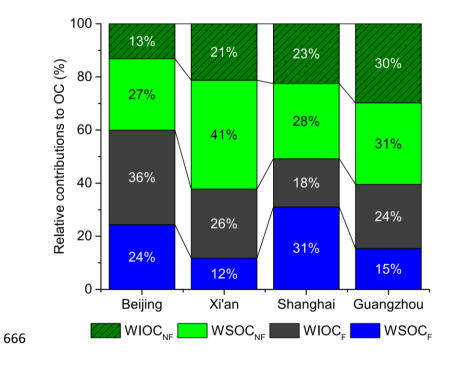




Figure 6. Relationship between the fraction of non-fossil WIOC and WSOC(a) and averaged
relative contribution (%) to OC from WSOC and WIOC from non-fossil and fossil sources (b).

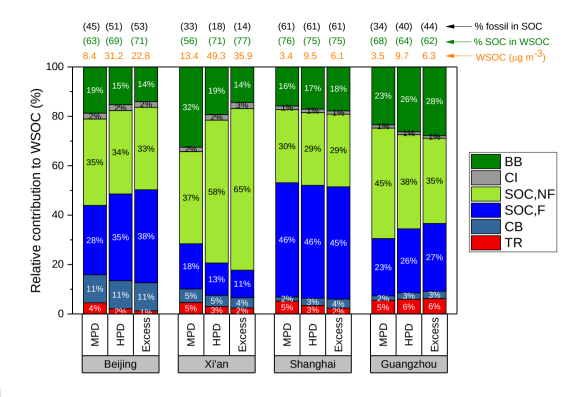
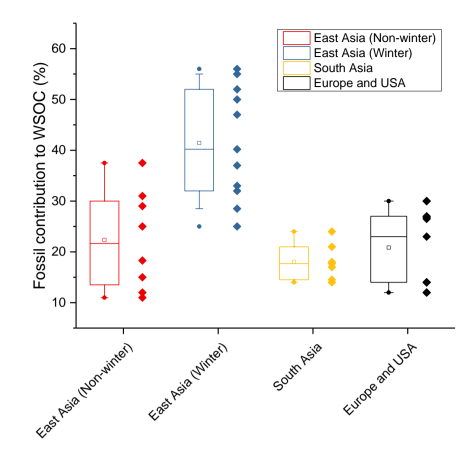




Figure 7. Relative contributions (%) to WSOC from biomass burning as well as secondary
organic carbon (SOC) from fossil and non-fossil sources (WSOC_{SOC,F} and WSOC_{SOC,NF},
respectively) in different cities during moderately polluted days (MPD) and heavily polluted
days (HPD) as well as their corresponding excess (Excess=HPD-MPD). The numbers above

the bars refer to the average WSOC concentrations and the SOC fractions (%) of WSOC.



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Figure 8. Box-plot of the fossil contribution (%) to the WSOC aerosols in East Asia, South Asia, USA and Europe. The box represents the 25th (lower line), 50th (middle line) and 75th (top line) percentiles; the empty square within the box represent the mean values; the end lines of the vertical bars represent the 10th (below the box) and 90th (above the box) percentiles; the solid dots represents the maximum and minimum values; the solid diamonds represent the individual data (Table 1). The data from East Asia is grouped by the winter and non-winter seasons.