2 Dear Editor,

Author's Response

We sincerely thank all reviewers for their helpful comments and guidance. We haveprovided responses to Referee #4 comment below in blue.

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6 Referee #4:

7 Almost all of my questions have been answered by authors, and I supposed that the manuscript

- 8 could be published after some minor revisions.
- 9 1.The abstract should be rewritten and be more concise.
- 10 Response:

Thanks. We have rewritten and shortened the abstract (from 487 words to 300 words) inthe revised MS to make it more concise as follows:

'Based on the published literatures and typical profiles from the source library of Nankai 13 University, a total of 3326 chemical profiles of the main primary sources of ambient 14 particulate matter across China from 1987 to 2017, are investigated and reviewed to trace 15 the evolution of their main components and identify the main influencing factors to the 16 17 evolution. In general, the source chemical profiles are varied with sources and influenced by different sampling methods. The most complicated profiles are likely attributed to coal 18 combustion and industrial emissions. The profiles of vehicle emissions are dominated by 19 organic carbon (OC) and elemental carbon (EC), and varied with the changing standards 20 of sulfur and additives in the gasoline and diesel as well as the sampling methods. In 21 addition to sampling methods, the profiles of biomass burning and cooking emissions are 22 also impacted by the biofuel categories and cooking types, respectively. The variations of 23 the chemical profiles of different sources, and the homogeneity of the sub-type source 24 profiles within the same source category were examined with uncertainty analysis and 25 cluster analysis. As a result, a relatively large variation has been found in the source 26 profiles of coal combustion, vehicle emissions, industry emissions and biomass burning, 27 indicating that these sources have the priority to establish the local profiles due to their 28 29 high uncertainties. The presented results highlight the need for increasing investigation of 30 more specific markers (e.g., isotopes, organic compounds and gaseous precursors) beyond routine measured components to discriminate sources. Although the chemical profiles of 31 main sources have been reported previously in literatures, it should be noted that some of 32 these chemical profiles are out of date currently, which needs to be updated immediately. 33 34 Additionally, specific focus should be placed on the sub-type of source profiles in the future, especially for local industrial emissions in China.' 35 2. There still are some grammar faults, which should be corrected. 36

37 Response:

Thanks. We have checked the MS again and found some grammar faults, such asgrammatical tense, singular & plural faults. All the corrections have been demonstrated in

40 the revision mode of the revised MS followed this response.

41 3.The marks of the figures are not clear, especially for Figure 2.

42 Response:

- 43 Thanks. We have plotted the figures again in the revised MS to make them clearer.
- 44

Characteristics of the main primary source profiles of particulate matter 45 across China: from 1987 to 2017 46

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54	
55	Abstract
56	Based on the published literatures and typical profiles from the source library of Nankai
57	University, a total of 3326 chemical profiles of the main primary sources of ambient
58	particulate matter across China from 1987 to 2017, including coal combustion, industrial
59	emissions, vehicle emissions, fugitive dust, biomass burning and cooking emissions, were
60	are investigated and reviewed to trace the evolution of their main components and identify
61	the main influencing factors to the evolution. In general, the source chemical profiles are
62	varied with sources and influenced by different sampling methods. As a result, tThe most
63	complicated profiles are likely attributed to coal combustion and industrial emissions,
64	which are evidently influenced by the decontamination processes and sampling techniques
65	as well as the coal properties and the boiler types. The profiles of vehicle emissions are
66	dominated by organic carbon (OC) and elemental carbon (EC), and varied with the

67	changing standards of sulfur and additives in the gasoline and diesel as well as the
68	sampling methods. The profiles of fugitive dust, such as soil dust and road dust, are
69	dominated by the crustal materials and influenced by the sampling methods to some
70	extent. In addition to sampling methods, Tthe profiles of biomass burning and cooking
71	emissions are also impacted mainly by the biomass biofuel categories and sampling
72	methods. As expected, the profiles of cooking emissions are impacted mainly by and the
73	cooking types, respectively and materials. The variations of the chemical profiles of
74	different sources, and uncertainty analysis and cluster analysis of all these source profiles
75	are conducted to reveal the variations the homogeneity of the different sub-type source
76	profiles in the within the same source category and evaluate the differences between
77	source categorieswere examined with uncertainty analysis and cluster analysis. As a result,
78	A <u>a</u> relatively large variation has been founded in the source profiles of coal combustion,
79	vehicle emissions, industry emissions and biomass burning, indicating that it is necessary
80	to establish the local profiles for these sources have the priority to establish the local
81	profiles due to their high uncertainties. While the profiles of road dust and soil dust
82	presented a less variation with the stable chemical characteristics among the different
83	profiles in the same category, suggesting that the profiles of these sources could be
84	referenced for the cities in China when such local profiles are not available. The presented
85	results highlight the need for increasing investigation of more specific markers (e.g.,
86	isotopes, organic compounds and gaseous precursors) beyond routine measured
87	components to discriminate sources. Although the chemical profiles of main sources have
88	been reported previously in literatures, it should be noted that some of these chemical

profiles are out of date currently, which needs to be updated immediately. Additionally,
specific focus should be placed on the sub-type of source profiles in the future, especially
for local industrial emissions in China, to support the air quality research communities in
their efforts to develop high resolution source apportionment for making more effective
control strategies. *Keywords*: Source profiles; particulate matter; source apportionment.

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97 **1. Introduction**

In light of preventing us from being exposure to high level of PM, source apportionment 98 technique is a critical tool to help us in quantitative recognition of the source contributions 99 of ambient particulate matter (PM) and developing efficient and cost-effective abatement 100 policy. Given the thousands of PM sources in real-world, localized source information is 101 crucial to accurate source identification and contribution estimation. The physical and 102 chemical characterization of primary sources, termed source profile, is of great 103 importance in the application of receptor models for source apportionment study as it 104 characterizing specific sources from the physicochemical point of view that revealing the 105 signatures of source emissions (Watson, 1984; Bi et al., 2007; Simon et al., 2010; Hopke, 106 107 2016). Since the real-world measurement of source samples is costly and tough, many 108 studies us<u>eding</u> factor analytical models (source-unknown models, such as positive matrix 109 factorization (PMF), principle component analysis (PCA) etc.) instead of chemical mass balance (CMB) model (source profiles need to be known a priori) to estimate source 110 111 contributions. However, the measurement of sources is essentially a very important basic work to help obtain source signatures and then makes source identification and 112

apportionment possible. It should be noted that the interpretation of factors deduced from PMF analysis is based on the available source profiles (Shi et al., 2009; Simon et al., 2010; Liu et al., 2017; Hopke, 2016). In addition to source apportionment study, source profiles have also played an important role in calculating source-specific emissions of individual compounds and converting total emissions from sources into the speciated emissions for air quality models, which can further provide effective strategies for environmental management (Reff et al., 2009; Simon et al., 2010).

In the past decades, source profiles of PM from a variety of source types were 120 121 substantially developed all over the world, especially in USA (Simon et al., 2010), Europe (Pernigotti et al., 2016) and East Asia (Liu et al., 2017). The time evolution of source 122 profiles is partly determined by the source apportionment techniques. In general, the 123 124 receptor model was developed based on the assumption of mass conservation (Winchester and Nifong, 1971; Miller et al., 1972). A mass balance equation represents that the 125 measured particle mass can be regarded as the linear sum of the mass of all chemical 126 127 components contributed from several sources (Cooper and Watson, 1980; Watson, 1984). 128 Initially, the mass balance equations were deployed for a couple of specific elements and source types in America (Miller et al., 1972; Hopke, 2016). Elements, ions and carbon 129 materials gradually tend to be the routine chemical species in the source apportionment of 130 131 PM. With the development of advanced sampling and chemical analysis techniques, more valuable information, such as organic compounds (Schauer and Cass, 2000; Simoneit et 132 133 al., 1999), isotopic measurement of radiocarbon (Wang et al., 2017), sulfur (Han et al., 2016) and nitrogen (Pan et al., 2016), high-resolution aerosol mass spectra (Zhang et al, 134

2011) and particle size distribution (Zhou et al., 2004) etc., have been explored to further 135 expand the existing or new profiles. These is information has have been proved to provide 136 source specificity capable of being incorporated into receptor models as new markers 137 (Zheng et al., 2002), constraining source contributions (Amato et al., 2009), and 138 developing new models (Ulbrich et al., 2012; Dai et al., 2019). For example, Dai et al. 139 (2019) developed a size-resolved CMB approach for source apportionment of PM based 140 on the size profiles of sources. The new valuable information improves the performance 141 of source apportionment models to obtain more precise and reliable results. 142

143 Since the 1980s, source profile studies were initially implemented in China (Dai et al., 1987). During the past three decades, hundreds of source profiles have been achieved 144 across China (Zhao et al., 2006; Bi et al., 2007; Zhao et al., 2007; Kong et al., 2011; Kong 145 146 et al., 2014; Qi et al., 2015; Wang et al., 2015; Zhang et al., 2015; Zhao et al., 2015; Pei et al., 2016; Tian et al., 2017; Guo et al., 2017). These profiles covered more than forty cities 147 and several source types. The main ubiquitous sources of atmospheric PM in China during 148 149 the past three decades can be roughly divided into coal combustion sources (CC, with sub-type sources of coal-fired power plants, coal-fired industrial boiler and residential 150 coal combustion), vehicle exhaust (VE, gasoline and diesel engines), industrial processes 151 emissions (IE), biomass burning (BB), cooking emissions (CE), fugitive dust (FD, with 152 sub-type sources of soil fugitive dust, construction dust and road dust) and other localized 153 specific sources. These available profiles have filled the gap of the knowledge of source 154 compositions and provided effective markers for the source apportionment studies. 155 However, the current state and potential issues of pre-existing primary source profiles of 156

157 PM in China are still unclear, it is time to overview these source profiles along the time158 line and add more profile knowledge to the atmospheric research community.

In fact, more real-world measured profiles in China were actually not published. A 159 database of particulate source profiles founded by Nankai University contains 2870 160 profiles measured across China since the 1980s. In this paper, the characteristics and time 161 evolution of the published primary profiles and some typical profiles of particulate matter 162 founded by Nankai University were discussed. To collect the potential published data 163 related to source profiles, a two-round literature search work covering literature from 164 165 1980 to 2018 was done in this work. In the first round of searching, two authors are responsible for the same source to ensure every source category has been searched twice 166 independently. The search keywords depend on source category. The following keywords 167 168 for each source were used individually or in combination. As for CC sources, the key words combustion/coal burning/coal-fired boiler/coal-fired are "coal power 169 plant/residential coal" and "source profile/chemical profile/particle composition". The key 170 171 words for other sources are shown as follows. IE: "industrial emission" and "source profile/particle composition"; "vehicle profile/chemical VE: emission/exhaust 172 emission/traffic emission/diesel engine/truck emission/gasoline engine/on-road 173 vehicle/tunnel experiment/chassis dynamometer/portable emission measurement system" 174 and "source profile/chemical profile/particle composition"; CE: "cooking emission" and 175 "source profile/chemical profile/particle composition"; BB: "biomass burning/bio-fuel 176 boiler" and "source profile/chemical profile/particle composition"; FD: "soil/fugitive 177 dust/crustal material/construction dust/road dust" and "source profile/chemical 178

profile/particle composition". Papers and dissertations in Chinese on China National 179 Knowledge Infrastructure (CNKI) and papers in English on the web of science were 180 searched using above keywords, respectively. The duplicated paper was then 181 double-checked and excluded. The papers with topic related to source profiles but without 182 providing any information of real-measured sources were also excluded. For example, 183 papers reported source apportionment results with the use of PMF and CMB but without 184 reporting local profiles were not taken into account. As a result, a total of 193 papers have 185 been collected from these efforts. In the second round of searching, the valid papers with 186 187 available source profile data and detailed source sampling and chemical analysis methods were counted and used for post-analysis. Finally, a total of 456 published source profiles 188 since the 1980s across China were collected.a total of 456 published source profiles, 189 190 coupled with the database of source profiles (2870 profiles) founded by Nankai University are reviewed in this work. 191

This review is based on the following ideas. In Section 2.1, we summarized the types and 192 193 the number of particulate source profiles in China published since the 1980s, and reviewed the technological development of the sampling and chemical analytical methods 194 for source samples. In Section 2.2, the characteristics and time evolutions of the 195 ubiquitous source profiles in China (CC, VE, IE, BB, CE and FD) in terms of the marker 196 species of each main source and the effect of various impact factors on source profiles 197 have been discussed. In section 2.3, the homogeneity of the sources within the same 198 source category and the heterogeneity between different source categories were further 199 investigated by using the coefficient of variation (CV, the standard deviation divided by 200

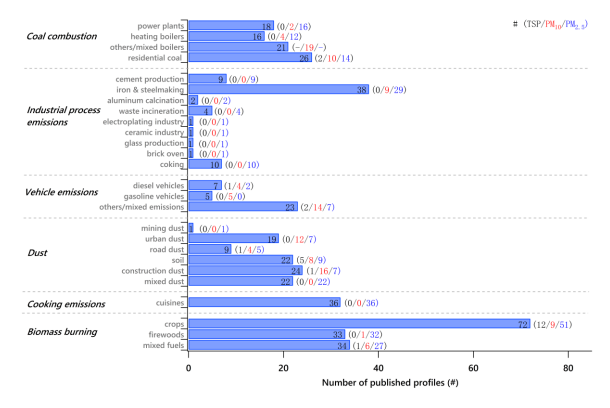
the mean) and cluster analysis, respectively. In Section 3, we summarized the main
findings and a few issues of current source profiles, as well as the future requirements for
the on-going development of source profiles in China.

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205 **2.** Overview of source profiles across China

After literature searching (peer-reviewed papers published in international and Chinese journals), a total of 456 published source profiles since the 1980s across China were collected. In general, all of these profiles were subjectively divided into the above six source categories, with 81 of them were attributed to CC, 67 to IE, 35 to VE, 98 to FD, 36 to CE, and 139 to BB. For the certain aerodynamic sizes, we obtained a total of 306 PM_{2.5} profiles, 123 PM₁₀ profiles, and 27 for other sizes. The overview of these profiles are shown in Fig. 1.

These published profiles were detected in different parts of China. In eastern China, there 213 are published profiles of 35 CC (excluded residential coal combustion), 14 IE, 14 VE, 18 214 215 BB, 2 CE, and 14 FD; in northern China, there are published profiles of 16 CC, 23 IE, 9 VE, 8 BB, 13 CE, and 62 FD; in western China, there are only profiles of 20 CC; in 216 southern China, there are published profiles of 10 VE, 10 CE, and 5 FD; in central China, 217 there are published profiles of 17 BB. The profiles of residential coal combustion are 218 mainly detected in the regions that have obvious activities of residential coal burning, 219 such as the northern and western China. The region of different parts of China was 220 221 defined by Zhu et al. (2018).



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Figure 1. Overview of the published source profiles across China.

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225 **2.1 Development of sampling and analysis techniques**

Sampling for source emissions. In the past three decades, the sampling techniques used in 226 the source apportionment research in China have been significantly improved to catch the 227 real-world emissions of particles from various complex primary sources. In the 1980s, CC 228 is the predominant sources of PM in China (Dai et al., 1987). The source measurement of 229 CC was mainly performed by collecting the dust directly from the precipitators. Source 230 samples of FD was-were collected from the surface of fugitive dust sources (soil, road 231 dust, et al.) (Dai et al., 1987; Qu, 2013). Apparently, such sampling method cannot catch 232 the real-world emissions from the sources to the ambient air, especially for the CC or 233 other emission sources with humid and high-heat fume. The compositions of the PM in 234 235 such fume appears to be changing due to the physical condensation and chemical

reactions during their dispersion process in the ambient air. Since the 1970s, dilution 236 tunnel sampling method (DTSM) has been developed to originally obtain source samples 237 238 from vehicle emissions that could be close to the real compositions from the sources (Hildemann et al., 1989). Subsequently, various dilution tunnels have been developed 239 with different tunnel materials, resident time, dilution ratios, diameter of effective mixing 240 lengths to collect particles emissions from stationary sources (Houck et al., 1982; Smith et 241 al., 1982; Hildemann, 1989). The development and application of such technique in China 242 was after 2000 (Ge et al., 2001; Ge et al., 2004), while it has been widely used nowadays 243 244 (England et al., 2000; Lind et al., 2003; Ferge et al., 2004; Zhou et al., 2006; Li et al., 2009; Wang et al., 2012). 245

As for fugitive dust, another problem is to collect the particle samples with certain 246 247 aerodynamic size from the dust samples. In the 1980s-1990s, the Barco particle size analyzer was used to obtain the size distributions from the source samples 248 (Kauppinen et al., 1991). Due to the low efficiency and potential safety risk of Barco 249 250 sampler, a new sampling technique called the resuspended chamber (RSM) was developed in the 1990s by Chow et al. (1994), which has been widely used since 2000 in 251 China. This method is capable of obtaining the particle sample with certain aerodynamic 252 sizes from the dust powder collected from the source field. Nowadays, most source 253 samples with the particle aerodynamic size of 2.5 µm or 10 µm of fugitive dust were 254 collected by the resuspended sampling method in China (Ho et al., 2003; Zhao et al., 255 2006). Although the resuspended chamber couldn't completely simulate the real 256 environment, it still is the best available choice for the collection of fugitive dust samples 257

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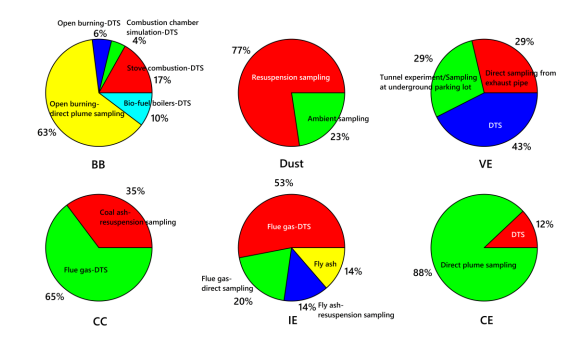


Figure 2. Share of sampling methods for the samples collection of each source type in China fromliteratures. DTS denotes dilution tunnel sampling method.

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Except for stationary sources, the moving sources like vehicle emissions are gradually becoming the dominant source in megacities of China. A variety of measurement methods for vehicle emissions have been developed over the world, such as directly measurements on the exhaust emissions of on-road vehicle and chassis dynamometer, portable emission measurement system as well as tunnel experiment.

As for biomass burning and residential coal combustion, with the use of DTSM, the measurements involved in different combustion ways. For example, indoor/lab simulation with fuel burned with stove/chamber, and open burning/field measurement. In addition to this, the biomass fuel can be burned in bio-fuel boilers, which is on the way to 273 industrialize recently.

- In the published profiles, 65% coal combustions, 53% industrial emissions, 12 cooking emissions, 43% vehicle emissions, and 37% biomass burning profiles were obtained with
- 276 DTSM (as shown in Fig. 2).
 277 *Chemical analysis*. The chemical an
- 277 *Chemical analysis*. The chemical analysis methods have been significantly improved 278 since the 1980s. A typical source profile from literature data usually contains elements 279 (e.g., Al, As, Ca, Cd, Cr, Cu, Fe, K, Mg, Mn, Na, Pb and Zn), organic carbon (OC), 280 elemental carbon (EC), and water-soluble ions (WSI, e.g., Cl⁻, NO₃⁻, SO₄²⁻, NH₄⁺, K⁺, 281 Na⁺-,_Mg²⁺ and Ca²⁺) in China. Detailed procedures in terms of the establishment of 282 different source profiles are available in previous publications (Chow et al., 1994;Chow et 283 al., 2004;Hou et al., 2008b;Pei et al., 2016).

284 PM samples collected on Teflon filters were mostly analyzed for elements by Inductively Coupled Plasma Optical Emission Spectrometer (ICP-OES) or Inductively Coupled 285 286 Plasma Atomic Emission Spectrometer (ICP-AES) in China. In recent years, Inductively 287 Coupled Plasma Mass Spectrometry (ICP-MS) and X Ray Fluorescence were also used, 288 with measurement systems which have lower threshold/higher accuracy and quick response, respectively (Tsai et al., 2004). The total carbon (TC) mass is typically 289 290 determined using thermal and or thermal-optical methods. With the use of thermal/optical carbon analyzer, there are two widely used approaches to divide organic carbon (OC) and 291 elemental carbon (EC) from TC, named DRI IMPROVE_A and NIOSH 5040, which are 292 operationally defined by the time-temperature protocols, the OC/EC split approaches by 293 optical reflectance/transmittance. (Chow et al., 1994;Ho et al., 2003;Chow et al., 294 295 2004;Zhang et al., 2007;Phuah et al., 2009). Quartz fiber filters were normally used for the determination of WSI by different types of Ion Chromatography (IC) with high-capacity 296 cation-exchange column and anion-exchange column (Qi et al., 2015). 297

298 Organic Ttracer species, a unique species that can be used as an indicator of a particular source, playing an important role in estimating source contributions. However, most of the 299 source profiles in China are reported with inorganic species, with only a few studies 300 providing information of organic compounds. Organic tracers are of great value in source 301 apportionment studies, as it provides more source-specific information in addition to 302 inorganic species. For example, leveglucosan is a well-known organic tracer represents 303 304 for biomass burning (Lee et al., 2008), azzaarenes as markers of inefficient coal combustion (Junninen et al., 2009; Bandowe et al., 2016), sterols, monosaccharide 305 306 anhydrides and amides as a marker of cooking emissions (Schauer et al., 1999; Schauer et al., 2002; He et al., 2004; Zhao et al., 2007a,b; Cheng et al., 2016;). Furthermore, for 307 better discriminating sources, Pb stable isotopes, which are not obviously influenced by 308 ordinary chemical, physical or biological fractionation processes (Gallon et al., 2005; 309 Cheng and Hu, 2010), were determined with an ICP-MS. Additionally, some other isotope 310 measurements, for example radiocarbon (Wang et al., 2017), sulfur (Han et al., 2016), and 311 nitrogen (Pan et al., 2016), as well as natural silicon (Lu et al., 2018), have also been 312 reported to be used as source indicators recently. 313

The above efforts indicate that the reported source profiles were collected by various sampling methods and chemically analyzed by different instruments, making the source profiles a high uncertainty of comparability. It is necessary to establish standards for the procedures of source sampling, chemical analysis and QA/QC to ensure the representativeness, validation and comparability of source profiles in China.

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320 **2.2** Characteristics and evolution of source profiles

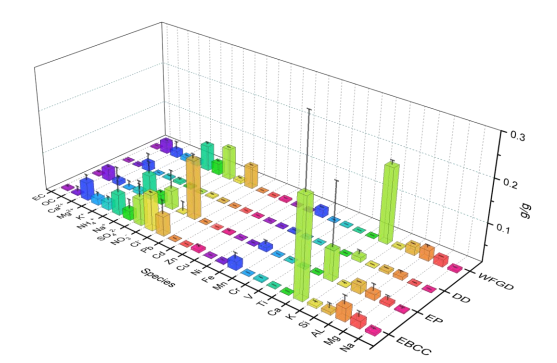
321 **2.2.1 Coal combustion**

322 Coal is the main fuel used in China which has been widely used in coal-fired power plants,

323 coal-fired industrial boilers and residential household stoves, on average accounted for more than 60% of the total energy consumed in China (CESY,2015). Thus, it was the 324 325 main cause of air pollution particularly during heating-season in Northern China. The source profiles of CC sources are influenced by many factors, such as coal type and 326 property, boiler/stove type and efficiency, burning conditions (burning rate and fuel 327 loading), decontamination devices etc. (Shen, 2010), making it appears to be the most 328 complicated type among the primary sources. The source profiles of CC in China are 329 mainly consisted of crustal materials, OC, EC, SO4²⁻ and trace metals, indicating the 330 331 chemical nature of coal burning.

Coal-fired power plants. Within the same sampling method (dilution tunnel sampling 332 method) and the same boiler type, the characteristics of the source profiles of coal-fired 333 334 power plants equipped with different dust removal and desulfurization facilities are compared (Fig. 3). OC, EC, and Cl⁻ in the profiles of the electrostatic precipitators (EP) 335 are higher than that in the electric bag compound dust collectors (EBCC), with average 336 values of 0.0289 ± 0.0342 , 0.0036 ± 0.0033 g/g and 0.1403 ± 0.1686 g/g, respectively. 337 Higher Ca, NO₃⁻, Ca²⁺in the source profiles obtained by the EBCC is found as well. 338 Comparing data from different desulfurization facilities (Fig. 3), SO₄²⁻ and Ca in PM_{2.5} 339 profiles from the wet flue gas desulfurization (WFGD) is much higher than that from dry 340 desulfurization (DD). It is reported that SO_4^{2-} is converted from SO_2 in the flue gas 341 through a limestone slurry washing reaction and then discharged with the fume (Ma et al., 342 343 2015). Ca is also infused in the fume when the flue gas went through the limestone washing process. OC in PM_{2.5} profiles from the WFGD is also higher than that from DD, 344

suggestinged that the possible conversion of gaseous or liquid organics to the particulate state in the lime slurry. NH_4^+ , Na^+ , and Cl^- are also higher in WFGD profiles than that in DD. The formation mechanism of these species in the WFGD needs further investigation.



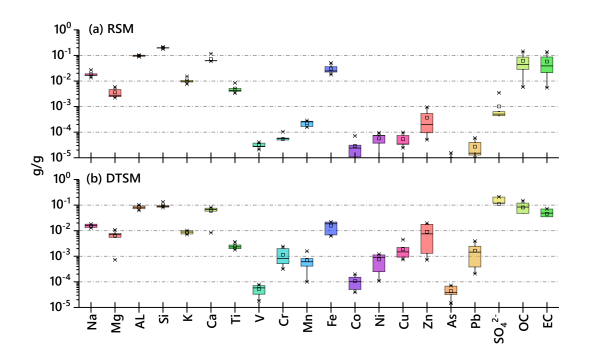
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Figure 3. Comparison of PM_{2.5} source profiles collected under different dust removal and
desulfurization facilities. EP denotes electrostatic precipitators, EBCC denotes electric bag
compound dust collectors, WFGD denotes wet flue gas desulfurization, DD denotes dry
desulfurization. Data from the source library of Nankai University) were counted.

To evaluate the impact of different sampling methods on the contents of source profiles, measurements with the coal ash resuspension sampling method (RSM) and the stack gas DTSM were simultaneously used for source sampling at a coal-fired power plant in Wuxi, China-were compared. The results of the obtained PM₁₀ source profiles are shown in Fig. 4. For RSM, the crustal elements (Si, Mg, Al and Ti) –are significantly higher than

DTSM, while the SO₄²⁻ fraction of DTSM is significantly higher than RSM, reaching 360 0.1643 g/g. V, Cr, Mn, Co, Ni, Cu, Zn, Pb and other trace metal fractions are strongly 361 enriched in DTSM, which is 1.7 to 60.7 times that in RSM, suggesting that these trace 362 metal elements have a low melting point and are easily liquefied or gasified during 363 combustion, and then condensed on the surface of the particles in the flue or after exiting 364 the flue (where small particles have a large specific surface area and are more prone to 365 enrichment) (Dai et al., 1987). The similar results were also reported earlier elsewhere 366 (Meij, 1994; Meij and Winkel, 2004; Zhang et al., 2009b). 367

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Figure 4. Characteristics of chemical profiles for PM₁₀ emitted from coal-fired power plant
obtained by different sampling methods in Wuxi city. RSM and DTSM denote resuspension
sampling method and the dilution tunnel sampling method, respectively. Data were from the
source library of Nankai University were counted.

Coal-fired industrial boiler. The coal-fired industrial boilers are used for providing hot 375 water or steam for industry or municipal heating. These boilers consumed about 1.1 376 billion tons of coal annually in China, accounting for 25% of the total coal consumption 377 and only have the average capacity of 2.7 MW (ERI, 2013). Comparing with the profiles 378 379 detected in coal-fired power plants, there are substantial differences in the source profiles 380 of the coal-fired industrial boilers. Fig.-5 shows the difference of the chemical compositions of source profiles between coal-fired industrial boilers with wet 381 desulfurization (IBW) and power plant boilers with wet desulfurization (PPW) with PM 382 samples collected using the same method. Mg, Al, Si, Ca, SO₄²⁻, NH₄⁺ and OC in the 383 profiles of PPW are higher than that of IBW, which was likely resulted from the 384 combustion efficiency and desulfurization efficiency, as PPW was required to operate 385 386 with high efficiency of desulfurization by the government while IBW was less under controlled. 387

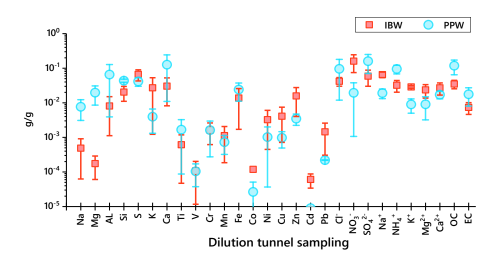




Figure 5. Average and standard deviation of chemical species in the source profiles of coal-fired industrial boilers equipped with wet desulfurization (IBW) and power plant boilers equipped with wet desulfurization device (PPW), respectively. Data were collected from the source library of

392 Nankai University.

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Residential Coal Combustion (RCC). In 2015, the total amount of coal consumption in 394 mainland China is about 3970.14 Mt with a total of 93.47 Mt coal consumed in residential 395 section (CESY, 2015). RCC is an important source of atmospheric PM in rural area, 396 particularly in heating-season (Duan et al., 2014; Tao et al., 2018; Chen et al., 2005; 397 Zhang et al., 2007; Chen et al., 2004). Contrary to industrial furnaces and boilers, coal 398 burned in household stoves has a significant impact on indoor and outdoor air quality in 399 400 terms of its low thermal efficiency, incomplete combustion and the lack of air pollutant control devices. It was reported that the emission factors of air pollutants for coal burned 401 in household stoves are more than two orders of magnitude higher than those burned in 402 403 industrial boilers and power plants (Li et al., 2017), thus pollutants emitted from RCC have drawn great concern in recent years. 404

In general, coals can be classified as anthracite and bituminous coals in the forms of raw 405 406 chunks and briquettes (Shen, 2015), burneding with a movable brick or cast-iron stoves 407 that has been used over centuries in China (Shen et al., 2010). There are many real-world 408 measurements on particle emissions from RCC aiming to investigateing its emission nature (Chen et al., 2005). Most studies have rather placed focus on the emission factors 409 410 than chemical composition as the emission factor of RCC is high uncertain for a given air pollutant. The chemical characteristics of RCC profiles are varied greatly with the 411 412 sampling techniques. Three decades ago, Dai et al (1987) reported the averaged elemental profile of 15 RCC particle samples in Tianjin in 1985, with the use of Barco analyzer to 413

cut fly ash (collected from the stack of RCC stove) into particles with aerodynamic 414 diameter less than 12 µm. As expected, this sampling technique resulted in a high fraction 415 416 of crustal elements in the chemical profile. The resuspension chamber has also been used to cut particle size from coal fly ash. However, the coal fly ash is not the particles 417 emission from stack. Thus, the accuracy of RCC source profile has been improved until 418 the DTSM has been introduced into China. As shown in Fig. 6, the fractions of crustal 419 elements (Mg, Al, Si, Ca, Ti) in the profile measured from coal ash are an order of 420 magnitude higher than that in the RCC profile sampled by using DTSM, while the 421 422 fraction of sulfate, nitrate and OC are two to three orders of magnitude lower in coal ash 423 PM_{2.5}.

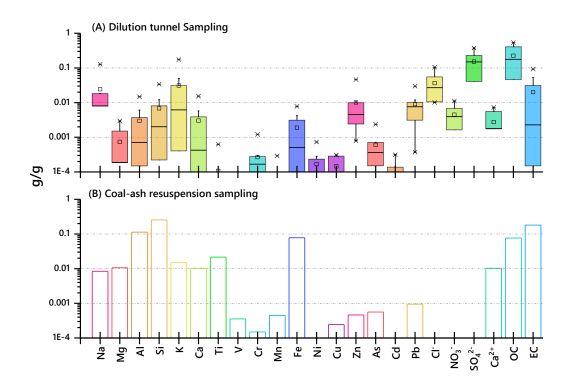


Figure 6. RCC Profiles of PM_{2.5} collected by dilution tunnel sampling method (A, data
were collected from available published profiles (Ge et al., 2004; Kong, 2014; Liu et al.,
2016; Liu et al., 2017; Yan et al., 2017; Dai et al., 2019)) and coal fly ash resuspension

428 sampling method (B, data were collected from Wang et al. (2016)).

429

430 Many efforts have been implemented in a national level to reduce pollutants emissions from RCC by introducing improved stoves and cleaner fuels since the 1990s, such as the 431 432 China National Improved Stove Program (Shen et al., 2015). The highly efficient stove is reported likely has a reduced emission load. Given the limited available data, it is unable 433 to compare the chemical profiles between the lowly and highly efficient stove in this work. 434 It is also reported that the emission factors of air pollutants from RCC varied widely 435 436 because of the variations in coal type and property, stove type and burning condition (Shen et al., 2010). As shown in Fig. 7, PM_{2.5} emissions from the burning of chunk coals 437 have a high fraction of OC, EC, sulfate, nitrate and ammonium, a low fraction of Na, Ca 438 439 and K (K⁺) than the burning of honeycomb briquette coals. Generally, OC and sulfur are the predominate species in $PM_{2.5}$ emitted by RCC. 440

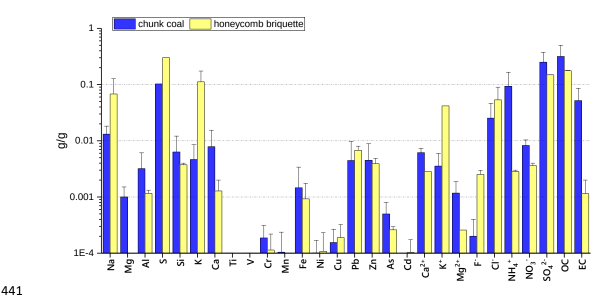


Figure 7. RCC profiles of PM_{2.5} emission from chunk coal and honeycomb briquette
coals. Data were collected from published data (Ge et al., 2004; Kong, 2014; Liu et al.,

444

2016; Liu et al., 2017; Yan et al., 2018; Dai et al., 2019).

445

As we mentioned above, there are many factors that affecting the profiles of coal
combustion sources. Therefore, local CC source profiles should be measured in the study
area to improve the accuracy and reliability of source apportionment results.

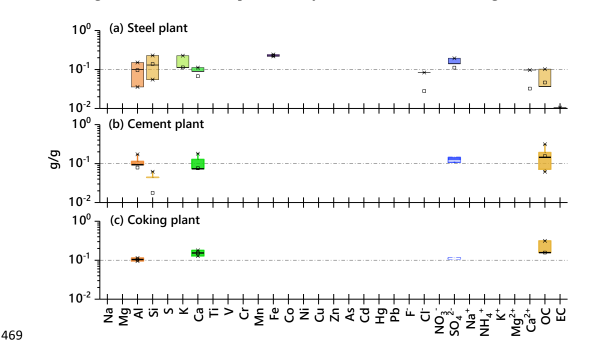
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450 **2.2.2 Industrial process emissions**

The industrial emissions are one of the most important sources in China (Zhu et al., 2018). 451 452 Particles from industrial emissions are mainly collected using DTSM (53%). The source profiles of industrial emissions could be influenced by several key factors, such as raw 453 materials used in industrial processes, manufacture processes, various sampling methods, 454 455 different sampling site, control measures taken by different factories and process operating conditions (Watson and Chow, 2001; Kong et al., 2011; Pant and Harrison, 2012; 456 Guo et al., 2017). There are great differences between the source profiles from different 457 458 industrial sources. Fig. 8 shows the chemical composition of China's main industrial emissions (cement plant, coking plant and steel plant) (Ma et al., 2015; Qi et al., 2015; 459 Yan et al., 2016; Zhao et al., 2015a). For cement industrial sources, Ca, Al, OC and SO₄²⁻ 460 461 are the most abundant species, with average value more than 0.-10 g/g. For coking industrial sources, Ca²⁺, Al and SO4²⁻ are elevated while OC displayed a somewhat 462 notable lower level. For steel industrial sources, the highest fraction species are Fe, Si, K 463 and SO_4^{2-} , while Cl⁻, Ca²⁺, EC and OC showed a lower content less than 0.0010 g/g. 464

465 In China, there are many industrial types with different emission characteristics. The

source profiles of industrial emissions are far from being fully understood so far. The
profiles of some important industrial sources, such as the glass melt kiln, non-ferrous
smelting, and ceramics, are reported rarely and needed further investigation in the future.



470 Figure 8. Characteristics of chemical profiles for particulate matter emitted from industrial
471 emissions. Data from the source library of Nankai University, Zhao et al. (2015), Qi et al. (2015),
472 Ma et al. (2015) and Yan et al. (2016) were counted.

473

474 2.2.3 Vehicle emissions

475 Vehicle emissions appears to be the predominant source of ambient $PM_{2.5}$ in urban areas

in China, particularly in megacities like Beijing and Shanghai (Cai et al., 2017b; Cui et al.,

- 477 2016; Zhang et al., 2015). It is reported that the contribution of vehicle emissions to PM_{2.5}
- 478 was in the range of 5% to 34% over China based on receptor models (Zhang et al., 2017b).
- 479 There are many factors affecting vehicle emissions such as fuel types, vehicle types,
- 480 emission control technologies, operating conditions, engine performances, sampling

methods and so on (Watson et al., 1990; Chen et al., 2017b; Maricq, 2007). The 481 representativeness of the source profiles of vehicle emissions is often controversial. Fig. 9 482 summarizes the PM₁₀ source profiles of different vehicle types obtained by direct 483 sampling method in China (Chen et al., 2017b). For both diesel and gasoline vehicles, 484 their emission profiles are dominated by OC, EC, NO₃⁻, NH₄⁺, SO₄²⁻, Ca, Fe and Zn. The 485 abundance of EC in diesel vehicle exhaust (particularly in heavy-duty diesel vehicle 486 exhaust) is higher than that in gasoline vehicles, which may due to the different 487 combustion completion rates between diesel and gasoline on account of the length of 488 489 hydrocarbon chains of them (Chen et al., 2017b). Since Mn has been used in the gasoline explosion-proof agent, the fraction of Mn in the particulate matter from the gasoline 490 vehicle emission is higher than that of diesel vehicle. 491

492

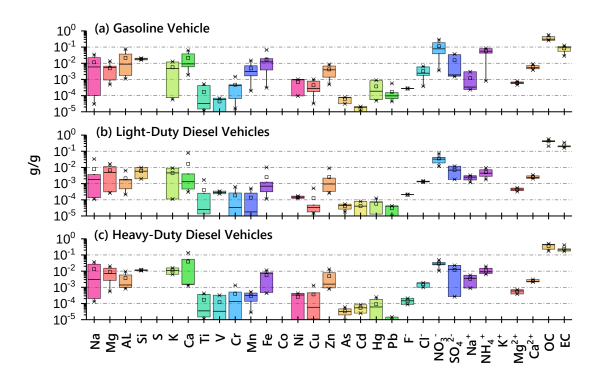




Figure 9. Chemical compositions of source profiles for PM₁₀ of different vehicle types obtained

495 by direct sampling method. Data from the source library of Nankai University and Chen et al.

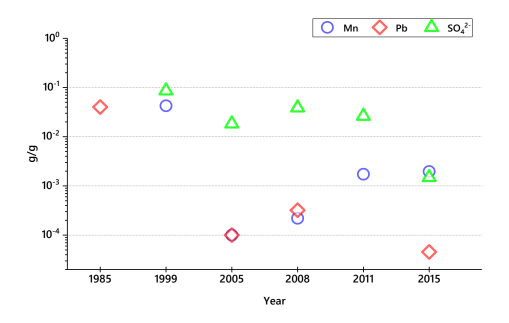
496 (2017) were counted.

497

Fig. S1 summarizes the characteristics of chemical profiles for particulate matter emitted from vehicles obtained by different sampling methods. Crustal elements (Si, Al, Ca, Mn) in the chemical profiles obtained by SDSM are higher than that of DSM, which may due to the influence of suspended road dust. NH_4^+ and NO_3^- in chemical profiles obtained by DSM are lower than that of SDSM, probably because their precursors are still in the gaseous state when the samples were are collected at a higher temperature by DSM (Kong and Bai, 2013).

The source profiles of the vehicle exhaust also varied with upgrades of the fuel. In China, 505 the oil used for vehicle has been upgraded for five times in the past eighteen years. The 506 evolutions of the fractions of Mn, Pb and SO_4^{2-} in particulate matter emitted by vehicle 507 from the past three decades are shown in the Fig. 10. Pb was a tracer of the gasoline 508 before 2000 while leaded gasoline was banned to be used in mainland China after 2000 509 510 (State Council of China, 1998). The standard value of sulfur in the car-used gasoline was $800 \ \mu\text{g/g}$ in 2000 and $10 \ \mu\text{g/g}$ in 2018 (Guo, 2013). The standard value of Mn was 0.018 511 g/L in 2000 and only 0.002 g/L in 2018 (Li, 2016). The similar trend could also be found 512 in the standard of diesel in China (Zhang et al., 2009a). All these changes in the oil 513 514 standard will definitely cause the evolution of source profiles of vehicle exhaust. With the government's request to stop producing, selling and using leaded gasoline, the fraction of 515 516 Pb in vehicle emissions decreased significantly. In 2005, the fraction of Pb in motor vehicle emissions dropped significantly as compared with 1985 (Dai et al., 1986;Han et 517

al., 2009). The fraction of Mn is also greatly reduced after 2000 (Bi et al., 2007;Han et al., 2009). Similarly, the fraction of SO_4^{2-} in vehicle emissions also showed a significantly decreasing trend since 2000, indicating a causal relationship with the reduction of sulfur in the car-used gasoline in China.



522

Figure 10. Time series of Mn, Pb and SO₄²⁻ of the particulate matters emitted from vehicles
obtained. Data were collected from the source library of Nankai University, Dai et al. (1986),
Zhang et al. (2000), Bi et al. (2007), Han et al. (2009), Zhang et al. (2009), Guo et al. (2013), Li et
al. (2016).

By comparing the main components of on-road vehicles $PM_{2.5}$ source profiles derived from local studies and EPA SPECIATE database, Xia et al. (2017) found that both the source profiles of motor vehicles in China and the United States were dominated by OC and EC, but with different proportions. In America, the gasoline, ethanol and methanol are added as the aerator, while such oxygen content of gasoline in China is smaller than America, which is an important reason for the difference in the OC content in the spectrums at home and abroad (Xia et al., 2017). In China, the fraction of SO_4^{2-} is 2.4 times higher than that of foreign motor vehicles (Wang et al., 2015; Xia et al., 2017), which may be related to the high<u>er</u> sulfur content in the fuels (Guo et al., 2013; Li et al., 2016).

537 **2.2.4 Fugitive dust**

Fugitive dust is founded to be one of the major sources of urban particulate matter (Chow 538 et al., 2003; Kong et al., 2011; Cao et al., 2012; Zhu et al., 2018), especially in northern 539 cities in China with dry climate and limited precipitation (Shen et al., 2016; Cao et al., 540 541 2008). Urban fugitive dust is not only influenced by soil properties with geographic 542 locations, it is actually but also the mixture of various dust-related sources. Therefore, fugitive dust is often referred to soil dust, road dust, construction dust (Doskey et al., 1999; 543 544 Kong et al., 2014). Fugitive dust samples were generally collected by using resuspension chamber. 545

As shown in Fig. 11, the primary species in soil dust are Si, Al, Ca, with mass fractions 546 547 ranged from 0.0500 to 0.2010 g/g. Si is the predominant species among the detected elements, followed by Al, Fe, Na and Mg. The main chemical components of road dust 548 are Si, OC and Ca, with fractions ranged from 0.0712 to 0.0855 g/g. Al, Fe and SO_4^{2-} are 549 the relatively lower species (less than 0.0005 g/g) in the chemical profiles of road dust. Si, 550 551 Ca, Al and Fe are all crustal elements, indicating that the soil dust has a greater impact on the composition of road dust. It also shows that OC and SO4²⁻ in the source profiles of 552 road dust are higher than that of soil dust, indicating that the road dust is also affected by 553 vehicle emissions or coal combustion and other anthropogenic sources (Ma et al., 2015). 554

In general, the total water–soluble ions accounts for 0.0248-0.0648 g/g of fugitive dust, suggested suggesting that insoluble matter is not the main component of fugitive dust.

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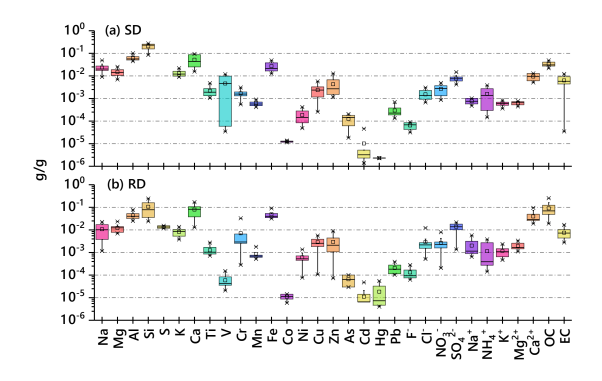


Figure 11. Characteristics of chemical profiles for particulate matter emitted from fugitive dust.
SD and RD denote soil dust and road dust, respectively. Data were collected from the source
library of Nankai University.

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Many studies have demonstrated that the ratios of different chemical components can be used as markers for fugitive dust (Alfaro et al., 2003; Arimoto et al., 2004). Kong et al. (2011) found that the Ca/Al ratio of paving road dust affected by construction activities was significantly different from that of soil dust. Zhang et al. (2014) reported that the heavy metals like Zn and Pb capable of being the tracers of urban fugitive dust, as they found Zn/Al and Pb/Al ratios in urban fugitive dust were 1.5 to 5 times those in desert,

Gobi, and loess soil samples. The NO_3^{-}/SO_4^{2-} ratio has been used to compare the relative importance of stationary sources vs mobile sources. Much high<u>er</u> NO_3^{-}/SO_4^{2-} ratio of road dust in Hong Kong has been reported by Ho et al. (2003), revealing the more important impact of vehicle emissions on the chemical composition of road dust as compared to coal combustion.

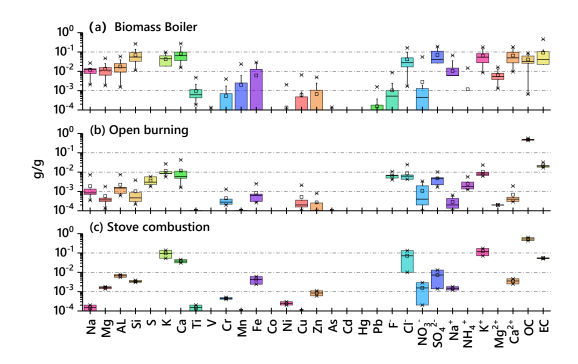
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575 2.2.5 Biomass burning

Traditionally, China is an agricultural-based country in the world (Bi et al., 2007). As an 576 577 effective way to eliminate plant residues, direct combustion by open burning and in domestic stove are the predominant and popular practice during the harvest seasons 578 (Andreae and Merlet, 2001; Ni et al., 2017; Cheng et al., 2013; Li et al., 2014b; Streets et 579 580 al., 2003). However, it releases a lot of pollutants into ambient air, and consequently impactsing air quality, human health and climate (Yao et al., 2017; Chen et al., 2017a). 581 Biofuel burned with boilers is also an important subtype of biomass burning (Tian et al., 582 583 2017). The wheat straw, corn stalks and rice straw represent 80% of the agricultural combustion in China (Ni et al., 2017), and there are also firewood, soybean and rape. The 584 biofuel types, sampling procedures and burning conditions result in great differences in 585 the levels and chemical properties of PM emissions from biomass burning (Tian et al., 586 2017; Vicente and Alves, 2018). 587

Biomass are usually burned in three ways in China, that is open burning (OB), residential stove combustion (RSC), and biofuel boiler burning (BBB). At present, there are two popular ways in the measurements of biomass burning: field combustion experiment

(FCE) and laboratory combustion simulation (LCS) (Hays et al., 2005; Li et al., 2014a; 591 Sanchis et al., 2014; De Zarate et al., 2000). Fig. 12 summarizes the biomass burning 592 profiles of PM_{2.5} from three burning styles obtained in China. The samples of biomass 593 boiler exhaust are obtained by resuspension sampling method. The main components in 594 the profiles of biomass burning are OC, EC, K⁺, Cl⁻, K and Ca (Fig. 12). The fraction of 595 EC is 4.2 times higher in BBB than RSC, which is potentially due to the uneven mixing of 596 the air in the biomass boiler that easy to make straw burning in anaerobic condition (Tian 597 et al., 2017). The high EC emissions can also happen if high temperature flaming burning 598 599 condition was dominant in the BBB. The oxygen content is relatively sufficient in OB, which leads to relatively higher OC emission. The fraction of Ca was higher in BBB 600 exhaust than OB (Fig. 12). For specific components emissions from the biomass burning, 601 602 EC emissions from firewood combustion was the highest, which is likely due to the high combustion temperature and flaming dominance burning condition, and the higher content 603 of lignin in wood (Tang et al., 2014), since lignin facilitates the formation of black carbon 604 605 (Wiinikka and Gebart, 2005).



607

Figure 12. Major chemical compositions of PM_{2.5} source profiles of biomass burning. Data were
 collected from the source library of Nankai University.

610

Chen et al. (2007) investigated the particulate emissions from wildland fuels burning in a laboratory combustion facility in the U.S., and found the percentage of total carbon (TC) of PM was $63.7\% \sim 100\%$, which was higher than that in China ($4.9\% \sim 68\%$). K ($0.4\% \sim 23.7\%$), Cl ($0.1\% \sim 9.6\%$) and S ($0.1\% \sim 2.9\%$) were important part of the remaining PM mass in the U.S, which is different from China due to the different biomass categories and combustion processes.

617

618 2.2.6 Cooking emissions

With the economic growing, the cooking styles and types of food ingredients on the table are becoming increasingly diverse. Since the 1990s, the variety of ingredients and cooking styles was also influenced by the foreign food culture. As China is famous for its 622 abundance of food culture, the cooking styles are varied with different regions, even in different cities. Thus, cooking is undoubtedly an important local source of ambient 623 particles. Given that there is no ubiquitous source profile for cooking emission, it is better 624 to measure source profile of cooking emissions in real-world in the study area. As one of 625 the essential cooking ingredients in the food and beverage industry, the types of edible 626 oils are changing in recent years (Pei et al., 2016). Soybean oil, rapeseed oil and peanut 627 oil are common edible oils for public dining in China. Due to changes in consumer 628 demand, other types of edible oils, such as olive oil, camellia oil and flaxseed oil, have 629 630 also been increasingly welcomed by the catering industry. Furthermore, Chinese-style cooking is characterized by high temperature stir-frying that releasing much more organic 631 matter than the cooking style of western food (Zhao et al., 2007b). 632

The chemical nature of $PM_{2.5}$ emitted from commercial cooking were investigated in many studies, with source profiles varied greatly with different factors such as cooking styles, cooking foods, seed oils and fuel (He et al., 2004;Zhao et al., 2007b;Hou et al., 2008b;Zhao et al., 2015b;Pei et al., 2016). Robinson et al. (2006) found that the contribution of cooking emission to OC in $PM_{2.5}$ calculated by chemical mass balance model using different source profiles yielded a difference by a factor of more than nine.

Previous studies founded that organic matter accounted for 66.9 % of the total suspended particles (TSP) mass emitted from cooking activities (Zhao et al., 2015b). OC is the major constituent and accounted for 36.2%~42.9% of the total mass, while the fraction of EC is much lower. Several water-soluble ions measured in the fine particles presented a relatively lower but a noticeable percentages, which made up of about 9.1%~17.5% of the total $PM_{2.5}$ mass (Anwar et al., 2004). Inorganic elements are found to be 7.3%~12.0% of the total $PM_{2.5}$ mass due to their greater presence in cooking oil and raw materials (He et al., 2004).

Fig. 13 shows the PM_{2.5} chemical profiles of cooking emissions including hot pot, 647 648 Chinese restaurant, barbecue and cafeteria (See and Balasubramanian, 2006; Taner et al., 2013; Zhang et al., 2017a). For elements, on average, the most abundant elements in 649 cooking profiles is Al, followed by Ca and Fe. The high levels of Ca and Fe are probably 650 651 emitted from raw material and cooking utensils (See and Balasubramanian, 2006; Taner et 652 al., 2013). The high level of Cr, originated from stainless steel grills, was observed in a barbeque restaurant (Taner et al., 2013). Overall, OC is the most abundant species in the 653 profiles of cooking emissions. 654

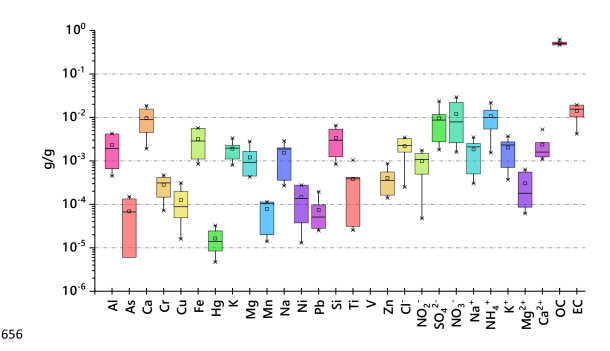
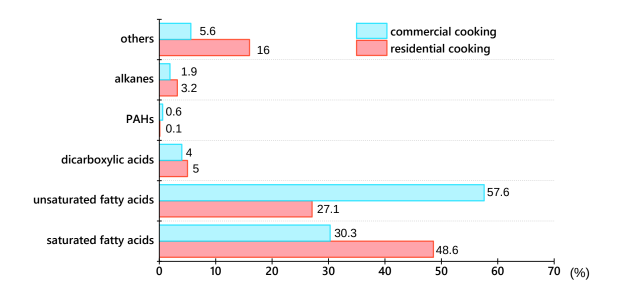


Figure 13. PM_{2.5} Chemical profiles of cooking emissions. Data from the source library of Nankai
University, Zhang et al. (2017), See et al. (2006) and Taner et al. (2013) were counted.

Organic matter (OM) is the predominant species in PM_{2.5} emitted from cooking activities 660 (He et al., 2004; Hou et al., 2008a; Pei et al., 2016). Many organic compounds, including 661 n-alkanes, dicarboxylic acids, polycyclic aromatic hydrocarbons (PAHs), saturated fatty 662 663 acids and unsaturated fatty acids, were quantified in the above mentioned studies. Fig. 14 shows the fractions of main organic compounds in the quantified OM emission from 664 residential cooking (Zhao et al., 2015b) and commercial cooking (Pei et al., 2016). 665 Among the quantified organic compounds, the predominant species is unsaturated fatty 666 acids (49.4%-77.8%), followed by saturated fatty acids (25.1%-43.8%). 667

668



669

Figure 14. Proportions of major organic compounds in quantified OM emission from commercial

671 cooking (Pei et al., 2016) and residential cooking (Zhao et al., 2015b).

672

In addition, except for biomass burning, many studies have reported that the levoglucosan

was also founded in the emissions from residential coal combustion (Yan et al., 2017) and 674 a variety of Chinese and western cooking styles (He et al., 2004; Zhao et al., 2007a, b). 675 Pei et al. (2016) also found Italian cooking style released the smallest amount of 676 monosaccharide anhydrides and the largest amount of cholesterol due to the lower ratio of 677 vegetables to meat used in the Italian cooking than Chinese cooking materials. Malay 678 cooking released higher PAHs concentrations than the Chinese and India methods (See et 679 al., 2006). Deep frying emitted more PAHs than other cooking methods because of the 680 higher temperature and more oil used during cooking. As far as we know, molecular 681 682 markers used for cooking included levoglucosan, galactosan and cholesterol (He et al., 2004; Zhao et al., 2007a, b) while cholesterol can be regarded as the best marker for meat 683 cooking (Schauer et al., 1999; Schauer and Cass, 2000; Schauer et al., 2002). 684

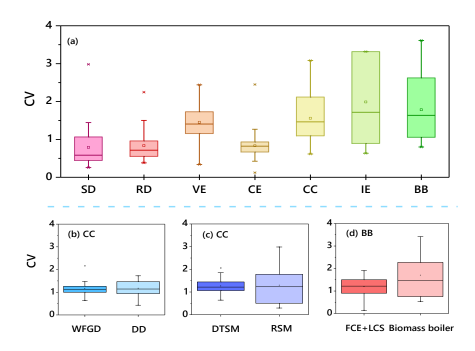
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686 2.3 Statistical analysis of the source categories

The chemical profile of a given source category is always established from profiles of several similar sources belonging to this category. Non-negligible uncertainties would be introduced in this process. To evaluate such uncertainties, the coefficient of variation (CV, the standard deviation divided by the mean) is used in this section to further characterize the homogeneity of sources within the same source category (Fig. 15).

The values of CV above three (Pernigotti et al., 2016) are observed in coal combustion, industry emissions and biomass burning, indicating these source profiles shows a great variations due to the effects of their influencing factors as described in above sections. The profiles of road dust and soil dust showed a less variations with stable chemical characteristics among the different profiles in the same category. However, the responses
of source profiles to various impact factors are different (Fig. 15(ab)-(ed)). For example,
the sampling methods have a notable effect on the source profile of coal combustion (the
variation of coal combustion source profiles obtained by resuspension sampling is greater
than that by DTSM), while the desulfurization methods have little-smaller impact.

701 Since source profiles owned local characteristic, it is important and necessary to establish and update local source profiles to reveal the real situation of source emissions (Zhang et 702 al., 2017b; Zhu et al., 2018). However, local source profiles are not always available in 703 704 some developing areas in the case of limited funds and instruments. According to the above statistical results, it can be inferred that the profiles of road dust and soil dust could 705 be references for the cities in China without such local profiles, while it is necessary to 706 707 establish the local profiles of the industrial emissions, vehicle emissions, coal combustion, and biomass burning. 708



710 **Figure 15.** Coefficients of variation calculated for each source category. SD denotes soil dust, RD

denotes road dust, VE denotes vehicle emissions, CE denotes cooking emissions, CC denotes coal
combustion, IE denotes industrial emissions, BB denotes biomass burning, WFGD denotes wet
flue gas desulfurization, DD denotes dry desulfurization, DTSM denotes the dilution tunnel
sampling method, RSM denotes resuspension sampling method, FCE denotes field combustion
experiment, LCS denotes laboratory combustion simulation.

In order to investigate the similarity of the real-world measured source profiles with 717 homogeneous chemical signature, cluster analysis was applied to the collected data by 718 719 using the package R pvclust (Suzuki and Shimodaira, 2006; Pernigotti et al., 2016). The significance test was performed with resampling the data via bootstrap method. This 720 function is expected to assign each cluster an approximated unbiased (AU) p-value by 721 722 hierarchic clustering (Shimodaira, 2002). Details on the operation steps of this method 723 areis discussed earlier by Pernigotti et al. (2016). The input source profiles involved in the cluster calculation must contain more than two common chemical species, including 724 725 elements, ions and OC/EC. In order to reduce the interference of different particle sizes, we used 226 source profiles of PM_{2.5} for the calculation. The result of cluster analysis and 726 additional information of the source profiles are shown in Fig. 16 and Table S1. As shown 727 in Fig. 16, clusters are marked if the AU p-value ≥ 90 (values were reported in red). It 728 729 shows that the source profiles are divided into (1) biomass burning, (2) and (4) coal combustion, (3) industrial emission, (5) soil dust, (6) road dust, (7) cooking emissions and 730 731 (8) vehicle emissions. These subjectively measured profiles are successfully classified by objectively method based on their chemical nature, though there are some different 732

733	sources mixed up (Fig. 16). This result indicates that the routine measured components
734	are not enough to distinguish all the source categories when in the case of the chemically
735	co-linear sources exist. Both the source profiles of cooking and vehicle emissions are
736	characterized by high OC, which makes them easy to be identified as the same source
737	type. The chemical collinearity of the source composition between coal combustion and
738	dust also makes it difficult to be distinguished. To solve the chemical co-linearity problem
739	between sources, more specific tracers, especially organics should be further explored.

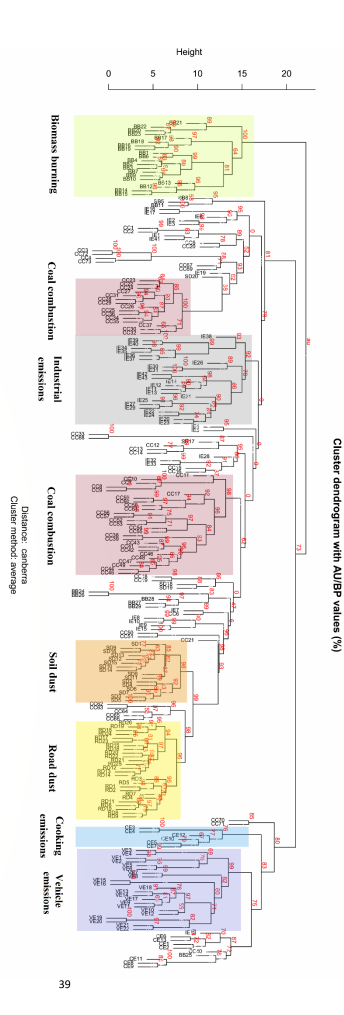


Figure 16. Result of cluster analysis on the profiles. AU p-values are reported in red as %.

741

742 **3.** Conclusion

743 The chemical profiles of main sources of particulate matter have been established in China since the 1980s. With the development of sampling and analysis techniques, the 744 dataset of source profiles has been gradually enlarged and could to able to reflect the real 745 emissions of the sources to the ambient air. A total of 456 published source profiles, 746 coupled with the database of source profiles (2870 profiles) founded by Nankai University 747 are reviewed in this work. Six source categories include coal combustion, industrial 748 process emissions, vehicle emissions, fugitive dust, biomass burning and cooking 749 emissions are investigated to characterize sources in chemical nature and explore the main 750 factors that influencing the chemical composition. 751

In general, coal combustion is the most complicated source in all source categories as it is 752 influenced by many factors from the fuel combustion processes to pollution-controlling 753 processes. Sulfate is the predominant species of fine particles emissions from coal 754 combustion stationary sources equipped with wet flue gas desulfurization device, and is 755 also the second largest species in fine particles emissions from residential coal 756 combustion. The source profiles of industrial emissions are mainly determined by the 757 components of the industrial products and its pollution-controlling techniques. With the 758 changing standards of gasoline and diesel oil since the 1980s, Pb and Mn are no longer 759 the tracers of emission from the gasoline vehicles. OC and EC are always the dominant 760 species of vehicle emissions from the 1980s despite the changing standards. The profiles 761 762 of the fugitive dust including the road dust and soil dust are characterized by the high levels of crustal elements, such as Si, Al and Ca. The profiles of the biomass burning are 763

determined by the biomass categories and the different combustion phases (smoldering and flaming), with K^+ and levoglucosan to be the common tracers. As for cooking emissions, the source profiles of the emissions from the different cooking types were all dominated by OC.

The result of uncertainty analysis showed that the relatively large variation in the source 768 profiles of industry emissions, vehicle emissions, coal combustion and biomass burning, 769 calling for establishing the local profiles for these sources due to their high uncertainties. 770 While the profiles of road dust and soil dust present a less variation, suggesting that the 771 772 profiles of these sources could be referenced for the cities in China when the local profiles are not available. Since source profiles owned local characteristic, it is important and 773 necessary to establish local source profiles to reveal the real situation of source emissions 774 775 and update it immediately.

The result of cluster analysis on the routine measured species of source profiles suggested 776 that some of the sources are difficult to be distinguished (cooking emissions vs vehicle 777 778 emissions), indicating that more chemical tracers, such as the isotopes and organic compounds, should be further explored in the source profiles to reduce the collinearity 779 among different source profiles. Current source profile database is still missing some 780 important source categories that have significant impacts on the air quality, and lacking 781 sufficient source profiles, especially for the industrial emissions, such as the glass melt 782 kiln, nonferrous metal smelting, bricks and tiles kiln. Thus, specific focus should be 783 784 placed on these important but overlooked sources in the future.

785

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