2	Insight into PM _{2.5} Sources by Applying Positive
3	Matrix Factorization (PMF) at an Urban and
4	Rural Site of Beijing
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31 ABSTRACT

32 This study presents the source apportionment of PM_{2.5} performed by PMF on data collected at an urban (Institute of Atmospheric Physics - IAP) and a rural site (Pinggu-PG) in Beijing as part of the 33 Atmospheric Pollution and Human Health in a Chinese megacity (APHH-Beijing) field campaigns. 34 The campaigns were carried out from 9th November to 11th December 2016 and 22nd May to 24th June 35 2017. The PMF included both organic and inorganic species, and a seven-factor output provided the 36 most reasonable solution for the PM_{2.5} source apportionment. These factors are interpreted to be 37 traffic emissions, biomass burning, road dust, soil dust, coal combustion, oil combustion and 38 secondary inorganics. Major contributors to PM_{2.5} mass were secondary inorganics (IAP: 22%; PG: 39 24%), biomass burning (IAP: 36%; PG: 30%), and coal combustion (IAP: 20%; PG: 21%) sources 40 during the winter period at both sites. Secondary inorganics (48%), road dust (20%) and coal 41 combustion (17%) showed the highest contribution during summer at PG, while PM_{2.5} particles were 42 mainly composed of soil dust (35%) and secondary inorganics (40%) at IAP. Despite this, factors that 43 were resolved based on metal signatures were not fully resolved and indicate a mixing of two or more 44 sources. PMF results were also compared with sources resolved from another receptor model (i.e. 45 46 CMB) and PMF performed on other measurements (i.e. online and offline aerosol mass spectrometry (AMS)) and showed a good agreement for some but not all sources. The biomass burning factor in 47 PMF may contain aged aerosols as a good correlation was observed between biomass burning and 48 oxygenated fractions ($r^2=0.6-0.7$) from AMS. The PMF failed to resolve some sources identified by 49 the CMB and AMS, and appears to overestimate the dust sources. A comparison with earlier PMF 50 source apportionment studies from the Beijing area highlights the very divergent findings from 51 application of this method. 52

53 Key words: Source apportionment; PM_{2.5}; Beijing; PMF; CMB; online AMS; offline AMS

55 1. INTRODUCTION

56 Atmospheric particulate matter (PM) is composed of various chemical components and can affect air quality (and consequently human health), visibility, and ecosystems (Boucher et al., 2013; Heal et al., 57 2012). Through absorption and scattering of solar radiation and by affecting clouds, PM also have a 58 major impact on the climate, and thus the hydrological cycle. PM with an aerodynamic diameter less 59 than 2.5 μ m (PM_{2.5}) is given special attention due to its adverse effects on human health as it can 60 penetrate deep into human lungs when inhaled. Several recent studies have indicated that many 61 adverse health outcomes, such as respiratory and cardiovascular morbidity and mortality, are related 62 to long-term exposure to PM (Lu et al., 2021; Wang et al., 2016; Xing et al., 2016; Xie et al., 2019). 63 In addition, over a million premature deaths per year are reported in China due to poor air quality 64 (GBD MAPS Working Group, 2016). Beijing, the capital city of China, is a megacity with 65 approximately 21 million inhabitants that are regularly exposed to severe haze events. For example, 66 77 pollution episodes (defined as two or more consecutive days where the average PM_{2.5} 67 concentration exceeds 75 µg m⁻³) were observed between April 2013 to March 2015 (Batterman et 68 al., 2016). PM_{2.5} concentrations have reached 1,000 µg m⁻³ in some heavily polluted areas of Beijing 69 70 (Ji et al., 2014). In addition, a study compared the number of cases of acute cardiovascular, cerebrovascular, and respiratory diseases in the Beijing Emergency Center and haze data from Beijing 71 Observatory between 2006 and 2013. Their results showed a rising trend, highlighting the average 72 number of cases per day for all three diseases was higher on hazy days than on non-hazy days (Zhang 73 et al., 2015). Therefore, major control measures were implemented to reduce PM_{2.5} pollution in 74 Beijing (Vu et al., 2019). Recently, one-third of Chinese cities in 2020 were kept under lockdown to 75 prevent the transmission of COVID-19 virus, which strictly curtailed personal mobility and economic 76 activities. The lockdown led to an improvement in air quality and managed to bring down the levels 77 of PM_{2.5}. Despite these improvements, PM_{2.5} concentrations during the lockdown periods remained 78 higher than the World Health Organization recommendations, suggesting much further effort is 79

needed (He et al., 2020; Le et al., 2020; Shi et al., 2021). A quantitative source apportionment
provides key information to support such efforts.

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Receptor models are widely used for source apportionment of PM_{2.5}. These methods include positive 83 matrix factorization (PMF) (Paatero, 1997; Paatero and Tapper, 1994), principal component analysis 84 (PCA) (Lee et al., 2011), chemical mass balance (CMB) (Watson et al., 1990), and UNMIX (Herrera 85 Murillo et al., 2012). Among these methods, PMF is a widely used multivariate method that can 86 resolve the dominant positive factors without prior knowledge of sources. Previous PMF studies, 87 88 based on high resolution Aerosol Mass Spectrometer data, have provided valuable information on the sources of PM in urban Beijing and its surrounding areas (Zhang et al., 2015; Huang et al., 2010; Sun 89 et al., 2010; Sun et al., 2013; Zhang et al., 2013; Zhang et al., 2014; Zhang et al., 2017; Zhang et al., 90 91 2016; Hu et al., 2016; Qiu et al., 2019). However, the factors that influence haze formation and related 92 sources remain unclear due to its inherent complexity (Tie et al., 2017; Sun et al., 2014). Filter-based PMF studies provide a valuable tool for identifying sources of airborne particles, by utilising size-93 94 resolved chemical information (Li et al., 2019; Ma et al., 2017a; Tian et al., 2016; Yu et al., 2013; Song et al., 2007; Song et al., 2006). These source apportionment studies have predominantly used 95 OC (organic carbon), EC (elemental carbon), water soluble ions and metals as the input data matrix 96 to explore the co-variances between species and their associated sources, but to the best of our 97 knowledge, the use of organic markers in PMF has not been explored extensively in Beijing. The use 98 99 of organic molecular markers in PMF has enhanced our understanding of the PM fraction as they can be source specific (Shrivastava et al., 2007; Jaeckels et al., 2007; Zhang et al., 2009; Wang et al., 100 2012; Srimuruganandam and Shiva Nagendra, 2012; Schembari et al., 2014; Laing et al., 2015; 101 102 Waked et al., 2014; Srivastava et al., 2018) and could potentially offer a clearer link between factors and sources. 103

This study presents the results obtained from the PMF model applied to a filter-based dataset collected 105 in the Beijing metropolitan area at two sites, urban and rural. The study provides source 106 apportionment results from both urban and rural locations in Beijing including their temporal and 107 108 spatial variations. In addition, the study also presents a short summary of previously published filterbased studies conducted in the Beijing metropolitan area and their major outcomes. A comparison of 109 110 the present PMF results was also made with other source apportionment approaches or applications of PMF to other datasets, with an aim to discuss the existing PM sources in the Beijing metropolitan 111 area, including focusing on the strengths and weaknesses of the source apportionment approach 112 employed. 113

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115 **2. METHODOLOGY**

Details about the sampling site, measurements, sample collection and analytical procedures are reported elsewhere (Shi et al., 2019; Xu et al., 2021; Wu et al., 2020), and hence only the essential information is presented in this section.

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120 **2.1 Sampling Site and Sample Collection**

The PM_{2.5} sampling was conducted simultaneously at the urban and rural sites from 9th November to 121 12th December 2016 and 22nd May to 24th June 2017 as part of the Atmospheric Pollution and Human 122 Health in a Chinese megacity (APHH-Beijing) field campaigns (Shi et al., 2019) (Figure S1). The 123 124 urban sampling site (116.39E, 39.98N) - the Institute of Atmospheric Physics (IAP) of the Chinese Academy of Sciences in Beijing, represents typical condition of central Beijing, there are various 125 roads nearby including a highway road approximately 200 m away. The rural Pinggu site (PG) 126 (40.17N, 117.05E) is located in Xibaidian village. This site is approximately 60 km to the north-east 127 of Beijing city centre and about 4 km north-west of the Pinggu town centre. The site is surrounded 128

by trees and farmland. In addition, residents mainly use coal and biomass for heating and cooking inindividual homes.

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132 24-hour $PM_{2.5}$ samples were collected every day on pre-baked quartz filters (Pallflex, 8×10 inch) and 133 47 mm PTFE filters (flow rate of 15.0 L min⁻¹) using high volume (Tisch, USA, flow rate of 1.1 m³ 134 min⁻¹) and medium volume (Thermo Scientific Partisol 2025i) air sampler. Field blanks were also 135 collected during the sampling campaign at both sites. The quartz filters were then analyzed for organic 136 tracers, OC/EC and ion species. PTFE filters were used for the determination of PM_{2.5} mass and 137 metals. Details on preparation and conservation of these filter samples have already been reported 138 elsewhere (Wu et al., 2020; Xu et al., 2021).

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Real time composition of non-refractory PM₁ particles (NR-PM₁) was measured using an Aerodyne aerosol mass spectrometer (AMS) at a time resolution of 2.5 min. The operational details on the AMS measurements have been given elsewhere (Xu et al., 2019). In addition, the measurements of gaseous species such as O₃, CO, NO, NO₂ and SO₂ were performed using gas analyzers. The meteorological parameters including temperature (T), relative humidity (RH), wind speed (WS), and wind direction (WD) were also measured at both sites.

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147 2.2 Analytical Procedure

In all 62 and 72 chemical species were quantified in each PM_{2.5} sample from IAP and PG, respectively. This included EC/ OC, 36 organic tracers, 7 major inorganic ions (Na⁺, K⁺, Ca²⁺, NH4⁺, Cl⁻, NO₃⁻ and SO₄²⁻) and 17 metallic elements (V, Cr, Co, Mn, Ni, Cu, Zn, As, Br, Sr, Ag, Cd, Sn, Sb, Ba, Hg and Pb) at IAP. Similarly, the identified species at PG included EC/OC, 51 organic tracers, 7 major inorganic ions and 12 metallic elements (V, Cr, Co, Mn, Ni, Cu, Zn, As, Sr, Sb, Ba, and Pb). EC and OC measurements were performed using a Sunset lab analyser (model RT-4) and DRI multiwavelength thermal-optical carbon (model 2015) analyser based on the EUSAAR2 (European
Supersites for Atmospheric Aerosol Research) transmittance protocol at both sites, IAP and PG,
respectively, following the procedure explained by Paraskevopoulou et al. (2014). Major inorganic
ions and metallic elements were analysed using an ion chromatograph (Dionex, Sunnyvale, CA,
USA) and Inductively coupled plasma-mass spectrometer (ICP-MS) at both sites, respectively. Major
crustal elements including Al, Si, Ca, Ti and Fe were determined by X-ray fluorescence spectrometer
(XRF).

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Organic tracers at IAP included n-alkanes 11 C₂₄-C₃₄, 2 hopanes, 17 PAHs, 3 anhydrous sugars 162 (levoglucosan, mannosan, galactosan), 2 fatty acids (palmitic acid, stearic acid) and cholesterol. 163 These organic tracers were analysed by gas chromatography mass spectrometry (GC/MS, Agilent 164 7890A GC plus 5975C mass-selective detector) coupled with a DB-5MS column (30 m \times 0.25 mm \times 165 0.25 µm) following the protocol explained in Xu et al. (2020). At PG, organic tracers were analysed 166 based on the method reported by Wu et al. (2020) using GC/MS (Agilent GC-6890N plus MSD-167 5973N) coupled with a HP-5MS column (30 m \times 0.25 mm \times 0.25 μ m). This included quantification 168 of similar species (12 n-alkanes C24-C35, 9 hopanes, 22 PAHs, 3 anhydrous sugars (levoglucosan, 169 mannosan, galactosan), 4 fatty acids (palmitic acid, stearic acid, linoleic acid, oleic acid) and 170 cholesterol) with few additional ones. Recoveries for the identified organic tracers ranged from 70-171 100% and 80-110%, at IAP and PG, respectively. Field blank were also analysed as part of quality 172 control and demonstrated very low contamination (<5%). 173

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175 In addition, one or two punches of $PM_{2.5}$ filter sample were also analysed offline using AMS to 176 investigate the water-soluble OA (WSOA) mass spectra following the procedure explained 177 previously (Qiu et al., 2019).

179 2.4 Positive Matrix Factorization

Detailed information on the receptor modelling methods used within this study can be found elsewhere (Paatero and Tapper, 1994; Hopke, 2016). Positive matrix factorization (PMF) is a multivariate factor analysis tool and based on a weighted least square fit, where the weights are derived from the analytical uncertainty. The best model solution was obtained by minimizing residuals obtained between modeled and observed input species concentrations Estimation of analytical uncertainties for the filter-based measurements was calculated using Eq. (1) (Polissar et al., 1998).

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$$\sigma_{ij} = \begin{cases} \frac{5}{6} LD_j & \text{if } X_{ij} < LD_j \\ \sqrt{(0.5 * LD_j)^2 + (EF_j X_{ij})^2} & \text{if } X_{ij} \ge LD_j \end{cases}$$
Eq. (1)

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where LD_j is the detection limit for compound *j* and EF_j is the error fraction for the compound *j*. The detection limits of all compounds used in the PMF model is given in Table S1 (SI). The U.S. Environmental Protection Agency (US-EPA) PMF 5.0 software was used in this work to perform the source apportionment.

194

Selection of the input data. The selection of species used as input data for the PMF analysis is 195 important and can significantly influence the model results (Lim et al., 2010). The following set of 196 197 criteria were used for the selection of the input species: signal to noise ratio (S/N) (Paatero and Hopke, 2003), major PM chemical species, compounds with maximum data points above the detection limit 198 and those being considered as specific markers of a given source (e.g., levoglucosan, picene) (Oros 199 and Simoneit, 2000; Simoneit, 1999) were selected. These steps were taken to limit the input data 200 201 matrix according to the total number of samples (n=133); some species were also not included if they belonged to a single source and correlated with another marker of this source. A total of 31 species 202 were used in the model (PM_{2.5}, OC, EC, K^+ , Na⁺, Ca²⁺, NH₄⁺, NO₃⁻, SO₄²⁻, Cl⁻, Ti, V, Mn, Ni, Zn, Pb, 203

Cu, Fe, Al, C26, C29, C31, $17\alpha(H)$ -22,29,30-trisnorhopane, $17\beta(H)$, $21\alpha(H)$ -30-norhopane,chrysene, benzo(b)fluoranthene, benzo(a)pyrene, picene, corene, levoglucosan, and stearic acid). The concentration of PM_{2.5} was included as a total variable in the model (with large uncertainties) to directly determine the source contributions to the daily mass concentrations.

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Selection of the final solution. As normally recommended, a detailed evaluation of factor profiles,
 temporal trends, fractional contribution of major species to each factor and correlations with external
 tracers, were investigated carefully to select the appropriate number of factors.

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A few constraints were also applied to the base run to obtain clearer chemical source profiles in the 213 final solution. The general framework for applying constraints to PMF solutions has already been 214 discussed elsewhere (Amato et al., 2009; Amato and Hopke, 2012). The changes in the Q values were 215 216 considered here as a diagnostic parameter to provide insight into the rotation of factors. All model runs were carefully monitored by examining the Q values obtained in the robust mode. To limit 217 218 change in the Q-value, only "soft pulling" constraints were applied. The change in the Q-robust was < 1%, which is acceptable as per PMF guidelines (< 5%) (Norris et al., 2014). Finally, three criteria 219 were used to select the optimal solution, including correlation coefficient (r) between the measured 220 221 and modelled species, bootstrap and t-test (two-tailed paired t-test) performed on the base and constraint runs, as explained previously (Srivastava et al., 2018). 222

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224 2.5 Back trajectories and Geographical Origins

The geographical origin of selected identified sources and pollutants was investigated using concentration-weighted trajectory (CWT), non-parametric wind regression (NWR), and cluster analysis methods. NWR combines ambient concentrations with co-located measurements of wind direction and speed and highlights wind sectors that are associated with high measured concentrations (Henry et al., 2009). The general principle is to smooth the data over a fine grid, so that a weighted

concentration could be estimated by any wind direction (ϕ) /wind speed (v) couple, where the 230 weighing coefficients are determined through Gaussian-like functions. CWT and cluster analysis 231 assess the potential transport of pollution over large geographical scale (Polissar et al., 2001). These 232 233 approaches combine atmospheric concentrations measured at the receptor site with back trajectories and residence time information and help to geographically evaluate air parcels responsible for high 234 235 concentrations. For this purpose, hourly 24-h back trajectories arriving at 200 m above sea level were calculated from the PC-based version of HYSPLIT v4.1 (Stein et al., 2015; Draxler, 1999). NWR, 236 CWT calculations and cluster analysis, were performed using the ZeFir Igor package (Petit et al., 237 2017). 238

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2.6 Other Receptor Modelling Approaches

Sources were also resolved at both sites separately using another receptor model known as chemical
mass balance (CMB) as well as PMF performed on high resolution AMS data collected at IAP. Details
on sources resolved using these approaches are reported elsewhere (Wu et al., 2020; Xu et al., 2021;
Sun et al., 2020).

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Briefly, CMB is based on a linear least squares approach and accounts for uncertainties in both, source 246 profiles and ambient measurements to apportion the sources of OC. The US EPA CMB8.2 software 247 was used for this purpose at both sites. The source profiles applied in the model were taken from local 248 studies to better represent the sources, including profiles of straw burning (Zhang et al., 2007), wood 249 burning (Wang et al. 2009), gasoline and diesel vehicles (Cai et al. 2017), industrial and residential 250 coal combustion (Zhang et al., 2008), and cooking (Zhao et al., 2015). Only the source profile for 251 vegetative detritus (Rogge et al. 1993; Wang et al., 2009) was not available from local studies. The 252 selected fitting species included EC, anhydrous sugar (levoglucosan), fatty acids, PAHs, hopanes and 253 alkanes 254

255 **3. RESULTS AND DISCUSSION**

256 3.1 Overview on PM Sources in Beijing based on the Current Source Apportionment Study

A seven-factor output provided the most reasonable solution for the PM_{2.5} source apportionment performed on the combined dataset from IAP and PG (Figures 1 and 2).

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Based on the factor profiles, we identified traffic emissions, biomass burning, road dust, soil dust, 260 coal combustion, oil combustion and secondary inorganics. For the same dataset, solutions with six 261 sources were less explanatory and some factors were mixed. Conversely, an increase in the number 262 of factors led to the split of meaningful factor profiles. In the final solution, the comparison of the 263 reconstructed PM_{2.5} contributions from all sources with measured PM_{2.5} concentrations for different 264 seasons at both sites showed good mass closure ($r^2 = 0.61-0.91$, slope = 0.99-1.12, p < 0.05, ODR 265 (orthogonal distance regression)). A low r^2 (0.61) value was observed for the summer period at IAP 266 (Figure S2). This may be due to the inability of PMF to model low concentrations observed for 267 sources such as biomass burning and coal combustion during the summer. In addition, most of the 268 species showed good agreement with measured concentrations (Table S2). Bootstrapping on the final 269 270 solution showed stable results with more than 95 out of 100 bootstrap mapped factors (Table S3). Finally, no significant difference (p>0.05) was observed in the factor chemical profiles between the 271 272 base and the constrained runs (Table S4).

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Overall, secondary inorganics, biomass burning, and coal combustion sources were the main contributors to the total PM_{2.5} mass during winter (Figure 3). These sources accounted for 22%, 36%, 20%, and 24%, 30%, 21% of PM_{2.5} mass at IAP and PG, respectively. Secondary inorganics, road dust and coal combustion showed the highest contribution during summer at PG, while PM_{2.5} particles were mainly composed of soil dust and secondary inorganics at IAP. Identified aerosol sources, factor profiles and temporal evolutions are discussed below. Note, PMF was carried out on the combined datasets and thus only provides a single set of factor profiles for both sites. Similar to previous studies (Li et al., 2019; Ma et al., 2017a; Tian et al., 2016; Yu et al., 2013; Liu et al., 2019; Zhang et al.,
2013), neither secondary organic aerosol nor cooking emissions were identified, and given the good
mass closure must be present within other source categories.

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Coal combustion. Coal combustion was identified based on it accounting for a high proportion of
PAHs (27-78%), especially picene (78%) as a specific marker of coal combustion (Oros and
Simoneit, 2000), together with significant amount of OC (45%) and EC (29%) (Figure 1). This factor
also made a substantial contribution to n-alkanes (28-58%), stearic acid (64%) and hopanes (53-56%),
as these compounds are also abundant in coal smoke (Bi et al., 2008; Zhang et al., 2008; Oros and
Simoneit, 2000; Guo et al., 2015).

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The coal combustion factor accounted for 20% of the PM mass (16.0 µg m⁻³) at the urban site IAP 292 during winter and followed typical seasonal variations. However, the contributions of this source to 293 PM_{2.5} mass were broadly similar (21% vs 17%, Figure 3) at PG during both seasons, while the average 294 concentrations were higher in winter than summer (19.4 μ g m⁻³ > 4.6 μ g m⁻³). Due to a lack of 295 infrastructure at the rural site PG, the residents still use coal for cooking and heating purposes at the 296 297 time of sampling (Shi et al., 2019). There is a reduction in coal usage for heating due to elevated temperatures in the summertime, leading to low levels of this factor. But the similar contribution at 298 the rural site could be linked to consistent cooking activities throughout the year (Figure 2) (Shi et 299 al., 2019; Tao et al., 2018). These results were in good agreement with previous observations reported 300 at the same urban site (18%) (Ma et al., 2017a; Tian et al., 2016). In addition, similar contributions 301 were also observed at other urban locations around Beijing (Wang et al., 2008; Liu et al., 2019). 302

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This factor also included significant contributions from levoglucosan (60%). Levoglucosan, a major pyrolysis product of cellulose, and has been proposed as molecular marker of biomass burning aerosols (Simoneit, 1999). A study conducted in China suggested that residential coal combustion can also contribute significantly to levoglucosan emissions, based on both source testing and ambient measurements (Yan et al., 2018). Therefore, it is expected that the contribution of levoglucosan is probably linked to residential coal use for cooking in this case. It is also possible that the high contribution of levoglucosan could also be linked to model bias as the PMF model only provides an average factor profiles for both sites irrespective of their nature (rural vs. urban) and different sampling periods (summer vs. winter).

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This was further supported as NWR and CWT analysis showed similar results, mostly dominated by a northerly flow during the winter period at both sites. High concentrations of this source and levoglucosan were observed at low wind speeds (Figure S3), indicating the significant role of local activities. Higher levels were observed at the rural site PG (19.4 μ g m⁻³ vs 16.0 μ g m⁻³ at the urban site). However, a south-westerly flow was dominant during summer and could be related to transport of air masses from the Hebei province where a large number of industries operate.

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Oil combustion. The oil combustion factor profile included high contributions to V (79%) and Ni 321 (70%) (see Figure 1). V and Ni are widely used markers for oil combustion in residential, commercial 322 and industrial applications (Viana et al., 2008; Mazzei et al., 2008; Pant et al., 2015; Huang et al., 323 2021). The V/Ni ratio obtained in this study was 0.9, close to the previously obtained ratio for residual 324 oil used in power plants (Swietlicki and Krejci, 1996). Results suggest this source might be attributed 325 to residual oil combustion linked to industrial activities as a large number of highly polluting 326 industries are still located in the Beijing neighbourhood (Li et al., 2019). CWT and NWR analysis 327 suggested the influence of regional transport at both sites, highlighting the dominance of south 328 westerly and south easterly flows during the winter and summer at both sites (Figure S4). 329

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The source did not show any seasonal pattern (Figure 2), and accounted for 2% (1.4 μ g m⁻³) and 6% (1.6 μ g m⁻³) at IAP, and 8% (7.1 μ g m⁻³) and 9% (2.1 μ g m⁻³) at PG of the PM_{2.5} mass during winter and summer, respectively (Figure 3). The contribution of this source to the PM mass was within the

similar range to the previous study conducted at the same urban site (contribution 4.7%) (Li et al.,
2019), which also found a high proportion of V attributed to the identified source.

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Biomass burning. The biomass burning factor was characterized by high contributions to Cl⁻ (74%), 337 338 $K^+(27\%)$ and levoglucosan (25%) (Figure 1). This factor also made significant contributions to PAHs (Chry (66%), B[b]F (66%), Cor (68%)) and followed a clear seasonal variation with a higher 339 contribution in winter (Figure 2). It accounted for 36 % (29.0 µg m⁻³) and 30% (27.3 µg m⁻³) of the 340 PM_{2.5} mass during the wintertime at IAP and PG, respectively (Figure 3), while the contribution 341 during the summertime was extremely low. This was expected due to elevated temperature during 342 the summer period and reduction in biomass burning activities. In addition, $NO_3^{-}(24\%)$, and NH_4^{+} 343 (24%) also contributed significantly to the biomass burning factor. Biomass burning is an important 344 natural source of NH₃ (Zhou et al., 2020) which rapidly reacts with HNO₃ to form NH₄NO₃ aerosols. 345 346 The presence of NH_4NO_3 aerosols in biomass burning plumes, has also been reported previously (Paulot et al., 2017; Zhao et al., 2020). 347

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It was unexpected to observe a low contribution of levoglucosan, a known biomass burning marker, 349 to this factor. However, model bias and the contribution of other relevant sources to levoglucosan 350 could cause such observations, as discussed above. K⁺ is also produced from the combustion of wood 351 lignin and has been used extensively as an inorganic tracer to apportion biomass burning contributions 352 to ambient aerosol (Zhang et al., 2010; Lee et al., 2008a). However, the contribution of K⁺ to this 353 factor was relatively low, possibly because K⁺ has also other sources, such as soil dust (Duvall et al., 354 2008). Cl⁻ can be emitted from both coal combustion and biomass burning, especially during the cold 355 356 period in Beijing (Sun et al., 2006). It is also important to note that high Cl⁻ levels observed in this factor could be associated with coal combustion as Cl⁻ has been used to represent coal combustion 357 activities in China (Wang et al., 2008). If we consider this, high Cl⁻ levels related to the coal 358 combustion factor should have also shown a significant contribution to PM mass during the 359

summertime at the rural site (PG) as residents near the rural site mostly use coal and biomass for cooking activities as discussed above, but they do not. Results suggest this factor can be attributed mainly to biomass burning aerosols in the Beijing metropolitan area, but some influence of coal combustion signals cannot be ignored. Back trajectory analysis also confirmed the local origin of this source during the wintertime at both sites (Figure S5).

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The source contribution reported in the present study was higher than that found in earlier studies in Beijing (11-20%) (Li et al., 2019; Ma et al., 2017a; Tian et al., 2016; Yu et al., 2013; Song et al., 2007; Song et al., 2006; Liu et al., 2019), suggesting some inclusion of coal burning. As both these sources follow a similar typical seasonal variation, i.e., high concentration during the cold period, it makes their separation difficult due to correlation.

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Secondary inorganics. Secondary inorganics were typically characterized by high contributions to 372 NO_3^{-} , SO_4^{2-} and NH_4^+ (55%, 56% and 56% of the total species mass, respectively) (Figure 1). This 373 factor showed a temporal variation, with remarkably high concentrations observed during the period 374 of high relative humidity (RH) and low ozone concentration in the winter (Figure S6). The 375 heterogeneous reactions on pre-existing particles in the polluted environment under high RH and low 376 377 ozone conditions have been shown to play a key role in the formation of secondary aerosols compared to gas-phase photochemical processes (Sun et al., 2013; Niu et al., 2016; Ma et al., 2017b). Therefore, 378 379 aqueous phase processes may be the major formation pathway for secondary inorganic aerosols in Beijing during the study period. Additionally, the factor showed a similar contribution (22-24%) to 380 PM mass in winter at both sites. This value is lower than the value reported by Liu et al. (2019) at the 381 other urban location (44%) in Beijing as a part of same APHH-Beijing campaign, although it should 382 be noted that the sampling site and dates of sampling differed. We also noticed the source profile 383 reported by Liu et al. (2019) contained a majority of all measured secondary inorganic species (>70%) 384 as well as 20% of OC while the factor identified in the present study only accounted for ~55% of 385

secondary inorganic species and 11% of OC with remaining fractions identified in other factors. Thus,
although the identification of the factor was "secondary" in both studies, they do not represent exactly
the same source. The modelled difference in the contribution of this factor to PM mass may also be
related to the uncertainties of the input species: a filter-based dataset was used in the present study
while Liu et al., (2019) used online measurements.

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The highest contribution to the PM mass was observed during the summertime, with an average concentration of 11.1 μ g m⁻³ (40%) and 13.2 μ g m⁻³ (48%) at IAP and PG, respectively (Figure 3).

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Traffic emissions. The traffic emissions factor showed relatively high contributions to metallic 395 elements, such as Zn (47%), Pb (57%), Mn (27%) and Fe (22%) (Figure 1). Zn is a major additive to 396 lubricant oil. Zn and Fe can also originate from tyre abrasion, brake linings, lubricants and corrosion 397 of vehicular parts and tailpipe emission (Pant and Harrison, 2012; Pant and Harrison, 2013; 398 Grigoratos and Martini, 2015; Piscitello et al., 2021). As the use of Pb additives in gasoline has been 399 banned since 1997 in Beijing, the observed Pb emissions may be associated with wear (tyre/brake) 400 rather than fuel combustion (Smichowski et al., 2007). These results suggest the contribution of both 401 exhaust and non-exhaust traffic emissions to this factor. Further insight on the type of non-exhaust 402 emissions is hard to predict as these metal concentrations vary according to several parameters, such 403 as traffic volume and patterns, vehicle fleet characteristics and the climate and geology of the region 404 (Duong and Lee, 2011). 405

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Traffic sources accounted for 9% and 6% of $PM_{2.5}$ mass during the winter and summertime at IAP (Figure 3), corresponding to an average concentration of 7.4 µg m⁻³ and 1.8 µg m⁻³, respectively. In addition, a low contribution (4%) was observed at PG during both seasons as PG experiences a low traffic volume. The contribution of the traffic source to the $PM_{2.5}$ mass was found to be low compared to other studies conducted in the Beijing area; (14-20%) (Li et al., 2019; Tian et al., 2016; Yu et al., 2013; Liu et al., 2019), with the exception of a study by Ma et al. (2017a) where a similar contribution

was reported. The observed low contribution was further supported as a recent study also confirms 413 that road traffic remains a dominant source of NO_x and primary coarse PM, however, it only accounts 414 a relatively small fraction of PM_{2.5} mass at urban locations in Beijing (Harrison et al., 2021). It should 415 416 be noted that nitrate that can be formed from NO_x emitted from road traffic is not included in this factor. Despite the low factor contribution, the resolved chemical profile of this source was consistent 417 with previously identified profiles linked to road traffic emissions in the Beijing area (Ma et al., 418 2017a; Yu et al., 2013). We noticed that OC/EC contribution in this factor is relatively low, while it 419 may be higher in traffic emissions. However, given the modern gasoline fleet in Beijing (Jing et al., 420 2016), it is not unexpected to observe low OC and EC contribution. We do expect higher OC and EC 421 contribution from diesel vehicles. In addition, there was no obvious seasonal variation as expected, 422 though slightly higher concentrations were observed in the cold period, probably due to the typical 423 atmospheric dynamics, and consequent poorer dispersion at this time of year. 424

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Metallic elements such as Mn, Fe and Zn were also used previously to indicate industrial activities 426 (Li et al., 2017; Yu et al., 2013). Back trajectory analysis reveals the influence of local emissions with 427 a slight regional transport during the winter period at both sites, dominated by north easterly flow 428 (Figure S7). Therefore, there is a possibility that these elements could also come from the Hebei 429 province where a large number of smelter industries are located. North-easterly and south-easterly 430 flows were dominant during the summer period at IAP and PG with possible regional influence. These 431 observations suggest that indeed this factor contains traffic aerosols, though a significant influence 432 of industrial emissions cannot be ruled out. 433

434

Road dust. This factor makes a major contribution to crustal species, such as Na⁺, Al and Fe (60%, 48% and 34% of species in this factor respectively) suggesting this factor may represent the characteristics of a dust related source as reported previously (Kim and Koh, 2020). Such a high contribution of Na⁺ in the identified factor was unexpected. Na is a major element of sea salt, seaspray and marine aerosols (Viana et al., 2008), and has also been found to be enriched in fine

particulates from coal combustion (Takuwa et al., 2006). It is notable that a high proportion of Na⁺ 440 was attributed to road dust in a previous study conducted at the same urban site (Tian et al., 2016), 441 and a crustal source seems likely, but has not been confirmed. In addition, the given factor also 442 443 included significant contributions to Mn, Pb and Zn (26%, 23% and 20% of species in this factor respectively), which are associated with brake and tyre wear as mentioned above (Pant and Harrison, 444 2012; Pant and Harrison, 2013; Grigoratos and Martini, 2015; Piscitello et al., 2021). High 445 concentrations of Zn and Pb have also been reported for particles emitted from asphalt pavement 446 (Canepari et al., 2008; Sörme et al., 2001). In addition, the ratio of Fe/Al observed in the factor 447 chemical profile was 1.26, much higher than the value observed in the earth's crust (0.6), suggesting 448 449 an anthropogenic origin of some Fe (Sun et al., 2005). This is likely as processes associated with vehicles, such as tyre/brake wear and road abrasion, can contaminate soil with metals, as the urban 450 sampling site is located close to roads suggesting the resolved factor is likely linked to road dust 451 emissions. These metals (Fe and Al) can also have industrial sources as already reported in the Beijing 452 area (Wang et al., 2008; Tian et al., 2016; Yu et al., 2013; Li et al 2019). The Beijing-Tianjin-Hebei 453 454 region is the largest urbanised megalopolis region in northern China, and home to many iron and steel making industries. Fe is characteristic components of iron and steel industry emissions (Li et al., 455 2019) while Al may also come from metal processing (Yu et al., 2013). However, disentangling the 456 457 influence of industrial emissions would require further investigation.

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This source also made significant contributions to OC, EC and SO_4^{2-} (11-19%) (Figure 1) and was consistent with the road dust source profiles observed previously in the Beijing area (Song et al., 2006; Song et al., 2007; Tian et al., 2016; Yu et al., 2013). This factor accounted for 20% of the PM_{2.5} mass during the summertime (5.5 µg m⁻³) with exceptionally low contribution (3%) during the cold period at PG (Figure 3). However, the factor contribution at IAP was similar during both seasons. In addition, the contribution to PM mass at IAP in this study was similar to that reported by Tian et al.

Crustal dust = 1.16(1.9Al + 2.15Si + 1.41Ca + 1.67Ti + 2.09Fe)467 468 A good correlation was observed between the estimated crustal dust and this factor during both 469 seasons at PG (rural, winter: $r^2 = 0.78$, m (slope) =0.9; summer: $r^2 = 0.94$, m=0.5) and IAP (urban, 470 winter: $r^2 = 0.51$, m=1.3; summer: $r^2 = 0.68$, m=1.2), highlighting that this may also contain a 471 significant fraction of crustal dust (Figure S8). This suggests that the identified factor is not resolved 472 cleanly and contains a mixed characteristic of road dust and crustal dust. 473 474 Soil dust. This factor mainly represents wind-blown soils and was typically characterized by a high 475 contribution to crustal elements, such as Ti (63%), Ca²⁺ (41%), Fe (27%) and Al (17%) (Figure 1). 476 In addition, the contributions to Mn and Zn in the factor profile (Mn=24%, Zn=15%) suggest that the 477 given source also included resuspended road dust but probably to a lesser extent. This source also 478 showed a significant contribution to n-alkanes (e.g., C29, C31), derived from epicuticular waxes of 479 higher plant biomass (Kolattukudy, 1976; Eglinton et al., 1962), with the highest contribution (37%) 480 to C31. This suggests the presence of plant derived organic matter in the soil dust, which is also 481 consistent with a high contribution to OC (15%). 482

(2016) and the studied urban site in both cases was the same. Crustal dust mass was also estimated

based on the concentrations of Al, Si, Fe, Ca, and Ti using the equation below (Chan et al., 1997).

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No clear seasonal variation was observed at PG. However, this factor showed a high contribution (35%, $9.8 \ \mu g \ m^{-3}$) to PM_{2.5} mass during the summertime at IAP, while the contribution during other seasons at both sites was less than 10% (Figure 3). The factor profile resolved here was similar to the profile reported by Ma et al. (2017a) for soil dust, but their soil dust factor only showed a 10 % contribution to PM_{2.5} mass. In addition, other previous studies (Yu et al., 2013; Zhang et al., 2013) also reported significant contribution of soil dust to PM_{2.5} mass, suggesting that soil dust is an important contributor to PM_{2.5} mass in the Beijing area. It is also expected as Beijing is in a semi-arid

region and there are sparsely vegetated surfaces both within and outside the city. This factor also 491 showed good agreement with the crustal fraction estimated from the element masses only during 492 winter at PG ($r^2 = 0.51$) and summer at IAP ($r^2 = 0.58$). This again highlights the probable mixing of 493 494 this source with other factors, or mis-attribution. Back trajectory analysis also indicates the influence of regional transport during the summer period at IAP, dominated by south easterly-westerly flow 495 (Figure S9) due to high windspeeds (3.6 m s^{-1}) . Therefore, there is a possibility that the high 496 497 contribution is linked to long-range transport in advected air masses. A recent study (Gu et al., 2020) 498 conducted in Beijing showed the high concentrations of more oxidised aerosols during summer due to enhanced photochemical processes; however, such type of source was not resolved due to a lack 499 500 of filter based markers. This suggests the given source may contain some influence from an unidentified/unresolved SOC fraction. Although the most plausible attribution appears to be to soil 501 dust, it is not fully resolved from other sources. 502

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The use of Si in PMF could provide a better understanding on these dust related sources. However, it 504 is not used in the present PMF input due to high number of missing data points. The sensitivity of 505 PMF results to the use of Si has also been investigated by adding Si to the input matrix and providing 506 high uncertainty to the missing data. No change was observed in the factor profile and temporal 507 variation of the resolved factors compared to the present one. In addition, we also noticed a good 508 509 correlation between Si and Al, where Al has been used in PMF (Figure S10). Several PMF runs were also made with inorganic data only, however the resolved factors were either mixed or hard to 510 identify. In addition, attempts to improve the PMF results by varying the input species and by 511 analysing data for the IAP and PG sites separately did not offer any advantage. 512

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516 3.2 Comparison of Filter-Based PMF Results with other Receptor Modelling Approaches 517 on the same Dataset

518 The source apportionment results from PMF were compared with those from CMB on the same filterbased composition data and PMF performed on other measurements (i.e. online AMS (PM1), offline 519 AMS $(PM_{2.5})$ to get a deeper insight into the identified PMF factors and their origins (Figsures 4, 5, 520 6 and 7). The CMB method resulted in the estimation of eight OC sources (i.e., vegetative detritus, 521 residential coal combustion (CC), industrial CC, cooking, diesel vehicles, gasoline vehicles, biomass 522 burning, other OC), including one secondary factor (Other OC) at both sites (IAP and PG) The online 523 AMS datasets allowed the identification of 6 OA (MOOOA (more oxidised oxygenated OA), 524 LOOOA (low more oxidised oxygenated OA), OPOA (oxidised primary OA), BBOA (biomass 525 burning OA), COA (cooking OA), CCOA (coal combustion OA) factors during winter at IAP, while 526 analyses on the offline AMS measurements resolved 4 OA (OOA, BBOA, COA, CCOA) factors. 527

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For these analyses, OC concentrations related to the online/offline AMS OA factors were further calculated by applying OC-to-OA conversion factors specific to each source, i.e., 1.35 for coal combustion organic carbon (Sun et al., 2016), 1.38 for cooking organic carbon, 1.58 for biomass burning organic carbon (Xu et al., 2019), and 1.78 for the oxygenated fraction (Huang et al., 2010) and used to evaluate the OC concentrations of relevant OA factors.

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Only OC equivalent concentrations were used to perform comparison for all approaches. OC mass closure was also verified at IAP during the wintertime by investigating the relation between: OC modelled by online AMS PMF vs filter based PMF ($r^2=0.7$, slope=1.17), OC measured vs OC modelled by filter based PMF ($r^2=0.7$, slope=1.07), OC measured vs OC modelled by online AMS PMF ($r^2=0.9$, slope=0.92), OC modelled by offline AMS PMF vs OC model by filter based PMF ($r^2=0.6$, slope=0.75), OC measured vs OC modelled by offline AMS PMF ($r^2=0.9$, slope=1.41), and OC measured vs WSOA (offline AMS) ($r^2=0.9$, slope=0.85) (Figure S11). The comparison of OC

modelled by PMF and CMB was also investigated at IAP ($r^2=0.8$, slope=1.05) and PG ($r^2=0.6$, 542 slope=1.78) (Figure S12). All source apportionment approaches showed a fairly good agreement in 543 reconstructing the total OC mass, justifying their direct comparison. In addition, it should be noted 544 545 that the difference in the sampling size cut-off between online AMS (NR-PM1) and filter measurements (PM_{2.5}) may contributes to the differences observed in the source apportionment 546 547 results. Therefore, we also compared the relation between NR-PM measured vs PM measured $(r^2=0.96, slope=0.92)$, and NR-PM measured vs PM modelled by filter based PMF $(r^2=0.9, r^2=0.9)$ 548 slope=1.29) (Figure S13). The agreements observed suggests that the most of the PM_{2.5} mass was 549 550 accounted for by the PM₁ fraction, indicating that the difference in the size-cut off is relatively small.

- 551
- 552 (a) With CMB results at IAP

Resolved CMB and PMF factors were compared including data from both seasons at IAP and PG 553 (Figure 4). A good correlation ($r^2=0.6$, n=68, p<0.05) was observed between biomass burning factors, 554 suggesting that this source was well resolved using both approaches (Figure 4). However, a slightly 555 higher concentration was reported by the CMB model (2.0 and 1.6 µg m⁻³ by CMB and PMF 556 respectively). Individual coal combustion factors (industrial/residential) did not shown any 557 significant correlation ($r^2 < 0.2$) with the coal combustion factor identified using PMF, although the 558 559 total coal combustion fraction from CMB, the sum of industrial and residential fractions, did show an improved correlation ($r^{2=}0.4$). Some improvement on the correlation was seen if two outlier 560 datapoints were removed (see Figure 4). A likely reason is that PMF did not resolve coal combustion 561 and biomass burning factors well as both factors presented a strong seasonal pattern with high 562 concentrations during the winter. Another possibility is the difficulty in resolving primary and 563 564 secondary fractions due to a lack of secondary organic markers used in the study. This was further supported by the fact noted above that the PMF biomass burning factor also contained some signal 565 from coal combustion activities. The sum of coal combustion and biomass burning factors from both 566

approaches showed a good correlation ($r^2=0.7$, n=68, p<0.05), suggesting a common emission pattern 567 (e.g., high in winter and low in summer), making it challenging to resolve them. Factors linked to 568 vehicle emissions did not show any correlation. A weak correlation ($r^2=0.3$, n=68, p<0.05) was 569 570 observed between Other OC from CMB, a proxy for the secondary organic fraction and the PMF secondary inorganics factor. In addition, other OC also weakly correlated with soil dust ($r^2=0.22$, 571 n=34, p<0.05) in summer, suggesting the mixing of unresolved secondary fraction with soil dust 572 profile and supports the hypothesis discussed above. It should be noted that other OC could also 573 contain unresolved primary fractions as PMF results indicated substantial influence of industrial 574 emissions and dust related sources. However, the source profiles related to industrial emissions and 575 576 dust were not accounted for in the CMB model (Xu et al., 2021).

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578 (b) With CMB results at PG

The comparison was also made using data from both seasons at PG (Figure 5). Biomass burning 579 aerosols showed a good correlation for both approaches ($r^2=0.7$, n=20, p<0.05) but a substantially 580 higher concentration was estimated by the CMB model (5.1 µg m⁻³ and 2.0 µg m⁻³ by CMB and PMF 581 respectively). A significant correlation was also seen between traffic related factors from CMB and 582 PMF (gasoline-CMB vs traffic ($r^2=0.6$, n=20, p<0.05), diesel-CMB vs traffic ($r^2=0.6$, n=20, p<0.05)), 583 indicating that traffic sources resolved using PMF at the PG site may have included signals from both 584 diesel and gasoline vehicles; however it was not conclusive at the IAP site, as discussed above. This 585 suggests the traffic source resolved using PMF may contain particles linked to traffic emissions, but 586 the influence of other sources is prominent at IAP and resulted in poor correlation. In addition, for 587 traffic related factors from CMB, both showed a higher concentration (gasoline-CMB= 0.8 ug m^{-3} . 588 diesel-CMB=4.5 µg m⁻³, traffic-PMF=0.2 µg m⁻³). As with IAP, no significant correlation was 589 observed between coal combustion factors from both approaches. The sum of coal combustion and 590 biomass burning factors from both approaches also did not present a good correlation ($r^2=0.3$, n=20, 591

p<0.05). This highlights the limitation of these methodologies to apportion sources when extreme 592 meteorological conditions may lead to high internal mixing of sources. Unfavourable dispersion 593 conditions have been previously observed in the Beijing region during severe haze events in winter 594 595 (Wang et al., 2014). A high correlation was observed between Other OC (CMB) and secondary inorganics (PMF) ($r^2=0.7$, n=20, p<0.05). In addition, Other OC also showed a very high correlation 596 with the biomass burning factor resolved from PMF ($r^2=0.9$, n=20, p<0.05). This suggests that the 597 598 biomass burning factor in PMF may contain a substantial amount of aged aerosols since carbon emitted during biomass burning is in some cases oxygenated and water soluble (Lee et al., 2008b) 599 and is subject to rapid oxidation in the atmosphere. 600

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602 (c) With online AMS PMF factors at IAP (winter)

BBOC (biomass burning OC) from PMF-AMS analysis agreed well with that from PMF ($r^2=0.7$. 603 n=27, p<0.05; 4.0 μ g m⁻³ and 3.1 μ g m⁻³ by online AMS and PMF, respectively) (Figure 6). Coal 604 combustion related factors showed a modest correlation (CCOC (coal combustion OC) vs coal 605 combustion-PMF, $r^2=0.4$, n=27, p<0.05) but the mass concentration of the coal combustion source 606 by PMF (11.3 μ g m⁻³) is significantly higher than by PMF-AMS (CCOC=4.7 μ g m⁻³). This may 607 partly be due to the different size cut offs used by these measurements (PM_1 for AMS vs $PM_{2.5}$). In 608 609 addition, significant improvement on the correlation was seen if two outlying points were removed $(r^2=0.8, \text{ see Figure 6})$. Oxygenated fractions from AMS, MOOOC (more oxidised oxygenated OC) 610 and LOOOC (low oxidised oxygenated OC) also exhibited a good correlation with secondary 611 inorganics (LOOOC vs secondary inorganics ($r^2=0.6$, n=27, p<0.05, LOOOC=2.9 µg m⁻³, secondary 612 inorganics=1.6 μ g m⁻³), MOOOC vs secondary inorganics (r²=0.7, n=27, p<0.05, MOOOC=4.4 μ g 613 m^{-3})). This was also confirmed by LOOOC and MOOOC showing a good correlation with NO₃⁻ and 614 SO4²⁻ previously (Cao et al., 2017). The formation of both secondary inorganic aerosol and 615 oxygenated organic aerosol is dependent upon largely the same set of oxidant species, notably but 616 not solely the hydroxyl and nitrate radicals. In both cases there are also both homogeneous and 617

heterogeneous (aqueous phase) pathways, so conditions which promote the formation of oxidised 618 organic aerosol will also favour formation of secondary inorganic aerosol, and hence a correlation is 619 to be expected, and is often observed (Hu et al., 2016; Zhang et al., 2011). In addition, both 620 621 oxygenated fractions were also found to be correlated with biomass burning aerosols (LOOOC vs biomass burning-PMF (r²=0.7, n=27, p<0.05), MOOOC vs biomass burning-PMF (r²=0.6, n=27, 622 p<0.05)). This further highlights a potentially important role of biomass burning activity in SOA 623 formation at IAP. A good correlation was also observed between OPOC (oxidised primary OC) and 624 secondary inorganics and biomass burning ($r^2=0.7$, n=27, p<0.05). 625

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(d) Offline AMS PMF factors at IAP (winter)

BBOC from PMF-offline AMS analysis showed a good correlation with that from PMF ($r^2=0.6$, n=32, 628 p < 0.05) (Figure 7) but the mass concentration of BBOC (4.6 µg m⁻³) is higher than biomass burning 629 (3.1 µg m⁻³) from PMF. This was also noticed above while comparing with BBOC resolved using 630 online AMS PMF, suggesting a potential uncertainty in estimating the source contribution from 631 632 biomass burning. The uncertainty in filter-based PMF analysis could be related to model error. This was further supported as biomass burning factor also made significant contributions to Ca^{2+} (15%). 633 Ni (30%), Cu (50%), and Al (35%), and these species are not necessarily from biomass burning 634 emissions but they were not resolved by PMF. In addition, the uncertainties linked to PMF-AMS 635 analysis could also contribute. A high correlation was noticed for secondary factors resolved using 636 both approaches (OOC (oxygenated OC) vs Secondary inorganics, $r^2=0.8$, n=32, p<0.05). OOC also 637 showed a good correlation with the biomass burning factor (OOC vs biomass burning-PMF, $r^2=0.7$, 638 n=32, p<0.05). This supports the hypothesis discussed previously on the origin of oxygenated 639 640 fractions.

641

642 Overall, the comparison of filter based PMF results was in broad agreement with other receptor 643 modelling approaches applied on the same dataset. However, large discrepancies were also observed

for some factors / sources. Common sources such as biomass burning and coal combustion were well 644 resolved using all approaches with some exceptions observed when using filter based PMF approach. 645 This could be linked to internal mixing of sources when the influence of climate and local 646 647 meteorology on both sources is predominant and making it challenging to resolve using PMF. A good agreement was also observed between secondary inorganic aerosols and secondary fractions resolved 648 649 using other approaches. However, sources identified based on metal signatures using PMF indicated some mixing or mis-attribution. For example, the influence of unresolved SOC on the soil dust profile 650 was observed during summer. 651

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653 3.3. Comparison with Previous PMF Source Apportionment Results in Beijing

In this section an attempt has been made to understand the PM sources identified in the Beijing 654 metropolitan area by previous studies. The goal was here to assess the previous PMF source 655 apportionment results and report any discrepancies noticed in the resolved sources using PMF. This 656 may provide useful insight on sources resolved in the present study and also in exploring the issues 657 associated with filter based PMF modelling in the Beijing metropolitan area. Details of the studies 658 659 conducted to evaluate PM sources using a PMF model applied to inorganic and organic markers in the Beijing metropolitan area are presented in Table S5 and the major outcomes are discussed 660 hereafter. 661

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Overall, these previous PMF studies provide insights on PM sources in the Beijing metropolitan area (Li et al., 2019; Ma et al., 2017a; Tian et al., 2016; Yu et al., 2013; Song et al., 2007; Song et al., 2006; Liu et al., 2019; Wang et al., 2008; Zhang et al., 2013). The major identified sources are dust, traffic emissions, coal combustion, industrial activity, secondary inorganic aerosols and biomass burning. Although there is a general issue of their inability to identify sources such as secondary organic aerosol and cooking emissions, similar to the present study, due to the lack of organic markers used in the PMF model to apportion these sources. However, beyond this, their PMF outcomes were not consistent. Large discrepancies between the sources were seen (Table S5) based on the sources
identified as well as their contribution to PM mass concentrations. Several factors could cause these
differences such as the chemical species used as input in the PMF model, the period of the study,
identification of sources based on chemical signatures and changes to the sources with time.

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Input species considered within the previous studies were combinations of water-soluble ions, metallic elements, OC and EC. Similar input species were used in all of these studies, with the exception of the studies by Yu et al. (2013) and Li et al. (2019) who used only metallic elements for the source apportionment. As shown in this study, including organic markers may help to resolve some of the primary sources.

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Another important parameter, the chemical species used for identifying sources were not always 681 consistent. For example, coal combustion was resolved based on high contribution of OC, EC, and 682 Cl present in the factor profile by Zhang et al. (2013), Wang et al. (2008), Song et al. (2007) and Song 683 et al. (2006), in accordance with source profiles determined in the laboratory (Zheng et al., 2005). 684 High Cl associated with fine aerosols in winter is a distinctive feature in Beijing and even around 685 inland China, which is ascribed to coal combustion (Wang et al., 2008). Contrarily, Tian et al. (2016) 686 identified coal combustion based on a high contribution of OC and EC, while the high contribution 687 of Cl was attributed to a biomass burning source, similar to another study (Ma et al., 2017a). In other 688 studies (Li et al., 2019; Yu et al., 2013; Liu et al., 2019) coal combustion was resolved based on the 689 presence of metallic elements such as V, Se, Co, Cd, As and Ni, where V and Ni are widely used 690 markers for oil combustion (Mazzei et al., 2008). High loadings of As and Se have also been reported 691 as a typical source characteristic of coal combustion (Vejahati et al., 2010). Similar to coal 692 693 combustion, biomass burning was often characterised using the presence of K (Li et al., 2019; Tian et al., 2016; Yu et al., 2013; Song et al., 2007; Song et al., 2006; Zhang et al., 2013; Liu et al., 2019; 694 Wang et al., 2008), a typical marker of biomass burning. Farming in Beijing's suburban districts has 695

been extensive in recent years. Burning of the crop remnants and fallen leaves by farmers in autumn 696 and winter results in the enhanced emissions of K. In addition, the contributions of Cl and Na were 697 also considered for the identification of these sources in some cases, depending on the species used 698 699 within the input (Song et al., 2007; Song et al., 2006; Tian et al., 2016). This highlights the fact that none of the studies have used organic markers such as picene and levoglucosan which are very 700 701 specific to these combustion sources as discussed before, which may cause uncertainty in the resolved 702 sources. However, in the present study the use of organic markers played a key role in the identification of these sources and their better apportionment. Despite this, some issues were observed 703 with these identified sources during winter due to extreme meteorological conditions as well as co-704 705 emission of these aerosols at the same time, probably indicative of poor performance of the PMF model under certain conditions. 706

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Other important sources linked to traffic emissions, industrial activities and dust, are commonly 708 resolved among all the studies. The characterisation of these sources was predominantly based on the 709 710 metallic elements. For example, Zn, Cu, and Pb including sometimes EC were most often used to characterise traffic emissions among all previous studies. Both Zn and Cu have been identified within 711 brake linings and tyre fragments (Thorpe and Harrison, 2008) and Pb has been used in the past within 712 713 gasoline as an anti-knock additive in China (Li et al., 2019). However, Cu and Zn can also serve as indicators for industrial sources (Li et al., 2017; Yu et al., 2013). Other metallic elements (e.g., Sb, 714 Cr, Mn, K, Br and Ba) were also considered in certain cases to trace traffic emissions (Ma et al., 715 2017a; Tian et al., 2016; Yu et al., 2013). However, a high contribution of Cr, Mn and sometimes Fe 716 to the given sources has also been attributed to industrial activities. Both Cr and Cr-containing 717 718 compounds are widely used in metallurgy, electroplating, pigment, leather and other industries (Dall'Osto et al., 2013). A previous study found that ferrous metallurgy could emit Mn (Querol et al., 719 2006). Furthermore, both Fe and Mn are characteristic components of iron and steel industry 720 emissions. In addition, Co, Mg, Al, Ca, Cd, Pb, Tl, Zn, V, Ni and Cu were also considered for the 721

apportionment of industrial sources (Tian et al., 2016; Yu et al., 2013). Zhang et al. (2013) identified a mixed source of traffic and incineration emissions, based on high loading of Cu, Zn, Cd, Pb, Sb, Sn, Mo, NO_3^- and EC. In the present study, the assignment of road traffic emissions was based on high loadings of Zn and Pb. It was also seen that the given source may contain some influence from industrial activities, as the industrial contribution was not resolved like previous studies and probably accounted in other factors. Thus, it is clear that these metals could belong to several sources and their proper assignment to respective sources is difficult in the complex environment.

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The same issue was observed with the assignments of dust type (dust/road dust/soil dust/mineral dust/yellow dust/local dust) sources. Although the dust type sources were often found to be composed of crustal elements (e.g., Ca, Mg, Si, Ti, Al, Fe), the attribution of crustal elements to a particular source was not consistent from one study to another previously. The two dust sources (road dust and soil dust) identified in the present study also indicated mixing with other factors.

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The identification of the secondary inorganic aerosol factors was often based on the high contribution 736 of water soluble ions (NO₃⁻, SO₄²⁻, NH₄⁺), consistent with other studies (Ma et al., 2017a; Song et al., 737 2007; Song et al., 2006; Tian et al., 2016; Liu et al., 2019; Wang et al., 2008; Zhang et al., 2013). 738 These results highlight the role of chemical species used in characterising source profiles and their 739 740 influence on the variability noticed in the Beijing metropolitan area. This issue arises because many of these species are not source specific, making it challenging to directly link PMF factors to sources. 741 Pant and Harrison (2012), reviewing receptor modelling studies from India, noted a tendency to 742 attribute metal-rich source profiles to "industry" in a rather casual manner without evidence of local 743 industrial sources. 744

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The change in sources and emissions over the course of time due to stringent emissions regulationscould also be considered plausible for the observed variability in the chemical profile and contribution

of identified sources. Li et al. (2019) showed levels of trace metals (V, Cr, Mn, As, Cd and Pb) 748 decreased more than 40% due to the emission regulations, while crustal elements decreased 749 considerably (4-45%), suggesting emissions from anthropogenic activities were suppressed. A 750 751 reduction in the contribution of sources such as dust and industrial activity was observed in the present 752 study and another recent study performed by Liu et al. (2019) relative to the previous ones, indicating 753 the effect of regulatory measures on the contribution of identified sources to PM_{2.5}. However, the 754 concentration of the majority of metallic elements (K, Cr, Mn, Fe, Co, Cu, Zn, As, Ag, Cd and Pb) increased when pollution levels changed from clean days to heavily polluted days. This highlights 755 that specific atmospheric conditions could also play a major role for the observed variability. Another 756 757 factor is the time of year when these studies were conducted as some of the identified sources (e.g., coal combustion and biomass burning) exhibit typical seasonal patterns. During a low concentration 758 period, PMF models may have difficulty in resolving sources, leading to mixing of factors. 759

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Overall, the present study provides a view of existing PM_{2.5} sources in the Beijing metropolitan area 761 by applying the PMF model to a filter-based dataset, which included water soluble ions, metals and 762 763 organic markers. Despite this, factors that were resolved based on metal signatures were not fully resolved and indicate a mixing of different sources. As a part of same campaign, also discussed above, 764 Liu et al. (2019) used a similar approach by applying PMF on high resolution (1-hour) data, which 765 included OC, EC, ions, and metals, and did not encounter any issue. However, previous filter based 766 PMF studies conducted in the Beijing region that mostly included ions and metals in their input 767 dataset often showed difficulty in the proper assignment of metals to their respective sources. Even 768 the use of metal signatures from one to another study was not consistent. This highlights that the low 769 770 temporal resolution of filter data could not capture fast occurring atmospheric processes in Beijing, 771 and may lead to a "blurring" of sources by the long averaging period. Atmospheric circulation and dynamic mechanisms play a key role in persistent haze events in Beijing during the cold period (Wu 772 et al., 2017; Feng et al., 2014). Such events are associated with the high pollution periods and will 773

offer opportunities for chemical and physical transformation within the aerosol that lead to contravention of the requirement of receptor models for preservation of chemical profiles between source and receptor.

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778 **4. CONCLUSION**

This study presents the outcomes of PMF performed on the combined dataset collected at two sites 779 (IAP and PG) in the Beijing metropolitan area, including their comparison with source apportionment 780 results from other approaches or based on different measurements. The PMF analysis resulted in the 781 identification of seven sources: coal combustion, biomass burning, oil combustion, secondary 782 inorganics, traffic emissions, road dust and soil dust. These results were in a good agreement with 783 previously published source apportionment results made using PMF. However, factors that were 784 resolved based on metal signatures were not fully resolved and indicate an internal mixing of different 785 sources. In particular, soil dust, road dust and some industrial sources have many elements in common 786 and are very difficult to distinguish. 787

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PMF results were compared with sources resolved from CMB and with PMF performed on other 789 measurements (online AMS, offline AMS). Results showed a broad agreement with some notable 790 exceptions. While this study provides some confirmatory evidence on PM_{2.5} source apportionment in 791 Beijing, it highlights weaknesses of the PMF method when applied in this locality, and the results 792 should be viewed in the context of studies using other methods such as CMB which appear able to 793 give a more comprehensive view of the key sources affecting air quality. No industrial source profiles 794 795 were used as inputs to the CMB model reported here, so CMB offers no further insights into possible contributions from industry. 796

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800 AUTHOR CONTRIBUTIONS

- 801 This study was conceived by ZS and RMH. DS performed the PMF analysis and wrote the paper with
- the help of Z.S. and R.M.H. T.V.V. and D.L. conducted the aerosol sampling and laboratory-based
- chemical analyses. X.W. and J.X. conducted the CMB modelling at PG and IAP sites, respectively.
- All authors discussed the paper and approved the final version for publication.

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806 COMPETING INTERESTS

807 The authors declare that they have no conflict of interest.

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This article is part of the special issue "In-depth study of air pollution sources and processes within Beijing and its surrounding region (APHH-Beijing) (ACP/AMT inter-journal SI)". It is not associated with a conference.

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1317 FIGURE LEGENDS

1318 1319	Figure 1.	Factor profiles for identified factors at IAP and PG. The bars show the composition profile (left axis) and the dots, the Explained Variation (right axis).
1320 1321	Figure 2.	Temporal variation of identified factors at IAP and PG. Solid and broken lines represent IAP and PG, respectively.
1322	Figure 3.	Contribution of different sources to PM _{2.5} mass at IAP and PG.
1323 1324	Figure 4.	Correlations observed between PMF and CMB results at IAP. *If two outlying points are removed from the coal combustion-PMF, correlations are markedly improved.
1325	Figure 5.	Correlations observed between PMF and CMB results at PG.
1326 1327 1328	Figure 6.	Correlations observed between PMF and online AMS PMF results at IAP (winter). *If two outlying points are removed from the coal combustion-PMF, correlations are markedly improved.
1220	F ! #	
1329	Figure 7.	Correlations observed between PMF and offline AMS PMF results at IAP (winter).



Figure 1. Factor profiles for identified factors at IAP and PG. The bars show the composition profile(left axis) and the dots, the Explained Variation (right axis).





Figure 2. Temporal variation of identified factors at IAP and PG. Solid and broken lines representIAP and PG, respectively.

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Coal combustion Road dust Traffic emissions Oil combustion Secondary inorganics Biomass burning Soil dust

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Figure 3. Contribution of different sources to PM_{2.5} mass at IAP and PG.





Figure 4. Correlations observed between PMF and CMB results at IAP. *If two outlying points are 1346 removed from the coal combustion-PMF, correlations are markedly improved. 1347

Coal combustion-PMF (µg m⁻³)



1350 Figure 5. Correlations observed between PMF and CMB results at PG.1351



Figure 6. Correlations observed between PMF and online AMS PMF results at IAP (winter). *If two
outlying points are removed from the coal combustion-PMF, correlations are markedly improved.



1358 Figure 7. Correlations observed between PMF and offline AMS PMF results at IAP (winter).