1	The characteristics of environmental particulate matter in the urban area of						
2	Beijing, China, during the 2008 Olympic Games						
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## 23 Abstract

Atmospheric particulate matter (PM) and street dust samples from the Chaoyang 24 25 District of eastern Beijing were studied over a period encompassing the 2008 Beijing Olympic Games. PM<sub>10</sub> concentration data are combined with trajectory clustering and 26 27 potential source contribution function (PSCF) methods to identify the principal transport pathways. Sources for high-concentration aerosol events and airflow from 28 the surrounding Hebei Province and Shandong Province to the southeast are found to 29 30 exert the most significant external influence on Beijing's air quality. China undertook 31 a number of initiatives to improve air quality for the Olympic Games and we show that PM<sub>10</sub> concentrations and magnetic susceptibility were significantly lower during 32 the Olympic period compared to the pre-Olympic period confirming that controlling 33 34 local sources in Beijing and shutting factories in surrounding provinces substantially improved air quality. On short timescales PM<sub>10</sub> shows an inverse correlation to 35 relative humidity and hence precipitation which acts to improve air quality. 36 37 Atmospheric PM and street dust remained high through the Olympic period probably due in part to redistribution of historical sources and implying that the aim of zero 38 pollution is not achievable in the short term. Analysis of the heavy metal content in 39 both PM and street dust identifies consistently high values of Zn, with Pb relatively 40 higher in the PM; a primary source in vehicular emissions therefore seems likely. 41

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43 Keywords: Atmospheric pollution; 2008 Olympic Games; Beijing; PM<sub>10</sub>; magnetic
44 susceptibility; heavy metals

45 **1. Introduction** 

Particulate matter (PM) in the atmosphere and in ground deposits can originate 46 47 from natural sources (dust blown into the air by wind, salts splashed into the air by sea spray and soot from volcanoes and forest fires) and from various anthropogenic 48 activities of which the biggest sources are vehicle and smokestack emissions, and the 49 creation of dust generated when vegetation has been removed for construction or 50 grazing purposes. When averaged globally, anthropogenic PM appears to account for 51 about 10% of the total aerosol amount (Perrino, 2010) but this figure varies greatly 52 53 from place to place as do the chemical compositions and inferred sources. PM can vary in size from sub-micron aerosols to visible dust particles: the coarse particles 54 rapidly removed from the air by sedimentation are of local impact only, whereas fine 55 56 particles can have a global reach (Perrino, 2010). Urban surfaces typically receive fine PM issued from remote sources through atmospheric transport as well as a wider 57 range of particle sizes from local human activities (Harrison et al., 1981; Thornton, 58 59 1991). Street dust can also be easily re-suspended back into the atmospheric aerosol by wind (Wise and Comrie, 2005) or vehicle movement (Almeidam et al., 2006). 60 Analysis is therefore complex but the integrated data from studies of PM in street dust 61 nevertheless provides the essential basis for understanding atmospheric pollution and 62 assessing effects on human health (e.g. Hien et al., 1999). 63

64 PM typically contains magnetic particles characterized by stable and intense 65 magnetic properties (Maher et al., 1999) with this magnetic fraction linked closely to 66 heavy metals such as zinc, cadmium and chrome (Georgeaud et al., 1997) but also to

mutagenic organic compounds (Morris et al., 1995), all of which are dangerous to 67 human health. Thus magnetic properties provide valuable proxies for deducing the 68 69 origin of PM and because of their control by the above factors they become a convenient signature of air pollution (Oiao et al., 2013). Routinely-measureable 70 71 magnetic parameters provide information on the concentration, domain state (or indirectly the magnetic grain size), and mineralogy of magnetic particles and 72 collectively these are related to original geological or subsequent environmental 73 processes (Liu et al., 2012). Statistical methods such as trajectory clustering have 74 75 been widely used to identify the pathways and sources of air pollution (e.g. Ashbaugh, 1983; Sirois and Bottenheim, 1995; Wang et al., 2006; Borge et al., 2007). In the 76 present investigation we study magnetic susceptibility in atmospheric fallout samples 77 78 and apply statistical clustering technique to a 5-month dataset of atmospheric trajectories to identify the particulate matter sources and long-range transport patterns 79 that can influence air pollution. 80

81 The 29th Olympic and Para-Olympic Games were held between 2008 August 8 and September 17, in Beijing, a densely populated city with more than 16 million 82 83 residents and 3 million motor vehicles. Traffic congestion and air pollution thus presented two major challenges to the organizers of the games. To improve air quality 84 and control traffic a series of measures were implemented which included the 85 relocation of industrial plants with large emissions outside of the city and the 86 implementation of new standards to reduce vehicular emission and limit their use. 87 Domestic controls included a progressive switching to clean fuels and low-sulfur coal 88

for household use before and during the Olympic period (Li et al., 2010; Zhou et al., 89 2010). Whilst most of these measures were intended to have a lasting effect, the 90 vehicular restrictions were largely temporary in nature and are therefore expected to 91 be detectable for only a limited time period. This research evaluating the impact of 92 these air pollution control measures is therefore classified as "before", "during" and 93 "after" the Games. The present study has had two objectives: firstly we have aimed to 94 determine the source regions influencing the air in Beijing in order that effective 95 source control strategies can be put into place in the longer term; secondly, we have 96 97 sought to evaluate the relationship between atmospheric PM and street dust. It is well known that the re-suspension of road dust particles from urban street surfaces is an 98 important source of atmospheric PM pollution (Amato et al., 2009; Martuzevicius et 99 100 al., 2011; Zhao et al., 2016), and the measurement of atmospheric deposits on street surfaces can be useful for studying deposition over a longer time intervals. 101

102 **2. Experimental Section** 

103 2.1. Sample collection

Atmospheric PM was determined by the gravimetric method at monthly intervals from June 2008 to March 2009 in the Chaoyang Distinct of eastern Beijing. The PM samples were collected in 15×30 cm cylindrical glass vessels containing glycol and the vessels placed on a 1.5 m sampling frame. PM samples were collected at two sampling sites: the first was a residential location at Sanlitun (SLT) near the Chaoyang Park (CY) containing the Olympic Site and the second was an industrial site, Fatou (FT) located near the Jing-Shen Highway. The Olympic Park is located at the

northwestern sector of Chaoyang district (Fig. 1), and street dust samples were 111 collected on roads with different traffic densities or pavements around the Olympic 112 Park between November 2007 and October 2008. The sampling sites were selected 113 from the traffic avenue and forest park inside the Olympic Park (AT) near to the north 114 5<sup>th</sup> ring road (Fig. 1) with samples collected using a nylon brush and non-magnetic 115 scoop from squares 0.5 to 1  $m^2$  in area prior to transfer to clean, self-sealing 116 polyethylene bags. To evaluate the relationship between PM and street dust over the 117 interval of enforcement measures, the samples were selected for analysis between 118 June 2008 and October 2008. The details of sampling sites and methods are described 119 in Qiao et al. (2011a, b). 120

121 2.2. Methods

122 All samples were air-dried and sieved through a mesh of size 500 µm to remove obvious refuses and the dust residue tightly packed into 10cc polyethylene cubes for 123 magnetic measurements in the Paleomagnetism and Geochronology Laboratory at the 124 Institute of Geology and Geophysics, Chinese Academy of Sciences. A total of 17 125 cubes from 4 sites were prepared for analysis in this way. Magnetic susceptibility 126 measurements using a Kappabridge KLY-3 were accompanied by determination of Fe, 127 V, Cr, Co, Ni, Cu, Zn and Pb heavy metal contents using inductively-coupled 128 spectrometry (ICP-MS); the latter technique employed the 129 plasma-mass DZ/T0223-2001 method with HR-ICP-MS (Element I) Finning MAT equipment 130 located in the Analytical Laboratory of the Beijing Research Institute of Uranium 131 Geology. The averaged mass amounts used for chemical measurement were 10 mg, 132

and for magnetic measurements were respectively 0.141g (PM samples) and 4.025g (street dust samples). Although the unit for mass specific susceptibility ( $\chi$ ) is more generally preferred, for the very small sample amounts involved with PM samples, the magnetic susceptibility is conveniently denoted by the volume susceptibility value ( $\kappa$ ).

The Geo-accumulation index,  $I_{geo}$ , is defined as  $I_{geo} = \log_2 (C_n/1.5B_n)$  where  $C_n$  is 138 the measured concentration of the heavy metals in the environment; B<sub>n</sub> is the 139 geochemical background value in soil (CNEMC, China National Environmental 140 141 Monitoring Center, 1990), and the factor 1.5 is introduced to minimize effects of possible variations in the background values as originally assessed from studies of 142 bottom sediments (Müller, 1969). This index provides a simple factor for assessing 143 144 the impact of natural geological processes on the natural background values and the influence of human activity responsible for the heavy metal pollution. The 145 geo-accumulation index consists of 7 classes or grades (Table 1), whereby the highest 146 147 class 6 reflects a 100-fold enrichment above the background values (Forstner et al., 1990). 148

To accommodate external processes 3-day back-trajectories arriving at Beijing (39.9N, 116.4E) 500m above ground level (a.g.l) were calculated using the Hybrid Single Particle Lagrangian Integrated Trajectory (HYSPLIT) model (Stein et al., 2015; Rolph, 2016) loaded into the geographic information system (GIS) based software, TrajStat (Wang et al., 2009). The NCEP (National Centers for Environmental Prediction) archive data downloaded from NOAA provided the meteorological data for input into the model. The  $PM_{10}$  concentrations for Beijing from June 2008 to October 2008 were calculated from the air quality index (AQI) reports for major Chinese cities and, for the purposes of statistically analyzing the data,  $PM_{10}$ concentrations at Beijing were assigned to corresponding trajectories using the Euclidean distance for the trajectory clusters.

- 160 **3. Results and discussion**
- 161 3.1. Transport Pathways and sources

Although PM sources in Beijing clearly contribute to PM concentrations in the 162 163 city, it has already been shown that surrounding areas contribute to Beijing's PM concentrations (Streets et al., 2007). Five clusters were therefore produced by the 164 clustering algorithm (a total of 152 trajectories classified into 5 clusters) to determine 165 166 the impact of different source regions on the PM concentration in Beijing; these cluster-mean trajectories are shown in Fig. 2 by different colors. The air masses 167 associated with clusters 1 and 2 have traveled over desert and semi-desert regions of 168 Inner Mongolia on the way to Beijing; cluster 3 air is anticipated to have initially 169 followed paths over Russia subsequent to passing southeasterly through desert and 170 semi-desert regions of Mongolia and Inner Mongolia before reaching Beijing. Each 171 year dust storms carry particulate matter from the deserts of Gobi and Teklimakan 172 towards Beijing, especially during the spring; other lithogenic sources of dust are bare 173 soils, coal heaps and construction sites occurring in and around Beijing. The air 174 masses associated with clusters 4 and 5 however, were from easterly and southerly 175 directions respectively that have passed over regions with a variable vegetated cover. 176

On the map showing results of the Potential Source Contribution Function 177 (PSCF) analysis (Fig. 3) high values of PSCF (>0.4) for PM<sub>10</sub> are found along Bohai 178 179 Bay and in Hebei and Shandong Provinces, where heavy industries are situated. Particles from these regions are mainly associated with transport paths recorded by 180 clusters 4 and 5 (Fig. 2). Based on the US Environmental Protection Agency's (EPA) 181 Models-3/Community Multi-scale Air Quality (CMAQ) model simulation over the 182 Beijing region, Streets et al. (2007) estimated that about 34% of PM<sub>2.5</sub> on average at 183 the Olympic Stadium site is attributable to sources outside Beijing. Correspondingly 184 185 our clustering results suggest that particle transport from Hebei (21.05%), Inner Mongolia (23.03%) and Shandong (23.03%) provinces had a significant impact on 186  $PM_{10}$  levels in Beijing between June and October 2008 (Fig. 2). Hence the local 187 188 control measures in Beijing were clearly insufficient for achieving air quality goals set for the Beijing Olympics. 189

190 3.2. Meteorology, particle concentration and magnetic susceptibility

Fig. 4 summarizes the record of temperature, relative humidity and PM<sub>10</sub> mass 191 concentration observations covering the time interval of the Olympic Games 192 compared with the experimental results of the present study for magnetic 193 susceptibility. The temperature (T) shows an inverse correlation with relative 194 humidity (RH), while the RH values remains high through the main interval, the PM<sub>10</sub> 195 mass concentration decreases to a minimum in the "during" period (Fig.4a-b). 196 Magnetic susceptibility ( $\kappa$ ) from the FT district adjoins the Jing-Shen Highway (see 197 Fig. 1) and displays some correspondence with the  $PM_{10}$  mass concentration in 198

Beijing dropping sharply in the "before" period and rising gently in the "after" period 199 (Fig.4c). Magnetic minerals in aerosols could be derived from combustion processes 200 201 related to industry, domestic heating or vehicles (Petrovský & Elwood, 1999). Therefore, magnetic parameters, notably magnetic susceptibility, are possible proxies 202 to monitor the relative changes of atmospheric PM pollution in an area over time. The 203 RH during the sampling periods in 2008 were almost at the same level as in 2009, but 204 the  $PM_{10}$  mass concentration was obviously lower (Fig. 4d). So the reduction in  $PM_{10}$ 205 and magnetic susceptibility is attributable to the source control measures implemented 206 207 during the Olympic and Paralympic Games; these included removing approximately one-half of the vehicles (~1.5 million cars) off the roads in Beijing on alternate days 208 under an even-odd license plate system, closing pollution-emitting factories, and 209 210 slowing down construction activities. The measures implemented were substantial and extended far beyond Beijing: in the neighboring Tianjin municipality, in Hebei, 211 Shanxi, and Shandong provinces, and in the Inner Mongolia Autonomous Region, 212 213 polluting factories were closed and high-emission cars removed from roads (Stone, 2008). 214

In our previous study of street dust in the Beijing Olympic Park we observed that magnetic compositions in street dust decreased significantly during the interval of the Olympic Games enabling magnetic measurements to serve as a tool for rapidly and efficiently monitoring the impact of control measures (Qiao et al., 2011a). The temporal distribution of magnetic concentration parameters in street dust showed a similar trend to the PM variation in the Chaoyang district. Nevertheless, the  $\chi$  value of street dust at some sampling sites during the Olympic period was actually higher than
the value at other times indicating a primary component of street dust originating
mainly from sources in the immediate proximity.

3.3. Comparison of the elemental composition of street dust and PM samples

225 The heavy metal concentration values determined from inductively-coupled plasma-mass spectrometry are listed in Table 2. The order for the mean concentrations 226 of eight heavy metals in the PM is Zn>Cr>Pb>Cu>V>Ni>Fe>Co; zinc and cobalt 227 remain the most and least abundant metals respectively in the street dust but the 228 229 intervening order is somewhat different: Zn>V>Cr>Cu>Pb>Fe>Ni>Co. The relative importance of these elements, particularly the consistent relatively-high amounts of 230 Zn, seems to have no straightforward explanation. Unfortunately vehicle emission 231 232 investigations focus on the gas content and group particulate matter is quoted without constituent metal analysis. To facilitate a comparison, normalized elemental 233 compositions of street dust and PM samples are compared in Table 2 where elemental 234 mass ratios have been normalized with respect to the (mainly crustal-derived) 235 elemental aluminum for clarity. As shown in Table 2, except for Cr and Pb the mean 236 and normalized concentrations of other heavy metals for street dust exceed the PM 237 values, especially the Zn. In part, the elemental concentrations at a specific location 238 will be determined by the distance from their sources and reflect emissions from 239 point-sources emitters. The FT site (Fig. 1) located close to the Jing-Shen Highway 240 was once the highest polluting industrial area of Beijing where Zn and perhaps Pb are 241 more likely to have been retained by refining processes in contrast to the other 242

elements found here in greater abundance. Metallurgical processes have been found to 243 produce the largest emissions of Cu, Ni and Zn (Pacyna, 1998), while exhaust 244 emissions from road vehicles also contain various amounts of these metals (Pacyna, 245 1986; Lee et al., 1999). Johansson et al. (2009) found that vehicle emissions with high 246 loadings of Cu and Zn were an important source of street dust. The mobility of each 247 element will also have a part to play here: with voluminous vehicle and industrial 248 emissions of polluting elements such as Zn and Pb, it is unsurprising that the results 249 identify higher concentrations of these elements compared with the largely-residential 250 251 area SLT to the northwest. The mass concentrations of all 8 elements (Fe, V, Cr, Co, Ni, Cu, Zn and Pb) for street dust in the traffic site are higher than those in the park 252 site and a primary origin in vehicular source emission is obvious; this is especially the 253 254 case for Zn indicating that vehicle exhaust is likely an important source of heavy metals in the Chaoyang District, although the full balance of sources responsible for 255 the high Zn is presently unclear. 256

Since leaded gasoline was phased out in Beijing in 1997, motor exhaust 257 emissions seem unlikely to be the dominant source of Pb in the aerosols unless they 258 result from erosion of engines and exhausts. Liu et al. (2005) suggest that when winds 259 from southern and southwestern directions prevail in Beijing, high concentrations of 260 vapor and air pollutants from power plants, refining industries and biomass burning 261 enhance the aerosol concentrations of Pb and Zn (found together in primary ores). 262 Re-suspended dust could be the possible sources of these metals and an additional 263 factor here is the construction and demolition activity before the Olympic Games 264

resulting in construction dust being an important contributor to PM and street dustprior to commencement of the Games.

The  $I_{\text{geo}}$  values for heavy metals in the PM and street dust from Chaoyang District are listed in Table 3 and plotted in Fig. 5. The Geo-accumulation index results reveal PM contents at the SLT and FT, traffic and park sites where street dust is uncontaminated to only moderately contaminated with respect to Cu, Zn and Pb although the  $I_{\text{geo}}$  value for Zn in the traffic area are especially high and classified as extremely contaminated (Table 3). In contrast the results of  $I_{\text{geo}}$  in the park area indicate little or no metal contamination.

The average  $I_{geo}$  values for heavy metals in PM are -1.15 for Fe, -1.46 for V, 274 -0.39 for Cr, -1.80 for Co, -0.76 for Ni, 0.44 for Cu, 0.96 for Zn and 1.45 for Pb 275 276 respectively. The maximum  $I_{geo}$  values for V, Fe and Co are less than zero, indicating that PM in the study areas is contaminated by other metals. The maximum  $I_{\rm geo}$  values 277 for V, Cr, Fe, Co and Ni in street dust are less than zero and ranked as 278 "uncontaminated". The  $I_{geo}$  values for the other metals such as Cu, Zn and Pb are all 279 greater than zero, indicating that street dust in Chaoyang District is polluted by these 280 metals to varying degrees (Fig.5b). The same phenomena is reported for road dusts 281 from Bulgaria and Greece (Bourliva et al., 2016; Jordanova et al., 2014). Zn in the 282 283 traffic area is recognized as extremely contaminated (c.f. Müller, 1979) whilst in the park area  $I_{geo}$  values are less than zero. As in the previous research (Qiao et al., 2011a) 284 285 the high Zn concentrations in traffic areas is attributable to vehicle emissions, and is probably ultimately sourced in corrosion of the vehicle engines and body work.  $I_{geo}$ 286

values for Cu, Zn and Pb are higher than zero for both PM and street dust samples indicating that dust in the Chaoyang District is contaminated by the metals derived from anthropogenic sources: in contrast the metals V, Fe and Co are evidently not readily shifted into the environment by engine usage. The high Pb fluxes associated with past emissions likely stored in soils that are now being remobilized by surface erosion processes to contribute to PM contamination.

### 293 **4.** Conclusions

Five atmospheric trajectories to aerosols arriving in Beijing during the period 294 295 from June 2008 to October 2008 spanning the interval of the 2008 Olympic Games were identified for application of cluster analysis methodology. Pathways associated 296 with trajectory clusters 4 and 5 passing over Shandong and Hebei provinces before 297 298 reaching Beijing account for 23.03% and 21.05% of all PM. PSCF analysis also identifies high PM values in these provinces as well as along Bohai Bay. Dust 299 particles in Beijing came mainly from distant transport from the south and southeast 300 301 and were then supplemented by local emissions.

The study clearly shows that levels of  $PM_{10}$  and  $\kappa$  were lower during the Olympic period compared with the pre-Olympic period, and there is no doubt that the measures planned to limit air pollution in Beijing greatly improved air quality during the interval of the 2008 Olympic Games although the limited post-Olympic record suggests that major improvement is unlikely to be sustained without the imposition of more restrictive measures. PM and street dust remains heavily contaminated with Cu, Zn and Pb and this is prominently the case for Zn where  $I_{geo}$  attains a value of 3.5 in

the street dust. Even with firm and widespread controls the levels of heavy metal 309 pollution remain high for historic reasons and imply that the limit of zero emissions 310 cannot be achieved in practice. Furthermore, the  $I_{geo}$  values for the street dust are 311 generally higher than those for PM of a given element and district. Thus, street dust 312 tends to be more heavily contaminated than the PM. The distribution of the heavy 313 metal concentration in street dust in the study area indicates that vehicle emissions are 314 mainly responsible for heavy metal pollution as shown by the highest heavy metal 315 concentrations found in the traffic site. 316

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## 440 **Figure Captions:**

441 Figure 1: Schematic map of the study area showing sampling locations of atmospheric442 PM and street dust.

443

444 Figure 2: Cluster-mean back trajectories for Beijing during the period covered by the445 Olympic Games (June 2008 and October 2008).

446

Figure 3: Potential source contribution function (PSCF) map for Beijing PM<sub>10</sub> for the
interval between June 2008 and October 2008.

449

Figure 4: Time series of (a) temperature and relative humidity (RH) between June 2008 and October 2008. (b) Relative humidity and  $PM_{10}$  mass concentration also between June 2008 and October 2008. (c)  $PM_{10}$  mass concentration variation and smoothed magnetic susceptibility ( $\kappa$ ) covering the same interval between June 2008 and October 2008. (d)  $PM_{10}$  mass concentration and relative humidity (RH) in 2008 and 2009 covering the broader interval before, during and after the period of Olympic Games.

457

Figure 5: Geo-accumulation index ( $I_{geo}$ ) for elements (a) in dusts of PM compared with (b) street dust from the Chaoyang District.





Figure 1



Figure 2





Figure 3



Figure 4



471 Table 1

472 The seven classes comprising the Geo-accumulation index.

	Class Value		Particulate matter quality						
	0	$I_{geo} \leq 0$	Practically uncontaminated						
	1	$0 < I_{geo} < 1$	Uncontaminated to moderately contaminated						
	2	$1 < I_{geo} < 2$	Moderately contaminated						
	3	$2 < I_{geo} < 3$	Moderately to heavily contaminated						
	4	$3 < I_{geo} < 4$	Heavily contaminated						
	5	$4 < I_{geo} < 5$	Heavily to extremely contaminated						
	6	5 <igeo< td=""><td>Extremely contaminated</td></igeo<>	Extremely contaminated						
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Table 2 Summary of the heavy metal content, normalized elemental compositions and magnetic susceptibility of atmospheric PM and street dust samples collected in the Chaoyang District. Values are given in mg/kg except for Fe where values are in g/kg. Magnetic susceptibility values are denoted as  $\kappa$  for PM and  $\chi$  for street dust.

Site ID		Fe	V	Cr	Со	Ni	Cu	Zn	Pb	magnetic
Site ID										susceptibility
	Max	26.21	61.32	447.95	8.91	80.28	70.78	283.52	104.55	28.24
SLT(n=3)	Min	19.96	40.21	40.78	3.74	17.01	27.48	226.93	67.16	4.07
	Mean±SD	23.12±3.12	48.59±11.20	179.36±232.65	7.08±2.89	38.24±36.40	46.78±22.03	263.94±32.07	87.82±19.01	12.17
	normalized	0.29	0.61	2.25	0.09	0.48	0.59	3.32	1.1	
	Max	20.29	43.27	238.66	9.24	42.91	67.87	414.03	125.21	72.65
FT(n=4)	Min	16.24	37.69	35.17	4.33	17.33	37.57	266.42	114.26	9.75
	Mean±SD	18.30±1.86	40.02±2.75	92.63±97.61	7.08±2.15	25.25±11.87	53.02±12.62	336.90±78.07	119.67±4.79	25.85
	normalized	0.23	0.50	1.16	0.09	0.32	0.67	4.23	1.50	
	Max	26.21	61.32	447.95	9.24	80.28	70.78	414.03	125.21	72.65
PM(n=7)	Min	16.24	37.69	35.17	3.74	17.01	27.48	226.93	67.16	4.07
	Mean±SD	20.36±3.41	43.70±8.16	129.8±157.97	7.08±2.26	30.82±23.67	50.35±15.89	305.63±70.08	106.02±20.54	19.01
	normalized	0.26	0.55	1.63	0.09	0.39	0.63	3.84	1.33	
	Max	40.70	108.00	88.90	15.00	43.80	146.00	70828.00	114.00	495.98
Traffic(n=5)	Min	34.36	90.00	73.40	12.30	31.30	63.20	15106.00	68.10	284.24
	Mean±SD	37.27±2.35	98.00±7.05	79.65±6.01	14.13±0.98	36.67±4.48	97.13±32.27	32968.17±19695.27	79.30±17.26	391.05
	normalized	0.47	1.23	1.00	0.18	0.46	1.22	414.17	0.99	
	Max	31.25	89.10	58.80	12.10	27.50	47.00	134.00	46.60	72.76
Park(n=5)	Min	27.44	81.90	50.00	11.30	22.10	27.50	72.10	27.10	25.65
	Mean±SD	29.43±1.33	84.78±2.77	54.90±3.06	11.73±0.29	25.25±1.82	36.02±7.98	106.27±24.10	38.70±7.16	56.94
	normalized	0.37	1.07	0.69	0.15	0.32	0.45	1.34	0.49	
Dust(n=10)	Max	40.70	108.00	88.90	15.00	43.80	146.00	70828.00	114.00	495.98
	Min	27.44	81.90	50.00	11.30	22.10	27.50	72.10	27.10	25.65
	Mean±SD	33.35±4.48	91.39±8.58	67.28±13.70	12.93±1.43	30.96±6.79	66.58±38.99	16537.22±21698.84	59.00±24.66	247.86
	normalized	0.42	1.15	0.85	0.16	0.39	0.84	207.75	0.74	

509 normalized mass concentration of element species in PM and street dust, relative to Al

510 The mean values for SLT, FT, traffic and park are means of monthly values at one location; the511 mean values for PM and dust are means of monthly values at different locations

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Site ID V Fe Cr Со Ni Cu Zn Pb SLT(n=3) -0.96 -1.31 -1.78 0.77 -0.08 -0.60 0.29 1.18 FT(n=4) -0.88 -1.29 -1.57 -0.62 -1.83 0.55 1.10 1.65 PM(n=7) -1.15 -1.46 -0.39 -1.80 -0.76 0.44 0.96 1.45 Traffic(n=5) -0.26 -0.28 -0.36 -0.73 -0.26 1.39 7.57 1.03 Park(n=5) 0.0002 -0.60 -0.49 -0.90 -0.99 -0.79 -0.004 -0.57 Dust(n=10) -0.38 -0.63 0.69 0.52 -0.43 -0.86 -0.52 3.50

Table 3 Geo-accumulation indices for elements in PM and street dust