Concentrations and solubility of trace elements in fine particles at a mountain site, southern China: regional sources and cloud processing

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13 Abstract

14 The concentrations and solubility of twelve trace elements in PM2.5 at Mt. Lushan, southern China, were investigated during the summer of 2011 and the spring of 2012. The average PM_{2.5} mass was 15 $55.2 \pm 20.1 \ \mu g \ m^{-3}$ during the observation period. Temporal variations of all trace elements including 16 17 total and water-soluble fractions with several dust storm spikes in total fractions of Al and Fe were 18 observed. The enrichment factor (EF) values were one order of magnitude higher for the 19 water-soluble fractions versus the total fractions of trace elements. Four major emission sources, 20 namely nonferrous metal mining and smelting (for Cr, As, Ba and parts of Zn), coal combustion (for 21 Pb, Zn, Se, Cu and Mn), crustal materials (for Al and Fe) and municipal solid waste incineration (for 22 Cd and Mo), were classified by principal component analysis (PCA). Trajectory cluster analysis and 23 the potential source contribution function (PSCF) consistently identified the Yangtze River Delta 24 (YRD), the Pearl River Delta (PRD), and the neighboring provinces of Mt. Lushan as the major source regions and transport pathways for anthropogenic elements. Northern China was identified as 25

1 a major source region for crustal elements. It should be noted that apart from the YRD, the area 2 around Mt. Lushan has become the most significant contributor to the solubility of most trace 3 elements. Element solubility can be partially determined by emission sources. However, enhanced 4 solubility of trace elements corresponding to increased concentrations of sulphate after the 5 occurrence of cloud events indicated significant effects of cloud processing on aerosol elements 6 dissolution. Metal particles mixed with sulphate in cloud droplet residues were further investigated 7 through transmission electron microscopy (TEM) analysis. Irreversible alteration of particle 8 morphology by cloud processing was confirmed to be highly responsible for the enhancement of 9 trace elements solubility. The findings from this study imply an important role of regional 10 anthropogenic pollution and cloud processing in the evolution of aerosol trace elements solubility 11 during transport in the troposphere.

12 **1** Introduction

13 Epidemiologic studies have associated long-term exposure to highly inhalable ambient fine 14 particulate matter ($\leq 2.5 \,\mu$ m, PM_{2.5}) with many adverse health outcomes such as cardiovascular and 15 respiratory morbidity and mortality, whether in urban or rural areas (Hoek et al., 2013;Cao et al., 16 2012; Weichenthal et al., 2014). Sustained exposure to high concentrations of PM air pollution has 17 been suggested to contribute to a decrease in life expectancy of approximately 3.0 y in China (Chen 18 et al., 2013). The long atmospheric lifetimes of days give fine particles opportunities to be subject to 19 long-range transboundary or intercontinental transport in the air and to deposition toward remote 20 areas, carrying abundant anthropogenic pollutants and affecting ecosystems (Mahowald, 2011). Fine 21 particles are also responsible for regional and global climate change and the hydrologic cycle 22 interference through radiative forcing directly by reflecting sunlight and indirectly by changing cloud 23 properties and precipitation acting as cloud condensation nuclei (Kaufman et al., 2002).

Although trace elements constitute only a small proportion of fine particle mass, their negative impacts on human health and ecosystems have attracted considerable attention because of their toxicity and bioaccumulation by inhalation and deposition. Toxicological and epidemiologic studies often suggest trace metals, particularly water-soluble metals which are more easily bioavailable, as the critical components harmful to the cardiopulmonary system through oxidative chemistry

1 (Cakmak et al., 2014;Costa and Dreher, 1997). Soluble trace elements, especially the transition 2 metals, are likely to be the primary drivers of the generation of reactive oxygen species, inducing 3 cellular inflammation (Charrier et al., 2014; Cheung et al., 2012; Shafer et al., 2010). Besides, trace 4 metal ions also play an important catalysis role in secondary aqueous-phase (Harris et al., 2013) and 5 particle-phase (Clements et al., 2013) sulphate formation, heterogeneous production and elimination 6 of gas-phase hydrogen peroxide (Guo et al., 2014a) as well as multiphase cloud chemistry 7 (Deguillaume et al., 2004). Importantly, most anthropogenic trace elements in ambient air are 8 considered to be mainly concentrated on fine particles, strengthening their toxicity. Therefore, 9 understanding both the concentrations and water solubility of fine particle trace elements is essential 10 for revealing their bioavailability in ambient environments and reactivity in atmospheric chemistry.

11 Diverse characteristics and sources of atmospheric trace elements have been investigated in various 12 areas. For example, regional background, urban and industrial sites in Spain were found to have 13 clearly different concentration levels and source origins of trace elements in PM (Ouerol et al., 2007); 14 technogenic metalliferous fine aerosols observed in urban Montseny demonstrated the 15 spatiotemporal variability of trace metal pollution (Moreno et al., 2011); traffic, static combustion and crustal were suggested as the main sources for Fe, Zn, Pb and Cu in PM_{2.5} in an urban 16 17 background area in Edinburgh (Heal et al., 2005); meanwhile, stationary industrial emissions from 18 coal combustion were confirmed as the contributor to substantial atmospheric lead pollution of PM_{2.5} 19 in Shanghai (Chen et al., 2008). The United States has already established a spatially gridded 20 national emissions inventory of PM_{2.5} trace elements in speciation profiles with the dominant sources 21 of crustal, biomass burning, coal combustion and industrial processing (Reff. et al., 2009). In Asia, 22 atmospheric metallic element pollution by human activities has been surveyed, with the highest 23 metal concentrations occurring in China (Fang et al., 2010). Coal combustion, nonferrous metal 24 smelting and iron and steel manufacturing are found to be the major disproportionate sources of 25 atmospheric hazardous trace elements in China according to the latest anthropogenic emissions 26 inventory, which highlighted an increasing trend in air pollution by toxic heavy metals (Cheng et al., 27 2015).

28 Recently, many field observations have found that even remote mountains and oceans have been

polluted by anthropogenic trace elements in fine particles, significantly confirming the contributions 1 2 of long-range transport (Deng et al., 2011;Schwab, 2004;Fomba et al., 2013). A few studies on dust 3 in Asia (Takahashi et al., 2011;Hsu et al., 2013) indicate that the water solubility of trace elements 4 are different for various emission sources and could be modified further during long-range transport, 5 probably by photochemical aging and cloud processing, etc. (Hsu et al., 2010). However, little 6 information about fine particle trace elements has been reported with regard to the identification of 7 source regions or pathways, and the main mechanism of trace element dissolution by atmospheric 8 processing is still unclear.

9 This paper provides a dataset of the concentration and solubility (defined as the proportion of 10 dissolved element concentration in total content) of trace elements in fine particles at Mt. Lushan, a 11 high, rural mountain within an acid precipitation area in southern China. We focused on identifying 12 the potential source region distributions for individual elements, aiming to evaluate the contributions 13 of local, regional and long-range transport. Regional contributions to element solubility were also 14 preliminarily investigated. One condensation-evaporation cloud cycle can increase aerosol solubility, 15 and atmospheric aerosols may experience up to ten cloud cycles before removal (Desboeufs et al., 16 2001; Spokes et al., 1994); hence, we intensively discussed the influence of actual cloud processing 17 on the alteration of fine particle trace element solubility, considering the rather frequent cloud events 18 at Mt. Lushan. The size, mixing state and chemical composition of individual metal particles in cloud 19 residues were detected using transmission electron microscopy (TEM) to investigate the effects of 20 microphysical properties on the evolution of element solubility.

21 2 Methodology

22 2.1 Site description

Mt. Lushan is one of the National Geoparks of China, with an area of 302 km², and is situated in northern Jiangxi Province, southern China, adjacent to the Yangtze River and Poyang Lake (Fig. 1). Mt. Lushan is significantly influenced by a subtropical monsoon and mountain climate, with frequent cloud events occurring from spring to autumn. The sampling site is located in Guniubei (115°59′E, 29°35′N, 1165 m a.s.l., see Fig. S1), the top of Guling town where most residents work in tourism or related services and little pollution is produced. There are intensive coal-fired power plants located at the eastern coastal area, and abundant mineral resources and many large non-ferrous industries exist in Jiangxi and in neighbouring provinces. Therefore, Mt. Lushan is a favourable platform for observing the long-range transport of regional air pollution and for evaluating the influence of atmospheric particles on the local ecological environment.

6 **2.2 Sample collection**

7 Daily PM_{2.5} samples were collected on quartz fibre filters (MK 360, 90 mm, Munktell, Switzerland) 8 during the summer of 2011 (8 August-23 September) and the spring of 2012 (18 March-20 May). 9 The filters were heated at 500 % for 3 h prior to sampling, subsequently balanced at 25 % and 50 \pm 10 5% relative humidity for at least 24 h and weighed three times before and after collection. The 11 sampling procedures were conducted during non-rainy and non-cloudy periods by employing a PM_{2.5} impactor (TH-150A, Tianhong Co., China) with a medium flow rate of 100 L min⁻¹. Seventy-six valid 12 13 ambient samples and 6 blank filters were obtained and stored in the dark and at -20 °C for laboratory 14 analysis.

15 Cloud water were collected by a improved single-stage Caltech Active Strand Cloudwater Collector 16 (CASCC2, Demoz et al., 1996). Air with cloud droplets are inhaled into the instrument with a sampling flow rate of 24.5 m³ min⁻¹ and impacted on vertical 508 µm diameter Teflon strands. The 17 18 droplets are then collected into a Teflon sample trough along the strands and flows into a 500 mL 19 high-density polyethylene bottle. The theoretical sampling efficiency for size cut of 3.5 µm droplet 20 diameter corresponds to 50%. In this study, the sampling interval of cloud water was 2–3 h. The 21 samples were immediately filtered and stored in brown glass bottles at 4 °C with preservation of 1% 22 v/v high pure hydrochloric acid.

The sampling system of cloud residues (Fig. 2, Li et al., 2011) was employed to obtain individual cloud residues in cloud events on 11 September 2011 and 18 April 2012. The moisture of the cloud droplets is removed by a diffusion dryer filled with desiccant (silica gel) to prevent the impact of the water vapor on TEM grids. Then, the individual residues are collected onto TEM grids using a single-stage cascade impactor with a 0.5 mm diameter jet nozzle. The calculated effective size d_{50} for cloud droplet is about 0.7 μ m and the collection efficiency is estimated to be 5-10%. For cloud residues at Mt. Lushan, a low sampling flow rate of 0.5 L min⁻¹ and an optimum sampling time of about 12 min were applied, neither destroying the TEM grids nor overlapping each individual residues. An optical microscopy with magnification from ×500 to ×1200 was operated on site to check the integrity of carbon film and particle distribution on the TEM grid. If suitable, the grids were sealed in a dry plastic tube and stored in desiccators at 25 °C and 20 ± 3% RH until the laboratory analysis. Otherwise, another satisfactory sample shall be collected and preserved.

8 2.3 Extraction procedure

9 2.3.1 Total fraction (Acid)

10 To obtain the total concentration of trace elements in fine particles with the least loss of semi-volatile 11 elements, one quarter of the samples and blank filters were digested with an acid mixture (5 mL 65% HNO₃ and 2 mL 30% H₂O₂, guaranteed reagent grade, Kemiou Co., China) in closed Teflon vessels 12 13 using a microwave digestion system according to a controlled gradient temperature procedure: 14 reaching 120 $^{\circ}$ C in 5 minutes and remaining there for 5 minutes, then reaching 160 $^{\circ}$ C in 5 minutes 15 and remaining there for 2 minutes, and finally reaching 185 °C in 5 minutes and remaining there for 16 15 minutes. After cooling to room temperature, the solutions were filtered and subsequently diluted 17 to 25 mL with high-purity deionised water (\geq 18 M Ω ·CM) in acid-cleaned brown glass bottles.

18 **2.3.2 Water-soluble fraction (Water)**

Half of the samples and blank filters were extracted with 50 mL high-purity deionised water (≥ 18 M Ω ·CM) using ultra-sonication for 1 h followed by filtration. Solutions of approximately 25 mL were preserved in acid-cleaned brown glass bottles with 1% v/v high pure hydrochloric acid to lessen adsorption of the elements by the glass walls. All solutions were stored at 4 °C until instrument analysis.

24 **2.4 Chemical and physical analysis**

25 The concentrations of the total and water-soluble fractions of twelve trace elements (including Al, Cr,

Mn, Fe, Cu, Zn, As, Se, Mo, Cd, Ba and Pb) were measured by Inductively Coupled Plasma Mass 1 2 Spectrometry (ICP-MS, Agilent 7500a) based on the EPA 200.8 method using internal standard 3 substances of Li, Sc, Ge, Y, In, Tb and Bi to eliminate matrix interference. National standard 4 materials (soil, GSS-4, China) were digested for calculating element recoveries by the acid extraction 5 method. All recoveries were found to be a little lower than 100% (Table S1), ranging from 93.0% for 6 Cd to 72.7% for Se, primarily ascribed to volatilisation losses and acid mixture that is not strong 7 enough to release all mineral constituents from the silica matrix into the solution. Reagent and filter 8 blanks were also prepared and analysed for background element content. The detection limits of the 9 ICP-MS analysis for the total fraction elements determined were calculated as three times the 10 standard deviation (3δ) of the six blank values (Table S1 in the Supplement). Trace elements 11 concentrations in cloud water samples were similarly determined by ICP-MS.

Individual metal particles in cloud residues on the TEM grids were analysed with a JEM-2100 TEM operating at 200 kV accompanied by a semi-quantitative determination of elemental composition by an energy-dispersive X-ray spectrometer (EDS) that can detect elements heavier than carbon. EDS spectra were collected for only 15 s to minimise radiation exposure and potential beam damage. Copper could not be analysed because of interference from the copper TEM grid.

17 2.5 Meteorology

Meteorology parameters during the observation periods and two cloud events were obtained from the local meteorological station (listed in Table S2). Generally, summer days exhibited higher temperature, relative humidity and wind speed than spring. In particular, the cloud events were characterized by high humidity (>99%), low wind speed (about 1.0 m s⁻¹) and bad visibility (nearly 0 km).

23 2.6 Backward trajectories and potential source contribution function (PSCF)

The Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model (PC Version 4.8) developed by the National Oceanic and Atmospheric Administration Air Resources Laboratory (NOAA-ARL) (Draxler and Rolph, 2014) was used to reconstruct the three-dimensional backward trajectories for the purpose of identifying the possible impacts of long-range transport. Three-day air
mass backward trajectories initiated every 12 h arriving at Mt. Lushan at the height of 1,165 m a.s.l.
were calculated, and five mean trajectory clusters were classified.

The potential source contribution function (PSCF) analysis developed by (Hopke et al., 1995) was computed using a geographical information system based software TrajStat (Wang et al., 2009) with the calculated backward trajectories and measured atmospheric pollutant concentrations, representing the conditional probability that air parcels may be responsible for concentrations higher than the criterion level during transport to the receptor site. The PSCF value for the *ij*th grid cell is defined as

9
$$\operatorname{PSCF}_{ij} = \frac{m_{ij}}{n_{ij}}$$
(1)

10 where n_{ij} is the total number of trajectory segment endpoints that fall in the *ij*th cell and m_{ij} is the 11 number of endpoints for the same cell with arrival times at the sampling site corresponding to 12 pollutant concentrations higher than an arbitrary criterion value. In this study, the total number of 13 endpoints was 18,998, and the geophysical domain (90-130 °E, 15-45 °N) was divided into 4,800 grid cells with a 0.5 $^{\circ} \times 0.5$ $^{\circ}$ resolution. The average concentrations were set as criteria for PM_{2.5} and 14 each trace element, except for Al, for which the 75th percentile was better to distinguish between 15 16 larger versus moderate regional sources. The PSCF values were multiplied by an arbitrary weight 17 function W_{ij} to reduce the effect of small values of n_{ij} and to better reflect the uncertainty in values 18 for these cells (Polissar et al., 2001). The weighting function reduced the PSCF values when the total 19 number of the endpoints in a particular cell (n_{ij}) was less than approximately three times the average 20 value (n_{Ave}) of the endpoints per each cell, defined as

$$21 \qquad W_{ij} = \begin{cases} 1.00 & n_{ij} > 3n_{Ave} \\ 0.70 & 1.5n_{Ave} < n_{ij} \le 3n_{Ave} \\ 0.42 & n_{Ave} < n_{ij} \le 1.5n_{Ave} \\ 0.17 & 0 < n_{ij} \le n_{Ave} \end{cases}$$

(2)

1 **3** Results and discussion

2 **3.1 Elemental composition**

3 **3.1.1 General characterisation**

The ambient atmospheric concentrations of PM2.5 and trace elements including total and 4 5 water-soluble fractions at Mt. Lushan are summarised in Table 1, and the comparison with typical mountains and megacities in China are listed in Table S3. The average PM_{2.5} concentration at Mt. 6 Lushan in spring (54.7 μ g m⁻³) was approximately that of summer (55.9 μ g m⁻³), more than twice the 7 WHO air quality guideline for daily $PM_{2.5}$ of 25 µg m⁻³, but still lower than many urban sites at 8 ground level such as Beijing (118.5 μ g m⁻³) and Guangzhou (81.7 μ g m⁻³) (Yang et al., 2011). It was 9 clearly contrary to Mt. Tai (Deng et al., 2011), where PM_{2.5} exhibited a concentration 2-3 times 10 higher in summer (123.1 μ g m⁻³) than in spring (46.6 μ g m⁻³), suggesting a less-pronounced seasonal 11 12 variation of PM_{2.5} at Mt. Lushan.

13 The total fraction concentrations of twelve trace elements in spring were comparable with those in summer at Mt. Lushan, with the exception of higher Al in spring. Overall, elements Al and Fe 14 contributed to the highest concentration, and the other elements decreased from 258.3 ng m^{-3} for Zn 15 to 2.0 ng m⁻³ for Mo. The dominant elements (Al, Fe and Zn) exhibited significantly lower 16 concentrations at Mt. Lushan than Mt. Tai (1200 ng m⁻³ for Al, 810 ng m⁻³ for Fe and 400 ng m⁻³ for 17 Zn) (Deng et al., 2011) and slightly higher than Mt. Gongga (295.8 ng m⁻³ for Al, 224.0 ng m⁻³ for Fe 18 and 154.6 ng m⁻³ for Zn) (Yang et al., 2009a), probably due to the geomorphology and climatology 19 20 biases. As shown in Table S3, all elements (except Ba) have lower concentration levels in 21 comparison with Mt. Dinghu, a national nature reserve influenced by atmospheric pollution (Yang et 22 al., 2009b) and with typical megacities in China such as Beijing, Guangzhou (Yang et al., 2011) and Shanghai (Chen et al., 2008). For water-soluble fractions, Zn transformed into the primary elements, 23 24 followed by Al and Fe, with higher concentrations in summer than spring; other trace elements (from 25 Pb to Mo) were still comparable between summer and spring.

1 3.1.2 Temporal variations

2 Figure 2 displays the temporal variations of trace element concentrations in PM_{2.5} at Mt. Lushan for 3 total and water-soluble fractions during the sampling campaigns. Most total fraction elements 4 exhibited more intense variances in spring than in summer, with unique temporal patterns such as the 5 highly variable pattern observed for aluminium and the relatively steady pattern of arsonium. Similar 6 temporal patterns were observed for the water-soluble and total fractions for most elements, except 7 for Al and Fe, which had drastic variations with many spike episodes for total fractions but lower 8 constant concentration for water-soluble fractions. The dependence of temporal variations of 9 water-soluble elements on their total fractions were also examined (Fig. S3). The concentrations of water-soluble Zn, Cu, Mn, As, Se and Cd were found to be rather consistent with their total fraction 10 according to their high correlations ($R^2 = 0.50-0.78$), whereas there was only a weak association 11 12 between the total concentrations and water-soluble fractions of Al, Fe and Cr.

13 Several spikes in element concentration were observed in specific periods. The first spike (Spike I) 14 for all elements from 11-13 August 2011 was ascribed to the short sampling time due to the 15 interruption by an unexpected thundershower. However, the significantly elevated total concentration 16 of Al, Fe and Mn during 23-25 March 2012 (Spike II) and 25-26 April 2012 (Spike III) with no 17 distinct elevation in the other elements were observed (Fig. 2), likely contributed by the dust storms 18 originating from the Gobi desert and the Taklimakan desert in spring according to the MODIS image, 19 the OMI Aerosol Index and air mass backward trajectory analysis (Fig. S4). These features were 20 similar with the Asian Dust observed at Huaniao Island in the East China Sea, where concentrations 21 of Al, Fe, Mn and Ba 3-4 times higher were observed with no significant difference for Zn, Cu, As 22 and Cd in the air compared with non-dust days (Guo et al., 2014b). In addition, the dust storms 23 contributed little to the water-soluble fractions of all trace elements.

24 **3.2 Source identification**

25 **3.2.1 Correlations between individual elements**

26 The Pearson correlation coefficients *r* between each element for both total fraction and water-soluble

1 fraction were examined (Table S4). Moderate significant associations of Fe with elements Al, Mn 2 and Cr are found to exist both in total and water-soluble fractions with comparable coefficients 3 between 0.40–0.68. Correlations for other trace elements in water-soluble fraction are stronger than 4 in total fraction, suggesting the higher possibilities of common sources for some water-soluble 5 elements. For example, the correlation levels of Zn-Cu, Zn-Se, Pb-Zn and Pb-Se are dramatically 6 high for water-soluble fractions (0.72 < r < 0.82) and moderate for total fractions (0.25 < r < 0.52), 7 which could be attributed to their common anthropogenic sources of coal-fired power plants and 8 industrial processes based on the suggestion of the spherical morphology and elemental composition 9 of individual metal particles via the TEM examination (Li et al., 2014).

10 **3.2.2** Anthropogenic pollution

11 The enrichment factor (EF) for individual trace elements was applied to identification of their 12 general crustal and anthropogenic sources and evaluate the degree of pollution using the following 13 formula:

14
$$\operatorname{EF}_{i} = (X_{i}/X_{R})_{\operatorname{aerosol}}/(X_{i}'/X_{R})_{\operatorname{crust}}$$
 (2)

where EF_i is the enrichment factor of element *i*; X_i and X_R are the concentrations of element *i* and 15 reference element R in aerosol, respectively; X_i and X_R are the background contents of elements in 16 17 Chinese soils (Wei et al., 1991); and Al is selected as the reference element for calculation. Element sources are classified into three groups with the following standard: EF < 10 is considered to be a 18 19 crustal origin without enrichment; 10 < EF < 100 comes from mixed origins (crustal and 20 anthropogenic sources); EF > 100 indicates air pollution from an anthropogenic origin. Figure 3 21 describes the EF values for both the total and water-soluble fractions of each element in a decreasing 22 order. Elements with EF > 100, including Se, Cd, Zn, Pb, As, Mo and Cu, were found to be highly 23 enriched, ranging from hundreds to tens of thousands in both fractions, indicating the severe 24 anthropogenic pollution of atmospheric trace elements. Cr, Ba and Mn were likely to be from mixed 25 origins because the majority of their EF values fell within 10-100. Fe was predominantly from a crustal origin based on its EF < 10. It is worth noting that the water-soluble fractions of most 26 27 elements (except Cr and Fe) have higher EF values than the total fractions by approximately one 1 order of magnitude.

2 **3.2.3 Emission source classification**

The principal component analysis (PCA) was performed in detail to classify the main sources of trace elements in PM_{2.5} at Mt. Lushan. The PCA results for trace elements in total fraction are shown in Fig. 4 with four major components identified.

6 The first principal component was considered primarily from nonferrous metal mining and 7 metallurgical smelting, accounting for 34.4% of the total variance with high loadings for Cr, As, and 8 Ba and considerable loadings for Zn and Mo. Although there were diverse EF values which 9 suggested anthropogenic origins mixed with crustal materials ranging from dozens for Cr and Ba to 10 thousands for As, Zn and Mo, these total fraction elements still presented considerably moderate 11 correlations. According to the provincial output statistics of nonferrous metals in 2010 (Cheng et al., 12 2015) and the Overall Planning of Mining Resources of 2008–2015 for Jiangxi and adjacent Hunan 13 Province, there are abundant nonferrous minerals (such as barite, realgar and sphalerite, etc.) and 14 many large-scale metallurgical industries within and near Jiangxi Province. In addition, high 15 concentrations of Pb and Zn in cloud drop residues at Mt. Lushan were suggested deriving from fine 16 particles contributed by nonferrous smelting in Jiangxi Province (Li et al., 2013).

17 The second principal component that contributed to high loadings for Pb, Cu, Se, Zn, and Mn was 18 identified as coal combustion. Coal combustion is representatively considered to be the source of 19 most atmospheric trace elements in China. All species in this component except Mn exhibited high 20 EF values, indicating their anthropogenic origins. The semi-volatile Se is usually used as a tracer for 21 determining atmospheric sources and even the pathways of coal combustion pollutants, because it is 22 primarily produced in high temperature combustion and can undergo long-range transport with fine 23 particles after the rapid gas-to-particle conversion (Husain et al., 2004;Wen and Carignan, 2007). 24 Coal combustion is also becoming the major contributor to atmospheric Pb with the nationwide phase-out of leaded gasoline in China (Xu et al., 2012). The high correlations between Se and the 25 other species in Component 2 clearly indicated the likely sources of coal combustion. 26

27 The third component in particular explained most of the Al and Fe which are the typical signatures of

1 crustal materials or soils, along with relatively high correlations and low EF values.

Municipal solid waste incineration was regarded as the primary sources of fine particle Cd and Mo at Mt. Lushan, explaining 8.4% of the variance. Both Cd and Mo had low concentrations but high EF values, with a strong association in total fraction (r = 0.71). Jakob et al., (1995) found that Cd in fly ash of municipal solid waste incinerators could evaporate into the atmosphere completely during heat treatment. The rapid increase in atmospheric Cd emissions from municipal solid waste incineration in recent years and the unbalanced large emissions in east-central and southeast China (Tian et al., 2012) were responsible for fine particle Cd at Mt. Lushan.

9 **3.3** Potential source region distributions

10 Three-day air mass backward trajectories were calculated to estimate the potential source regions and 11 pathways of air pollutants, making full use of the geographical advantages of the high-altitude 12 observation site and the long-range transport capacity of fine particles. The air masses associated 13 with PM_{2.5} samples were classified into five clusters (Fig. 5a). The air masses of the Southwest (SW) 14 cluster originated from the South China Sea and passed through the Pearl River Delta (PRD), Hunan and Jiangxi Province with the highest occurrence frequency of 28.9%. Cluster East (E) mainly came 15 16 from the East China Sea and partly from the Circum-Bohai Sea Region, traversing the Yangtze River 17 Delta (YRD) and Anhui Province at a frequency of 27.6%. Air masses of cluster Local (L) were 18 found to be advancing toward or revolving around Mt. Lushan slowly from ground-level within areas 19 covering distances of approximately 200 km away from the observation site, contributing 22.4% of 20 the occurrence frequency. Clusters Northwest (NW) and North (N) both went through the China 21 Loess Plateau and accounted for 13.2 and 7.9% of the occurrence frequency, respectively. The 22 primary difference was that the former mainly originated from Taklimakan and the Gobi Desert, 23 whereas the latter was from remote Siberia and Mongolia with a much higher wind speed.

Figure 5b shows the averaged $PM_{2.5}$ concentration of different clusters. Cluster E exhibited the highest $PM_{2.5}$ concentration of 67.0 µg m⁻³, followed by cluster L (59.6 µg m⁻³) and the similar clusters NW and N, whereas cluster SW had the lowest concentration (45.3 µg m⁻³). The order of $PM_{2.5}$ concentration seemed to be consistent with the PSCF result (Fig. 5c) which integrated $PM_{2.5}$

1 concentration with air mass occurrence frequency. As shown in Fig. 5c, the YRD and southern Anhui 2 Province were the most likely source regions of $PM_{2.5}$ at Mt. Lushan. These economically developed 3 regions exhibit massive energy consumption and industrial production, becoming important 4 anthropogenic source regions in eastern China. Local Jiangxi, the neighbouring southern Anhui and 5 eastern Hunan were also significant PM_{2.5} source regions, where many nonferrous metal smelting 6 industries are widely distributed. However, the contributions to PM_{2.5} at Mt. Lushan by long-range 7 transport of air masses from northern China (Inner Mongolia and the Loess Plateau) were much less 8 than from eastern China and local regions, suggesting nearly negligible impacts of mineral dusts or 9 soil transport on fine particle mass concentration at Mt. Lushan compared with regional 10 anthropogenic pollution. In addition, the PM_{2.5} concentration in five clusters or different source 11 regions were highly dependent on the inorganic ions, especially the secondary ions (Li et al., 2014). 12 Interestingly, the PRD contributed much less concentrations to fine particles in spite of its high 13 occurrence frequency, probably due to particle scavenging by frequent rainfall during the Asian 14 summer monsoon.

15 Figure 6a-l illustrates the distinctive potential source region distributions for individual elements in 16 fine particles at Mt. Lushan. Crustal elements Al and Fe exhibited similar source region distributions 17 (Fig. 6a and b). The highest PSCF values in northern China including the Loess Plateau, Henan and 18 eastern Hubei Province indicated that these areas were very important source regions and pathways 19 for fine particle Al and Fe, with vast contributions of long-range transport of mineral dusts or soils. 20 The PRD and eastern Hunan Province were likely to be other important source regions for Al and Fe 21 probably due to the influence of mining activities, whereas the YRD in eastern China and the local 22 Jiangxi Province made much lower contributions.

Fine particle As (Fig. 6c) had an almost parallel pattern of source region distribution with Fe, except the discordance that the PRD, eastern Guangxi, Hunan and Jiangxi Province were identified as the most important source regions and pathways, attributable to the emissions of the nonferrous metals mining and smelting industries. Northern China also tended to be an important potential source region, suggesting the possible contribution of natural sources for As. Similar to As, the PRD, Hunan and Jiangxi Province, as well as northern China were also potential source regions for fine particle Mn (Fig. 6d). However, the YRD area was characterised by extremely high PSCF values for Mn,
 indicating an important anthropogenic source region.

3 Fine particle Pb, Zn, Cu and Se (Fig. 6e-h), which were identified primarily from coal combustion 4 by the PCA analysis, had similar source region distributions with some discrepancies, generally 5 coinciding with the geographical distribution of coal-fired power plants in China which were 6 aggregated within coastal areas such as the YRD, the PRD and Shandong Province (Fig. S5a). The 7 YRD area including Jiangsu, Zhejing and southern Anhui Province was an especially pronounced 8 source region for Pb, Zn, Cu and Se at Mt. Lushan. The significantly high PSCF values in the YRD 9 region were attributable to the large coal consumption by intensive power plants in Jiangsu, Zhejiang 10 and even Shandong Province, and to the differences in trace element contents in raw coal such as 11 high Se contents in Anhui Province, both of which made great contributions to emissions of 12 atmospheric trace elements (Tian et al., 2014). Meanwhile, the PRD, Hunan and the local area of 13 Jiangxi Province also played a remarkable role in source region distributions of Zn, Cu and Se at Mt. 14 Lushan, for which coal-fired power plants and nonferrous metals smelting industries in these areas 15 were primarily responsible. Unexpectedly, northern China rather than the PRD area was also identified as a potential source region of fine particle Pb, possibly attributed to the partial 16 17 contributions of long-range transport of mineral dusts.

18 As shown in Fig. 6i and j, the YRD, the PRD, eastern Hunan and western Jiangxi were identified as 19 the common likely source regions and pathways for fine particle Cd and Mo at Mt. Lushan. The 20 identified source regions corresponded very well with the uneven regional distribution of municipal 21 solid waste (MSW) incineration plants in China (Fig. S5b), with 183.5, 175.1 and 146.6 kg of 22 atmospheric Cd emitted by MSW incineration from Jiangsu, Zhejiang and Guangdong, respectively, 23 the top three largest emitting provinces in 2010 (Tian et al., 2012). Figure 6k shows an apparently 24 more homogeneous source region distribution for Ba, with the PRD and areas surrounding Mt. 25 Lushan as potential source regions which have a high natural background content of barite. On the 26 contrary, few potential source regions were identified for Cr (Fig. 61) besides the PRD, parts of 27 Hunan Province and a small region in central Zhejiang with low PSCF values.

28 It should be noted that the potential source region distributions for trace elements at Mt. Lushan were

1 very different with PM_{2.5}. The YRD, the PRD and Hunan Province were likely to be the major source 2 regions for most anthropogenic elements (except Cu within eastern Jiangxi), and northern China was 3 likely for crustal elements (Al and Fe). However, the contribution probabilities of areas near Mt. 4 Lushan for most elements were obviously lower than other source regions. This result revealed that 5 long-range transport and regional anthropogenic sources, rather than local sources, made much more 6 significant contributions to fine particle trace elements at Mt. Lushan, which could be ascribed to the 7 different types and quantities of emissions as well as the protection policies for Mt. Lushan and its 8 adjacent Poyang Lake.

9 In addition, the order of mass concentration of individual trace elements for five air mass clusters 10 ending at Mt. Lushan (Fig. S6) was in accordance with the distributions of likely source regions 11 identified by PSCF values, further suggesting greater contributions by regional sources and 12 long-range transport to fine particle trace elements at Mt. Lushan. For example, clusters N and NW 13 contributed to the highest concentration for crustal Al and Fe and anthropogenic Zn and As, followed 14 by the highest cluster SW, E and the lowest cluster L; exceptionally, cluster L made the highest 15 contribution to the combustion element Cu.

16 **3.4 Trace element solubility**

17 **3.4.1** Comparison of individual element solubility

18 In this study, element solubility is calculated as the percentage of water-soluble concentration of an 19 element divided by its total concentration. Figure 7 compares the average water solubility of 20 individual elements in PM_{2.5} at Mt. Lushan with other sites in decreasing order. The most soluble 21 elements were As, Mn, Cu, Zn and Se with a solubility reaching up to 60-70%; followed by Cd, Pb, 22 Ba and Mo with moderate solubility, approximately 40-50%; whereas the least soluble elements were Fe, Al and Cr with a solubility less than 30%. Compared with TSP over the East China Sea 23 24 (Hsu et al., 2010), some elements (Mn, Cu, Ba, Mo, Fe and Al) were significantly highly soluble in PM_{2.5} at Mt. Lushan, whereas other species (As, Zn, Se and Cd) showed the opposite with the 25 exception of comparable Pb and Cr. Similarly, compared with the PM_{2.5} at Edinburgh, UK (Heal et 26

al., 2005), the elements Mn, Cu and Fe exhibited higher water solubility, whereas the elements As,
Zn and Pb were comparable, and the elements Cd and Cr were lower at Mt. Lushan. A large gap in
anthropogenic elements (As, Mn, Cu and Pb) between Mt. Lushan and Nanjing (Hu et al., 2012) was
also observed. Experimental conditions (compared in Supplement, Table S5) such as digestion
methods or analysis instruments may result in some solubility discrepancies at different sites, but
they are far from the determinant.

7 3.4.2 Effect of emission sources on element solubility

8 Element solubility is associated with four major emission sources identified by EF value and PCA 9 analysis and displayed in Fig. 8. Crustal elements with mean EF < 10 had low average solubility (< 10 30%) besides the highly dissolved Mn that was possibly impacted by a combustion source; the 11 solubility of elements related to mining and metallurgy varied between < 10% - 70% with a 10-1000 12 range in EF value; coal combustion elements characterised by high EF values over 100 exhibited a 13 high solubility of approximately 45%–75%, contributable to the combustion process during which 14 oxidisable elements in coal are mostly altered to easily soluble species (Quispe et al., 2012); 15 elements produced by waste incineration were also highly soluble (40%-50%) coupled with EF > 16 100. The roughly increasing tendency of solubility along with EF value suggested that element 17 solubility was closely related to the emission sources, likely indicating that crustal elements with low 18 EF values often showed low solubility, whereas polluted elements featuring high EF values were 19 usually highly soluble. This result coincides well with the conclusion that the dissolvable portion of 20 aerosol elements is controlled by their dominant crustal or anthropogenic origins (Hsu et al., 2005). 21 However, there are also some exceptions that could not be explained by emission sources. For 22 instance, the low solubility of atmospheric Cr at Mt. Lushan could probably be ascribed to the stable 23 crystalline structure in minerals such as chromite, with the reduced Cr (III) being the dominant 24 species in ambient fine particles (Werner et al., 2007). Figure S7 further shows a hyperbolic or 25 negative logarithmic relationship between element concentration and the solubility for and only for Al, Fe, Cr and Mn, similar to many inverse relationships for aerosol Fe, Al and other species 26 27 reviewed by (Schulz et al., 2012), indicating the unneglected effect of total concentration on crustal trace element solubility rather than anthropogenic elements. Thus, trace element solubility may be
 initially determined by emission sources with various interferences.

3 **3.4.3** Source region contributions to element solubility

Figure 9 illustrates the potential source regions for individual element solubility in PM_{2.5} at Mt. Lushan with the expanding application of PSCF analysis to evaluate the local and regional contributions. It can be seen that almost all of the elements had quite different solubility distributions with their total concentrations, characterised by the highest PSCF values for element solubility in local Jiangxi and the YRD region followed by lower PSCF values in parts of the PRD region and Hunan Province.

10 In this study, we used fine particle Fe for detailed explanation because there are many investigations 11 on the solubility of aerosol Fe, which plays a critical role in the global iron cycle among terrestrial 12 dust, the atmosphere and the ocean, as well as in the effects on climate (Jickells et al., 2005). As 13 shown in Fig. 9a, northern China, identified as the major source region for crustal Fe, made little 14 contribution to its solubility. The significantly low solubility of fine particle Fe in northern China, 15 much lower than the PSCF criteria value, was consistent with the very low soluble Fe (solubility less 16 than 2%) from desert dusts or soils (Baker et al., 2006; Jickells et al., 2005). The experiment 17 performed by (Mackie et al., 2006) suggests that the amount of readily released Fe does not increase 18 after the uplift and abrasion of soils in areas with little atmospheric pollution; thus, the low solubility of the Fe from northern China could be attributed to the initial absence of dissolved Fe in dusts or 19 20 soils derived from the Gobi desert or Loess Plateau. On the contrary, the relatively much higher 21 PSCF values for Fe in the local Jiangxi, southern Anhui and the YRD indicated greater contributions 22 of anthropogenic pollution in these regions to the solubility of fine particle Fe at Mt. Lushan, because 23 the high solubility of Fe in the YRD and Anhui Province was just in accordance with the large 24 amounts of anthropogenic emissions of SO_2 and NO_x in these regions (Zhang et al., 2009), which 25 could significantly promote the dissolution of aerosol Fe acting as acid precursors (Schulz et al., 2012;Solmon et al., 2009). A significant correlation (0.629) of iron solubility and sulphate 26 27 concentration in our fine particle samples was further verified, similar to the correspondence of high iron solubility to the sulphur content in ambient fine particles observed by (Oakes et al., 2012).
Moreover, the outstanding enhancement of fine particle Fe solubility in local regions close to Mt.
Lushan was principally recognised to be an effect of cloud processing, an important atmospheric
process controlling Fe dissolution in aerosols (discussed in Sect. 3.4.4). The regional distributions for
the solubility of crustal Al was similar with Fe, except for the lower PSCF values near Mt. Lushan
(Fig. 9b).

7 Anhui, Jiangsu and Jiangsi Province were identified as the most important contributors to the high 8 solubility of fine particle As at Mt. Lushan (Fig. 9c), ascribed to the large amount of atmospheric As 9 emissions from anthropogenic sources (approximately 200 tons in 2010) in these provinces (Cheng 10 et al., 2015); in contrast, little contributions were made to As solubility by the PRD and Hunan 11 Province where the highest As concentrations were provided. For the elements from coal combustion 12 (including Mn, Pb, Zn, Cu and Se, Fig. 9d-h) and MSW incineration (including Cd and Mo, Fig. 9i 13 and j), the highest contributions of element solubility were uniformly focused within Jiangxi 14 Province, the YRD and parts of Fujian Province, generally coinciding with their significantly high 15 concentrations emitted from anthropogenic sources in eastern China. The solubility pattern for Ba 16 (Fig. 9k) was rather similar to its concentration distribution, whereas the pattern of high solubility for 17 Cr in Anhui Province (Fig. 91) was very different to the concentration pattern.

18 It can be concluded that the solubility of fine particle trace elements from local and regional source 19 regions were much higher than that by long-range transport from south-western China and northern 20 China. The high contributions for particle element solubility in the YRD region should be attributed 21 to the serious air pollution. However, on account of the relatively less air pollution released from 22 local sources, the frequent cloud processing was considered as the significant contributor to the high 23 solubility of trace elements in local regions near Mt. Lushan.

Additionally, the statistics on the solubility of individual fine particle elements at Mt. Lushan for five air mass categories are shown in Fig. S8. In general, the fine particle elements derived from cluster L exhibited the highest solubility whether for crustal or anthropogenic elements, closely followed by cluster E; elements from cluster SW showed moderate solubility, ascribed to the wash-out of soluble constituents by frequent rainfall in summer monsoon; elements from cluster NW and N were the least soluble, primarily impacted by crustal materials from northern China. It was demonstrated that
 the statistical results for most elements fairly agreed with the solubility distributions identified by
 PSCF analysis.

4 3.4.4 Evolution of element solubility during cloud processing

5 Cloud processing might be of great importance in affecting aerosol dissolution at Mt. Lushan where 6 frequent cloud events occurred. To determine the evolution of element solubility during cloud 7 processing, cloud events were selected with the following conditions: (1) non-raining cloud to avoid 8 scavenging of particles by precipitation, (2) short duration and time interval of cloud processing and 9 particle sampling to minimise the interference of external aerosols, (3) identical air mass trajectory of 10 the cloud and particles with low wind speed to ensure consistent origins of the air masses. Figure 10 11 and Table S6 show the alteration of individual element solubility in fine particles after two selected 12 cloud events on 11 September 2011 and on 18 April 2012.

13 It was evident that after cloud processing on 11 September 2011, most fine particle elements 14 exhibited a drastic increase in solubility. For instance, post-cloud particle Al became much more 15 soluble compared with the pre-cloud particle, with solubility changing from 1.7 to 28.7%. Other 16 elements, such as Cu, Zn, As, Se, Cd and Ba, also showed an approximately one- to two-fold 17 increase in solubility. The cloud processing on 18 April 2012 also brought about a significantly large number of solubility increments for the crustal elements Al and Fe. The less increments for 18 19 anthropogenic elements such as Cu, As, Se and Pb should be due to their already higher solubility in 20 pre-cloud particles (Table S6) which might have undergone intensive cloud cycles during the 21 prolonged residence time in the air. Generally, the evidently higher solubility for the post-cloud trace 22 elements after the two cloud events, overpassing their overall average solubility, corroborated the 23 remarkable dissolution efficiencies of aerosol elements by cloud processing at Mt. Lushan.

Interestingly, the increase in aerosol element solubility after the occurrence of cloud processing was coupled with the elevation of aerosol sulphate concentration as shown in Fig. 10 and Table S6. For example, post-cloud sulphate particles (4 h) exhibited a sharp elevation of 10 μ g m⁻³ after the cloud processing (3 h) on 11 September 2011. Based on the very short intervals between cloud events and

post-cloud particle sampling, the increased sulphate particles were most likely contributed by cloud 1 2 water sulphate, which could be generated by the rapid atmospheric aqueous-phase oxidation of S(IV) with a rate greater than 100% h⁻¹ (Seinfeld and Pandis, 2012), much quicker than the gas-phase 3 oxidation of SO₂ with a rate of approximately 1% h⁻¹ (Newman, 1981). In addition, the higher 4 5 sulphate concentrations in cloud water and post-cloud particles on 11 September 2011 (41.70 mg L⁻¹ and 24.72 µg m⁻³, respectively) from cluster E compared with those on 18 April 2012 (26.78 mg L⁻¹ 6 and 17.34 μ g m⁻³, respectively) from cluster L were likely to be associated with the larger amount of 7 8 anthropogenic SO₂ gases from the YRD in eastern China because acidic substances, such as sulphate 9 formed by their precursor gases (e.g., SO₂), are prone to exist in fine particles and lead to 10 acidification of aerosols (Ren et al., 2011); and the pH of cloud droplets modified by these dissolved 11 acidic substances in particles could in turn influence the dissolution of aerosol trace elements 12 (Deguillaume et al., 2005). As a consequence, the enhancement of trace element solubility during 13 cloud processing were likely dependent on the corresponding heterogeneous formation of sulphate.

14 To further understand the effect of cloud processing on aerosol elements dissolution, individual cloud 15 droplet residues were collected during the two cloud events, and the micro-morphology and composition of metal particles were examined by TEM-EDS analysis. Representative TEM images 16 17 and the EDS spectra of major metal particles are shown in Fig. 11. Plenty of Pb-rich, Fe-rich and fly 18 ash particles were observed to be the dominating metal particles in the two cloud events, consistent 19 with the major types of metal-associated particles (Pb-rich, fly ash, Fe-rich and Zn-rich) in cloud 20 droplets at Mt. Lushan (Li et al., 2013). Almost all of metal particles were found to be internally 21 mixed within S-rich cloud residues with the encapsulation of cloud water and presented a nearly 22 spherical shape or aggregation, suggesting the likely sources to be high-temperature coal combustion 23 by coal-fired power plants and industries, during which industrial gases containing abundant SO₂ and 24 metals were released, followed by the generation of metal-sulphate particles (Gier éet al., 2006).

Figure 11 also demonstrates the small diameters of the metal particles in cloud residues much smaller than 1 µm, especially for the aggregation of spherical Fe nanoparticles embedded in aged S-rich residues (Fig. 11c). The unique, large specific surface of nano-sized Fe particles is able to effect a higher absorption intensity and more complicated surface chemistry during cloud processing,

generating greater quantities of surface complexes with acidic anions such as sulphate, and 1 2 improving the dissolution rates of iron particles (Rubasinghege et al., 2010). Shi et al. (2009) 3 suggested that the variations in pH during cloud processing would induce the formation of 4 amorphous Fe nanoparticles and an increase in iron solubility (reactivity) in Saharan dusts. More 5 importantly, the solubilisation of aerosol elements caused by the change in particle morphology 6 during cloud processing is irreversible (Deguillaume et al., 2005). Finally, along with the dissipating 7 of cloud water in droplets, the Fe nanoparticles and sulphate residues remain in post-cloud particles, 8 resulting in an attractive enhancement of Fe solubility. Therefore, we can conclude that irreversible 9 changes or alterations of particle morphology such as specific surface area, which may result from 10 the acidification and heterogeneous reactions during cloud processing, should be the critical factor 11 for the significant enhancement of solubility not only for Fe, but also for many other trace elements 12 in cloud-processed fine particles.

13 4 Conclusions

14 The characteristics of fine particle trace elements at the summit of a mountain in southern China were investigated. The PM_{2.5} concentration (55.2 \pm 20.1 µg m⁻³) was twice the WHO guideline but 15 16 much lower than that in urban sites, with little seasonal variation. The total fractions of Al and Fe and 17 the water-soluble fraction of Zn were the dominant elements. Similar temporal variations between 18 the total and water-soluble fractions for all species along with several spikes contributed by dust 19 storms for Al and Fe were observed. The enrichment factor analysis showed notable air pollution to 20 aerosol trace elements at Mt. Lushan, especially to the water-soluble fractions. Nonferrous metal 21 mining and smelting, coal combustion, crustal materials and municipal solid waste incineration were 22 further classified as the major emission sources, which could initially determine the trace element 23 solubility with interferences such as total concentrations. The YRD including Jiangsu, Zhejiang and 24 parts of Anhui provinces were indicated as the most important source regions of PM_{2.5} by trajectory 25 cluster analysis and PSCF results, which was discrepant with the source distributions for individual 26 trace elements. The YRD, the PRD, eastern Hunan and western Jiangxi were identified as the 27 primary source regions and transport pathways for combustion-related elements (Mn, Pb, Zn, Cu, Se, 28 Cd and Mo) which were emitted by regional coal-fired power plants and municipal solid waste

1 incineration; whereas the long-range transport of mineral dusts from northern China was the major 2 contributor to crustal Al and Fe. Interestingly, local regions contributed relatively less to trace 3 elements concentrations. However, the YRD region and Jiangxi province were likely to make the 4 greatest contributions to element solubility rather than long-range transport from northern or 5 southern China. Significant enhancement in trace element solubility and corresponding increase in 6 aerosol sulphate were observed during two selected cloud events. TEM-EDS analysis of cloud 7 residues showed the metal particles were mostly mixed with sulphate. It was concluded that the 8 irreversible alteration of particle morphology by heterogeneous reactions or acidification with 9 sulphate during cloud processing could crucially increase trace elements solubility.

10 This study highlighted the contributions of regional anthropogenic pollution and cloud processing to 11 trace element solubility of $PM_{2.5}$ at Mt. Lushan. We expect that fine particles would undergo several 12 cloud-processing cycles during long-range transport with massive clouds covering the surface of the 13 earth, which can promote aerosol elements dissolution and increase health and ecological risks.

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	Overall		Summer,2011		Spring,2012	
	Total	Water soluble	Total	Water soluble	Total	Water soluble
PM _{2.5}	55.2 ±20.1		55.9 ±21.8		54.7 ±18.9	
Al	449.1 ±441.1	73.3 ±93.5	369.1 ±464.4	108.0 ± 121.7	515.1 ±415.2	48.9 ±57.3
Fe	331.1 ±236.2	71.5 ±70.5	330.4 ±250.9	101.9 ± 70.5	331.6 ±227.3	15.7 ±12.9
Zn	258.3 ±162.8	172.9 ± 105.8	274.3 ±129.5	211.9 ±104.4	245.3 ±186.1	142.3 ±97.5
Pb	68.2 ±49.3	29.4 ±23.8	65.4 ±47.3	37.8 ±27.5	70.5 ±51.4	22.7 ±18.3
Ba	63.8 ±54.2	23.4 ±17.0	66.0 ± 71.1	19.3 ±15.0	61.9 ±33.6	26.7 ± 18.0
Mn	22.2 ± 12.2	15.5 ±8.1	24.7 ± 12.5	19.0 ± 8.8	20.2 ± 11.6	12.8 ± 6.3
As	$21.5\ \pm 19.6$	14.8 ± 11.9	22.3 ± 16.3	12.40 ± 6.1	20.9 ± 22.2	17.0 ± 15.1
Cr	13.7 ± 17.2	0.7 ± 0.5	$18.2\ \pm 19.5$	0.8 ± 0.6	9.6 ±13.8	0.3 ± 0.1
Cu	12.4 ±9.6	9.9 ± 8.9	13.2 ± 6.4	10.7 ±6.9	11.7 ±11.6	9.3 ±10.2
Se	7.0 ±3.3	5.0 ±2.9	7.6 ± 2.9	5.2 ± 2.9	6.5 ± 3.5	4.8 ±2.9
Cd	$2.5~\pm1.8$	1.3 ± 0.9	2.8 ± 1.9	1.3 ± 0.6	2.2 ± 1.6	1.2 ± 1.1
Mo	2.0 ± 2.0	0.6 ± 0.4	2.6 ±2.2	0.7 ± 0.5	1.5 ±1.7	0.5 ± 0.3

Table 1. Concentrations (mean \pm SD) of PM_{2.5} (µg m⁻³) and trace elements (ng m⁻³) including total

2 and water soluble fraction at Mt. Lushan.



Figure 1. The location of Mt. Lushan and adjacent Yangtze River and Poyang Lake.



Figure 2. Temporal variations of individual trace element concentrations (ng m^{-3}) in PM_{2.5} at Mt.

- 3 Lushan. Boxes indicate the concentration spikes.



Figure 3. Enrichment factor (EF) values of fine particle trace elements at Mt. Lushan in total and
water soluble fraction.



Figure 4. Loadings of (a) Component 1, (b) Component 2, (c) Component 3 and (d) Component 4
for trace elements in total fraction in PM_{2.5}.



Figure 5. (a) Three-days mean trajectories arriving at Mt. Lushan, (b) averaged PM_{2.5} concentration
of five clusters, and (c) the likely source regions of PM_{2.5} identified using PSCF plots during
observation period.



Figure 6. Likely source regions of individual elements in PM_{2.5} at Mt. Lushan, identified using PSCF
plots. Note the criteria are 75th percentile for Al and mean concentration for others.



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Figure 7. Comparison of element solubility for aerosol at Mt. Lushan ($PM_{2.5}$, this study), the East China Sea (TSP, NE_{NAD} group, (Hsu et al., 2010)), Nanjing, China ($PM_{2.5}$, (Hu et al., 2012)) and Edinburgh, UK ($PM_{2.5}$, urban background site, (Heal et al., 2005)).



Figure 8. Scatter plot of aerosol element solubility vs. logarithmic EF value, computed with
geometric mean in 99% confidence interval. Colored shade areas represent different sources
classified by EF value and PCA analysis.



Figure 9. Solubility distributions in likely source regions for individual elements in PM_{2.5} at Mt.
Lushan, identified by PSCF with the mean solubility values as criteria.



Figure 10. The increments of trace elements solubility and sulphate concentrations by cloud processing. It is plotted as the ratios of post- to pre-cloud solubility or concentrations. Ratio > 1 (blue dashed line) indicates an increase of solubility or concentration.



Figure 11. Typical TEM images and corresponding EDS spectra of metal particles embedded in
individual S-rich cloud residues collected during cloud events. (a) Pb-S. (b) fly ash-S. (c) Fe-S. The
dotted red circles indicate the examined area of EDS. Cu* signals result from copper TEM grid.