



# Characteristics and source apportionment of fine haze aerosol in Beijing during the winter of 2013

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**Abstract.** For PM<sub>2.5</sub> filter samples collected daily at the Chinese Research Academy of Environmental Sciences (Beijing, China) from December of 2013 to February of 2014 (the winter period), chemical characteristics and sources were investigated with an emphasis on haze events in different alert levels. During the 3 months, the average PM<sub>2.5</sub> concentration was 89 µg m<sup>-3</sup>, exceeding the Chinese national standard of 75 µg m<sup>-3</sup> in 24 h. The maximum PM<sub>2.5</sub> concentration was 307 µg m<sup>-3</sup>, which characterizes developed-type pollution (PM<sub>2.5</sub>/PM<sub>10</sub>>0.5) in the World Health Organization criteria. PM<sub>2.5</sub> was dominated by SO<sub>4</sub><sup>2-</sup>, NO<sub>3</sub><sup>-</sup>, and pseudo-carbonaceous compounds with obvious differences in concentrations and proportions between non-haze and haze episodes. The non-negative matrix factorization (NMF) analysis provided reasonable PM<sub>2.5</sub> source profiles, by which five sources were identified: soil dust, traffic emission, biomass combustion, industrial emission, and coal combustion accounting for 13, 22, 12, 28, and 25 % of the total, respectively. The dust impact increased with northwesterlies during non-haze periods and decreased under stagnant conditions during haze periods. A blue alert of heavy air pollution was characterized by the greatest contribution from industrial emissions (61 %). During the Chinese Lantern Festival, an orange alert was issued and biomass combustion was found to be the major source owing to firework explosions. Red-alert haze was almost equally contributed by local traffic and transported coal combustion emissions from the vicinity of Beijing (approximately 40 % each) that was distinguished by the highest levels of NO<sub>3</sub><sup>-</sup> and SO<sub>4</sub><sup>2-</sup>, respectively. This study

also reveals that the severity and source of haze are largely dependent on meteorological conditions.

## 1 Introduction

With the increasing PM<sub>2.5</sub> concentration in northern China, winter haze occurrences increased from 3 to 16 days per year during 2000–2012 (Wang and Chen, 2016). The frequency of haze events during winter is enhanced by meteorological conditions; the minimum daily temperatures typically reach −15 to −20 °C (Wu et al., 2012) and the boundary layer height becomes shallow, to less than 100 m (Zheng et al., 2015). Moreover, the combustion of fossil fuel increases at low temperatures (Zhang and Samet, 2015). As the air quality deteriorated, China released its third revision of the “The National Ambient Air Quality Standards” (NAAQS) in 2012 (GB3095–2012), which stipulated safe PM<sub>2.5</sub> levels for the first time (Zhang and Cao, 2015). However, the worst haze events in the major cities of China were recorded during the winter of 2012–2013. During January of this period, Beijing experienced almost daily haze and the hourly PM<sub>2.5</sub> concentration reached 855 µg m<sup>-3</sup> (Zheng et al., 2015). In Beijing, winter haze episodes lasted 5 days (Zheng et al., 2015, 2016). The long duration of haze with high PM<sub>2.5</sub> concentration triggers a red alert for air pollution (Liu et al., 2017), which is the highest level of the heavy air pollution warning system issued in the “Emergency plan for heavy air pollution in Beijing” (revised in 2016; in Chi-

nese: <http://zhengce.beijing.gov.cn/zhengce/197/1861/1921/1005141/1283071/index.html>).

The concentrations of SO<sub>2</sub>, NO<sub>x</sub>, and volatile organic compounds (VOCs), which are important precursors of PM<sub>2.5</sub>, vary by emission and policy implementations. Related particulate compositions (sulfate, nitrate, and organic matter) comprise two-thirds of PM<sub>2.5</sub> (Huang et al., 2014; Hu et al., 2015). Over a 7-year period (2000–2006), SO<sub>2</sub> emission increased by 53 %, consistent with the increases in power plant emissions from 10.6 to 18.6 Tg (Lu et al., 2010). Particularly in northern China, the emissions from power plants have increased by 85 % over this period. In contrast, SO<sub>2</sub> levels have significantly decreased since 2006, when stricter SO<sub>2</sub> regulations, such as the use of flue-gas desulfurization systems or scrubbers, were imposed (Van der A et al., 2016). The reduction was particularly rapid during 2008–2009. On the other hand, the NO<sub>x</sub> concentration increased from 2000 to 2012 (Hong et al., 2016; Cao et al., 2011). This increase is in agreement with the increased number of vehicles, which contribute 90 % of the total NO<sub>x</sub> emissions in Beijing (Hendrick et al., 2014; Wu et al., 2012). Meanwhile, the continuous increase in VOC emissions (from 13 Tg yr<sup>−1</sup> in 2000 to 26 Tg yr<sup>−1</sup> in 2012) was mainly driven by industrial processes (~ 70 %; Hong et al., 2016). Coal combustion (especially that of raw coal) from households is underestimated in the southern and eastern rural areas of Beijing. Rural coal combustion comprises approximately 75 % of Beijing's total coal combustion (Cheng et al., 2017). After the 2008 Olympic Games, residential coal combustion emitted large amounts of SO<sub>2</sub>, NO<sub>x</sub>, and VOCs (70, 17, and 43 kt, respectively). In 2013, these amounts had increased 2-fold to 132, 33, and 81 kt, respectively. At the end of 2013, China issued the Air Pollution Prevention and Control Action Plan (CAAC, 2013), which greatly reduced the precursor emissions in 2014 (Wang et al., 2015).

Under the strict regulations on boiler and industrial emissions, SO<sub>2</sub> concentrations in Beijing significantly decreased during the winter of 2013 and the fuel sulfur was reduced by more than 80 % in 2014 (relative to its 2013 levels; CAAC, 2013, 2015). Over the same period, the NO<sub>x</sub> levels were reduced by 6.7 % over the nation, but exceeded the standard by 42 % in Beijing, where local traffic emissions remained high. Meanwhile, the PM<sub>2.5</sub> pollution is the most severe in the region of southern Beijing, where the annual average concentration reached 150 µg m<sup>−3</sup> during 2014–2015. The level is comparable to the national standard of PM<sub>10</sub> (CAAC, 2015; Zhang and Cao, 2015).

Since the 2008 Olympics and 2013 CAAC, heavy industries have been relocated and high-quality fuel has been introduced. Both actions have reduced the concentrations of gaseous precursors (Wang et al., 2009; Van der A et al., 2016), although these reductions are in contrast to the frequent hazes currently observed in Beijing. In recent studies, the PM<sub>2.5</sub>, dust, and SO<sub>2</sub> concentrations in Beijing have been mainly attributed to regional transport (Wang et al., 2014,

2011; Yang et al., 2013). Considering the extreme haze situation in Beijing, researchers have sought the crucial factors of haze formation, usually by identifying the emission sources of PM<sub>2.5</sub>. The source apportionment of PM<sub>2.5</sub> is commonly analyzed by source receptor models such as positive matrix factorization (PMF) and non-negative matrix factorization (NMF; Reff et al., 2007; Kfoury et al., 2016). These models have implicated coal and industries as major sources of PM<sub>2.5</sub> in Beijing (Huang et al., 2014; Zhang and Cao, 2015; Zhang et al., 2013).

Following the severe and frequent haze occurrences in January of 2013, the chemical characteristics and sources of PM<sub>2.5</sub> in Beijing were extensively investigated (Jiang et al., 2015; Zheng et al., 2015; Zhang et al., 2015; Chen et al., 2017). However, few studies have investigated the winter season of 2013–2014, which immediately followed the enactment of the 2013 CAAC in China. In particular, the source apportionment of Beijing's haze remains unknown (Wu et al., 2016). In the present study, we thoroughly examine the chemical compositions of PM<sub>2.5</sub> in Beijing during the winter of 2013–2014, and accordingly, diagnose the haze occurrence, probe the local and transported influence on haze, and quantify the critical source contributions.

## 2 Experiments

Filtered samples of PM<sub>10</sub> and PM<sub>2.5</sub> were collected on the roof of a three-story container (~ 15 m above ground level) at the Chinese Research Academy of Environmental Sciences (CRAES) in Beijing, China (40.04° N, 116.42° E), from December of 2013 to February of 2014. The site is located near a four-way intersection of a residential area located between the 5th and 6th ring roads of Beijing.

Aerosols were collected for 24 h (from 19:00 to 19:00 the following day) on a 90 mm polypropylene filter using a medium volume sampler at a flow rate of ~ 100 L min<sup>−1</sup> (2030, Laoying, China). Seventy PM<sub>2.5</sub> samples were collected and analyzed. The water-soluble ions (Cl<sup>−</sup>, NO<sub>2</sub><sup>−</sup>, CO<sub>3</sub><sup>2−</sup>, SO<sub>4</sub><sup>2−</sup>, NO<sub>3</sub><sup>−</sup>, Na<sup>+</sup>, NH<sub>4</sub><sup>+</sup>, K<sup>+</sup>, Mg<sup>2+</sup>, and Ca<sup>2+</sup>) were measured by ion chromatography (IC25, Dionex, USA) with a detection limit between 0.01 and 0.06 µg m<sup>−3</sup>. The ionic measurement method is detailed in Lim (2009). For trace elemental analysis, the samples were digested by a mixture of acids as described in Zhang et al. (2014). A quarter of each filter was placed into a polytetrafluoroethylene flask and digested with 8 mL of HNO<sub>3</sub>/H<sub>2</sub>O<sub>2</sub> (6/2 v/v, super-pure grade, Merck, Darmstadt, Germany) at 180 °C for 8 h. The solution was separated by centrifugation and diluted to 25 mL with ultrapure water. The concentrations of trace metals (21 species, including Si) were determined by inductively coupled plasma optical emission spectrometry (Prodigy 7, Teledyne Leeman, USA). The mass concentration of PM<sub>10</sub> was also determined for comparison with that of PM<sub>2.5</sub>.

The total concentrations of the water-soluble ions and trace elements were subtracted from the  $\text{PM}_{2.5}$  mass, to provide a measure that likely represents the carbonaceous components that were not directly measured. In this study, therefore, it was referred as the pseudo-carbonaceous components and used for the following discussion. The concentrations of these pseudo-carbonaceous components were comparable to those of  $\text{PM}_{2.5}$  concentrations observed in Beijing (Ji et al., 2016). A meteorological suite of relative humidity, temperature, and visibility was collected by CRAES from a sharing network of the China Meteorological Data Service Center (CMDC; <http://data.cma.cn/en/?r=data/detail&dataCode=A.0012.0001>). The gaseous species  $\text{NO}_x$ ,  $\text{SO}_2$ ,  $\text{CO}$ , and  $\text{O}_3$  were measured using commercial analyzers (42i, 43i, 48i, 49i, Thermo Fisher, USA) in CRAES.

The  $\text{PM}_{2.5}$  source was identified by non-negative matrix factorization (NMF) analysis. Introduced by Lee and Seung (1999, 2001), NMF operates similarly to positive matrix factorization (PMF). Both analysis methods find two matrices (**W** and **H**, termed the contribution matrix and the source profile matrix, respectively) that best reproduce the input data matrix (**V**) using the same factorization approach ( $\mathbf{V} = \mathbf{WH}$ ) as a positive constraint. However, while PMF is a generalized, alternative least-squares method, NMF minimizes the conventional least-squares error and the generalized Kullback–Leibler divergence. The uncertainties in NMF analysis were estimated as 0.3 plus the analytical detection limit (Xie et al., 1999a, b).

In addition to NMF analysis, the origin of air masses was traced by trajectory analysis. For air masses arriving at 500 m altitude, backward trajectories were computed for 72 h using HYSPLIT model with GDAS data in SplitR (Stein et al., 2015, <https://github.com/rich-iannone/SplitR>).

### 3 Characteristics of winter $\text{PM}_{2.5}$

#### 3.1 $\text{PM}_{2.5}$ and $\text{PM}_{10}$ mass variations

During the 2013–2014 winter period in Beijing, the mass concentrations of  $\text{PM}_{2.5}$  and  $\text{PM}_{10}$  varied in a similar pattern (Fig. 1). Zheng et al. (2015) reported a similar trend between the  $\text{PM}_{2.5}$  and  $\text{PM}_{10}$  concentrations. In this study, the average  $\text{PM}_{10}$  concentration was  $142 \mu\text{g m}^{-3}$ , comparable to the Chinese national standard of  $150 \mu\text{g m}^{-3}$  in 24 h (GB3095–2012). However, the mean  $\text{PM}_{2.5}$  concentration was  $89 \mu\text{g m}^{-3}$ , exceeding the standard of  $75 \mu\text{g m}^{-3}$  in 24 h. The  $\text{PM}_{2.5}$  standard was most severely exceeded in February 2014, when the average concentration ( $133.5 \mu\text{g m}^{-3}$ ) reached the highest winter concentration in Beijing during 2005–2015 (Lang et al., 2017).

Based on the criteria of the World Health Organization (WHO; 2006), the wintertime air pollution of Beijing was classified as developed type, meaning that the  $\text{PM}_{2.5}/\text{PM}_{10}$  ratio exceeded 0.5 in 70 % of the samples (Table 1). The

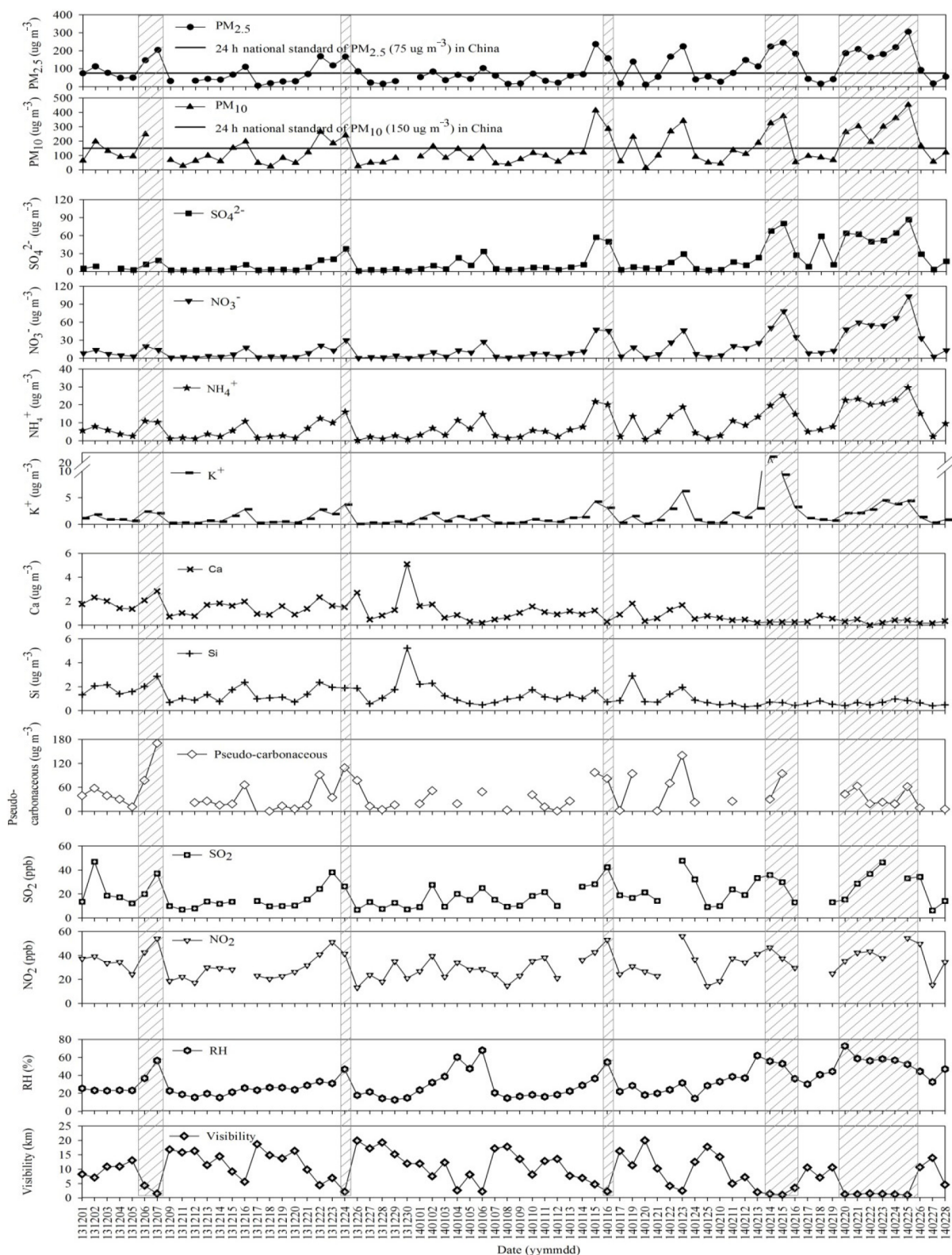
**Table 1.** Statistics of  $\text{PM}_{2.5}$  mass concentrations.

$\text{PM}_{2.5}$ mass classification	Number	$\text{PM}_{2.5}^a$ [ $\mu\text{g m}^{-3}$ ]
Chemical and NMF analysis	70	
Comparison with $\text{PM}_{10}$ mass	67	89
$\text{PM}_{2.5}/\text{PM}_{10} > 0.5$	47	113
$\text{PM}_{2.5} > 75 \mu\text{g m}^{-3}$ and $\text{PM}_{10} > 150 \mu\text{g m}^{-3}$	23	168
$\text{PM}_{2.5} > 75 \mu\text{g m}^{-3}$ and $\text{PM}_{10} < 150 \mu\text{g m}^{-3}$	5	113
$\text{PM}_{2.5} < 75 \mu\text{g m}^{-3}$ and $\text{PM}_{10} < 150 \mu\text{g m}^{-3}$	19	39
$\text{PM}_{2.5}/\text{PM}_{10} \leq 0.5$	20	31
Haze days <sup>a</sup>	13	198
Red alert	6	218
Orange alert	3	216
No/blue alert	4	168

<sup>a</sup> Average concentration. <sup>b</sup> Heavy air pollution alert.

mean  $\text{PM}_{2.5}$  concentration of these samples ( $113 \mu\text{g m}^{-3}$ ) was 4 times higher than that in developing-type pollution ( $31 \mu\text{g m}^{-3}$ ). In approximately half of the developed-type samples, the  $\text{PM}_{2.5}$  and  $\text{PM}_{10}$  mass concentrations exceeded the national standards, all of which were collected during haze events. The average  $\text{PM}_{2.5}$  concentration over 13 haze days reached  $198 \mu\text{g m}^{-3}$  and the visibility was significantly reduced to  $\sim 1$  km (Fig. 1). In contrast, the  $\text{PM}_{2.5}$  concentration exceeded the standard without violating the  $\text{PM}_{10}$  concentration on only a few days. These results well reflect the wintertime characteristics of  $\text{PM}_{2.5}$  levels in Beijing, which are largely related to haze episodes. The average  $\text{PM}_{2.5}$  concentration of the developed type was comparable to that of the developing-type unless the  $\text{PM}_{2.5}$  concentration exceeded the standard.

On 12 out of 13 haze days, the pollutant levels met the criteria of heavy air pollution alerts stipulated in the “Emergency plan for heavy air pollution in Beijing” (revised in 2016). In the lowest level of the four-tier warning system, blue alert, the daily average air quality index (AQI) exceeded 200 on only 1 day. In Table 1, the one no alert and three blue-alert haze days are defined as no/blue-alert haze events. The average  $\text{PM}_{2.5}$  concentration on these days was  $168 \mu\text{g m}^{-3}$  (Table 1). During the red-alert period (February 20–25), the daily  $\text{PM}_{2.5}$  concentration peaked at  $306 \mu\text{g m}^{-3}$ . A red alert is declared when the air pollution is heavy and severe. During a red alert, AQI exceeds 200 on 4 consecutive days, 2 consecutive days of which with AQI exceeding 300. Although the daily average AQI remained higher than 300 during the February 14–16 period, this event prompted an orange alert because it continued for only 3 days. The AQI data can be found at <http://www.tianqihoubao.com/aqi/beijing-201402.html> (in Chinese). Here, we describe episodes in terms of alerts defined in the heavy air pollution system rather than in the haze alert system, because the former definition is based on the daily averaged AQI, whereas the three-tier haze warnings depend on the hourly meteorological parameters (rel-



**Figure 1.** Variations in mass and chemical composition of  $\text{PM}_{2.5}$ ,  $\text{PM}_{10}$  mass, gaseous precursors, and meteorological parameters measured from 1 December 2013 to 28 February 2014. Horizontal lines indicate the Chinese national standards of PM concentrations in 24 h and the vertically shaded regions denote the 13 haze days.

**Table 2.** Sources identified by NMF analysis.

Factor	Contribution	Sources
Factor 1	13 %	soil dust
Factor 2	22 %	traffic emission
Factor 3	12 %	biomass combustion
Factor 4	28 %	industrial emission
Factor 5	25 %	coal combustion

ative humidity and visibility) or  $\text{PM}_{2.5}$ . Because we measured the daily concentrations, the heavy air pollution alert was suitable for our purpose.

### 3.2 Chemical composition

Throughout the wintertime, the average  $\text{PM}_{2.5}$  concentration remained close to  $90 \mu\text{g m}^{-3}$ , 20 % above the national standard. The major  $\text{PM}_{2.5}$  components were  $\text{SO}_4^{2-}$ ,  $\text{NO}_3^-$ ,  $\text{NH}_4^+$ , and pseudo-carbonaceous compounds, with average concentrations of 18.8, 16.9, 8.5, and  $38.6 \mu\text{g m}^{-3}$ , respectively. Collectively, these four compositions comprised 83 % of the  $\text{PM}_{2.5}$  mass (Fig. 2). On the 57 non-haze days, the fractional chemical compositions and concentrations of  $\text{SO}_2$  and  $\text{NO}_2$  were comparable to those of the entire period (70 days). In contrast, the portions of soil minerals such as  $\text{Ca}^{2+}$  and trace elements (including Si) were 3–4 times higher on non-haze days than on haze days. The  $\text{Ca}^{2+}$  and Si concentrations were highly correlated ( $r^2 = 0.8$ ) and were more related to the  $\text{PM}_{10}$  ( $r^2 = 0.6$ ) levels than to  $\text{PM}_{2.5}$  levels. This reflects the significant impact of soil dust on non-haze days (Fu et al., 2012). On haze days, the particle masses, compositions,  $\text{SO}_2$ , and  $\text{NO}_2$  varied widely among the different alert levels.

### 3.3 Source profiles

The  $\text{PM}_{2.5}$  sources were identified in an NMF analysis of the measurement data. The data included 8 water-soluble ions, 13 trace elements, and pseudo-carbonaceous compounds. After comparison through principal component analysis, the principal factors were determined. Finally, five critical factors were distinguished: soil dust, traffic emission, biomass combustion, industrial emission, and coal combustion (Table 2). The five source profiles are presented in Fig. 3. Despite their clear signatures, the contributions of dust and traffic emissions were approximately half of those of biomass combustion, industrial emission, and coal combustion (Table 2).

Factor 1 (soil dust) is confirmed by high  $\text{Ca}^{2+}$ , Si, Fe,  $\text{Cl}^-$ , and  $\text{Na}^+$  contents (Fu et al., 2012). The high concentrations of  $\text{Cl}^-$  and  $\text{Na}^+$  likely originate from dry lake deposits (Abuduwaili et al., 2015), which are spread over the northern area of Beijing. Elevated heavy metals suggest the presence of fugitive dust mixed with industry or traffic

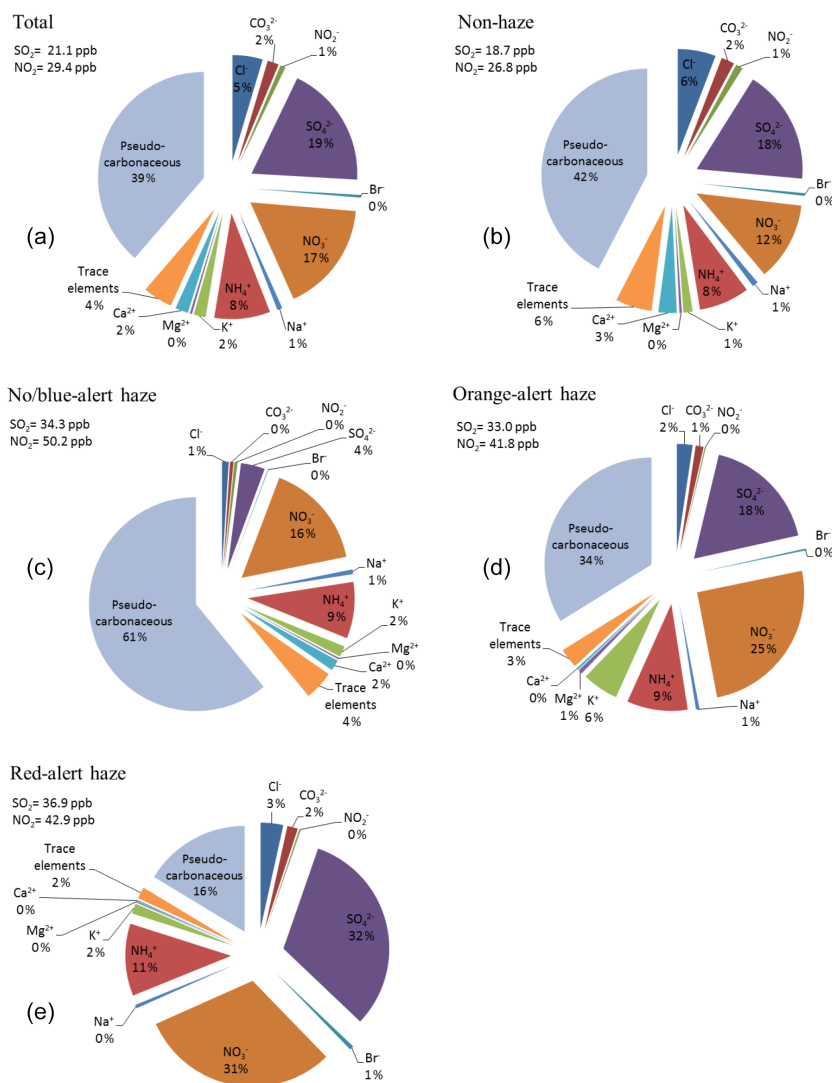
emissions (Wan et al., 2016). The high loadings of  $\text{NO}_3^-$  and  $\text{NH}_4^+$  in Factor 2 indicate traffic emissions (He et al., 2016). As is well known,  $\text{NH}_3$  is emitted from three-way catalytic converters in vehicles (Chang et al., 2016). Factor 3 (biomass combustion) emits large amounts of  $\text{K}^+$  and  $\text{NH}_4^+$  (Balasubramanian et al., 1999), along with the elements that give exploding fireworks their color (namely Mg, Fe, Al, Ti, Cu, and Si; Baranyai et al., 2015). The concentrations of these firework indicators are most significantly elevated during the Chinese Lantern Festival (14, 15, and 16 February; Fig. 1). Factor 4 (industrial emissions) is distinguished by high pseudo-carbonaceous materials and heavy metals. Factor 5 (coal combustion) is characterized by high  $\text{Cl}^-$ ,  $\text{SO}_4^{2-}$ , and  $\text{NO}_3^-$  contributions, which are absent in Factor 4. Although both Factors 4 and 5 represent the influence of industrial emissions near Beijing, Factor 5 is more clearly sourced from industries requiring high energy, such as iron and steel, cement, and power plants (Tan et al., 2016; Zhang et al., 2013). In contrast, Factor 4 indicates emissions from industrial processes using VOCs as raw materials such as furniture manufacturing, petroleum refining, machinery equipment manufacturing, and printing (Wu et al., 2015).

In a previous study, source apportionment by NMF or PMF analysis distinguished 7–8 factors (Zhang et al., 2013), including a secondary formation source. The secondary source was not separated as an individual factor in the present study. As a typical secondary species,  $\text{SO}_4^{2-}$  dominates in Factor 5. However, a  $\text{NO}_3^-$  signature appears in all factors except Factor 4. As a megacity, Beijing is surrounded by large satellite cities with industrial complexes. Thus, Beijing is susceptible to emissions from these areas in addition to its own emissions when certain meteorological conditions occur. This will be discussed in detail in the following section. In fact, the atmospheric conditions facilitated haze occurrence, leading to the major sources and the degree of aging for aerosols being intimately coupled. Therefore, these five factors primarily represent direct emission sources with secondary sources being implicitly included. In addition,  $\text{NO}_2$  is more likely sourced from local emissions, but  $\text{SO}_2$  is expected to be transported from nearby regions.

## 4 Characteristics of winter haze

### 4.1 Chemical and meteorological characteristics

The chemical compositions of  $\text{PM}_{2.5}$  clearly differed on haze in contrast to non-haze days in terms of secondary ions and pseudo-carbonaceous compounds (Fig. 2). The largest fraction of pseudo-carbonaceous compounds (61 %) was accompanied by the smallest proportion of  $\text{SO}_4^{2-}$  (4 %) on no/blue-alert days, suggesting low coal consumption by high-VOC-emitting industries. On orange-alert haze events, the  $\text{NO}_3^-$  fraction was twice that on non-haze days, and the  $\text{K}^+$  and  $\text{Mg}^{2+}$  proportions were maximized (at 6 and 1 %, respec-



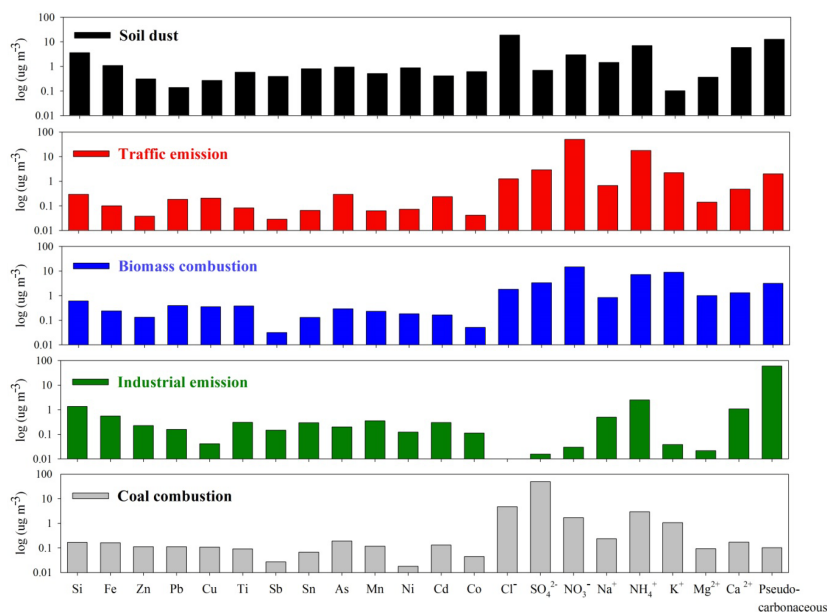
**Figure 2.**  $\text{PM}_{2.5}$  mass contributions of water-soluble ions, trace elements, and pseudo-carbonaceous matter during the entire period (a), non-haze days (b), and haze days at blue-alert (c), orange-alert (d), and red-alert (e) warning levels. The average  $\text{SO}_2$  and  $\text{NO}_2$  concentrations are also given.

tively), implying biomass-combustion emission during the Lantern Festival in China. The concentrations of  $\text{SO}_4^{2-}$  and  $\text{NO}_3^-$  were comparable with the greatest contribution in red-alert haze events. In addition, these species were closely related to the  $\text{Cl}^-$  ( $r^2 = 0.8$ ) and  $\text{NH}_4^+$  ( $r^2 = 0.9$ ) concentrations, respectively, suggesting large contributions by coal combustion and vehicle emission. It is also noteworthy that the  $\text{SO}_4^{2-}$  fraction varied more widely than the  $\text{NO}_3^-$  fraction (Table S1 in the Supplement). Among the three levels of haze events,  $\text{SO}_4^{2-}$  varied from 4 to 32 %, whereas  $\text{NO}_3^-$  varied from 16 to 31 % and  $\text{NH}_4^+$  from 9 to 11 %. Similarly, although both  $\text{SO}_2$  and  $\text{NO}_2$  concentrations were the highest in red-alert haze,  $\text{SO}_2$  enhancement (relative to non-haze days) was 20 % larger than  $\text{NO}_2$  enhancement. Because the sulfur compounds were much more elevated than the nitrogen com-

pounds on haze days (particularly in red-alert haze events), it was concluded that coal combustion, which emits sulfur compounds, was the largest contributor to the winter haze in Beijing. Furthermore, coal emissions are mostly transported from near Beijing (Hendrick et al., 2014).

To examine the meteorological conditions favorable for haze occurrence and clarify the emission source regions, surface weather maps combined with daily average backward trajectories at 500 m were compared during non-haze and haze events. Previous studies also reported that weather conditions were critical for haze formation. In eastern China, migratory anticyclones and weak pressure gradients were the prerequisites of winter haze from 1980 to 2012 (Peng et al., 2016). High  $\text{PM}_{2.5}$  episodes in Beijing usually began with weak southerly winds and ended with strong northerly winds





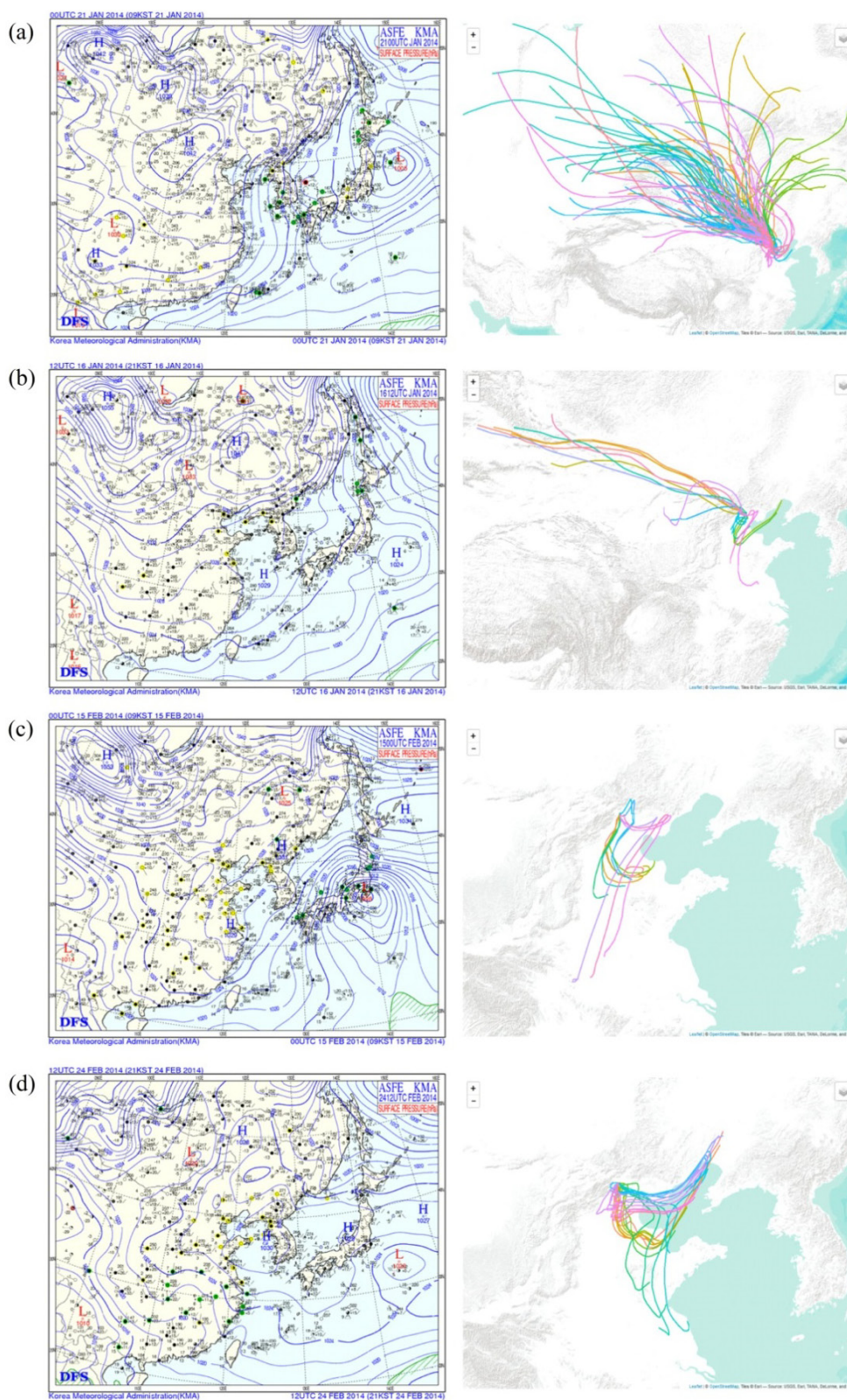
**Figure 3.** Composition profiles of the five factors identified in NMF analysis.

(Guo et al., 2014). In this study, the air mass was usually transported from the northwest. As the high-pressure system expanded, however, it was transported from the west, southwest, and southeast. Throughout this process, weather conditions became increasingly stagnant (Fig. 4) and the haze alert level increased gradually. The recent study also emphasized the effect of meteorological conditions on the severity of haze in Beijing (Cai et al., 2017). When air masses were rapidly transported from the northern desert area (Fig. 4a), mineral species such as  $\text{Ca}^{2+}$  and Si were enriched on non-haze days and the  $\text{PM}_{10}$  mass was high. In the western regions of Beijing (Fig. 4b), where various industries manufacture food, drink, furniture, pharmaceuticals, and other products from VOCs (<http://www.berkeleysg.com/2016/06/china-manufacturing-distribution-map/>), the fraction of pseudo-carbonaceous compounds rose to its maximum as the air mass slightly lingered over the region. During 14–16 February, firework explosions caused a spike in  $\text{K}^+$ ,  $\text{Mg}^{2+}$ , and  $\text{NH}_4^+$  concentration under the stagnant weather conditions, in which the air mass moved very slowly from the southwestern areas, where population density is the highest (Cheng et al., 2017). As the air mass moved eastward toward the high-energy-requiring regions (<http://berc.berkeley.edu/energy-access-developing-parts-china/>; Fig. 4d), such as Tianjin and Tangshan, where coal consumption is high for industrial use and residential heating (Cheng et al., 2017), the  $\text{PM}_{2.5}$  and  $\text{SO}_4^{2-}$  ( $\text{SO}_2$ ) concentrations reached their maxima.

## 4.2 Source profiles

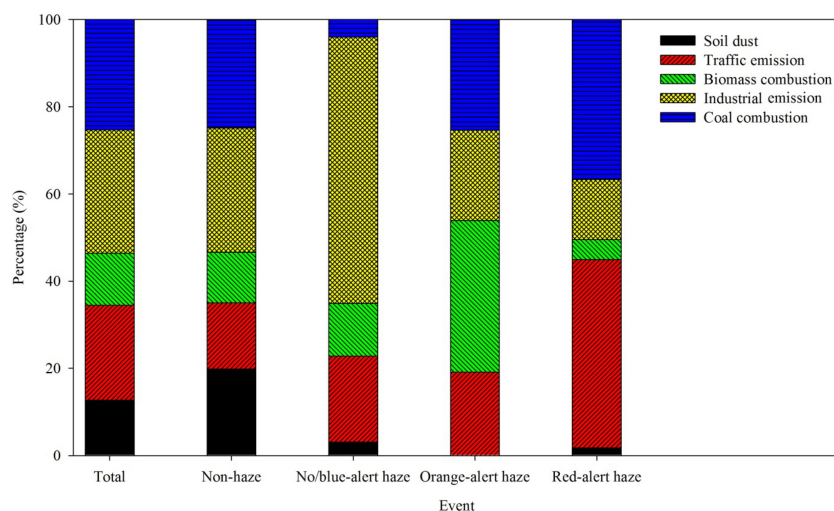
To quantify each source contribution during the winter haze in Beijing, daily samples were analyzed by NMF and the source profiles during haze and non-haze episodes were compared (Fig. 5). In all samples, the main contributions were industrial, traffic, and coal combustion emissions (22–28 %), followed by soil dust and biomass combustion (13 and 12 %, respectively). However, soil dust loading, which is associated with elevated fractions of Ca, Si, and pseudo-carbonaceous matters (Fig. 2), was enhanced to 20 % during non-haze events. Meanwhile, the local traffic contribution decreased as the air mass was rapidly transported from the northwestern desert areas, as mentioned in Sect. 4.1.

The three types of haze episodes exhibited strong contrasts not only in their chemical species and source regions, as mentioned above, but also in their source profiles (Fig. 5). No/blue-alert haze was dominated by industrial emissions (61 %) as the airflow passed over the industrial regions manufacturing products from raw VOCs. Consequently, the pseudo-carbonaceous concentration increased. During orange- and red-alert haze events, the dust contribution was negligible and the anthropogenic fraction increased sharply. During the Chinese Lantern Festival (which triggered an orange alert), a biomass signature with the highest  $\text{K}^+$  concentration was observed in the air mass transported from the southwestern populated areas of Beijing. The  $\text{K}^+$  contribution (35 %) was 3 times higher than that on non-haze days. During February 20–25, the outflow of the high coal-consuming eastern region enhanced the proportion of coal combustion products to 37 %. Simultaneously, the traffic contribution was the highest at 43 %. The coal and traffic



**Figure 4.** Surface weather maps and 72 h backward trajectories on days of (a) non-haze (57 days), (b) no/blue-alert haze (4 days), (c) orange-alert haze (3 days), and (d) red-alert haze (6 days). Trajectories were calculated twice a day at 18:00 and 06:00 UTC for non-haze days in (a) and every 6 h at 12:00, 18:00, 24:00, and 06:00 UTC for haze days in (b, c, d).





**Figure 5.** Comparison of source contributions (left to right) over the entire winter, during non-haze events, and during no/blue-alert, orange-alert, and red-alert haze events.

effects were accompanied by 2-fold increases in  $\text{SO}_4^{2-}$  and  $\text{NO}_3^-$  in  $\text{PM}_{2.5}$ .

## 5 Policy implications

During the 2013–2014 winter period in Beijing, the average  $\text{PM}_{2.5}$  concentration exceeded the standard by 20 %, and in February reached its highest level over 2005–2015 (Lang et al., 2017). The  $\text{PM}_{2.5}$  mass closure and concentration of gaseous precursors during the 57 non-haze days were comparable to those of the entire winter period. Mineral dust is an important source of  $\text{PM}_{2.5}$  and elevates the  $\text{PM}_{10}$  concentration on non-haze days. The average  $\text{PM}_{2.5}$  concentrations increased significantly from  $65 \mu\text{g m}^{-3}$  on non-haze days to  $168 \mu\text{g m}^{-3}$  on no/blue-alert days and to  $218 \mu\text{g m}^{-3}$  on red-alert days.

When weather conditions stagnate under weak pressure gradients, the alert levels of heavy air pollution upgrade on haze days. The migratory anticyclones also shift the air masses, causing wide variations in chemical species and emission sources. During haze days, the  $\text{NO}_2$  and  $\text{NO}_3^-$  concentrations exceed those of  $\text{SO}_2$  and  $\text{SO}_4^{2-}$ , respectively, but the sulfur-containing species vary more widely than the nitrogen species. The sulfur compounds are particularly enhanced in stagnant air masses transported from the vicinity of Beijing, including the southern and eastern regions, leading to the large sulfur variation with little change in nitrogen. These results highlight the significant influence of emissions from industries requiring high energy and using coal and from local vehicles in the vicinity of Beijing on winter haze formation, which agrees with findings from previous studies (Hendrick et al., 2014; Wang et al., 2016). To abate the severe haze in Beijing, therefore, it is necessary to reduce vehicle emissions in Beijing and sulfur emissions from in-

dustrial complexes and uncontrolled coal combustion in surrounding cities. For cost effectiveness, the weather forecast needs to be incorporated into the policy implementation.

## 6 Conclusions

This study investigated the chemical characteristics of  $\text{PM}_{2.5}$  during the 2013–2014 winter period in Beijing and identified its sources with an emphasis on haze events by measuring the particle masses, water-soluble ions, and trace elements in filtered samples. Finally, policy implications for controlling haze occurrences in Beijing were deduced from the analysis.

The samples were collected daily at CRAES, Beijing, China, from December of 2013 to February of 2014. During the winter period, the overall average  $\text{PM}_{2.5}$  concentration in Beijing was  $89 \mu\text{g m}^{-3}$ , exceeding the Chinese national standard of  $75 \mu\text{g m}^{-3}$  in 24 h. The excess was linked to high occurrence of haze events in February of 2014. The high  $\text{PM}_{2.5}$  episodes were concurrent with  $\text{PM}_{10}$  exceedance. Seventy percent of the samples were identified as developed type in the WHO criteria; that is, their  $\text{PM}_{2.5}/\text{PM}_{10}$  ratios exceeded 0.5. All 13 recognized haze events in this study were included in the developed type.

The chemical compositions showed that secondary ions were doubled on haze days relative to non-haze days, but mineral species were halved during haze events. For the 70 daily  $\text{PM}_{2.5}$  samples, NMF analysis was performed and the source profiles were compared between haze and non-haze days. The analysis identified five principal sources, of which industrial emission, coal combustion, and traffic emission comprised similar fractions of 28, 25, and 22 %, respectively. The soil-dust and biomass-combustion sources were well distinguished and contributed 13 and 12 %, respectively. Comparing the source profiles between non-haze and haze

events, the impact of soil dust was most noticeable on non-haze days, when the air masses were transported rapidly from northwestern desert areas and brought high concentrations of  $\text{Ca}^{2+}$  and Si into Beijing. However, nearby transport of industrial, biomass combustion, and coal combustion emissions, along with local traffic emission, contributed to haze events under stagnant weather conditions. The contributions of these four sources increased by up to 61, 35, 37, and 43 % in no/blue-alert, orange-alert, and red-alert days, respectively. The industries that are mainly located to the west of Beijing use VOCs as raw materials, elevating the pseudo-carbonaceous components in  $\text{PM}_{2.5}$ . Biomass combustion increases during the firework displays of the Lantern Festival (14–16 February). At that time, the  $\text{K}^+$  and  $\text{Mg}^{2+}$  concentrations are maximized. When a red alert was issued for 6 days in 2014, the contribution of  $\text{SO}_4^{2-}$  and  $\text{NO}_3^-$  increased by factors of 3 and 2, respectively, from their non-haze levels. Overall, the sulfur compounds ( $\text{SO}_2$  and  $\text{SO}_4^{2-}$ ) varied much more widely than the nitrogen compounds ( $\text{NO}_2$  and  $\text{NO}_3^-$ ) through haze events, implying the substantial contribution of industrial emissions from coal combustion in surrounding cities. The high level of nitrogen compounds suggests local vehicle emissions as a main source of winter haze in Beijing. This study also emphasizes the role of weather conditions in haze formation by building up stagnant conditions that facilitate the transport of industrial emissions from the vicinity of Beijing. These findings will be applicable to policy making.

**Data availability.** Measurement data and analysis results are available upon request to the corresponding author (zhangkai@craes.org.cn or meehye@korea.ac.kr).

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