Effectiveness evaluation of temporary emission control action in 2016 winter in Shijiazhuang, China

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Abstract.

To evaluate the environmental effectiveness of the control measures for atmospheric pollution in Shijiazhuang of China, a large-scale controlling experiment for emission sources of atmospheric pollutants (i.e., a temporary emission control action, TECA) was designed and implemented during November 1, 2016 to January 9, 2017. Under the unfavorably meteorological conditions, compared to the no control action and heating period (NCAHP), the mean concentrations of PM$_{2.5}$, PM$_{10}$, SO$_2$, NO$_2$, and chemical species (Si, Al, Ca$^{2+}$, Mg$^{2+}$) in PM$_{2.5}$ during the control action and heating period (CAHP) still decreased by 15 %, 26 %, 5 %, 19 %, 30.3 %, 4.5 %, 47.0 % and 45.2 %, respectively, indicating that the control measures for atmospheric pollution were effective and was in a right direction. The effects of control measures in suburbs were better than those in urban area, especially for the control effects of particulate matter sources. The control effects for emission sources of carbon monoxide (CO) were not apparent during the TECA period, especially in suburbs, likely due to the increasing usage of domestic coal in suburbs along with the temperature decreasing.

The results of PMF analysis showed that crustal dust, secondary sources, vehicle emissions, coal combustion and industrial emissions were main PM$_{2.5}$ sources. Compared to the whole year (WY) and the no control action and no heating period (NCANHP), the contribution concentrations and proportions of coal combustion to PM$_{2.5}$ increased significantly during other stages of TECA period. The contribution concentrations and proportions of crustal dust and vehicle emissions to PM$_{2.5}$ decreased apparently during the CAHP compared to other stages of TECA period. The contribution concentrations and proportions of industrial emissions to PM$_{2.5}$ during the CAHP decreased apparently compared to the NCAHP. The pollutants’ emission sources during the CAHP were in effective control, especially for crustal dust and vehicles. While the necessary coal heating for cold winter and the unfavorably meteorological conditions had an offset effect on the control measures for emission sources to some degree. The results also illustrated that the discharge of pollutants might be still enormous even under such strict control measures.

The backward trajectory and potential source contribution function (PSCF) analysis in the light of atmospheric pollutants suggested that the potential sources-areas mainly involved in the surrounding regions of Shijiazhuang, i.e., south of Hebei, north of Henan and Shanxi. The regional nature of the atmospheric pollution in Northern China Plain revealed that there is an urgent need for making cross-boundary control policy except for local control-measures given the high background
The TECA is an important practical exercise but it can’t be advocated as the normalized control measures for atmospheric pollution in China. The direct cause of atmospheric pollution in China is the emission of pollutants exceeds the air environment’s self-purification capacity, and the essential reason is unreasonable and unhealthy pattern for economic development of China.

Keywords: Atmospheric pollutants; Effectiveness evaluation; Control action; PMF; PSCF
1 Introduction

As a consequence of rapid industrialization and urbanization, China has been suffering from air quality degradation in recent years (Fu et al., 2014; Gao et al., 2015; Han et al., 2014; Hao et al., 2017; Zhao et al., 2011). Frequently occurred severe haze is featured by long duration, extensive coverage and sharply-increasing particulate concentration (Jiang and Xia, 2017; Tao et al., 2014; Wang et al., 2016a; Zhang et al., 2015a). It has been suggested that severe haze pollution increases the risk of respiratory and cardiovascular diseases (Chen et al., 2013; Gao et al., 2015; Pan et al., 2014; Zhang et al., 2014a; Zhou et al., 2015). On the basis of previous statistics, there are four haze-prone city clusters in China, including Beijing-Tianjin-Hebei region, Yangtze River Delta, Pearl River Delta and Sichuan Basin (Bi et al., 2014; Chen et al., 2016a; Fu et al., 2014; Fu and Chen, 2017; Li et al., 2016b; Tao et al, 2013a; Wang et al., 2015b; Wu et al., 2008; Zhang et al., 2015b).

In recent years, the role of particulates in hazy events has been becoming more and more prominent. The particulates can be discharged from varieties of sources or formed by physicochemical/aqueous-oxidation reactions between gaseous precursors, which have significant negative effects on climate, atmospheric visibility and public health (Chen et al., 2015; Fu and Chen, 2017; Lee et al., 2015; Quinn and Bates, 2003; Shen et al., 2015; Tai et al., 2010; Zhang et al., 2010). The high observed concentrations of fine particles and prolonged haze events have occurred frequently during autumn and winter, and covered large regions in China. In some cases, the instantaneous mass concentration of PM$_{2.5}$ had reached up to 1000 μg/m$^3$ (Qin et al., 2016; Zhang et al., 2014b), which caused the extensive concern from citizens and government agencies.

Confronted with severe air pollution and degradation of air quality, the government has taken a variety of control measures in recent years, including the odd-and-even license plate rule (http://www.sjz.gov.cn/col/1274081553614/2016/11/17/1479391129628.html), the mandatory installation of desulfurization, denitrification and other pollution-controlling facilities in factories (Liu et al., 2017a; Ma et al., 2015; Peng et al., 2017) and the on-line monitoring system structure plan in construction sites, etc. The atmospheric quality in China has been notably improved so far. From 2013 to 2016, the concentrations of atmospheric pollutants in China showed a decreased trend, and the annual mean concentrations of PM$_{2.5}$, PM$_{10}$, SO$_2$ and NO$_2$, in 2016 reached up to 50 μg/m$^3$, 85 μg/m$^3$, 21 μg/m$^3$ and 39 μg/m$^3$, respectively, and significantly lower than those in 2013 (http://www.zhb.gov.cn/hjzl/zghjzkgb/lnzghjzkgb/). However, the annual mean concentrations of
PM\textsubscript{2.5} and PM\textsubscript{10} in 2016 were still 1.4 and 1.2 times higher than the national ambient air quality standard (NAAQS) (GB3095-2012 Grade II, PM\textsubscript{2.5}: 35 μg/m\textsuperscript{3}, PM\textsubscript{10}:70 μg/m\textsuperscript{3}). Note that the concentrations of PM\textsubscript{2.5} and PM\textsubscript{10} during Beijing-Tianjin-Hebei region were up to 71 μg/m\textsuperscript{3} and 119 μg/m\textsuperscript{3} in 2016, and 2.0 and 1.7 times higher than the NAAQS, respectively. Therefore, China still has a lot of work to do to improve the national air quality.

Over the last decade, Chinese government has implemented stricter control-measures for emission sources during multiple international events held in China than normal times (Chen et al., 2016b; Guo et al., 2013; Liu et al., 2013; Sun et al., 2016; Wang et al., 2010; Wang et al., 2017). For instance, the first attempt took place during the Beijing 2008 Olympic Games (Guo et al., 2013). Drastic control actions were executed to cut down the emissions of atmospheric pollutants from motor vehicles, industries and building construction activity (UNEP, 2009; Wang et al., 2009a; Wang et al., 2010). UNEP (2009) suggested that the concentration of PM\textsubscript{10} in Beijing was reduced by 20 % due to the emission reduction measures. Liu et al. (2013) reported that the concentrations of SO\textsubscript{2}, NO\textsubscript{2}, PM\textsubscript{10} and PM\textsubscript{2.5} were reduced by 66.8 %, 51.3 %, 21.5 % and 17.1 %, respectively, during the 2010 Asian Games in Guangzhou of China, and during which stricter control measures for emission sources were implemented. Furthermore, further stricter controls for emission sources were implemented in both Beijing and its surrounding regions during the 2014 Asia-Pacific Economic Cooperation (APEC) summit and Parade. Compared to no-control during APEC and Parade, a decreasing trend with 51.6–65.1 % and 34.2–64.7 % of PM\textsubscript{2.5} concentrations during the control period was reported (Wang et al., 2017). Eventually, all the efforts led to a blue-sky days during the APEC, which was acknowledged as “APEC Blue” (Wang et al., 2016b). As we can see that the air quality can be improved in response to stricter emission controls in international events held in China. However, once these stricter control-measures of emission sources were repealed, and the air quality would be deteriorated subsequently (http://www.mep.gov.cn/gkml/hbb/qt/201412/t20141218_293152.htm), indicating that the prevention and control of air pollution in China still had a long way to go.

Shijiazhuang (38.03° N, 114.26° E), a hinterland city of Northern China Plain with a high population density, is an important city in Beijing-Tianjin-Hebei region (Sun et al., 2013). The rapid industry development has a great contribution to this city’s economic growth and degradation of air quality at the same time (Du et al., 2010; Li et al., 2015; Yang et al., 2015, 2016a). Shijiazhuang has
been one of the cities with the most serious air pollution in the world (https://www.statista.com/chart/4887/the-20-worst-cities-worldwide-for-air-pollution/), and deteriorating air quality poses a great risk to public health (http://www.who.int/ceh/risks/cehair/en/), as well as drags on the expansion of economy. The government of Shijiazhuang has adopted a variety of control measures (http://www.sjzhb.gov.cn/), however, it seems that the improvement in air quality of Shijiazhuang is not go into effect so far, and the atmospheric pollution is still heavy. In 2016, the annual concentrations of PM$_{2.5}$ and PM$_{10}$ in Shijiazhuang reached up to 70 $\mu g/m^3$ and 123 $\mu g/m^3$, respectively, which were 2.0 and 1.8 times higher than the NAAQS (GB3095-2012 Grade II) (http://www.zhb.gov.cn/hjzl/tj/201706/t20170606_415527.shtml). Especially in the heating period in winter, the degree of atmospheric pollution in Shijiazhuang was even more serious.

The effectiveness of control measures has been queried in recent years. Therefore, based on previous examples of APEC, Parade and the Asian Games, etc., a large-scale controlling experiment for atmospheric pollutants sources (i.e., TECA) was designed and implemented to investigate whether control measures in Shijiazhuang are effective for the atmospheric pollution. The experiment was carried out in Shijiazhuang during November 1 2016 to January 9 2017, during which more stringent control measures of atmospheric pollution than usual were put into practice. Then, by combining of the changes of atmospheric pollutants concentrations, emission source contributions and other factors such as meteorological conditions, regional transmission, etc., the effectiveness of control measures was evaluated before and after the control measures were taken.

2 Materials and Methods

2.1 Site description

Shijiazhuang city is located in the east of Taihang Mountain in north of China (Fig. 1), and the urban area is 15848 km$^2$, with a population of more than 10 million in 2016. Shijiazhuang is a large industrial city that is famous for raw materials, energy production and steel, power, and cement industries. The number of vehicles is more than 2.0 million until 2016. Shijiazhuang has a typical temperate and monsoonal climate with four clearly distinct seasons, with northeasterly, southeasterly and northwesterly winds prevailed during the TECA period (Fig. S1). The mean wind speed was 0.6 m/s, and the average temperature was 14.9 $^\circ$C during the TECA period. The mean relative humidity was up to 76.5 %, and the mean height of mixed layer was 509 m during the TECA period. The meteorological conditions during the four stages of the TECA period in Shijiazhuang
were shown in Table 1.

The seven monitoring sites including Twenty-second Middle School (TSMS), High-tech Zone (HTZ), Great Hall of the People (GHP), Century Park (CP), Water Source Area in the Northwest (WSAN), University Area in the Southwest (UAS) and Staff Hospital (SH) are located in urban area of Shijiazhuang. While other seventeen sites including Fenglong Mountain (FLM), Gaoyi (GY), Gaocheng (GC), Xingtang (XT), Jinzhou (JZ), Jingxing Mining District (JXMD), Lingshou (LS), Luquan (LQ), Luancheng (LC), Pingshan (PS), Shenze (SZ), Wuji (WJ), Xinle (XL), Yuanshi (YS), Zanhuang (ZH), Zhaoxian (ZX) and Zhengding (ZD) are suited in suburbs of Shijiazhuang. The more details were shown in Table S1.

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Fig. 1. Maps of the online monitoring stations and the filter membrane sampling sites in Shijiazhuang. The 24 online monitoring stations mainly include Twenty-second Middle School (TSMS), Fenglong Mountain (FLM), High-tech Zone (HTZ), Great Hall of the People (GHP), Century Park (CP), Water Source Area in the Northwest (WSAN), University Area in the Southwest (UAS), Staff Hospital (SH), Gaoyi (GY), Gaocheng (GC), Xingtang (XT), Jinzhou (JZ), Jingxing Mining District (JXMD), Lingshou (LS), Luquan (LQ), Luancheng (LC), Pingshan (PS), Shenze (SZ), Wuji (WJ), Xinle (XL), Yuanshi (YS), Zanhuang (ZH), Zhaoxian (ZX) and Zhengding (ZD). The filter membrane sampling sites are mainly located in TSMS, LQ and LC.

Table 1. The meteorological conditions during the four stages (NCANHP, NCAHP, CAHP and ACA) of the TECA period in Shijiazhuang.

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2.2 Sampling and Analysis

2.2.1 Sampling

From November 1, 2016 to January 9, 2017, the concentrations of PM$_{2.5}$, PM$_{10}$, SO$_2$, NO$_2$, CO, O$_3$ and synchronous meteorological conditions (temperature, relative humidity, wind speed and wind direction) were monitored in the 24 monitoring sites belonged to national, provincial and city controlling points (Fig. 1). The more details about monitoring instruments were described in Table S2. The heights of mixed layer were measured with a lidar scanner (AGHJ-I-LIDAR (HPL)), which was set at an atmospheric gradient monitoring station in Shijiazhuang near CP site (Fig. 1), and more details were shown in supplemental material. The PM$_{2.5}$ filter membrane samples were collected in TSMS, LQ, and LC sites from November 24, 2015 to January 9, 2017. Three sampling sites were set on the rooftops of buildings at 12-15 meters above ground level. Meanwhile, the parallel samples and the field blanks were also collected at each site. More details about filter membrane sampling were shown in Table S3. Before sampling, the quartz filter membranes (47 mm
in diameter, Whatman, England) and polypropylene filter membranes (47 mm in diameter, Beijing Synthetic Fiber Research Institute, China) were baked in the oven at 500 °C and 60 °C, respectively. All the filter membranes after sampling were stored at 4 °C before subsequent gravimetric and chemical analysis to improve the accuracy of experimental results.

2.2.2 Gravimetric and Chemical analysis

A 24-hour equilibrium process of PM$_{2.5}$ filter membranes was performed at a condition of constant temperature (20 ± 1 °C) and humidity (45-55 %) before gravimetric analysis. For the gravimetric analysis, all the filter membranes were weighted twice on a microbalance with resolution of 0.01 mg (Mettler Toledo, XS105DU) before and after sampling. An electrostatic eliminating device was applied to ensure the accuracy of gravimetric results.

After the gravimetric analysis, the quartz filter membranes which carried atmospheric particulates were used to analyze water-soluble ions by Ion chromatography (Thermo Fisher Scientific, Dionex, ICS-5000+). One-eighth of the filter membrane was cut up and put into a 25 mL glass tube with 20 mL ultrapure water. After 1-hour ultrasonic extraction and 3 minutes centrifugalization, the supernatant was filtered with disposable filter head (0.22 μm) for subsequent instrumental analysis. The ions analyzed included SO$_4^{2-}$, NO$_3^-$, Cl$^-$, NH$_4^+$, K$^+$, Ca$^{2+}$, Na$^+$ and Mg$^{2+}$, and more details were shown in Figs. S2 and S3. Prior to the ions detection, standard solutions were prepared and detected for over three times and low relative standard deviations (RSD) were obtained. Analytical quantification was carried out by using calibration curves made from standard solutions prepared.

Polypropylene filter membranes were used for elemental analysis by inductively coupled plasma–mass spectrometry (ICP-MS, Agilent 7700x). Perchloric acid-nitric acid digestion method was applied for the pretreatment of filter membranes. Aggregately, 10 elemental species (Al, Si, Ti, Cr, Mn, Fe, Cu, Zn, As and Pb) were determined. The detection limits of all the elements were shown in Table S4. For quality assurance and quality control (QA/QC), standard reference materials were pre-treated and analyzed with the same procedure, with the recovered values for all the target elements falling into the range or within 5 % of certified values.

The OC and EC were determined on a 0.558 cm$^2$ quartz filter membrane punch by Desert Research Institute (DRI) Model 2001 Thermal/Optical Carbon Analyzer with IMPROVE A thermal/optical reflectance (TOR) protocol. The quartz filter membrane was heated stepwise to
temperatures of 140 °C, 280 °C, 480 °C and 580 °C in a non-oxidizing helium (He) oven to analyse
OC1, OC2, OC3 and OC4, respectively. Then, the oven was added to an oxidizing atmosphere of
2 % oxygen (O2) and 98 % He, and the quartz filter membrane was gradually heated to 580 °C,
780 °C and 840 °C to analyse EC1, EC2 and EC3, respectively. The POC is defined as the carbon
combusted after the initial introduction of oxygen and before the laser reflectance signal achieves
its original value and the POC is specified as the fraction of OC. According to the IMPROVE A
protocol, OC is defined as OC1+OC2+OC3+OC4+POC, and EC is defined as
EC1+EC2+EC3−POC. For QA/QC, we carried out the measurement with the field blank filter
membranes, standard sucrose solution and repeated analysis in the study. During each season, the
field blanks were sampled and the particulate samples have been corrected by the average
concentration of the blanks. For checking the precision of instrument, a replicate sample was
analysed for every 10 samples, and the standard deviation <± 5 % was accepted. The method
detection limits (MDLs) of OC and EC are 0.45 and 0.06 μg/cm², respectively.

2.3 PMF model

PMF model can decompose a matrix of sample data (X) into two matrices: source profile (F)
and source contribution (G), in terms of observations at the sampling sites (Paatero and Tapper,
1994). The principle of PMF model can be described by:

\[ X_{ij} = \sum_{k=1}^{p} g_{ik} f_{kj} + e_{ij} \]  

where \( X_{ij} \) represents concentration of the \( j \)th species in the \( i \)th sample, \( g_{ik} \) represents the contribution
of the \( k \)th source to the \( i \)th sample, \( f_{kj} \) represents the source profile of \( j \)th species from the \( k \)th source,
\( e_{ij} \) represents the residual for the \( j \)th species in the \( i \)th sample, and \( p \) represents the number of sources.

PMF can identify emission sources of PM2.5 without source profiles. Data below MDLs are
retained for using in PMF model with the related uncertainty adjusted in terms of the characteristics
that PMF model admits data to be signally weighed. To assess the stability of the solution, the object
function Q can be allowed to review the distribution of each species, which is expressed by:

\[ Q = \sum_{i=1}^{n} \sum_{j=1}^{m} \left( x_{ij} - \sum_{k=1}^{p} g_{ik} f_{kj} \right)^2 \mu_{ij} \]  

where \( \mu_{ij} \) represents the uncertainty of \( j \)th species in the \( i \)th sample, which is applied to weight the
observations that include the sampling errors, missing data, detection limits and outliers.

The purpose of PMF model was to minimize the function (Eq. (2)). Data below MDLs were
retained and their uncertainties were set to 5/6 of the MDLs. Missing values were replaced by the median concentration of a given species, with an uncertainty of four times the median (Brown et al., 2015). Values that were larger than the MDLs, the calculation of uncertainty was in terms of a user supplied fraction of the concentration and MDLs, and the error fraction was suggested as 10 % by Paatero (2000). Uncertainty was described by:

\[
\text{Uncertainty} = \sqrt{\left(\text{Error Fraction} \times \text{concentration}\right)^2 + \left(0.5 \times \text{MDL}\right)^2}
\]  

(3)

In this study, EPA PMF 5.0 model was used to identify the PM$_{2.5}$ sources in Shijiazhuang city. Based on the field investigation and change of $Q$ values, and finally, five factors were chosen in PMF analysis. When five factors were chosen and input in PMF model, and the calculated $Q$ value (5162) from PMF model was close to theoretical values (5045). The observed PM$_{2.5}$ concentrations and calculated PM$_{2.5}$ concentrations from PMF model showed high correlations ($r = 0.96$) (Fig. S4). S/N is the signal-to-noise ratio, which is used to address weak and bad variables when running PMF model (Paatero and Hopke, 2003). The signal vector is identified as S and the noise vector is identified as N. Next, S/N is defined as Eq. (4). Variables with S/N ≤ 0.2 were removed from the analysis, while weak variables (0.2 ≤ S/N ≤ 2.0) were down-weighted (Ancelet et al., 2012). S/N of As, Ti and Cr were lower than 1.0 in this study, and these species were set as weak variables.

\[
S/N = \sqrt{\sum n_i^2 / \sum n_i^2}
\]  

(4)

where $i$ represents the chemical species in PM$_{2.5}$.

2.4 Backward trajectory and PSCF model

In this study, the 72-h backward trajectory arriving in Shijiazhuang (38.05° N, 55.2° E) was calculated at 1-h intervals during the CAHP by the Hybrid Single Particle Lagrangian Integrated Trajectory (HYSPLIT) model. The final global analysis data were produced from the National Center for Environmental Prediction's Global Data Assimilation System wind field reanalysis (http://www.arl.noaa.gov/). The model was run 4 times per day at starting times, i.e., 0:00, 06:00, 12:00, 18:00 LT; the starting height was set at 100 m above the ground. The PSCF model was used to identify the potential sources-areas in terms of the HYSPLIT analysis. The study region was divided into $i \times j$ small equal grid cells. The trajectory clustering and PSCF model were performed by using the GIS-based software TrajStat (Liu et al., 2017a; Wang et al., 2009b). The PSCF value was defined as:
PSCF = \frac{m_{ij}}{n_{ij}} \quad (5)

where \( i \) and \( j \) were the latitude and longitude indices, \( n_{ij} \) represented the number of endpoints that fell in the \( ij \) cell, and \( m_{ij} \) was the number of endpoints in the same cell that were related to the samples that were greater than the threshold criterion.

Based on the NAAQS (GB3095-2012 guideline value (24 h) of Grade II), the criterion values of PM$_{2.5}$, PM$_{10}$, NO$_2$, CO were set to 75 \( \mu \)g/m$^3$, 150 \( \mu \)g/m$^3$, 80 \( \mu \)g/m$^3$ and 4 mg/m$^3$, respectively. The criterion values of SO$_2$ and O$_3$ were set to 68 \( \mu \)g/m$^3$ and 15 \( \mu \)g/m$^3$ respectively, in terms of the average during the CAHP. When \( n_{ij} \) is smaller than three times the grid average number of trajectory endpoint (\( n_{ave} \)), a weighting function \( W(n_{ij}) \) was used to reduce uncertainty in cells (Dimitriou et al., 2015). The weighting function was defined by:

\[
W_{PSCF_{ij}} = \frac{m_{ij}}{n_{ij}} * W(n_{ij}) \quad (6)
\]

\[
W(n_{ij}) = \begin{cases} 
1.00 & n_{ave} \leq n_y \\
0.70 & 1.5n_{ave} < n_y \leq 3n_{ave} \\
0.40 & n_{ave} < n_y \leq 1.5n_{ave} \\
0.20 & n_y \leq n_{ave}
\end{cases} \quad (7)
\]

The studying field ranged from 33° N to 51° N, and 97° E to 121° E, and the region that was covered by the backward trajectories was divided into 432 grid cells of 1.0° × 1.0°. The total number of endpoints during the CAHP was 12672. Accordingly, there was an average of 5 trajectory endpoints in per cell (\( n_{ave} = 5 \)).

2.5 Measures taken in the controlling experiment

The measures taken in the controlling experiment began on November 18, 2016 and ended on December 31, 2016 in Shijiazhuang (http://www.sjz.gov.cn/col/1274081553614/2016/11/17/1479391129628.html). The measures taken in the control action were mainly aimed at controlling emission sources of atmospheric pollutants in Shijiazhuang, which mainly included five aspects: (1) reduce the usage of coal, (2) decrease industrial production, (3) inhibition of dust emission, (4) driving restriction, and (5) prohibit open burning. The more details were described in supplemental material.

Actually, a total of 1543 enterprises were shut down in the whole city of Shijiazhuang during the control action period, including pharmaceutical, steel, cement, coking, casting, glass, ceramics, calcium and magnesium, sheet, sand and stone processing, stone processing and other industries.
The situation of specific closed-enterprises in different districts and counties is shown in Table S5. In closed enterprises in Shijiazhuang, the number of mining and stone processing enterprises was the largest, which was up to 733 and account for 48 % of all the closed enterprises. The numbers of casting and building materials enterprises were up to 297 and 227, respectively, accounting for 19 % and 15 % of the all, respectively. In addition, 64 enterprises related to pharmaceutical industry were halted only for the VOC technology, and the 17 enterprises related to chemical industry must stop production. The numbers of closed enterprises for cement and calcium/magnesium industry were up to 49 and 40, respectively. The number of closed factories related to furniture and tanneries was 43, and the numbers of closed steel and coking enterprises were up to 4 and 7, respectively.

The average value of daily social-electricity consumption from November 18 to December 31, 2016 was 103,470,000 kW • h (Fig. S5), which declined 10 % compared to that of daily social-electricity consumption from November 1 to 17, 2016, and declined 6 % compared to that of daily social-electricity consumption during the same period in 2015. Restriction of motor vehicles based on odd-and-even license plate rule in urban area of Shijiazhuang resulted in the decrease of the average traffic-flow on arterial roads, which reduced about 30 % compared to before the control action (Fig. S6). The dust emission can be reduced about 390 tons per day by a series of dust control-measures. Compared to before the control action, the daily emissions of SO$_2$, NO$_x$, smoke dust and VOCs reduced about 20 %, 33 %, 15 % and 7 %, respectively, during the control action period, on the basis of the statistics on pollutants emission inventories.

3 Results and discussion

3.1 Variations of atmospheric pollutants concentrations

3.1.1 Temporal trend

The time series of atmospheric pollutants concentrations during the TECA period are shown in Fig. 2. The average concentrations of PM$_{2.5}$ and PM$_{10}$ during the TECA period in Shijiazhuang were up to 181 μg/m$^3$ and 295 μg/m$^3$, respectively, which were 5.2 and 3.2 times than the Grade II limit values in the NAAQS. The ratio of PM$_{2.5}$/PM$_{10}$ reached up to 0.62 during the TECA period, indicating that the fine particulate dominated on the particulate pollution in Shijiazhuang. The mean concentration of PM$_{2.5}$ during the TECA period was significantly higher than those of winter in Beijing (95.50 μg/m$^3$), Tianjin (144.6 μg/m$^3$), Hangzhou (127.9-144.9 μg/m$^3$), Heze (123.6 μg/m$^3$) and Xinxiang (111 μg/m$^3$) (Cheng et al., 2015; Gu et al., 2011; Liu et al., 2015; Liu et al., 2017a;
Feng et al., 2016), and lower than those of winter in Handan (240.6 μg/m³) and Xian (266.8 μg/m³) (Meng et al., 2016; Zhang et al., 2011). Additionally, the NAAQS (GB3095-2012 Grade II) values of SO₂, NO₂, CO and O₃ were 60 μg/m³, 40 μg/m³, 4 mg/m³ and 160 μg/m³, respectively. During the TECA period, the average concentration of SO₂ (60 μg/m³) could meet the NAAQS, and that of NO₂ (81 μg/m³) was far exceed the NAAQS; while those of CO (3.4 mg/m³) and O₃ (15 μg/m³) were less than the NAAQS.

As well known, the date of coal-fired heating in Shijiazhuang began in November 15, 2016 (http://www.sjz.gov.cn/col/1451896947837/2016/10/28/1477635691926.html). Depending on the changes of atmospheric pollution sources and meteorological conditions (Table 1), the timeline of the TECA was divided into four stages: stage 1: no control action and no heating period (NCANHP), ranging from November 1 to 14, 2016; stage 2: no control action and heating period (NCAHP), ranging from November 15 to 17, 2016; stage 3: control action and heating period (CAHP), ranging from November 18 to December 31, 2016; stage 4: after control action (ACA), ranging from January 1 to 9, 2017.

During the TECA period, the variations of atmospheric pollutants concentrations were mainly affected by the heating for cold winter and the control measures of the control action except for the meteorological conditions. Therefore, we defined the following equations to evaluate the effects of the heating and control action, respectively, based on the atmospheric pollutants concentrations during the different stages of TECA (i.e., NCANHP, NCAHP, CAHP and ACA).

\[ P_{i-heating} = \frac{(C_{i-CAHP} - C_{i-NCANHP}) \times 100}{C_{i-NCANHP}} \]  
\[ P_{i-action} = \frac{(C_{i-NCAP} - C_{i-CAHP}) \times 100}{C_{i-NCAP}} \]  

where \( P_{i-heating} \) represents the increasing percentage (%) of atmospheric pollutant concentration because of the combined effects of heating for cold winter and meteorological conditions; \( P_{i-action} \) represents the decreasing percentage (%) of atmospheric pollutant concentration because of the combined influences of control action and meteorological conditions; \( C_{i-NCANHP} \) represents the concentration (μg/m³, CO: mg/m³) of atmospheric pollutant during the no-control action and no-heating period; \( C_{i-NCAHP} \) represents the concentration (μg/m³, CO: mg/m³) of atmospheric pollutant during the no-control action and heating period; \( C_{i-CAHP} \) represents the concentration (μg/m³, CO: mg/m³) of atmospheric pollutant during the control action and heating period.
During the NCANHP, the mean concentrations of PM$_{2.5}$ and PM$_{10}$ were 156 μg/m$^3$ and 253 μg/m$^3$ in Shijiazhuang, respectively. With the beginning of heating, the mean concentrations of PM$_{2.5}$ and PM$_{10}$ increased 44 μg/m$^3$ and 64 μg/m$^3$ during the NCAHP, respectively, and the $P_{PM_{2.5}}$-heating and $P_{PM_{10}}$-heating values were up to 28 % and 25 % (Fig. 3 and Fig. 4). However, during the CAHP, the mean concentrations of PM$_{2.5}$ and PM$_{10}$ were 185 μg/m$^3$ and 291 μg/m$^3$, respectively, which decreased by 15 % and 26 % compared to the NCAHP. And the $P_{PM_{2.5}}$-heating and $P_{PM_{10}}$-heating values were up to 8 % and 8 %, respectively. The mean height of mixed layer, the mean wind speed and temperature during the CAHP were lower than those during the NCAHP (Table 1). Unfavorably meteorological conditions during the CAHP had an offset effect on the control measures for emission sources. In view of Eq. (9), it can be seen that the positive values for $P_{PM_{2.5}}$-heating and $P_{PM_{10}}$-heating are more able to show that control action was effective. During the ACA, the concentrations of PM$_{2.5}$ and PM$_{10}$ were 227 μg/m$^3$ and 383 μg/m$^3$, respectively, which increased significantly by 42 μg/m$^3$ and 92 μg/m$^3$ compared to the CAHP. The variations of SO$_2$ and NO$_2$ concentrations during different stages of TECA were similar to those of PM$_{2.5}$ and PM$_{10}$ concentrations. The $P_{SO_{2}}$-heating and $P_{NO_{2}}$-heating values were 50 % and 33 %, respectively, and the $P_{SO_{2}}$-heating and $P_{NO_{2}}$-heating values were 5 % and 19 %. Note that the mean concentration of CO in Shijiazhuang city varied from 2.2 mg/m$^3$ during the NCANHP to 5.5 mg/m$^3$ during the ACA period, which showed an increasing tendency (Fig. 3). Because CO was mainly produced from the uncompleted combustion of fossil fuels, so the usage of domestic coal might be increasing with the gradual decrease of temperature from the NCANHP (8.4 °C) to the ACA period (0.7 °C) (Table 1). Meanwhile, it can also be inferred that the control of domestic coal during the TECA period in Shijiazhuang city performed little efficiency. Because of the lack of emission inventories for domestic coal or small-boiler coal in Shijiazhuang, so that the control measures were less targeted. Additionally, the concentrations of O$_3$ during different stages of TECA were lower compared to other pollutants (Figs. 2 and 3). Overall, the control measures of emission sources in Shijiazhuang during the TECA period were go into effect, while the coal heating for cold winter and the unfavorably meteorological conditions during the CAHP had an offset effect on the efforts of control measures for pollutant sources to some extent. The average wind speed during the CAHP (0.4 m/s on average) was lower than those during the other stages of the TECA period (0.5-0.7 m/s on average) (Table 1), and the wind directions were changeable (Fig. S1), which was in favor of the accumulation of atmospheric pollutants, and thus
causing the concentrations of atmospheric pollutants to increase during the CAHP. Note that the heights of mixed layer showed an apparently decreasing tendency from the NCANHP (540 m on average) and the NCAHP (590 m on average) to the ACA (431 m on average), and the height of mixed layer during the CAHP was only 474 m on average (Table 1). The decrease in the height of mixed layer can cause the concentrations of atmospheric pollutants near the ground to be compressed significantly and enhanced subsequently. In addition, during the CAHP, the multidirectional air-masses that were mainly originated from the Beijing-Tianjin-Hebei and its surrounding areas (e.g. Henan, Shandong and south of Hebei) displayed an overlap with each other in Shijiazhuang (Fig. S7), and further aggravate the level of air pollution in Shijiazhuang.

3.1.2 Spatial variation

The concentrations variations of PM$_{2.5}$, PM$_{10}$ and related gaseous pollutants (SO$_2$, NO$_2$, CO and O$_3$) during four stages (NCANHP, NCAHP, CAHP and ACA) in urban area and suburb in Shijiazhuang are shown in Figs. 3 and 5. During the NCANHP, the average concentrations of PM$_{2.5}$ in urban area and suburb were 166 μg/m$^3$ and 152 μg/m$^3$, respectively. The concentrations of PM$_{2.5}$ in urban area and suburb increased significantly during the NCAHP (t-test, $p<0.01$). The meanly increased concentration of PM$_{2.5}$ (46 μg/m$^3$) in urban area was higher than that of in suburb (43 μg/m$^3$), but the value of P$_{PM2.5-heating}$ in suburb (29 %) was higher than that in urban area (27 %) (Fig. 4). Note that the mean concentration of PM$_{2.5}$ in urban area was up to 243 μg/m$^3$ during the CAHP, which showed an increasing tendency, and the P$_{PM2.5-action}$ value was -15 % (Fig. 4), likely due to the unfavorably meteorological conditions such as lower wind speed (0.4 m/s) and lower height of mixed layer (474 m), etc. (Table 1 and Fig. S7). Conversely, compared to the NCAHP, the concentrations of PM$_{2.5}$ in suburb (a mean of 161 μg/m$^3$) decreased significantly during the CAHP (t-test, $p<0.01$), and the P$_{PM2.5-action}$ was up to 18 % (Fig. 4), indicating the control measures of PM$_{2.5}$ sources in suburb might be more effective than urban area. The tendency of SO$_2$ concentrations
during different stages of TECA (except the ACA period) was similar to that of PM$_{2.5}$. The $P_{\text{SO}_2}$-heating and $P_{\text{SO}_2}$-action values in urban area were up to 58% and -4%, respectively, and were up to 47% and 8% in suburb during the TECA period (Fig. 4). However, the concentrations of SO$_2$ in urban area and suburb decreased remarkably during the ACA compared to the CAHP (t-test, $p<0.01$), probably due to the effective control measures.

During the NCANHP, the average concentrations of PM$_{10}$ in urban area and suburb were 280 and 242 $\mu$g/m$^3$, respectively. Then, the meanly increased concentrations in urban area and suburb were up to 65 and 64 $\mu$g/m$^3$ during the NCAHP, which were comparable with each other. Nevertheless, the mean $P_{\text{PM}_{10}}$-heating value in suburb was higher (26%) than that in urban area (23%) (Fig. 4). During the CAHP, the meanly decreased concentration of PM$_{10}$ in urban area was 1 $\mu$g/m$^3$, and apparently lower than that of suburb (36 $\mu$g/m$^3$), as well as the mean $P_{\text{PM}_{10}}$-action values in urban area and suburb were 0.4% and 12%, respectively (Fig. 4). It can be seen that the control of PM$_{10}$ sources in suburb was more effective compared to urban area, in case of exclusion of unfavorably meteorological conditions (Table 1 and Fig. S7), probably related to more than 700 enterprises closed down which mainly carried out ore mining and stone processing in suburb (Tables S1 and S5). The tendency of NO$_2$ concentrations in urban area and suburb was similar to that of PM$_{10}$ during different stages of TECA period. The mean $P_{\text{NO}_2}$-heating values in urban area and suburb were up to 31% and 34%, respectively; while the mean $P_{\text{NO}_2}$-action values in urban area and suburb were up to 17% and 21%, respectively. Note that the concentrations of CO in urban area and suburb showed an increasing tendency from the NCANHP (2.1-2.4 mg/m$^3$) to the ACA period (5.5 mg/m$^3$) (Fig. 3). The $P_{\text{CO}}$-heating and $P_{\text{CO}}$-action values in urban area were 22% and -15%, respectively, while those in suburb were 32% and -20% during the TECA period. In addition, as shown in Fig. 5, the concentrations of CO in the eastern and northern suburb in Shijiazhuang were significantly higher than those of urban areas (t-test, $p<0.01$). Note that the concentrations of O$_3$ in urban area and suburb were lower during different stages of TECA (Fig. 5). Overall, during the TECA period, the effect of control measures for atmospheric pollutants sources in suburb was better than in urban area, especially for the effect of control measures for particulate matters sources. The effect of control measures for CO was not notable during the TECA period, especially in suburb, likely due to the increasing usage of domestic coal in suburb along with the temperature decreasing (Table 1).
3.2 Variations of chemical species in PM$_{2.5}$

The average concentrations of chemical species in PM$_{2.5}$ in Shijiazhuang during the whole sampling period are shown in Fig. 6. The annual mean concentrations of OC, SO$_2^-$, NO$_3^-$ and NH$_4^+$ in PM$_{2.5}$ were 43.1 μg/m$^3$, 39.0 μg/m$^3$, 33.6 μg/m$^3$ and 25.6 μg/m$^3$, respectively, and their contributions to PM$_{2.5}$ were up to 23.1 %, 20.0 %, 17.3 % and 12.3 %, respectively. The annual mean concentrations of EC and Cl$^-$ were 11.7 μg/m$^3$ and 7.7 μg/m$^3$, respectively, which accounted for 5.9 % and 4.1 % of PM$_{2.5}$. Note that the annual mean concentrations of elements in PM$_{2.5}$ were relatively lower, which varied from 0.03 to 2.6 μg/m$^3$, accounting for 0.02-2.4 % of PM$_{2.5}$.

Compared to other elements, the annual mean concentrations of Si (2.6 μg/m$^3$) and Al (1.4 μg/m$^3$) were relatively higher during the whole sampling period, which accounted for 2.4 % and 1.2 % of PM$_{2.5}$, respectively. In this study, the annual mean concentrations of OC, SO$_2^-$, NO$_3^-$ and NH$_4^+$ in Shijiazhuang were clearly higher than Beijing (Gao et al., 2016), Tianjin (Wu et al., 2015), Jinan (Gao et al., 2011), Shanghai (Wang et al., 2016c), Chengdu (Tao et al., 2013b), Xian (Wang et al., 2015a), Hangzhou (Liu et al., 2015) and Heze (Liu et al., 2017a).

The values of $P_{\text{heating}}$ and $P_{\text{action}}$ of different chemical species in PM$_{2.5}$ were calculated by using the Eq. (8) and (9). The variations of chemical species in PM$_{2.5}$ at four stages of the TECA and the values of $P_{\text{heating}}$ and $P_{\text{action}}$ in Shijiazhuang are shown in Figs. 7 and 8. Compared to the NCANHP, the concentrations of chemical species during the NCAHP showed a significantly increased tendency (t-test, $p<0.01$), the concentrations of SO$_2^-$, Cl$^-$, OC, EC, Si, Al, Ca$^{2+}$ and Mg$^{2+}$ increased by 7.9, 3.7, 6.7, 3.2, 1.6, 0.6, 0.4 and 0.1 μg/m$^3$, respectively, and the $P_{\text{heating}}$ values of these species were up to 30.0 %, 40.2 %, 14.6 %, 22.1 %, 78.8 %, 63.5 %, 47.4 % and 45.9 %, respectively, during the NCAHP. As these species (i.e., SO$_2^-$, Cl$^-$, OC, EC, Si, Al, Ca$^{2+}$ and Mg$^{2+}$) were closely associated with coal combustion (Cao et al., 2011; Liu et al., 2015; Liu et al., 2016; Liu et al., 2017a, c), therefore, coal combustion for heating in winter probably had a great impact on increasing of these chemical species in PM$_{2.5}$. Furthermore, compared to the NCANHP, the concentrations of Cr, Cu, Fe, Mn, Ti, Zn and Pb increased by 0.02, 0.02, 0.34, 0.02, 0.02, 0.28 and 0.07 μg/m$^3$, respectively, and the $P_{\text{heating}}$ values of these species were 72.7 %, 33.1 %, 34.4 %,
During the NCAHP, the concentrations of Cr, Cu, Fe, Mn, Ti, Zn and Pb were closely related to industrial sources (Liu et al., 2015; Kabala and Singh, 2001; Morishita et al., 2011; Mansha et al., 2012; Yao et al., 2016), thus, the industrial emissions might have a higher influence on PM$_{2.5}$ during the NCAHP than that during the NCANHP. Also, it might be closely associated with the unfavorably meteorological factors (Table 1 and Fig. S7).

Compared to the NCAHP, the concentrations of SO$_4^{2-}$, Cl$^-$, OC and EC during the CAHP increased by 16.8, 0.3, 19.8 and 14.6 μg/m$^3$, respectively, and the P$_{1-action}$ values of which were up to -48.8 %, -2.0 %, -37.3 % and -83.0 %, respectively, during the CAHP. As coal combustion was an important source of SO$_4^{2-}$, Cl$^-$, OC and EC (Cao et al., 2011; Liu et al., 2015; Liu et al., 2016; Liu et al., 2017a, c), so it can be inferred that the influence of coal combustion might increase apparently during the CAHP compared to the NCAHP, which was likely due to the increased usage of the coal for domestic heating with the reduction of temperature during winter (Table 1). Additionally, unfavorably meteorological conditions during the CAHP can have an offset effect on the control measures for coal-combustion sources. As also Fig. 5 shown that the concentrations of CO during the CAHP were higher than those during the NCAHP, especially in rural areas. Furthermore, OC and EC were associated with the vehicle exhaust (Liu et al., 2016; Liu et al., 2017a), thus, the effect of motor vehicle management and control measures during the CAHP might be offset by the unfavorably meteorological conditions to some extent during the CAHP (Table 1 and Fig. S7). However, compared to the NCAHP, the concentrations of Si, Al, Ca$^{2+}$ and Mg$^{2+}$ during the CAHP decreased by 1.1, 0.1, 0.6 and 0.1 μg/m$^3$, respectively, and the P$_{1-action}$ values of which were up to 30.3 %, 4.5 %, 47.0 % and 45.2 %, respectively. As Si, Al, Ca$^{2+}$ and Mg$^{2+}$ were mainly originated from the crustal dust (Liu et al., 2016; Shen et al., 2010; Wang et al., 2015a; Yang et al., 2016b), therefore, the influence of crustal dust on PM$_{2.5}$ during the CAHP might decrease clearly compared to the NCAHP. That’s closely related to the control measures of inhabitation of dust emission during the TECA period (as shown in section 2.5). In general, from the view of the variation of PM$_{2.5}$ speciation, there was no doubt that the TECA had a certain positive environmental effect on the improvement of air quality. However, the ambient pollutant concentration was impacted by not only the emission sources, but also the meteorological conditions, regional background level and distant transportation, it was understandable that the concentration of CO had a “rebound” effect during the CAHP as the height of mixing layer was only 474 m and a low wind.
speed of 0.4 m/s.

Fig. 6. The average concentrations and percentages of chemical species in PM$_{2.5}$ in Shijiazhuang during the whole sampling period: November 24, 2015 to January 9, 2017.

Fig. 7. The variations of chemical species in PM$_{2.5}$ during the four stages (NCANHP, NCAHP, CAHP and ACA) of the TECA period.

Fig. 8. The $P_{\text{heating}}$ and $P_{\text{action}}$ of chemical species in PM$_{2.5}$ during the TECA period in Shijiazhuang.

3.3 Variations of PM$_{2.5}$ sources contributions

The filter membrane samples of PM$_{2.5}$ were collected in three sites (LQ, LC and TSMS) in Shijiazhuang from November 24, 2015 to January 9, 2017, and source apportionment was carried out by using EPA PMF 5.0, as well as five factors were identified during the period (Figs. 9 and 10).

The chemical profile of factor 1 was mainly represented by Si (72.3 %), Ca$^{2+}$ (74.0 %), Mg$^{2+}$ (43.9 %) and Al (71.3 %), which were derived mainly from crustal dust (Liu et al., 2016; Shen et al., 2010; Wang et al., 2015a). Thus, factor 1 was viewed as crustal dust. The contribution proportions of factor 1 to PM$_{2.5}$ decreased from 19.5 % (38.5 μg/m$^3$) during the WY, 18.7 % (42.1 μg/m$^3$) during the NCANHP, 16.9 % (48.0 μg/m$^3$) during the NCAHP to 15.0 % (40.3 μg/m$^3$) during the CAHP, and increased up to 16.3 % (48.3 μg/m$^3$) during the ACA. The main species of factor 2 were SO$_4^{2-}$ (53.9 %), NO$_3$ (89.8 %) and NH$_4^+$ (75.0 %). Therefore, it was easily identified as secondary sources (Liu et al., 2015, 2016, 2017a; Santacatalina et al., 2010; Srimuruganandam and Nagendra, 2012). The contribution proportions of factor 2 to PM$_{2.5}$ ranged from 29.5 % (66.4 μg/m$^3$) during the NCANHP, 30.8 % (87.9 μg/m$^3$) during the NCAHP, 31.6 % (84.8 μg/m$^3$) during the CAHP to 32.7 % (64.6 μg/m$^3$) during the WY, and decreased to 28.8 % (85.2 μg/m$^3$) during the ACA. Factor 3 was represented by the relatively high loadings of OC (55.9 %), EC (70.9 %), Cu (26.9 %) and Zn (26.5 %). Given that the OC and EC are generally predominant in the reported source profile of vehicle exhaust (Liu et al., 2016, 2017a; Yao et al., 2016), and Zn is widely used as an additive for lubricant in two-stroke engines, and Cu is closely associated with brake wear (Begum et al., 2004; Canha et al., 2012; Lin et al., 2015; Liu et al., 2017a). Therefore, factor 3 was identified as vehicle emissions. The contribution proportions of factor 3 to PM$_{2.5}$ decreased from 14.2 % (32.0 μg/m$^3$) during the NCANHP, 13.4 % (26.4 μg/m$^3$) during the WY, 13.3 % (37.8 μg/m$^3$) during the NCAHP to 10.6 % (28.5 μg/m$^3$) during the CAHP, and increased to 14.1 % (41.7 μg/m$^3$) during the ACA.
Factor 4 was characterized by the high contributions of Ca\(^{2+}\) (26.0 %), Mg\(^{2+}\) (31.0 %), Si (13.3 %), As (84.9 %), Cl\(^{-}\) (38.6 %), OC (20.2 %) and SO\(_{4}\)^{2-} (26.7 %), and the combination of these species in factor 4 inferred they were co-emission from coal combustion (Cao et al., 2011; Liu et al., 2015, 2016, 2017a,c; Zhang et al., 2011). Therefore, factor 4 was identified as coal combustion. The contribution proportions of factor 4 to PM\(_{2.5}\) increased from 26.2 % (51.7 \(\mu g/m^3\)) during the WY, 28.0 % (63.2 \(\mu g/m^3\)) during the NCANHP, 29.5 % (84.0 \(\mu g/m^3\)) during the NCAHP to 31.7 % (85.2 \(\mu g/m^3\)) during the CAHP, and lightly increased to 32.6 % (96.3 \(\mu g/m^3\)) during the ACA. Factor 5 was identified as industrial emissions, with high loadings of Cr (66.7 %), Cu (63.7 %), Fe (83.2 %), Mn (51.3 %), Ti (70.0 %), Zn (69.2 %), Pb (42.1 %) and Cl\(^{-}\) (41.0 %) (Almeida et al., 2015; Liu et al., 2015, 2016; Morishita et al., 2011; Mansha et al., 2012; Yao et al., 2016). The contribution proportions of factor 5 to PM\(_{2.5}\) ranged from 5.0 % (11.3 \(\mu g/m^3\)) during the NCANHP, 5.1 % (10.0 \(\mu g/m^3\)) during the WY to 5.9 % (16.7 \(\mu g/m^3\)) during the NCAHP, and decreased to 5.3 % (14.2 \(\mu g/m^3\)) during the CAHP and 4.9 % (14.4 \(\mu g/m^3\)) during the ACA. Note that the contribution of industrial emissions to PM\(_{2.5}\) was relatively lower than other sources (Fig. 10).

In general, crustal dust, secondary sources, vehicle emissions, coal combustion and industrial emissions were identified as PM\(_{2.5}\) sources in Shijiazhuang (Fig. 9). Compared to the WY and NCANHP, the contribution concentrations and proportions of coal combustion to PM\(_{2.5}\) increased significantly during other stages of TECA period (Fig. 10), which was closely associated with the coal heating for cold winter (Liu et al., 2016), and the unfavorably meteorological conditions (Table 1 and Fig. S7). The contribution concentrations and proportions of crustal dust and vehicle emissions to PM\(_{2.5}\) decreased apparently during the CAHP compared to other stages of TECA period (Fig. 10). It indicated that the control effects of motor vehicles and crustal dust were remarkable during the CAHP, even under unfavorably meteorological conditions (Table 1), and the results were consistent with the above analysis. The contribution proportions of secondary sources to PM\(_{2.5}\) during the CAHP showed little change compared to other stages of TECA period (Fig. 10). However, compared to the WY and NCANHP, the contribution concentrations of secondary sources to PM\(_{2.5}\) increased significantly during the NCAHP, CAHP and the ACA (Fig. 10), likely due to high concentrations of gaseous precursors (i.e., SO\(_2\) and NO\(_2\)) (Fig. 5), unfavorably meteorological conditions (Table 1), and frequent hazy events during these periods, when there were significant secondary reactions (Han et al., 2014; Li et al., 2016a). In addition, it also illustrated that the
discharge of atmospheric pollutants might be still enormous even under such strict control measures.

Note that the contribution concentrations and proportions of industrial emissions to PM$_{2.5}$ during the CAHP decreased apparently compared to the NCAHP (Fig. 10), indicating that the control of industrial emissions was also effective during the CAHP.

Chen et al. (2016b) reported that the concentrations of particles during the 2014 Youth Olympic Games (YOG) period (August) were much lower than before-Games period (July) and after-Games period (September); and fugitive dusts, construction dusts and secondary sulfate aerosol decreased obviously in YOG, which means mitigation measures have played an effective role in reduction of particulate matter. Wang et al. (2017) found that the contributions of vehicles, industrial sources, fugitive dust, and other sources decreased 13.5-14.7 %, 10.7-11.2 %, 4.5-5.6 % and 1.7-2.7 %, respectively, during the Asia-Pacific Economic Cooperation (APEC) and China’s Grand Military Parade (CGMP), compared to the period before the control actions. Guo et al. (2013) found that primary vehicle contributions were reduced by 30 % at the urban site and 24 % at the rural site, compared with the non-controlled period before the Beijing 2008 Olympics. The reductions in coal combustion contributions were 57 % at PKU site and 7 % at Yufa site. As we can see that these control-actions of the strict measures taken for emission sources during the international events held in China, including the TECA in Shijiazhuang, were all very important practical exercises and rarely scientific experiments. However, it cannot be advocated as the normalized control measures for atmospheric pollution in China. These strict measures taken during these periods are temporary, and there is a normal recovery of all the emissions of sources after the operation. Once adverse weather conditions occur, and the hazy events may continue to happen eventually. In short, the direct cause of the severe atmospheric pollution in China is that the emission of pollutants beyond the air environment’s self-purification capacity, and the essential reason is unreasonable and unhealthy pattern for economic development of China.
### 3.4 Backward trajectory and PSCF analysis

The backward trajectory analysis was used to identify the transport pathways of the air mass during the CAHP. In terms of the directions and travelled areas, these trajectories were divided into the five groups (Fig. 11). Trajectory clusters 1, accounting for 31.3% of the total, originated from Shanxi province and passed over North of Hebei before arriving at Shijiazhuang. Trajectory cluster 1 reflected the features of small-scale, short-distance air mass transport (Fig. 11). The higher concentrations of PM$_{10}$ (358 μg/m$^3$), PM$_{2.5}$ (237 μg/m$^3$) and CO (3.9 μg/m$^3$) might be due to the variety of emission sources and the accumulation of pollutants from surrounding areas, since the moving speed of air mass in cluster 1 was much lower than other trajectories (Fig. 11 and Table 2).

Trajectory cluster 2, 3 and 4 accounted for 58.0% of the total trajectories, and began from the northwest of China, passed through the Inner Mongolia and Shanxi, showing the features of large-scale, long-distance air transports. The relative lower concentrations of PM$_{10}$ (189-290 μg/m$^3$), PM$_{2.5}$ (119-181 μg/m$^3$), SO$_2$ (50-67 μg/m$^3$), NO$_2$ (58-78 μg/m$^3$) and CO (2.1-3.0 mg/m$^3$) were closely associated with high moving speeds of air mass (Fig. 11 and Table 2), and relatively less anthropogenic emission sources in the northwest of China. Trajectory cluster 5 was mainly originated from Ningxia province, passed over Shaanxi, Shanxi and Hebei before arriving at Shijiazhuang, accounting for 10.8% of the total, showing the features of small-scale, short-distance air transport significantly elevated levels of PM$_{10}$ (451 μg/m$^3$), PM$_{2.5}$ (303 μg/m$^3$), SO$_2$ (83 μg/m$^3$), NO$_2$ (104 μg/m$^3$) and CO (4.8 mg/m$^3$) with trajectory cluster 5 were associated with the sources and the accumulation of pollutants from surrounding areas. As well known that the Beijing-Tianjin-Hebei region was one of the severest polluted areas in China (Bi et al., 2014; Chen et al., 2013; Gu et al., 2011; Wang et al., 2014; Zhao et al., 2012), it might be an important reason why the concentrations of pollutants were higher with trajectory clusters 1 and 5 (Fig. 11 and Table 2).

In this study, PSCF model was used to analyze the potential sources-areas of atmospheric pollutants by combining backward trajectories and the concentrations of atmospheric pollutants in Shijiazhuang during the CAHP, and the results were shown in Fig. 12. The values of weighted potential source contribution function (WPSCF) of CO were higher in the north of Shaanxi, south of Shanxi and central and southern Inner Mongolia, which were mainly potential sources-areas of CO concentrations in Shijiazhuang (Fig. 12 (a)). The WPSCF values of NO$_2$ were higher in north of Henan and Shaanxi, Hebei, Shanxi, and central and southern Inner Mongolia, which were mainly
potential sources-areas of NO$_2$ concentrations in Shijiazhuang (Fig. 12 (b)). The WPSCF values of O$_3$ and SO$_2$ were higher in the north of Henan and Shaanxi, Shanxi, and south of Hebei, which were distinguished as major potential sources-areas of O$_3$ and SO$_2$ concentrations in Shijiazhuang (Fig. 12 (c) and (d)). Moreover, the southwest of Shandong was also identified as mainly potential sources-areas of SO$_2$ concentrations in Shijiazhuang. As for PM$_{2.5}$ and PM$_{10}$, the WPSCF values were higher in south of Hebei, and east of Shanxi, which were identified as mainly potential sources-areas of PM$_{2.5}$ and PM$_{10}$ concentrations in Shijiazhuang (Fig. 12 (e) and (f)). Overall, the potential sources-areas of the atmospheric pollutants in Shijiazhuang mainly concentrated in the surrounding regions of Shijiazhuang, including south of Hebei, north of Henan and Shanxi. Previous studies also reported that Shanxi, Hebei and Henan provinces had serious air pollution problems (Feng et al., 2016; Kong et al., 2013; Meng et al., 2016; Zhu et al., 2011), revealing the regional nature of the atmospheric pollution in Northern Plain of China. Therefore, there is an urgent need for making cross-boundary control policy except for local control-measures given the high background level of pollutants.

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**Fig. 11.** Five clusters of the 72-h air mass backward trajectories during the CAHP. Red star represents Shijiazhuang city.

**Fig. 12.** Potential sources areas of atmospheric pollutants obtained from PSCF model during the CAHP. Red star represents Shijiazhuang city. The colors represent potential sources-areas influenced on the atmospheric pollutants, and the red color could be determined to be relatively important sources-areas while the blue color means unimportant potential sources-areas.

**Table 2.** The average concentrations of atmospheric pollutants in different clusters during the CAHP.

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### 4 Conclusions

The control measures of atmospheric pollution in Shijiazhuang were effective and was in a right direction. Under unfavorably meteorological conditions, the mean concentrations of PM$_{2.5}$, PM$_{10}$, SO$_2$, NO$_2$, and chemical species (Si, Al, Ca$_{2+}$, Mg$_{2+}$) in PM$_{2.5}$ during the CAHP significantly decreased compared to the NCAHP. Overall, the effects of control measures in suburbs were better than in urban area, especially for the effects of control measures for particulate matters sources. The effects of control measures for CO emission sources were not apparent during the CAHP, especially in suburbs.

The pollutant’s emission sources during the CAHP were in effective control, especially for
crustal dust and vehicles. While the necessary coal heating for cold winter and the unfavorably meteorological conditions had an offset effect on the control measures for emission sources to some extent. The discharge of pollutants might be still enormous even under such strict control measures.

The backward trajectory and PSCF analysis in the light of atmospheric pollutants suggested that the potential sources-areas mainly concentrated in surrounding regions of Shijiazhuang, i.e., south of Hebei, north of Henan and Shanxi. The regional nature of the atmospheric pollution in Northern China Plain revealed that there is an urgent need for making cross-boundary control policy except for local control-measures given the high background level of pollutants.

The TECA is an important practical exercise but it can’t be advocated as the normalized control measures for atmospheric pollution in China. The direct cause of atmospheric pollution in China is the emission of pollutants exceeds the air environment’s self-purification capacity, and the essential reason is unreasonable and unhealthy pattern for economic development of China.

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Reference


Dimitriou, K., Remoundaki, E., Mantas, E., and Kassomenos, P.: Spatial distribution of source areas of PM$_{2.5}$ by Concentration Weighted Trajectory (CWT) model applied in PM$_{2.5}$ concentration and composition data, Atmos. Environ., 116, 138–145, 2015.


Jiang, B. F., and Xia, D. H.: Role identification of NH$_3$ in atmospheric secondary new particle


Srimuruganandam, B., and Nagendra, S. M. S.: Application of positive matrix factorization in characterization of PM$_{10}$ and PM$_{2.5}$ emission sources at urban roadside, Chemosphere, 88, 120–130, 2012.

Sun, X., Yin, Y., Sun, Y. W., Sun, Y., Liu, W., and Han, Y.: Seasonal and vertical variations in aerosol distribution over Shijiazhuang, China, Atmos. Environ., 81, 245–252, 2013.

Tai, A. P. K., Mickley, L. J., and Jacob, D. J.: Correlations between fine particulate matter (PM$_{2.5}$) and meteorological variables in the United States: implications for the sensitivity of PM$_{2.5}$ to climate change, Atmos. Environ., 44, 3976–3984, 2010.


Yang, H. N., Chen, J., Wen, J. J., Tian, H. Z., and Liu, X. G.: Composition and sources of PM$_{2.5}$


Fig. 1. Maps of the online monitoring stations and the filter membrane sampling sites in Shijiazhuang. The 24 online monitoring stations mainly include Twenty-second Middle School (TSMS), Fenglong Mountain (FLM), High-tech Zone (HTZ), Great Hall of the People (GHP), Century Park (CP), Water Source Area in the Northwest (WSAN), University Area in the Southwest (UAS), Staff Hospital (SH), Gaoyi (GY), Gaocheng (GC), Xingtang (XT), Jinzhou (JZ), Jingxing Mining District (JXMD), Lingshou (LS), Luquan (LQ), Luancheng (LC), Pingshan (PS), Shenze (SZ), Wuji (WJ), Xinle (XL), Yuanshi (YS), Zanhuang (ZH), Zhaoxian (ZX), and Zhengding (ZD). The filter membrane sampling sites are mainly located in TSMS, LQ and LC.

Fig. 2. The variations of atmospheric pollutants concentrations during the four stages (NCANHP, NCAHP, CAHP and ACA) of the TECA period in Shijiazhuang.
Fig. 3. The concentrations variations of PM$_{2.5}$, PM$_{10}$ and gaseous pollutants during the four stages (NCANHP, NCAHP, CAHP and ACA) of the TECA period in Shijiazhuang. Error bar represented standard deviation.

Fig. 4. The P$_{\text{heating}}$ and P$_{\text{action}}$ of PM$_{2.5}$, PM$_{10}$ and gaseous pollutants (SO$_2$, NO$_2$, CO and O$_3$) calculated by equation (8) and (9) in urban area and suburb in Shijiazhuang.
Fig. 5. The spatial variations of atmospheric pollutants (PM$_{2.5}$, PM$_{10}$, SO$_2$, NO$_2$, CO and O$_3$) during the four stages (NCANHP, NCAHP, CAHP and ACA) of the TECA period in Shijiazhuang. The pictures were produced by ArcGIS based kriging interpolation method.
Fig. 6. The average concentrations and percentages of chemical species in PM$_{2.5}$ in Shijiazhuang during the whole sampling period: November 24, 2015 to January 9, 2017. Error bar represented standard deviation.

Fig. 7. The variations of chemical species in PM$_{2.5}$ during the four stages (NCANHP, NCAHP, CAHP and ACA) of the TECA period. Error bar represented standard deviation.
Fig. 8. The $P_{i\text{-heating}}$ and $P_{i\text{-action}}$ of chemical species in PM$_{2.5}$ during the TECA period in Shijiazhuang.

Fig. 9. Source profiles obtained with the PMF for PM$_{2.5}$. Filled bars identify the species that mainly characterize each factor profile.
Fig. 10. Source contributions of PM$_{2.5}$ during different stages in Shijiazhuang. WY represents whole year: November 24, 2015 to January 9, 2017.

Fig. 11. Five clusters of the 72-h air mass backward trajectories during the CAHP. Red star represents Shijiazhuang city.
Fig. 12. Potential sources areas of atmospheric pollutants obtained from PSCF model during the CAHP. Red star represents Shijiazhuang city. The colors represent potential sources-areas influenced on the atmospheric pollutants, and the red color could be determined to be relatively important sources-areas while the blue color means unimportant potential sources-areas.
Table 1. The meteorological conditions during the four stages (NCANHP, NCAHP, CAHP and ACA) of the TECA period in Shijiazhuang.

<table>
<thead>
<tr>
<th></th>
<th>NCANHP</th>
<th>NCAHP</th>
<th>CAHP</th>
<th>ACA</th>
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<tr>
<td></td>
<td>Ave.</td>
<td>S.D.</td>
<td>Ave.</td>
<td>S.D.</td>
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<tr>
<td>Temperature (℃)</td>
<td>8.4</td>
<td>3.6</td>
<td>7.4</td>
<td>2.4</td>
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<td>Relative humidity (%)</td>
<td>77.7</td>
<td>17.0</td>
<td>73.4</td>
<td>15.7</td>
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<td>Wind speed (m/s)</td>
<td>0.7</td>
<td>1.2</td>
<td>0.6</td>
<td>0.6</td>
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<tr>
<td>Height of mixed layer (m)</td>
<td>540</td>
<td>144</td>
<td>590</td>
<td>274</td>
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Ave. represents average value, S.D. represents standard deviation. NCANHP represents the no control action and no heating period, NCAHP represents the no control action and heating period, CAHP represents the control action and heating period, and ACA represents after control action.

Table 2. The average concentrations of atmospheric pollutants in different clusters during the CAHP.

<table>
<thead>
<tr>
<th>Clusters</th>
<th>Probability of occurrence (%)</th>
<th>SO₂</th>
<th>NO₂</th>
<th>O₃</th>
<th>CO(mg/m³)</th>
<th>PM₁₀</th>
<th>PM₂.₅</th>
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<tr>
<td>1</td>
<td>31.3</td>
<td>68</td>
<td>88</td>
<td>14</td>
<td>3.9</td>
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<td>237</td>
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<tr>
<td>2</td>
<td>14.2</td>
<td>67</td>
<td>78</td>
<td>24</td>
<td>3.0</td>
<td>290</td>
<td>181</td>
</tr>
<tr>
<td>3</td>
<td>27.3</td>
<td>65</td>
<td>69</td>
<td>20</td>
<td>2.8</td>
<td>232</td>
<td>152</td>
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<tr>
<td>4</td>
<td>16.5</td>
<td>50</td>
<td>58</td>
<td>27</td>
<td>2.1</td>
<td>189</td>
<td>119</td>
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<tr>
<td>5</td>
<td>10.8</td>
<td>83</td>
<td>104</td>
<td>16</td>
<td>4.8</td>
<td>451</td>
<td>303</td>
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