Development of a High Temporal-Spatial Resolution Vehicle Emission Inventory Based on NRT Traffic Data and Its Impact on Air Pollution in Beijing

Part 2: Impact of vehicle emission on urban air quality

J. J. He¹, L. Wu¹, H. J. Mao¹, H. L. Liu², B. Y. Jing¹, Y. Yu³, P. P. Ren¹, C. Feng⁴, and X. H. Liu⁴

¹The College of Environmental Science & Engineering, Nankai University, Tianjin, China
²Chinese Academy of Meteorological Sciences, China Meteorological Administration, Beijing, China
³Clod & Arid Regions Environmental & Engineering Research Institute, Chinese Academy of Sciences, Lanzhou, China
⁴Tianjin Vehicle Emission Control Center, Tianjin, China

Correspondence to: H. J. Mao (hongjun_mao@hotmail.com); H. L. Liu (liuhongli@cams.cma.gov.cn)

Abstract

In a companion paper (Jing et al., 2015), a high temporal–spatial resolution vehicle emission inventory (HTSVE) for 2013 in Beijing has been established based on near real time (NRT) traffic data and bottom up methodology. In this study, based on the sensitivity analysis method of switching on/off pollutant emissions in the Chinese air quality forecasting model CUACE, a modeling study was carried out to evaluate the contributions of vehicle emission to the air pollution in Beijing main urban areas in the periods of summer (July) and winter (December) 2013. Generally, CUACE model had good performance of pollutants concentration simulation. The model simulation has been improved by using HTSVE. The vehicle emission contribution (VEC) to ambient pollutant concentrations not only changes with seasons but also changes over moment. The mean VEC, affected by regional pollutant transports significantly, is 55.4 and 48.5 % for NO₂, while 5.4 and 10.5 % for PM₂.₅ in July and December 2013, respectively. Regardless of regional transports, relative vehicle emission contribution (RVEC) to NO₂ is 59.2 and 57.8 % in July and December 2013, while 8.7 and 13.9 % for PM₂.₅. The RVEC to PM₂.₅ is lower than PM₂.₅ contribution rate for vehicle emission in total emission, which may be caused by easily dry deposition of PM₂.₅ from vehicle emission in near-
1 Introduction

In recent years, the serious atmospheric environment problems in China attract special attention from government, publics and researchers. Due to the control of coal combustion, the type of air pollution is changing from smoke to vehicle exhaust and mixed sources, and the secondary aerosols and regional transports play an important role in severe haze episodes (Zhang et al., 2006; Huang et al., 2014), which make it more difficult to control air pollution. Air pollution caused by traffic emission has become the main concern of pollution control, especially in metropolitan cities. Direct emission pollutants from road traffic include nitrogen oxides (NOx), carbon monoxide (CO), hydrocarbon (HC), particulate matter (PM) and so on (Zhou et al., 2005; Song and Xie, 2006). Based on RAINS-ASIA computer model, five sectors direct emissions of sulfur dioxide (SO2), nitrogen oxides (NOx) and carbon monoxide (CO) including industry, power, domestic, transportation and biofuels in 1990, 1995 and 2020 were estimated for China by Streets and Waldhoff (2000), the transportation sector contributed approximately 1 and 2% to total SO2 emissions, 9 and 12% to total NOx emissions, 14 and 22% to total CO emissions in 1990 and 1995. Traffic emission has a significant contribution to urban air pollution in many cities in China (Qin and Chan, 1993; Fu et al., 2001), while more stringent vehicle emission standards lead to simultaneous reduction of surface ozone (O3) and fine particulate matter (PM2.5) concentrations (Saikawa et al., 2011).

Beijing, as the capital of China, is one of the most important metropolitan cities in the world, providing a habitat for a population over 21 million. The number of vehicle in Beijing increased rapidly during the last decades and hit 5.5 million in 2014, putting an immense pressure on environment. A lot of researches on the impact of vehicle emission in Beijing have completed from different perspective. Hao et al. (2001) developed vehicle emission inventory and investigated the contribution of traffic on atmospheric pollutant concentrations utilizing a Gaussian dispersion model in 1995, and vehicle emission contributed 76.8 and 40.2% to total CO and NOx emissions, 76.5 and 68.4% to ambient CO and NOx concentrations. During the Sino-African summit in 2006, the number concentrations of the particles and accumulation modes seemingly reduced by 20–60% due to the strict traffic restrictions (Cheng et al., 2008). Zhang et al. (2011) evaluated the effectiveness of air
pollution control through traffic restriction measure in August 2007 and discovered road mobile
sources were more effective on dust elements than anthropogenic elements of PM. Based on positive
matrix factorization (PMF), Liu et al. (2014) investigated the source apportionment of ambient fine
particle and found the vehicle emission was mainly responsible for particles in the size range 10–50
nm and accounted for 47.9 % of particle number concentration during summertime in 2011. A series
of emission control measurements and atmospheric observations during the 2008 Beijing Olympic
Games created a valuable case to research the effectiveness of control measures on mitigating air
pollution. It was illustrated that the black carbon (BC) concentration after traffic control during
Olympic decreased 74 %, and diesel trucks were a major contribution to the ambient summertime BC
levels (X. Wang et al., 2009). With the 32.3 % traffic flow reduction, numerical simulation revealed
the average reduction rate of PM$_{10}$, CO, and NO$_2$ were 28, 19.3 and 12.3 % respectively, but an
increase rate of O$_3$ was 25.2 % (Wang and Xie, 2009). Compared with uncontrolled period, on-road
air pollutant concentrations during the Olympics air pollution control period, which is concluded from
versatile mobile laboratory moving along Beijing’s Fourth Ring Road, decreased significantly, by up
to 54 % for CO, 41 % for NOx, 70 % for SO$_2$ and 12 % for BC. (M. Wang et al., 2009). Hence, there
is a certain controversy between previous studies and a significant fluctuation of pollutant
concentration contribution in different periods. Further researches should be conducted in traffic
emission effect on Beijing’s air quality resulted from air pollution and pollutants emission
characteristics changes in recent years and later on.

In a companion paper (Jing et al., 2015), based on NRT traffic data, high temporal–spatial resolution
vehicle emission inventory for 2013 in Beijing was established via a bottom up methodology. This
part (Part 2) utilizes Chinese Unified Atmospheric Chemistry Environment (CUACE) model to
simulate ambient pollutant concentrations and evaluate the contributions of vehicle emission in
Beijing main urban areas in periods of summer and winter 2013 based on the sensitivity analysis
method of switching on/off pollutant emissions. In Sect. 2, the details of the methods, datasets and
model setup are shown. CUACE model evaluation and the effect of new vehicle emission inventory
are presented in Sect. 3. The main conclusions are presented in Sect. 4.
2 Data and Method

2.1 Model description

Developed by China Meteorological Administration (CMA), CUACE model is used to simulate air quality for Beijing in this study. CUACE model is a unified chemical weather numerical forecasting system which is independent with weather and climate model. It consists of four functional blocks: anthropogenic and natural emissions; atmospheric gaseous chemical mechanisms; atmospheric aerosol chemical mechanisms; numerical assimilation system. Gaseous chemical block is based on the Regional Acid Deposition Model (RADM) covering 66 gaseous species (Stockwell et al., 1990; Wang et al., 2015). Aerosol module includes mixing scheme, clear-sky processes, dry deposition, below-cloud scavenging, in-cloud processes. Seven aerosol species, i.e. sulfates (SF), soil dust (SD), black carbon (BC), organic carbon (OC), sea salts (SS), nitrates (NI), and ammonium salts (AM) are considered in aerosol chemical module. The first six aerosol components were divided into 12 bins with diameter ranging between 0.01 and 40.96 μm. Based on the mixing assumptions, the ambient size and density of aerosols in a size bin are evaluated. The optical properties of these aerosols are readily computed when the mixing state, composition and ambient size are determined. The details of sulphur chemistry, cloud chemistry, coagulation, nucleation, condensation etc. were depicted by Gong et al. (2003). CUACE is online coupled to fifth-generation Penn State/NCAR mesoscale model (MM5) and Global/Regional Assimilation and PreDiction System (GRAPSE). MM5 is selected to simulate mesoscale meteorological fields in this study. For different research target and application purpose, CUACE is designed with open interface to make it easily being integrated to different time and spatial scale models. A more detailed description can refer to Gong et al. (2009). The performance of CUACE was evaluated by many researchers. Wang et al. (2010) simulated dust weather occurred in April 2006 and indicated CUACE model could predict the outbreak, development, transport and depletion processes of sand and dust storms accurately over China and the East Asian region. Li et al. (2014) evaluated air quality prediction by CUACE model over Urumqi and acquired a quite accurate forecasting on air quality levels, especially for NO$_2$ and PM$_{10}$ levels. Given the good performance in air quality prediction, CUACE model has been used for haze forecasting in National Meteorological
Center of CMA and some local environmental protection agencies.

2.2 Numerical simulation design

In this study, MM5-CUACE model is configured to have three nested domains to reduce spurious boundary effects in the inner domain with horizontal resolution of 27 km covering North China and the surrounding areas, 9 km covering Jing-Jin-Ji (Beijing, Tianjin and Hebei) areas and 3 km-resolution covering Beijing city and surrounding areas (Fig. 1). In the vertical, there are a total of 35 full eta levels extending to the model top at 10 hPa, with 16 levels below 2 km.

Two periods: July and December in 2013 are selected for model integration to evaluate different seasonal impact (summer and winter respectively) of vehicle emission on air quality. The time steps of MM5 and CUACE model are 15 s and 150 s respectively. Driving field provides the initial, lateral and surface boundary conditions and transmits the weather background information to MM5. However, for large domain or long term simulations, the large-scale weather situation simulated by MM5 may diverge from that of the driving field. The methods to constrain MM5 to the driving field involve frequent re-initialization, analysis nudging, spectral nudging, and scale-selective bias correction (Bowden et al., 2013). 36 h re-initialization run is executed to simulate meteorological conditions and air quality, and the former 12 h simulation is discarded as spin up time, which is the same as Zhang et al. (2012). The initial and boundary meteorological conditions are from T639 reanalysis data with 30×30 km spatial resolution and 6 h temporal resolution supplied by CMA (Xiao et al., 2010). The initial and boundary chemical conditions of the first simulation segment are based on averages from several field studies over eastern Pacific Ocean (McKeen et al., 2002) which was used as the default profiles in WRF-Chem, and other segment initial and boundary conditions are derived from previous simulation segment. The extra 10 day run (i.e. 21st June to 30th June, 21st November to 30th November) was conducted to reduce the effect of chemical initial and boundary conditions.

Two real simulations which based on default emission of CUACE and the improved emission with high temporal–spatial resolution vehicle emission (hereafter refer to HTSVE) are carried out to evaluate the accuracy of pollutant concentrations simulated by CUACE and analyze the influence of
HTSVE on Beijing air quality, and hereafter refer to SIM1 and SIM2 respectively. The contribution rate to ambient pollution level (or source apportionment) based on air quality numerical model includes source sensitivity simulations using the brute force method (also referred as zero-out method) or the decoupled direct method (DDM), air pollution tagged method, and the adjoint method (An et al., 2015; Burr and Zhang, 2011; Zhang et al., 2015). With comprehensible physical and chemical process, adjoint method has a significant advantage in source apportionment compared to sensitivity simulations or tagged method. However, the development of adjoint model is facing a challenge due to complicated mathematics and a large amount of data processing and programming, which results in less available regional scale air quality adjoint model. At recently, An et al. (2015) developed an adjoint of the aerosol module in the CUACE. The development of gaseous adjoint module of CUACE is needed for more widely application in source apportionment or source assimilation. The tagged method tracks contribution of pollutant from specific source and undergo the explicit atmospheric processes, but it is not able to simulate indirect effects and oxidant-limiting effects. With the ability in simulating indirect effects and relative simple model run, source sensitivity analysis is widely used in source attribution. However, significant source variation may result in misunderstanding due to non-linearity and atmospheric background concentrations change. In pervious study, the impact of Beijing local emission on air pollution is almost linear via source sensitivity analysis (An et al., 2007). Sensitivity analysis is suitable to investigate the contribution of vehicle emission in Beijing due to limited change of emission in this study. The vehicle emission contribution (VEC) to ambient pollutant concentration is computed based on the sensitivity analysis method of switching on (SIM2) and off (here after refer to SIM3) vehicle emission in Beijing. This method keeps atmospheric background pollution level basically steady which has a significant effect on the chemical conversion because of relative limited change of emission. Meanwhile the effect of vehicle emission on secondary pollution, e.g. secondary aerosol which becomes the important components of PM in Beijing (Huang et al., 2015) was considered. The formula of VEC is shown as follows:

$$\text{VEC} = \frac{C_{\text{SIM2}} - C_{\text{SIM3}}}{C_{\text{SIM2}}} \times 100\%$$

(1)

where C represents pollutant concentration. In fact, the regional transports of pollutants has obviously effect on VEC, and we calculate relative vehicle emission contribution (RVEC) which does not
consider pollutant regional transports, as shown in Eq. (2):

\[ RVEC = \frac{C_{SIM2} - C_{SIM3}}{C_{SIM2} - C_{SIM4}} \times 100\% \]  

(2)

where SIM4 represents the simulation of switching off all emission sources in Beijing. All simulation test schemes are listed in Table 1.

2.3 Emission inventory

CUACE model has an independent pollution emission module, which contains natural and anthropogenic emissions including many gas and particle matter emissions (Gong et al., 2009). Anthropogenic emissions of SO\(_2\), NO\(_x\), CO, VOCs, PM\(_{2.5}\), PM\(_{10}\), BC, OC, etc. used in emission module were developed by CMA based on INTEX-B inventory, the emissions database for global atmospheric research (EDGAR) and environmental statistics database. Gridded INTEX-B inventory covers 22 countries and regions in East Asia with a resolution of 0.5°×0.5°, and is classified into industry emission, power station emission, residential emission and vehicle emission (Zhang et al., 2009). The EDGAR is a joint project of the European Commission Joint Research Centre and the Netherlands Environmental Assessment Agency. The environmental statistics database is supplied by Environmental Protection Agency. Some old data was corrected or updated according to the variation rate of anthropogenic emissions from environmental statistics database. Finally emission inventory was pretreated by SMOKE for detailed temporal and spatial distribution. Hourly emissions were obtained for CUACE model input. The emission inventory is a key factor to air quality numerical simulation. Annual emissions of CO, NO\(_x\), SO\(_2\) and PM\(_{2.5}\) in CUACE model in Beijing are 3149.5, 173.8, 158.2 and 79.0 kt respectively. By comparing the different researches (Table 2) found that there are many uncertainties of inventories, especially for CO and NO\(_x\) emissions, but it is difficult to identify which one is more accurate. With rapid economic development and the adjustment of energy structure, anthropogenic emissions have a significant variation in recent years in North China. However, the database of emission inventory in previous studies (Table 2) is before 2010, which is the main reason for the differences between CUACE emission and others. For example, the Beijing municipal government has taken a strict traffic restriction since 2008. The amount of vehicle in Beijing increases about 8% in 2013. The change of vehicle emission maybe responsible for NO\(_2\)
emission variation. Except for date of basic data, the methods of establishing inventory, emission factors, basic data source would result in significant difference of emission inventory.

This study focus on vehicle source and its influence. HTSVE based on NRT traffic data was used to replace the vehicle emission in CUACE emission module to analyze its effects on air quality simulation. The detailed description of high temporal–spatial resolution vehicle emission and comparison with vehicle emission in CUACE emission module were presented in part 1. The contribution of major species from vehicle emission is presented in Table 3. The vehicle emission of NO, NO\textsubscript{2} and HC from HTSVE is higher, while CO and PM\textsubscript{2.5} is lower than that from CUACE.

2.4 Observational data

2.4.1 Meteorological data

The accuracy of mesoscale meteorological fields simulated by MM5 has a significant effect on air quality simulation, and it should be evaluated with observation data firstly. In this study, the observed near-surface meteorological fields including 2 m temperature, 2 m specific humidity and 10 m wind speed are obtained from Meteorological Information Comprehensive Analysis and Process System (MICAPS) of CMA. MICAPS surface data has eight conventional observation times everyday (00:00, 03:00, 06:00, 09:00, 12:00, 15:00, 18:00, 21:00 UTC) and 20 meteorological stations located in study region (Fig. 1a).

2.4.2 Air quality data

To evaluate simulated air quality by CUACE, hourly near-surface average concentrations of NO\textsubscript{2} and PM\textsubscript{2.5} from 9 atmospheric environment monitoring stations in Beijing (shown in Fig. 1b) in simulation periods were acquired from China National Environment Monitoring Centre. The monitoring stations distributed in study region could reflect different area pollution level and capture overall air quality in Beijing city.
3 Results and discussions

3.1 Model evaluation and the impact of new vehicle emission inventory

The accuracy of air quality simulation based on numerical model greatly relates to mesoscale meteorological simulation. Although the good performance of MM5 has obtained in many studies, the MM5’s results is verified firstly as the different accuracy of meteorological fields in different study domains, seasons and physical parameterizations. Based on statistical analysis, 2 m temperature root mean square error (RMSE) and correlation coefficient ($R$) are 3.4 K and 0.81 in July, 3.8 K and 0.87 in December. MM5 can capture temporal and spatial variation of near-surface temperature effectively. 2 m specific humidity RMSE and $R$ are 2.4 g kg$^{-1}$ and 0.56 in July, 0.9 g kg$^{-1}$ and 0.82 in December, which indicates that basic temporal and spatial variation of near-surface specific humidity is simulated by MM5. 10 m wind speed RMSE and $R$ are 1.4 m s$^{-1}$ and 0.37 in July, 1.7 m s$^{-1}$ and 0.57 in December. The RMSE was 1–4 K for 2 m temperature, 1–2 g kg$^{-1}$ for 2 m specific humidity and 1–4 m s$^{-1}$ for 10 m wind speed in most studies (Han et al., 2008; He et al., 2013; He et al., 2014; Jiménez-Guerrero et al., 2008; Kioutsioukis et al., 2016; Papalexiou and Moussiopoulos, 2006; Miao et al., 2008). In this study, MM5 presents the essential features of the local circulation over Beijing as seen from above analysis and its performance observed here is comparable to other studies generally. The details of meteorological evaluation are provided in supplement file. The statistic parameters could refer to He et al. (2014).

NO$_2$ and PM$_{2.5}$ are the major concerns as they are susceptible to vehicle emission. Interval of simulated and observed daily mean near-surface NO$_2$ and PM$_{2.5}$ concentrations averaged over 9 sites during two periods are shown in Fig. 2. CUACE model underestimates the NO$_2$ concentration significantly, especially during serious pollution periods. Due to the increasing emission of HTSVE (Table 2), the NO$_2$ concentration from SIM2 increases 31.8 and 11.1 % in July and December respectively, resulting in significant improvement to the previous underestimates. The RMSEs of NO$_2$ daily mean concentration decrease 17.6 and 10.9 % in two periods when HTSVE is used. Temporal correlation coefficients of NO$_2$ daily mean concentrations for SIM1 and SIM2 are 0.80 and 0.79 respectively in December, which indicates CAUCE can reproduce NO$_2$ time trends accurately.
However, low correlation (0.21 and 0.12 for SIM1 and SIM2 respectively) in July reflects the complexity of air quality numerical simulation. Simulated PM$_{2.5}$ daily mean concentration is basically consistent with observed value. Minor difference of PM$_{2.5}$ concentration is observed between SIM1 and SIM2 due to less vehicle emission change (Table 3). Based on temporal correlation analysis, SIM2 improves PM$_{2.5}$ time trends slightly, with correlation coefficients of 0.75 and 0.77 in two periods for SIM1, 0.76 and 0.78 for SIM2. Compared with SIM1, the RMSE of PM$_{2.5}$ daily mean concentration has slightly decrease for SIM2. It is obviously that simulated PM$_{2.5}$ concentration is more accurate than simulated NO$_2$ concentration in July, similar phenomena was found in previous studies (Roustan et al., 2011; Wu et al., 2011). CUACE’s ability is evaluated through the comparison of model grid and site station values, however, this method has several uncertainties because the local information is involved. It should be noted that the lifetime of ambient NO$_2$ is shorter than that of ambient PM$_{2.5}$ due to the different chemical processes, and local characteristics are more significantly for NO$_2$. The grid average concentration of NO$_2$ simulated by CUACE weakens the sub-grid local characteristics, and results in poor performance of NO$_2$ simulation compared with PM$_{2.5}$. The uncertainty of emission inventory increases with the spatial resolution of numerical model. Although vehicle emission was replaced with HTSVE, the uncertainty of emission inventory of other sectors in Beijing and all emissions in surrounding areas is still an important reason for the bias of pollutant concentrations. Seasonal difference of CUACE model performance is found in this study, with accurately simulation in winter, and this may relate to meteorological condition, especially on wind field bias as mentioned above. The uncertainty of photochemical reaction which is more significant in summer might result in large bias compared to the performance of NO$_2$ in winter. Overall, the performance of CUACE model is comparable with other studies in Beijing (Gao et al., 2011; Wu et al., 2011). As better performance acquired by SIM2, it is made as a baseline scenario in the following analysis.

Spatial distribution of pollutant concentration relates to pollutant emission distribution and meteorological condition. The spatial distribution of pollutant concentration from CUACE is basically consistent with sites observation (Fig. 3). The mean wind in Beijing urban region is the southwest wind in July, and drives local pollutant transports from southwest to northeast. The high
NO2 concentration is located in northeastern city, while two high PM2.5 concentration regions appear in west and center city (Fig. 3a and b). The spatial distribution of NO2 is different from that of PM2.5 because of emission sources distribution difference with one high emission area inner 5th ring road for NO2 and two high emission areas in west 6th ring road and inner 3rd ring road for PM2.5 (Fig. 4). High concentrations present in high emissions or its downwind. The mean concentrations of NO2 and PM2.5 are 29.8 and 91.3 μg m⁻³ in July. Beijing urban region is dominated by northwest wind in December, and pollutant concentration distribution is obviously different from that in July. NO2 concentration is high in southeast city, and gradually decreases outward (Fig. 3c). High PM2.5 concentration is mostly located in west and southeast city (Fig. 3d). It is found that significant difference presents in NO2 distribution between July and December while slightly difference for PM2.5 due to the combined effect of wind fields and emission distributions. The mean concentrations of NO2 and PM2.5 are 42.8 and 136.4 μg m⁻³ in December respectively.

### 3.2 The effect of vehicle emission on urban air quality

VEC on ambient pollutant concentration is analyzed through comparison simulation with and without vehicle emission (SIM2 and SIM3 respectively). Probability density function (PDF) is a good way to describe the total representation. The PDF of instantaneous VEC in two periods is shown in Fig. 5. The maximum frequencies of VEC to NO2 in July and December are appeared in 55–60 % and 50–55 % respectively. The frequencies of VEC to NO2 from 15 to 60 % in December are larger than that in July (Fig. 5a), which indicates large contribution presents in summer while small contribution presents in winter. Based on one-way analysis of variance, the difference of VEC to NO2 in summer and winter is significant. This may relates to seasonal differences of meteorological condition and pollutant emission. In summer, high temperature and strong solar radiation lead to strong atmosphere oxidation ability, and therefore it is easy to convert from NO to NO2, which results in large contribution to NO2 concentration. Meanwhile, the high rate of NO2 emission from vehicle (Table 3) is another reason for large contribution to ambient NO2 concentration in summer. The VEC to PM2.5 is considerably lower than that to NO2. The maximum frequencies of VEC to PM2.5 in July and December are appeared in 0–5 % and 5–10 % respectively. Different from NO2, the mean VEC to
PM\textsubscript{2.5} in summer is smaller than that in winter, with a significant difference from one-way analysis of variance. Relative humidity in summer is larger than that in winter, and high relative humidity is conductive to gas-particle conversion processes of other emission sources (Yao et al., 2014), which may be one of the reason for small VEC to PM\textsubscript{2.5} in summer. The strong turbulence mixing in summer makes rapidly vertical exchange and transport of pollutant in boundary layer, and finally results in small VEC to PM\textsubscript{2.5} in summer. Wind field variation is another reason for seasonal change of VEC to PM\textsubscript{2.5}, which will be investigated in the following part.

As the local transports of pollutants, the VEC in Beijing city depends on wind field and spatial distribution of vehicle emission. Wind dependency map of VEC to NO\textsubscript{2} and PM\textsubscript{2.5} are shown in Fig. 6. High VEC to NO\textsubscript{2} in July is appeared in south wind with 3–4 m s\textsuperscript{-1}, while north wind with 6–7 m s\textsuperscript{-1} for that in December. Due to the difference of lifetime between NO\textsubscript{2} and PM\textsubscript{2.5}, the wind dependency map to PM\textsubscript{2.5} is quite different from that to NO\textsubscript{2}. High VEC to PM\textsubscript{2.5} in July and December appeared in north wind due to many vehicle emission of particle matter in northeast city (Jing et al., 2015). The dominant wind is southwest wind in July and northwest in December (Fig. 3), which brings a small VEC to PM\textsubscript{2.5} in summer. Significant regional transport which is analyzed in next section is one of the reason for relative small VEC to PM\textsubscript{2.5} in summer.

Figure 7 shows time series of VEC to NO\textsubscript{2} and PM\textsubscript{2.5} daily mean concentrations in main urban areas (within the 6\textsuperscript{th} ring road) in two periods. The VEC not only changes with seasons, which is consistent with Cheng et al. (2007), but also changes with time. Time series of regional mean VEC is 49.8–60.0 \% to ambient NO\textsubscript{2} concentration in July, with a mean contribution rate of 55.4 \%. In December, regional mean contribution on NO\textsubscript{2} concentration decreases to 28.5–57.9 \% at different days, with a mean contribution rate of 48.5 \%. VEC to ambient PM\textsubscript{2.5} concentration is less than 10.3 and 13.6 \% at different times, with mean contribution rate of 5.4 and 10.5 \% in July and December respectively.

The change of VEC to PM\textsubscript{2.5} between July and December is most caused by meteorological condition in two periods. With different lift time of PM\textsubscript{2.5} and NO\textsubscript{2}, PM\textsubscript{2.5} concentration is more affected by regional transports, while NO\textsubscript{2} concentration is more affected by local emissions. Therefore the contribution with time variation for PM\textsubscript{2.5} is different from that for NO\textsubscript{2}. Except for wind field, pollution level is an important factor to VEC. It is obviously that low VEC presents in serious
pollution, while high VEC presents in low pollution concentration level, especially for NO\textsubscript{2} (Fig. 8). The absolute contribution of vehicle emission increases in severe pollution mostly because of adverse dispersion condition. However, pollutant regional transport is enhanced in severe pollution, which results in negatively correlation between VEC and pollution concentration level. The VEC has a significant spatial variation, previous study pointed that PM\textsubscript{2.5} had larger contribution from vehicle emission (13.0–16.3 % vs. 5.1 %) in urban as compared to that in suburban (S. W. Wu et al., 2014). Figure 9 shows the spatial distribution of mean contribution rate of vehicle emission in two periods. Vehicle emission contributes 26.0–76.4 % and 22.9–66.4 % of NO\textsubscript{2} at different regions in July and December. Significant effect of vehicle emission on ambient NO\textsubscript{2} concentration level is found in southeast and northeast city. VEC to PM\textsubscript{2.5} is 1.2–15.4 % and 2.4–24.4 % in July and December. The large contribution appears in northeast city in both summer and winter, which is widely different from the distribution of NO\textsubscript{2} contribution.

As can be seen from Table 4, receptor source apportionment and numerical sensitivity analysis are two main methods to compute VEC on ambient pollutant concentration, and VEC has significantly uncertainties from previous studies. In summary, vehicle emission contributes 4–17 and 22 % to PM\textsubscript{2.5} concentration based on receptor source apportionment and numerical simulation methods, and 56–74 % to NO\textsubscript{x} concentration based on numerical simulation method. The difference of the vehicle emission contribution to PM\textsubscript{2.5} with the different methods is relatively large. The uncertainties of VEC are related to sampling or simulation time, the location, analysis method and weather conditions. The results from receptor source apportionment (CMB, PMF etc.) only represent the characteristics of receptor point and can be applied for primary pollutants (Cheng et al., 2015), however it is different from numerical sensitivity analysis which normally describes the regional characteristics and applies for primary and secondary pollutants. The uncertainty of emission source in numerical model may be the main reason for significant difference to VEC in previous numerical studies. Though relatively short simulation in this study, our results are comparable with previous studies, and meanwhile keep the difference which comes from analyzing periods and method.

In this study, the rates of NO\textsubscript{2} and PM\textsubscript{2.5} from vehicle emission in total emission takes account for 55.1 and 22.3 % in July and 53.9 and 20.6 % in December (Table 3) of total emission. Because of the
effect of pollutant regional transports, the contribution rate of vehicle emission on ambient pollutant concentration is lower than the rate of vehicle emission in total emissions. The difference between these two rates became significantly larger with more contribution of outside emission, which implies the importance of weather condition. In order to avoid the effect of weather situation on analysis results, the relative contribution of vehicle emission on pollutant concentrations is analyzed in following section.

The chemical components of PM$_{2.5}$ represents the characteristics of emission source and complexity chemical processes of pollutant in atmosphere. Based on sensitivity test, the VECs of BC, OC and NI are large, while relative small for SF, and AM (Table 5). The VECs of BC and OC in December are approximately twice of that in July. Seasonal changes for the rates of BC and OC from vehicle emission in total emission are inapparent which indicates that it is not the reason for seasonal change of VECs. Beijing is controlled by southerly wind dominantly, which results in significant regional transport. And it causes small (large) VECs of BC and OC in summer (winter). Atmospheric chemical processes and dispersion conditions are also the reason for seasonal change of different components VECs. Using MM5-CMAQ model simulation, Cheng et al. (2013) investigated the VEC to the PM$_{2.5}$ and found the VEC of BC was 32.3% and 30.7% in summer and winter respectively. Our results are comparable with Cheng et al. (2013) in winter, while show some difference in summer.

3.3 Relative contribution of vehicle emission

Air pollution in Beijing is attributed not only from local emissions but also from regional transports. Using the CMAQ model, An et al. (2007) investigated the contribution to pollutant concentrations in Beijing by using emission switch on/off method, the contribution of non-local emission accounted for 15–53 % of PM$_{2.5}$. Wu et al. (2011) studied the contribution to air pollution during CAREBeijing-2006, and local emission in Beijing accounted for 65 % of SO$_2$, 75 % of PM$_{10}$ and 90 % of NO$_2$ concentrations. Pollutant regional transport depends on atmospheric circulation and regional emission characteristics. By comparing pollutant concentrations between SIM2 and SIM4, local emissions in Beijing contributes 93.6 % and 62.6 % to NO$_2$ and PM$_{2.5}$ concentrations in July, and 83.8 % and 76.1 % to NO$_2$ and PM$_{2.5}$ concentrations in December, which have a profound effect on RVEC.
Figure 10 depicts the spatial distribution of RVEC to NO$_2$ and PM$_{2.5}$ in July and December, and similar distribution is found in two periods. The RVEC to NO$_2$ is large in southeast and northeast main urban areas, while small in west main urban areas. Time series of regional mean RVEC to NO$_2$ in main urban areas range from 52.3 to 63.4 %, and 49.4 to 61.2 %, with the mean of 59.2 and 57.8 % in July and December respectively. Different from NO$_2$, the RVEC to PM$_{2.5}$ is large in northeast of main urban areas in two periods. Time series of regional mean RVEC to PM$_{2.5}$ range from 5.7 to 11.3 % and 9.9 to 16.1 %, with the mean of 8.7 and 13.9 % in July and December respectively. The differences of RVECs to NO$_2$ and PM$_{2.5}$ in July and December are significant based on one-way analysis of variance. The spatial distribution of RVEC are tremendously affected by vehicle emissions, as they are mostly consistent with the rate of vehicle emission in total emission (Fig. 4). As pointed by Jing et al., (2015), the uncertainty of HTSVE is very small through multiple comparison with statistical data and real time observation. But the uncertainty of other sector emissions has a negative influence on the precision of RVEC, which need more improvement for accurate environmental management.

Local circulation also determines the spatial distribution of RVEC. High PM$_{2.5}$ emission from vehicle is found between north Fourth Ring Road and north Five Ring Road (See Part. 1, Fig. 9). Controlled by southwest wind, PM$_{2.5}$ from vehicle is easily transferred out of the main urban areas, which results in low RVEC in July. However, the most of PM$_{2.5}$ from vehicle stay in east main city controlled by northwest wind, which results in high RVEC in December. Based on zero out method, Cheng et al. (2013) found the contribution rates to pollutant concentrations were higher than those to the emissions because near-surface emission from vehicle facilitated greater contribution to local pollutant concentrations on the ground level. Regardless of regional transports, the contribution of vehicle emission to ambient PM$_{2.5}$ concentration is substantial lower than the rate of vehicle emission in total emission in this study. Our finding is seemingly in conflict with Cheng et al. (2013), but may be more reasonable for following reasons. Different from elevated emission, PM$_{2.5}$ from vehicle emission in near-surface layer easily descends to the ground or is absorbed by vegetation, which leads to low contribution rate to PM$_{2.5}$ concentration. Secondary aerosol generated by photochemical reaction is different for different sector emissions. The VEC to SF is low in Beijing (Table 5), which indirectly causes low VEC to PM$_{2.5}$. Furthermore, pollutant regional transport and the background concentration...
may result in lower VEC to PM$_{2.5}$ than the rate of emission.

4 Conclusion

Air quality simulation has been improved by using HTSVE. In summer (July), high NO$_2$ concentration was located in the northeastern part of city, while two high PM$_{2.5}$ concentration regions appeared in west and center of the city. In winter (December), NO$_2$ concentration was high in southeast city, then gradually decreased outward, while high PM$_{2.5}$ concentration was mostly located in west and southeast part of city. The VEC in Beijing city depends on wind field, spatial distribution of vehicle emission and air pollution level. High VEC to NO$_2$ in July appeared along with south wind and low pollution concentration level, while north wind and low pollution concentration level for that in December. High VEC to PM$_{2.5}$ in July and December appeared along with north wind and low pollution concentration level.

Seasonal change of VEC was observed in this study. The mean VECs to NO$_2$ were 55.4 and 48.5 %, while the mean VECs to PM$_{2.5}$ were 5.4 and 10.5 % in July and December respectively. Regional pollutants transport was one of the most important reason for small contribution rate for ambient pollutant concentrations compared with contribution rate for pollutant emission in Beijing. Sensitivity analysis indicated that all local emissions in Beijing contributed 93.6 and 62.6 % to NO$_2$ and PM$_{2.5}$ concentrations in July, and 83.8 and 76.1 % to NO$_2$ and PM$_{2.5}$ concentrations in December, which had an important effect on RVEC. Regardless of regional transports, the RVEC to NO$_2$ was large in the southeast and northeast main urban areas, and northeast main urban areas for PM$_{2.5}$. The mean RVECs to NO$_2$ were 59.2 and 57.8 %, while the mean RVECs to PM$_{2.5}$ were 8.7 and 13.9 % in July and December respectively. The RVEC to PM$_{2.5}$ was lower than PM$_{2.5}$ contribution rate for vehicle emission, which was caused by easily dry deposition of PM$_{2.5}$ from vehicle emission in near-surface layer.

Acknowledgments

This work was supported by China’s National 863 program (2012AA063303), the National Science and Technology Infrastructure Program (2014BAC16B03), and the Opening Research Foundation of the Key Laboratory of Land Surface Process and Climate Change in Cold and Arid Regions, Chinese Academy of Sciences (LPCC201405).
Reference

An, X., Zhu, T., Wang, Z., Li, C., and Wang, Y.: A modeling analysis of a heavy air pollution episode occurred in Beijing, Atmos. Chem. Phys., 7, 3103-3114, doi:10.5194/acp-7-3103-2007, 2007.

An, X. Q., Zhai, X. S., Jin, M., Gong, S. L., Wang, Y.: Tracking influential haze source areas in North China using an adjoint model, GRAPES-CUACE, Geosci. Model Dev. Discuss., 8, 7313-7345, doi:10.5194/gmd-8-7313-2015, 2015.

Bowden, J. H., Nolte, C. G., and Otte, T. L.: Simulating the impact of the large-scale circulation on the 2-m temperature and precipitation climatology, Clim. Dynam., 40, 1903-1920, doi:10.1007/s00382-012-1440-y, 2013.

Burr, M., and Zhang, Y.: Source apportionment of fine particulate matter over the Eastern U.S. Part I: source sensitivity simulations using CMAQ with the brute force method. Atmospheric Pollution Research, 2, 299-316, 2011.

Cao, G. L., Zhang, X. Y., Gong, S. L., An, X. Q., and Wang, Y. Q.: Emission inventories of primary particles and pollutant gases for China, Chinese Sci. Bull., 56, 781-788, doi:10.1007/s11434-011-4373-7, 2011.

Cheng, L., Xu, X., Zhang, L.: Overview of receptor-based source apportionment studies for speciated atmospheric mercury, Atmos. Chem. Phys. Discuss., 15, 5493–5536, doi:10.5194/acpd-15-5493-2015, 2015.

Cheng, S. Y., Chen, D. S., Li, J. B., Wang, H. Y., and Guo, X. R.: The assessment of emission-source contributions to air quality by using a coupled MM5-ARPS-CMAQ modeling system: A case study in the Beijing metropolitan region, China, Environ. Modell. Softw., 22, 1601-1616, doi:10.1016/j.envsoft.2006.11.003, 2007.

Cheng, S. Y., Lang, J. L., Zhou, Y., Han, L. H., Wang, G., and Chen, D. S.: A new monitoring-simulation-source apportionment approach for investigating the vehicular emission contribution to the PM2.5 pollution in Beijing, China, Atmos. Environ., 79, 308-316, doi:10.1016/j.atmosenv.2013.06.043, 2013.

Cheng, Y. F., Heintzenberg, J., Wehner, B., Wu, Z. J., Su, H., Hu, M., and Mao, J. T.: Traffic restrictions in Beijing during the Sino-African Summit 2006: aerosol size distribution and visibility compared to long-term in situ observations, Atmos. Chem. Phys., 8, 7583-7594, doi:10.5194/acp-8-7583-2008, 2008.

Fu, L. X., Hao, J. M., He, D. Q., and He, K. B.: Assessment of vehicle pollution in China, J. Air Waste Manage., 51, 658-668, 2001.

Gao, Y., Liu, X., Zhao, C., and Zhang, M.: Emission controls versus meteorological conditions in determining aerosol concentrations in Beijing during the 2008 Olympic Games, Atmos. Chem. Phys., 11, 12437–12451, doi:10.5194/acp-11-12437-2011, 2011.

Gong, S. L., Barrie, L. A., Blanchet, J. P., von Salzen K., Lohmann, U., Lesins, G., Spacek, L., Zhang, L. M., Girard, E., Lin, H., Leaitch, R., Leighton, H., Chylek, P., and Huang, P.: Canadian aerosol module: a size-segregated simulation of atmospheric aerosol processes for climate and air quality models 1. Model development, J. Geophys. Res., 108, 4007, doi:10.1029/2001JD002002, 2003.

Gong, S. L., Zhang, X. Y., Zhou, C. H., Liu, H. L., An, X. Q., Niu, T., Xue, M., Cao, G. L., and Cheng, Y. L.: Chemical weather forecasting system CUACE and application in China’s regional haze forecasting, in: Proceeding of the 26th Annual Meeting of Chinese Meteorological Society,
Han, Z. W., Ueda, H., and An, J. L.: Evaluation and intercomparison of meteorological predictions by five MM5-PBL parameterizations in combination with three land-surface models, Atmos. Environ., 42, 233-249, doi:10.1016/j.atmosenv.2007.09.053, 2008.

Hao, J. M., Wu, Y., Fu, L. X., He, K. B., and He, D. Q.: Motor vehicle source contributions to air pollutants in Beijing. Environ. Sci., 22, 1-6, 2001.

Hao, J. M., Wang, L. T., Li, L., Hu, J. N., and Yu, X. C.: Air pollutants contribution and control strategies of energy-use related sources in Beijing, Sci. China Ser. D, 48, 138-146, 2005.

He, J. J., Yu, Y., Liu, N., Zhao, S. P.: Numerical model-based relationship between meteorological conditions and air quality and its implication for urban air quality management, Int. J. Environ. Pollut., 53, 265-286, 2013.

He J. J., Yu, Y., Liu, N., Zhao, S. P., Chen J. B.: Impact of land surface information on WRF’s performance in complex terrain area, Chinese J. Atmos. Sci., 38, 484-494, doi:10.3878/j.issn.1006-9895.2013.2014.

Huang, R. J., Zhang, Y. L., Bozzetti, C., Ho, K. F., Cao, J. J., Han, Y. M., Daellenbach, K. R., Slowik, J. G., Platt, S. M., Canonaco, F., Zotter, P., Wolf, R., Pieber, S. M., Bruns, E. A., Crippa, M., Ciarelli, G., Piazzalunga, A., Schwikowski, M., Abbaspade, G., Schnelle-Kreis, J., Zimmermann, R., An, Z. S., Szidat, S., Baltensperger, U., Haddad, I. E., and Prévôt, A. S. H.: High secondary aerosol contribution to particulate pollution during haze events in China, Nature, 514, 218-222, doi:10.1038/nature13774, 2014.

Jiménez-Guerrero, P., Jorba, O., Baldasano, J. M., and Gassó, S.: The use of a modeling system as a tool for air quality management: annual high-resolution simulation and evaluation, Sci. Total Environ., 390, 323-340, doi:10.1016/j.scitotenv.2007.10.025, 2008.

Jing, B. Y., L. Wu, H. J. Mao, S. L., Gong, J. J. He, C. Zou, G. H. Song, and X. Y. Li: Development of a High Temporal-Spatial Resolution Vehicle Emission Inventory Based on NRT Traffic Data and Its Impact on Air Pollution in Beijing, Part A: Development and evaluation of vehicle emission inventory, Atmos. Chem. Phys. Discuss., 15, 26711-26744, doi:10.5194/acp-15-26711-2015, 2015.

Kioutsoukis, I., de Meij, A., Jakobs, H., Katragkou, E., Vinuesa, J., and Kazantzidis, A.: High resolution WRF ensemble forecasting for irrigation: Multi-variable evaluation, Atmos. Res., 167, 156-174, doi:10.1016/j.atmosres.2015.07.015, 2016.

Li, M., Zhang, Z. Y., Liu, S. J., Yu, X. J., and Ju, C. X.: Verification of CUACE air quality forecast in Urumqi, Desert and Oasis Meteorol., 8, 63-68, 2014.

Liu, Z. R., Hu, B., Liu, Q., Sun, Y., and Wang, Y. S.: Source apportionment of urban fine particle number concentration during summertime in Beijing, Atmos. Environ., 96, 359-369, doi:10.1016/j.atmosenv.2014.06.055, 2014.

McKeen, S. A., Wotawa, G., Parrish, D. D., Holloway, J. S., Buhr, M. P., Hubler, G., Fehsenfeld, F. C., and Meagher, J. F.: Ozone production from Canadian wildfires during June and July of 1995, J. Geophys. Res., 107, 4192, doi:10.1029/2001JD000697, 2002.

Miao, S. G., Chen, F., Lemone, M. A., Tewari, M., Li, Q. C., and Wang, Y. C.: An observational and modeling study of characteristics of urban heat island and boundary layer structures in Beijing, J. Appl. Meteorol. Clim., 48, 484-501, doi:10.1175/2008JAMC1909.1, 2008.
Papalexiou, S. and Moussiopoulos, N.: Wind flow and photochemical air pollution in Thessaloniki, Greece. Part II: Statistical evaluation of European Zooming Model’s simulation results, Environ. Modell. Softw., 21, 1752-1758, doi:10.1016/j.envsoft.2005.09.004, 2006.

Qin, Y. and Chan, L. Y.: Traffic source emission and street level air pollution in urban areas of Guangzhou, South China (P.R.C.), Atmos. Environ., 27B, 275-282, 1993.

Roustan, Y., Pausader, M., and Seigneur, C.: Estimating the effect of on-road vehicle controls on future air quality in Paris, France. Atmos. Environ., 45, 6828-6836, doi:10.1016/j.atmosenv.2010.10.010, 2011.

Saikawa, E., Kurokawa, J., Takigawa, M., Borken-Kleefeld, J., Mauzerall, D. L., Horowitz, L. W., and Ohara, T.: The impact of China’s vehicle emissions on regional air quality in 2000 and 2020: a scenario analysis, Atmos. Chem. Phys., 11, 9465–9484, doi:10.5194/acp-11-9465-2011, 2011.

Stockwell, W. R., Middleton, P., Chang, J. S., and Tang, X.: The second generation regional acid deposition model chemical mechanism for regional quality modeling, J. Geophys. Res., 95, 16343-16376, 1990.

Song, X. Y. and Xie, S. D.: Development of vehicle emission inventory in China, Environ. Sci., 27, 1041-1045, 2006.

Song, Y., Xie, S. D., Zhang, Y. H., Zeng, L. M., Salmon, L. G., and Zheng, M.: Source apportionment of PM2.5 in Beijing using principal component analysis/absolute principal component scores and UNMIX, Sci. Total. Environ., 372, 278-286, doi:10.1016/j.scitotenv.2006.08.041, 2006.

Streets, D. G. and Waldhoff, S. T.: Present and future emissions of air pollutants in China: SO2, NOx, and CO, Atmos. Environ., 34, 363-374, doi:10.1016/S1352-2310(99)00167-3, 2000.

Wang, H. L., Zhuang, Y. H., Wang, Y., Sun, Y., Yuan, H., Zhuang, G. S., and Hao, Z. P.: Long-term monitoring and source apportionment of PM2.5/PM10 in Beijing, China, J. Environ. Sci., 20, 1323-1327, doi:10.1016/S1001-0742(08)62228-7, 2008.

Wang, H., Gong, S. L., Zhang, H. L., Chen, Y., Shen, X. S., Chen, D. H., Xue, J. S., Shen, Y. F., Wu, X. J., and Jin, Z. Y.: A new-generation sand and dust storm forecasting system GRAPES_CUACE/Dust: Model development, verification and numerical simulation, Chinese Sci. Bull., 55, 635-649, 2010.

Wang, H., Xue, M., Zhang, X. Y., Liu, H. L., Zhou, C. H., Tan, S. C., Che, H. Z., Chen, B., and Li, T.: Mesoscale modeling study of the interactions between aerosols and PBL meteorology during a haze episode in Jing-Jin-Ji (China) and its nearby surrounding region-Part 1: Aerosol distributions and meteorological features, Atmos. Chem. Phys., 15, 3257-3275, doi:10.5194/acp-15-3257-2015, 2015.

Wang, M., Zhu, T., Zheng, J., Zhang, R. Y., Zhang, S. Q., Xie, X. X., Han, Y. Q., and Li, Y.: Use of a mobile laboratory to evaluate changes in on-road air pollutants during the Beijing 2008 Summer Olympics, Atmos. Chem. Phys., 9, 8247–8263, doi:10.5194/acp-9-8247-2009, 2009.

Wang, T. and Xie, S.: Assessment of traffic-related air pollution in the urban streets before and during the 2008 Beijing Olympic Games traffic control period, Atmos. Environ., 43, 5682-5690, doi:10.1016/j.atmosenv.2009.07.034, 2009.

Wang, X., Westerdahl, D., Chen, L. C., Wu, Y., Hao, J. M., Pan, X. C., Guo, X. B., and Zhang, K. M.: Evaluating the air quality impacts of 2008 Beijing Olympic Games: On-road emission factors and black carbon profiles, Atmos. Environ., 43, 4535-4543, doi:10.1016/j.atmosenv.2009.06.054,
Wu, Q., Wang, Z. F., Gbaguidi, A., Gao, C., Li, L. N., and Wang, W.: A numerical study of contributions to air pollution in Beijing during CAREBeijing-2006, Atmos. Chem. Phys., 11, 5997–6011, doi:10.5194/acp-11-5997-2011, 2011.

Wu, Q., Xu, W., Shi, A., Li, Y., Zhao, X., Wang, Z., Li, J., and Wang, L.: Air quality forecast of PM10 in Beijing with Community Multi-scale Air Quality Modeling (CMAQ) system: emission and improvement, Geosci. Model. Dev., 7, 2243-2259, doi:10.5194/gmd-7-2243-2014, 2014.

Wu, S. W., Deng, F. R., Wei, H. Y., Huang, J., Wang, X., Hao, Y., Zheng, C. J., Qin, Y., Lv, H. B., Shima, M., and Guo, X. B.: Association of cardiopulmonary health effects with source-appointed ambient fine particulate in Beijing, China: a combined analysis from the healthy volunteer natural relocation (HVNR) study, Environ. Sci. Technol., 48, 3438-3448, doi:10.1021/es404778w, 2014.

Xiao, D., Deng, L. T., Chen, J., and Hu, J. K.: Tentative verification and comparison of WRF forecasts driven by data from T213 and T639 models, Torrent. Rain and Disast., 29, 20-29, 2010.

Yao, Q., Cai, Z. Y., Han, S. Q., Liu, A. X., and Liu, J. L.: Effects of relative humidity on the aerosol size distribution and visibility in the winter in Tianjin, China Environ. Sci., 34, 596-603, 2014.

Yu, L. D., Wang, G. F., Zhang, R. J., Zhang, L. M., Song, Y., Wu, B. B., Li, X. F., An, K., and Chu, J. H.: Characterization and source apportionment of PM2.5 in an urban environment in Beijing, Aerosol Air Qual. Res., 13, 574-583, doi:10.4209/aaqr.2012.07.0192, 2013.

Zhang, J. P., Zhu, T., Zhang, Q. H., Li, C. C., Shu, H. L., Ying, Y., Dai, Z. P., Wang, X., Liu, X. Y., Liang, A. M., Shen, H. X., and Yi, B. Q.: The impact of circulation patterns on regional transport pathways and air quality over Beijing and its surrounding, Atmos. Chem. Phys., 12, 5031–5053, doi:10.5194/acp-12-5031-2012, 2012.

Zhang, L., Liu, L.C., Zhao, Y.H., Gong, S.L., Zhang, X.Y., Henze, D., Capps, S., Fu, T., Zhang, Q., Wang, Y.X.: Source attribution of particulate matter pollution over North China with the adjoint method. Environ. Res. Lett., 10, 084011, doi:10.1088/1748-9326/10/8/084011, 2015.

Zhang, M. G., Pu, Y., Zhang, R., and Han, Z.: Simulation of sulfur transport and transformation in East Asia with a comprehensive chemical transport model, Environ. Modell. Softw., 21, 812-820, 2006.

Zhang, Q., Streets, D. G., Carmichael, G. R., He, K. B., Huo, H., Kannari, A., Klimont, Z., Park, I. S., Reddy, S., Fu, J. S., Chen, D., Duan, L., Lei, Y., Wang, L. T., and Yao, Z. L.: Asian emissions in 2006 for the NASA INTEX-B mission, Atmos. Chem. Phys., 9, 5131–5153, doi:10.5194/acp-9-5131-2009, 2009.

Zhang, R. J., Shen, Z. X., Zhang, L. M., Zhang, M. G., Wang, X., and Zhang, K.: Element composition of particles during periods with and without traffic restriction in Beijing: the effectiveness of traffic restriction measure, Sient. Onl. Lett. Atmos., 7, 61-64, doi:10.2151/sola.2011-016, 2011.

Zhang, R., Jing, J., Tao, J., Hsu, S.-C., Wang, G., Cao, J., Lee, C. S. L., Zhu, L., Chen, Z., Zhao, Y., and Shen, Z.: Chemical characterization and source apportionment of PM2.5 in Beijing: seasonal perspective, Atmos. Chem. Phys., 13, 7053-7074, doi:10.5194/acp-13-7053-2013, 2013.

Zhao, B., Wang, P., Ma, J. Z., Zhu, S., Pozzer, A., and Li, W.: A high-resolution emission inventory of primary pollutants for the Huabei region, China, Atmos. Chem. Phys., 12, 481-501, doi:10.5194/acp-12-481-2012, 2012.
Seasonal trends in PM2.5 source contributions in Beijing, China, Atmos. Environ., 39, 3967–3976, doi:10.1016/j.atmosenv.2005.03.036, 2005.

Zhou, Y., Fu, L. X., Yang, W. S., and Wang, Y.: Analysis of vehicle emission in Beijing by remote sensing monitoring, Tech. Equip. Environ. Poll. Contr., 6, 91-94, 2005.
| Numerical simulation | Emission source                                      |
|----------------------|------------------------------------------------------|
| SIM1                 | Default emission of CUACE                           |
| SIM2                 | Improved emission with Beijing HTSVE                |
| SIM3                 | Switch off Beijing vehicle emission                 |
| SIM4                 | Switch off Beijing anthropogenic emission           |
Table 2. Emission of major anthropogenic species in Beijing (unit: $10^3$ t yr$^{-1}$).

| Source                      | CO  | NOx | SO$_2$ | PM$_{2.5}$ |
|-----------------------------|-----|-----|--------|------------|
| CUACE emission              | 3149.5 | 173.8 | 158.2 | 79.0 |
| CUACE emission*             | 3119.3 | 183.2 | 158.2 | 78.8 |
| An et al. (2007)            | 1021.8 | 227.0 | 211.3 | 53.4 |
| Zhang et al. (2009)         | 2591.0 | 327.0 | 248.0 | 90.0 |
| Cao et al. (2011)           | 1998.0 | 437.0 | 172.0 | 162.0 |
| Wu et al. (2011)            | 236.2 | 172.5 |       | 67.9 |
| Zhao et al. (2012)          | 2580.0 | 309.0 | 187.0 | 90.0 |
| Q.Z. Wu et al. (2014)       | 1793.8 | 200.0 | 78.8  | 59.1 |

* represents CUACE emission with replaced vehicle emission by HTSVE.
Table 3. The rate of major species from vehicle emission in total emission (unit: %).

|       | CO  | NO  | NO₂ | HC  | PM₂.₅ |
|-------|-----|-----|-----|-----|-------|
| CUACE<sup>a</sup> | 29.8 | 32.1 | 30.4 | 80.0 | 23.4  |
| CUACE<sup>b</sup> | 31.1 | 35.5 | 33.6 | 49.0 | 25.3  |
| HTSVE<sup>a</sup> | 23.8 | 47.9 | 55.1 | 84.0 | 22.3  |
| HTSVE<sup>b</sup> | 21.3 | 46.6 | 53.9 | 55.8 | 20.6  |

<sup>a</sup> and <sup>b</sup> represent July and December.
Table 4. The contributions of traffic emission on ambient pollutant concentrations in Beijing.

| Source                        | Period   | Contribution (%) | Method                                      |
|-------------------------------|----------|------------------|---------------------------------------------|
| Hao et al. (2001)             | 1995     | NOx: 68.4; CO: 76.5 | Numerical simulation based on ISCST3        |
| Hao et al. (2005)             | 1999     | NOx: 74; PM$_{10}$: 14 | Numerical simulation based on ISCST3        |
| Zheng et al. (2005)           | 2000     | PM$_{2.5}$: 6.7   | Chemical mass balance receptor model (CMB)  |
| Song et al. (2006)            | 2000     | PM$_{2.5}$: 6.0–10.8 | PCA/APCS and UNMIX                         |
| Cheng et al. (2007)           | 2002     | PM$_{10}$: 28.7–42.9 | MM5-APRS-CMAQ                              |
| Wang et al. (2008)            | 2001-2006| PM$_{2.5}$: 5.9; PM$_{10}$: 8.4 | Positive matrix factorization (PMF)         |
| Zhang et al. (2009-2010)      | 2009-2010| PM$_{2.5}$: 4     | PMF                                        |
| Yu et al. (2013)              | 2010     | PM$_{2.5}$: 17.1  | PMF                                        |
| S. W. Wu et al. (2010-2011)   | 2010-2011| PM$_{2.5}$: 12.0  | PMF and mixed-effects models                |
| Cheng et al. (2011)           | 2011     | PM$_{2.5}$: 22.5±3.5 | MM5-CMAQ and source apportionment methods   |
| Liu et al. (2014)             | 2011     | PM(NC): 47.9      | PMF                                        |
| Huang et al. (2014)           | 201301   | PM$_{2.5}$: 5.6   | CMB and PMF                                |
Table 5. The VEC of chemical components in PM$_{2.5}$ in Beijing urban region (Unit:%).

|     | BC  | OC  | Ni(NO$_3^-$) | SF(SO$_4^{2-}$) | AM(NH$_4^+$) |
|-----|-----|-----|--------------|-----------------|--------------|
| Jul.| 12.3| 12.4| 13.4         | 1.8             | 2.1          |
| Dec.| 24.3| 25.8| 15.1         | 7.6             | 4.3          |
Figure 1. The model simulation domain (a) and observation station distribution (circle represents meteorological station, triangle represents environmental station) in inner domain (b).
Figure 2. The comparison of site average NO$_2$ and PM$_{2.5}$ concentrations between SIM1, SIM2 and observation in July (a, b) and December (c, d) 2013.
Figure 3. The spatial distribution of near-surface NO$_2$ and PM$_{2.5}$ mean concentration from SIM2 in July (a, b) and December (c, d) 2013 respectively. Black lines represent the main traffic arteries in Beijing, scatter represents the mean concentrations of sites observation, white arrows represent near-surface mean wind field.
Figure 4. Annual mean emissions and the rate of vehicle emission in total emission for NO$_2$ (a, c) and PM$_{2.5}$ (b, d) respectively. Black lines represent the main traffic arteries in Beijing.
Figure 5. The probability density function (PDF) of instantaneous VEC for NO$_2$ (a) and PM$_{2.5}$ (b).
Figure 6. Wind dependency map of VEC to NO$_2$ and PM$_{2.5}$ in July (a, b) and December (c, d) 2013. Wind speeds are shown from 0 m s$^{-1}$ to 7.5 m s$^{-1}$. 
**Figure 7.** Time series of daily mean and standard deviation of vehicle emission contribution rate on NO$_2$ and PM$_{2.5}$ concentrations of Beijing main urban areas in July (a, b) and December (c, d) 2013.
Figure 8. The scatter of daily mean concentration vs VEC for NO₂ and PM₂.₅ in July (a, b) and December (c, d).
Figure 9. The spatial distribution of mean contribution rate of vehicle emission on NO$_2$ and PM$_{2.5}$ in July (a, b) and December (c, d) 2013. Black lines represent the main traffic arteries in Beijing, white arrows represent near-surface mean wind field.
Figure 10. The spatial distribution of vehicle emission contribution in local emission to NO$_2$ and PM$_{2.5}$ in July (a, b) and December (c, d) 2013. Black lines represent the main traffic arteries in Beijing, white arrows represent near-surface mean wind field.