Using a mobile laboratory to characterize the distribution and transport of sulfur dioxide in and around Beijing

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Abstract

Megacities are places with intensive human activity and energy consumption. To reduce air pollution, many megacities have relocated energy supplies and polluted industries to their outer regions. However, regional transport then becomes an important source of air pollution in megacities. To improve air quality before and during the 2008 Beijing Olympics, a wide range of control strategies were implemented, including the relocation of polluting industries. High sulfur dioxide (SO$_2$) concentrations were occasionally observed during this period. Potential sources from southern regions of Beijing were indicated by backward trajectories and urban/rural stationary measurements, but direct evidence was lacking. Here we used a mobile laboratory to characterize the spatial distribution and regional transport of SO$_2$ to Beijing during the Campaign for Air Quality Research in Beijing and the Surrounding Region (CAREBEIJING)-2008. Among the five days chosen for the case studies during the Olympic air pollution control period, four had high SO$_2$ concentrations (6, 20 August and 3, 4 September 2008) while one had low SO$_2$ concentration (11 September 2008). The average values of SO$_2$ during the low SO$_2$ concentration day were 3.9 ppb, much lower than during the high concentration days (7.8 ppb). This result implied an impact by regional transport from outside Beijing. During these days, we captured transport events of SO$_2$ from areas south of Beijing, with a clear decrease in SO$_2$ concentrations southeast of the 6th to 4th Ring Roads around Beijing and along the 140 km highway from Tianjin to Beijing. The influx of SO$_2$ through the 4th to 6th Ring Roads ranged from 2.07 to 4.64 kg s$^{-1}$ on 4 September and 0.21 to 1.56 kg s$^{-1}$ on 20 August 2008. Locally emitted SO$_2$ from a source located along Jingshi Highway outside the southwest section of the 5th Ring Road of Beijing was identified using wind field data generated by the Weather Research and Forecasting model and the measured particle size distribution, with an estimated flux of 0.11 kg s$^{-1}$ to Beijing.
1 Introduction

Sulfur dioxide (SO\textsubscript{2}) is one of the most important precursors of secondary aerosols in the atmosphere. It is responsible for severe air pollution, leading to degraded visibility, changes in the radiation budget, and acid rain (Wang et al., 2008; Zhang et al., 2004a; Ramanathan and Crutzen, 2003). In addition, SO\textsubscript{2} is harmful to human health (Brunekreef and Holgate, 2002) and may cause increased respiratory diseases, reduced pulmonary function, low birth weight, and mortality (Xu et al., 1998).

Rapid economic development, industrialization, and urbanization have occurred within and around the megalcity of Beijing, increasing demands for fossil fuel consumption (Ohara et al., 2007), which is the main source of SO\textsubscript{2}. In 2008, the annual average energy consumption levels in Beijing, Tianjin, and the surrounding provinces of Heibei, Shanxi, and Shandong were 63.4 (Beijing Statistical Yearbook, 2009), 53.6 (Tianjin Statistical Yearbook, 2009), 242.2 (Hebei Statistical Yearbook, 2009), 268.8 (Shanxi Statistical Yearbook, 2009), and 125.1 (Shandong Statistical Yearbook, 2009) million tons of standard coal, respectively. To improve air quality and maintain clean air throughout the 2008 Beijing Olympics Games, Beijing municipal government implemented comprehensive long- and short-term air pollution control measures. The measures included moving heavy polluters out of the city, using low sulfur coal and high standard fuel, reducing the number of on-road vehicles, and freezing construction activities before and during the Olympics Games. While significant decreases in SO\textsubscript{2} were reported during the Olympics (Wang et al., 2009a, b; Qin et al., 2009), periods with relatively high SO\textsubscript{2} concentration occasionally occurred during the Olympic period, suggesting an important role of the regional transport of SO\textsubscript{2} emitted outside Beijing.

Previous studies have reported that SO\textsubscript{2} in Beijing is contributed by both local emission and regional transport (Zhang et al., 2004b; Xu et al., 2004; Sun et al., 2004). Rural/urban stationary (Liu et al., 2008; Guo et al., 2010) and tower observations (Sun et al., 2009) in Beijing have revealed that high wind speeds from southern areas might play a vital role in the increase of SO\textsubscript{2} concentrations in Beijing. An et al. (2007) used
the Community Multiscale Air Quality (CMAQ) model to simulate the regional transport of \( \text{SO}_2 \) and its flux pathway during a heavy pollution episode. They estimated that the southeast and southern areas of Beijing contributed 26\% and 18\% of \( \text{SO}_2 \) to the city. However, due to uncertainties in the \( \text{SO}_2 \) emission inventory outside Beijing, it is difficult to model the dispersion and transport of \( \text{SO}_2 \) accurately on small scales (Matsui et al., 2009). Thus, direct evidence from spatial distribution measurements is required.

Aircraft-based measurement with fast response instruments is a suitable approach to both studying the spatial distribution and quantifying the regional transport flux of \( \text{SO}_2 \) (Wang et al., 2006; Matvev et al., 2002; Beryrich et al., 1998). In northern China (where Beijing is located), research on \( \text{SO}_2 \) transport using aircraft measurements has focused on large-scale processes (Li et al., 2010; Ma et al., 2010; Ding et al., 2009) in the upper boundary layer or in the free troposphere. Given the high cost of aircraft measurement and difficulties in obtaining navigation approval, on-road measurement from mobile laboratories is an optimal method for ground-based spatial distribution measurement. On-road measurement platforms have been specifically designed and used for two types of applications: examination of the temporal-spatial variations in air pollutants (Bukowiecki et al., 2002; Weijers et al., 2004) and quantification of local emissions and regional transport flux (Johansson et al., 2009; Rivera et al., 2009). Johansson et al. (2008) and Li et al. (2009) calculated \( \text{SO}_2 \) fluxes from local emissions in Beijing using mobile-based mini-differential optical absorption spectroscopy (DOAS) around the 5th Ring Road of Beijing. However, these calculations were only based on column concentrations of \( \text{SO}_2 \). There is hence a lack of information on the ground level transport and concentrations of other pollutants.

Here we used an on-road mobile laboratory to measure the spatial distribution of \( \text{SO}_2 \) concentrations and identify transport processes in the southern part of Beijing. These observations were part of the Campaign for Air Quality Research in Beijing and the Surrounding Region-2008 (CAREBEIJING)-2008. The flux of \( \text{SO}_2 \) from both local emission and regional transport was estimated using wind field data from the WRF model version 3.1.1.
2 Methodology

2.1 Mobile laboratory and driving routes

Details of the setup and performance of the instruments installed in the mobile laboratory were described in our previous paper (Wang et al., 2009a). Briefly, a diesel vehicle (IVECO Turin V) was selected as the mobile platform. To minimize the loss of particles in the sampling inlet, an isokinetics inlet system was designed to enhance the sampling efficiency. The instruments onboard provided data on the concentrations of gaseous pollutants \((\text{NO}-\text{NO}_2-\text{NO}_x, \text{SO}_2, \text{CO}, \text{CO}_2, \text{O}_3)\) (ECOTECH, Australia), black carbon (MAAP, THERMO, USA), particle surface area (Nanoparticle surface area monitor, TSI, USA), particle size distribution (SMPS, DMA 3080 and CPC 3550, TSI, USA), and volatile organic compounds (VOCs; PTR-MS, IONICON, Austria). In this paper only the data of \(\text{SO}_2\) and SMPS were used (Table 2). The time resolutions of \(\text{SO}_2\) and SMPS data were 10 s and 2 min, respectively. Additional instruments included a Global Positioning System (GPS) and meteorological parameters (temperature, humidity, and pressure). The driving speed was \(60 \pm 5 \text{ km h}^{-1}\).

To characterize the spatial distribution of \(\text{SO}_2\) and to investigate the regional transport progress of \(\text{SO}_2\) to Beijing, routes were specially designed at local scale around the south area of Beijing, and at regional scale between Beijing and Tianjin megacities (Fig. 1). The details of routes and monitoring information are listed in Table 1. In general, five days were chosen for the measurements, with four days (6, 20 August and 3, 4 September 2008) showing high \(\text{SO}_2\) concentrations and one day showing low \(\text{SO}_2\) concentration (11 September 2008). We first selected routes 1 and 2 along the southeast of the Ring Roads to map the \(\text{SO}_2\) spatial distribution within the city of Beijing. The Ring Roads covers a wide scope from the urban area to the city center and thus is preferable to investigate the transport of \(\text{SO}_2\) at local scale. Because of the limited battery power of the instruments installed on the platform, observations in the mornings and afternoons were conducted on different days. To illustrate the spatial distribution and transport of \(\text{SO}_2\) between two megacities (Beijing and Tianjin), we
selected a newly built highway (Jingjintang II) between the two cities. The highway was far from industrialized areas, and because it was newly built, it had few cars driving on it during the Olympics. Thus the anthropogenic contribution from vehicular and industrial emissions was low, and the air pollutant levels were comparable to those of background air (Wang et al., 2009a). This unique feature was favorable for studying the air mass transport.

Continuous measurement was also conducted along the southwest of the 5th Ring Road and part of the 6th Ring Road in the southern area, where high-density industry is located, as shown in Fig. 1b (blue line). The characteristics of the SO\textsubscript{2} distribution and its local and regional emission sources surrounding the southwest of Beijing were reported previously (Wang et al., 2009a). Here we added wind field and particle size distribution data for a better understanding the distribution and emission sources of SO\textsubscript{2}.

### 2.2 Ground-based meteorological and SO\textsubscript{2} measurements

Concentrations of SO\textsubscript{2} were simultaneously measured at three intensive monitoring stations before, during, and after the Olympics air pollution control period. The PKU station was in an urban station located at Peking University, Beijing (39.99° N, 116.31° E); YuFa (YF, 39.51° N, 116.31° E) and YongLeDian (YLD, 39.75° N, 116.73° E) were rural stations representing the regional background. They were located approximately 50 km to the south and southeast of Beijing. In addition, 11 meteorological stations (blue stars in Fig. 1) of the Chinese Academy of Meteorological Sciences (CAMS) were selected for meteorological data analysis. Table 2 shows measurement parameters and instruments at each station.

To ensure that data between different stations were comparable, the SO\textsubscript{2} analyzers (Ecotech, 9850B, Australia) in all the stations were automatically calibrated between 00:00 and 01:00 every day using the same certified calibration standard (50 ppb, accuracy 3 %, diluted with N\textsubscript{2}, Beijing Huayuan Gas Chemical Industry Co., Ltd.). Calibration at five different concentrations (0 %, 20 %, 40 %, and 80 % of the detection
range) was performed each time. Meanwhile, calibrations of the SO\textsubscript{2} analyzer on the mobile laboratory were conducted before and after each sampling trip using similar procedures as at the stations. Intercomparison of SO\textsubscript{2} concentrations between the mobile laboratory and PKU station was also performed. The difference between the SO\textsubscript{2} concentrations was within 15\%, with a correlation coefficient of 0.82 (Wang et al., 2009a).

2.3 Wind Field

2.3.1 Lagrangian trajectory simulation

The Hybrid Single Particle Lagrangian Integrated Trajectory model (HYSPLIT, version 4.9), developed by US National Oceanic and Atmospheric Administration (NOAA) Air Resources Laboratory (ARL), was used to calculate the forward and backward trajectories from or to the stations in Beijing (Draxler and Rolph, 2003). Backward trajectories were computed once every 6 h for 24 h (1 day) at a selected height of 10 m above ground level for 45 days from 1 August to 15 September, 2008 and were grouped by cluster analysis of the model. A Global Data Assimilation System (GDAS) archive meteorological database was chosen to run the trajectory model, which had 1°×1° horizontal resolution. The horizontal resolution of the model is 1°×1°, which is enough to distinguish the original regions of the air masses in our study.

2.3.2 Weather Research and Forecasting (WRF) simulation

Direct measurements of wind speed and wind direction from the mobile lab during driving may lead to large deviations in the estimation of pollutant fluxes (Johansson, 2009). Here we used the WRF model version 3.1.1 to conduct mesoscale meteorological simulations for high-resolution wind fields and planetary boundary layer height (PBL) during the measurement periods. The WRF modeling system is a next-generation mesoscale numerical weather forecast and simulation system that includes...
the Advanced Research dynamics solver. A detailed description of the WRF model can be found on the WRF web-site http://www.wrf-model.org/index.php, as well as in our Supplement.

Here the horizontal resolution of the model was 1 km × 1 km. Four domains were used for the WRF calculation (Fig. S1). The averaged concentrations of SO\(_2\) were calculated based on each grid unit (1×1 km in Figs. 5 and 7, 2×2 km in Fig. 6). The bottom level reaching 18–20 m above the surface was selected for meteorological analysis. WRF model can simulated backward trajectories with high spatial, however, to identify the main original locations of the air masses during the measurement period and classified them into regions, it is not necessary to use WRF model for such a high resolution tracks which may take much longer time than the Hysplit model.

Comparing the WRF modeling results with the observations from meteorological stations involved in each domain during the measurement periods, we found close agreement in the wind directions, with \( R = 0.83 \). The wind speed had a relatively low correlation coefficient \( (R = 0.66) \) (Fig. S2). This could be contributed to the complicated land use in Beijing in which the local turbulences were not only influence the accuracy of the measured data in the meteorological stations (near the ground level where high buildings surround) but also challenge the predictive power of the model. However, given that most of the mobile routes were conducted in rural area, then we used just the rural sites to calculate the correlation between model and measurements (Fig. S2), and found the correlation coefficient increased significantly both for the wind speed \( (R = 0.83) \) and wind direction \( (R = 0.89) \).

The WRF model applied the newly updated land use initialized from MODIS and contains high densities of the wind fields in the vertical level (10 layers in 1000 m) and horizontal level (1 by 1 km). These improvements of the model lead us to believe that the model is able to produce a reliable spatial distribution of the wind field.

Hourly averaged WRF predicted PBL were compared with the temporal variations of aerosol extinction coefficient retrieved from Lidar measurement (Figs. S3, S4). We selected three days (6, 20 August and 11 September, 2008) with clear sky well mixed
convective condition; the measurement during these days could provide sufficient vari-
ation distributions of extinction coefficients. Figures S3 and S4 show that the model calculated PBL was generally in agreement with with the top boundary of the extinction coefficients during the measurement period, even though difference remained, this could be the major source of the flux uncertainty.

2.4 Assessment of the regional influx of SO$_2$

Quantitative assessment of SO$_2$ flux has been well established by the design of specified routes encircling target sources (Wang et al., 2006; Rivera et al., 2009; Johansson et al., 2008) or crossing frontiers perpendicular to the horizontal wind direction (Beryrich et al., 1998; Matvev et al., 2002). In this study, we carefully designed sector-routes on 20 August (route 1) and 4 September (route 2), so that the mobile tracks could traverse the wind direction. To calculate the transport flux outside Beijing, a simple formula based on the above measurement approach with the wind vector towards Beijing was used:

\[
\text{Flux (kg s}^{-1}) = \sum_{i=1}^{n} C_i \cdot V_i \cdot \text{Sin} \theta_i \cdot H_i \cdot d_i \times 10^{-9} \text{ (kg µg}^{-1})
\]

where

- \( \text{Flux} \) – the total SO$_2$ flux across the ring road with \( n \) \( 1 \times 1 \) km$^2$ unit cells \( \text{ (kg s}^{-1}) \)
- \( C_i \) – the mean concentration of SO$_2$ \( \text{ (µg m}^{-3}) \) in the \( i \)-th grid \( (1 \times 1 \) km$^2$)\)
- \( \theta_i \) – the angle between wind direction and the driving route in the \( i \)-th grid \( (1 \times 1 \) km$^2$)\)
- \( V_i \) – wind speed \( \text{ (m s}^{-1}) \) in the \( i \)-th grid \( (1 \times 1 \) km$^2$) generated by WRF output.
- \( H_i \) – mixing layer height \( \text{ (m)} \) in the \( i \)-th grid \( (1 \times 1 \) km$^2$) generated by WRF output.
3 Results and discussion

3.1 SO₂ concentration time series

Figure 2 shows the time series of SO₂ concentration and the air pollution index (API) from the three stations during the air pollution control period. The API data were from the Beijing Municipal Environmental Protection Bureau (EPB) and can be downloaded at http://www.bjepb.gov.cn/. These data gave the air pollution levels of particulate matter (PM₁₀), SO₂, and nitrogen dioxide (NO₂). API levels under 50 and 100 meant that the air quality met National Ambient Air Quality Standard Levels I and II. An API over 100 implies potentially adverse effects on the population.

In general, SO₂ concentrations were low during the Olympic air pollution control period (8 to 24 August 2008) at all stations, with average values of 5.8 ppb in the city and 3.2 ppb in rural areas. The mean API varied around 50. However, there were three periods with high SO₂ concentration at all stations: 3∼10 August, 19∼30 August, and 2∼8 September (Fig. 2). During these periods SO₂ concentration varied dramatically, with diurnal peak levels higher than 15 ppb. These trends were in agreement with API variations, for which mean values in the three periods were 80, 75, and 66, respectively. The maximum API level was around 100. During the 2008 Beijing Olympics period, most of the SO₂ emission sources were strictly controlled within and around
Beijing (Wang et al., 2009a); therefore, we assumed that local emission within Beijing city was not a major source of the increase in SO$_2$. Hence regional transport of SO$_2$ under appropriate meteorological conditions, e.g., wind speed and wind direction, should have been the major cause.

To further investigate the possible source of the observed SO$_2$, back trajectories were used to examine the air masses arriving in Beijing during the measurement period. A total of 184 trajectories (4 per day) were generated from the center of Beijing and then classified into four groups by cluster analysis. This revealed four typical source regions. Figure 3 shows the three clusters that derived from regional transport outside Beijing. The first group of trajectories (blue) originated from the Bohai Sea in the southeast of China and then moved over the Tianjin and Tangshan areas. The second group (green) was southerly from inland China, passing over Henan and Hebei provinces. The third group (red) was from the northwest of Inner Mongolia and the northeast regions of China. The last group (not shown here) travelled locally around Beijing city. Among the 184 trajectories, clusters 1, 2, 3, and 4 accounted for 22%, 28%, 26%, and 24% of all the trajectories, respectively. The 24 hour trajectories of clusters 1 and 2 were short, due to the slow wind, and thus SO$_2$ might have accumulated in Beijing. In contrast, the long trajectories of cluster 3 indicated strong winds from the northern area, which brought clean air and were favorable to air dispersion. Figure 4 shows the mean SO$_2$ concentration and API value in Beijing for each trajectory cluster. As expected, high SO$_2$ concentrations and API values were observed on the same days as the trajectory clusters 1 and 2. The average SO$_2$ concentrations ranged from 5.7 to 7.8 ppb, while the API values varied from 64.9 to 75.2. Trajectory clusters 3 and 4 were associated with low SO$_2$ concentrations and API values, which ranged from 3.9 to 4.8 ppb and 39 to 45, respectively.

Apparently, regional transport played an important role in increasing SO$_2$ concentrations in Beijing. To further investigate the SO$_2$ transport into Beijing, we used three mobile laboratory measurements to provide spatial distributions of SO$_2$ and to quantify its regional influx to Beijing.
3.2 Case studies of SO₂ spatial distribution

3.2.1 Southeastern area: routes 1 and 2

Figure 5a and b overlap the SO₂ concentration measured by the mobile laboratory and the wind field generated with the WRF model on 20 August and 4 September. The prevailing winds were from the SE and SSE with speeds of $1.2 \pm 0.2 \text{ m s}^{-1}$ and $3.6 \pm 1.2 \text{ m s}^{-1}$, respectively. The average mixing boundary layer height was $1027 \pm 232 \text{ m}$ on 20 August and $1223 \pm 48 \text{ m}$ on 4 September (Fig. S4). On both days, SO₂ concentrations had apparent horizontal gradients, increasing from urban to rural areas (Fig. 7) and clearly reflecting the transport patterns of SO₂ from the source regions in south area. It is noted that the temporal variations of SO₂ emissions may be convoluted with the spatial variability due to the driving speed of the mobile. However, since the wind speeds in both the measuring days were not fast, assuming that the SO₂ emissions during measurement period were stable and the air masses were large enough, this influence on the SO₂ distributions was low.

Figure 5 shows that high SO₂ concentrations were observed for approximately 30 km along the 5th and 6th Ring Roads, implying large-scale air mass transport. The mean value of SO₂ on 4 September was $14.9 \pm 3.2 \text{ ppb}$, twice as high as the $6.8 \pm 2.1 \text{ ppb}$ observed on 20 August. The 48 h backward trajectory on 20 August started in the Bohai Sea and encircled the suburban and urban areas of Tianjin before reaching Beijing (Fig. S6). On 4 September (Fig. S6), however, the 48 h backward trajectory originated from the mid-eastern mainland of China, traveling northward across Hebei, Henan, and even Shandong provinces. These areas contain numerous thriving industrialized cities where anthropogenic pollutants, especially SO₂, are generated (Streets et al., 2007). Liu et al. (2010) reported the neighboring provinces around Beijing such as Shandong, Hebei and Shanxi province contribute a large amount of SO₂ emissions of China (details please see Sect. 3.4). Air masses flowing across these regions usually pick up heavy pollutants downwind, leading to high concentrations of SO₂.
Figure 5c and d show the distributions of SO\(_2\) influx through the Ring Roads based on (1) and the high-resolution wind field. The flux of SO\(_2\) on 4 September was four times higher than that on 20 August. The SO\(_2\) in Beijing on 4 September was obviously imported from the southern region, while the pathway on 20 August was possibly from the southeast.

### 3.2.2 Southeastern area: routes 3 and 4 between Beijing and Tianjin

Figure 6a shows consistent decreasing trends of SO\(_2\) concentration from Tianjin to Beijing on 3 September, under southeasterly prevailing winds, indicating the regional transport of SO\(_2\) from relatively distant sources. The average SO\(_2\) concentrations along route 3 were modestly high, about 23 ppb. The maximum SO\(_2\) concentration over 40 ppb was observed in the industrial centers of Tianjin, Tanggu, and Hangu (Fig. 7). In contrast, SO\(_2\) concentrations had a very different distribution on 11 September (Fig. 6b), when the northwesterly wind dominated. The SO\(_2\) concentrations were low, with an average concentration of around 5 ppb along route 4 (Fig. 7), while SO\(_2\) concentrations in Beijing and Tianjin were relatively high. This suggests that SO\(_2\) in the cities came mainly from local SO\(_2\) emission sources.

While the wind field on 3 September was similar to that on 20 August and 4 September, the mean SO\(_2\) concentrations on the southeast of the 5th ring road differed between the three days, e.g. 7.5 ppb, 13.2 ppb, and 16.2 ppb on 20 August, 3 September and 4 September, respectively. This difference of the SO\(_2\) concentrations can be explained by the reason that 20 August was still in the Olympic period, when air pollution was strictly controlled. SO\(_2\) emissions on August 20 were expected less than that on 3 and 4 September. This difference can also be found from the stationary measurements as well as API variations in Fig. 1, in which the SO\(_2\) concentrations and API values on 20 August were lower than that on 3 and 4 September.

Figure 6c and d show the distributions of the SO\(_2\) flux along route 3 from Beijing to Tianjin on 3 September and route 4 from Tianjin to Beijing on 11 September. Since wind directions in both days were mostly towards Beijing or Tianjin, we simplify the flux
formula by using the product of mean SO₂ concentration in each cell and the measured wind speed to represent the SO₂ flux through the individual grid unit (2 × 2 km²). The wind speeds on both days were similar: 2.38 ± 0.27 m s⁻¹ and 2.04 ± 0.15 m s⁻¹, respectively. The boundary layer was high on 3 September (1220 ± 150 m) and relatively low on 11 September (833 ± 267 m). The flux of SO₂ imported to Beijing on 3 September was 124.3 g s⁻¹ in 2 × 2 km² grid units, much higher than the flux to Tianjin on 11 September (24.2 g s⁻¹ in 2 × 2 km² grid units).

### 3.2.3 Southwestern area: route 5 surrounding Beijing

We previously reported high concentrations of SO₂ and other pollutants in the Shijingshan district and along Jingshi Highway on 6 August 2008 and suggested this pollution was possibly from both local emission and regional transport (Wang et al., 2009a), but we had no further evidence to support this suggestion. In the particle number size distributions (Fig. 8c), we found bimodality with a remarkably high peak of ultrafine particle number concentrations in the 14.9–100 nm range and a smaller peak in the range larger than 100 nm, along Jingshi Highway. A similar observation was found in the Shijingshan district, but with lower particle number concentration peaks. Local emissions sources such as traffic exhaust, industrial emission, and biomass burning were identified as the major contributors to the particle number concentrations in the small range, while the larger size particles were “aged”, having formed during regional transport. Thus, we believe that the increase in SO₂ recorded along Jingshi Highway (Fig. 8b) was caused by both local emissions and regional transport.

To further investigate the high SO₂ concentrations in the Shijingshan district, we used the high-resolution wind field from the WRF simulation. As shown in Fig. 8a, on 6 August the observation along Jingshi Highway was dominated by winds flowing toward the northeast area, with the Shijingshan district located precisely downwind. Pollutants dispersed along the downwind direction from Jingshi Highway, leading to a high peak in SO₂ concentration across the Shijingshan district (see Fig. 8b dashed boxes).
The first peak of SO$_2$ at Shijingshan section shows clearly an overlay of the local emission and regional transported SO$_2$. We used a simple spline interpolation to separate the SO$_2$ from local emission (shaded area of the first Shijianshan peak in Fig. 8b) and regional transport (underneath the red dots of the first Shijianshan peak in Fig. 8b). The SO$_2$ from local emission was estimated by extracting the regional background of SO$_2$ from the measured SO$_2$ concentration. With this, we estimated the local emission from Jingshi Highway to the Shijingshan district represented 16% of the measured SO$_2$ concentrations.

### 3.3 Estimate of SO$_2$ influx to Beijing

Table 3 lists the flux of SO$_2$ through the Ring Roads on 6 and 20 August and 4 September calculated with (1). The SO$_2$ flux on the 5th Ring Road on 6 August was determined as 1.64 kg s$^{-1}$, with 2.9 m s$^{-1}$ wind speed and 1140 ± 233 m boundary layer height. The flux of local emission on Jingshi Highway was assessed by calculating the downwind SO$_2$ traverse at the southwestern 5th Ring Road in the Shijingshan district, generating a SO$_2$ flux of 0.11 kg s$^{-1}$. The flux of SO$_2$ increased from the 4th to 6th Ring Roads, from 0.21 to 1.56 kg s$^{-1}$ on 20 August and 2.07 to 4.64 kg s$^{-1}$ on 4 September. The large difference between the estimated fluxes of SO$_2$ could be explained by the higher SO$_2$ concentrations and stronger winds on 4 September compared with those on 20 August.

Uncertainty in estimates of the SO$_2$ flux in this study may came from the uncertainties in both horizontal and vertical distributions of wind profiles, the height of boundary layers, and the time interval across the plume (Wang et al., 2008). The horizontal differences in wind speed and direction found by the WRF model were 19% and 7.5% with regard to the station monitoring data. Zhu et al. (2010) observed a homogeneous vertical distribution of SO$_2$ in daytime in Beijing with DOAS measurements, which suggested our assumption of homogeneous vertical distribution of SO$_2$ is justified. However, uncertainties in vertical wind profile and PBL were difficult to estimate because...
of the lack of measurement data. Given the large spatial variation of SO\textsubscript{2} emission in neighboring regions of Beijing (Sect. 3.4), we are confident that our flux calculations are reasonable with acceptable uncertainty.

### 3.4 Comparison of measured SO\textsubscript{2} fluxes with emission inventory

Table 4 shows a comparison of our results with published emission inventory from the provinces surrounding Beijing. Given a rapid increase of the emissions in China, we choose the most recent report of the emission inventory in 2007 (Liu et al., 2010), which was based on the INTEX-B inventory. The sum of the emissions from the power plant, industry, domestic and traffic sources in provinces surrounding Beijing were listed in Table 4.

As shown in Table 4, Beijing and Tianjin have the lowest emission of SO\textsubscript{2}, while Shandong province has the highest SO\textsubscript{2} emission, followed by Hebei and Shanxi provinces. The SO\textsubscript{2} emission from Shandon was tenfold higher than that in Beijing and could contributed to the SO\textsubscript{2} concentration in Beijing under south or southeast winds. The Hebei province is in proximity to Beijing with high SO\textsubscript{2} emission, and may contribute more to Beijing SO\textsubscript{2} concentrations, due to shorter distance compared to Shandon province. The emissions from Tianjin city was much lower than the emissions from the other provinces, which may partially account for the relatively low SO\textsubscript{2} concentrations on 20 August when southeastern wind dominated.

We extrapolate the SO\textsubscript{2} flux measured on 20 August and 4 September at the ring roads, to the annual SO\textsubscript{2} flux into Beijing. Table 4 shows that the annual influxes of SO\textsubscript{2} through the southeast of the 6th ring road to Beijing could be as high as 146.3 Gg yr\textsuperscript{-1} and 65.3 Gg yr\textsuperscript{-1} on the 4th ring road. This is about one third or equivalent to the SO\textsubscript{2} emission in Beijing, suggesting the transport of SO\textsubscript{2} is of pivotal importance to the SO\textsubscript{2} concentration in Beijing. Apparently, high uncertainties are associated with the SO\textsubscript{2} flux measurements and this extrapolation, but our results gives a relative importance of the SO\textsubscript{2} transport similar as previously reported model estimation of a heavy pollution episode (An et al., 2007).
4 Conclusions

We used a mobile laboratory to identify and assess the transport processes and emission sources of \( \text{SO}_2 \) surrounding Beijing in combination with the wind field distribution. The measurements were conducted as a part of the CAREBeijing-2008 campaign from August to September 2008 and covered the southeast and southwest areas outside Beijing.

Continuous measurements at the PKU urban station and the YuFa and YongLeDian rural stations were performed before, during, and after the comprehensive control period. Three potential polluted periods were identified, with short backwards trajectories from the south and southeast areas outside Beijing.

Mobile monitoring was conducted to investigate the regional transport of \( \text{SO}_2 \) from the southern mainland of China. We designed three routes. The route along the southeast of the Ring Roads reflected the wide scope of the polluted plumes, while the “straight line” between Beijing and Tianjin further supported the regional transport of \( \text{SO}_2 \) over long distances. We also noticed that the plumes from the center of mainland China usually picked up pollutants along the way, leading to high concentrations on 4 September compared to those on 20 August. On 6 August, regional transport from southeast areas as well as local emissions on Jingshi Highway and their dispersion over the downwind district of Shijingshan were identified and demonstrated by WRF simulations and particle size-distribution measurements.

Using a simple method assuming constant wind speed and directions within 1 h and homogeneous vertical profiles of the well-mixed boundary layer, we found that the flux of \( \text{SO}_2 \) transported to Beijing was high on 4 September and low on 20 August. From the 4th to the 6th Ring Roads, the \( \text{SO}_2 \) flux ranged from 2.07 to 4.64 kg s\(^{-1}\) on 4 September and 0.21 to 1.56 kg s\(^{-1}\) on 20 August. Influx from the southwest on 6 August across the 5th Ring Road was 1.64 kg s\(^{-1}\). Local emission on Jingshi Highway was roughly 0.11 kg s\(^{-1}\). In summary, mobile monitoring is a useful approach to evaluate temporal
and spatial variations in SO$_2$ transport processes. The uncertainty of the SO$_2$ flux estimation can be reduced by using concurrent measured boundary layer height.

Supplementary material related to this article is available online at: http://www.atmos-chem-phys-discuss.net/11/16465/2011/acpd-11-16465-2011-supplement.pdf.

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References


Wang, P., Richter, A., Bruns, M., Burrows, J. P., Scheele, R., Junkermann, W., Heue, K.-P.,


Table 1. Mobile measurement dates and route types.

<table>
<thead>
<tr>
<th>Cruise Route</th>
<th>Starting date time (LT)</th>
<th>Ending date time (LT)</th>
<th>Route type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Route 1</td>
<td>20 Aug 2008 (07:58)</td>
<td>20 Aug 2008 (12:50)</td>
<td>Southeast 2th to 6th Ring RD</td>
</tr>
<tr>
<td>Route 2</td>
<td>4 Sep 2008 (14:13)</td>
<td>4 Sep 2008 (18:22)</td>
<td>Southeast 4th to 6th Ring RD</td>
</tr>
<tr>
<td>Route 3</td>
<td>3 Sep 2008 (10:05)</td>
<td>3 Sep 2008 (15:13)</td>
<td>Beijing to Tianjin</td>
</tr>
<tr>
<td>Route 4</td>
<td>11 Sep 2008 (10:09)</td>
<td>11 Sep 2008 (14:37)</td>
<td>Beijing to Tianjin</td>
</tr>
<tr>
<td>Route 5</td>
<td>6 Aug 2008 (12:00)</td>
<td>6 Aug 2008 (14:50)</td>
<td>Southwest area outside Beijing</td>
</tr>
</tbody>
</table>
Table 2. Instrumentation performed in this study.

<table>
<thead>
<tr>
<th>Station name</th>
<th>Instruments</th>
<th>Measurement species</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mobile Laboratory</td>
<td>SO₂ analyzer (Ecotech 9850A, Australia) (10 s) SO₂ scanning mobility particle sizer (SMPS) (3080, TSI Inc.) (2 min)</td>
<td>SO₂</td>
</tr>
<tr>
<td>PKU station</td>
<td>SO₂ analyzer (Ecotech 9850B, Australia) (1 h)</td>
<td>Wind speed, wind direction</td>
</tr>
<tr>
<td></td>
<td>Mete One (1~3 h)</td>
<td></td>
</tr>
<tr>
<td>YuFa station</td>
<td>SO₂ analyzer (Ecotech 9850B, Australia) (1h)</td>
<td>Wind speed, wind direction</td>
</tr>
<tr>
<td></td>
<td>Mete One (1~3 h)</td>
<td></td>
</tr>
<tr>
<td>YongLeDian station</td>
<td>SO₂ analyzer (Ecotech 9850B, Australia) (1 h)</td>
<td>Wind speed, wind direction</td>
</tr>
<tr>
<td></td>
<td>Mete One (1~3 h)</td>
<td></td>
</tr>
<tr>
<td>CAMS stations</td>
<td>Mete One (1~3 h)</td>
<td>Wind speed, wind direction</td>
</tr>
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</table>
Table 3. SO$_2$ import fluxes and local emission derived from mobile laboratory in different routes surrounding Beijing.

<table>
<thead>
<tr>
<th>Date</th>
<th>4th Ring RD SE</th>
<th>5th Ring RD SE</th>
<th>6th Ring RD SE</th>
<th>5th Ring RD SW</th>
<th>Shijingshan</th>
</tr>
</thead>
<tbody>
<tr>
<td>6 Aug 2008</td>
<td>0.21</td>
<td>0.41</td>
<td>1.56</td>
<td></td>
<td>1.64</td>
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<tr>
<td>20 Aug 2008</td>
<td>2.07</td>
<td>4.02</td>
<td>4.64</td>
<td></td>
<td></td>
</tr>
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<td>4 Sep 2008</td>
<td></td>
<td></td>
<td></td>
<td></td>
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Table 4. Comparison of the 2007 emission flux of SO$_2$ in provinces surrounding Beijing with the estimated SO$_2$ annual flux into Beijing.

<table>
<thead>
<tr>
<th>Province</th>
<th>Ring roads</th>
<th>SO$_2$ emission (Gg yr$^{-1}$)</th>
<th>SO$_2$ flux (Gg yr$^{-1}$) 20 Aug/4 Sep</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beijing</td>
<td>194</td>
<td></td>
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<td>Liu et al. (2010)</td>
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<tr>
<td>Tianjin</td>
<td>323</td>
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<td>Liu et al. (2010)</td>
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<td>Hebei</td>
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<td>Liu et al. (2010)</td>
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<tr>
<td>Shandong</td>
<td>4373</td>
<td></td>
<td></td>
<td>Liu et al. (2010)</td>
</tr>
<tr>
<td>Shanxi</td>
<td>1738</td>
<td></td>
<td></td>
<td>Liu et al. (2010)</td>
</tr>
<tr>
<td>Beijing 4th Ring RD</td>
<td>6.6/65.3</td>
<td></td>
<td>This study</td>
<td></td>
</tr>
<tr>
<td>Beijing 5th Ring RD</td>
<td>12.9/126.7</td>
<td></td>
<td>This study</td>
<td></td>
</tr>
<tr>
<td>Beijing 6th Ring RD</td>
<td>49.2/146.3</td>
<td></td>
<td>This study</td>
<td></td>
</tr>
</tbody>
</table>
Fig. 1. Maps of the mobile monitoring areas in CAREBeijing-2008. The green track in (a) shows route 3 and 4 in Table 1. The red, yellow and blue tracks in (b) show the route 1, 2 and route 5, respectively. The red stars on the maps represent the stationary sites on SO$_2$ measurements and the blue stars show the CAMS meteorological stations.
Fig. 2. Temporal variations of SO$_2$ concentrations at PKU, YuFa and YongLeDian stations and API values reported by Beijing EPB from 1 August to 15 September 2008. The red arrows show the wind vectors at PKU station. The dashed line of level 1 and 2 indicates the maximum of national air quality criteria I (50) and II (100), respectively. Period I, II and III show the potential SO$_2$ pollution periods with increase trends between 3–10 August, 19–30 August and 2–8 September.
Fig. 3. Air mass backward trajectories at 10 m above ground level at Beijing from 1 August to 15 September 2008. The red, green and blue trajectories show the plumes from north, south and southeast regions.
Fig. 4. The average concentrations of SO$_2$ and API measured at PKU (Haidian for API), YuFa (YuFa) and YongLeDian (Tongzhou) sites subdivided on the basis of four directions of backtrajectories from 1 August to 15 September 2008. The bars show the standard deviations of SO$_2$ concentrations and API.
Fig. 5. The spatial distributions of SO\textsubscript{2} concentrations and wind field by WRF with 1 × 1 km grids resolution in southeastern areas of Beijing on (a) 20 August and (b) 4 September 2008, as well as the corresponding flux on 20 August (c) and 4 September 2008 (d).
Fig. 6. The spatial distributions of SO$_2$ concentrations from Beijing to Tianjin on (a) 3 September and (b) 11 September 2008; the corresponding flux on 3 September and 11 September 2008 are shown in (c) and (d).
Fig. 7. Boxplots of the \( \text{SO}_2 \) concentrations on the ring roads in southeast of Beijing during the measurement periods. The small block indicates the mean value and the upper, middle and bottom layers of the box show the 75, 50, 25-th percentiles of the dataset. The bars are determined by the 5 and 95th percentiles of the dataset and “×” show the maximum and minimum values, respectively.
Fig. 8. (a) SO$_2$ spatial distributions and wind field by WRF with $1 \times 1$ km$^2$ grids resolution in southwestern areas of Beijing on 6 August 2008. The dashed areas on the left and middle of the figure represent Jingshi Highway and Shijingshan district where high peak values were observed; (b) time series of SO$_2$ variations on 6 August, where the red plots and the black shades show the regional background and local emission of SO$_2$ across the 5th Ring Road. (c) SMPS particle size distributions measured in separated areas of the route on 6 August. The red and blue arrows display the Jingshi Highway and Shijingshan districts in (a) corresponding to the SO$_2$ peak values in (b) and the particle size distributions in (c).