# Response to Interactive comments from Anonymous Referee #1

Referee comments are in black. Author responses are in blue.

General Remarks: The manuscript present the results of observing NO2 emission measurements in Beijing based on the Car MAX-DOAS technology. Through 19 times of city-circle-around Car-MAX-DOAS experiments, the author showed the potential of Car MAX-DOAS measurement technology in atmospheric monitoring. This observation method can be effectively used for dynamic monitoring of urban NO2 emissions. However, the database the authors use for the conclusions is relatively weak. So some revisions are needed to consider this manuscript for publication in ACP.

We thank the anonymous referee for his/her insightful and constructive comments. Below are our point-to-point responses in detail.

# Major concerns:

1. Line 24-25, "typically larger NO2 VCD at the southern parts of the 6th Ring Road than at the northern parts". According to Figures 5 and 6, the NO2 VCD at the southern parts and northern parts were not typically different in January.

The sentence has been deleted in the revised version of the manuscript.

2. Since each measurement time was different, from 2 to 2.5 hours (sometimes nearly 3 hours), the author should introduce the traffic situation during the measurement and analyze the impact on the measurement results.

Traffic jam did not occur during the measurement and the impact of traffic situation should be negligible. Different people might drive at different speeds although we had suggested them to drive at a low speed and stable level. Only after the experiments, we realized that driving slowly might also cause a problem for the emission estimate since the change in wind field might be more pronounced with a longer experimental time.

3. The author should introduce the NOx emission sources in Figure 1 and analyze the influence on the measurement.

We have added the related descriptions in lines 198-199 and analyze the influence on the NO2 VCD measurements in lines 396-398 of the revised manuscript.

4. The results of Car Max- DOAS measurement show that NOx emission in heating season is nearly three times as much as that in non heating season, which is obviously higher than that calculated by MEIC inventory estimate. Since central heating is adopted in Beijing urban area, the author should analyze heating season NOx sources in detail, and evaluate the contributions to the measurement results.

Many thanks for suggestions. Central heating from power plant and home heating increase emissions in heating season compared to non-heating season. We calculated the average NO<sub>X</sub> emission rates of four sectors including industry, power, residential, and transportation from the MEIC within the 6th Ring Rd of Beijing in January,

September, and October 2012, and the ratio of  $NO_X$  emission rates in Jan. and the average in Sep. and Oct (Table S1). The  $E_{NOX}$  from power and residential in January are remarkably higher than other two months, especially  $E_{NOX}$  from residential in January are about 5 times those in other months. Corresponding descriptions are added in 371-372 and lines 437-445 of the revised version. We cannot retrieve the specific positions of heating season NOx sources from MAX-DOAS measurements by the method used in this study. But we agree that it is a significative scientific issue and will use the source apportionment model to investigate it in the future study.

# Minor comments:

1. Abstract, here it is more appropriate to use "different months" instead of "different seasons".

Corrected.

- 2. Line 43, "less than" instead of "smaller than" Corrected.
- 3. Line 54-56, "NO and NO2 (together denoted as NOX) form primarily in combustion processes, and the conversion between NO and NO2 in the atmosphere is very rapid" is well known and meaningless here

The sentence has been changed to "the studies on the spatiotemporal variation of NO and NO2 (together denoted as NOX), with the latter being a precursor of nitrate aerosols, are very important for understanding the aerosol formation and its influencing factors" in lines 59-61 of the revised manuscript.

4. Please unify the format of "Car-MAX-DOAS" in the manuscript. For example: line 90 "car-MAX-DOAS", line 94 "Car-MAX-DOAS", line 101 "car MAX-DOAS", line 190 "car MAX-DOAS" et al.

Done.

5. Line 147, "the roof of a car" instead of "the roof a car". Corrected.

# Retrieving tropospheric NO<sub>2</sub> vertical column densities around the city of Beijing and estimating NO<sub>X</sub> emissions based on carcar MAX-DOAS measurements

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Abstract. We carried out 19 city-circle-around ear-car MAX-DOAS experiments on the 6th Ring Road of Beijing in January, September, and October 2014. The tropospheric vertical column densities (VCDs) of NO<sub>2</sub> were retrieved from measured spectra by the Multi-Axis Differential Optical Absorption Spectroscopy (MAX-DOAS) technique and used to estimate the emissions of  $NO_x$  ( $\equiv NO + NO_2$ ) from urban Beijing during the experimental periods. The offline LAPS-WRF-CMAQ model system was used to simulate the wind fields by assimilation of observational data and calculate the NO2-to-NOx concentration ratios, both of which are also needed for the estimation of NO<sub>x</sub> emissions. The NO<sub>X</sub> emissions in urban Beijing for the different months seasons derived from the ear-Car MAX-DOAS measurements in this study were compared to the multi-resolution emission inventory in China for 2012 (MEIC 2012). Our ear-Car MAX-DOAS measurement results showed higher NO<sub>2</sub> VCD in January than in the other two months and typically larger NO2 VCD at the southern parts of the 6th Ring Road than at the northern parts. The wind field had obvious impacts on the spatial distribution of NO<sub>2</sub> VCD, with the mean NO2 VCD along the 6th Ring Road typically being higher under the south wind than under the north wind. In addition to the seasonal difference, the journey-to-journey variations of estimated NO<sub>X</sub> emissions rates (E<sub>NOX</sub>) were large even within the same month, mainly due to uncertainties in the calculations of wind speed, the ratio of NO<sub>2</sub> and NO<sub>X</sub> concentration, and the decay rate of NO<sub>X</sub> from the emission sources to the measured positions under different meteorological conditions. The ranges of  $E_{\rm NOX}$  during the heating and non-heating periods were  $22.59 \times 10^{25}$  to  $31.28 \times 10^{25}$  28.7 ×  $10^{25}$  to  $60.0 \times 10^{25}$  molecules s<sup>-1</sup> and  $9.61 \times 10^{25}$  to  $11.96 \times 10^{25}$  7.7 ×  $10^{25}$  to  $24.8 \times 10^{25}$  molecules s<sup>-1</sup>, respectively. The average  $E_{NOX}$  values in the heating and non-heating periods were  $26.94 \pm 6.14 \times 10^{25}$  molecules s<sup>-1</sup> and  $10.99 \pm 1.23 \times 10^{25}$  molecules s<sup>-1</sup>,  $43.0 \times 10^{25}$  molecules s<sup>-1</sup> and  $13.9 \times 10^{25}$  molecules s<sup>-1</sup>, respectively. The uncertainty range of  $E_{NOX}$  was 19.52 - 52.01% 16.4 - 33.2%. The monthly emission rates from MEIC 2012 are found to be lower than the estimated  $E_{NOX}$ , particularly in January. Our results provide important information and datasets for the validation of satellite products and also show how ear Car MAX-DOAS measurements can be used effectively for dynamic monitoring and updating of the NO<sub>x</sub> emissions from megacities such as Beijing.

#### 1. Introduction

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Over the past decade, serious haze events have weather has occurred frequently in autumn and winter in Beijing due to massive anthropogenic emissions from the combustion consumption of fossil fuels and other sources (He et al., 2013; Zhang et al., 2013). High concentrations of aerosol particulate matter with dynamic diameter less smaller than 2.5 µm (PM<sub>2.5</sub>) threaten public health (Cao et al., 2014), disturb traffic operation by affecting visibility, and result in perturbations changes to the weather and climate because of by scattering and absorption of solar radiation (Liao et al., 2015; Cheng et al., 2017). Measurements have shown that organic matter (OM), sulfate, nitrate, and ammonium made up more than 78% of the PM<sub>2.5</sub> in January 2013 in Beijing (Huang et al., 2014). Fractions of nitrate in PM<sub>2.5</sub> have obviously-increased recently with the control of industry and coal in the Beijing-Tianjin-Hebei region, which has reduced SO<sub>2</sub> emissions and the ratio of sulfate in PM<sub>2.5</sub>, while traffic emissions are still at high levels. A recent study based on the aerosol observations at the campus of Peking University in 2014 revealed that aerosol pollution is nitrate-driven in spring and early fall and OM-driven in late fall and winter (Tan et al., 2018). - Recent research (Tan et al., 2018) based on the aerosol observations at the campus of Peking University in 2014 revealed that aerosol pollution is nitrate driven in spring and early fall and OM driven in late fall and winter. The researchers-study suggested that nitrate formation was more significant than sulfate formation during severe pollution episodes in Beijing. Therefore, studies on the spatiotemporal variation of NO and NO<sub>2</sub> (together denoted as NO<sub>X</sub>), with the latter being a precursor of nitrate aerosols, are very important for understanding the aerosol formation and its influencing factors the spatiotemporal variation of gaseous precursors of nitrate are very important for understanding

the aerosol formation and its influencing factors. NO and NO<sub>2</sub> (together denoted as NO<sub>X</sub>) form primarily in combustion processes, and the conversion between NO and NO<sub>2</sub> in the atmosphere is very rapid.

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Emission inventories are usually developed by the so-called bottom-up approach, which is based on combinations of activity statistics (such as energy consumption and industrial production) and source- or region-specific emission factors (Hao et al., 2002; Zhang et al., 2007; Zhao et al., 2012; Streets et al., 2013). However, there are large high uncertainties in bottom-up emissions inventories associated with the statistics, emissions factors, temporal allocation profiles, and grid allocation factors (Ma and Van Aardenne, 2004; Zhao et al., 2012). Moreover, estimating "current" emissions by the bottom-up methodology is fundamentally difficult because publication of basic statistics is generally a couple of years behind. The "top-down" emission estimatetop down constraint is a useful supplement to bottomup estimates, which are subject to uncertainties in emissions factors and emissions activities (Streets et al., 2013). Inverse modeling, in which emissions are optimized to reduce the differences between simulated and observed data, is a powerful method that solves the problems of the bottom-up approach. Recently, its application to the estimation of NO<sub>X</sub> emissions has been widely reported. NO<sub>X</sub> emission rates are derived by constraining satellite observations using the relationship between model-simulated NO<sub>2</sub> vertical column density (VCD) and primary NO<sub>X</sub> emission estimates from the bottom-up approach (Martin, 2002; Jaegle' et al., 2005; Konovalov et al., 2006; Wang et al., 2007; Lin et al., 2012; Zyrichidou et al., 2015). Nevertheless, errors and uncertainties still exist in the retrieval of satellite data, and thesewhich leads to a large decrease in the accuracy of estimated emissions, precision, particularly in highly polluted regions such as Beijing and its surroundings (Ma et al., 2013a; Jin et al., 2016). Uncertainties can arise from noise, surface albedo, cloud blocks, profile shape, interference from ozone absorption, correlations with other retrieved parameters, fitting wavelength window, and so forth (Jin et al., 2016; Ma et al., 2013a; Shaiganfar et al., 2011, 2017). Air mass factor (AMF) errors can produce additional errors during the conversion process from the slant to vertical columns. Therefore, comprehensive ground-based measurements of the tropospheric columns and vertical profiles of NO2 are quite important and necessary to evaluate and validate satellite retrieval products.

The Multi-Axis Differential Optical Absorption Spectroscopy (MAX-DOAS)MAX DOAS (Multi-Axis Differential Optical Absorption Spectroscopy) is a new-ground-based remote sensing technique developed during the last two decades. It makes use of the scattered sunlight measured from horizontal through zenith pointing direction to retrieve the VCD and vertical profiles of trace gases and aerosols

with relatively high sensitivity in the lower atmosphere (Hönninger et al., 2004; Wagner et al., 2004; Platt and Stutz, 2008). MAX-DOAS has been extensively used to derive tropospheric column information of NO<sub>2</sub> and some other pollutants in various regions (Wittrock et al., 2004; Brinksma, et al., 2008; Irie et al., 2008; Vlemmix et al., 2010; Li et al., 2013; Hendrick et al., 2014). Mobile- (or earCar-) MAX-DOAS measurements have been used to quantify NO<sub>X</sub> emissions from cities and regions such as Beijing (Johansson et al., 2008), Mexico (Johansson et al., 2009), Mannheim and Ludwigshafen (Ibrahim et al., 2010), Deli (Shaiganfar et al., 2011), Shanghai (Wang et al., 2012), North China (Wu et al., 2018). Compared to ground-based observations at a fixed site, earCar-MAX-DOAS measurements can provide information on the horizontal spatial distribution of pollutants, which is important for explaining the urban/regional representativeness of satellite observations and validating the NO<sub>2</sub> VCDs and NO<sub>x</sub> emission estimates from the new, high pixel resolution measurements by the TROPOMI instrument on the Sentinel-5P over megacities such as Beijing. Moreover, due to the rapid expansion of urban area and increasing energy consumption, both locations and strength of emission sources in Beijing might may have changed significantly. Therefore, intensive Car-MAX-DOAS measurement campaigns are still needed to estimate the emissions of NO<sub>X</sub> in Beijing. Mean wind speed and wind direction along the ring road during the sampling periods were are usually used to estimate NO<sub>X</sub> emissions in the previous studies. Since wind field changes rapidly due to local circulation and then results in uncertainties in quantification of NO<sub>X</sub> emissions (Johansson et al., 2008; Shaiganfar et al., 2011, 2017; Davis, et al., 2019), refined and accurate simulations of wind fields are needed for the accurate emission estimate.

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In this study, we estimated the total NO<sub>X</sub> emissions from urban Beijing based on the VCD of NO<sub>2</sub> obtained from intensive ear-Car MAX-DOAS measurements on the 6th Ring Road of Beijing in January, September, and October of 2014. The offline LAPS-WRF-CMAQ model system with data assimilation method was used to derive wind speed, wind direction, and NO<sub>2</sub>/NO<sub>X</sub> concentration ratios, which are needed to estimate total urban NO<sub>X</sub> emissions based on ear-Car MAX-DOAS measurements. We attempted to accurately estimate the NO<sub>X</sub> emission rates and the seasonal difference and deeply investigate the uncertainties and appropriate meteorological conditions for the estimation based on Car MAX-DOAS measurements. This paper is organized as follows: Section 2 describes the intensive ear Car MAX-DOAS experiments and the retrieval method for deriving tropospheric NO<sub>2</sub> VCD, the model system used to simulate the wind fields and the ratios of NO<sub>2</sub> and NO<sub>X</sub>, and the method used to quantify total NO<sub>X</sub> emissions. Section 3 presents the results of the NO<sub>2</sub> VCD and the estimated NO<sub>X</sub> emissions as

well as their uncertainties due to simulated errors in the wind field. Conclusions are provided in Section 4.

# 125 2. Theory, experimental, and method

#### 2.1 Formula to estimate urban NOx emissions

The complete NO<sub>2</sub> flux  $F_{NO_2}$  across the urban Beijing area encircled by the driving route S is estimated according to the closed integral method (CIM) the method of Ibrahim et al. (2010).

$$F_{NO_2} = \oint_s VCD_{NO_2}(s) \cdot \vec{w} \cdot \vec{n} \cdot \vec{w} \cdot \vec{n} \cdot ds \tag{1}$$

- Here  $VCD_{NO_2}(s)$  is the NO<sub>2</sub> VCD at the sampling position within the driving route;  $\vec{n}$  indicates the normal vector parallel to the Earth's surface and orthogonal to the driving direction at the position of the driving route;  $\vec{w}$  is the average wind vector within the NO<sub>2</sub> layer, which is denoted by wind speed at the height of 10 m. We carried out <u>ear-Car</u> MAX-DOAS measurements along closed driving routes around large emissions sources, i.e., the 6th Ring Road of Beijing (Fig. 1).
- We averaged the wind vector data from the WRF model between surface and 1000m altitude weighted by the winter exponentially decreasing profiles according to the method of Shaiganfar et al. (2017).

$$\underline{\vec{w}} = \frac{\sum_{i} w(z_i) \cdot e^{-\frac{z_i}{z_0}}}{\sum_{i} e^{-\frac{z_i}{z_0}}}$$
(2)

Hear  $w(z_i)$  is the wind vector at altitude  $z_i$ , and  $z_0$  indicates the assumed scale height of 300m for winter.

According to the ealculation method of Ibrahim et al. (2010) CIM, the complete NO<sub>X</sub> emissions from the encircled areas are determined considering the partitioning between NO and NO<sub>2</sub>( $c_L$ ) and the finite lifetime of NO<sub>X</sub>( $c_\tau$ ).

$$E_{NO_X} = c_L \cdot c_\tau \cdot F_{NO_2} \tag{23}$$

$$c_L = \frac{c_{NO_X}}{c_{NO_2}} \tag{34}$$

Here  $c_L$  is simply the ratio of  $NO_X(C_{NO_X})$  and  $NO_2(C_{NO_2})$  bulk concentration in the polluted layer which are simulated by the CMAQ model in this study. It is a function of the Leighton ratio ( $L\underline{c}=[NO]/[NO_2]$ ),  $c_L=1+L\underline{c}$ . To analyze whether there is or not the impact of VOCs on lifetime of  $NO_X$ , we also calculate another Leighton ratios,  $L_T$ , referring to the method of Davis et al. (2019).

$$L_r = \frac{j_{NO_2[NO_2]}}{k_B[NO][O_2]} \tag{5}$$

Where  $j_{NO_2}$  is the NO<sub>2</sub> photolysis rate,  $k_8$  is the temperature-dependent rate constant for the reaction between NO and O<sub>3</sub>. We calculate  $j_{NO_2}$  according to the method of Dickerson et al., 1982

 $c_{\tau}$  describes the decay of NO<sub>X</sub> from the emission sources to measured positions.  $c_{\tau}$  can be estimated from the NO<sub>X</sub> lifetime  $\tau$ , which is the reciprocal of the product of reaction rate <u>constant coefficient k</u>, OH concentration ( $C_{OH}$ ) and air density (M)(Ma et al., 2013), and transport time t, which is the distance between emission source and sampling point r divided by the wind speed  $\overline{w}$ . w.

$$c_{\tau} = e^{\frac{t}{\tau}} = e^{\frac{r}{/_{\overline{\mathbf{ww}}}}}$$

$$(4\underline{6})$$

$$\tau = \frac{1}{k \cdot C_{OH} \cdot M} \tag{57}$$

We firstly calculated averaged simulated wind speed and direction, the ratio of  $NO_X$  and  $NO_2$ , and the  $NO_X$  lifetime from surface to 1000m at every sampling position on the 6th Ring Rd of Beijing for each journey, and computed the distance between the sampling position and the center of the city of Beijing for r. Then, we computed the  $c_\tau$ ,  $F_{NO2}$ , and  $F_{NOX}$ . The lifetime  $\tau$  was calculated with simulated average OH concentration and air density from surface to 1000m at each sampling position for each journey.

We averaged our model simulated quantities over the urban area for the NO<sub>x</sub> lifetime  $\tau$ , used the simulated wind speed at sampling position as w, and computed the distance between the sampling position and the center of the city of Beijing for r.

#### 2.2 Car MAX-DOAS measurements

# 2.2.1 Instrument and experiment

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We measured and retrieved tropospheric NO<sub>2</sub> VCD along the sixth ring road of Beijing (hereafter referred to as 6th Ring Rd) in January, September, and October of 2014 using a Mini MAX-DOAS instrument mounted settled on the vehicle.

The instrument, manufactured at Hoffmann Messtechnik GmbH, Germany, is a fully automated, light-weighted spectrometer designed for the spectral analysis of scattered sunlight by the MAX-DOAS technique (Hönninger et al., 2004; Davis et al., 2019). The same type of instrument was used in previous studies, including long-term site measurements in Beijing (Ma et al., 2013a)and a ear-Car MAX-DOAS observational journey in Europe (Wagner et al., 2010a). The instrument consists of a hermetically sealed

metal box of approximately 3 liter volume containing entrance optics, fiber coupled spectrograph and all electronics. A spectrometer with the model Ocean Optics USB2000+ is used. A stepper motor, adjusted outside the box, rotates the whole instrument to control the elevation viewing angle. The spectrograph covers the range 292-436 nm and its entrance slit is 50 µm wide. A Sony ILX511 CCD (charged coupled device) detects the light in 2048 individual pixels. The whole spectrograph is cooled by a Peltier stage to guarantee a stable temperature of the optical setup and a small dark current signal. For this study, the instrument was mounted on the roof of a car. Inside the car, two 12V DC batteries alternatively supplied electronic power for the running of instruments and a laptop computer, with a script run by the DOASIS software (Kraus, 2001b) to control the measurement process and the recording of spectra. The temperature of the spectrograph was set to be maintained at -5°C in January and at 0°C in September and October, well below the ambient temperatures during the experimental days of the study. The signal spectra of dark current and electronic offset were measured each day before and after the field experiment on the road, with 10000 msec and 1 scan for dark current measurements and 3 msec and 1000 scans for electronic offset measurements. Measurements were made alternatively at 30° and 90° elevation angles. with every 30° measurement immediately followed by a 90° measurement. Each elevation angle measurement had an integration time of about 1 min, including typically 300-400 scans for an average spectrum., with an integration time of about 1 min for each elevation angle.

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The instrument onboard the car was operated to measure scattered sunlight from the driving forward direction. There were no high buildings on both sides of the 6th Ring Rd., and the measurements were made at a wide-field view. The driving speed was typically controlled at 80–90 km h<sup>-1</sup>, and it generally took about 2.0–2.5 h to complete one circle (about 187 km) around the 6th Ring Rd. Figure 1 shows the driving route of the ear-Car MAX-DOAS experiment on a map of Beijing and distribution of large point sources of NOx from the MEIC inventory within the 6th Ring Rd. For this study, the field experiments were carried out on 14 selected days, with one or two circling circle-journeys each day. In total, there are 19 circling journeys available. The sampling periods in this experiment and the meteorological conditions are listed in Table 1. In most cases, the meteorological conditions changed slightly within one circling journey period. The average wind speeds for experimental days in January, September, and October were 2.5, 2.5, and 2.4 m s<sup>-1</sup>, the corresponding total cloud fractions were 4.9, 7.5, and 4.2, and the mean planetary boundary layer (PBL) heights were 192, 188, and 238 m, respectively. The dominant wind directions in the three months were much more variable, including north, south, and other directions.

other directions, and static wind field. Since variations of wind field can affect the estimation of E<sub>NOX</sub>, we synthetically analyze distribution of wind field using simulations from the WRF model and reanalysis data with a spatial resolution of 0.125° × 0.125° every three hours from the European Centre for Medium-Range Weather Forecasts (ECMWF). In some cases, the wind direction changed slightly within one circling journey period which is marked as south (S) or north (N) type in Table 1. However, the wind field during some journeys was convergent or divergent in some areas of Beijing which is marked as other type (O), and the wind speed was very low in three journeys which is marked as static type (St). To estimate the NOx emissions accurately using the CIM, the wind speed needs to be sufficiently high so that the transport across the encircled area is fast compared to the atmospheric lifetime of the trace gas (Ibrahim et al., 2010). In this study, we only consider the circling journeys with consistent wind field (S or N type) and relatively high wind speed to estimate the NOx emissions. The primary information for all the journeys, including 11 selected ones for emission estimation, is given in Table 1.

#### 2.2.2 Spectral retrieval

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The retrieval of NO<sub>2</sub> slant column densities (SCDs) is based on the DOAS method (Platt, 1994). The WinDOAS software (Fayt and Van Roozendael, 2011) was adopted to analyze the spectra in the 400-431 nm range on a daily basis. The Fraunhofer reference spectrum (FRS) was selected among the measured spectra at the 90° elevation angle each day by two steps: first, a spectrum measured around noon was chosen; second, the spectrum corresponding to the minimum NO<sub>2</sub> SCD derived in the preliminary analysis using the FRS from the first step was finally selected. The absorption cross sections of NO<sub>2</sub> at 294 K (Vandaele et al., 1998), O<sub>3</sub> at 221 K (Burrows et al., 1999), and the Oxygen dimer dimmer-O<sub>4</sub> at 298 K (Greenblatt et al., 1990), as well as a FRS, a Ring spectrum calculated from the FRS by DOASIS (Kraus, 2001a) and a polynomial of third order were included in the spectral fitting process. Figure 2 shows an example of our spectral analysis for a measurement on 18 January 2014, 11:39:38 BJT. As shown in the figure, the atmospheric NO<sub>2</sub> absorption structure can be clearly extracted from the measured spectra.

# 2.2.3 Derivation of tropospheric NO<sub>2</sub> VCD

The trace gas VCD in the troposphere can be calculated using its SCD divided by the air mass factor (AMF) at an elevation angle,  $\alpha$ :

$$VCD_{trop} = \frac{SCD_{trop}(\alpha)}{AMF_{trop}(\alpha)}$$
 (68)

For the <u>in-situ</u> <u>site</u> MAX-DOAS measurements, a FRS from the same elevation sequence was used in most cases, and the stratospheric absorption can be assumed to be the same during one elevation sequence. Therefore, the VCD<sub>trop</sub> can be calculated by extending Eq. <u>1-8</u> to Eq. <u>2-9</u> using the so-called differential tropospheric slant column density (DSCD<sub>trop</sub>( $\alpha$ ) = SCD<sub>trop</sub>( $\alpha$ )-SCD<sub>trop</sub>(90°)) divided by the differential air mass factor (DAMF<sub>trop</sub>( $\alpha$ ) = AMF<sub>trop</sub>( $\alpha$ )-AMF<sub>trop</sub>(90°)):

$$VCD_{trop} = \frac{DSCD_{trop}(\alpha)}{DAMF_{trop}(\alpha)} = \frac{DSCD_{meas}(\alpha)}{DAMF_{trop}(\alpha)}$$
(79)

with  $DSCD_{meas}(\alpha) = SCD_{meas}(\alpha) - SCD_{ref}$  (Wagner et al., 2010b; Ma et al., 2013a).

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For the <u>car\_Car\_MAX-DOAS</u> measurements, the trace gas concentrations can change significantly during one measurement sequence and thus the dependence of retrieved trace gas DSCDs on the elevation angle may not be so regular as for the <u>in-situ site-measurements</u>. Therefore, it would be a better choice to use a single FRS for the analysis of all the spectra measured along the driving route (Wagner et al., 2010b). According to Wagner et al. (2010b), Eq.<u>1-8</u> can be further extended to

$$VCD_{trop} = \frac{DSCD_{meas}(\alpha) - DSCD_{offset}(SZA)}{AMF_{trop}(\alpha)}$$
 (108)

where  $DSCD_{offset}$  depends on the solar zenith angle (SZA) and thus local time,  $t_i$ . For each elevation sequence i during the individual measurement day,  $DSCD_{offset}$  is calculated from a single pair of measurements with

$$DSCD_{offset}(t_i) = \frac{AMF_{trop}(90^\circ) \cdot DSCD_{meas}(\alpha, t_i) - AMF_{trop}(\alpha) \cdot DSCD_{meas}(90^\circ, t_i)}{AMF_{trop}(\alpha) - AMF_{trop}(90^\circ)}$$
(119)

The time series of the calculated  $DSCD_{offset}(t_i)$  in this study could be fitted by a low-order polynomial, e.g.,  $P(x) = a_0 + a_1 \cdot x + a_2 \cdot x^2$ , as a function of time. The fitted polynomial then represents the best guess for  $DSCD_{offset}$  and can be used to calculate the  $VCD_{trop}$  from Eq. <u>103</u>. In this study, the AMF was calculated by the <u>geometry approximation (Brinksma et al., 2008; Wagner et al., 2010b)</u>, that is:

$$AMF_{trop}(\alpha) \approx \frac{1}{\sin(\alpha)}$$
 \_ (1240\_)

As an illustration, Figure 3 shows the changes of individual NO<sub>2</sub> DSCD<sub>meas</sub> and DSCD<sub>offset</sub> for 30° elevation angle of each sequence as a function of time on 18 January 2014. As shown in Fig. 3, aA second order polynomial fitted from individual DSCD<sub>offset</sub> data points as shown in Fig. 3 tends to be stable and

can be used to represent an average value of DSCD<sub>offset</sub>. tends to converge against a much more stable average DSCD<sub>offset</sub> value.

# 2.2.4 Calculation of monthly average NO2 VCD

# 2.3 LAPS-WRF-CMAQ model simulation

#### 2.3.1 Model setup and data

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To quantify the NO<sub>X</sub> emissions in Beijing more accurately, refined simulations of the wind field and NO<sub>2</sub> to NOx concentration ratio were needed. In this study, we utilized the offline LAPS-WRF-CMAQ model system with high spatiotemporal resolution and data assimilation technique to obtain the refined wind speed and wind direction and an accurate ratio of NO<sub>2</sub> and NO<sub>X</sub> concentration during the ear-Car MAX-DOAS experiments. The aforementioned model system includes three components: the LAPS model (Albers et al., 1996), the WRF model (Michalakes et al., 2004), and the CMAQ model (Dennis et al., 1996). Simulation of wind speed and direction is improved by the LAPS-WRF model, which assimilates observed data at the surface and high layers using the one-dimensional and three-dimensional variational assimilation method (Albers et al., 1996). The CMAQ model is used to simulate temporal-spatial distribution of NO2 and NO concentration. The Local Analysis and Prediction System (LAPS), developed by the NOAA Earth System Research Laboratory, is used in many numerical weather forecast centers around the world. It is a mesoscale meteorological data assimilation tool that employs a suite of observations to generate a realistic, spatially distributed, time-evolving, three-dimensional representation of atmospheric features and processes (McGinley et al., 1991). The three-dimensional realistic meteorological analyses field can be used as the initial condition of the WRF model and improve the simulation of wind field. WRF is a mesoscale numerical weather prediction system designed for both atmospheric research and operational forecasting needs. CMAQ is an air-quality model developed by the U.S. Environmental Protection Agency's Atmospheric Science Modeling Division. It consists of a suite of computer programs for modeling air quality issues, including reactive gases such as NO<sub>2</sub>, NO, SO<sub>2</sub>, O<sub>3</sub>, and others, particulate matter (PM), air toxics, acid deposition, and visibility degradation.

This study focused on Beijing at a horizontal resolution of 4 km × 4 km with 31 vertical layers of varying thickness (between the surface and 50 hPa) using a triple-nested simulation technique. The horizontal resolutions of the three sets of grids were 36 km, 12 km, and 4 km, respectively (Fig. S1a), and the output temporal interval was 1 h. The LAPS-WRF simulations were driven by FNL/NCEP

analysis data every 6 h during the car-Car MAX-DOAS experiments, with a spatial resolution of 1° × 1°. In addition, to improve the simulation of wind field and NO<sub>2</sub> and NO concentrations, many meteorological data of the same periods, such as wind speed, wind direction, air temperature, and relative humidity, observed at 2400 surface weather stations and by 120 radiosonde stations were assimilated into the initial field of the WRF model using the one-dimensional and three-dimensional variational assimilation method in the LAPS model. The CMAQ model uses the multi-resolution emission inventory in China for the year 2012 (MEIC 2012) MEIC 2012 with 0.25° × 0.25° resolution (Zhang et al., 2009; Li et al., 2017).- Hourly gridded MEIC emission datasets at a horizontal resolution of 4 km × 4 km for the CMAQ model were generated by the Sparse Matrix Operator Kernel Emissions (SMOKE) modeling system (UNC, 2014) using reasonable temporal and spatial allocation coefficients (Cheng et al., 2017). Meteorological outputs from the WRF simulations were processed to create model-ready inputs for CMAQ using the Meteorology-Chemistry Interface Processor (MCIP) (Otte and Pleim, 2010). The chemical mechanism is CB05, and the boundary conditions of trace gases consist of idealized, Northern Hemispheric, mid-latitude profiles based on results from the NOAA Agronomy Lab Regional Oxidant Model. The model simulation was started one day before the first day of the experiment to avoid the spinup problem and improve the simulation accuracy.

# 2.3.2 Validation of simulated surface wind and NO2

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Modelled wind speeds and directions were validated by observation data from four weather stations in Beijing. Figures S2, S3 and S4show the scatter distribution between simulated wind speed and observation, wind rose of modelled wind direction and measurements, and their time serials. We adopted the The\_observed hourly wind speed and direction data at the meteorological stations, shown in Figures S2 and S3, were obtained from the China Meteorological Administration. The four stations are the from Nanjiao (NJ), Tongzhou (TZ), Mentougou (MTG), and Shunyi (SY) meteorological stations, which represent the south, east, west, and north areas of Beijing, respectively. It was shown that the temporal variation in simulated wind speed at the four stations were consistent with the observations from the perspective of time serial of wind speed, but the simulations were higher than the observations due to impacts of the complex topography and limited observation data assimilated to the LAPS-WRF model (Fig. S2 and Fig. S4a)(Fig. S2). To calculate the E<sub>NOX</sub> accurately, we corrected the simulated wind speed using the observation data from the four weather stations in order to reduce the systemic error.

Specifically, we computed the relative error of the modeled wind speed based on measurements at four weather stations for each journey and then added the error bar to simulated wind speed at every sampling position during the same journey, we computed the relative error of modelled wind speed during every journey and then used it to correct the simulated wind speed at all sampling points for every journey. The correlation coefficient between simulated and observed wind speeds at the four stations was is 0.47, and the result passespassed the 99.9% significance test. The root mean square error (RMSE) was is small, with a value of 1.18 m s<sup>-1</sup>. Except for the MTG station, simulated wind directions at the other three stations are were in accordance with the observations, particularly for the primary wind direction (Fig. S3 and Fig.S4b). In general, simulated wind direction are also coincident with observations from the perspective of time serial of wind direction, and simulations are larger than measurements during some periods at some stations due to the effects of the complex topography and limited observation data assimilated to the model (Fig. S4b). The primary wind direction and its frequency at the MTG station were are not consistent with the observations because these are affected by the complex topography near the Taihang and Yanshan mountains. In general, the corrected wind speed and wind direction data are reliable for estimation of the NO<sub>X</sub> emissions, and the uncertainty of E<sub>NOX</sub> due to the variation of wind field will be discussed in Section 3.3. Hence, the simulations of wind speed and wind direction were reliable for estimation of the NO<sub>X</sub> emissions.

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Figure S4 presents the temporal variation in simulated and observed NO<sub>2</sub> concentration from January 18 to October 13, 2014. The hourly measurements of NO<sub>2</sub> concentrations (shown in Fig. S1b) were obtained from the National Environment Monitoring Station in China. In general, the temporal variation in the NO<sub>2</sub> simulation was is consistent with the observation. The simulated values were are close to the observations, except for January 21–24, September 19, and October 9–10, when NO<sub>2</sub> simulations were are higher than the observations. The correlation coefficient between simulated and observed NO<sub>2</sub> concentrations was is 0.73, and the result passes passed the 99.9% significance test (Fig. S5). The RMSE and mean absolute error (MAE) were are 16.14 and 19.21 μg m<sup>-3</sup>, respectively. Because The observed NO<sub>2</sub> might include the some NO<sub>2</sub> component, it can lead leading to a systematical biases (underestimation) of NO<sub>2</sub> by model compared to observation (Ma et al., 2012). Thus, the simulated NO<sub>2</sub> concentrations and hence the ratio of NO<sub>2</sub> and NO<sub>3</sub> were are reliable for estimating NO<sub>3</sub> emissions.

#### 2.4 Selection of the journeys for estimating NOx emissions

To estimate the NO<sub>2</sub> fluxes ( $F_{NO2}$ ) and  $F_{NOX}$  accurately, we firstly selected six journeys with the RMSEs of simulated wind speeds at the four weather stations smaller than 1.5 m s<sup>-1</sup> from the primary selected 11 journeys. Then we assessed whether the meteorological and chemical conditions meet the criteria of Shaiganfar et al. (2017) for each of these six journeys. It should be pointed out that we cannot assess the problem of large partitioning ratio due to the absence of the whole seasonal simulated or observed data in autumn and winter. The assessment results of other four problems are listed in Table 2. We excluded the journeys in which more than two problems occurred. It needs to be noted that lifetime correction coefficients  $c_{\tau}$  on October 12 and 13 are slightly larger than 1.5, which is the criteria of large lifetime correction (Shaiganfar et al., 2017), so we also adopted the data on October 12 and 13 to estimate the  $E_{NOX}$ . Lastly, NO<sub>2</sub> VCD measurements outside of the 6th Ring Rd during five selected journeys were not used to quantify  $F_{NO2}$  and  $E_{NOX}$ .

#### 3. Results and discussion

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# 3.1 Tropospheric NO<sub>2</sub> VCD

Figure 4 presents the <u>journey-to-journey</u> temporal variation in the tropospheric NO<sub>2</sub> VCD on the 6th Ring Rd of Beijing in January, September, and October, 2014. In general, the NO<sub>2</sub> VCD in January was higher than that in other months. The highest values falling between 8 × 10<sup>16</sup> and 13 × 10<sup>16</sup> molecules cm<sup>-2</sup> occurred on January 19, 23, and 24. The mean, maximum, and minimum NO<sub>2</sub> VCD during the sampling periods were all larger in January than in the other two months. The mean NO<sub>2</sub> VCD ranged mostly from 4.5±1.83 × 10<sup>16</sup> to 9.0±1.24 × 10<sup>16</sup> 4.5 × 10<sup>16</sup> to 9 × 10<sup>16</sup> molecules cm<sup>-2</sup> in January, but values were all lower than 4.5 × 10<sup>16</sup> molecules cm<sup>-2</sup> in September and October. The NO<sub>2</sub> VCD values during the mornings of January 23 and October 13 were 9.05 × 10<sup>16</sup> and 1.23 × 10<sup>16</sup> molecules cm<sup>-2</sup>, corresponding to the maximum and minimum values, respectively, during the 19 circling journeys. This result may might be caused by higher emissions from coal fired heating (Table S1) and lower photolysis of NO<sub>2</sub> in winter, and some meteorological conditions that were unfavorable for dispersion and transport of pollutants in winter. Lower PBL height and lower wind speed suppress horizontal and vertical diffusion and transport of NO<sub>2</sub>. Southwest and east winds are favorable for the transport of air pollutants from the south and east areas to the city of Beijing. Higher cloud cover is unfavorable for photolysis of NO<sub>2</sub>. A similar pattern of seasonal variation in tropospheric NO<sub>2</sub> VCD was found previously by site MAX-DOAS

measurements in Beijing(Ma et al., 2013a; Hendrick et al., 2014).

To investigate the differences in the spatial distribution of NO<sub>2</sub> VCD among the three months, we computed the monthly average NO<sub>2</sub> VCD for every sampling point along the 6th Ring Rd of Beijing in January, September, and October, 2014. Firstly, we used the locations of all sampling points on the morning of September 23 as the reference point for the calculation of NO<sub>2</sub> VCD monthly average, with the most sampling sites (98 points) for all observation periods. Then, we calculated the monthly average value at each reference point using the data of the nearest sampling point. The distance from the nearest sampling point to a reference point was less than 1.5 km. Figure 5 shows that the monthly average NO<sub>2</sub> VCD values at most sampling points on the 6th Ring Rd were obviously higher-larger in January than in the-other two months (by a factor of two in most cases). The spatial distribution characteristics of NO<sub>2</sub> VCD in September were similar to those in October. In addition, the NO<sub>2</sub> VCD values at the northern and southern parts of the 6th Ring Rd were all larger than those in other areas for all three months. The high NO<sub>2</sub> VCD in the southern region was related to strong local emissions to the south of Beijing and transport from central and southern Hebei and the city of Tianjin (Meng et al., 2018). As shown in Fig. 4, the maximum journey-averaged NO<sub>2</sub> VCD occurred on the morning of January 23, and the minimum occurred on the morning of October 13.

as shown in Fig. 6. The NO<sub>2</sub> VCD values on the 6th Ring Rd in the morning of January 23 were all large, particularly in the north and southwest areas, with magnitudes of  $10 \times 10^{16}$  to  $12 \times 10^{16}$  molecules cm<sup>-2</sup>. On October 13, high NO<sub>2</sub> VCD was located in at the southern areas and it might be related to the two southern emission sources, which were closer to the 6th Ring Rd with higher emission rates compared to the northern ones with lower emission rates. but values were lower in the northern areas. The spatial distribution differences between these two journeys were related to the high emission during the heating season in January (see section 3.2) and the impacts of the wind field. To investigate the impact of the wind field on the spatial distribution of NO<sub>2</sub> VCD, weWe used thin-grid ECWMF reanalysis data for January 23 and October 13 with a spatial resolution of  $0.125^{\circ} \times 0.125^{\circ}$  to investigate the impact of the wind field on the spatial distribution of NO<sub>2</sub> VCD. Figure 7 shows the distribution difference of wind fields at 8:00 and 14:00 BJT on these two days, respectively. The NO<sub>2</sub> VCD was large with weak south wind and with convergence of southeast and northwest wind in Beijing and its surrounding area, but its values were far smaller with strong north wind. Weak south wind and a breeze or calm wind resulted in

We investigated the spatial distribution differences in NO<sub>2</sub> VCD between these two circling journeys,

the transport of NO<sub>2</sub> from the south area in Hebei Province and its accumulation on January 23. Strong north wind suppressed the transport of NO<sub>2</sub> from the south area on October 13. These results indicate that the wind field has large impacts on the spatial distribution of NO<sub>2</sub> VCD in Beijing.

Figure 8 presents the spatial distributions of wind and NO<sub>2</sub> VCD averaged for the three\_two different wind fields. The mean NO<sub>2</sub> VCD at most sampling points along the 6th Ring Rd was obviously higher under the south wind field than under the north wind. High NO<sub>2</sub> emission in the three months was sources were located within the 5th Ring Rd of Beijing in the three months (Fig. 10), and the background concentrations of NO<sub>2</sub> VCD in the north and south areas were remarkably different due to the impacts of emission sources from to the south areas of Beijing, such as Hebei Province. Hence, southerly wind can transfer air pollutants from the southern area to Beijing and lead to high NO<sub>2</sub> flux and NOx emission, whereas impacts of north wind on NO<sub>2</sub> flux and NOx emission are smaller because the background concentration of NO<sub>2</sub> VCD in to the north of Beijing is were lower. Convergence of the wind field in the south parts of the 6th Ring Rd is favorable to the accumulation of NO<sub>2</sub> from the surrounding area to the southern parts of the ring road.

# 3.2 Quantification of NOx emissions

Figure 9 shows the journey-to-journey variation in estimated  $F_{NO2}$  and  $E_{NOX}$  over Beijing for five circling journeys in January, September, and October, 2014. To estimate the NO<sub>2</sub> fluxes ( $F_{NO2}$ ) and  $F_{NOX}$  accurately, we used the data from 10 circling journeys (Table 1), for which the RMSEs of simulated wind speeds at the four weather stations were all less than 1.5 m s<sup>-1</sup>. In addition, NO<sub>2</sub> VCD measurements at the sampling points outside of the 6th Ring Rd during 11 circling journeys were not used to quantify  $F_{NO2}$  and  $F_{NOX}$ . Figure 9 shows the journey to journey variation in estimated  $F_{NO2}$  and  $F_{NOX}$  over Beijing for the 10 circling journeys in January, September, and October, 2014. The  $F_{NO2}$  fell in between  $F_{NO2}$  molecules  $F_{NOX}$  molecules  $F_{NOX$ 

Specifically, the total E<sub>NOX</sub> was higher in January than in the other two months. The average E<sub>NOX</sub> in the heating period was about 3.12.5 times as much as those in the non-heating period. The coal fired heating in Beijing included central heating in urban area and scattered coal combustion in suburb or rural area for the year 2014. We calculated the average NO<sub>X</sub> emission rates of four sectors including industry, power, residential, and transportation from the MEIC within the 6th Ring Rd of Beijing in January, September, and October 2012, and the ratio of each specific NO<sub>X</sub> emission rate in January to the corresponding average value in September and October (Table S1). The E<sub>NOX</sub> from the power and residential section were remarkably higher in January than in other two months, and especially E<sub>NOX</sub> from the residential was about 5 times more in January than in other months. In general, central heating in urban area are from power plant and residential use the scattered coal combustion in suburb or rural area.

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In addition to the seasonal differences, the journey-to-journey variation in estimated  $E_{NOX}$  were is large even within the same month, mainly due to uncertainties in the calculations of wind speed, ratio of NO<sub>2</sub> and NO<sub>3</sub> concentration, and decay rate of NO<sub>3</sub> from the emission sources to the measured positions under different meteorological conditions. In addition to the NO<sub>2</sub> VCD, wind speed, and wind direction at the sampling points, the estimated NO<sub>3</sub> emission rate is obviously affected by the Leighton ratio of NO and NO<sub>2</sub> concentration and the lifetime of NO<sub>3</sub> (Valin, et al., 2013). Thus, the estimated NO<sub>3</sub> emission rate could be very large even if the NO<sub>2</sub> VCD was small, such as in the case of September 14January 27. It should be noted that the low mean wind speed on September 14 leads January 27 was relatively small and led to higher  $c_{\tau}$ , so the E<sub>NO3</sub> for this journey is not so low although the E<sub>NO2</sub> was very low. In addition, if both  $c_{\tau}$  and  $c_{L}$  are large, high E<sub>NO3</sub> can be derived, meanwhile, the ratios of NO<sub>3</sub> and NO<sub>2</sub> were relatively large, so E<sub>NO3</sub> on January 27 was large although F<sub>NO2</sub> was relatively small. Thus, if  $c_{\tau}$  and  $c_{L}$  are simultaneously larger, higher E<sub>NO3</sub> occurs. However, if only one factor is larger and the other is smaller, such as higher  $c_{\tau}$  and lower  $c_{L}$  as on January 18, the morning and afternoon of September 14, and the morning of October 13, E<sub>NO3</sub> is lower.

# 3.3 Comparisons with MEIC inventory and other estimates

We compared the estimated  $NO_X$  emission with the multi-resolution emission inventory in China (MEIC) released by Tsinghua University for 2012 (MEIC 2012) (Zhang et al., 2009;Zhang et al., 2012). The horizontal resolution of MEIC 2012 is  $0.25^{\circ} \times 0.25^{\circ}$ , and five sectors, <u>i.e.</u>, agriculture, industry, power, <u>residentialresidents</u>, and transportation, are included.

Figure 10 presents the spatial distributions of  $NO_X$  emission rates over Beijing in January, September, and October, 2012, from MEIC. A high  $NO_X$  emission zone was located within the 5th Ring Rd of Beijing, and a low emissions zone occurred in its surroundings in other areas. The  $NO_X$  emissions in January were obviously larger than those in the other two months. The concentrated distribution of  $NO_X$  emission sources within the 5th Ring Rd of Beijing indirectly indicates the applicability of Eq. (1) to estimate the

NOx emission rates from the car MAX DOAS measurements on the 6th Ring Rd in this study.

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Figure 11 shows the journey to journey estimated NO<sub>x</sub> emission rates from ear Car MAX-DOAS measurements for each selected journey (see Sect. 2.4) in January, September, and October, 2014 (denoted as  $E_{NOX}$ ), and the corresponding monthly averaged NO<sub>x</sub> emission rates from the MEIC 2012 for the same region within the 6th Ring Rd of Beijing (hereafter expressed as MEIC Month). The MEIC Month is obviously lower than the estimated E<sub>NOX</sub> in January. While the two emission estimates are very close in September, the MEIC Month is slightly smaller than the E<sub>NOX</sub> in October. In most cases, the MEIC Month was lower than the estimated E<sub>NOX</sub>, particularly in January. The differences between the estimated E<sub>NOX</sub> and the MEIC Month during some journeys were remarkably large. The differences between the E<sub>NOX</sub> and MEIC 2012 during the 10 journeys may be caused by (1) the interannual variations differences in emission inventory, (2) the different timescales of the two emission estimates, (3) the uncertainty of the estimated E<sub>NOX</sub> and MEIC 2012, (4) inconsistency of wind field during the period of measurements, (5) extra transfers from source areas other than urban Beijing, and so on. Firstly, the  $E_{NOX}$  in this study was is estimated for the year 2014, whereas the MEIC MonthMEIC 2012 was established for the year 2012. Secondly, our results represented only the conditions during a few measurements during daytime, whereas the MIEC 2012 denoted monthly average conditions. Thirdly, the uncertainty of MEIC 2012 is large, particularly in autumn and winter (Li et al., 2017; Meng et al., 2018). There are also large uncertainties in the estimated E<sub>NOX</sub> caused by, e.g., the inconsistency of wind field during a circling journey and the transfer of NO<sub>2</sub> from other source areas than urban Beijing. Fourthly, the emission estimation method used in this study The CIM assumes that the wind field is constant during the period of measurements period and that the wind speed is also sufficiently high that the transport across the encircled area is fast compared to the atmospheric lifetime of the trace gas (Ibrahim et al., 2010). However, the wind field during some journeys (January 27 and October 12,13) might could have changed systematically and been convergent or divergent in some areas of Beijing, as

changes during the period of measurements period can have large impacts on become important to the emission estimate, particularly if measurements with high trace gas VCD are accompanied by strong deviations of the actual wind speed (or direction) from the assumed average values. For example, on the morning afternoon of January 27, the highest high NO<sub>2</sub> VCD was measured, and the wind field changed during the measurement journey. In such cases, the systematic changes in wind speed and direction can lead to additional uncertainties in E<sub>NOX</sub>. Moreover Finally, because the southerly wind can bring NO<sub>x</sub> emitted in the south-central regions of Hebei Province to Beijing, the E<sub>NOX</sub> from ear-Car MAX-DOAS measurements will be overestimated under south wind conditions, e.g., on January 26.

#### 3.4 Uncertainty analysis of estimated emissions.

We calculated the uncertainty of  $E_{NOX}$  according to the error transfer formula of relative deviation based on the errors of measured NO<sub>2</sub> VCD, simulated wind speed and direction, and  $c_{\tau}$  and  $c_{\tau}$ . Figure 11 also shows the uncertainty of  $E_{NOX}$ , calculated from the errors of measured NO<sub>2</sub> VCD, simulated wind speed,  $c_{\tau}$  and  $c_{\tau}$  according to the error transfer formula of relative deviation. The standard deviation (STD) of wind speed over a period of time can provide a bound for the related uncertainties of the emission estimate (Ibrahim *et al.*, 2010). Therefore, we first computed the uncertainty of  $F_{NO2}$  based on the STD of simulated wind speed after correction the corrected wind speed and the measurement error of NO<sub>2</sub> VCD (about ± 10%, Ma *et al.*, 2013a) for each journey. Then, we calculated STD of  $c_{\tau}$  according to the first derivative of equation (4) and the monthly-STD of  $c_{L}$  using different NOx lifetime and the ratios of NO<sub>X</sub> and NO<sub>2</sub> at sampling position on the 6th Ring Rd of Beijing for each journey. Its regional average data within the 6th Ring Rd of Beijing during all journeys in each month. We used the identical STD of  $c_{L}$  for each journey in the same month to calculate of the uncertainty of  $E_{NOX}$ -Figure 11 shows the uncertainties of  $E_{NOX}$  for five journeys and The results showed that the STD ranges of wind speed,  $c_{L}$  and  $c_{T}$  were 0.13-1.30 m s<sup>-1</sup>, 0.11-0.37, and 0.17-1.97, respectively. The the uncertainty range of  $E_{NOX}$  for five journeys molecules  $s^{-1}$  (19.52–52.01%), was 16.4-33.2%.

We also give the spatial variation in the  $NO_X/NO_2$  ratio and  $NO_X$  lifetime at the entire route for the emission calculation during five journeys (Fig. S7 and S8), and estimate the error contribution of five factors including  $NO_2$  VCD, wind speed, wind direction, the  $NO_X/NO_2$  ratio, and the  $NO_X$  lifetime to the total uncertainty of  $E_{NOX}$  (Table 3). In general, there are obvious seasonal and regional difference in the  $NO_X/NO_2$  ratio and  $NO_X$  lifetime and it is necessary to use specific ratios and lifetime value to estimate

the E<sub>NOX</sub> for each journey. Specifically, the NO<sub>X</sub>/NO<sub>2</sub> ratio and NO<sub>X</sub> lifetime are larger in January than in September and October, and they are larger at the southern part of the 6th Ring Rd than at other parts for most journeys. Among error contributions of five factors, the impacts of wind speed and direction are the largest for most journeys except for September 14. For January 26 and 27, error contributions of wind speed to the uncertainty of E<sub>NOX</sub> are larger than other four factors. For September 14, uncertainty of E<sub>NOX</sub> is mainly caused by the errors of NO<sub>X</sub> lifetime and wind direction. For October 12 and 13, error contributions of the NO<sub>X</sub>/NO<sub>2</sub> ratio are also remarkable. Thus, it is important to obtain the accurate wind vector profiles, NO<sub>X</sub>, NO<sub>2</sub>, and OH concentration data except for NO<sub>2</sub> VCD to reduce the uncertainty of E<sub>NOX</sub> estimation using the CIM.

We also calculate the Leighton ratios,  $L_r$ , to assess impacts of VOCs on the NOx lifetime. The  $L_r$  during five journeys is 0.85, 0.80, 1.04, 1.19, and 1.33 on January 26 and 27, September 14, October 12 and 13, respectively. Results show that the NOx lifetime for the three journeys on September and October are affected by VOCs and lead to extra errors of  $E_{NOX}$ . While VOCs can't cause the deviation in the NOx lifetime and estimation of the  $E_{NOX}$  in January.

# 4. Conclusions

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We carried out 19 city-circle-around ear-Car MAX-DOAS experiments on the 6th Ring Rd of Beijing in January, September, and October, 2014. The VCD of NO<sub>2</sub> was retrieved and the temporal and spatial distributions were investigated. Then the NO<sub>X</sub> emission rates in urban Beijing were estimated using the measured NO<sub>2</sub> VCD together with the refined wind fields, NO<sub>2</sub> to NO<sub>X</sub> ratios, and NO<sub>2</sub> lifetimes simulated by the LAPS-WRF-CMAQ model system, and the results were compared to the emission rates from the MEIC inventory 2012.

The mean, maximum, and minimum-NO<sub>2</sub> VCD values averaged for each experimental journeyduring the sampling periods in January were all larger than those in the other two months, mainly due to higher emissions in winter. The measured mean-NO<sub>2</sub> VCD was typically larger at the southern parts of the 6th Ring Road than at the northern parts because weak south wind resulted in the transport and accumulation of NO<sub>2</sub> from southern areas in Hebei Province and strong north wind suppressed the transport of NO<sub>2</sub> from the southern area. Such inhomogeneous distributions of tropospheric NO<sub>2</sub> VCD bring a challenge for the validation of satellite products for Beijing as well as other megacities.

The journey-to-journey variation in estimated E<sub>NOX</sub> were large, even within the same month, mainly

due to uncertainties in the calculation of wind speed, the ratio of NO<sub>2</sub> and NOx concentration, and the decay rate of NO<sub>X</sub> from the emission sources to the measured positions under different meteorological conditions. The ranges of  $E_{NOX}$  during the heating and non-heating periods were  $28.7 \times 10^{25}$  to  $60.0 \times 10^{25}$  molecules s<sup>-1</sup> and  $7.7 \times 10^{25}$  to  $24.8 \times 10^{25}$  molecules s<sup>-1</sup>, respectively. The average  $E_{NOX}$  values in the heating and non-heating periods were are estimated to be  $26.94 \pm 6.1443.0 \times 10^{25}$  molecules s<sup>-1</sup> and  $10.99 \pm 1.23 \pm 3.9 \times 10^{25}$  molecules s<sup>-1</sup>, respectively, with an. The uncertainty range of  $E_{NOX}$  was  $19.52 - 52.01\% \pm 16.4 \pm 33.2\%$ . The monthly emission rates in the area within the 6th Ring Rd of Beijing from MEIC 2012 are were lower than the estimated  $E_{NOX}$ , particularly in January. The differences between the  $E_{NOX}$  and the monthly emission rates from MEIC 2012 may can be attributed attributable to the interannual differences in the emissions inventory, the different timescales and uncertainties of two kinds of inventory, inconsistencies of wind field during the period of measurements, and extra transfers from source areas other than urban Beijing.

Our results showed that ear-Car MAX-DOAS measurements can be used effectively for dynamic monitoring and updating of the  $NO_x$  emissions from megacities such as Beijing. To estimate the  $E_{NOX}$  by ear-Car MAX-DOAS accurately in Beijing and other similar megacities, appropriate meteorological conditions, such as smaller fluctuations of the wind field, relatively larger wind speed, and suitable wind direction, need to be selected to avoid the impact of extra transfers of large emission sources from surrounding areas. In addition to the  $NO_2$  VCD, simultaneous observations of wind speed, wind direction, and surface NO and  $NO_2$  concentrations are recommended to reduce the uncertainties of  $E_{NOX}$   $e_{\pi}$  and  $e_{E}$ .

Data availability. The NCEP-FNL reanalysis and ECMWF are publicly available at <a href="http://rda.ucar.edu/datasets/ds083.2/">http://rda.ucar.edu/datasets/ds083.2/</a> and <a href="https://www.ecmwf.int/en/forecasts/datasets">https://www.ecmwf.int/en/forecasts/datasets</a>, respectively. The NO<sub>2</sub> - measurements and meteorological observations including wind speed and wind direction data are available at <a href="http://l13.108.142.147:20035/emcpublish">http://l13.108.142.147:20035/emcpublish</a> and <a href="http://data.cma.cn/">http://data.cma.cn/</a>, respectively. The tropospheric NO<sub>2</sub> VCD data derived from this study are available on the request.

Author contributions. JM and XC designed the research. JM, JJ, JG, MQ, QX, and PY contributed to the measurements, and JM performed the spectral analysis and retrieval. XC and JP designed the model experiment and performed the model simulations. XC, YL, JP, and XM contributed to the data processing and analyses. XC and JM analyzed the results and wrote the paper with inputs from all authors.

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Table 1. Sampling periods of the <u>car\_Car\_MAX-DOAS</u> experiment and corresponding meteorological conditions over Beijing in January, September, and October, 2014.

Journey	Date	Time (BJT)	Wind speed (m/s)	Type of wind field*	Total cloud fraction	PBL Height (m)
1**	2014/1/18	10:48-13:09	2	<u>St</u> O	0	564
2	2014/1/19	13:31-15:40	1	O	7	167
3 <u>**</u>	2014/1/21	13:15-15:32	3	S	0	163
4	2014/1/23	10:39-12:25	3	O	7	187
5	2014/1/23	13:07-15:12	2	O	7	163
6 <u>**</u>	2014/1/24	10:42-12:03	2	N	8	39
7 <u>**</u>	2014/1/24	13:03-15:09	3	N	8	39
8**	2014/1/26	10:21-12:13	5	S	5	341
9 <u>**</u>	2014/1/27	09:11-11:38	2	<u>St</u> O	7	75
10**	2014/1/27	13:30-15:28	2	O	0	178
11**	2014/9/14	09:40-12:52	4	N	10	173
12**	2014/9/14	15:02-17:17	2	N	10	226
13**	2014/9/17	09:07-11:42	2	<u>St</u> O	7	173
14 <u>**</u>	2014/9/19	09:09-11:50	2	S	3	178
15	2014/10/9	13:04-14:44	1	<u>St</u> S	7	43
16 <u>**</u>	2014/10/10	09:52-12:28	2	S	7	663
17**	2014/10/12	14:02-16:42	3	N	7	167
18**	2014/10/13	09:12-11:59	3	<u>N</u> O	0	186
19 <del>**</del>	2014/10/13	13:11-16:27	3	O	0	130

<sup>\*</sup>Four types of wind filed are South (S), North (N), Other (O), and Static (St).

<sup>\*\*</sup>The data are preliminarily selected to estimate the NO<sub>X</sub> emissions.

<sup>\*</sup>Three types of wind filed are South (S), North (N) and Other (O).

<sup>\*\*</sup>The data from ten circling journeys are used to estimate the NOx emission.

Table 2. Overview of the problems for the six circling journeys.

<u>Date</u>	Time (BJT)	Large wind variability *	Large lifetime correction*	Gap/route close to the centre*	Small difference between influx and outflux*	Multiple problems **
2014/1/26***	10:21-12:13	<u>N</u>	<u>N</u>	<u>N</u>	<u>Y</u>	<u>N</u>
2014/1/27***	13:30-15:28	<u>Y</u>	<u>N</u>	<u>N</u>	<u>N</u>	<u>N</u>
2014/9/14	09:40-12:52	<u>Y</u>	$\underline{\mathbf{Y}}$	<u>Y</u>	<u>N</u>	<u>Y</u>
2014/9/14***	<u>15:02-17:17</u>	<u>N</u>	<u>Y</u>	<u>N</u>	<u>N</u>	<u>N</u>
2014/10/12***	14:02-16:42	<u>Y</u>	<u>Y</u>	<u>N</u>	<u>N</u>	<u>Y</u>
2014/10/13***	09:12-11:59	<u>Y</u>	<u>Y</u>	<u>N</u>	<u>N</u>	<u>Y</u>

<sup>\*</sup>Whether the condition meets the criteria of Shaiganfar et al. (2017) or not, Y and N denote Yes and No respectively.

\*\* Multiple problems mean whether more than two conditions can meet the criteria or not.

\*\*\* The data of five circling journeys are ultimately used to estimate the NOx emission.

Table 3. Error contributions (%) of multiple factors to the uncertainties in estimated  $E_{NOX}$  during five circling journeys.

<u>Factors</u>	Jan. 26, AM	Jan. 27, PM	Sep. 14, PM	Oct. 12, PM	Oct. 13, AM
$\underline{ ext{VCD}_{ ext{geo}}}$	<u>10</u>	<u>10</u>	<u>10</u>	<u>10</u>	<u>10</u>
Wind speed	<u>27.02</u>	<u>26.83</u>	<u>7.97</u>	<u>33.10</u>	3.68
Wind direction	<u>10.97</u>	<u>16.50</u>	<u>20.54</u>	<u>33.78</u>	38.37
NOx/NO <sub>2</sub> ratio	12.21	<u>13.46</u>	<u>7.82</u>	<u>29.33</u>	29.48
<u>Lifetime</u>	3.63	7.67	<u>48.60</u>	<u>15.22</u>	<u>19.02</u>

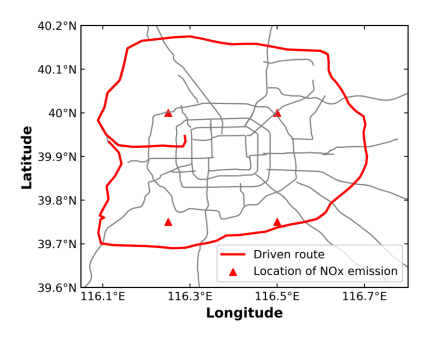
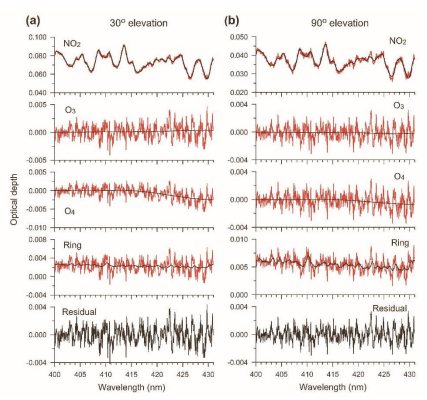
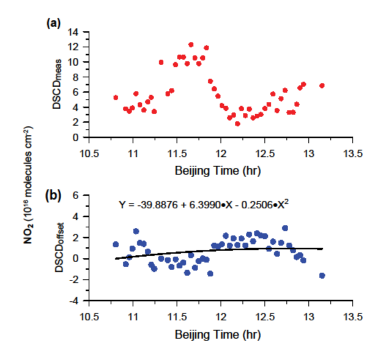


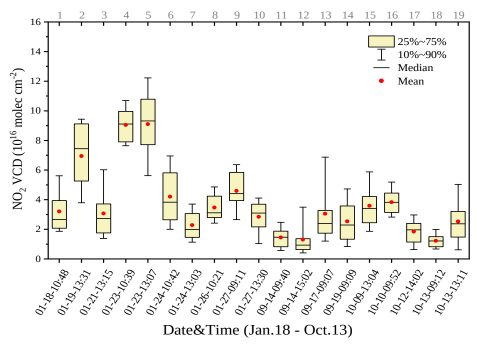
Fig. 1 Driving routes (red line) of the <u>car-Car MAX-DOAS</u> experiment <u>and distribution of four large</u> <u>point NO<sub>X</sub> emission sources within on-</u>the 6th Ring Rd of Beijing.



**Fig. 2** Examples of the NO<sub>2</sub> retrieval from two successive spectra measured (**a**) at a 30° elevation angle (with NO<sub>2</sub> differential slant column density (DSCD) of  $1.23 \times 10^{17}$  molecules cm<sup>-2</sup>) and (**b**) at a 90° elevation angle (with NO<sub>2</sub> DSCD of  $6.22 \times 10^{16}$  molecules cm<sup>-2</sup>) on January 18, 2014, at around 11:40 BJT.



**Fig. 3** Time series of the NO<sub>2</sub> (**a**) DSCD<sub>means</sub> (red dots) and (**b**) DSCD<sub>offset</sub> (black dots) (units of 10<sup>16</sup> molecules cm<sup>-2</sup>) for the 30° elevation angle of each sequence on January 18, 2014. The black curve represents a second-order polynomial fit from individual DSCD<sub>offset</sub> data points.



**Fig. 4** Time series of the tropospheric NO<sub>2</sub> vertical column density (VCD) for 19 circling journeys on the Sixth Ring Road of Beijing in January, September, and October, 2014. Lower (upper) error bars and yellow boxes are the 10th (90th) and 25th (75th) percentiles of the data of each journey, respectively. Hyphens inside the boxes are the medians, and red circles are the mean values. The numbers of each journey are labeled at the top axis. See Table 1 for detailed information about each journey.

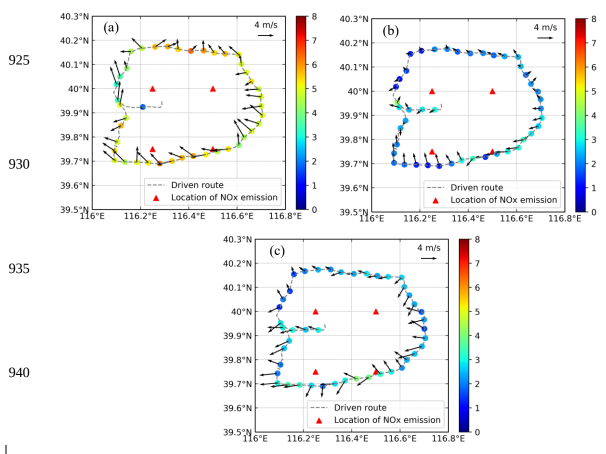


Fig. 5 Distributions of the monthly average NO<sub>2</sub> VCD (10<sup>16</sup>E16 molecules cm<sup>-2</sup>) on the 6th Ring Rd of Beijing in (a) January, (b) September, and (c) October, 2014.

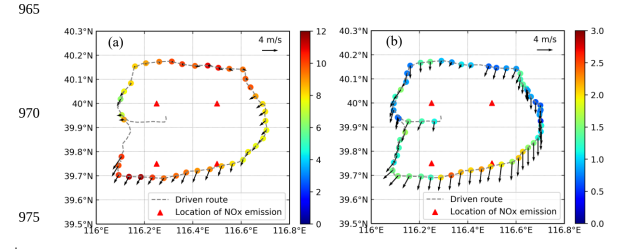
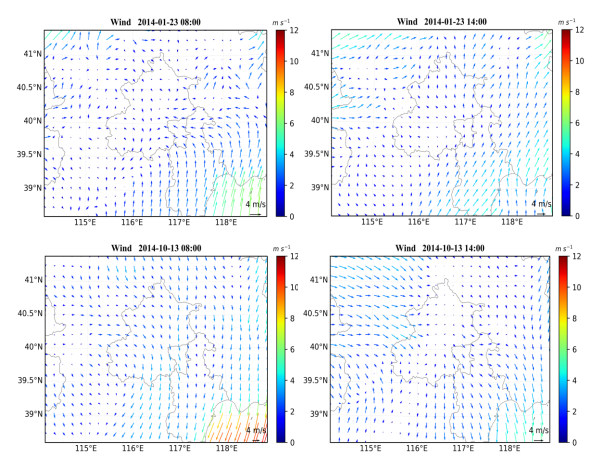
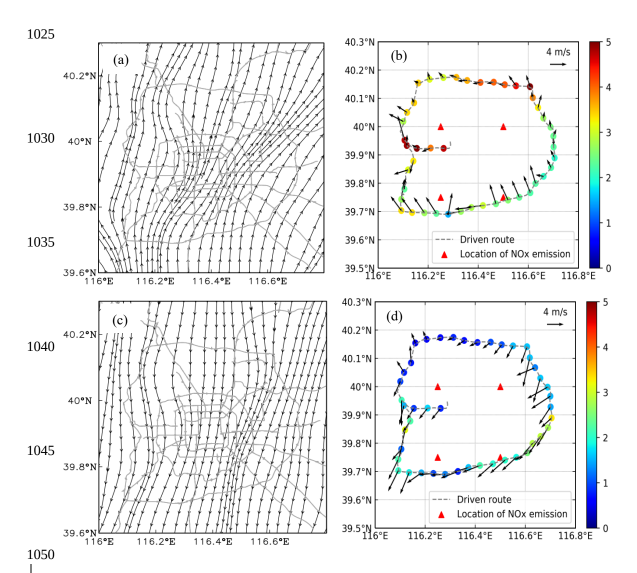


Fig. 6 Distributions of the maximum and minimum NO<sub>2</sub> VCD (<u>10<sup>16</sup>E16</u> molecules cm<sup>-2</sup>) on the 6th Ring Rd of Beijing on the morning of (a) January 23 and (b) October 13, 2014.



**Fig. 7** Wind fields in Beijing and the surrounding area from ECWRF at 08:00 (left column) and 14:00 (right column) BJT on January 23 and October 13, 2014.



**Fig. 8** Average wind stream and NO<sub>2</sub> VCD (E16 molecules cm<sup>-2</sup>) distributions under the three-two different types of wind field over Beijing: (a) south wind, (b) NO<sub>2</sub> VCD under south wind, (c) north wind, and (d) NO<sub>2</sub> VCD under north wind.

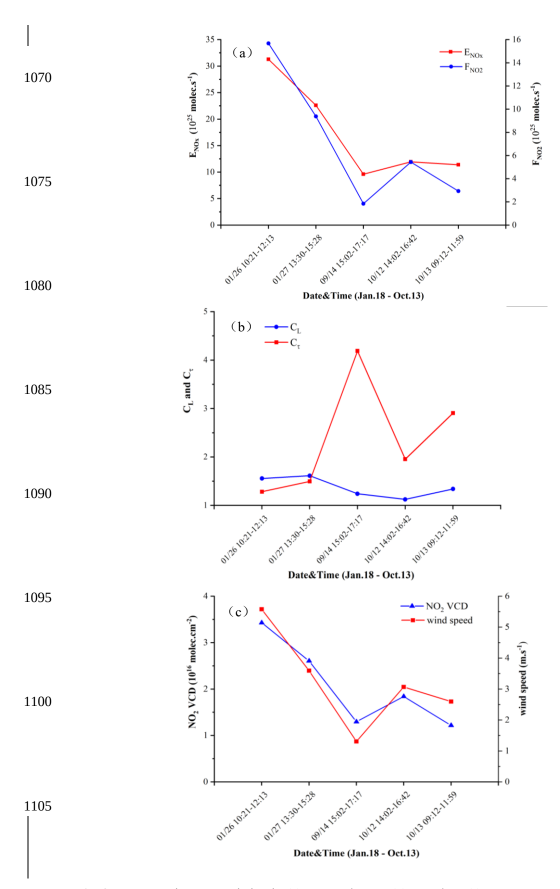


Fig. 9 Journey-to-journey variation in (a)  $F_{NO_2}$  and  $E_{NO_2}$ , (b)  $C_{\tau}$  and  $C_L$ , (c) NO<sub>2</sub> VCD and mean wind speed for five 10 circling journeys on the 6th Ring Rd of Beijing in January, September, and October,

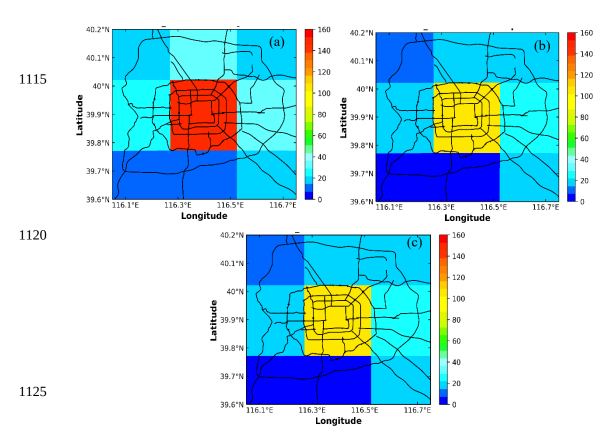


Fig. 10 Spatial distributions of  $NO_X$  emissions (mole km<sup>-2</sup> h<sup>-1</sup>) over Beijing based on the MEIC inventory in (a) January, (b) September, and (c) October 2012.



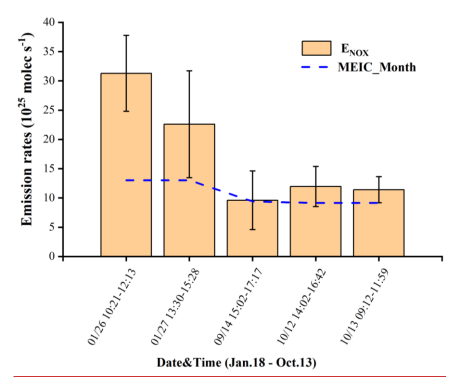
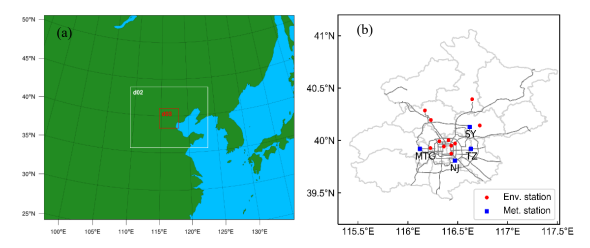
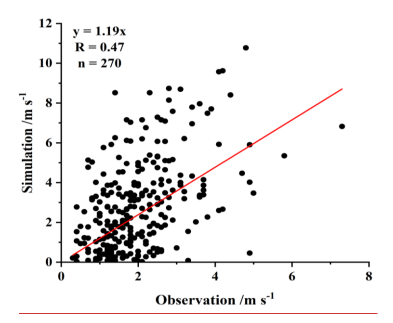


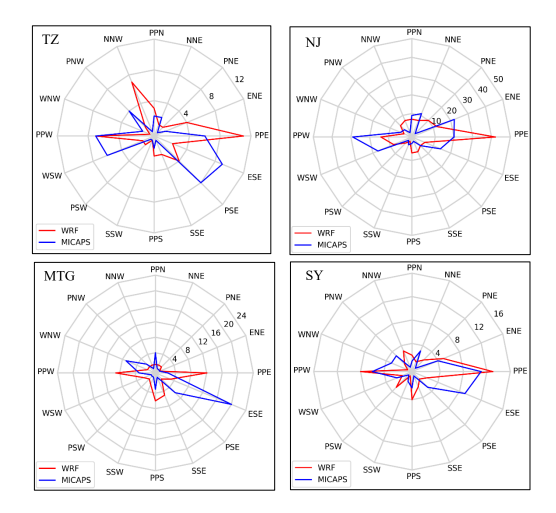
Fig. 11 Journey-to-journey variation in estimated  $E_{NO_x}$  and corresponding monthly emissions rates from the MEIC inventory (MEIC\_Month) within the 6th Ring Rd of Beijing in January, September, and October 2014. Error bars represent the uncertainties in estimated  $E_{NO_X}$ 



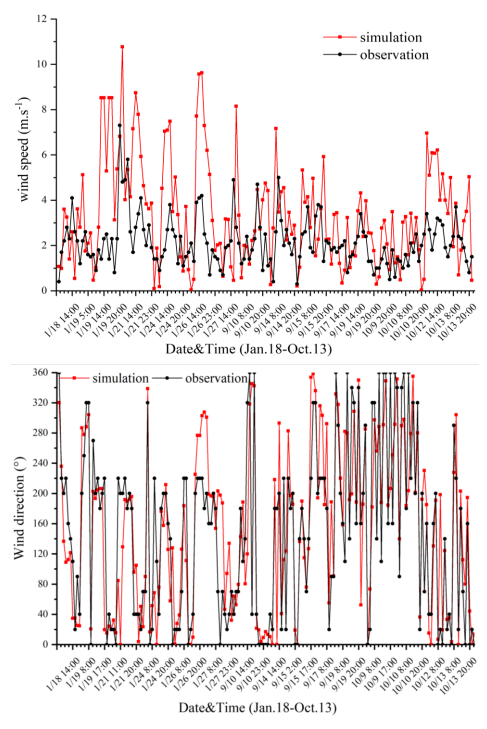
<u>Fig.S1</u> <u>Fig.1S</u> Triple-nested domains of (a) the LAPS-WRF-CMAQ model system and (b) the distribution of meteorological/environmental monitoring stations.



<u>Fig.S2</u> <u>Fig.2S</u> Scatterplot of simulated wind speed and observations at four stations in Beijing. <u>The standard deviation of the slope is 0.002.</u>



<u>Fig.S3Fig.3S</u> Wind rose of simulated wind direction and observations from MICAPS datasets at four stations in Beijing.



**Fig. S4** Time serial of simulated wind speed and direction, and observations during car MAX-DOAS experiments at four weather stations in Beijing.

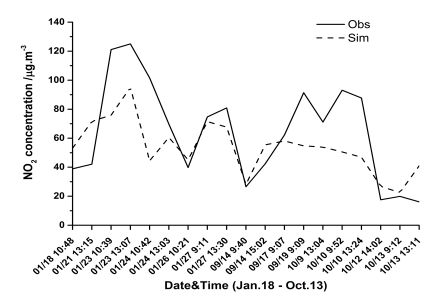


Fig.S5Fig.4S Time series of regional average simulation and in situ observation of NO<sub>2</sub> concentration at 12 stations in Beijing.

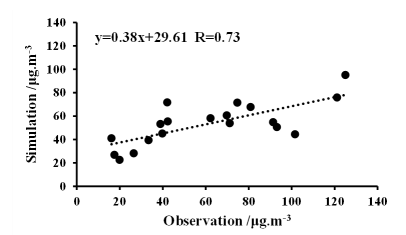


Fig.S6Fig.5S Scatter plot between regional average simulation and observation of NO<sub>2</sub> concentration at 12 stations in Beijing.

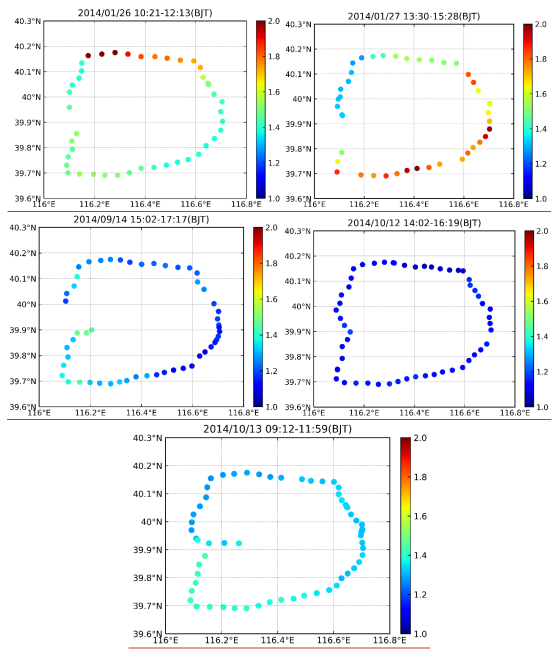


Fig. S7 Distributions of the ratio of NOx and NO<sub>2</sub> on the 6th Ring Rd of Beijing during five journeys.

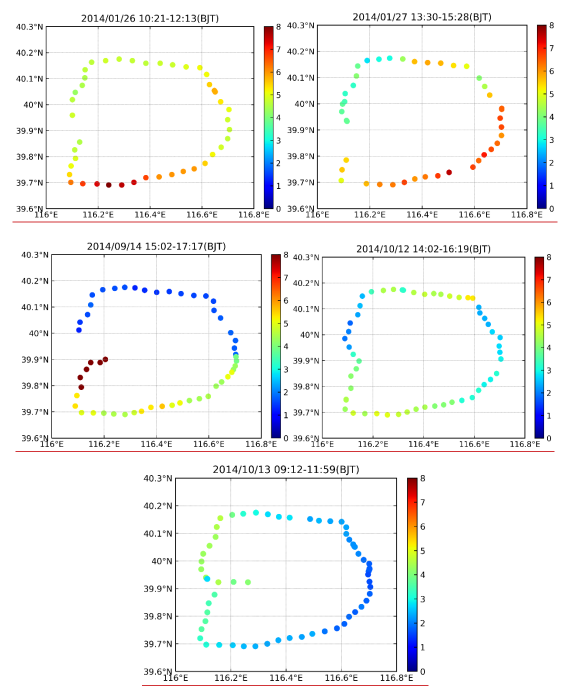


Fig. S8 Same to figure S7, except for the lifetime of NOx (h).

Table S1. Four types of monthly  $E_{NOX}$  from the MEIC inventory within the 6th Ring Rd of Beijing in January, September, and October 2012, and the ratio of  $E_{NOX}$  in Jan. to the average in Sep. and Oct.

	<u>industry</u>	power	<u>resident</u>	transport	<u>total</u>
<u>January</u>	5.78	<u>1.92</u>	1.39	<u>3.94</u>	13.02
<u>September</u>	4.06	<u>1.15</u>	0.25	3.93	9.40
October	4.03	0.93	0.26	3.93	9.15
Ratio	1.43	<u>1.84</u>	<u>5.43</u>	1.00	<u>1.40</u>