# Modeling study of the 2010 regional haze event in the North China Plain

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## 1 Abstract

2 The online coupled Weather Research and Forecasting-Chemistry (WRF-Chem) model was applied to simulate a haze event that happened in January 2010 in the North China Plain (NCP), 3 and was validated against various types of measurements. The evaluations indicate that WRF-4 Chem provides reliable simulations for the 2010 haze event in the NCP. This haze event was 5 mainly caused by high emissions of air pollutants in the NCP and stable weather conditions in 6 7 winter. Secondary inorganic aerosols also played an important role and cloud chemistry had important contributions. Air pollutants outside Beijing contributed about 64.5% to the PM<sub>2.5</sub> 8 9 levels in Beijing during this haze event, and most of them are from south Hebei, Tianjin city, 10 Shandong and Henan provinces. In addition, aerosol feedback has important impacts on surface 11 temperature, Relative Humidity (RH) and wind speeds, and these meteorological variables affect aerosol distribution and formation in turn. In Shijiazhuang, Planetary Boundary Layer (PBL) 12 decreased about 278.2m and  $PM_{2.5}$  increased more than  $20\mu g/m^3$  due to aerosol feedback. 13 Feedbacks associated to Black Carbon (BC) account for about 65.7 % of the PM<sub>2.5</sub> increases and 14 15 59.9% of the PBL decreases in Shijiazhuang, indicating more attention should be paid to BC from both air pollution control and climate change perspectives. This contribution decreased 16 17 from about 60% to 50% after deceasing BC emissions by 50% and the uncertainty can be additionally reduced by improving the model performance in simulating sulfate and OC. 18 19 20 21 22 23 24

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#### 1 1 Introduction

2 The North China Plain (NCP) is one of the most densely populated areas in the world and it has been the Chinese center of culture and politics since early times. Beijing, the capital of China, 3 4 Tianjin, Shijiazhuang and other big cities with active economic developments are located in the NCP. This region is experiencing heavy haze pollution with record-breaking high concentrations 5 of particulate matters (L. T. Wang et al., 2014). Haze is defined as an air pollution phenomenon 6 7 where horizontal visibility is less than 10 km caused by aerosol particles, such as dust and Black Carbon (BC), suspended in the atmosphere (Tao et al., 2012). Its formation is highly related to 8 meteorological conditions, emissions of pollutants and gas-to-particle conversion (Sun et al., 9 10 2006; Watson, 2002). Haze has attracted much attention for its adverse impacts on visibility and 11 human health. During haze periods, reduced visibility affects land, sea and air traffic safety and the fine particles can directly enter the human body and adhere to lungs to cause respiratory and 12 13 cardiovascular diseases (Liu et al., 2013). Moreover, haze affects climate and ecosystems via aerosol-cloud-radiation interactions (Sun, et al., 2006; Liu et al., 2013). 14

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Because haze influences visibility, human health and climate (Gao et al., 2015), numerous 16 17 studies have used multiple methods to investigate physical, chemical and seasonal characteristics of aerosols during haze. The increase of secondary inorganic aerosols is considered to be an 18 attribute of the haze pollution in east China (Tan et al., 2009; Zhao et al., 2013). Tan et al. (2009) 19 20 studied the characteristics of aerosols in non-haze and haze days in Guangzhou, China and found that secondary pollutants (OC, SO4<sup>2-</sup>, NO<sup>3-</sup> and NH<sup>4+</sup>) were the major components of haze 21 aerosols and they showed a remarkable increase from non-haze to haze days. Similar conclusions 22 23 were drawn by Zhao et al. (2013) after studying the chemical characteristics of haze aerosols in the NCP. Secondary Organic Aerosol (SOA) formation can also be significant during haze (Tan 24 et al., 2009; Zhao et al., 2013). Studies of aerosol optical properties show that fine-mode aerosols 25 26 were dominant during haze (Yu et al., 2011; Li et al., 2013). In addition, contributions of diverse 27 factors to haze formation, such as biomass burning and regional transport, have been investigated. 28 Chen et al. (2007) used MM5-CMAQ to reproduce the haze pollution in September 2004 in the Pearl Region Delta (PRD) region and discovered that sea-land breeze played an important role. 29 Wang et al. (2009) discovered that almost 30-90 percent of the organics during the haze 30

1 happened in June 2007 in Nanjing were from wheat straw burning. Cheng et al. (2014)

- 2 concluded that biomass burning could cause haze issues and they found biomass burning
- 3 contributed 37% of PM<sub>2.5</sub>, 70% of Organic Carbon (OC) and 61% of Elemental Carbon (EC)
- 4 based upon both modeling and measurement results of case study in summer 2011 in the

5 Yangtze River Delta (YRD) region. These biomass burning events mainly occurred in summer

6 and autumn in east and south China (Cheng et al., 2013, 2014; Li et al., 2010; Wang et al., 2007,

7 2009). To evaluate regional contributors to the haze in southern Hebei, Wang et al. (2012)

8 simulated the time period from 2001 to 2010 and concluded that Shanxi province and the

9 northern Hebei were two major contributors, and winter was the worst season, followed by

10 autumn and summer.

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12 X. Han et al. (2014) pointed out that the haze formation mechanism in winter in Beijing was different from that in summer and mass concentrations of PM<sub>2.5</sub> in winter were relatively higher 13 14 and the compositions were different than in summer. The extreme winter haze in the NCP has attracted enormous scientific interests. It has been found the stagnant meteorological conditions 15 16 (weak surface wind speed and low Planetary Boundary Layer (PBL) height) and secondary 17 aerosol formation are the main causes of winter haze formation (S. Han et al., 2014; He et al., 2014b; K. Huang et al., 2014; Sun et al., 2014; Wang et al., 2014a; Zhao et al., 2013; Zheng et al., 18 2014, 2015). Other causes proposed include high local emissions (He et al., 2014b; Zheng et al., 19 20 2014), enhanced coal combustion in winter (K. Huang et al., 2014; Sun et al., 2014), 21 heterogeneous chemistry (He et al., 2014a; X. Huang et al., 2014; Quan et al., 2014; Wang et al., 2014a, b; Zheng et al., 2014, 2015) and regional transport (Tao et al., 2014; Sun et al., 2014; L. T. 22 Wang et al., 2014; Z. Wang et al., 2014; Zheng et al., 2014). It was also pointed out that fog 23 processing (K. Huang et al., 2014), aerosol-radiation interactions (J. Wang et al., 2014; Z. Wang 24 25 et al., 2014; B. Zhang et al., 2015) and nucleation events (Guo et al., 2014) may play important roles in winter haze formation. 26

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The complex haze formation mechanisms need further studies. Li et al. (2015) emphasized that

regional transport of  $PM_{2.5}$  is a major cause of severe haze in Beijing, but R. Zhang et al. (2015)

30 pointed out that the evidence provided by Li et al. (2015) is insufficient and regional transport

should be evaluated using chemical transport models. Furthermore, the contribution of aerosol 1 2 feedbacks to PM<sub>2.5</sub> levels remains unquantified. Therefore, the roles of regional transport and 3 aerosol-radiation interactions in haze events need to be better understood. In this study, the 4 online coupled model WRF-Chem, which is capable of simulating aerosols' effects on meteorology and climate, is used to reproduce the severe haze event that happened in the NCP 5 from 16 to 19 January 2010. During this haze event, the highest hourly PM<sub>2.5</sub> concentration 6 reached 445.6 and 318.1µg/m<sup>3</sup> in Beijing and Tianjin and the areas with low visibility covered 7 most eastern China regions (Zhao et al., 2013). In this study, we address the following important 8 questions: (1) what is the performance of the model configurations in representing the 9 meteorological variables, and the physical and chemical characteristics of the aerosols during the 10 selected study period?; (2) How does the haze build up and dissipate?; (3) How do the chemical 11 species of PM<sub>2.5</sub> change during haze period?; (4) Does regional transport play an import role in 12 the 2010 haze event in Beijing?; (5) What is the contribution of aerosol feedback mechanisms to 13  $PM_{2.5}$  levels during the haze event?; and (6) What is the role of BC absorption in the feedback 14 mechanism? In Sect. 2, we describe the model we use and model configuration, including 15 16 emissions and used parameterization schemes. In Sect. 3, surface meteorological, chemical observations, atmospheric sounding products, as well as remote sensing products are used to 17 18 evaluate the model performance. In Sect. 4, questions from (2) to (6) are answered in detail. Conclusions are provided in the Sect. 5. 19

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#### 21 2 Model description and configuration

22 The WRF-Chem model version 3.5.1 was employed to simulate the 2010 haze event in the NCP region and aerosol-radiation interactions were included (Chapman et al., 2008; Fast et al., 2006). 23 Domain settings are the same as those of Jing-Jin-Ji modeled area of Yu et al. (2012). Three 24 domains with two-way nesting were used and grid resolutions were 81km  $\times 81$ km (domain 1), 25 26 27km  $\times 27$ km (domain 2) and 9km  $\times 9$ km (domain 3) (see supporting information Figure S1). The number of vertical grids used was 27 and the number of horizontal grids was 81×57, 49×49, 27 28 and 55×55, respectively. The first domain covers most areas of the East Asia region, including China, Korea, Japan and Mongolia. Beijing was set to be the center of the innermost nested 29 domain. The chemical and aerosol mechanism used was gas-phase chemical mechanism CBMZ 30

(Zaveri and Peters, 1999) coupled with the 8-bin sectional MOSAIC model with aqueous
chemistry (Zaveri et al., 2008). MOSAIC treats all the important aerosol species, including
sulfate, nitrate, chloride, ammonium, sodium, BC, primary organic mass, liquid water and other
inorganic mass (Zaveri et al., 2008). Some of the physics configuration options include Lin
cloud-microphysics (Lin et al., 1983), RRTM long wave radiation (Mlawer et al., 1997),
Goddard short wave radiation (Chou et al., 1998), Noah land surface model, and the Yonsei
University planetary boundary layer parameterization (Hong et al., 2006).

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Emissions are key factors in the accuracy of air quality modeling results. The monthly 2010 9 10 Multi-resolution Emission Inventory for China (MEIC) (http://www.meicmodel.org/) was used 11 as the anthropogenic emissions. This inventory includes emissions of sulfur dioxide (SO<sub>2</sub>), nitrogen oxides (NO<sub>x</sub>), Carbon Monoxide (CO), non-methane volatile organic compounds 12 (NMVOC), NH<sub>3</sub>, BC, organic carbon (OC), PM<sub>2.5</sub>, PM<sub>10</sub>, and carbon dioxide (CO<sub>2</sub>) by several 13 sectors (power generation, industry, residential, transportation, etc.). Biogenic emissions were 14 calculated on an online way by the MEGAN model (Guenther et al., 2006). Meteorological 15 initial and boundary conditions were obtained from the National Centers for Environmental 16 Prediction (NCEP) Final Analysis (FNL) data set. Chemical initial and boundary conditions were 17 taken from MOZART-4 forecasts (Emmons et al., 2010). The period from 11 to 24 January 2010 18 was chosen as the modeling period, covering the 2010 NCP haze period (from 16 to 19 January 19 20 2010). To overcome the impacts of initial conditions, three days were simulated and considered as spin-up time. 21

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#### 23 3 Model Evaluation

## 24 **3.1 Observation data sets and evaluation metrics**

25 Model evaluation was conducted in terms of both temporal variation and spatial distribution.

26 Table 1 gives a summary of the observation data and variables used in the model evaluation. The

27 meteorological variables, including 2 meter temperature (T2), 2 meter relative humidity (RH2)

and 10 meter wind speed (WS10), at four stations (Beijing, Tianjin, Baoding and Chengde) were

used. Surface concentrations of PM<sub>2.5</sub>, NO<sub>2</sub>, SO<sub>2</sub> at three sites (Beijing, Tianjin and Xianghe, 1 2 shown in Figure S1), and Aerosol Optical Depth (AOD) at four sites (Beijing city, Beijing forest, 3 Baoding city, Cangzhou city) were also used in the evaluation against measurements. PM<sub>2.5</sub> and AOD are typical variables to represent severity of haze pollution. To evaluate how model 4 performs in simulating horizontal and vertical distributions of meteorological and chemical 5 variables, soundings of temperature and RH at Beijing, and AODs derived from CALIPSO were 6 used in this study. The statistical metrics calculated include correlation coefficient R, mean bias 7 (MB), mean error (ME), the root mean square error (RMSE), the normalized mean bias (NMB), 8 the normalized mean error (NME), the mean fractional bias (MFB) and the mean fractional error 9 (MFE). The definitions of these metrics can be found in Morris et al. (2005) and Willmott and 10 Matsuura (2005). 11

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## 13 **3.2 Meteorology simulations**

Figure 1 shows the temporal variations of simulated and observed 24-h average temperature (a-14 15 d), relative humidity (e-h) and wind speed (i-l) at Beijing, Tianjin, Baoding and Chengde stations. These observations were collected from the China Meteorological Data Sharing Service System 16 (CMDSSS) data set. From normal days to haze days (gray shaded), temperature and relative 17 humidity increased and wind speeds decreased. Generally, the variations of surface temperature, 18 19 RH and wind speeds are captured by model, although overestimations of wind speed occur at the 20 Chengde station throughout the whole period. Model mean, observation mean, MB, ME and RMSE were calculated and summarized in Table 2. The MB and RMSE for surface temperature 21 vary from -2.0 to 2.0 K and from 1.5 to 3.2 K, respectively. The model underestimates 22 temperature at Beijing, Tianjin and Baoding stations, and overestimates temperature at the 23 24 Chengde station. RH agrees well with observations, with MB varying from -4.4% to 8.1% and 25 RMSE varying from 6.4% to 11.1%. The magnitudes of MB and RMSE are comparable with 26 those of Wang et al. (2014b). The model shows good performance in simulating wind speed, with RMSE ranging from 1.1 to 1.6 m/s at Beijing, Tianjin and Baoding stations, below the level 27 of "good" model performance criteria for wind speed prediction proposed by Emery et al. (2001). 28 29 Wind speeds at the Chengde station were overestimated, with RMSE larger than the proposed 30 criteria (2m/s).

2 Figure 2 compares simulated and observed vertical temperature profiles at 0800 and 2000 (CST) 3 from January 15 to January 20 at Beijing city. These atmospheric sounding data are from the NCAR Earth observing laboratory atmospheric sounding data set. The model captures the 4 vertical profiles of temperature well. Obvious strong temperature inversions existed during the 5 haze period (from 01/16 08:00 to 01/19 20:00) and the lapse rate during this period was about 5-6 7 15°C/km, indicating unfavorable conditions for diffusion of pollutants. The model captures the general vertical profiles of RH, although the performance is not as good as for temperature (see 8 9 supporting information Figure S2).

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#### **3.3 Chemical simulations**

12 Figure 3(d-f) shows variations of simulated and observed hourly PM<sub>2.5</sub>, NO<sub>2</sub> and CO at the SDZ station. The haze event started from 16 January with rapid increase of PM<sub>2.5</sub>, NO<sub>2</sub>, and CO 13 concentrations and ended on 20 January. The relationships between meteorological condition and 14 pollution levels are clearly shown. Both the observation and the model show that temperature 15 16 and relative humidity increase, wind speeds are low, and pollution levels build up (Figure 3). The 17 magnitudes and trends over time of the simulated PM<sub>2.5</sub>, NO<sub>2</sub> and CO are generally consistent with measurements, although overestimation of PM2.5 and underestimations of NO2 and CO exist 18 19 during the haze days. Figure 4 shows the temporal variations of the simulated and observed PM<sub>2.5</sub>, NO<sub>2</sub> and SO<sub>2</sub> at Beijing (a-c), Tianjin (d-f) and Xianghe (g-i) stations. The observations 20 21 and the model predictions show that the buildups of pollution during the haze event were similar 22 at these three sites, occurring over a large geographical region at the same time.  $SO_2$  was 23 overestimated in Beijing, but other simulations agree well with observations, especially for PM<sub>2.5</sub>. Observation mean, model mean, MB, ME, NMB, NME, MFB, and MFE were calculated for 24-24 25 h average simulated and observed PM<sub>2.5</sub> at these three stations and summarized in Table 3. As shown in Table 3, the model underestimates PM<sub>2.5</sub> concentrations at all stations. NMBs for PM<sub>2.5</sub> 26 27 are -8.5%, -26.9% and -39.1% at Beijing, Tianjin and Xianghe, respectively. MFBs at these three stations range from -21.8% to 0.4% and MFEs range from 26.3% to 50.7%. They are all within 28 29 the criteria proposed by Boylan et al. (2006) that model performance is "satisfactory" when MFB is within  $\pm 60\%$  and MFE is below 75%. Although the model performance for PM<sub>2.5</sub> is 30

satisfactory, biases still exist, especially during severe haze days. Reasons for the biases might be errors in meteorological variables, large uncertainties of emission inventory, effects of horizontal and vertical resolutions, and incomplete treatments of atmospheric chemistry. Many atmospheric chemistry reactions have been and are being proposed for PM formation in winter haze. For example, He et al. (2014a) proposed that mineral dust and NO<sub>x</sub> could promote the formation of sulfate in heavy pollution days. The sensitivity of the simulations to some of these factors will be discussed in future studies.

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#### 9 **3.4 Simulations of optical properties**

In WRF-Chem, aerosol optical properties are calculated at four specific wavelengths, 300nm,
400nm, 600nm, and 1000nm, while AOD observations from CSHNET, CALIPSO are not at
these four wavelengths. To evaluate model performance of simulating AOD, we derived AOD at
observation wavelengths based on Angstrom exponent relation (Schuster et al., 2006). In severe
haze days, AOD could not be retrieved, so the observerd AOD data in some days are missing.
Model agrees very well with the CSHNET AOD observations at all four stations (supporting
informaiton Figure S3).

17 CALIPSO retrievals provide vertical curtains of aerosol and clouds. Figure 5 shows paths of the CALIPSO satellite, simulated extinction coefficient and observed plume top, and simulated 18 AOD and CALIPSO retrieved AOD at 532nm at three moments: January 14 12:00(CST) (a-c), 19 January 21 02:00(CST) (d-f), and January 21 12:00(CST) (g-i), respectively. There were no 20 21 retrievals in the NCP during haze days. Figure 5(a), (d) and (g) show that the CALIPSO satellite 22 passed over the NCP region at these three moments. Simulated extinction coefficient matches 23 observed plume top (Figure 5(b), (e) and (h)), indicating that the model captures the vertical distributions of aerosols. The model also has good performance in simulating AOD at 532nm, 24 25 although underestimations happen around latitude 36°N (Figure 5(c), (f) and (i)).

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27 The model is shown to be capable of simulating the major meteorological and chemical

evolution of this haze event. As spatial and vertical profiles of the haze period are incomplete or

29 missing in the satellite retrievals and ground stations only provide point estimates, we can use the

model to understand the haze spatial, vertical and temporal evolution, as discussed in thefollowing sections.

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#### 4 4 Results and Discussions

#### 5 4.1 Meteorological conditions and evolution of air pollutants

6 The evolution of the spatial distributions of the haze event is shown in Figure 6, where the horizontal distributions of PM2.5 and wind vectors are plotted every 12 hours from January 14 7 00:00 to January 21 00:00. In the second plot (January 14 12:00), air flows converged at the NCP 8 surface areas, resulting in a small increase of PM2.5 concentration. From January 14 00:00 to 9 January 16 00:00,  $PM_{2.5}$  concentration over the NCP was generally below  $120\mu g/m^3$ . From 10 January 16 to January 18, Beijing and surrounding areas were controlled by a weak high pressure 11 system (Zhao et al., 2013). During this period, large amounts of emissions in the NCP 12 13 accumulated and the persistent southerly winds brought some air pollutants northward to Beijing and southern Hebei areas. The weak high pressure system was replaced by a low pressure system 14 15 that lasted until January 20, and this weather condition was not conductive for dispersion of air pollutants (Zhao et al., 2013). On January 19, the NCP haze was in the worst state, with PM<sub>2.5</sub> 16 concentrations above  $350 \mu \text{g/m}^3$  in south NCP. From January 20, strong northerly winds 17 dispersed the accumulated air pollutants and the haze ended. 18

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To illustrate the vertical structure of the haze, vertical cross sections of PM<sub>2.5</sub> concentration and clouds are presented in Figure 7. The cross section diagonally cuts the region with the lower left corner of 34N, 110E to the upper corner at 44N, 122E (see supporting information Figure S4). There were two highly polluted points (around latitudes 35 and 39) and they started merging as one from January 18 12:00 (Figure 7). At that time, southerly winds blew air pollutants northwards (Figure 6) and the polluted region was expanded. On January 19, there were fog and/or clouds near the surface and the impacts of fog and/or clouds will be discussed in Sect. 4.2.

Further details of the evolution of the haze are shown in the temporal variations of  $PM_{2.5}$ 1 2 concentrations in Shijiazhuang, Tianjin and Chengde (marked in Figure S1) in Figure 8. All three 3 sites show similar temporal variations. Around noon of January 15, PM<sub>2.5</sub> concentrations in Shijiazhuang, Chengde and Beijing increased at nearly the same time, labeled by red arrow in 4 Figure 8. Air pollutants started accumulating when the NCP was controlled by the weak and 5 stable weather conditions. Compared to Shijiazhuang and Beijing, the capital city of Hebei 6 7 province and the capital of China, PM<sub>2.5</sub> concentrations in Chengde were lower (Figure 8). It was estimated that there are more than 8100 coal-fired boilers and industrial kilns in Shijiazhuang 8 9 city (Peng et al., 2002), resulting in high intensity of emissions in Shijiazhuang. On January 20, Chengde was the first to show sharp decrease of PM<sub>2.5</sub> concentrations, followed by Beijing and 10 Shijiazhuang, corresponding to the northerly wind impacts discussed above. 11

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13 To better understand the relationships between meteorological factors and pollution levels, time series of different pairs of variables are shown in Figure 9. CO shows very high correlation with 14  $PM_{2.5}$  (Figure 9(a)), which is consistent with the observation and modeling results in Santiago, 15 16 Chile (Perez et al., 2004; Saide et al., 2011), and shows the large contribution of primary sources (including gaseous precursors) to PM<sub>2.5</sub>. Secondary aerosol formation also plays a role as PM<sub>2.5</sub> 17 peaks on the 19<sup>th</sup> while CO peaks on the 18<sup>th</sup>. RH and wind speed are two important factors 18 affecting the concentrations of aerosols. RH has similar variations as PM2.5 concentration (shown 19 20 in Figure 9(a) and 5(b)). The NCP is close to the sea and under the slow southerly flows, 21 temperature and RH increase along with PM<sub>2.5</sub>. During the haze event, RH values were generally above 40% and wind speeds were below 2 m/s (Figure 9(b)). Low wind speed is unfavorable for 22 the dilution of air pollutants and high RH would accelerate the formation of secondary species, 23 such as sulfate and nitrate, to aggravate the pollution level (Sun et al., 2006). NO<sub>X</sub> concentrations 24 show similar variations as PM<sub>2.5</sub>, indicating the buildup of concentrations during the wind speed 25 stagnation. Ozone shows lower concentrations during haze event (Figure 9(c)) because high 26 aerosol loadings produce low photochemical activity due to decrease in UV radiation. The 27 concentrations have an inverse relationship with PBL Height (PBLH) as shown in Figure 9(d). 28 Diurnal maximums of PBLHs were mostly below 400m and PBL collapsed at night during the 29 30 haze event, indicating aerosols were trapped near the surface. On January 21 and 22, PBLHs 31 were between 800 and 1000 meters, which helped diffuse and dilute the air pollutants, resulting

in a decrease in concentration. The relationships between these variables are further discussed
 with respect to the influences of aerosol feedback mechanism in Sect. 4.4.

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4 Figure 10 shows the temporal variations of vertical profiles of simulated PM<sub>2.5</sub> concentration (a), temperature (b), RH (c) and wind speeds (d) at the Beijing site. PM<sub>2.5</sub> was accumulated below 5 500m and concentrations reached peak values around January 18 00:00 (Figure 10(a)), when a 6 strong temperature inversion happened over Beijing (Figure 10(b)), which inhibited vertical 7 8 atmospheric mixing. A strong temperature inversion also happened on January 19 (Figure 10(b)). 9 From January 16 to 19, RH was mostly higher than 50% and reached a peak on the night of 10 January 19 (Figure 10(c)). As a result, air pollutants released into the atmosphere were trapped in 11 the moist atmosphere and accumulated as near surface horizontal winds were very weak (below 12 1.5m/s) during the haze period (Figure 10(d)). As mentioned above, the high RH enhances the formation of secondary species, which will be discussed in the following section. 13

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#### 15 4.2 Evolution of aerosol composition during haze

16 As shown above, during haze events, aerosols build up due to low mixing heights and low wind 17 speeds. An important question is what is the role of secondary aerosol formation during such 18 events? Previous measurement studies have found that the increase of secondary inorganic 19 pollutants could be considered as a common property of haze pollution in East China (Zhao et al., 2013). The observed and simulated chemical species of  $PM_{2.5}$  in Beijing are shown in Figure 20 11(a) and 11(b), respectively. Observed secondary inorganic aerosols (SIA) (NH<sub>4</sub><sup>+</sup>, SO<sub>4</sub><sup>2-</sup>, NO<sup>3-</sup>) 21 22 increased significantly during the haze episode and accounted for 37.7% of PM<sub>2.5</sub> mass 23 concentration (Zhao et al., 2013). Primary OC, BC, sulfate, nitrate and ammonium accounted for 24 the major parts of the simulated  $PM_{2.5}$  during haze. Table 4 summarizes the mean concentrations of primary aerosols (primary OC and BC) and SIA (NH<sub>4</sub><sup>+</sup>, SO<sub>4</sub><sup>2-</sup>, NO<sup>3-</sup>) in non-haze days, and 25 in the most serious haze day. The primary aerosols increased by a factor of 4.0 from non-haze 26 days to haze days. The SIA also increased from non-haze days to haze days, which agrees with 27 the observation (Tan et al., 2009; Zhao et al., 2013). The SIA increased by a factor of 7.6 from 28 non-haze days to haze days. The increasing factors for observed primary aerosols and SIA are 29

2.9 and 6.9, which are close to those factors from simulations. However, the amounts of sulfate 1 2 are underestimated by WRF-Chem, compared with the observation in Figure 11(a) from Zhao et 3 al. (2013). Tuccella et al. (2012) pointed out that the underestimation of simulated sulfate could be due to the underestimation of  $SO_2$  gas phase oxidation, errors in nighttime boundary layer 4 height predicted by WRF-Chem, and/or the uncertainties in aqueous-phase chemistry. It could 5 also be caused by the missing heterogeneous sulfate formation in current model (He et al., 2014a; 6 7 Wang et al., 2014d; Zheng et al., 2014a). As discussed earlier, the SO<sub>2</sub> gas phase concentrations at this site were overestimated. Adding reaction pathways to produce sulfate aerosol would 8 improve both the predictions of sulfate (increase) and SO<sub>2</sub> (decrease) (He et al., 2014a; Wang et 9 al., 2014c; Zheng et al., 2014a). 10

11

12 We investigated the role of aqueous phase chemistry during the haze event. The aqueous phase pathway can reach a level of over 50  $\mu$ g/m<sup>3</sup> around the Beijing area, accounting for a significant 13 part (about 14.3%) of total PM<sub>2.5</sub> concentration (see supporting information Figure S5). As 14 shown in Figure 7, fog/clouds existed near the surface on January 19 and this corresponds to the 15 16 PM<sub>2.5</sub> difference on that day due to aqueous phase pathway. The sulfate production in aqueous phase may be higher than shown in this study after adding missing aqueous-phase reactions. The 17 impacts of heterogeneous reactions on sulfate production will be investigated in future studies. 18 As shown in Figure 11(a) and 7(b), the model underestimates OC. To evaluate the formation of 19 Secondary Organic Aerosol (SOA) during the haze event, the RADM2/MADE-SORGAM model 20 was used. The CBMZ/MOSAIC version used is not capable of simulating SOA formation 21 because CBMZ was hard-wired with a numerical solver in WRF-Chem and thus SOA 22 23 condensable precursors could not be directly added into it (Zhang et al., 2012). RADM2 is an upgrade of RADM1 and it gives more realistic predictions of H<sub>2</sub>O<sub>2</sub> (Stockwell et al., 1990), and 24 25 Schell et al., (2001) incorporated SOA into the Modal Aerosol Dynamics Model for Europe 26 (MADE) (Ackermann et al., 1998) by means of the Secondary Organic Aerosol Model (SORGAM). SORGAM treats anthropogenic and biogenic aerosol precursors separately and 27 28 eight SOA compounds are considered, of which four are anthropogenic and the other four are 29 biogenic (Schell et al., 2001). Predicted Anthropogenic SOA (ASOA), biogenic SOA (BSOA) 30 and Primary Organic Aerosol (POA) in Beijing are shown in Figure 11(c). SOA indeed shows a

marked increase from non-haze days to haze days, but the amount of SOA is very small 1 2 compared with POA. The highest SOA concentrations in China are usually found in summer and 3 in Central China (Jiang et al., 2012). In addition, almost all of the simulated SOA are ASOA. 4 Jiang et al. (2012) also concluded that in winter, the fractions of ASOA are larger than 90% in north China. Biogenic emissions are usually controlled by solar radiation and temperature, and 5 solar radiation is weaker and temperature is lower in winter compared with summer. Moreover, 6 7 the high isoprene, API (a-pinene and other cyclic terpenes with one double bond) and LIM (limonene and other cyclic diene terpenes) emissions are located below 30 °N and in Northeast 8 China (Jiang et al., 2012), not in the NCP, so the SOA concentrations are not high in this winter 9 haze event period in the NCP. As shown in Table 4, the mean SOA concentration in non-haze 10 days is  $0.15\mu g/m^3$  and in the most serious haze day is  $8.2\mu g/m^3$ . The factor increase of SOA from 11 non-haze days to haze day is 8.2, which is lower than that of primary aerosols and much lower 12 than that of SIA. The SOA formation in winter has not been well studied and it might be 13 underestimated by the model as it could have missing pathways to SOA formation. Further work 14 is needed to improve the underestimation of SOA formation in the winter. 15

16

#### 17 4.3 Impacts of surrounding areas on haze in Beijing

Previous studies found that both local emissions and regional transport have significant
contributions to the high fine particle levels in Beijing (Yang et al., 2011). A sensitivity
simulation was conducted to quantify the contributions of surrounding areas to haze in Beijing,
when Beijing local emissions were turned off. The ratio of PM<sub>2.5</sub> in Beijing when Beijing
emissions are turned off to PM<sub>2.5</sub> in Beijing when Beijing emissions are on represents the nonlocal contributions. It can reach above 80% during haze (see supporting information Figure S6)
and the average contribution is about 65% from January 16 to January 19.

To figure out the dominant transport paths, FLEXPART-WRF (Stohl et al., 1998; Fast and
Easter, 2006) was used to generate 72-hour backward dispersions around the Beijing area. 50000
particles were released backwards from a box (1 degree×1 degree×400m), the center of which is
Beijing urban area, from January 19 00:00. The number concentrations of particles were plotted
at 6 hours before, 12 hours before, 24 hours before and 48 hours before the released time (Figure
12). For 12 hours, Beijing was influenced by sources to the south, including sources from south

1 Hebei, Tianjin and Shandong. For 2 days, more sources contributed to the haze buildup in

2 Beijing, including sources from Henan and Inner Mongolia. A number of coal mines are located

3 in Hebei, Shandong and Henan provinces and Inner Mongolia areas have high emissions of

4 primary aerosols.

5

# 6 **4.4 The impact of aerosol feedback**

7 Aerosols affect weather and climate through many pathways, including reducing downward solar radiation through absorption and scattering (direct effect), changing temperature, wind 8 9 speed, RH and atmospheric stability due to absorption by absorbing aerosols (semi-direct effect), serving as cloud condensation nuclei (CCN) and thus impacting optical properties of clouds (first 10 indirect effect), and affecting cloud coverage, lifetime of clouds and precipitation (second 11 12 indirect effect) (Zhang et al., 2010; Forkel et al., 2012). The feedback mechanisms are complex and many aspects of them are not well understood. Although previous studies have investigated 13 aerosol-radiation-meteorology interactions (Y. Zhang et al., 2010; R. Forkel et al., 2012), the 14 studies on short time scale events with high aerosol loadings, such as haze events, are limited. 15 This section focuses on evaluating the impacts of aerosol feedback mechanism on meteorology 16 17 and air quality. The feedback discussed in this paper only includes aerosols' direct and semidirect effects. 18

19

# 20 4.4.1 Impact of feedback on meteorology and PM<sub>2.5</sub> distribution

Figure 13(a) shows the observed daily maximum surface solar radiation and simulated surface 21 solar radiation for the with feedback (WF) and without feedback (NF) scenarios in Beijing. 22 Simulated daily maximum surface shortwave radiation values for the NF scenario are higher than 23 24 observations and the overestimations are reduced by implementing aerosol feedback (Figure 13(a)). For the NF case, the correlation coefficient R between simulated and observed daily 25 maximum surface shortwave radiation is 0.84 in Beijing; for the WF scenario, the correlation 26 27 coefficient increased to R=0.93, and the haze reduced the shortwave radiation values by 30 to 80%. 28

1 The changes in radiation have impacts on the environment. Simulated PBLH and PM<sub>2.5</sub>

2 concentration at Shijiazhuang for the WF and NF scenarios are shown in Figure 13(b) and 13(c).

3 In non-haze days, PBLH differences between the two scenarios are negligible due to low aerosol

4 loadings. In haze days, PBLHs in the WF scenario are generally lower (by up to 60%) than in the

5 NF scenario. As shown in Figure 13(c),  $PM_{2.5}$  concentration at Shijiazhuang in WF scenario is

6 higher than it in the NF scenario and the difference reaches about  $50\mu g/m^3$  on January 19.

7 Aerosols affect PBLHs in two ways: (1) radiation is scattered back to sky and absorbed, and as a

8 result, radiation reaching the surface is reduced (Figure 13(a)) and temperature is lowered; and (2)

9 suspended aerosols like BC absorb radiation to heat the upper PBL (Ding et al., 2013). Both of

10 these ways increase temperature inversion and atmospheric stability, and thus exacerbate  $PM_{2.5}$ 

11 pollution.

12 Figure 14 shows temporal variations of vertical profiles of (a) PM<sub>2.5</sub> (c) RH (e) temperature (g)

13 wind speeds differences in Beijing between WF and NF scenarios. When aerosol feedback is

14 included, PM<sub>2.5</sub> concentrations near Beijing surface are mostly increased, except on the morning

of January 17, on the afternoon of January 18 and on January 19 (Figure 14(a)). The increases of

16  $PM_{2.5}$  are caused by the above mentioned decrease of temperature gradient from surface to aloft

17 (shown in Figure 14(e)) and atmospheric stability. Apart from these,  $PM_{2.5}$  concentrations are

also affected by RH and wind speeds. In WF scenario, RH is generally increased near the surface,

19 especially on January 19 (Figure 14(c)), while horizontal wind speeds are also increased on

20 January 19, which is the main cause of decreases of  $PM_{2.5}$  concentrations in Beijing.

21

To evaluate the impact of aerosol feedback on horizontal meteorological fields and PM<sub>2.5</sub> 22 23 distributions, averaged differences of PM<sub>2.5</sub> concentrations, temperature, PBLHs and horizontal winds between WF and NF scenarios at 2p.m. and 2a.m. in haze days (from January 16 to 19) 24 25 were calculated and are shown in Figure 15. Figure 15(c) shows that PBLHs are reduced in almost all NCP areas when aerosol feedbacks are considered at 2p.m.. At 2p.m., PM<sub>2.5</sub> 26 concentrations increase about  $21.9\mu$ g/m<sup>3</sup> at Shijiazhuang (114.53°E, 38.03°N). In a few locations 27 (the areas to the south of Beijing (Figure 15(a)), PM levels decrease although PBLHs are 28 suppressed in those areas. The decreases of PM<sub>2.5</sub> concentrations in the areas south of Beijing are 29 30 due to big horizontal wind changes, shown in Figure 15(g). When aerosol feedback is included,

surface temperature decreases in areas where there are high aerosol loadings (Figure 15(e)). 1 2 Figure 15(d) shows that PBLHs are enhanced in east and southwest NCP areas at 2a.m. with 3 aerosol feedback. Aerosol feedback mechanism at night time is more complex compared to it at 4 day time. At night, there is no incoming shortwave radiation from the sun and major radiation is the long wave radiation emitted from the earth. The presence of clouds and some kinds of 5 aerosols can trap outgoing long wave radiation, and as a result, the surface atmosphere is 6 7 warmed. Different aerosols show different effects on long wave radiation. Greenhouse gases (GHGs) absorb long wave radiation, while large particles like dust scatter long wave radiation. 8 As a result, the upper atmosphere temperature is likely to be warmer or cooler than surface 9 10 atmosphere temperature. If the upper atmosphere is warmer than the surface, a stable PBL will form. This can explain why aerosol feedbacks increase PBL heights in some regions and 11 12 decrease in some other regions of NCP. Changes of PM<sub>2.5</sub> concentrations at 2a.m. are mainly caused by changed PBLHs (Figure 15(b)), showing decreasing trends in areas where PBLHs are 13 enhanced, because changes of winds are relatively small (Figure 15(h)). Temperature changes at 14 2a.m. are similar to it at 2p.m., but the magnitudes are smaller. 15

16

#### 17 **4.4.2** Impact of BC absorption on meteorology and PM<sub>2.5</sub> distribution

To investigate BC's influence on meteorology and air quality, sensitivity tests were conducted by 18 removing BC absorption in WRF-Chem (i.e., imaginary refractive index set to zero). Figure 14 19 20 shows temporal variations of vertical profiles of (b) PM<sub>2.5</sub> (d) RH (f) temperature and (h) wind speeds differences in Beijing between WF and NBCA scenarios. The differences between WF 21 22 and NBCA can be used to represent impacts of BC absorption since in WF scenario both scattering and absorbing are considered while in the NBCA scenario only scattering is 23 considered. It is obvious from Figure 14(f) that the upper atmosphere is heated by BC, especially 24 25 at 1.5km, which increases temperature inversion and atmospheric stability. BC absorption's 26 impacts on PM<sub>2.5</sub>, RH and wind speeds are similar to the impacts of both scattering and 27 absorption, but the magnitudes are smaller (Figure 14(b), (d) and (g)).

- Figure 16 is similar to Figure 15 except that the differences are between WF and NBCA
- 29 scenarios. At 2p.m.,  $PM_{2.5}$  concentration is increased about 14.4µg/m<sup>3</sup> in Shijiazhuang (114.53°E,
- $38.03^{\circ}$ N), accounting for about 65.7% of PM<sub>2.5</sub> changes due to the total aerosol feedback (Figure

16(a)). At 2p.m., the maximum decrease in PBLH is about 166.6m (Figure 16(c)), accounting for
about 59.9% of the maximum decrease in PBLH in Figure 15(c). At 2p.m., surface temperature
in high aerosol loading areas are decreased about 0-2 °C (Figure 16(e)), while the temperature
decreases in the same areas are above 2°C in Figure 16(e). At 2a.m., changes of PM<sub>2.5</sub>, PBLHs,
surface temperature and wind speeds are similar to Figure 15, with smaller magnitudes.

6

7 The contribution of BC absorption in aerosol feedbacks depends on the model performance in simulating BC and scattering aerosols (sulfate, OC). As shown in Figure 11, BC was 8 9 overestimated, and sulfate and OC were underestimated in Beijing. The overestimation could be 10 as large as a factor by 2 in some days. As a result, the relative contributions of BC absorption in aerosol feedbacks are uncertain. To explore the uncertainties of the BC absorption contribution, 11 12 we conducted a simulation by reducing BC emissions by 50%. The changes of PBLH and PM<sub>2.5</sub> concentrations at 2p.m. due to aerosol feedbacks and BC absorption after BC emission changes 13 14 are shown in Figure 17. The domain maximum increases of PM<sub>2.5</sub> concentrations because of aerosol feedbacks and BC absorption are  $19.1\mu g/m^3$  and  $10.2\mu g/m^3$ , respectively for the base and 15 50% BC emission cases. The domain maximum decreases of PBLH due to aerosol feedbacks and 16 BC absorption are 235.7m and 114.2m, respectively. These numbers are smaller than before 17 because BC emissions were reduced by 50%. Due to 50% perturbation in BC emissions, the 18 contribution of BC absorption in aerosol feedbacks decreased from about 60% to 50%. This 19 20 number can be additionally reduced if OC and sulfate concentrations are simulated well. In the 21 future, we can get more accurate estimations of BC absorption in aerosol feedbacks after the performances of simulating BC, OC and sulfate are improved. 22

23

## 24 **5 Conclusions**

25 In this study, the online coupled WRF-Chem model was used to reproduce the haze event

happened in January, 2010 in the NCP. The model was evaluated against multiple observations,

27 including surface observations of meteorological variables and air pollutants, atmospheric

sounding products, surface AOD measurements, and satellite AOD measurements. The

29 correlation coefficients between simulated and observed PM<sub>2.5</sub> concentrations in Beijing, Tianjin

and Xianghe stations are 0.77, 0.75 and 0.69, indicating that WRF-Chem provides reliable
 representation for the 2010 haze event in the NCP.

3

4 This haze event is mainly caused by high emissions of air pollutants in the NCP region and 5 stable weather conditions in winter. The haze built up almost simultaneously in major cities in 6 the NCP and dissipated from north to south. During haze days, horizontal wind speeds and 7 mixing heights were low, temperature inversion happened above surface and RH values were 8 above 40%. Photochemistry was not significant during haze days due to weak UV radiation. In 9 addition, secondary inorganic aerosols played an important role in the haze event. The role of 10 cloud chemistry in this haze event cannot be ignored.

11

The contribution of non-local sources to PM<sub>2.5</sub> in Beijing was also studied. The average contribution was about 64.5% in haze days. The FLEXPART model was implemented to investigate the sources of the non-local contributions and results show that air pollutants from south Hebei, Tianjin city, Shandong and Henan provinces are the major contributors to the PM<sub>2.5</sub> in Beijing.

17

Impacts of high aerosols in haze days on radiation, boundary layer heights and PM<sub>2.5</sub> have been 18 demonstrated. When aerosol feedback is considered, simulated surface radiation agrees well with 19 20 observations. In haze days, aerosol feedback has important impacts on surface temperature, RH 21 and wind speeds, and these meteorological variables affect aerosol distribution and formation in 22 turn. The role of BC in aerosol feedback loop has also been investigated. It can account for about as high as 65.7% of the PM<sub>2.5</sub> increases, and 59.9% of the PBLH decreases in Shijiazhuang. 23 24 More attention should be paid to BC from both air pollution control and climate change perspectives. Due to the underestimation of sulfate and OC, and overestimation of BC in the 25 current model, the contribution of BC absorption in aerosol feedbacks may have been 26 overestimated. We decreased the BC emission by 50%, and found that the contribution decreased 27 from about 60% to 50%. The uncertainty of this contribution can be additionally reduced by 28 29 improving the model performance in simulating sulfate and OC.

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- 10 chemical data are available at <u>http://www.acd.ucar.edu/wrf-chem/mozart.shtml</u>. Contact M. Gao
- 11 (meng-gao@uiowa.edu) or G.R. Carmichael (gcarmich@engineering.uiowa.edu) for data
- 12 requests.
- 13

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2	Data sets <sup>a</sup>	Variables <sup>b</sup>	Data frequency	Number of sites used	Data sources				
	CMDSSS	T2, RH2, WS10	Daily	4	http://cdc.cma.gov.cn/home.d				
					0				
	Atmospheric Sounding	T, RH	12 hours	1	http://weather.uwyo.edu/uppe				
	Sounding				<u>rair/sounding.html</u>				
	CARE- China	PM <sub>2.5</sub> , NO <sub>2</sub> , SO <sub>2</sub>	Hourly	3					
	CSHNET	AOD	Hourly	4					
	SDZ	T1.5, RH1.5, WS10, PM <sub>2.5</sub> , NO <sub>2</sub> ,	Hourly	1	Zhao et al. (2013)				
		CO							
	CALIPSO	AOD	N/A	N/A	http://www-				
					calipso.larc.nasa.gov/				
	MODIS	AOD	Daily	N/A	http://ladsweb.nascom.nasa.g				
					ov/data/search.html				

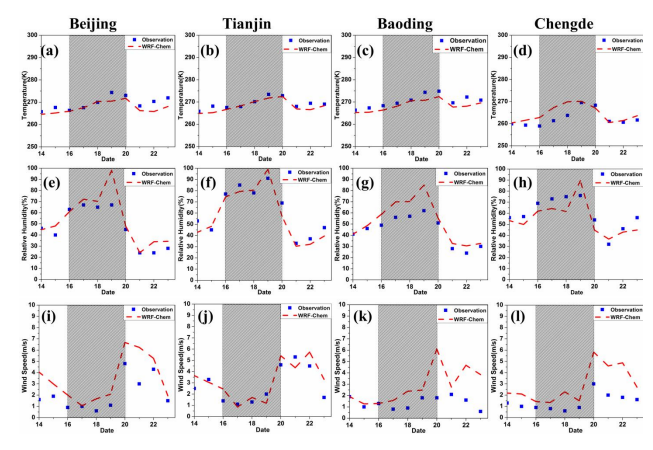
# Table 1. Observation Data and Variables Used in This Study

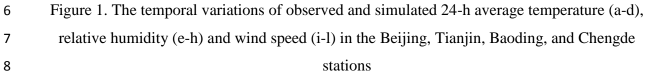
<sup>a</sup>CMDSSS—China Meteorological Data Sharing Service System; CARE-China—Campaign on the atmospheric
 Aerosol Research network of China; CSHNET—Chinese Sun Hazemeter Network; SDZ—Observation data at
 Shangdianzi site are extracted from paper Zhao et al. (2013); CALIPSO—The Cloud-Aerosol Lidar and Infrared
 Pathfinder Satellite Observation; MODIS—the Moderate Resolution Imaging Spectroradiometer. <sup>b</sup>T2— temperature

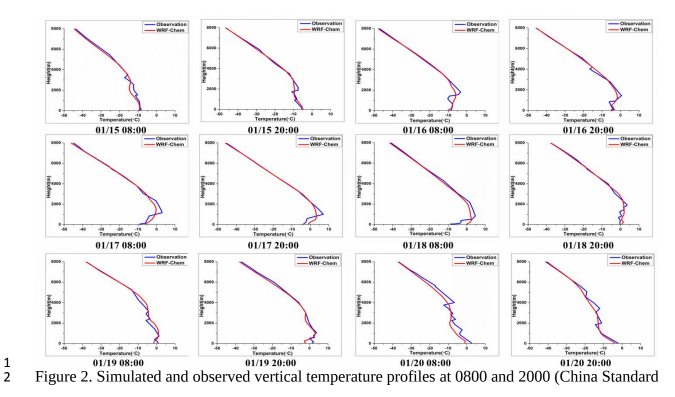
1	at 2m; RH2—relative humidity at 2m; WS10—wind speed at 10m; T1.5—temperature at 1.5m; RH1.5—relative																			
2	humidity at 1.5m; AOD—Aerosol Optical Depth.																			
3 4 5																				
6 Variables	Beijing Tianjin							Baoding					Chengde							
	Obs.	Mod.	MB	ME	RMSE	Obs.	Mod.	MB	ME	RMSE	Obs.	Mod.	MB	ME	RMSE	Obs.	Mod.	MB	ME	RMSE
T2(K)	269.5	267.6	-1.9	2.0	2.5	269.3	268.1	-1.1	1.2	1.5	270.4	268.5	-2.0	2.0	2.3	262.5	264.5	2.0	2.4	3.2
RH2 (%)	46.9	53.4	6.6	7.2	11.1	61.5	58.4	-3.1	5.9	6.4	44.4	52.5	8.1	8.1	10.4	59.4	55.0	-4.4	8.0	8.8
WS10(m/s)	2.1	3.4	1.3	1.3	1.6	2.8	3.2	0.4	1.0	1.1	1.4	2.8	1.4	1.4	2.1	1.4	2.9	1.5	1.5	1.8
7																				
8																				
9						Tabl	e 3. P	erfor	man	ce Stati	stics c	of PM <sub>2</sub>	2.5							
			Ob	s.	Mod	el	R	M	IB	MI	Ξ	NMB		NME	E M	FB	MF	Ξ		
			$(\mu g/m^3)$	(µg/n	n <sup>3</sup> )		(µg	/m <sup>3</sup> )	(µg/r	n <sup>3</sup> )	(%)		(%)	(9	%)	(%)				
	Beij	ing	111	.7	122.	1 0	).77	-10.	4	30.	4	-8.5		24.9	0	).4	26.3	;		
	Tiar	njin	103	3.3	141.	2 (	).75	-3	7.9	56.	1	-26.9		39.7	-7	7.8	49.6	)		
	Xian	ghe	93	.0	152.	6 (	).69	-5	9.7	68.	0	-39.1		44.5	-2	1.8	50.7	,		
10																				
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Table 4. Primary Aerosol, SIA and SOA ( $\mu$ g/m<sup>3</sup>) during Haze Days and Non-haze Days in Beijing

	Primary	SIA	SOA
Haze days	56.4	81.9	1.1
Non-haze days	14.2	10.8	0.3
Ratio	4.0	7.6	3.7









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Time, CST) from 15 January to 20 January

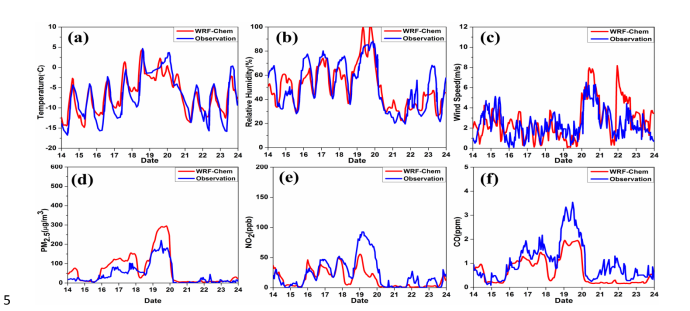


Figure 3. Simulated and observed hourly temperature, RH, wind speed, PM<sub>2.5</sub>, NO<sub>2</sub> and CO in 6 the Shangdianzi (SDZ) station 7

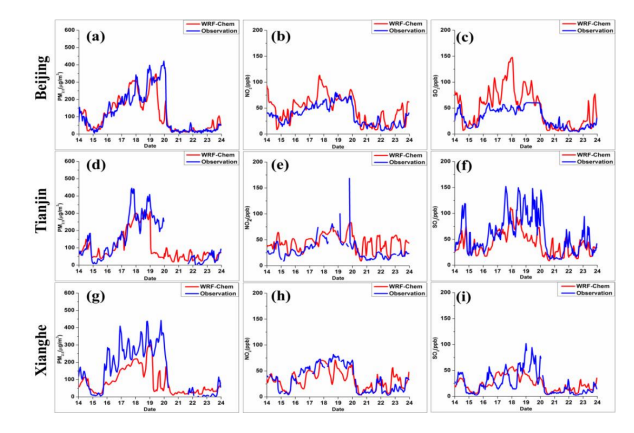


Figure 4. Temporal variations of the simulated and observed PM<sub>2.5</sub>, NO<sub>2</sub> and SO<sub>2</sub> at Beijing (a-c),
 Tianjin (d-f) and Xianghe (g-i) stations

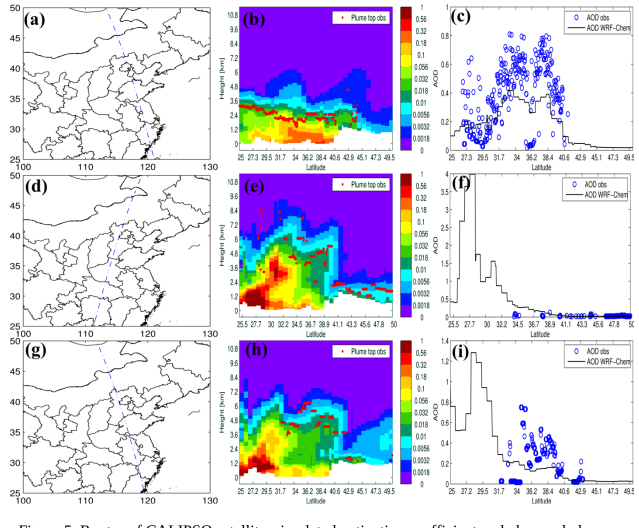
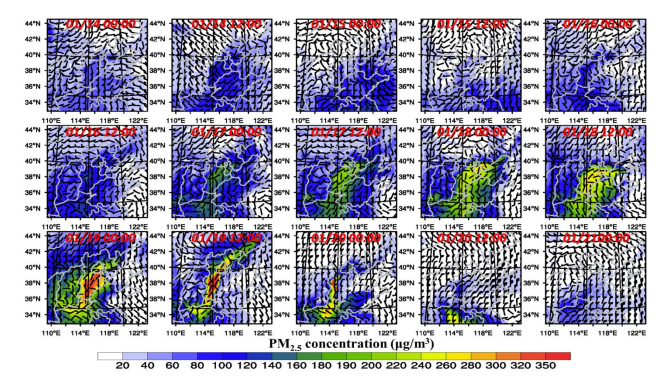
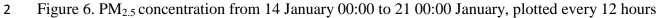
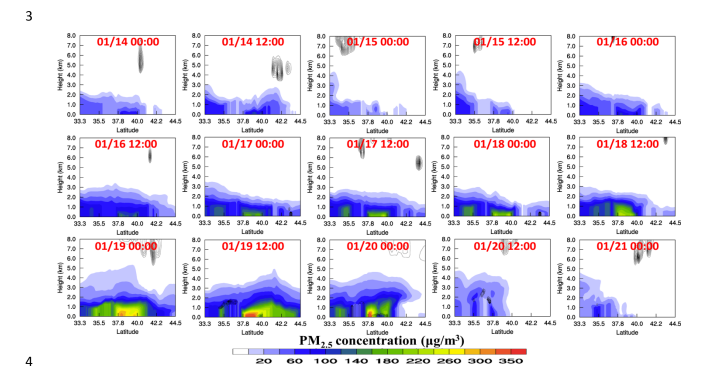


Figure 5. Routes of CALIPSO satellite, simulated extinction coefficient and observed plume
 top, and simulated AOD and CALIPSO retrieved AOD at 532nm at three moments: January 14
 12:00(CST) (a-c), January 21 02:00(CST) (d-f), and January 21 12:00(CST) (g-i)



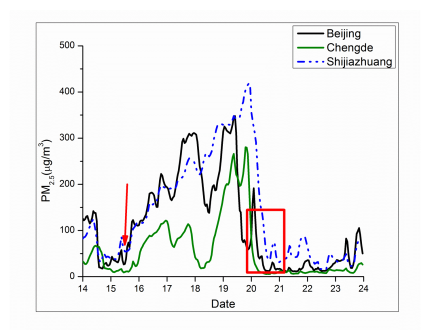




5 Figure 7. Cross section plots of PM<sub>2.5</sub> concentration and clouds from 14 January 00:00 to 21

00:00 January every 12 hours

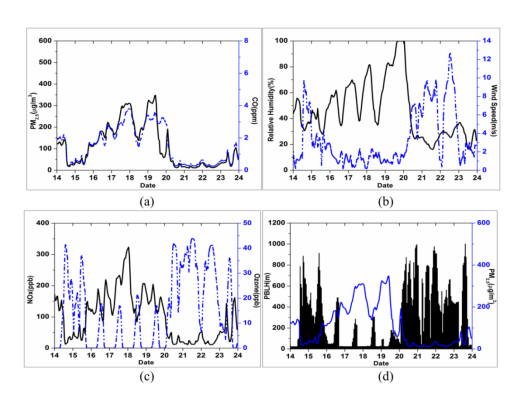
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2 Figure 8. Temporal variations of simulated PM<sub>2.5</sub> at Shijiazhuang, Beijing and Chengde





5 Figure 9. Simulated temporal variations of meteorological and chemical variables in Beijing

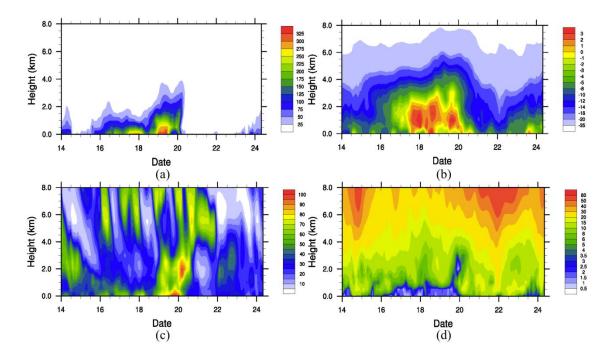


Figure 10. Temporal variations of vertical profiles of simulated (a) PM<sub>2.5</sub> (unit: μg/m<sup>3</sup>) (b)
 temperature (unit: °C) (c) RH (unit: %) (d) wind speeds (unit: m/s) in Beijing

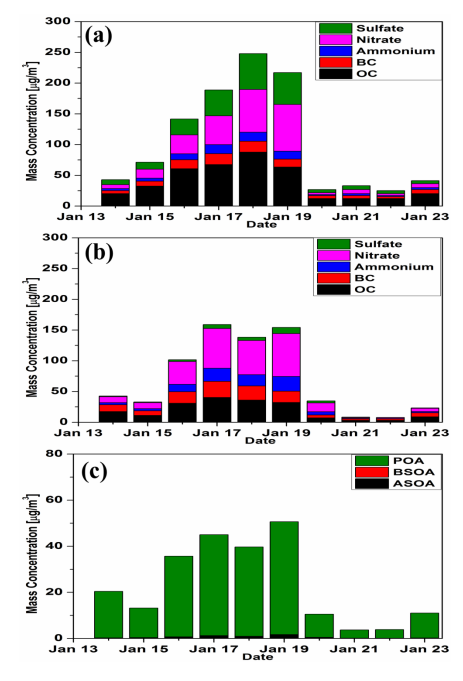


Figure 11. Observed (a) and simulated (b) chemical species of PM<sub>2.5</sub> and simulated SOA (c) in
 the Beijing site

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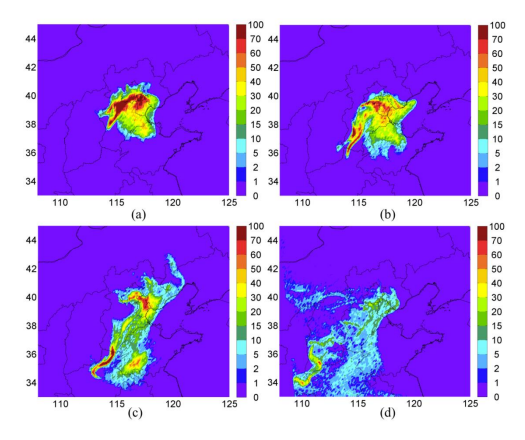


Figure 12. Backward dispersion of particles released on January 19 00:00, plotted 6, 12, 24, and
4 48 hours before being released (unit: number/grid cell)

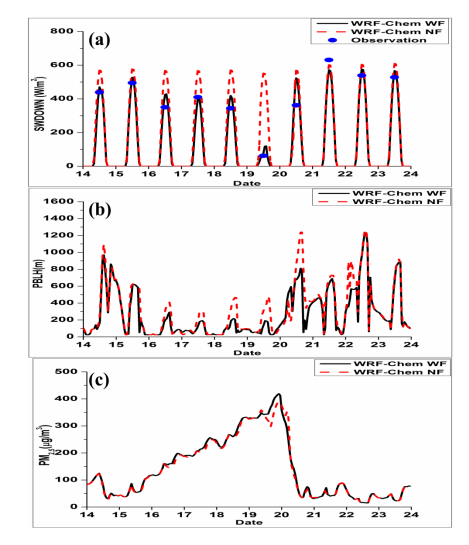


Figure 13. Observed daily maximum surface solar radiation and simulated surface shortwave
 radiation for the with feedback (WF) and without feedback (NF) scenarios in Beijing (a),
 simulated PBLH (b) in WF and NF scenarios at Shijiazhuang, and simulated PM<sub>2.5</sub> concentration
 (c) in WF and NF scenarios at Shijiazhuang

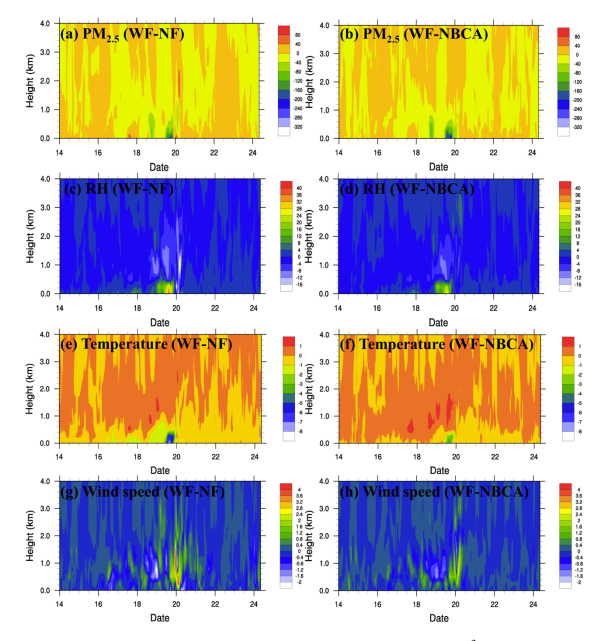


Figure 14. Temporal variations of vertical profiles of (a) PM<sub>2.5</sub> (unit: μg/m<sup>3</sup>) (c) RH (unit: %) (e)
temperature (unit: °C) (g) wind speeds (unit: m/s) differences in Beijing between WF and NF
scenarios; (b), (d), (f) and (h) are PM<sub>2.5</sub>, RH, temperature and wind speeds differences in Beijing
between WF and NBCA (BC absorptions are teased out) scenarios

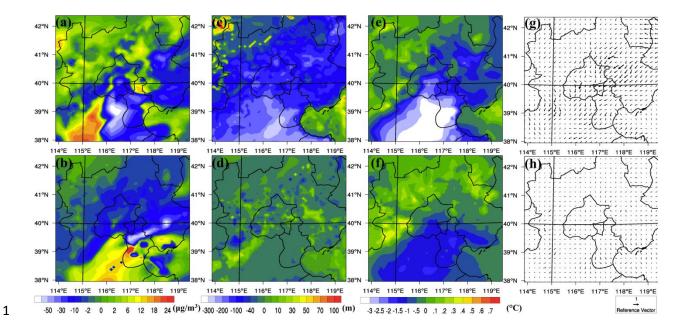


Figure 15. Differences of PM<sub>2.5</sub> concentration (unit: μg/m<sup>3</sup>), temperature (unit: °C), PBLH (unit:
m) and horizontal wind (unit: m/s) at 2p.m. (a, c, e, g) and 2a.m. (b, d, f, h) between WF and NF
scenarios

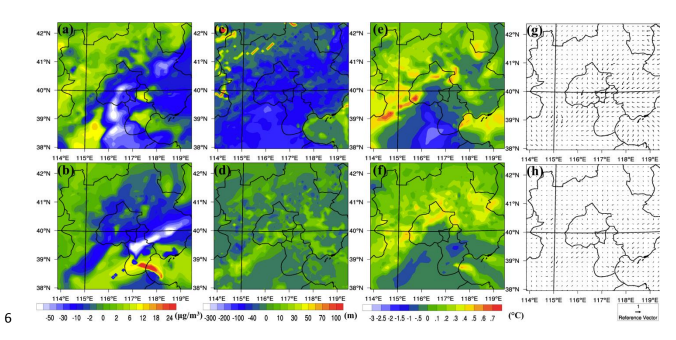


Figure 16. Differences of PM<sub>2.5</sub> concentration (unit: μg/m<sup>3</sup>), temperature (unit: °C), PBLH (unit:
m) and horizontal wind (unit: m/s) at 2p.m. (a, c, e, g) and 2a.m. (b, d, f, h) between WF and
NBCA scenarios

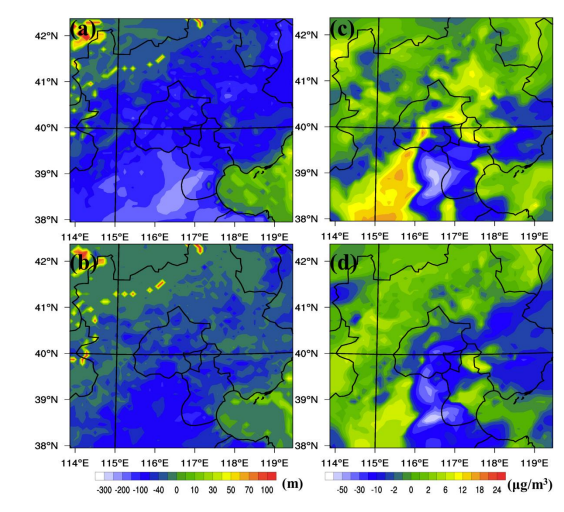


Figure 17. Differences of PBLH (unit: m) and PM<sub>2.5</sub> concentration (unit: μg/m<sup>3</sup>) at 2p.m.
between WF and NF scenarios (a, c) when BC emissions were reduced by half; differences of
PBLH (unit: m) and PM<sub>2.5</sub> concentration (unit: μg/m<sup>3</sup>) at 2p.m. between WF and NF scenarios (b,
d) when BC emissions were reduced by half