Aerosol radiative effects and feedbacks on boundary layer meteorology and 1 PM<sub>2.5</sub> chemical components during winter haze events over the 2 **Beijing-Tianjin-Hebei region** 3 4 Jiawei Li<sup>1</sup>, Zhiwei Han<sup>\*1,2</sup>, Yunfei Wu<sup>1</sup>, Zhe Xiong<sup>1</sup>, Xiangao Xia<sup>3</sup>, Jie Li<sup>1,2</sup>, Lin Liang<sup>1,2</sup>, 5 Renjian Zhang<sup>1</sup> 6 7 <sup>1</sup> Key Laboratory of Regional Climate-Environment for Temperate East Asia, Institute of 8 Atmospheric Physics, Chinese Academy of Sciences, Beijing 100029, China 9 <sup>2</sup> University of Chinese Academy of Sciences, Beijing 100049, China 10 <sup>3</sup> Key Laboratory of Middle Atmosphere and Global Environment Observation, Institute of 11 Atmospheric Physics, Chinese Academy of Sciences, 12 Beijing100029, China 13

14 Correspondence to: Zhiwei Han (hzw@mail.iap.ac.cn)

15

# 16 Abstract

An online-coupled regional chemistry/aerosol-climate model (RIEMS-Chem) was developed 17 and utilized to investigate the mechanisms of haze formation and evolution and aerosol 18 radiative feedback during winter haze episodes in February-March 2014 over the 19 Beijing-Tianjin-Hebei (BTH) region in China. Model comparison against a variety of 20 observations demonstrated a good ability of RIEMS-Chem in reproducing meteorological 21 variables, PBL heights, PM<sub>2.5</sub> and its chemical components, as well as aerosol optical 22 properties. The model performances were remarkably improved for both meteorology and 23 chemistry by taking aerosol radiative feedback into account. The domain average aerosol 24 radiative effects (AREs) were estimated to be -57 W m<sup>-2</sup> at the surface, 25 W m<sup>-2</sup> in the 25 atmosphere and -32 W m<sup>-2</sup> at the top of atmosphere (TOA), respectively, during a severe haze 26 episode (20-26 February), with the maximum hourly surface ARE reaching -384 W m<sup>-2</sup> in 27 southern Hebei province. The average feedback-induced changes in 2-m air temperature (T2), 28 10-m wind speed (WS10), 2-m relative humidity (RH2) and planetary boundary layer (PBL) 29 height over the BTH region during the haze episode were -1.8 °C, -0.5 m s<sup>-1</sup>, 10.0% and -184 30

m, respectively. The BTH average changes in PM<sub>2.5</sub> concentration due to the feedback were 31 estimated to be 20.0 µg m<sup>-3</sup> (29%) and 45.1 µg m<sup>-3</sup> (39%) for the entire period and the severe 32 haze episode, respectively, which demonstrated a significant impact of aerosol radiative 33 feedback on haze formation. The relative changes in secondary aerosols were larger than 34 those in primary aerosols due to enhanced chemical reactions by aerosol feedback. The 35 feedback-induced absolute change in PM2.5 concentrations was largest in haze persistence 36 stage, followed by those in growth stage and dissipating stage. Process analyses on haze 37 events in Beijing revealed that local emission, chemical reaction and regional transport 38 mainly contributed to haze formation in the growth stage, whereas vertical processes 39 (diffusion, advection and dry deposition) were major processes for PM<sub>2.5</sub> removals. Chemical 40 processes and local emissions dominated the increase in PM2.5 concentrations during the 41 severe haze episode, whereas horizontal advection contributed to the PM2.5 increase with a 42 similar magnitude to local emissions and chemical processes during a moderate haze episode 43 on 1-4 March. The contributions from physical and chemical processes to the 44 feedback-induced changes in PM2.5 and its major components were explored and quantified 45 46 through process analyses. For the severe haze episode, the increase in the change rate of  $PM_{2.5}$  (9.5 µg m<sup>-3</sup> h<sup>-1</sup>) induced by the feedback in the growth stage was attributed to the larger 47 contribution from chemical processes (7.3  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>) than that from physical processes (2.2 48  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>), whereas, during the moderate haze episode, the increase in the PM<sub>2.5</sub> change rate 49  $(2.4 \ \mu g \ m^{-3} \ h^{-1})$  in the growth stage was contributed more significantly by physical processes 50  $(1.4 \text{ µg m}^{-3} \text{ h}^{-1})$  than by chemical processes  $(1.0 \text{ µg m}^{-3} \text{ h}^{-1})$ . In general, the aerosol-radiation 51 feedback increased the accumulation rate of aerosols in the growth stage through weakening 52 vertical diffusion, promoting chemical reactions, and/or enhancing horizontal advection. It 53 enhanced the removal rate through increasing vertical diffusion and vertical advection in the 54 dissipation stage, and had little effect on the change rate of PM<sub>2.5</sub> in the persistence stage. 55

# 56

# 57 **1 Introduction**

58 Aerosols affect radiation transfer by scattering or absorbing solar and infrared radiation, 59 by acting as cloud condensation nuclei (CCN) to modify cloud properties, and by heating the

atmosphere to alter cloud formation, termed as the aerosol direct radiative effect, indirect 60 effect, and semi-direct effect (Twomey, 1974; Albrecht, 1989; Ramanathan et al., 2001), 61 respectively. In addition, there exists a set of interactions between chemistry, radiation and 62 meteorology (Dawson et al., 2007; Zhang, 2008; Isaksen et al., 2009; Baklanov et al., 2014; 63 Cai et al., 2017), which is highly complex and nonlinear and is currently one of the least 64 understood mechanisms in atmospheric science community. The above interactions are 65 traditionally not included or simplified in meteorological or chemical models, but have now 66 been considered and treated with different degrees of complexity in a few online coupled 67 models along with the advances in our knowledge and computer power, and the coupling of 68 meteorology and chemistry and its feedbacks remains one of the most challenging issues in 69 air quality and climate change (Zhang, 2008; Baklanov et al., 2014). 70

Rapid and continuous growth of economy and energy consumption in the past decades 71 has greatly elevated aerosol levels in China (Chan and Yao, 2008; Zhang et al., 2012; Li et al., 72 2017a), resulting in serious air pollution problem and potentially significant influence on 73 radiation and climate at multi-scales. Although emission control strategies have been 74 75 gradually implemented in recent years, haze events still often occur in east China, especially in the BTH region in wintertime due to both higher anthropogenic emissions and poorer 76 meteorological conditions. The haze issue has attracted wide attentions from public, 77 government and scientific community in China and a lot of monitoring and modeling studies 78 have been carried out to explore the sources, characteristics, formation and evolution 79 mechanisms of haze events at both urban and regional scales (Chan and Yao, 2008; Zhang et 80 al., 2012; Che et al., 2014; Guo et al., 2014; Huang et al., 2014; Sun et al., 2014; Zheng et al., 81 2015; Cheng et al., 2016; Ding et al., 2016; Li and Han, 2016a; Cai et al., 2017; Fu and Chen, 82 2017; Li et al., 2017b; Wang et al., 2017; Huang et al., 2018; Zhang et al., 2018a; Zhong et al., 83 2018a; Zhong et al., 2018b; An et al., 2019; Li et al., 2019a), through which our 84 understanding on haze issue has been promoted. However, there is still a large gap in our 85 knowledge formation 86 about haze mechanism, in particular the role of aerosol-radiation-meteorology feedback in haze formation and evolution (Fu and Chen, 2017; 87 Zhong et al., 2018a; An et al., 2019). 88

89

The aerosol radiative feedbacks on air quality and meteorology have ever been studied in

90 American and Europe with regional online coupled meteorology-chemistry models, such as 91 WRF-Chem (Zhang et al., 2010; Forkel et al., 2012), which demonstrates an important role of 92 the feedback in both air quality and meteorology. Carslaw et al. (2010) also pointed out the 93 complexity and significance of natural aerosol interactions and feedbacks within the Earth 94 system.

95 In east Asia, Han et al. (2013) revealed a significant feedback of mineral dust on dust deflation and transport, atmospheric dynamics, cloud and precipitation in spring and an 96 97 improvement of model prediction for PM (particulate matter) concentration and surface meteorology by the inclusion of the feedback effect into an online coupled 98 chemistry-aerosol-climate model. In recent years, given the increasing concerns on severe 99 PM pollution during haze days, some modeling studies have been conducted to investigate 100 the effect of aerosol radiative feedback on meteorology and near surface PM<sub>2.5</sub> concentration, 101 with focus on winter haze events in north China (Wang et al., 2014a; Wang et al., 2014b; 102 Zhang et al., 2015; Gao et al., 2015; Gao et al., 2016; Qiu et al., 2017; Zhao et al., 2017; 103 Zhang et al., 2018b; Chen et al., 2019; Wu et al., 2019). Most of the model results exhibited a 104 105 positive feedback which tended to increase PM<sub>2.5</sub> level, but the magnitude of such feedback differs largely, with the mean fractional change in PM2.5 concentration varying from just a 106 few percentage (Kajino et al., 2017; Wu et al., 2019) to around 30% (Wang et al., 2014a). 107 Some studies even show a negative feedback on PM<sub>2.5</sub> in Beijing (Zhang et al., 2015; Gao et 108 al., 2016). Recently, Gao et al. (2020) reported that the aerosol-radiation feedback-induced 109 daytime changes in PM<sub>2.5</sub> concentrations were less than 6% during haze days in the BTH 110 2010 from six applications of different online coupled region in January 111 meteorology-chemistry models under the international framework of the MICS-Asia (Model 112 Inter Comparison Study for Asia) Phase III. There existed some differences in the above 113 modeling studies in terms of study period and haze level, although they were all for winter 114 haze events in the BTH region. Zhong et al. (2018a) reported that over 70% of PM<sub>2.5</sub> increase 115 during cumulative explosive stage of haze event in Beijing in winter can be attributed to the 116 feedback effect based on integrated analysis of observations. The above studies highlight the 117 importance and large uncertainties in the aerosol radiative feedback, which require further 118 model development and investigation. 119

The diversity in the feedback effect among models could be associated with the 120 differences in the predictions of aerosol chemical components and aerosol optical properties, 121 the assumption of mixing state and hygroscopic growth scheme, as well as meteorological 122 fields, all of which determine the direction and magnitude of the feedback effect. Majority of 123 previous model studies underpredicted PM concentrations in the north China Plain, especially 124 for aerosol components, such as sulfate, nitrate and SOA concentrations, mainly due to 125 incomplete understanding and unrealistic treatment of secondary aerosol formation through 126 multi-phase chemical processes. Gao et al., (2018) reported that most of the participating 127 models (including WRF-Chem) in the MICS-Asia (Model Inter Comparison Study for Asia) 128 project underpredicted inorganic and organic aerosol concentrations by up to a factor of three. 129 Besides aerosol mass concentration, the unrealistic representation of aerosol properties, such 130 as composition, size distribution, mixing state, hygroscopic growth would also lead to model 131 biases in aerosol optical properties and direct radiative effects. The low biases in the 132 predicted aerosol compositions may lead to underpredictions of aerosol optical depth (AOD) 133 and consequently of aerosol radiative effects and feedback. Che et al. (2014) reported a 134 reduction of solar radiation by aerosols exceeding 200 W m<sup>-2</sup> during a severe haze event in 135 the north China Plain, much stronger than the estimations from models (around -100 W m<sup>-2</sup>). 136 Therefore, a realistic treatment and an accurate representation of aerosol processes and 137 properties are crucial to the estimation of aerosol radiative effects and feedback. 138

It has been well recognized that high aerosol loadings can apparently reduce incoming 139 solar radiation at the surface, leading to surface cooling and inversion associated with 140 reduced wind speed and vertical diffusivity, and consequently increase in surface aerosol 141 concentrations. However, while we have gained considerable knowledge on the overall 142 feedback effect of aerosols, the detailed processes involved in the feedback mechanism are 143 still poorly understood and barely quantified, for example, how does the aerosol radiative 144 effect modify meteorological variables? how do the radiative and meteorological changes 145 affect physical and chemical processes and in turn affect the magnitude and distribution of 146 aerosol components? how to quantify the relative contributions from various physical and 147 chemical processes to the feedback effect? 148

149

In this study, an online coupled regional climate-chemistry-aerosol model (RIEMS-Chem)

was developed and applied to explore the formation and evolution of haze events during 150 February-March 2014, in which a week-long haze episode with the daily maximum PM<sub>2.5</sub> 151 concentration up to 400 µg m<sup>-3</sup> (hourly mean up to 483 µg m<sup>-3</sup>) was observed. A wide variety 152 of field measurements of aerosol chemical components, optical properties, as well as 153 meteorological variables were conducted and applied to develop, constrain and validate the 154 model. The mechanisms of haze formation and evolution, aerosol radiative effects and 155 feedback on meteorology and chemistry were investigated and assessed. The overall aerosol 156 feedback on PM<sub>2.5</sub> and its aerosol compositions and the individual contributions to the 157 feedback from physical and chemical processes (advection, diffusion, deposition, chemistry, 158 etc.) during haze events were interpreted and quantified by a process analysis approach 159 incorporated in the model. The results from this study are expected to provide new insights 160 into the mechanism of aerosol-radiation-meteorology feedback, which is currently the source 161 of one of the largest uncertainties in haze formation and evolution. 162

163

### 164 **2 Model and Data**

#### 165 2.1 Model description

An online-coupled regional atmospheric chemistry/aerosol-climate model RIEMS-Chem 166 was used in this study, which was developed based on the Regional Integrated Environmental 167 Model System (RIEMS) (Fu et al., 2005; Wang et al., 2015). A series of modules and 168 parameterizations were adopted to represent major physical processes, including a modified 169 Biosphere-Atmosphere Transfer Scheme (BATS; Dickinson et al., 1993) to simulate land 170 surface process, the Medium-Range Forecasts scheme (MRF) to represent the planetary 171 boundary layer process (Hong and Pan, 1996), the cumulus convective parameterization 172 scheme from Grell (1993), and a modified radiation package of the NCAR Community 173 Climate Model, version CCM3 (Kiehl et al., 1996) to represent radiation transfer process 174 including aerosol effect. RIEMS had been applied to investigate East Asian monsoon climate 175 and the interactions among physical, biological and chemical processes (Xiong et al., 2009; 176 Zhao, 2013; Wang et al., 2015). RIEMS had participated in the Regional Climate Model 177 Intercomparison Project (RMIP) for Asia and it was one of the best models in predicting air 178

temperature and precipitation over east Asia (Fu et al., 2005).

The online-coupled model RIEMS-Chem has been developed in recent years by 180 incorporating major atmospheric chemistry/aerosol processes into the host model. Transport 181 of pPollutants are driven by meteorological fields provided byfrom RIEMS and changes of 182 pollutants exert feedbacks to-on the existing dynamic and physical modules (Han, 2010; Han 183 et al., 2012). Major atmospheric processes including emission, advection, diffusion, 184 185 multi-phase chemistries chemistry, dry deposition and wet scavenging of pollutants are considered. The advection and diffusion for pollutants are treated with the same scheme for 186 substances (such as moisture). Gas phase chemistry is represented by an updated 187 Carbon-bond mechanism (CB-IV; Gery et al., 1989). The aerosol effect on photolysis rate is 188 considered by using the Tropospheric Ultraviolet-Visible (TUV) radiation model (Lee-Taylor 189 and Madronich, 2007). Thermodynamic processes are calculated by the ISORROPIA II 190 model (Fountoukis and Nenes, 2007). Dry deposition velocity of aerosol is calculated by a 191 size-dependent scheme which is expressed as the inverse of the sum of resistance plus a 192 gravitational settling term, while below-cloud wet scavenging of aerosol is parameterized as a 193 194 function of precipitation rate and collision efficiency of particle by hydrometeor (Han et al., 2004). Heterogeneous reactions between gases and mineral dust and sea salt aerosols have 195 also been incorporated into RIEMS-Chem (Li and Han, 2010; Li et al., 2018a). SOA 196 formation is parameterized by a two-product model (Odum et al., 1997). 197

Current atmospheric chemistry models generally tend to underpredict sulfate 198 concentrations, especially in source regions during wintertime, such as north China, which 199 could be due to uncertainties in the treatment of chemical formation mechanism. Recent 200 model studies suggested that heterogeneous reactions could be an important pathway in 201 sulfate formation during winter haze episodes in north China (Li et al., 2017c; Li et al., 202 2018b). Therefore, heterogeneous reactions concerning the conversion of SO<sub>2</sub> to sulfate on 203 pre-existed hydrated aerosols were incorporated in RIEMS-Chem. The method of Li et al. 204 (2018b) was adopted, in which the uptake coefficient ( $\gamma_{so2}$ ) was a stepwise function 205 determined by the aerosol liquid water content (ALWCawe) which was predicted by the 206 ISORROPIA II model. Accordingly, the upper bound of awe-ALWC was set to 300 µg m<sup>-3</sup> 207  $(\gamma_{so2}=1\times10^{-4})$  while the lower bound was 30 µg m<sup>-3</sup>  $(\gamma_{so2}=1\times10^{-6})$ .  $\gamma_{so2}$  was linearly 208

interpolated between the upper and lower bounds in terms of <u>aweALWC</u>.

RIEMS-Chem treats 9 aerosol types including sulfate, nitrate, ammonium, black carbon 210 (BC), primary organic aerosol (POA), secondary organic aerosol (SOA), anthropogenic 211 primary PMs (PM<sub>2.5</sub> and PM<sub>10</sub>), dust and sea salt. The size distribution of the different types 212 of aerosols is previously prescribed based on the OPAC database (Optical Properties of 213 Aerosols and Clouds) (Hess et al., 1998). In this study, measurements in Beijing are used to 214 represent aerosol size distribution more realistically and to constrain the model. During the 215 study period, a scanning mobility particle sizer (SMPS; TSI, Inc., Shoreview, MN, USA) was 216 used to measure aerosol size distribution (Ma et al., 2017) and the geometric mean radius of 217 inorganic, black carbon and organic carbon aerosols were estimated to be 0.1 µm, 0.05 µm 218 and 0.1 µm, with standard deviations of 1.65, 1.6, 1.65, respectively. The above aerosol size 219 information was incorporated into RIEMS-Chem. The deflation of mineral dust is represented 220 by the scheme of Han et al. (2004) with 5 size bins (0.1-1.0, 1.0-2.0, 2.0-4.0, 4.0-8.0, 8.0-221 20.0 µm). Primary PMs from anthropogenic are also assigned to the 5 size bins. 222

Recent observational analyses of aerosol mixing state in Beijing (Ma et al., 2012; Wu et 223 224 al., 2016) indicated that more than 80% aerosols were internally mixed with BC during haze days, whereas about 70% of aerosols were externally mixed with BC in clean days, so an 225 internal mixing assumption was adopted for model simulation because this study focuses on 226 haze events. Recent measurements also exhibited that the geometric mean radius of dry 227 aerosol internal mixture during haze evolution from light-moderate to severe pollution stages 228 just increased slightly from 0.10 µm to 0.12 µm (Ma et al., 2017), so an average of 0.11 µm is 229 chosen for the geometric mean radius of internal mixture, with standard deviation of 1.65. 230

Aerosol optical parameters including extinction coefficient, single scattering albedo and asymmetry factor were calculated by a Mie-theory based method developed by Ghan and Zaveri (2007). In this method, the optical properties of different types of aerosols are pre-calculated by Mie theory and fitted by Chebyshev polynomials, which are functions of aerosol geometric mean diameter and refractive index:

236 
$$Q = \exp\left[\sum_{k=1}^{10} A_k T_k(x)\right],$$
 (1)

237 
$$x = \frac{2\log(D_p) - \log(D_{\min}) - \log(D_{\max})}{\log(D_{\max}) - \log(D_{\min})},$$
 (2)

where Q represents the aerosol optical properties (such as scattering efficiency).  $T_k(x)$  are the Chebyshev polynomial of order k, which is related to particle size,  $A_k$  are the Chebyshev coefficients which is related to refractive index,  $D_p$  is the geometric mean diameter,  $D_{min}$  and  $D_{max}$  are the minimum and maximum  $D_p$  for obtaining the Chebyshev polynomials, with values of 0.001 µm and 10 µm, respectively. It has been proved that 40 groups of  $D_p$  in the range from  $D_{min}$  and  $D_{max}$  are sufficient to control errors below 10% compared with classical Mie code calculation.

The effect of water uptake is treated by the  $\kappa$ -Köhler parameterization (Petters and Kreidenweis, 2007), which calculates aerosol wet diameter due to hygroscopic growth under different relative humidity. The bulk  $\kappa$  for internal mixture of aerosols is derived by the volume-weighted average of  $\kappa$  of each aerosol component, while the  $\kappa$  values for inorganic aerosols, BC, POA, SOA, dust and sea salt were set to 0.65, 0, 0.1, 0.2, 0.01 and 0.98, respectively, according to previous observational and modeling studies (Riemer et al., 2010; Liu et al., 2010a; Westervelt et al., 2012):

252 
$$\kappa = \sum_{j} \frac{V_{j}}{V_{a}} \kappa_{j}, \qquad (3)$$

where  $V_a$  is the total volume of dry aerosols,  $V_j$  is the volume of each aerosol component j. The refractive index of internally mixed aerosols is calculated using the Maxwell-Garnett mixing rule:

256 
$$R_{w}^{2} = R_{s}^{2} \left[ \frac{R_{i}^{2} + 2R_{s}^{2} + 2f_{i}(R_{i}^{2} - R_{s}^{2})}{R_{i}^{2} + 2R_{s}^{2} - f_{i}(R_{i}^{2} - R_{s}^{2})} \right],$$
(4)

$$f_i = \frac{V_i}{V},\tag{5}$$

where  $R_w$  is the refractive index of the internal mixture,  $R_i$  and  $R_s$  are the refractive index of insoluble components (BC and POA) and soluble components (inorganic aerosols, SOA and water), respectively.  $V_i$  represents the volume of insoluble components, V represents the total volume of wetted aerosols.

After obtaining the wet diameter  $(D_p)$  and refractive index of the internally mixed

aerosols ( $R_w$ ), the aerosol optical properties (Q) can be derived from formula (1) with the Chebyshev fitting coefficients table. Then, aerosol optical parameters, such as extinction coefficient can be obtained through multiplying Q by aerosol mass concentration from chemical module. The advantage of this optical module is the computational speed is much faster than that from the traditional Mie calculation, with a similar level of accuracy. This module has been successfully used in estimations of aerosol optical properties and direct radiative effects over East Asia (Han et al., 2011a; Li and Han, 2016b; Li et al., 2019b).

An empirical method from Hegg (1994) is applied to link cloud droplet number 270 concentration N<sub>c</sub> to mass concentration of hydrophilic aerosols (sulfate, nitrate, hydrophilic 271 BC and OC) to represent the first indirect effect, while the parameterization of Beheng (1994) 272 is used to represent the second indirect effect, in which the autoconversion rate converting 273 from cloud water to rain water depends on Nc and cloud liquid water content WL. The cloud 274 effective radius re is calculated based on Nc, WL and the cube of the ratio of the mean volume 275 radius and the effective radius of the cloud-droplet spectrum following Martin et al. (1994). 276 The effect of aerosols on ice nuclei and convective cloud is not treated yet in this model 277 because of the complexity and limitation in knowledge. 278

The aerosol optical parameters and N<sub>c</sub> due to aerosol activation calculated above are 279 transferred into radiation module to account for the perturbation of radiation and atmospheric 280 heating rate due to aerosol direct and indirect effects. The following subsequently called land 281 surface module and boundary layer module account for calculate the changes in land-air 282 fluxes of heat and moisture, turbulent diffusion coefficients and meteorological variables in 283 the boundary layer in response to the radiation change, and then air temperature tendency is 284 calculated in terms of the altered atmospheric heating rate and radiation, which further lead to 285 changes in meteorological variables, and in turn affect physical and chemical processes and 286 concentrations of aerosols and their precursors represented in the chemical module. All the 287 physical modules are called every 2.5 minutes and the transfer of variables between chemical 288 module and radiation/meteorological modules is made every 30 minutes. 289

RIEMS-Chem has been successfully applied in previous modeling studies of
 anthropogenic aerosols, mineral dust and marine aerosols regarding spatial-temporal
 distributions, physical and chemical evolutions, radiative and climatic effects over east Asia

(Han et al., 2011b; 2012; 2013; 2019; Li et al., 2014; Li and Han, 2016b; 2016c; Li et al.,
2019b). RIEMS-Chem <u>has have</u> been participating in the international model comparison
project Model Inter Comparison Study for Asia phase III (MICS-Asia III) and shows a good
ability in predicting PM<sub>2.5</sub> concentration and AOD over East Asia (Gao et al., 2018).

297

298 2.2 Process analysis

In RIEMS-Chem, a time-splitting scheme based on continuity equation is applied to 299 predict species concentrations; therefore, the species concentrations are the net results of 300 successive changes in concentration emissions, dynamic transport, and physical and chemical 301 processes different atmospheric physical and chemical processes, and the changes in species 302 concentration by each process can be recorded, allowing the quantification of individual 303 contribution of each process to species variation. In this study, a process analysis (PA) 304 scheme, which calculates the Integrated Process Rates (IPR) at each time step and each grid, 305 was embedded in RIEMS-Chem to identify the contributions of physical and chemical 306 processes to aerosol evolution. At each time step, the IPR for a certain process was calculated 307 308 by subtracting the species concentrations at the beginning of this process from the ones after the process. The IPR method has ever been applied to study the formation and fate of 309 particulate and gaseous pollutants in North America and China (e.g. Yu et al., 2008; Zhang et 310 al., 2009; Liu et al., 2010b). The processes involved in aerosol evolution include emissions of 311 primary species, advections (horizontal and vertical), diffusions (horizontal and vertical), dry 312 313 deposition, chemical processes (gas-phase chemistry, aqueous chemistry, thermodynamic equilibrium and heterogeneous reactions), cloud processes and wet deposition. Here cloud 314 process represents the effects of cloud attenuation of photolysis rate, aqueous-phase 315 chemistry and in-cloud mixing. In this study, PA is applied not only to quantify the 316 contributions of individual physical and chemical processes to haze evolution, but also to 317 help interpret the processes involved in aerosol radiative feedback. In addition, different from 318 the previous PA application, chemical processes are further classified into gas phase, 319 thermodynamic and heterogeneous reactions to provide more details on chemical pathways of 320 secondary aerosol formation. The mass balance of IPR has been examined, assuring that the 321 change in species concentration during one time step is equal to the sum of IPRs by each of 322

323 the processes.

324

#### 325 2.3 Emission inventories

Monthly mean anthropogenic emissions of sulfur dioxide  $(SO_2)$ , nitrogen  $(NO_x)$ , 326 ammonia (NH<sub>3</sub>), non-methane volatile organic compounds (NMVOC), carbon monoxide 327 (CO), black carbon (BC), primary organic carbon (POA), other anthropogenic primary PM<sub>2.5</sub> 328 and primary PM<sub>10</sub> in China for the year 2014 were obtained from the MEIC inventory 329 (Multi-resolution Emission Inventory for China) which was developed by Tsinghua 330 University (http://meicmodel.org). Anthropogenic emissions outside China were taken from 331 the MIX inventory which was developed to support the Model Inter-Comparison Study for 332 Asia phase III (MICS-Asia III) and the Hemispheric Transport of Air Pollution (HTAP) 333 projects (Li et al., 2017a). Both inventories of MEIC and MIX have the horizontal resolution 334 of 0.25 degree. Biomass burning emissions of aerosols and gas precursors for the year 2014 335 with a horizontal resolution of 0.25 degree were derived from the fourth version of the Global 336 Fire Emissions Database (GFED4) (Giglio et al., 2013). Monthly mean biogenic emissions of 337 338 isoprene and monoterpene were derived from Global Emissions Inventory Activity (GEIA, http://www.geiacenter.org/). The above emission data were bilinearly interpolated to the 339 lambert projection of RIEMS-Chem. 340

341

# 342 2.4 Model configuration and numerical experiments

RIEMS-Chem was configured on a lambert conformal projection with horizontal 343 resolution of 60 km, covering most areas of China, the Korean Peninsula, Japan and part of 344 the Indo-China Peninsula (Figure 1). 16 vertical layers distributed vertically and unevenly in 345 the terrain-following sigma coordinate, with the lowest 8 layers within the boundary layer. 346 This study focused on the Beijing-Tianjin-Hebei (BTH) region with more attentions to the 347 Beijing metropolitan. The study period was from 10 February to 12 March, 2014, 348 encountering several haze episodes. The first 7 days were taken as model spin-up and the 349 results from 17 February to 12 March were used for analysis. 350

Initial and boundary conditions for meteorological variables were provided by the final reanalysis data (FNL) with 1°×1° resolution and 6-hourly interval from the National Centers for Environmental Prediction (NOAA/NCEP, 2000). Lateral boundary conditions of chemical
species at 6-hourly interval were derived from the simulations of the global chemical model
MOZART-4 (Model for Ozone and Related chemical Tracers, version 4; Emmons et al.,
2010).

To investigate the aerosol radiative effects and its potential feedback on solar radiation, meteorological variables, planetary boundary layer (PBL) and aerosol concentrations in the study domain, two simulations were designed. The FULL simulation (with aerosols) considered all aerosol direct and indirect effects and feedbacks, and the <u>NoAer–NoAFB</u> simulation shuts off aerosol direct radiative effects and removes anthropogenic aerosols in aerosol indirect effects. In both simulations, the driving meteorological data, emissions and model settings were exactly the same.

364

365 2.5 Observational data

366 Several observational datasets for meteorological variables, aerosol concentrations and 367 aerosol optical parameters were obtained and used for model comparison and analysis.

In-situ 3-hourly observations of temperature at 2 meter (T2), wind speed at 10 meter (WS10) and relative humidity at 2 meter (RH2) from three meteorological monitoring sites around Beijing (Figure 1) were collected from the China Meteorological Data Service Center (CMDS) (http://data.cma.cn/).

To evaluate the model ability in reproducing evolution of planetary boundary layer 372 (PBL), high-frequency sounding data measured around 14:00 LST at the Xianghe station 373 (39°45'N, 116°58'E; approximately 63 km southeast of Beijing downtown) were collected, 374 from which the PBL height can be determined based on the vertical gradients of virtual 375 potential temperature and water mixing ratio according to the method from Heo et al. (2003). 376 This sounding dataset provided a good indicator of mixing layer height because the sounding 377 was launched at 14:00 LST and lasts for about one hour. The meteorological sounding was 378 launched in Xianghe once a week (every Tuesday) and totally four soundings were available 379 during the study period (18 and 25 February, 4 and 11 March). Fortunately, the four 380 soundings encountered one severe haze episode, one moderate haze episode, and two clean 381 days, providing robust evidences on day-to-day variation of mixing layer height under 382

various atmospheric conditions. Hourly downward shortwave radiation flux (SWDOWN) at
the surface was measured simultaneously at the Xianghe station by a pyranometer with sun
shield and was used in this study.

The measurements of mass concentrations of PM<sub>2.5</sub> and its components and aerosol 386 optical parameters were carried out at the tower division of the Institute of Atmospheric 387 Physics (IAP), Chinese Academy of Sciences (CAS) in Beijing (39°58'N, 116°22'E) from 17 388 February to 12 March, 2014. Real-time hourly PM<sub>2.5</sub> mass concentrations were online 389 measured by a hybrid beta attenuation particulate monitor (Model 5030 SHARP, Thermo 390 Scientific, USA). PM<sub>2.5</sub> samples were collected in parallel by an R&P Partisol®Model 2025 391 dichotomous sequential PM air sampler (Thermo, USA) and a MiniVol TAS PM sampler 392 (Airmetrics, USA) between 24 February and 12 March, 2014. Samples were collected twice 393 per day with one during the daytime (from 7:00 to 19:00 LST) and the other at night (from 394 19:00 to 7:00 of the next day). Totally 33 half-day samples were collected. Aerosol chemical 395 compositions including sulfate  $(SO_4^{2-})$ , nitrate  $(NO_3^{-})$ , ammonium  $(NH_4^{+})$ , BC and OC were 396 analyzed by ion chromatography (Dionex ICS-90 for cations and ICS-1500 for anions) and a 397 398 DRI-2100A carbonaceous aerosol analyzer. Real-time hourly aerosol extinction coefficient and aerosol absorption coefficient at dry condition (RH=10%) were synchronously measured 399 by a nephelometer (Aurora3000) and an aethalometer (AE-31), respectively. Detailed 400 information about this experiment including the sampling site, instruments, measurement 401 procedures and sample analysis were well documented in Ma et al. (2017). The mass 402 concentration of SOC was estimated using a revised EC tracer method (Zhao et al., 2013). 403

Measurements of AOD at the 4 sites (Nanjiao, Tianjin, Gucheng and Shangdianzi) in the 404 BTH region were obtained from the China Aerosol Remote Sensing Network (CARSNET) 405 (Che et al., 2014). Nanjiao is an urban site located in southern Beijing. Tianjin site is located 406 in the center of Tianjin city, about 120 km to the southeast of Beijing. Gucheng, a suburban 407 site in Hebei province, is about 130 km to the southwest of Beijing downtown. Shangdianzi is 408 located 150 km to the northeast of Beijing, which is a background station since it is far away 409 from anthropogenic sources. Daily mean AOD was derived by temporally averaging the raw 410 data measured by sunphotometer during daytime. To compare with the model output, AOD at 411 550 nm was used. 412

413

## 414 **3 Model validations**

415 3.1 Meteorological variables

Wind speed, temperature and relative humidity are key meteorological factors affecting 416 physical and chemical processes of atmospheric pollutants. The statistics for comparison 417 between in-situ observation and the FULL simulation for WS10, T2 and RH2 are presented in 418 Table 1. At the 3 sites (Beijing, Tianjin and Tanggu), the model performances were 419 reasonably good, although wind speeds were somewhat overpredicted. The overall 420 correlation coefficient (R) and normalized mean bias (NMB) at the 3 sites were 0.83 and -2% 421 for T2, 0.61 and -1% for RH2 and 0.47 and 31% for WS10. In all, RIEMS-Chem was able to 422 reasonably reproduce the meteorological variables during the study period. The statistics for 423 424 NoAerNoAFB simulation are also list in Table 1. It is noteworthy that the statistics for the FULL simulation are overall better than those for NoAerNoAFB simulation, such as the 425 warm bias in the simulated air temperature and positive bias in wind speed are apparently 426 reduced. This demonstrates the inclusion of aerosol radiative effects does improve 427 meteorological prediction in this study. 428

The observed hourly SWDOWN in Xianghe was compared with model simulation 429 (Figure 2a). In general, the FULL case well reproduced SWDOWN in clean days and 430 light-moderate polluted days, but tended to underpredict observations in heavy haze days, 431 such as the period from 20 to 26 February. Underpredictions of cloud amount and PM 432 concentrations could be reasons for the low bias. For the entire study period, the observed 433 and simulated (FULL) mean SWDOWN were 136.0 W m<sup>-2</sup> and 188.4 W m<sup>-2</sup>, respectively, 434 with R of 0.91 (Figure 2a). If only days with low cloud covers were considered, the 435 SWDOWNs were 183.3 W m<sup>-2</sup> and 213.7 W m<sup>-2</sup> from observation and the FULL case, 436 respectively, with the NMB of 16%. In contrast, the NoAerNoAFB case failed to capture the 437 decreasing tendency of SWDOWN during haze days, resulting in a larger bias (NMB of 72%) 438 than the FULL case. 439

440 To examine the model performance for meteorology in the vertical direction, we 441 collected meteorological sounding data at Beijing observatory from the website of University

of Wyoming (http://weather.uwyo.edu/upperair/sounding.html). Figure S1 and S2 present the 442 average observed and simulated vertical profiles of air temperature, wind speed and relative 443 humidity at 08:00 LST and 20:00 LST during the two haze episodes of 20-26 February and 444 1-4 March 2014 and the corresponding comparison statistics for these variables in the 445 troposphere and at altitudes below 3 km are listed in Tables S1 and S2. In general, the model 446 is able to generally capture the major features of vertical distribution of key meteorological 447 variables, although the model tends to predict higher relative humidity in the middle-upper 448 troposphere. Such overpredictions are also found for the same region in previous studies, 449 such as WRF-Chem simulation (Gao et al., 2016). The statistics indicate that the model 450 simulated vertical distribution of meteorological variables are within an acceptable accuracy 451 range of current meteorological model predictions. 452

453

#### 454 3.2 Planetary boundary layer (PBL) height

Figure 2b shows the simulated PBL heights at 14:00 LST from the FULL case and 455 NoAerNoAFB case during the study period and the observed PBL heights at 14:00 LST 456 457 determined from air soundings on 18 February (clean), 25 February (severe haze), 4 March (clean) and 11 March (haze), 2014, respectively. There was large variation in PBL height in 458 the afternoon, with higher PBL height in clean days and lower one in haze days, inversely 459 related to the PM<sub>2.5</sub> level. The FULL case well reproduced the very low PBL height during 460 the most severe haze episode on 25 February, with the observed and simulated PBL heights to 461 be 569 m and 587 m, respectively. In clean days, the much higher mixing layer was also well 462 captured, such as, on 4 March, the observed and simulated PBL heights were 2305 m and 463 2535 m, respectively. It is noteworthy that the simulated PBL heights in the NoAerNoAFB 464 465 case were consistently higher than those in the FULL case, and the PBL height simulation from the FULL case (considering aerosol radiative effects) was apparently in a better 466 467 agreement with observation than that from the NoAerNoAFB case, except for that on 18 February. 468

469

470 3.3 Mass concentrations of PM<sub>2.5</sub> and aerosol components

471

Figure 2c shows the hourly  $PM_{2.5}$  mass concentrations observed at the IAP site and those

472 from the FULL simulation and NoAerNoAFB simulation. The study period was characterized by three haze episodes, which was the episode 1 on 20–26 February, the episode 2 from 1 to 4 473 March, and the episode 3 from 8 to 11 March. The first episode experienced the most severe 474 pollution with the maximum hourly  $PM_{2.5}$  concentration exceeding 480  $\mu g\ m^{-3}$  on 25 475 February. The second and third ones were moderately polluted in terms of magnitude and 476 lasting time. In general, the model reproduced the hourly variation of PM<sub>2.5</sub> concentrations 477 reasonably well in the FULL case, although the peaks were somewhat underpredicted in 478 some days, which could be partly due to the overprediction of wind speed (Table 1) and 479 potential uncertainties in emission inventories. The low bias in PM2.5 concentrations could 480 also contribute to the overprediction of SWDOWN during the first haze episode (20-26 481 February) discussed in section 3.1. The average PM<sub>2.5</sub> concentrations during the study period 482 were 142.0 µg m<sup>-3</sup> and 131.4 µg m<sup>-3</sup> from observation and the FULL simulation, respectively, 483 with R of 0.8 and NMB of -7% (Table 2), which demonstrates a good model performance for 484 PM<sub>2.5</sub> predictions for the winter haze periods. A remarkable feature shown in Figure 2 is the 485 significant negative correlation between PM<sub>2.5</sub> concentration and PBL height and SWDOWN. 486

The comparison between the simulated daily mean surface aerosol components (sulfate 487  $(SO_4^{2-})$ , nitrate  $(NO_3^{-})$ , ammonium  $(NH_4^{+})$ , BC and OC) and observations at the IAP site are 488 presented in Figure 3. The daily mean observation in the figure is an average of the half-day 489 samples, while the original half-day samples are used for statistics calculation in Table 2. The 490 model (from the FULL case) generally exhibits a good performance for inorganic aerosol 491 (sulfate, nitrate and ammonium) concentrations in terms of both daily variation and 492 magnitude (Figure 3a - 3c). It is encouraging that the maximum values on 25 February during 493 the first haze episode and the moderate values on 3 March in the second haze episode are 494 well reproduced, although some low biases occurred in the last few days. On average, the 495 model simulations of 20.3 µg m<sup>-3</sup>, 24.3 µg m<sup>-3</sup> and 13.9 µg m<sup>-3</sup> are very close to the 496 observations of 21.0 µg m<sup>-3</sup>, 26.0 µg m<sup>-3</sup> and 14.1 µg m<sup>-3</sup> for sulfate, nitrate and ammonium, 497 respectively, with Rs of 0.92, 0.88 and 0.91 and NMBs of -4%, -6% and -2%, respectively 498 499 (Table 2). Most of the online coupled models tended to underpredicted sulfate concentration (Gao et al., 2016; Qiu et al., 2017; Gao et al., 2018), which led to an underestimation of 500 aerosol optical depth and radiative effect. The model in this study improves the simulation of 501

inorganic aerosols, mainly through the inclusion of heterogeneous chemical reactions forinorganic aerosols.

The model also reproduced the temporal variation and magnitude of BC (Figure 3d) and 504 OC (Figure 3e) concentrations in Beijing reasonably well. However, the model tended to 505 underpredict the peak OC values on 24-25 February and to overpredict BC concentrations 506 from late February to early March. The low bias in OC simulation during the haze episodes 507 could be attributed to the underprediction of SOC (Figure 3f) due to potentially missing 508 chemical pathways. Uncertainties in the emission inventory could also be a reason. Li et al., 509 (2017a) reported the uncertainties in BC and OC emissions for China could be  $\pm 200\%$ , larger 510 than those of emissions for gases (<70%) and primary PMs (~130%). The period mean BC 511 concentrations from observation and simulation were 5.2 µg m<sup>-3</sup> and 6.7 µg m<sup>-3</sup>, respectively, 512 with R of 0.92 and NMB of 28% (Table 2). The period mean simulated and observed POC 513 concentrations were 18.4 µg m<sup>-3</sup> and 15.5 µg m<sup>-3</sup>, respectively, with R of 0.93, whereas the 514 simulated SOC concentration was 9.9 µg m<sup>-3</sup>, lower than observation (13.6 µg m<sup>-3</sup>) by 27%, 515 with a correlation coefficient of 0.56. For OC (sum of POC and SOC), the simulated value 516 (28.3  $\mu$ g m<sup>-3</sup>) was very close to the observation (29.1  $\mu$ g m<sup>-3</sup>), with R of 0.88 and NMB of 517 -3%, respectively, which indicated a generally good model performance for the total OC 518 concentration. 519

It is noteworthy that by considering aerosol radiative effects, the model apparently improved simulations for both  $PM_{2.5}$  and its chemical compositions, which is illustrated by comparing model results between the FULL and NoAerNoAFB cases (Figure 2c, Figure 3 and Table 2). Another important finding is that the duration of haze episode was prolonged by about 2–3 hours by the aerosol radiative feedback compared with that without aerosol feedback (Figure 2c).

To evaluate the overall model performance on PM<sub>2.5</sub> and its gas precursors in the BTH 526 region, we also collected observations at 80 surface stations in 13 cities of the BTH from the 527 of **CNEMC** 528 website (China National Environmental Monitoring Center) (http://www.cnemc.cn/) and made a detailed comparison between observations and model 529 simulations for PM<sub>2.5</sub>, O<sub>3</sub>, NO<sub>2</sub> and SO<sub>2</sub>. The observed and simulated hourly mass 530 concentrations of these species in some typical cities are presented in Figure S3 - S8 and the 531

18

statistics for each city and for all the cities are presented in Table S3. The overall model
ability is generally satisfactory, with Rs of 0.87, 0.81, 0.60 and 0.74, NMBs of -0.4%, -11%,
-17% and 0.5% for PM<sub>2.5</sub>, O<sub>3</sub>, SO<sub>2</sub> and NO<sub>2</sub>, respectively, for all the sites in the BTH region.

535

## 536 3.4 Aerosol optical parameters

Figure 4a and 4b show the measured and simulated hourly aerosol extinction coefficient 537 (EXT) and aerosol absorption coefficient (ABS) at an RH of 10% at the IAP site during the 538 study period. It clearly showed that the model was able to well reproduce the magnitudes and 539 temporal variations of EXT and ABS under dry condition in the FULL case, although the 540 model tended to predict higher ABS in some days possibly due to the overprediction of BC 541 concentration. Single scattering albedo (SSA) which is defined as the ratio of scattering 542 coefficient (EXT minus ABS) to extinction coefficient is also given in Figure 4c. The FULL 543 case generally simulated high SSA values during haze episodes, such as 0.92 on 20-26 544 February, 0.85–0.9 from 1 to 4 March and 0.8–0.9 on 8–11 March, suggesting a dominant 545 role of light scattering aerosols in haze days. It is encouraging that the model reproduced SSA 546 547 during the severe haze episode (on 20–26 February) quite well, with both the simulation and observation being approximately 0.92. However, SSA observation in clean days (such as on 548 5-7 March) was lower than that in haze days, and the model tended to overpredict SSA in 549 clean days, which could be attributed to uncertainties in measurement. In clean days, both the 550 denominator (EXT) and numerator (EXT minus ABS) were small, a subtle perturbation in 551 EXT and/or ABS can result in a large variation in SSA. A previous observational study in 552 Beijing suggested that SSA observation was more uncertain in clean days than in polluted 553 days because the observed aerosol extinction coefficient was too low in clean days (Jing et al., 554 2015). On average, the observed EXT, ABS and SSA values were 0.51 km<sup>-1</sup>, 0.048 km<sup>-1</sup> and 555 0.85, respectively, whereas, the corresponding FULL simulations were 0.53 km<sup>-1</sup>, 0.052 km<sup>-1</sup> 556 and 0.88, with Rs of 0.8, 0.7 and 0.7 and NMBs of 4%, 10% and 5%, respectively (Table 2). 557 The above comparison demonstrates a good ability of the model in estimating aerosol optical 558 properties during the study period, which could be attributed to both the good performance 559 for aerosol compositions and the realistic representation of aerosol properties (aerosol size 560 distribution, mixing state, hygroscopic growth etc.), which is based on real-time 561

562 measurements in Beijing.

Besides EXT and ABS measured under dry condition, measurements of AOD at the four 563 CARSNET sites around Beijing (Nanjiao, Tianjin, Gucheng and Shangdianzi) were also used 564 to evaluate the model ability in simulating aerosol optical parameters in real atmosphere 565 (Figure 5). At the Nanjiao site, which is about 50km southeast of Beijing downtown (Figure 566 5a), AOD measurement was unavailable in most days during the first haze episode (20 to 26 567 February), with only two data (around 4.8) available on 25 February. The simulated daily 568 AOD from the FULL case varied from 3.1 to 4.0 during 24 - 26 February, somewhat lower 569 than the observation. The model tended to simulate lower AOD during the third haze episode 570 (8 to 11 March), which can be partly attributed to the predicted lower aerosol concentrations. 571 The measured AODs in Gucheng (southwest to Beijing) and Tianjin were similar in terms of 572 variation and magnitude (Figures 5b and 5c), showing high values during pollution periods 573 with the maximum daily AOD exceeding 4.0 in Gucheng and 3.5 in Tianjin. The FULL case 574 reproduced the AOD variations and magnitudes reasonably well at the two sites although low 575 biases still occurred during 8 to 11 March in Gucheng. For the regional background site 576 Shangdianzi (Figure 5d), the magnitude and variation of AOD were similar to those in 577 Nanjiao, suggesting that the haze episodes were regionally distributed because the temporal 578 variations and magnitudes of AOD were generally consistent at the four sites. 579

Table 3 summaries the performance statistics for daily mean AOD. In general, the model 580 reproduced the temporal variation and magnitude of AOD around Beijing reasonably well 581 with the overall R of 0.81 (0.67-0.90) and NMB of -8.6% (-15.6%-6.2%). The 582 underestimation is mainly contributed by the low biases during the third haze episode (8 to 11 583 March) when inorganic aerosol concentrations were underestimated (Figure 3a-3c). In 584 addition, the limitation in AOD samples during the severe haze episode in Nanjiao and 585 Shangdianzi could amplify the negative bias. At the Gucheng and Tianjin sites where more 586 samples were available, the mean measured AODs were 1.7 and 1.4, respectively, agreeing 587 well with the simulated values of 1.5 and 1.3 from the FULL case. 588

In summary, the above comparisons demonstrate that RIEMS-Chem was capable in reproducing the spatial distribution and temporal variation of meteorological variables (air temperature, wind speed, surface shortwave radiation, PBL height etc.), concentrations of total PM<sub>2.5</sub> mass and its chemical compositions and aerosol optical properties during the winter haze periods around Beijing. It is also noteworthy that the inclusion of aerosol radiative effects apparently improved the overall model performance for both meteorological variables and aerosol physical and chemical properties, highlighting the necessity to develop online coupled chemistry-meteorology model for both air quality and climate research. The good agreement above increases confidence in the reliability of the following model results on aerosol radiative effects and feedback.

599

# 600 4 Aerosol radiative effects and feedbacks Model results

# 4.1 Distributions of meteorological variables and near surface PM<sub>2.5</sub> concentration

The period-mean distributions of near-surface wind speed (WS10), temperature (T2), 602 relative humidity (RH2), PBL height and PM<sub>2.5</sub> concentration are shown in Figures 6a to 6e. 603 During the study period, westerly winds dominated the northwestern parts of the BTH region 604 while southeasterly prevailed over the southeastern parts, as a result, the near-surface wind 605 606 speeds were fairly weak over the convergence zone from southern Hebei province to Beijing (Figure 6c). Such wind pattern indicated that pollutants from southern parts of the domain 607 (such as Shandong and Henan provinces) can be transported northward to Beijing, Tianjin 608 and Hebei, and air pollutants over the weak-wind regions were easily accumulated to high 609 level. Near-surface temperature showed an apparent south-to-north gradient, with surface air 610 temperature in a range of 4 °C to 6 °C over the southern BTH region, -2 °C to 2 °C in the 611 vicinity of Beijing and parts of central Hebei, and lower than -2 °C in northern parts of the 612 domain (Figure 6a). Relative humidity was higher (~65% to 75%) over northern areas and 613 lower (~55% to 65%) over southern areas (Figure 6b). PBL height also exhibited an apparent 614 gradient in spatial distribution (Figure 6d), ranging from 800-1000 m in northern Hebei and 615 Inner Mongolia to about 600-700 m in southern Beijing, Tianjin and southern Hebei. A belt of 616 high PM<sub>2.5</sub> concentration spread from southwest to northeast (Figure 6e), with the maximum 617 value up to 150 µg m<sup>-3</sup> in the vicinity of Shijiazhuang and Beijing and Tianjin. The regions 618 with high PM<sub>2.5</sub> concentrations generally corresponded well to the weak-wind areas shown in 619 Figure 6c. 620

Averaged over the BTH region and the entire study period, the simulated T2, WS10, RH2, 621 PBL height and PM<sub>2.5</sub> concentration from the FULL case were 0.61 °C, 1.2 m s<sup>-1</sup>, 67.0%, 622 698.4 m and 90.0 µg m<sup>-3</sup>, respectively. According to the "Technical Regulation on Ambient 623 Air Quality Index" prescribed by Chinese Ministry of Environmental Protection in 2012, a 624 pollution event occurs when 24-hr mean PM<sub>2.5</sub> concentration  $\geq$  75 µg m<sup>-3</sup>. Totally, there were 625 11 days with domain and daily average  $PM_{2.5}$  concentration exceeding 75 µg m<sup>-3</sup> in the BTH 626 region, with the maximum exceeding 136  $\mu$ g m<sup>-3</sup>, indicating the severity of air pollution 627 during the study period. 628

629

# 4.2 Distributions of AOD, SSA and aerosol direct-radiative effects

Figure 6f shows that high AODs mainly distributed from northern Beijing to southwestern Hebei, southern Shanxi and northern Henan provinces, with the maximum up to 1.1. As AOD was determined by vertical profiles of aerosol compositions and RH, the spatial distribution of AOD was somewhat different from that of  $PM_{2.5}$  concentration. During the study period, the regional mean AOD in the BTH region was 0.78 (Table 4), about twice the long-term observed value of about 0.4 in February and March in the same region (Song et al., 2018).

The simulated SSAs were above 0.88 in the BTH region (Figure 6g), with relatively lower values (0.88 - 0.9) in the areas of high PM<sub>2.5</sub> concentration and higher ones (0.92 - 0.98)in the relatively clean areas. On average, the simulated SSA in the BTH was 0.91 (Table 4), within the range of 0.87 to 0.95 measured in the same region in January 2013 (Che et al., 2014) but slightly lower than the model simulated annual mean of 0.95 over eastern China (Zhuang et al., 2013).

All-sky aerosol radiative effects at the surface (ARE<sub>surf</sub>), at the top of atmosphere (ARE<sub>TOA</sub>) and in the atmosphere (ARE<sub>atm</sub>) under all-sky condition are presented in Figures 6h to 6j. During the study period, aerosols induced a negative ARE both at the surface and TOA and a positive ARE in the atmosphere over the BTH. The distribution of ARE resembles that of AOD, generally showing stronger effects over southwestern Hebei, Shanxi and northern Henan provinces where high AOD occurred. Moderate AREs appeared over Beijing, Tianjin and central Hebei, while relatively weak AREs appeared over the northern domain. The domain average AREs in the BTH region during the period were estimated to be  $-37 \text{ W m}^{-2}$ , 19 W m<sup>-2</sup> and  $-18 \text{ W m}^{-2}$  at the surface, in the atmosphere and at the TOA, respectively (Table 4). The indirect radiative effect was also estimated to be about  $-2 \text{ W m}^2$  at the surface and the TOA on average, much smaller than the direct radiative effect; therefore, the total <u>aerosol</u> radiative feedback is predominated by direct radiative effect during the study period.

The domain average all-sky AREs during the first haze episode (20-26 February) were -57 W m<sup>-2</sup>, 25 W m<sup>-2</sup> and -32 W m<sup>-2</sup> at the surface, in the atmosphere and at the TOA, respectively, and the values were further enhanced to -123 W m<sup>-2</sup>, 53 W m<sup>-2</sup> and -70 W m<sup>-2</sup> in terms of daytime mean. The maximum AREs at the surface and at TOA reached -384 W m<sup>-2</sup> and -231 W m<sup>-2</sup>, respectively, at 13:00 LST on 23 February in the vicinity of Shijiazhuang.

In Beijing, the estimated mean AREs were -70 W m<sup>-2</sup>, 32 W m<sup>-2</sup> and -38 W m<sup>-2</sup> at the surface, in the atmosphere and at the TOA, respectively, during the first haze episode, whereas the maximum ARE at the surface reached -304 W m<sup>-2</sup> at 13:00 LST on 22 February, which was associated with the high PM<sub>2.5</sub> concentration (453  $\mu$ g m<sup>-3</sup>) at that time.

Based on in-situ surface measurements, Che et al. (2014) estimated that during haze 665 periods in January 2013, the mean daytime AREs at Nanjiao and Xianghe were 666 approximately -42 W m<sup>-2</sup> and -50 W m<sup>-2</sup> at TOA, and -120 W m<sup>-2</sup> at the surface at both sites. 667 In this study, the daytime AREs averaged over the severe haze period (20-26 February) at 668 TOA were estimated to be -77 W m<sup>-2</sup> and -74 W m<sup>-2</sup> at Nanjiao and Xianghe, while the 669 corresponding AREs at the surface were -146 W m<sup>-2</sup> and -140 W m<sup>-2</sup>, respectively. Che et al. 670 (2014) also reported the maximum daily mean surface ARE of -220 W m<sup>-2</sup> at Nanjiao during 671 a severe haze episode in January 2013, in this study, the corresponding ARE was estimated to 672 be approximately -200 W m<sup>-2</sup> at the same site during the severe haze episode in February 673 2014. Therefore, the magnitudes of AREs during haze episodes simulated from this study 674 agreed favorably with the above observational based estimations around Beijing, despite the 675 different time period. 676

677

4.3 Impacts of aerosol radiative<u>The</u> feedback <u>effects</u> on meteorological variables and aerosols
 Figure 7a–7e shows the mean differences in T2, RH2, wind speed, PBL height and near
 surface PM<sub>2.5</sub> concentration induced by the radiative feedback due to all aerosols (FULL

681 minus <u>NoAerNoAFB</u>) in the domain during the study period.

The aerosol radiative effects led to a reduction in surface shortwave radiation and thus 682 surface air temperature in the entire domain. The magnitude of T2 variation decreased from 683 south to north of the BTH, with -1.6 °C to -2 °C in southern Hebei and -1.2 °C to -1.8 °C in 684 southern Beijing, respectively. Correspondingly, RH2 increased by 10%-16% in the above 685 regions. The changes in wind speed showed a patchy pattern, with decreases by  $\sim 0.1 \text{ m s}^{-1}$  in 686 southern Hebei, increases by ~0.2 m s<sup>-1</sup> in central Hebei, and decreases in most parts of 687 Beijing. Wind vector shows an anomalous northerly wind of  $\sim 0.5$  m s<sup>-1</sup> in the BTH region. 688 Due to the reduction in surface shortwave radiation, PBL height decreased over the entire 689 region, with the maximums up to 240 m in southern Hebei and northern Tianjin. The changes 690 in PBL height varied from -210 m in southern Beijing to -90m in northern Beijing. PM<sub>2.5</sub> 691 concentrations were consistently enhanced over the entire region, with the maximum increase 692 up to 33 ug m<sup>-3</sup> in southern Hebei and portions of Beijing and Tianjin. In most of the BTH 693 region, the percentage increase of PM<sub>2.5</sub> exceeded 25%, with the maximum increase 694 exceeding 33% in the vicinity of Shijiazhuang. It is of interest that the regions with the 695 maximum increase of PM<sub>2.5</sub> generally corresponded to those with the maximum decrease in 696 PBL height. The presence of aerosols reduced solar radiation reaching the ground surface, 697 resulting in decreases in surface air temperature and PBL height and an increase in relative 698 humidity, all of which favored accumulation and formation of aerosols due to weakened 699 vertical mixing and enhanced secondary aerosol formation. 700

The aerosol feedback during the first haze episode was further explored due to the much 701 higher PM<sub>2.5</sub> level than the period average. Figure 7f-7j show the mean changes in 702 meteorological variables and PM<sub>2.5</sub> concentrations during the first haze episode (20-26 703 February). In general, the changes induced by aerosol feedback were larger during the severe 704 haze episode than those over the entire study period. T2 decreased by 1.8 °C to 2.7 °C along 705 with an increase up to 20% in RH in southern Hebei and southern parts of Beijing and Tianjin. 706 Different from the entire period average, wind speed decreased consistently in the BTH, with 707 a maximum decrease of 1 m s<sup>-1</sup>. PBL height decreased by  $\sim$ 300 m in southern Hebei, 708 corresponding to the areas with large air temperature decrease. This resulted in a consistent 709 increase in PM<sub>2.5</sub> concentrations in the study domain, with the maximum increases exceeding 710

50% around Shijiazhuang and approximately 40% in Beijing and Tianjin, apparently higher than the entire period averages. If for daytime mean, the percentage changes of  $PM_{2.5}$  in the above areas increased to 70% and 60%, respectively (figure not shown). It is striking that the simulated maximum increase in hourly  $PM_{2.5}$  concentration can be up to 372 µg/m<sup>3</sup> (186%) in the vicinity of Shijiazhuang at about 10:00 LST on 24 February during the first haze episode, which demonstrates the substantial impact of the radiative feedback on  $PM_{2.5}$ concentration and haze formation.

It is worthwhile to further explore the effect of aerosol feedback during haze evolution. 718 We divided haze episode into three stages, the growth stage is defined as the time period of 719 PM<sub>2.5</sub> increase from clean condition to heavy pollution level, the persistence stage means the 720 duration period of haze and the dissipation stage means the period with a sharp decrease in 721 PM<sub>2.5</sub> concentration usually along with a cold front passage. During the first heavy haze 722 episode (20-26 February) in Beijing, aerosol radiative feedback caused the increases in PM<sub>2.5</sub> 723 concentration of 55  $\mu$ g m<sup>-3</sup>, 84  $\mu$ g m<sup>-3</sup> and 40  $\mu$ g m<sup>-3</sup>, with the fractional changes of 31%, 41% 724 and 67%, respectively, during the growth, persistence and dissipation stages. The larger 725 726 fractional change of PM<sub>2.5</sub> in the dissipation stage is due to the relatively large feedback-induced increase and the lowest PM2.5 concentration in the NoAerNoAFB case in 727 this stage. During the second haze episode (1-4 March), the increases in PM<sub>2.5</sub> concentration 728 due to aerosol feedback were 25  $\mu g$  m^-3, 45  $\mu g$  m^-3 and 24  $\mu g$  m^-3, with the fractional changes 729 of 21%, 35% and 34%, respectively, which are lower than the feedback effect during the first 730 haze episode. So, in terms of magnitude, the largest feedback effect on PM<sub>2.5</sub> occurred in the 731 persistence stage, followed by that in the growth stage, although the fractional change of 732 PM<sub>2.5</sub> was larger in the dissipation stage. 733

Table 5 summarized the average feedback-induced changes in meteorological variables and PM<sub>2.5</sub> concentrations over the BTH region during the entire and the first haze periods. During the study period, due to the radiative feedback by all aerosols (FULL minus NoAer<u>NoAFB</u>), surface air temperature and wind speed decreased by 1.4 °C and 0.04 m s<sup>-1</sup>, respectively, with RH increased by 8.7% in the BTH. PBL height was reduced by 160 m (or a percentage change of -18.6%) on average, along with a reduction of 3.3 m<sup>2</sup> s<sup>-1</sup> (-27.0%) in vertical diffusivity coefficient (K<sub>z</sub>), resulting in an increase of PM<sub>2.5</sub> level by 20.0  $\mu$ g m<sup>-3</sup> (28.6%). It is noticed that the above changes were strengthened during the severe haze episode on 20–26 February, with the 7-day average decreases in T2, WS10, PBL height and K<sub>z</sub> being up to -1.8 °C, -0.5 m s<sup>-1</sup>, -183.6 m (-31.0%) and 3.9 m<sup>2</sup> s<sup>-1</sup> (-48.8%), respectively, and the PM<sub>2.5</sub> concentration increased by 45.1  $\mu$ g m<sup>-3</sup> with a percentage increase of 38.7%. Because aerosols affect solar radiation in daytime, in term of daytime mean, the 7-day mean changes in T2, WS10 and PBL height were estimated to be -2.5 °C, -0.6 m s<sup>-1</sup> and -307.3 m (-37.6%), respectively, leading to an increase of 49.3  $\mu$ g m<sup>-3</sup> (48.5%) in PM<sub>2.5</sub> concentration.

The impact of aerosol radiative feedback in Beijing (Table 6) was stronger than the regional mean. During the first haze episode, the 7-day average changes in T2, WS10, RH2, PBL and PM<sub>2.5</sub> were estimated to be -2.1 °C, -0.6 m s<sup>-1</sup>, 17.0%, -195.6 m (-35.9%) and 68.0  $\mu$ g m<sup>-3</sup> (39.1%), respectively, and the daytime mean change in PM<sub>2.5</sub> concentration increased to 83.2  $\mu$ g m<sup>-3</sup> (60%), respectively.

Table 7 presents the average changes in major aerosol components (BC, sulfate and 753 nitrate) in PM<sub>2.5</sub> induced by the feedback effect. Over the BTH region, the feedback caused 754 the average increases in sulfate and nitrate by 5.0  $\mu$ g m<sup>-3</sup> (46.4%) and 6.8  $\mu$ g m<sup>-3</sup> (37.3%), 755 respectively, for the entire period, and by up to 12.6  $\mu$ g m<sup>-3</sup> (66.9%) and 14.6  $\mu$ g m<sup>-3</sup> (40.9%), 756 for the first haze episode. The feedback-induced increases in BC was 0.9  $\mu$ g m<sup>-3</sup> (25.1%) and 757 1.9 µg m<sup>-3</sup> (32.9%), respectively, for the entire period and the first haze episode. It was 758 noticed that the feedback-induced changes in sulfate and nitrate concentrations were larger 759 than that in BC concentration. This was because that the concentrations of secondary aerosols 760 were increased not only by weakened vertical diffusivity but also by enhanced chemical 761 reactions due to the radiative feedback, which will be discussed in detail in section 5.2. 762

The above analysis demonstrates a significant impact of aerosol feedback on PM<sub>2.5</sub> 763 concentration during winter haze episodes in the BTH region. Previous modeling studies 764 reported different degrees of aerosol radiative feedback in east China. Gao et al. (2015) 765 simulated an increase of near surface  $PM_{2.5}$  concentrations to be 10-50 µg m<sup>-3</sup> or 5-25% in the 766 BTH during a severe haze episode on 10-15 January 2013 by using WRF-Chem. For the 767 similar time period and region, Wang et al. (2014a) reported an increase in PM<sub>2.5</sub> 768 concentrations by 15-50 µg m<sup>-3</sup> or 10-30% by using a regional coupled model NAQPMS. Wu 769 et al. (2019) used WRF-Chem to investigate a haze episode from 5 December 2015 to 4 770

January 2016 in the North China Plain and found that the aerosol radiative effects can enhance near-surface  $PM_{2.5}$  concentration by 10.2 µg m<sup>-3</sup> (7.8%) on average.

The results from this study demonstrate a stronger aerosol-radiation feedback than 773 previous modeling studies, with an average increase in PM<sub>2.5</sub> concentration by up to 45.1 µg 774 m<sup>-3</sup> (38.7%) during a severe haze episode and further to 49.3 µg m<sup>-3</sup> (48.5%) for daytime 775 mean over the BTH region. This study also highlights that the aerosol feedback effect can 776 result in an increase of hourly  $PM_{2.5}$  concentrations by up to 372 µg m<sup>-3</sup> (186%) in the 777 vicinity of Shijiazhuang during the severe haze episode. The stronger feedback effect in this 778 study than previous model simulations is mainly due the predicted higher concentration of 779 aerosol components (especially inorganic aerosols) and aerosol optical properties, which are 780 also in a better agreement with observations. It is noticed that a recent study (Zhong et al., 781 2018a) reported that the aerosol feedback effect contributed over 70% to PM2.5 increase 782 during the cumulative explosive stage of haze event in winter Beijing based on integrated 783 analysis of observations from 2013 to 2016, which suggested a dominant role of the feedback 784 effect in haze formation. 785

786

# 787 **5** Process analysis of haze evolution and aerosol radiative feedback

The process analysis (PA) method calculates the Integrated Process Rates (IPRs) and is 788 applied to quantify the individual contributions of different physical and chemical processes 789 to variations of PM<sub>2.5</sub> and its chemical components. These processes include emission, 790 horizontal and vertical advections (HADV and VADV), horizontal and vertical diffusions 791 (HDIF and VDIF), dry deposition (DDEP), cloud (CLD, including aqueous chemistry and 792 wet scavenging), gas chemistry (GAS), thermodynamic chemistry (Thermo) and 793 794 heterogeneous chemistry (HET). The focus of this study is Beijing, so the model grid cell near the surface having Beijing is selected for analysis. 795

796

5.1 The mechanism of haze evolution related to various processes

5.1.1 Haze evolution during 20–26 February

There was a severe haze event lasting for about 7 days, with the maximum hourly  $PM_{2.5}$ 

up to 482  $\mu$ g m<sup>-3</sup> on 26 February. This haze was initially formed on 20 February, with the observed surface PM<sub>2.5</sub> concentration less than 50  $\mu$ g m<sup>-3</sup> on 19 February, rapidly increased to 343  $\mu$ g m<sup>-3</sup> on 20 February, and reached 482  $\mu$ g m<sup>-3</sup>, followed by rapid haze dissipation on 26 February due to the arrival of a cold front.

PA was used to provide insights into the evolution mechanism of the haze episode, which 804 was divided into the clean, growth, persistence and dissipation stages in this study. Figure 8 805 shows the average process budgets for changes in PM<sub>2.5</sub> (which is the sum of sulfate, nitrate, 806 ammonium, BC, OC, SOC and primary PM<sub>2.5</sub>) and its major components in Beijing during 807 the four stages of the first haze period (Figure 8) from the FULL simulation. Figure 8a shows 808 the hourly IPRs of PM<sub>2.5</sub> by physical and chemical processes. The emission of primary 809 aerosols was the largest contributor to the PM2.5 mass with a constant IPR of 29.8 µg m<sup>-3</sup> h<sup>-1</sup> 810 (not shown in Figure 8a for clarity) due to the use of a monthly based emission inventory. 811 Chemical processes (GAS, Thermo and HET) also contributed largely to PM<sub>2.5</sub>, with 812 generally larger contributions in the growth and persistence stages. Thermodynamic 813 equilibrium processes and gas chemistry accounted for over 2/3 of the chemical contributions, 814 with the former process mainly accounting for the formation of nitrate and ammonium and 815 the latter one for sulfate formation. The contribution from heterogeneous reactions was 816 generally small, but when conditions were favorable (such as high RH and high aerosol 817 concentration providing sufficient reaction surfaces), its contribution would also be 818 significant, such as on the morning of 22 February, at nighttime from 23 to 24 February, and 819 on the mornings of 25 and 26 February. Vertical diffusion and dry deposition consistently 820 removed PM<sub>2.5</sub> from the atmosphere. In general, the larger IPRs from both VDIF and DDEP 821 during the clean and dissipation stages resulted in lower PM2.5 concentrations, whereas the 822 lower IPRs from VDIF and DDEP in the growth stage favored aerosol accumulation. In the 823 persistence stage, the IPRs of VDIF and DDEP were generally small. It should be noted that 824 on every midday, when PBL was fully developed, the vertical diffusion reached the daily 825 maximum, producing distinctly large negative IPRs of VDIF. Advections (HADV and VADV) 826 and horizontal diffusion either contributed to the accumulation or loss of PM2.5. During this 827 severe haze episode, horizontal diffusion served as a sink of PM<sub>2.5</sub>, producing a negative IPR 828 of HDIF through the event. Horizontal advection served as a sink of PM2.5 in most of the time, 829

leading to a negative IPR of HADV, however, when the removal of PM2.5 by vertical 830 diffusion was strong at the midday, aerosols were advected to Beijing from surrounding areas 831 due to mass balance, resulting in a positive IPR of HADV. The positive IPR of VADV during 832 the growth and persistence stages of this event indicated that the downward transport of 833 aerosols from upper levels also contributed to the PM2.5 increase, such as on the mornings of 834 22 and 25 February. In general, the IPRs (represented the net effect of all processes, denoted 835 by the red line in Figure 8a) exhibited small positive values from evening to next morning on 836 every day, indicating a gradually increasing PM<sub>2.5</sub> concentration, whereas on every midday, 837 relatively large negative IPRs occurred, indicating an apparent decrease in PM<sub>2.5</sub> 838 concentration at that time. It should be mentioned that even in the persistence stage, the 839 diurnal variation of PM<sub>2.5</sub> occurred although the change rates were generally weaker than 840 those in the growth and dissipation stages. 841

Figure 8b to 8f show the mean IPRs for PM<sub>2.5</sub> and its major chemical components as well as the key meteorological variables averaged over each stage to help interpret the formation and evolution mechanism of this severe haze episode.

In the clean stage, emission and chemistry were the two major processes for PM<sub>2.5</sub> 845 production (Figure 8b). Emission contributed predominately to PM2.5 production (IPRs of 846 29.8  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>), whereas the contributions of gas (9.2  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>) and thermodynamic 847 chemistry (7.3  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>) were comparable. The most influential process for PM<sub>2.5</sub> removal 848 was vertical diffusion, with the IPRs of  $-30.3 \ \mu g \ m^{-3}h^{-1}$ , comparable to that of emission. Dry 849 deposition was the second most important process for PM<sub>2.5</sub> loss (-12.2 µg m<sup>-3</sup> h<sup>-1</sup>), followed 850 by horizontal diffusion. Advection had a negligible effect on PM<sub>2.5</sub> in this stage. In the growth 851 stage, it is noteworthy that the contributions from vertical diffusion (VDIF) and dry 852 deposition (DDEP) to PM<sub>2.5</sub> removal decreased markedly from -30.3  $\mu$ g m<sup>-3</sup> h<sup>-1</sup> and -12.2  $\mu$ g 853  $m^{-3} h^{-1}$  in the clean stage to -21.6 µg  $m^{-3} h^{-1}$  and -9.2 µg  $m^{-3} h^{-1}$ , respectively (Figure 8b), 854 mainly due to the decrease in wind speed and the increase in stability indicated by the 855 reduced vertical diffusivity coefficient K<sub>z</sub> (Figure 8f), leading to increases in concentrations 856 of all species. It is impressive that the contributions from chemical processes 857 (GAS+Thermo+HET) increased apparently compared with those in the clean stage, with the 858 IPRs from gas, thermodynamic and heterogeneous chemistry increase to 12.1 µg m<sup>-3</sup> h<sup>-1</sup>, 16.0 859

 $\mu g m^{-3} h^{-1}$  and 5.4  $\mu g m^{-3} h^{-1}$ , respectively. The increase in the contribution from 860 heterogeneous chemistry was mainly attributed to the increase in relative humidity and 861 aerosol surfaces, upon which heterogeneous reactions took place. It is noticed that the 862 contribution of thermodynamic chemistry increased with increasing relative humidity as well 863 along with haze formation (Figure 8f). The increase in the contribution of thermodynamic 864 chemistry was remarkable (with IPR from 7.3 to 16 µg m<sup>-3</sup> h<sup>-1</sup>), because gas precursors of 865 aerosols increased due to weakened vertical diffusivity and higher relative humidity during 866 haze period favored condensation from gas to aerosol phase. It is of interest that vertical 867 advection also contributed to PM<sub>2.5</sub> production (IPR of 5.4 µg m<sup>-3</sup> h<sup>-1</sup>) in this stage, which 868 indicated a potential downward import of PM<sub>2.5</sub> from upper layer. It is also noticed that 869 horizontal advection contributed to PM<sub>2.5</sub> loss (-12.8 µg m<sup>-3</sup> h<sup>-1</sup>). This is because the strong 870 gradient between the increased PM<sub>2.5</sub> level in Beijing caused by weakened vertical diffusivity 871 and the relatively lower PM<sub>2.5</sub> level in the surrounding areas, which led to an outflow of 872 PM<sub>2.5</sub>. In the growth stage, the net variation rate (IPR) of PM<sub>2.5</sub> concentration was 14.1 µg 873 m<sup>-3</sup> h<sup>-1</sup>, in which emissions, chemical processes (GAS+Therm+HET) and physical processes 874 (HADV+VADV+HDIF+VDIF+DDEP) contributed 29.8 µg m<sup>-3</sup> h<sup>-1</sup>, 33.5 µg m<sup>-3</sup> h<sup>-1</sup> and -49.2 875  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>, respectively. In the persistence stage, chemical production rate of PM<sub>2.5</sub> changed 876 slightly, and the production and loss rates of PM<sub>2.5</sub> were similar, leading to an approximately 877 zero IPR in this stage (Figure 8b). In the dissipation stage, the contribution of vertical 878 diffusion and dry deposition to PM2.5 loss increased largely, while the total chemical 879 production rate decreased, which resulted in a net IPR of  $-34.8 \ \mu g \ m^{-3} \ h^{-1}$ , indicating a 880 substantial decrease in PM2.5 concentration (Figure 8b). It was also noticed that HADV 881 contributed to PM<sub>2.5</sub> production in this stage, which was due to mass import to Beijing from 882 upwind areas by northwesterlies. 883

It should be mentioned that the contribution of emission was unchanged because the monthly based emission inventory from MEIC was used, and the contribution of cloud process was generally negligible throughout the period because there was little cloud and precipitation during the study period.

888 We further use PA to interpret evolution processes of primary (BC) and secondary 889 (sulfate and nitrate) aerosols.

Black carbon is considered to be inert and chemical inactive, so it is governed solely by 890 physical processes. In the clean stage, BC production was contributed solely by emission (5.7 891 µg m<sup>-3</sup> h<sup>-1</sup>), whereas vertical diffusion and dry deposition contributed equally to BC loss (-2.7 892  $\mu g m^{-3} h^{-1}$ ), and other processes were negligible (Figure 8c). In the growth stage, the 893 contribution of vertical diffusion and dry deposition to BC loss decreased to -2.0 µg m<sup>-3</sup> h<sup>-1</sup> 894 and -1.7  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>, respectively, and the net rate of change was 0.7  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>, indicating a 895 rapid increase of BC concentration in this stage (Figure 8c). In the persistence stage, the loss 896 rate by vertical diffusivity and dry deposition further increased mainly due to the increased 897 BC concentration (Figure 8c). It is noticed that horizontal advection somewhat contributed to 898 the loss of BC ( $-0.7 \text{ ug m}^{-3} \text{ h}^{-1}$ ), which indicated an increasing outflow of BC to surrounding 899 areas. The IPR was near zero, indicating a balance of production and loss rate in this stage. In 900 the dissipation stage, BC loss via vertical diffusion and dry deposition processes increased 901 largely, mainly due to increasing wind speed and vertical diffusivity, and the net IPR became 902 -1.6  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>. This absolute value was larger than that in the growth stage (0.7  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>), 903 which indicated a faster decrease in BC concentration than the BC increase in the growth 904 905 stage (Figure 8c).

As for secondary aerosols, like sulfate, contribution from direct emission was near zero. 906 In the clean stage, gas chemistry (5.9  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>) was the predominant process for sulfate 907 production, and vertical diffusion contributed most to the loss (-5.2  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>) (Figure 8d). In 908 the growth stage, contribution from vertical diffusion was reduced to  $-3.9 \ \mu g \ m^{-3} \ h^{-1}$  mainly 909 due to the decreased vertical diffusivity (Figure 8f), whereas positive contribution from gas 910 chemistry increased to 6.6 µg m<sup>-3</sup> h<sup>-1</sup>, which was resulted from competitive processes. For 911 sulfate formation from gas chemistry (SO<sub>2</sub>+OH $\rightarrow$ H<sub>2</sub>SO<sub>4</sub>, followed by nucleation or 912 condensation into particulate phase), the oxidation of SO<sub>2</sub> to sulfate was weakened because of 913 914 decreasing OH radical due to increasing aerosol attenuation of solar radiation, however,  $SO_2$ increased due to weakened vertical diffusivity, leading to a slight net increase of sulfate 915 concentration compared with the clean stage. It is noteworthy that the sulfate production rate 916 from heterogeneous reactions increased to 2.7  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>, mainly due to the increases in SO<sub>2</sub>, 917 aerosol surfaces and RH (as well as aerosol water content). All the processes led to a net 918 sulfate production rate of 2.7  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>, in which chemistry played a predominant role (IPR 919

of 9.3  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>). In the persistence stage, the contribution of gas and heterogeneous 920 processes further increased to 7.4  $\mu$ g m<sup>-3</sup> h<sup>-1</sup> and 4.3  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>, indicating an increasing 921 sulfate production through chemical processes (Figure 8d). It is interesting to note that 922 vertical diffusion contributed more to sulfate loss than in the growth stage, which was mainly 923 due to the higher sulfate level than in the growth stage while vertical diffusivity coefficients 924 were almost the same. The net IPR in this stage was just 0.2  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>, which indicated an 925 approximate balance of production and loss. In the dissipation stage, increasing vertical 926 diffusivity was the dominant process for sulfate loss, and chemical contribution decreased. It 927 is noticed a positive contribution to sulfate from horizontal advection (IPR of 4.3  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>). 928 which was due to an import of sulfate from upwind areas of Beijing by northwesterly winds, 929 like those for PM<sub>2.5</sub> and BC. 930

For nitrate, in the clean stage, thermodynamic process (4.5  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>) was the largest 931 contributor to nitrate production (Figure 8e). During the growth stage, the contribution of 932 thermodynamic processes (10.2 µg m<sup>-3</sup> h<sup>-1</sup>) increased by over a factor of two and was larger 933 than the contribution from heterogeneous process (Figure 8e). The substantial increase in the 934 contribution of thermodynamic processes to nitrate production was due to the combined 935 effects of the increased level of nitrate precursors (HNO<sub>3</sub> and NH<sub>3</sub>) resulting from weakened 936 diffusivity and the increased RH along with the decreased air temperature, which were 937 favorable for gas to aerosol conversion. The contribution of heterogeneous reactions 938 increased as well due to the increased aerosol surface and relative humidity. The net rate of 939 nitrate change in this stage was 5.3  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>. In the persistence stage, the contribution from 940 heterogeneous reactions changed slightly while the contribution from thermodynamic process 941 somewhat reduced (Figure 8e). This is because more NH<sub>3</sub> was consumed to neutralize the 942 increased sulfate, leaving less NH<sub>3</sub> to react with HNO<sub>3</sub>, and thus producing fewer nitrate. The 943 near zero net IPR of nitrate in this stage also indicated a balance of production and loss. In 944 the dissipating stage, the contribution of chemical processes was almost the same as that in 945 the clean stage, while physical processes dominated the loss and the net IPR of nitrate (Figure 946 947 8e).

948

# 949 5.1.2 Haze evolution during 1–4 March

We also investigate another haze period of 1–4 March using PA (Figure 9). The hourly 950 IPRs by different processes are shown in Figure 9a. An apparent difference between this 951 episode and the first one was the positive IPRs of HADV during this episode, especially in 952 the growth stage from 21:00 (LST) on 1 March to 9:00 (LST) on 2 March, which indicated 953 that horizontal transport contributed to the haze formation. Another difference is that the 954 chemical processes, especially heterogeneous reactions contributed less to the PM<sub>2.5</sub> mass 955 during the persistence stage, such as from 10:00 (LST) on 2 March to 3:00 (LST) on 4 March, 956 which will be discussed below. 957

The IPRs for PM<sub>2.5</sub> and its components and meteorological variables averaged over each 958 stage during this episode are calculated and presented in Figure 9b to 9f. For BC (Figure 9c), 959 the most evident difference from the first haze episode occurred in the growth stage, in which 960 horizontal advection contributed 1.5 µg m<sup>-3</sup> h<sup>-1</sup> to BC production, which was comparable in 961 magnitude to the negative contributions from vertical diffusion and dry deposition (-1.3 µg 962 m<sup>-3</sup> h<sup>-1</sup>), suggesting the import of BC into Beijing from surrounding areas. The wind direction 963 in the south of Beijing at this stage was southerly and wind speed was about 2-3 m s<sup>-1</sup>, so the 964 transport of pollutants from southern Hebei apparently contributed to the increase of BC level 965 in Beijing. Differently, during the first haze event on 20-26 February, wind direction was 966 easterly, bringing less polluted air mass from the Bohai Sea and northern Tianjin, so 967 horizontal advection contributed less to BC in Beijing. This transport feature was also 968 reflected in the change rates of sulfate (Figure 9d), nitrate (Figure 9e) and PM<sub>2.5</sub> (Figure 9b) 969 concentrations. An observational study for the same haze period in Beijing (Ma et al., 2017) 970 also suggested the important role of regional transport from the south of Beijing in haze 971 formation. 972

For sulfate (Figure 9d), although chemical processes still contributed most to sulfate production in the growth stage (6.0  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>), it is noticed that gas chemistry (5.9  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>) accounted for most of the sulfate production, whereas contribution from heterogeneous reactions was smaller than that in the first haze episode mainly due to lower relative humidity. In the growth stage, the net IPR was 1.9  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>, 30% smaller than that for the first haze, indicating a weaker secondary aerosol formation during this haze episode. In the persistence stage, sulfate production from gas phase oxidation was almost balanced by the loss from dry 980 deposition and vertical diffusion, resulting in a net IRP of -0.1  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>, indicating a small 981 variation of sulfate concentration during this stage on average.

For nitrate, in the growth stage, it is of interest to note that heterogeneous reactions (5.5 982  $\mu g m^{-3} h^{-1}$ ) dominated over thermodynamic processes (2.7  $\mu g m^{-3} h^{-1}$ ) in nitrate formation, 983 which could be due to the low RH in this stage. Fountoukis and Nenes (2007) indicated that 984 nitrate aerosol is hardly formed in the ISORROPIA II model when RH is below 40%. The 985 average RH is about 37% during this haze episode, resulting in more nitrate formed by 986 heterogeneous reactions. The net IPR in the growth stage was 3.7 µg m<sup>-3</sup> h<sup>-1</sup>, approximately 987 30% smaller than that in the first haze episode. In the persistence stage when relative 988 humidity increased to 51%, nitrate formation via thermodynamic processes became important, 989 and due to competition, nitrate formation from heterogeneous reactions was reduced. 990

For PM<sub>2.5</sub> (Figure 9b), in the growth stage, the IPR of PM<sub>2.5</sub> concentration was 13.0 µg 991 m<sup>-3</sup> h<sup>-1</sup>, in which emission, chemical processes (GAS+Therm+HET) and physical processes 992 (HADV+VADV+HDIF+VDIF+DDEP) contributed 29.8 µg m<sup>-3</sup> h<sup>-1</sup>, 23.9 µg m<sup>-3</sup> h<sup>-1</sup> and -40.7 993  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>, respectively. It is noteworthy that horizontal advection process (HADV) 994 contributed 22.4  $\mu$ g m<sup>-3</sup> h<sup>-1</sup> to PM<sub>2.5</sub> production in this episode, which was comparable to the 995 total chemical production of 23.9  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>. This reveals the comparable contributions to 996 PM<sub>2.5</sub> in Beijing from local sources and regional transport during this haze episode. In the 997 persistence stage, because of the change in wind direction and lower wind speed, the regional 998 transport of PM<sub>2.5</sub> became weak. The IPRs were -4.0  $\mu$ g m<sup>-3</sup> h<sup>-1</sup> for HADV and 1.2  $\mu$ g m<sup>-3</sup> h<sup>-1</sup> 999 for VADV, respectively, which were obviously smaller than those in the first haze episode. In 1000 1001 the dissipation stage, physical processes except HADV all contributed to the loss of PM<sub>2.5</sub>. Compared with the first haze episode, the negative IPR of VADV decreased mainly due to the 1002 larger wind speeds in this episode, as more PM<sub>2.5</sub> was removed by VADV, the remaining 1003 PM<sub>2.5</sub> loss by vertical diffusion decreased, consequently a weakened VDIF. The positive IPR 1004 of HADV increased as well due to larger wind speed than that in the first episode in this 1005 1006 stage.

1007 The above process analyses reveal that for the first haze episode (20–26 February) in 1008 Beijing, local emissions and chemical processes were the main contributors to the formation 1009 and persistence of the haze event. However, for the second haze (1–4 March), regional transport or horizontal advection played a more important role in haze formation, with a similar magnitude to local emissions and chemical productions in the growth stage. In all, for both episodes, local emission, chemical reaction and horizontal advection were major processes contributing to  $PM_{2.5}$  increase, whereas vertical processes (diffusion, dry deposition and advection) were major processes for  $PM_{2.5}$  removal. As the pollution level increased, the contribution of secondary aerosols through chemical formation to  $PM_{2.5}$  increased apparently in Beijing.

1017

1018 5.2 Contributions of physical and chemical processes to the aerosol feedback

1019 5.2.1 The first haze episode (20–26 February)

Figure 10 shows the contributions of each process to the feedback-induced difference in the change rates of  $PM_{2.5}$  and its major components ( $\Delta IPR$ ) during the first haze episode (20– 26 February), which were derived from the difference between cases with and without aerosol radiative effects (FULL minus NoAerNoAFB).

1024 The definition of the four stages during haze evolution is the same as that in section 5.1.1. 1025 For BC (Figure 10b) in the clean stage, the aerosol feedback caused a decrease in vertical diffusion and advection (Figure 10e), leading to an increase in BC concentration with the 1026  $\Delta$ IPR of 0.40 µg m<sup>-3</sup> h<sup>-1</sup> from VDIF+VADV, concurrently, the feedback caused an increased 1027 loss of BC through horizontal diffusion (HDIF) and advection (HADV) and dry deposition 1028 (DDEP) due to the increased BC concentration, with the  $\Delta$ IPR of -0.39 µg m<sup>-3</sup> h<sup>-1</sup> from 1029 HADV+HDIF+DDEP (Figure 10b). The net  $\Delta$ IPR was near zero, which indicated a 1030 negligible feedback effect during the clean stage. In the growth stage, the feedback caused a 1031 pronounced decrease in vertical diffusivity, advection, as well as dry deposition velocity, 1032 1033 leading to apparent increases in BC level, with the contributions to AIPRs from VDIF, VADV, and DDEP being 0.50 µg m<sup>-3</sup> h<sup>-1</sup>, 0.50 µg m<sup>-3</sup> h<sup>-1</sup> and 0.20 µg m<sup>-3</sup> h<sup>-1</sup>, respectively (Figure 1034 10b). The increase in BC concentration consequently led to an increase in outflow via HADV 1035 and HDIF, with the  $\Delta$ IPRs of -0.63 µg m<sup>-3</sup> h<sup>-1</sup> and -0.12 µg m<sup>-3</sup> h<sup>-1</sup>, respectively, which tended 1036 to reduce BC concentration. The total effect by summing the processes exhibited a net 1037 positive  $\Delta$ IPR of 0.44 µg m<sup>-3</sup> h<sup>-1</sup>, which indicated an apparent increase in BC concentration 1038 due to the feedback. In the persistence stage, the sign of  $\Delta$ IPR for each process was the same 1039

as that in the growth stage, and the  $\Delta$ IPR by vertical processes (0.84 µg m<sup>-3</sup> h<sup>-1</sup> from 1040 VADV+VDIF+DDEP) was generally balanced by that of horizontal processes (-0.80 µg m<sup>-3</sup> 1041  $h^{-1}$  from HADV+HDIF) and led to a net  $\Delta$ IPR of 0.04 µg m<sup>-3</sup> h<sup>-1</sup> (Figure 10b), which 1042 1043 indicated the difference in the BC change rate between the FULL and NoAerNoAFB cases 1044 was small in this stage. In the dissipating stage, the  $\Delta$ IPRs were negative for all the processes except for HADV. This was because of the higher BC levels due to the feedback, which 1045 caused more BC to be removed than without feedback, although the vertical diffusion 1046 1047 coefficient was smaller due to the feedback. The positive  $\Delta$ IPR from HADV suggested the enhanced BC import into Beijing from upwind regions due to the feedback. The sum of these 1048 processes produced a net  $\Delta$ IPR of -1.20 µg m<sup>-3</sup> h<sup>-1</sup>, which indicated a larger decreasing rate of 1049 BC concentration (from haze to clean level) due to aerosol feedback in this stage. 1050

For sulfate (Figure 10c), in the clean stage, the feedback-induced changes were as small 1051 as those for BC. In the growth stage, besides the positive  $\Delta$ IPRs by VDIF, VADV and DDEP 1052 as those for BC, the most impressive feature was the larger contributions from GAS and HET, 1053 with the  $\Delta$ IPRs being 0.29 µg m<sup>-3</sup> h<sup>-1</sup> and 1.73 µg m<sup>-3</sup> h<sup>-1</sup>, respectively, much larger than those 1054 (0.11  $\mu$ g m<sup>-3</sup> h<sup>-1</sup> and 0.23  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>) in the clean stage because of the increased gas 1055 precursors, aerosol surfaces and RH due to the feedback effect, which enhanced chemical 1056 formation (Figure 10c, 10e). The sum of the  $\Delta$ IPRs by all the processes was 1.92 µg m<sup>-3</sup> h<sup>-1</sup>, 1057 indicating an apparent increase in sulfate concentration due to the feedback effect. In the 1058 persistence stage, the  $\Delta$ IPRs by GAS and HET increased. However, the  $\Delta$ IPR of VDIF 1059 became negative, which could be explained by the increased sulfate concentration due to 1060 aerosol feedback caused more sulfate to be removed through vertical diffusion, leading to a 1061 negative  $\Delta$ IPR of VDIF, although the vertical diffusion coefficient was reduced by the 1062 1063 feedback. In the dissipation stage, the  $\Delta$ IPR by HET decreased because the feedback-induced differences in the concentrations of precursors and aerosols became smaller. The large 1064 negative AIPR by VDIF indicated a larger decreasing rate in sulfate concentration from the 1065 1066 persistence to clean stages due to the feedback.

1067 For nitrate (Figure 10d), the feedback-induced IPR changes in the clean stage were 1068 similar to those for sulfate. In the growth stage, remarkable increases in nitrate formation 1069 from Thermo and HET processes occurred, with the  $\Delta$ IPRs of 3.30 µg m<sup>-3</sup> h<sup>-1</sup> and 0.50 µg m<sup>-3</sup>

h<sup>-1</sup>, respectively (Figure 10d). The increased gas precursors and RH due to the aerosol 1070 feedback reinforced chemical formation processes. In this stage, the overall  $\Delta$ IPR was 3.90 1071 µg m<sup>-3</sup> h<sup>-1</sup>, suggesting a faster increasing rate in nitrate concentration in consideration of 1072 aerosol feedback. In the persistence stage, the  $\Delta$ IPR by Thermo was smaller than that in the 1073 growth stage (Figure 10d). This could be explained that the apparent increase in sulfate 1074 concentration via HET and GAS due to the feedback (Figure 10c) in this stage consumed 1075 more ammonia, which inhibited the formation of nitrate ammonium via thermodynamic 1076 1077 processes. The net  $\Delta$ IPR by all the processes in this stage was near zero, which indicated that the radiative feedback exerted little effect on the change rate of nitrate concentration during 1078 this stage. In the dissipation stage, the attenuation of solar radiation by aerosols was 1079 weakened because of the decrease in aerosol concentration, meanwhile, the concentrations of 1080 1081 gas precursors (NO<sub>x</sub>) were elevated due to the feedback, the combined effect resulted in an increase of photochemical production of HNO3; in addition, RH was increased due to the 1082 feedback as well, as a result, nitrate formation via thermodynamic process was enhanced, 1083 leading to a positive  $\Delta$ IPR of 3.73 µg m<sup>-3</sup> h<sup>-1</sup> by Thermo in this stage. 1084

For PM<sub>2.5</sub>, the net  $\Delta$ IPR due to aerosol feedback in the clean stage was 0.30 µg m<sup>-3</sup> h<sup>-1</sup>, in 1085 which 1.22 µg m<sup>-3</sup> h<sup>-1</sup> was from chemical processes (GAS+Thermo+HET) and -0.90 µg m<sup>-3</sup> 1086 h<sup>-1</sup> from physical processes (HADV+VADV+HDIF+VDIF+DDEP) (Figure 10a). In the 1087 growth stage, the net  $\Delta$ IPR was 9.50 µg m<sup>-3</sup> h<sup>-1</sup>, which meant in every hour, approximate 9.50 1088 µg m<sup>-3</sup> of PM<sub>2.5</sub> mass was elevated in Beijing due to the feedback effect. The above 1089 feedback-induced difference in the change rate of  $PM_{2.5}$  ( $\Delta IPR$ ) resulted from a combined 1090 effect from chemical processes (7.27  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>) and physical processes (2.23  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>), 1091 which suggested that chemical processes contributed more to the PM<sub>2.5</sub> increase than physical 1092 1093 processes. However, it was noted that the increased contribution from chemical processes was related to increasing gas precursors, which was partly associated with physical processes. 1094 It was noteworthy that the positive  $\Delta$ IPRs were contributed by both chemical processes (GAS, 1095 Thermo and HET) and vertical movements (VADV, VDIF and DDEP) (Figure 10a). The sum 1096 of positive  $\Delta$ IPRs was 22.88 µg m<sup>-3</sup> h<sup>-1</sup>, in which 7.27 µg m<sup>-3</sup> h<sup>-1</sup> was from chemical 1097 processes and 15.61 µg m<sup>-3</sup> h<sup>-1</sup> from vertical movements. This suggested a larger 1098 feedback-induced PM<sub>2.5</sub> increase through vertical movements than via chemical processes. 1099

However, the outflow (HADV+HDIF) of PM2.5 was also enhanced due to the increased PM2.5 1100 level by aerosol feedback, producing a negative  $\Delta IPR$  (-13.38 µg m<sup>-3</sup> h<sup>-1</sup>), and partly 1101 offsetting the positive  $\Delta$ IPR (15.61 µg m<sup>-3</sup> h<sup>-1</sup>) by vertical movements, resulting in a net  $\Delta$ IPR 1102 of 2.23  $\mu$ g m<sup>-3</sup> h<sup>-1</sup> from all the physical processes. In the persistence stage, the sign of  $\Delta$ IPRs 1103 by different processes generally resembled those in the growth stage except that of VDIF 1104 whose  $\Delta$ IPR was negative, which indicated more removal though VDIF mainly due to the 1105 increased secondary aerosol concentrations by aerosol feedback. The net  $\Delta$ IPR by all the 1106 processes was 0.40  $\mu$ g m<sup>-3</sup> h<sup>-1</sup> in this stage, indicating a small influence of aerosol feedback 1107 on the change rate of  $PM_{2.5}$  concentration. In the dissipating stage (Figure 10a), the large 1108 negative AIPR from VDIF indicated more PM2.5 mass was removed via vertical diffusion 1109 while considering aerosol feedback, although the feedback induced a smaller vertical 1110 diffusivity coefficient. The net  $\Delta$ IPR of -24.60 µg m<sup>-3</sup> h<sup>-1</sup> indicated a larger decreasing rate of 1111 PM<sub>2.5</sub> concentration in the FULL case than in the NoAerNoAFB case. 1112

1113

#### 1114 5.2.2 The second haze episode (1–4 March)

1115 For BC in the second haze episode (1–4 March), the most obvious difference from the first episode was in the growth stage, in which the  $\Delta$ IPR by horizontal advection (HADV) 1116 was 0.70  $\mu$ g m<sup>-3</sup> h<sup>-1</sup> (Figure 11b). The radiative feedback led to a weakened vertical 1117 diffusivity and a decreased PBL height (Figure 11e), which favored the accumulation of BC 1118 and caused a positive  $\Delta$ IPR of 0.40 µg m<sup>-3</sup> h<sup>-1</sup> from VDIF. The wind direction in the growth 1119 stage was southerlies as discussed above, bringing aerosols from the south to Beijing. The 1120 1121 aerosol feedback enhanced BC concentration in source regions through reducing vertical diffusivity, leading to an increased northward flux of BC and a positive  $\Delta$ IPR from HADV. The 1122 1123 higher BC concentration due to the feedback via HADV and VDIF consequently led to an increase in BC outflow out of Beijing via vertical advection (VADV) and horizontal diffusion 1124 (HDIF), with the  $\Delta$ IPRs of -0.60 µg m<sup>-3</sup> h<sup>-1</sup> and -0.20 µg m<sup>-3</sup> h<sup>-1</sup>, respectively. In this stage, the 1125 net  $\Delta$ IPR of BC was 0.20 µg m<sup>-3</sup> h<sup>-1</sup>, in which 0.50 µg m<sup>-3</sup> h<sup>-1</sup> was from horizontal 1126 (HADV+HDIF) and -0.30 µg m<sup>-3</sup> h<sup>-1</sup> from vertical movements 1127 movements (VADV+VDIF+DDEP), indicating that the feedback effect strengthened the contribution of 1128 horizontal movements to surface BC concentration in Beijing. In the persistence stage (Figure 1129

1130 11b), the net  $\Delta$ IPR was also near zero (-0.02 µg m<sup>-3</sup> h<sup>-1</sup>), indicating that the BC change rate 1131 was merely affected by the feedback in this stage. In the dissipation stage (Figure 11b), the 1132  $\Delta$ IPRs were negative for all the processes except for VDIF. This could be attributed to the 1133 higher BC levels due to the feedback, which caused more BC to be removed than without 1134 feedback through these processes. The net  $\Delta$ IPR was -0.17 µg m<sup>-3</sup> h<sup>-1</sup>, the same as that in the 1135 growth stage, but with opposite sign.

For sulfate (Figure 11c), in the growth stage, different from the relatively large positive 1136  $\Delta$ IPR by chemical processes in the first haze episode, the feedback caused small IPR changes 1137 via chemical production because SO<sub>2</sub> concentration in this episode was lower than that in the 1138 first one and sulfate was mainly formed in upwind regions and transported to Beijing. 1139 Consequently, relatively large sulfate increases through HADV and VDIF in this episode. In 1140 this stage, the feedback caused a slight increase in sulfate concentration by GAS with  $\Delta$ IPR 1141 of 0.17  $\mu$ g m<sup>-3</sup> h<sup>-1</sup> due to slightly elevated precursors, however, because of the low relative 1142 humidity (mean RH was 38%) and competitive processes, heterogeneous reactions were 1143 depressed. In terms of physical processes, due to the feedback effect, horizontal transport 1144 (HADV) was strengthened ( $\Delta$ IPR of 1.0 µg m<sup>-3</sup> h<sup>-1</sup>) due to the increased sulfate concentration 1145 to the south of Beijing, meanwhile, the weakened vertical diffusivity caused an increase in 1146 sulfate concentration by VDIF and DDEP, with the  $\Delta$ IPRs of 1.0 µg m<sup>-3</sup> h<sup>-1</sup> and 0.57 µg m<sup>-3</sup> 1147 h<sup>-1</sup>, respectively, consequently, the outflow of sulfate out of Beijing was also increased via 1148 vertical advection (VADV) and horizontal diffusion (HDIF). The net  $\Delta$ IPR in the growth 1149 stage was 0.90 µg m<sup>-3</sup> h<sup>-1</sup>, indicating an apparent increase in sulfate concentration due to the 1150 feedback. In the persistence stage, the  $\Delta$ IPRs by GAS and HET changed slightly compared 1151 with those in the growth stage. The negative  $\Delta$ IPR by VDIF indicated more loss of sulfate by 1152 vertical diffusion while considering aerosol feedback. The net  $\Delta$ IPR in this stage was 0.02 µg 1153 m<sup>-3</sup> h<sup>-1</sup>, indicating a negligible feedback effect on sulfate change rate in this stage. In the 1154 dissipation stage, the feedback-induced higher sulfate concentration caused more removal of 1155 sulfate via physical processes except HADV, resulting in a net  $\Delta$ IPR of -0.64 µg m<sup>-3</sup> h<sup>-1</sup>. The 1156 positive  $\Delta$ IPR from HADV was due to the strengthened import from upwind areas due to the 1157 feedback. 1158

1159

9 For nitrate, in the growth stage, the feedback also induced an increase in nitrate

concentration via horizontal advection like sulfate (Figure 11d). The increases in gas 1160 precursors and aerosol surfaces due to the feedback enhanced nitrate formation, resulting in 1161 nitrate increases via Thermo and HET, with the  $\Delta$ IPRs of 0.88 µg m<sup>-3</sup> h<sup>-1</sup> and 0.46 µg m<sup>-3</sup> h<sup>-1</sup>, 1162 respectively. To the persistence stage, the chemical production of nitrate increased largely 1163 caused by the feedback, with the  $\Delta$ IPR of Thermo being 4.30 µg m<sup>-3</sup> h<sup>-1</sup>. The reason could be 1164 the low RH in the growth stage (38% shown in Figure 9f) left most of nitric acid remained in 1165 gas phase together with the increase in RH due to the feedback (13.2% shown in Figure 11e) 1166 1167 drove its conversion from gas to aerosol phase. Due to the enhanced thermodynamics production, nitrate formation via heterogeneous reactions was depressed in this stage. The 1168 increased nitrate concentration via Thermo led to larger removal via vertical diffusion, 1169 resulting in a negative  $\Delta$ IPR of -4.80 µg m<sup>-3</sup> h<sup>-1</sup> by VDIF, and a net  $\Delta$ IPR of -0.10 µg m<sup>-3</sup> h<sup>-1</sup>. 1170 In the dissipation stage, like that in the first haze episode, the reduced aerosol attenuation of 1171 solar radiation and increased RH induced by aerosol feedback led to an increase in nitrate via 1172 thermodynamic process, with the  $\Delta$ IPR of 1.80 µg m<sup>-3</sup> h<sup>-1</sup> by Thermo. Consequently, 1173 heterogeneous reactions were depressed due to competitive processes ( $\Delta$ IPR of -0.97 µg m<sup>-3</sup> 1174  $h^{-1}$  by HET). In this stage, because of the higher nitrate concentration, the feedback led to 1175 larger removal by vertical processes (the  $\Delta$ IPR of VADV+VDIF+DDEP was -3.23 µg m<sup>-3</sup> h<sup>-1</sup>), 1176 with a net  $\Delta$ IPR of -1.78 µg m<sup>-3</sup> h<sup>-1</sup>, similar to the  $\Delta$ IPR in the growth stage but with opposite 1177 sign. 1178

For PM<sub>2.5</sub> (Figure 11a), the net  $\Delta$ IPR due to aerosol feedback in the growth stage was 1179 2.40 μg m<sup>-3</sup>  $h^{-1}$ , with 1.40 µg  $m^{-3}$  $h^{-1}$ from physical processes 1180 (HADV+VADV+HDIF+VDIF+DDEP) and 1.0 µg m<sup>-3</sup> h<sup>-1</sup> from chemical processes 1181 (GAS+Thermo+HET), which indicated that the feedback-induced increase in PM<sub>2.5</sub> 1182 1183 concentration per hour was produced through larger contributions from physical processes than chemical processes in this episode. HADV contributed most to the PM2.5 increase (with 1184  $\Delta$ IPR of 10.20 µg m<sup>-3</sup> h<sup>-1</sup>), followed by VDIF (with  $\Delta$ IPR of 2.90 µg m<sup>-3</sup> h<sup>-1</sup>). As mentioned 1185 above, the weakened vertical diffusivity caused by the feedback enhanced aerosol 1186 concentrations in the entire BTH region, meanwhile, the feedback induced a southeast wind 1187 anomaly with a slight change in wind speed in the regions south of Beijing. The combined 1188 effect of the elevated aerosol concentrations and southeast wind anomaly brought more 1189

aerosols to Beijing. In the persistence stage, the feedback increased PM<sub>2.5</sub> concentration 1190 mainly through chemical processes, with the  $\Delta$ IPR of 6.05 µg m<sup>-3</sup> h<sup>-1</sup>, which was mainly 1191 resulted from the enhanced thermodynamic production of ammonium nitrate, and such 1192 increase in aerosol mass due to feedback led to more aerosols to be diffused than that without 1193 feedback, leading to the  $\Delta$ IPR of -7.30 µg m<sup>-3</sup> h<sup>-1</sup> by VDIF. It is noticed that the signs of the 1194  $\Delta$ IPRs by VDIF were opposite between the growth and persistence stages even though the 1195 vertical diffusivities were both decreased. In the growth stage, the PM<sub>2.5</sub> concentration was 1196 1197 gradually increasing, the effect of the weakened vertical diffusivity was dominated, resulting in a positive  $\Delta$ IPR by VDIF which favored further accumulation of aerosols; in the 1198 persistence stage, the aerosol concentration had already been elevated to a high level, the 1199 effect of higher concentration surpassed that of weakened vertical diffusivity due to the 1200 feedback and led to a negative  $\Delta$ IPR, which meant the feedback caused more loss of PM<sub>2.5</sub> 1201 via VDIF. In the persistence stage, the net  $\Delta$ IPR was 0.44 µg m<sup>-3</sup> h<sup>-1</sup>, in which -5.6 µg m<sup>-3</sup> h<sup>-1</sup> 1202 from physical processes and 6.05  $\mu$ g m<sup>-3</sup> h<sup>-1</sup> from chemical processes, which indicated the 1203 feedback-induced overall changes in the change rate of PM<sub>2.5</sub> concentration in this stage were 1204 relatively small. In the dissipating stage, the removal of PM<sub>2.5</sub> was enhanced by the feedback 1205 through all the processes except HADV mainly due to the increased PM2.5 concentration, the 1206 positive  $\Delta$ IPR by HADV was caused by the enhanced import from upwind areas due to the 1207 feedback. In this stage, the feedback effect enhanced the removal of PM<sub>2.5</sub>, which was 1208 reflected by the net negative  $\Delta$ IPR of -4.30 µg m<sup>-3</sup> h<sup>-1</sup>. 1209

The above analyses quantify the key processes contributing to the aerosol radiative 1210 1211 feedback in Beijing during the two haze episodes. In the growth stage of the first haze episode, the feedback-induced PM2.5 enhancement was attributed to the positive contributions 1212 1213 from chemical processes and vertical movements, but partly offset by the increased outflow of PM<sub>2.5</sub> via horizontal advection, resulting in a larger increase in PM<sub>2.5</sub> through chemical 1214 processes than that from physical processes. Differently, duirng the second haze episode, the 1215 feedback-induced PM<sub>2.5</sub> enhancement in the growth stage was larger by physical processes 1216 than that by chemical processes, and horizontal advection contributed most to the PM<sub>2.5</sub> 1217 enhancement. In all, the radiative feedback increased the cumulative rate of aerosols in the 1218 growth stage via promoting chemical formations, weakening vertical diffusions and/or 1219

enhancing regional transport by horizontal advection. For both episodes, the radiative feedback exerted small effect on the change rate of  $PM_{2.5}$  concentration during the persistence stage and reinforced the decreasing rate of  $PM_{2.5}$  in the dissipation stage.

1223

# 1224 6 Conclusions

Several severe haze events occurred in the winter of 2014, with the most severe one on 1225 20-26 February. An online-coupled regional atmospheric chemistry/aerosol-climate model 1226 (RIEMS-Chem) was developed and utilized to investigate the mechanisms of haze formation 1227 and aerosol radiative feedback in the Beijing-Tianjin-Hebei (BTH) region. The heterogeneous 1228 chemical reactions were treated in the model and the measured size distribution and mixing 1229 state of aerosols in Beijing were used to constrain the model. Two numerical experiments, 1230 with and without aerosol effects were conducted to explore the aerosol radiative effects 1231 (AREs) and feedbacks on meteorological fields and aerosol distributions. Processes analysis 1232 technique was implemented in RIEMS-Chem to quantify the individual contributions from 1233 1234 various physical and chemical processes to aerosol evolution and radiative feedback. Model performance was comprehensively evaluated by comparing with a variety of observations for 1235 meteorological variables, surface shortwave radiation, PBL heights, PM<sub>2.5</sub> and its chemical 1236 components, as well as aerosol optical properties in the BTH region. The comparisons 1237 demonstrated that RIEMS-Chem was able to represent the magnitudes and variations of the 1238 above variables reasonably well, in particular, improving the simulation of inorganic aerosols 1239 and AOD, which was often underpredicted in current on-line coupled models. It is 1240 1241 encouraging that by considering the aerosol radiative effects, the model apparently improved predictions for meteorological variables, PM<sub>2.5</sub> and its chemical compositions and aerosol 1242 optical properties in the BTH region, suggesting the importance and necessity for developing 1243 chemistry-climate online coupled models in both air quality and climate research. 1244

During the study period, the meteorological conditions were characterized by weak southerly winds, high RH and low PBL height, which favored aerosol accumulation and haze formation in the BTH region. The average T2, WS10, RH2, PBL height and  $PM_{2.5}$ concentration from the FULL case were simulated to be 0.6 °C, 1.2 m s<sup>-1</sup>, 67.0%, 698.4 m and 90.0  $\mu$ g m<sup>-3</sup>, respectively, over the BTH region during the study period.

The distribution pattern of AOD generally resembled that of PM<sub>2.5</sub>, with the domain 1250 mean value of 0.78 and the maximum up to 1.1 during the study period. It was noteworthy 1251 that the simulated SSA averaged over the BTH region and the study period was 0.91, which 1252 indicated the dominance of scattering aerosols. The domain and period average AREs at the 1253 surface, in the atmosphere and at the TOA were estimated to be -37 W m<sup>-2</sup>, 19 W m<sup>-2</sup> and -18 1254 W m<sup>-2</sup>, respectively, and they were enhanced to -57 W m<sup>-2</sup>, 25 W m<sup>-2</sup> and -32 W m<sup>-2</sup> during 1255 the most severe haze episode (20–26 February). It was striking that the maximum hourly 1256 AREs at the surface and at TOA reached -384 W m<sup>-2</sup> and -231 W m<sup>-2</sup> around noon time in the 1257 vicinity of Shijiazhuang during the first haze episode. The magnitude of the model simulated 1258 AREs during the haze episode in this study agreed favorably with previous observational 1259 based estimates. 1260

The aerosol radiative effects generally led to a reduction in surface air temperature in the 1261 entire domain with larger decrease in southern BTH (-1.2 °C to -2 °C), accompanied by an 1262 increase in RH2 (10% to 16%) and a decrease in PBL height (-240 m to -210 m). The 1263 1264 changes in these meteorological variables were strengthened during the severe haze episode. Noticeably, PM<sub>2.5</sub> concentrations were consistently increased over the BTH region due to the 1265 aerosol feedback, with the maximum average increase exceeding 33  $\mu$ g m<sup>-3</sup> (33%) in southern 1266 Hebei and portions of Beijing and Tianjin during the study period, and the maximum hourly 1267 increase was up to 372  $\mu$ g m<sup>-3</sup> (186%) in the vicinity of Shijiazhuang during the severe haze 1268 episode. In terms of domain and period average, the feedback-induced changes were -1.4 °C 1269 for T2, -0.04 m s<sup>-1</sup> for WS10, 8.7% for RH2, -3.3 m<sup>2</sup> s<sup>-1</sup> for vertical diffusion coefficient, 1270 -160.0 m (-19%) for PBL height and 20.0 µg m<sup>-3</sup> (29%) for PM<sub>2.5</sub> concentration. The 1271 magnitude of the above changes were enhanced during the severe haze episode, with the 1272 7-day mean changes in T2, WS10, RH2, PBL height and PM<sub>2.5</sub> concentration being -1.8 °C, 1273 -0.5 m s<sup>-1</sup>, 9.8%, -183.6 m (-31%) and 45.1 µg m<sup>-3</sup> (39%), respectively, which demonstrated 1274 the significant aerosol radiative feedback on PM<sub>2.5</sub> accumulation and haze formation. The 1275 changes in sulfate and nitrate concentrations were larger than that in BC concentration 1276 because secondary aerosols were increased not only by weakened vertical diffusivity but also 1277 by enhanced chemical reactions caused by the feedback. 1278

1279 The magnitude of the feedback effect varied remarkably during haze evolution. The 1280 absolute change in  $PM_{2.5}$  concentration caused by the feedback was largest in the persistence 1281 stage, followed by those in the growth stage and in the dissipating stage. In Beijing, the 1282 feedback-induced increases in  $PM_{2.5}$  concentration were 55 µg m<sup>-3</sup>, 84 µg m<sup>-3</sup>, 40 µg m<sup>-3</sup>, 1283 respectively, during the growth, persistence and dissipation stages of the severe haze episode.

1284 PA method was applied to calculate the IPRs for quantifying the individual contributions from physical and chemical processes to variations of PM2.5 and its chemical components 1285 during haze episodes in Beijing. Two haze episodes were analyzed and compared to elucidate 1286 the mechanism of haze formation and evolution. For the first haze episode, the net IPR for 1287  $PM_{2.5}$  was 14.1 µg m<sup>-3</sup> h<sup>-1</sup> in the growth stage, in which emissions, chemical processes and 1288 physical processes contributed 29.8  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>, 33.5  $\mu$ g m<sup>-3</sup> h<sup>-1</sup> and -49.2  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>, 1289 respectively, which indicated a remarkable PM<sub>2.5</sub> increase contributed by chemical processes 1290 1291 in this stage. The most influential processes for PM<sub>2.5</sub> loss and production were vertical diffusion and thermodynamic processes, respectively. Compared with the clean stage, the 1292 losses by vertical diffusion and dry deposition reduced largely, and the production by 1293 1294 chemical processes increased, both leading to an evident increase in surface PM<sub>2.5</sub> concentrations in the growth stage. In the persistence stage, the production and loss of PM<sub>2.5</sub> 1295 were almost equal, resulting in an approximately zero IPR in this stage. In the dissipation 1296 stage, the loss of PM<sub>2.5</sub> by vertical diffusion and dry deposition increased greatly, leading to a 1297 net IPR rate of -34.8  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>, which meant a substantial decrease in PM<sub>2.5</sub> concentration. 1298

For the second haze episode, the net IPR for  $PM_{2.5}$  was 13.0 µg m<sup>-3</sup> h<sup>-1</sup> in the growth stage, in which emissions, chemical processes and physical processes contributed 29.8 µg m<sup>-3</sup> h<sup>-1</sup>, 23.9 µg m<sup>-3</sup> h<sup>-1</sup> and -40.8 µg m<sup>-3</sup> h<sup>-1</sup>, respectively. It was noteworthy that the contribution of horizontal advection to  $PM_{2.5}$  was of a similar magnitude to the contributions from local emissions and chemical processes, with the mean IPR of 22.4 µg m<sup>-3</sup> h<sup>-1</sup>, which indicated the important contribution of regional transport to haze formation in Beijing. Process analysis for the changes in  $PM_{2.5}$  components during haze evolution was also conducted.

1306 The contribution of each physical and chemical process to the feedback-induced changes 1307 in PM<sub>2.5</sub> and its major components were explored and quantified. For the first haze episode, 1308 the fast increase in PM<sub>2.5</sub> ( $\Delta$ IPR of 9.5 µg m<sup>-3</sup> h<sup>-1</sup>) due to aerosol feedback in the growth stage

was mainly attributed to the changes in vertical movements (VDIF and VADV) and chemical 1309 processes, but the increased outflow via horizontal advection (HADV) partly offset the 1310 increased PM<sub>2.5</sub> due to vertical movements, which caused a larger contribution to the PM<sub>2.5</sub> 1311 increase from chemical processes ( $\Delta$ IPR of 7.27 µg m<sup>-3</sup> h<sup>-1</sup>) than that from physical processes 1312 ( $\Delta$ IPR 2.23 µg m<sup>-3</sup> h<sup>-1</sup>). However, during the second haze episode, the feedback-induced 1313 PM<sub>2.5</sub> increase ( $\Delta$ IPR of 2.4 µg m<sup>-3</sup> h<sup>-1</sup>) in the growth stage was mainly contributed by 1314 physical processes ( $\Delta$ IPR of 1.40 µg m<sup>-3</sup> h<sup>-1</sup>) rather than that by chemical processes ( $\Delta$ IPR of 1315 1.0  $\mu$ g m<sup>-3</sup> h<sup>-1</sup>), and among physical processes, the PM<sub>2.5</sub> increase was mainly attributed to the 1316 increased horizontal advection ( $\Delta$ IPR of 10.2 µg m<sup>-3</sup> h<sup>-1</sup>). In general, in the growth stage of 1317 haze episodes, the feedback increased the accumulation rate of aerosols mainly through 1318 enhancing chemical formations, weakening vertical diffusions and/or enhancing regional 1319 transport by advections. The feedback-induced changes in the change rate of PM<sub>2.5</sub> 1320 concentration were small during the persistence stage, and the feedback enhanced the 1321 removal rate of PM<sub>2.5</sub> in the dissipation stage mainly through increasing vertical diffusion 1322 and/or vertical advection. 1323

1324 The results from this study demonstrated a significant impact of aerosol radiative feedback on meteorology, chemistry, aerosol distribution and evolution during winter haze 1325 events in the BTH region. The mechanism and processes through which the feedback affected 1326 haze formation and evolution were elucidated and quantified. This study is still subject to 1327 some uncertainties: 1.) An internal mixing was assumed for aerosol mixing in this study, but 1328 the mixing state of aerosols is always changing, while this assumption is generally realistic 1329 for haze days, it may overestimate the feedback effect for clean days. 2.) A typical size 1330 distribution measured during haze days was used, whereas the size of aerosol internal mixture 1331 could change to some extent with aging processes. These uncertainties require further 1332 development of model treatment for evolution of aerosol mixing state and size distribution, 1333 which is poorly represented in current online coupled models. 3.) Direct aerosol radiative 1334 effect dominated the feedback effect in this study, so more cases in different regions and 1335 seasons, when indirect effect could be more important are needed to elucidate the complete 1336 feedback mechanism at different spatial and temporal scales. 4.) Finer model grid resolution 1337 is expected to be applied to look into details of the feedback effect at urban scale along with 1338

finer emission inventory (Tao et al., 2020), vertical observations (Wilcox et al., 2016; Wang et al., 2018) and higher computational efficiency when available in the future. Finally, this study pointed out the significance and necessity of developing online coupled model for exploring chemistry/aerosol-weather/climate interactions and for improving meteorological and chemical predictions in both air quality and climate research in the future.

1344

## 1345 Author Contributions

1346 ZH designed the study, JL performed the model simulation, JL and ZH processed and 1347 analyzed the modeling data, ZH and JL wrote the paper, JL and ZX contributed to the model 1348 development, YW provided and analyzed the chemical observation data, XX provided the 1349 meteorological sounding and aerosol optical observation data, JL and LL processed and 1350 analyzed the observational data, RZ synthesized and analyzed the observation.

1351

# 1352 Data availability.

- 1353 The observational data can be accessed through contacting the corresponding authors.
- 1354

## 1355 Competing interests.

- 1356 The authors declare that they have no conflict of interests.
- 1357

#### 1358 Special issue statement.

This article is part of the special issue "Regional assessment of air pollution and climate
change over East and Southeast Asia: results from MICS-Asia Phase III". It is not associated
with a conference.

1362

## 1363 Acknowledgement.

This study was supported by the National Natural Science Foundation of China (no. 91644217), the National Key R&D Program of China (2019YFA0606802) and the Jiangsu Collaborative Innovation Center for Climate Change.

1367

#### 1368 **References**

Albrecht, B.: Aerosols, cloud microphysics, and fractional cloudiness, Science, 245, 1227,
 https://doi.org/10.1126/science.245.4923.1227, 1989.

An, Z., Huang, R., Zhang, R., Tie, X., Li, G., Cao, J., Zhou, W., Shi, Z., Han, Y., Gu, Z., and
Ji, Y.: Severe haze in northern China: A synergy of anthropogenic emissions and

- 1373 atmospheric processes, P. Natl. Acad. Sci. USA, 116,18, 8657–8666, 2019.
- Baklanov, A., Schlunzen, K., Suppan, P., Baldasano, J., Brunner, D., Aksoyoglu, S., 1374 Carmichael, G., Douros, J., Flemming, J., Forkel, R., Galmarini, S., Gauss, M., Grell, G., 1375 Hirtl, M., Joffre, S., Jorba, O., Kaas, E., Kaasik, M., Kallos, G., Kong, X., Korsholm, U., 1376 Kurganskiy, A., Kushta, J., Lohmann, U., Mahura, A., Manders-Groot, A., Murizi, A., 1377 Moussiopoulos, N., Rao, S.T., Savage, N., Seigneur, C., Sokhi, R.S., Solazzo, E., 1378 Solomos, S., Sorenson, B., Tsegas, G., Vignati, E., Vogel, B., and Zhang, Y.: Online 1379 coupled regional meteorology chemistry models in Europe: current status and prospects, 1380 Atmos. Chem. Phys., 14, 317-398, doi:10.5194/acp-14-317-2014, 2014. 1381
- Beheng, K. D.: A parameterization of warm cloud microphysical conversion processes,
  Atmos. Res., 33, 193–206, 1994.
- Cai, W., Li, K., Liao, H., Wang, H., and Wu, L.: Weather conditions conducive to Beijing
  severe haze more frequent under climate change, Nature Clim. Change, 7, 257–262, doi:
  10.1038/nclimate3249, 2017.
- Carslaw, K. S., Boucher, O., Spracklen, D. V., Mann, G. W., Rae, J. G. L., Woodward, S., and
  Kulmala, M.: A review of natural aerosol interactions and feedbacks within the Earth
  system, Atmos. Chem. Phys., 10, 1701–1737, doi:10.5194/acp-10-1701-2010, 2010.
- 1390 Chan, C. and Yao, X.: Air pollution in megacities in China, Atmos. Environ., 42, 1-42, 2008.
- Che H., Xia, X., Zhu, J., Li, Z., Dubovik, O., Holben, B., Goloub, P., Chen, H., Estelles, V., 1391 Cuevas-Agulló, E., Blarel, L., Wang, H., Zhao, H., Zhang, X., Wang, Y., Sun, J., Tao, R., 1392 Zhang, X., and Shi, G.: Column aerosol optical properties and aerosol radiative forcing 1393 during a serious haze-fog month over North China Plain in 2013 based on ground-based 1394 measurements, Chem. 1395 sunphotometer Atmos. Phys., 14, 2125-2138, doi:10.5194/acp-14-2125-2014, 2014. 1396
- Chen, L., Zhu, J., Liao, H., Gao, Y., Qiu, Y., Zhang, M. G., Liu, Z. R., Li, N., and Wang, Y. S.:
  Assessing the formation and evolution mechanisms of severe haze pollution in the
  Beijing–Tianjin–Hebei region using process analysis, Atmos. Chem. Phys., 19, 10845–
  10864, 2019.
- Cheng, Y., Zheng, G., Wei, C., Mu, Q., Zheng, Bo., Wang, Z., Gao, M., Zhang, Q., He, K.,
  Carmichael, G., Pösch, U., and Su, H.: Reactive nitrogen chemistry in aerosol water as a
  source of sulfate during haze events in China, Sci. Adv., 2, e1601530, 2016.
- Dawson, J. P., Adams, P. J., and Pandis, S. N.: Sensitivity of PM<sub>2.5</sub> to climate in the Eastern
  US: a modeling case study, Atmos. Chem. Phys., 7, 4295–4309, 2007.
- Dickinson, R.E., Henderson-Sellers, A., and Kennedy, P.J.: Biosphere-Atmosphere Transfer
  Scheme (BATS) Version 1e as coupled to NCAR Community Climate Model, NCAR
  Technical Note, NCAR/TN-387+STR, p.72, 1993.
- Ding, A., Huang, X., Nie, W., Sun, J., Kerminen, V. M., Petäjä, T., Su, H., Cheng, Y., Yang,
  X., Wang, M., Chi, X., Wang, J., Virkkula, A., Guo, W., Yuan, J., Wang, S., Zhang, R.,
  Wu, Y., Song, Y., Zhu, T., Zilitinkevich, S., Kulmala, M., and Fu, C.: Enhanced haze
  pollution by black carbon in megacities in China, Geophys. Res. Lett., 43, 2873-2879,
  10.1002/2016gl067745, 2016.
- Emmons, L. K., Walters, S., Hess, P. G., Lamarque, J. F., Pfister, G. G., Fillmore, D., Granier,
  C., Guenther, A., Kinnison, D., Laepple, T., Orlando, J., Tie, X., Tyndall, G.,
  Wiedinmyer, C., Baughcum, S. L., and Kloster, S.: Description and evaluation of the

- Model for Ozone and Related chemical Tracers, version 4 (MOZART-4), Geosci. Model.
  Dev., 3, 43-67, doi:10.5194/gmd-3-43-2010, 2010.
- Forkel, R., Werhahn, J., Hansen, A. B., McKeen, S., Peckham, S., Grell, G., and Suppan, P.:
  Effect of aerosol-radiation feedback on regional air quality a case study with
  WRF/Chem, Atmos. Environ., 53, 202–211, 2012.
- 1422Fountoukis, C. and Nenes, A.: ISORROPIA II: a computationally efficient thermodynamic1423equilibrium model for  $K^+-Ca^{2+}-Mg^{2+}-NH_4^+-Na^+-SO_4^{2-}-NO_3^--Cl^--H_2O$  aerosols,1424Atmos. Chem. Phys., 7, 4639-4659, 2007.
- Fu, C. B., Wang, S. Y., Xiong, Z., Gutowski, W. J., Lee, D., Mcgregor, J. L., Sato, Y., Kato,
  H., Kim, J., and Suh, M.: Regional climate model intercomparison project for Asia, Bull.
  Amer. Meteor. Soc., 86, 257-266, 2005.
- Fu, H. and Chen, J.: Formation, features and controlling strategies of severe haze-fog
  pollutions in China, Sci. Total. Environ., 578, 121–138, 2017.
- Gao, M., Carmichael, G. R., Wang, Y., Saide, P. E., Yu, M., Xin, J., Liu, Z., and Wang, Z.:
  Modeling study of the 2010 regional haze event in the North China Plain, Atmos. Chem.
  Phys., 16, 1673-1691, 10.5194/acp-16-1673-2016, 2016.
- Gao, M., Han, Z., Liu, Z., Li, M., Xin, J., Tao, Z., Li, J., Kang, J., Huang, K., Dong, X.,
  Zhuang, B., Li, S., Ge, B., Wu, Q., Cheng, Y., Wang, Y., Lee, H., Kim, C., Fu, J. S.,
  Wang, T., Chin, M., Woo, J., Zhang, Q., Wang, Z., and Carmichael G. R.: Air Quality
  and Climate Change, Topic 3 of the Model Inter-Comparison Study for Asia Phase III
  (MICS-Asia III), Part I: overview and model evaluation, Atmos. Chem. Phys., 18, 4859–
  4884, https://doi.org/10.5194/acp-18-4859-2018, 2018.
- Gao, M., Han, Z. W., Tao, Z. N., Li, J. W., Kang, J.-E., Huang, K., Dong, X. Y., Zhuang, B. L.,
  Li, S., Ge, B. Z., Wu, Q. Z., Lee, H.-J., Kim, C. H., Fu, J. S., Wang, T. J., Chin, M., Li,
  M., Woo, J.-H., Zhang, Q., Cheng, Y. F., Wang, Z. F., and Carmichael, G. R.: Air quality
  and climate change, Topic 3 of the Model Inter-Comparison Study for Asia Phase III
  (MICS-Asia III) Part 2: aerosol radiative effects and aerosol feedbacks. Atmospheric
  Chemistry and Physics, 20, 1147–1161, 2020.
- Gao, Y., Zhang, M., Liu, Z., Wang, L., Wang, P., Xia, X., Tao, M., and Zhu, L.: Modeling the
  feedback between aerosol and meteorological variables in the atmospheric boundary
  layer during a severe fog-haze event over the North China Plain, Atmos. Chem. Phys.,
  148 15, 4279-4295, 10.5194/acp-15-4279-2015, 2015.
- Gery, M.W., Whitten, G.Z., Killus, J.P., and Dodge, M.C.: A photochemical kinetics
  mechanism for urban and regional scale computer modeling, J. Geophys. Res., 94,
  12925-12956, 1989.
- Ghan, S. and Zaveri R.A.: Parameterization of optical properties for hydrated internally
  mixed aerosol, J. Geophys. Res., 112, D10201, doi:10.1029/2006JD007927, 2007.
- Giglio, L., Randerson, J. T., and van der Werf, G. R.: Analysis of daily, monthly, and annual
  burned area using the fourth generation Global Fire Emissions Database (GFED4), J.
  Geophys. Res.: Biogeosciences, doi:10.1002/jgrg.20042, 2013.
- Grell, G. A.: Prognostic evaluation of assumptions used by cumulus parameterizations, Mon.
  Weather. Rev., 121, 764-787, 1993.
- Guo, S., Hu, M., Zamora, M. L., Peng, J., Shang, D., Zheng, J., Du, Z., Wu, Z., Shao, M.,
  Zeng, L., Molina, M. J., and Zhang, R. : Elucidating severe urban haze formation in

- 1461
   China, P.
   Natl.
   Acad.
   Sci.
   USA.,
   111,
   17373–17378,

   1462
   https://doi.org/10.1073/pnas.1419604111, 2014.

   17373–17378,
- Han, X., Zhang, M. G., Han, Z. W., Xin, J. Y., and Liu, X. H.: Simulation of aerosol direct
  radiative forcing with RAMS-CMAQ in East Asia, Atmos. Environ., 45, 6576-6592,
  2011a.
- Han, Z. W., Ueda, H., Matsuda, K., Zhang, R. J., Arao, K., Kanai, Y., and Hasome, H.: Model
  study on particle size segregation and deposition during Asian dust events in March
  2002, J. Geophys. Res., 109, D19205, doi: 10.1029/2004jd004920, 2004.
- Han, Z. W.: Direct radiative effect of aerosols over East Asia with a Regional coupled
  Climate/Chemistry model, Meteorol. Z., Vol. 19, No. 3, 287-298, 2010.
- Han, Z. W., Xiong, Z., and Li, J. W.: Direct climatic effect of aerosols and interdecadal
  variations over East Asia investigated by a regional climate/chemistry model,
  Atmospheric and Oceanic Science Letters, 4(6), 299-303, 2011b.
- Han, Z. W., Li, J. W., Xia, X. A., and Zhang, R. J.: Investigation of direct radiative effects of
  aerosols in dust storm season over East Asia with an online coupled regional
  climate-chemistry-aerosol model, Atmos. Environ., 54, 688-699, 2012.
- Han, Z. W., Li, J. W., Guo, W. D., Xiong, Z., and Zhang, W.: A study of dust radiative
  feedback on dust cycle and meteorology over East Asia by a coupled regional
  climate-chemistry-aerosol model, Atmos. Environ., 68, 54-63, 2013.
- Han, Z. W., Li, J. W., Yao, X. H., and Tan, S. C.: A regional model study of the characteristics
  and indirect effects of marine primary organic aerosol in springtime over East Asia,
  Atmos. Environ., 197, 22–35, 2019.
- Hegg D. A.: Cloud condensation nucleus-sulfate mass relationship and cloud albedo. J.
  Geophys. Res. Atmos., 99, D12, 25903-25907, 1994.
- Hess, M., Koepke, P., and Schuit, I.: Optical properties of aerosols and clouds: the software
  package OPAC, Bull. Amer. Meteor. Soc., 79, 831-844 ,1998.
- Heo, B.-H., Jacoby-Koaly, S., Kim, K.-E., Campistron, B., Benech, B., and Jung, E.-S.: Use
  of the Doppler Spectral Width to Improve the Estimation of the Convective Boundary
  Layer Height from UHF Wind Profiler Observations, J. Atmos. Ocean. Technol., 20,
  408-424, 2003.
- Hong, S. and Pan, H.: Nonlocal boundary layer vertical diffusion in a medium-range forecast
  model, Mon. Weather Rev., 124, 2322–2339, 1996.
- Huang, R. J., Zhang, Y., Bozzetti, C., Ho, K. F., Cao, J. J., Han, Y., Daellenbach, K. R.,
  Slowik, J. G., Platt, S. M., Canonaco, F., Zotter, P., Wolf, R., Pieber, S. M., Bruns, E. A.,
  Crippa, M., Ciarelli, G., Piazzalunga, A., Schwikowski, M., Abbaszade, G.,
  Schnelle-Kreis, J., Zimmermann, R., An, Z., Szidat, S., Baltensperger, U., El Haddad, I.,
  and Prevot, A. S.: High secondary aerosol contribution to particulate pollution during
  haze events in China, Nature, 514, 218-222, 10.1038/nature13774, 2014.
- Huang, X., Wang, Z. L., Ding, A. J.: Impact of Aerosol-PBL Interaction on Haze Pollution:
  Multiyear Observational Evidences in North China, Geophys. Res. Lett., 45, 8596-8603,
  2018.
- Isaksen I.S.A., Granier, C., Myhre, G., Berntsen, T.K., Dalsøren, S.B., Gauss, M., Klimont, Z.,
  Benestad, R., Bousquet, P., Collins, W., Cox, T., Eyring, V., Fowler, D., Fuzzi, S.,
  Jo<sup>°</sup>ckel, P., Laj, P., Lohmann, U., Maione, M., Monks, P., Prevot, A.S.H., Raes, F.,

- Richter, A., Rognerud, B., Schulz, M., Shindell, D., Stevenson, D.S., Storelvmo, T.,
  Wang, W.-C., van Weele, M., Wild, M., and Wuebbles, D.: Atmospheric composition
  change: Climate–Chemistry interactions, Atmos. Environ., 43, 5138–5192, 2009.
- Jing, J., Wu, Y., Tao, J., Che, H.Z, Xia, X., Zhang, X., Yan, P., Zhao, D.M., and Zhang, L.M.:
  Observation and analysis of near-surface atmospheric aerosol optical properties in urban
  Beijing, Particuology, 18, 144-154, 2015.
- Kajino, M., Ueda, H., Han, Z. W., Kudo, R., Inomata, Y., Kaku, H.: Synergy between air
  pollution and urban meteorological changes through aerosol-radiation-diffusion
  feedback—A case study of Beijing in January 2013, Atmos. Environ., 171, 98-110,
  2017.
- Keihl, J.T., Hack, J.J., Bonan, G.B., Boville, B.A., Briegleb, B.P., Williamson, D.L., and
  Rasch, P.J.: Description of the NCAR Community Climate Model (CCM3), NCAR
  Technical Note, NCAR/TN-420+STR, p.152, 1996.
- Lee-Taylor, J., Madronich, S.: Climatology of UV-A, UV-B, and Erythemal Radiation at the
  Earth's Surface, 1979-2000, NCAR Technical Note, NCAR/TN-474+STR, pp 1-52.
  2007.
- Li, G.H., Bei, N.F., Cao, J.J., Huang, R.J., Wu, J.R., Feng, T., Wang, Y.C., Liu, S.X., Zhang,
  Q., Tie, X.X., and Molina, L.T.: A possible pathway for rapid growth of sulfate during
  haze days in China, Atmos. Chem. Phys., 17, 3301-3316, 2017c.
- Li, J., Chen, X.S., Wang, Z.F., Du, H.Y., Yang, W.Y., Sun, Y.L., Hu, B., Li, J.J., Wang, W.,
  Wang, T., Fu, P.Q., and Huang, H.L.: Radiative and heterogeneous chemical effects of
  aerosols on ozone and inorganic aerosols over East Asia, Sci. Total. Environ., 622–623,
  1327–1342, 2018b.
- Li, J. W. and Han, Z. W.: A modeling study of the impact of heterogeneous reactions on
  mineral aerosol surfaces on tropospheric chemistry over East Asia, Particuology, 8,
  433-441, 2010.
- Li, J. W., Han, Z. W., and Zhang, R. J.: Influence of aerosol hygroscopic growth
  parameterization on aerosol optical depth and direct radiative forcing over East Asia,
  Atmos. Res., 140-141, 14-27, 2014.
- Li, J. W. and Han, Z. W.: A modeling study of severe winter haze events in Beijing and its neighboring regions, Atmos. Res., 170, 87–97, 2016a.
- Li, J. W. and Han, Z. W.: Aerosol vertical distribution over east China from RIEMS-Chem
  simulation in comparison with CALIPSO measurements, Atmos. Environ., 143, 177-189,
  2016b.
- Li, J. W. and Han, Z. W.: Seasonal variation of nitrate concentration and its direct radiative
  forcing over East Asia, Atmosphere, 7(8), 105, 2016c.
- Li, J. W., Han, Z. W., and Yao, X. H.: A modeling study of the influence of sea salt on
  inorganic aerosol concentration, size distribution, and deposition in the western Pacific
  Ocean, Atmos. Environ., 188, 157-173, 2018a.
- Li, J. W., Han, Z. W., Yao, X. H., Xie, Z. X., and Tan, S. C.: The distributions and direct radiative effects of marine aerosols over East Asia in springtime, Sci. Total. Environ., 651, 1913–1925, 2019b.
- Li, M., Zhang, Q., Kurokawa, J.-I., Woo, J.-H., He, K., Lu, Z., Ohara, T., Song, Y., Streets, D.
  G., Carmichael, G. R., Cheng, Y., Hong, C., Huo, H., Jiang, X., Kang, S., Liu, F., Su, H.,

- and Zheng, B.: MIX: a mosaic Asian anthropogenic emission inventory under the
  international collaboration framework of the MICS-Asia and HTAP, Atmos. Chem.
  Phys., 17, 935-963, 10.5194/acp-17-935-2017, 2017a.
- Li, X., Wu, J., Elser, M., Tong, S., Liu, S., Li, X., Liu, L., Cao, J., Zhou, J., El-Haddad, I.,
  Huang, R., Ge, M., Tie, X., André S. H. Prévôt, and Li, G.: Wintertime secondary
  organic aerosol formation in Beijing–Tianjin–Hebei (BTH): contributions of HONO
  sources and heterogeneous reactions, Atmos. Chem. Phys., 19, 2343–2359,
  https://doi.org/10.5194/acp-19-2343-2019, 2019a.
- Li, Z., Guo, J., Ding, A., Liao, H., Liu, J., Sun, Y., Wang, T., Xue, H., Zhang, H., and Zhu, B.:
  Aerosol and boundary-layer interactions and impact on air quality, Natl. Sci. Rev., 4,
  810–833, 10.1093/nsr/nwx117, 2017b.
- Liu, X. H. and Wang, J.: How important is organic aerosol hygroscopicity to aerosol indirect forcing? Environ. Res. Lett., 5(4), 044010, http://iopscience.iop.org/1748-9326/5/4/044010, 2010a.
- Liu, X. H., Zhang, Y., Xing, J., Zhang, Q., Wang, K., Streets, D., Jang, C., Wang, W.-X., and
  Hao, J.-M.: Understanding of regional air pollution over China using CMAQ, part II.
  Process analysis and sensitivity of ozone and particulate matter to precursor emissions,
  Atmos. Environ., 44, 3719-3727, 2010b.
- Ma, N., Zhao, C. S., Müller, T., Cheng, Y. F., Liu, P. F., Deng, Z. Z., Xu, W. Y., Ran, L., Nekat,
  B., van Pinxteren, D., Gnauk, T., Müller, K., Herrmann, H., Yan, P., Zhou, X.J., and
  Wiedensohler, A.: A new method to determine the mixing state of light absorbing
  carbonaceous using the measured aerosol optical properties and number size
  distributions, Atmos. Chem. Phys., 12, 2381–2397, 2012.
- Ma, Q. X., Wu, Y. F., Zhang, D. Z., Wang, X. J., Xia, Y. J., Liu, X. Y., Tian, P., Han, Z. W.,
  Xia, X. A., Wang, Y., and Zhang, R. J.: Roles of regional transport and heterogeneous
  reactions in the PM<sub>2.5</sub> increase during winter haze episodes in Beijing, Sci. Total.
  Environ., 599–600, 246–253, 2017.
- Martin, G. M., Johnson, D. W., and Spice, A.: The Measurements and Parameterization of
  Effective Radius of Droplets in Warm Stratocumulus Clouds, J. Atmos. Sci., 51, 1823–
  1842, 1994.
- NOAA/NCEP.: NCEP FNL Operational Model Global Tropospheric Analyses, continuing
   from July 1999. Research Data Archive at the National Center for Atmospheric Research,
   Computational and Information Systems Laboratory, Dataset.
   https://doi.org/10.5065/D6M043C6, Last accessed, 12 April 2019, 2000.
- Odum, J.R., Jungkamp, T.P.W., Griffin, R.J., Flagan, R.C., and Seinfeld, J.H.: The
  atmospheric aerosol-forming potential of whole gasoline vapor, Science. 276, 96–99,
  1997.
- Petters, M.D. and Kreidenweis, S.M.: A single parameter representation of hygroscopic
  growth and cloud condensation nucleus activity, Atmos. Chem. Phys., 7, 1961-1971,
  2007.
- Qiu, Y., Liao, H., Zhang, R., and Hu, J.: Simulated impacts of direct radiative effects of scattering and absorbing aerosols on surface layer aerosol concentrations in China during a heavily polluted event in February 2014, J. Geophys. Res. Atmos., 122, 5955-5975, 10.1002/2016jd026309, 2017.

- Ramanathan, V., Crutzen, P. J., Kiehl, J. T., and Rosenfeld, D.: Aerosols, climate, and the
  hydrological cycle, Science, 294, 2119-2124, 10.1126/science.1064034, 2001.
- Riemer, N., West, M., Zaveri, R., and Easter, R.: Estimating black carbon aging time-scales
  with a particle-resolved aerosol model, J. Aerosol Sci., 41, 143-158, 2010.
- Song, Z.J., Fu, D.S., Zhang, X.L., Wu, Y.F., Xia, X.A., He, J.X., Han, X.L., Zhang, R.J., and
  Che, H.Z.: Diurnal and seasonal variability of PM<sub>2.5</sub> and AOD in North China plain:
  Comparison of MERRA-2 products and ground measurements, Atmos. Environ., 191,
  70-78, 2018.
- Sun, Y., Jiang, Q., Wang, Z., Fu, P., Li, J., Yang, T., and Yin, Y.: Investigation of the Sources
  and Evolution Processes of Severe Haze Pollution in Beijing in January 2013, J.
  Geophys. Res., 119, 4380–4398, 2014.
- Tao, Z. N., Chin, M., Gao, M., Kucsera, T., Kim, D.-C., Bian, H. S., Kurokawa, J.-I., Wang, Y.
  S., Liu, Z. R., Carmichael, G. R., Wang, Z. F., and Akimoto, H.: Evaluation of NU-WRF
  model performance on air quality simulation under various model resolutions an
  investigation within the framework of MICS-Asia Phase III. Atmos. Chem. Phys., 20,
  2319–2339, 2020.
- 1609 Twomey, S.: Pollution and the planetary albedo, Atmos. Environ., 8, 1251–1256, 1974.
- Wang, J., Wang, S., Jiang, J., Ding, A., Zheng, M., Zhao, B., Wong, D. C., Zhou, W., Zheng,
  G., and Wang, L.: Impact of aerosol-meteorology interactions on fine particle pollution
  during China's severe haze episode in January 2013, Environ. Res. Lett., 9, 094002,
  https://doi.org/10.1088/1748-9326/9/9/094002, 2014b.
- Wang, S. Y., Fu, C. B., Wei, H. L., Qian, Y., Xiong, Z., Feng, J.M., Zhao, D. M., Dan, L., Han,
  Z.W., Su, B.K., Zhao, M., Zhang, Y.C., Tang, J.P., Liu, H.N., Wu, J., Zeng, X.M., Chen,
  M., and Wang, L.Z.: Regional integrated environmental modeling system: development
  and application, Climate Change, 129, 499-510, 2015.
- Wang, Y., Bao, S., Wang, S., Hu, Y., Shi, X., Wang, J., Zhao, B., Jiang, J., Zheng, M., Wu, M.,
  Russell, A. G., Wang, Y., and Hao, J.: Local and regional contributions to fine particulate
  matter in Beijing during heavy haze episodes, Sci. Total. Environ., 580, 283-296,
  10.1016/j.scitotenv.2016.12.127, 2017.
- Wang, Z. L., Huang, X., Ding, A. J.: Dome effect of black carbon and its key influencing
  factors: a one-dimensional modelling study, Atmos. Chem. Phys., 18, 2821–2834, 2018.
- Wang, Z., Li, J., Wang, Z., Yang, W., Tang, X., Ge, B., Yan, P., Zhu, L., Chen, X., Chen, H.,
  Wand, W., Li, J., Liu, B., Wang, X., Wand, W., Zhao, Y., Lu, N., and Su, D.: Modeling
  study of regional severe hazes over mid-eastern China in January 2013 and its
  implications on pollution prevention and control, Sci. China. Earth. Sci., 57, 3-13,
  10.1007/s11430-013-4793-0, 2014a.
- Westervelt, D. M., Moore, R. H., Nenes, A. and Adams, P.J.: Effect of primary organic sea
  spray emissions on cloud condensation nuclei concentrations, Atmos. Chem. Phys., 12,
  89–101, 2012.
- Wilcox, E. M., Thomas, R. M., Praveen, P. S., Pistone, K., Bender, F. A., and Ramanathan, V.:
  Black carbon solar absorption suppresses turbulence in the atmospheric boundary layer.
  Proc. Natl. Acad. Sci. USA, 113(42), 11794–11799, 2016.
- Wu, J., Bei, N., Hu, B., Liu, S., Zhou, M., Wang, Q., Li, X., Liu, L., Feng, T., Liu, Z., Wang,
  Y., Cao, J., Tie, X., Wang J., Molina, L.T., and Li, G.: Aerosol-radiation feedback

- deteriorates the wintertime haze in the North China Plain, Atmos. Chem. Phys., 19,
  8703–8719, https://doi.org/10.5194/acp-19-8703-2019, 2019.
- Wu, Y. F., Zhang, R. J., Tian, P., Tao, J., Hsu, S.-C., Yan, P., Wang, Q. Y., Cao, J. J., Zhang, X.
  L., and Xia, X. A.: Effect of ambient humidity on the light absorption amplification of
  black carbon in Beijing during January 2013, Atmos. Environ., 124, 217-223, 2016.
- Xiong, Z., Fu, C. B., and Yan, X. D.: Regional Integrated environmental model system and its
   simulation of East Asia summer monsoon, Chinese. Sci. Bull., 54(22), 4253-4261, 2009.
- Yu, S., Mathur, R., Schere, K., Kang, D., Plein, J., Young, J., Tong, D., Pouliot, G., Mckeen,
  S.A., and Rao, S. T.: Evaluation of real-time PM<sub>2.5</sub> forecasts and process analysis for
  PM<sub>2.5</sub> formation over the eastern United States using the Eta-CMAQ forecast model
  during the 2004 ICARTT study, J. Geophys. Res., 113, D06204, doi:
  10.1029/2007JD009226, 2008.
- Zhang, B., Wang, Y., and Hao, J.: Simulating aerosol-radiation-cloud feedbacks on meteorology and air quality over eastern China under severe haze conditions in winter, Atmos. Chem. Phys., 15, 2387-2404, 10.5194/acp-15-2387-2015, 2015.
- Zhang, X. Y., Wang, Y. Q., Niu, T., Zhang, X. C., Gong, S. L., Zhang, Y. M., and Sun, J. Y.:
  Atmospheric aerosol compositions in China: spatial/temporal variability, chemical
  signature, regional haze distribution and comparisons with global aerosols, Atmos.
  Chem. Phys., 12, 779–799, https://doi.org/10.5194/acp-12-779-2012, 2012.
- Zhang, X., Zhong, J., Wang, J., Wang, Y., and Liu, Y.: The interdecadal worsening of weather
  conditions affecting aerosol pollution in the Beijing area in relation to climate warming,
  Atmos. Chem. Phys., 18, 5991–5999, https://doi.org/10.5194/acp-18-5991-2018, 2018a.
- Zhang, X., Zhang, Q., Hong, C., Zheng, Y., Geng, G., Tong, D., Zhang, Y. and Zhang, X.:
   Enhancement of PM<sub>2.5</sub> Concentrations by Aerosol-Meteorology Interactions Over China,
   J. Geophys. Res. Atmos., 123, https://doi.org/10.1002/2017JD027524, 2018b.
- Zhang, Y.: Online coupled meteorology and chemistry models: history, current status, andoutlook, Atmos. Chem. Phys., 8, 2895-2932,2008.
- Zhang, Y., Wen, X.-Y., Wang, K., Vijayaraghavan, K., and Jacobson, M. Z.: Probing into
  regional O<sub>3</sub> and PM pollution in the U.S., PartII. An examination of formation
  mechanisms through a process analysis technique and sensitivity study, J. Geophys.
  Res.,114 (D22305), doi:10.1029/2009JD011900, 2009.
- Zhang, Y., Wen, X. Y., and Jang, C. J.: Simulating chemistry-aerosol-cloud-radiation-climate
   feedbacks over the continental U.S. using the online-coupled weather research
   forecasting model with chemistry (WRF/Chem), Atmos. Environ., 44, 3568–3582, 2010.
- Zhao, B., Liou, K. N., Gu, Y., Li, Q., Jiang, J. H., Su, H., He, C., Tseng, H. R., Wang, S., Liu,
  R., Qi, L., Lee, W. L., and Hao, J.: Enhanced PM<sub>2.5</sub> pollution in China due to
  aerosol-cloud interactions, Sci. Rep., 7, 4453, 10.1038/s41598-017-04096-8, 2017.
- Zhao, D. M.: Performance of Regional Integrated Environment Modeling System (RIEMS) in
   precipitation simulations over East Asia, Clim. Dynam. 40, 1767-1787, 2013.
- 1676 Zhao, P., Dong, F., Yang, Y., He, D., Zhao, X., and Zhang, W.: Characteristics of
  1677 carbonaceous aerosol in the region of Beijing, Tianjin, and Hebei, China, Atmos.
  1678 Environ., 71, 389–398, 2013.
- 1679 Zheng, B., Zhang, Q., Zhang, Y., He, K. B., Wang, K., Zheng, G. J., Duan, F. K., Ma, Y. L.,
  1680 and Kimoto, T.: Heterogeneous chemistry: a mechanism missing in current models to

explain secondary inorganic aerosol formation during the January 2013 haze episode in
North China, Atmos. Chem. Phys., 15, 2031–2049, doi:10.5194/acp-15-2031-2015,
2015.

- Zhong, J., Zhang, X., Dong, Y., Wang, Y., Liu, C., Wang, J., Zhang, Y., and Che, H.: Feedback
  effects of boundary-layer meteorological factors on cumulative explosive growth of
  PM<sub>2.5</sub> during winter heavy pollution episodes in Beijing from 2013 to 2016, Atmos.
  Chem. Phys., 18, 247–258, https://doi.org/10.5194/acp-18-247-2018, 2018a.
- Zhong, J., Zhang, X., Wang, Y., Liu, C., and Dong, Y.: Heavy aerosol pollution episodes in
  winter Beijing enhanced by radiative cooling effects of aerosols, Atmos. Res., 209, 59–
  64, https://doi.org/10.1016/j.atmosres.2018.03.011, 2018b.
- Zhuang, B. L., Li, S., Wang, T. J., Deng, J. J., Xie, M., Yin, C. Q., and Zhu, J. L.: Direct
  radiative forcing and climate effects of anthropogenic aerosols with different mixing
  states over China, Atmos. Environ., 79, 349-361, 2013.

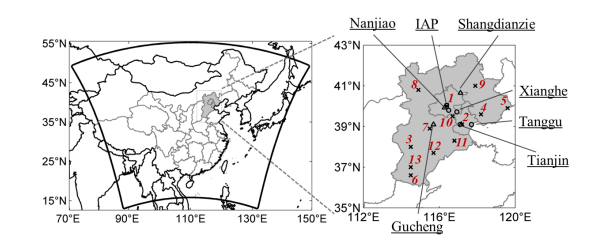




Figure 1. The model study domain. The shaded areas indicate the Beijing-Tianjin-Hebei 1698 (BTH) region. Markers are observation sites (square: IAP, observations of PM<sub>2.5</sub>, its chemical 1699 components, aerosol extinction coefficient (EXT) and aerosol absorption coefficient (ABS); 1700 circles: observations of meteorological variables; triangles: aerosol optical depth. The 1701 Xianghe site provides meteorological soundings and hourly surface shortwave radiation 1702 1703 (SWDOWN) measurements; the Tianjin site provides both meteorological variables and AOD). Hourly O<sub>3</sub>, SO<sub>2</sub>, NO<sub>2</sub> and PM<sub>2.5</sub> measurements at 13 cities from the CNEMC (China 1704 National Environmental Monitoring Center, http://www.cnemc.cn/) are labeled by crosses 1705 with numbers (1-Beijing, 2-Tianjin, 3-Shijiazhuang, 4-Tangshan, 5-Qinhuangdao, 6-Handan, 1706 1707 7-Baoding, 8-Zhangjiakou, 9-Chengde, 10-Langfang, 11-Cangzhou, 12-Hengshui and 13-Xingtai). 1708

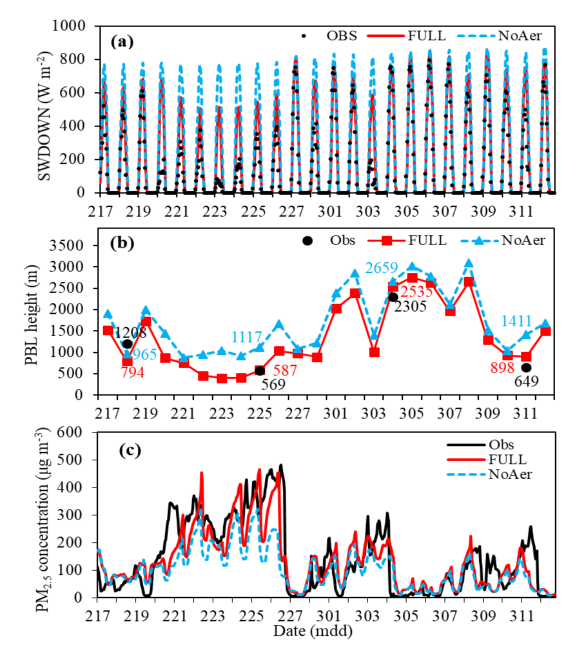


Figure 2. The model simulated and observed (a) hourly SWDOWN at Xianghe, (b) hourly
PBL height at 14:00 (LST) at Xianghe (note observations are available in the 4 days, numbers
are observations and corresponding simulations) and (c) hourly PM<sub>2.5</sub> concentration at IAP in
Beijing.

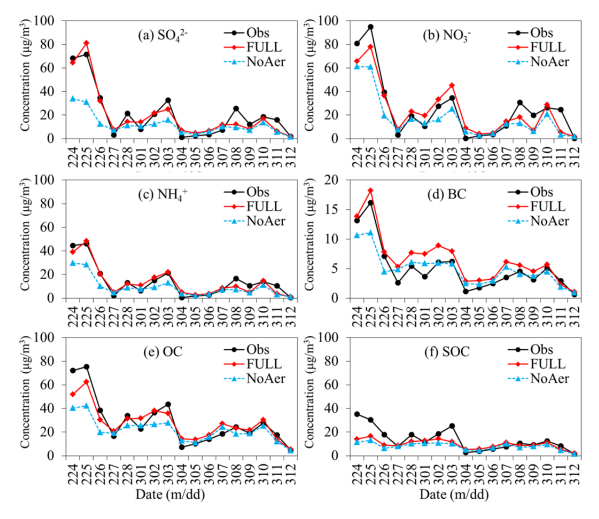
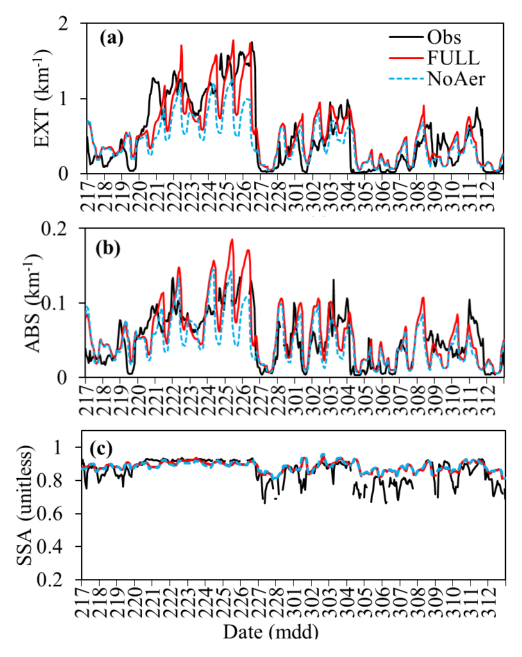


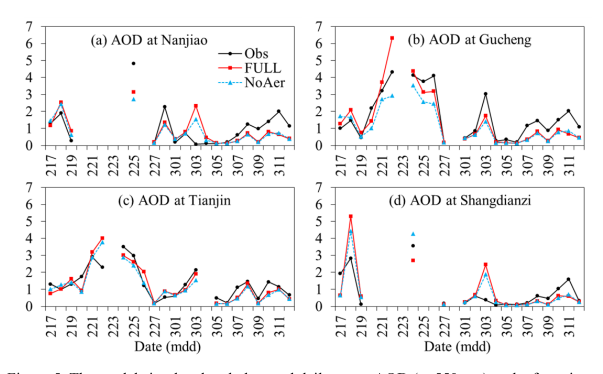
Figure 3. The model simulated and observed daily mean concentrations of aerosolcompositions in PM<sub>2.5</sub> at the IAP site in Beijing.



1718 Figure 4. The model simulated and observed hourly (a) aerosol extinction coefficient (EXT),

1719 (b) absorption coefficient (ABS) and (c) single scattering albedo (SSA) at the IAP site in

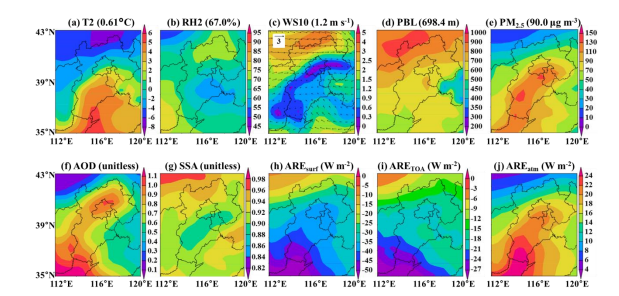
1720 Beijing under dry condition (RH=10%).



1721

1722Figure 5. The model simulated and observed daily mean AOD (at 550 nm) at the four sites of

1723 CARSNET.



1725

Figure 6. The model simulated (a) air temperature (T2), (b) relative humidity (RH2), (c) wind speed (WS10), (d) PBL height, (e) PM<sub>2.5</sub> concentration, (f) AOD, (g) SSA, (h) all-sky ARE at the surface, (i) all-sky ARE at the top of atmosphere and (j) all-sky ARE in the atmosphere from the FULL case. Numbers in the parentheses are averages over the BTH region during the entire study period.

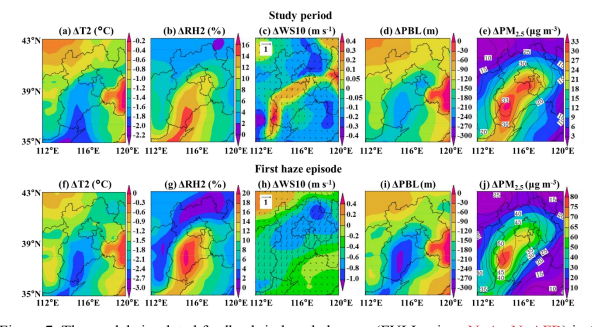


Figure 7. The model simulated feedback-induced changes (FULL minus NoAerNoAFB) in (a,
f) air temperature (T2), (b, g) relative humidity (RH2), (c, h) wind speed (WS10), (d, i) PBL
height and (e, j) PM<sub>2.5</sub> concentration averaged over the entire study period (a-e) and over the

1735 first haze episode (20–26 February) (f-j). Units are given in the parentheses.

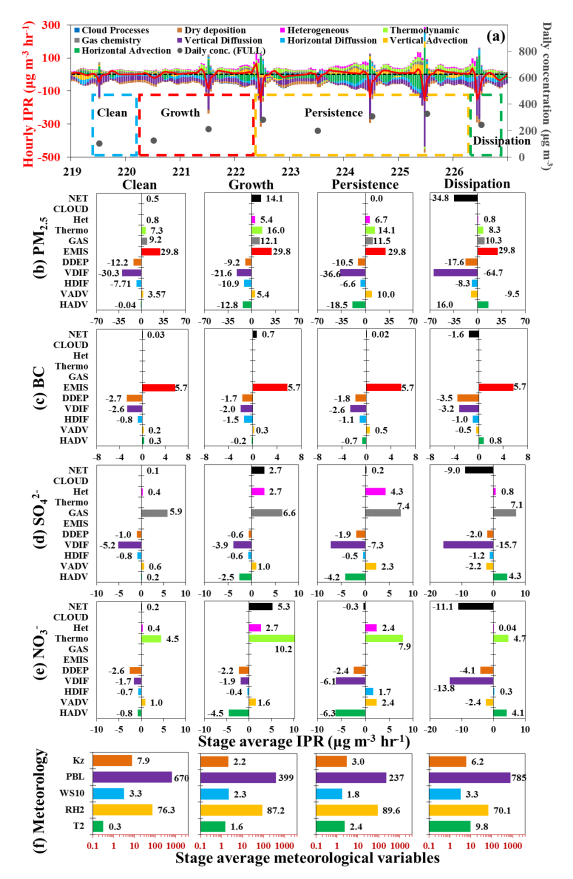


Figure 8. The model calculated integrated process rates (IPR) for the first haze episode (20–26 February) in Beijing. (a) hourly IPR, daily PM<sub>2.5</sub> concentration and the division of the four stages. The constant IPRs of emissions are not shown for clarity. The mean IPRs for (b) PM<sub>2.5</sub>, (c) BC, (d) sulfate (SO<sub>4</sub><sup>2-</sup>), nitrate (NO<sub>3</sub><sup>-</sup>), and (f) mean meteorological variables in the four stages. Note that zero IPR values are not listed. Units of T2, RH2, WS10, PBL and K<sub>z</sub> are °C, %, m s<sup>-1</sup>, m and m<sup>2</sup> s<sup>-1</sup>, respectively.

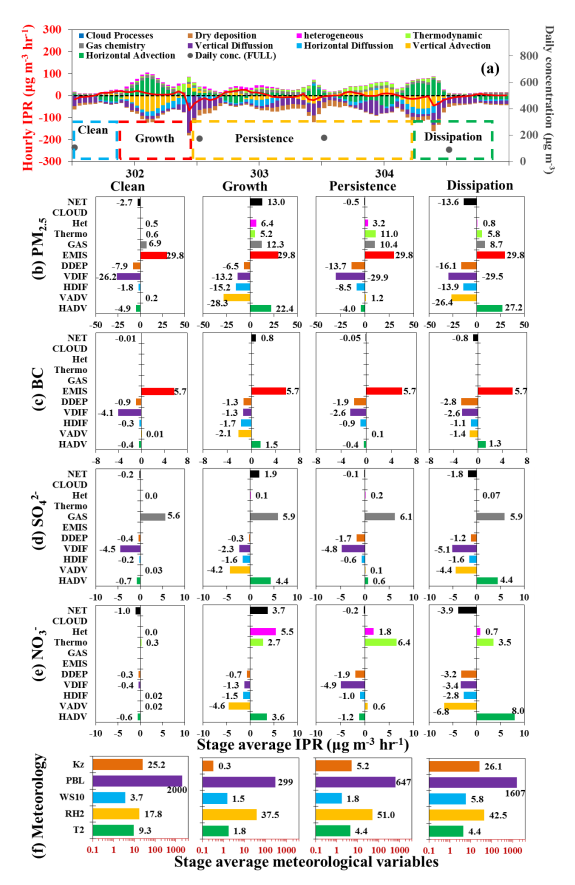


Figure 9. Same as Figure 8 but for the second haze episode (1–4 March).

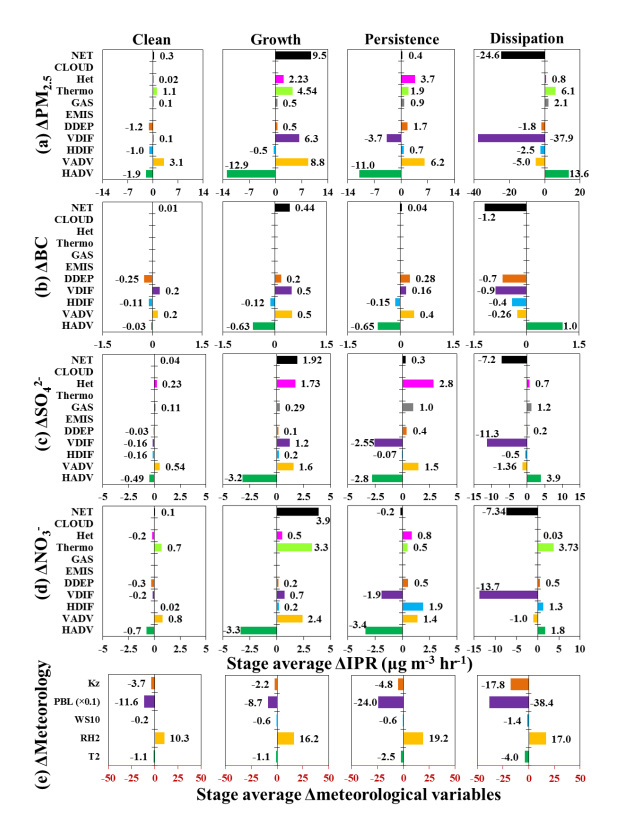


Figure 10. The feedback-induced mean changes in IPRs (FULL minus NoAerNoAFB) for PM<sub>2.5</sub> and its chemical components and meteorological variables during the first haze episode (20–26 February) in Beijing.  $\Delta$ IPRs for PM<sub>2.5</sub> and its chemical components and  $\Delta$ meteorological variables are averages over the four stages. Note that zero  $\Delta$ IPR values (no change) are not shown and the  $\Delta$ PBL heights are scaled by 0.1. The division of the four stages and units are the same as those in Figure 8.

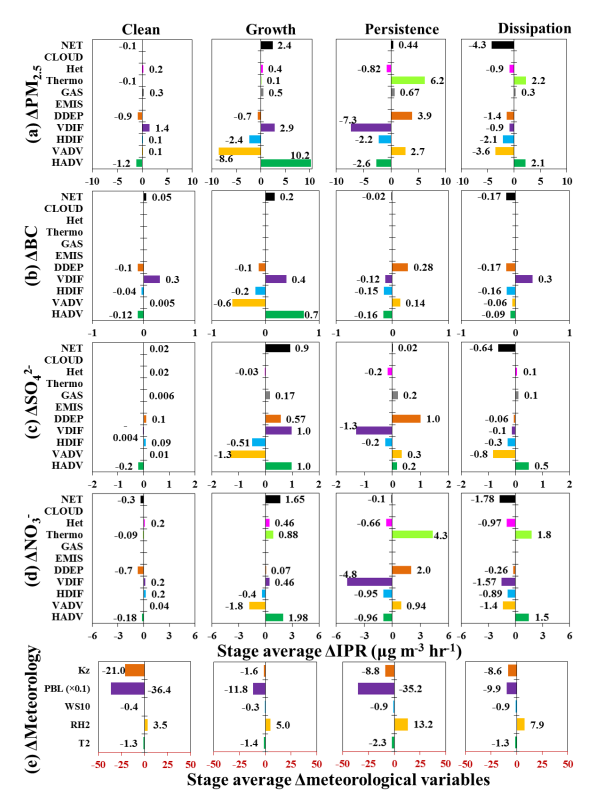


Figure 11. Same as Figure 10 but for the second haze episode (1–4 March). The division of the four stages are the same as that in Figure 9.

Table 1. Performance statistics for meteorological variables at observation sites in the BTH region. Mean observation (Obs), mean simulation (Sim), correlation coefficient (R) and normalized mean bias (NMB in %) are given. WS10, T2 and RH2 are wind speed at 10 meter, air temperature at 2 meter and relative humidity at 2 meter, respectively. All the sample numbers are 207.

Sites	Longitude	Latitude	WS	510 (r	n s <sup>-1</sup> )		-	Г2 (° <b>(</b>	C)		R	CH2 (	%)		SW	DOW	N (W	′ m <sup>-2</sup> )
			Obs	Sim	R	NMB	Obs	Sim	R	NMB	Obs	Sim	R	NMB	Obs	Sim	R	NMB
FULL																		
Beijing	39°48'N	116°30'E	2.3	2.9	0.53	28%	3.0	2.5	0.77	-16%	53.4	62.6	0.72	17%				
Tianjin	39°6'N	117°6'E	2.6	3.1	0.53	23%	3.5	3.8	0.89	8%	62.9	59.2	0.68	-6%				
Tanggu	39°6'N	117°42'E	2.4	3.4	0.36	42%	3.0	3.1	0.84	2%	69.3	61.3	0.49	-12%				
Total			2.4	3.1	0.47	31%	3.2	3.1	0.83	-2%	61.9	61.0	0.61	-1%				
Xianghe	39°45'N	116°58'E													136.0	188.4	0.91	38%
NoAer <u>NoAF</u>	B																	
Beijing	39°48'N	116°30'E	2.3	3.4	0.48	48%	3.0	4.1	0.74	37%	53.4	51.1	0.68	-4%				
Tianjin	39°6'N	117°6'E	2.6	3.6	0.48	39%	3.5	5.3	0.88	51%	62.9	47.8	0.65	-24%				
Tanggu	39°6'N	117°42'E	2.4	3.8	0.28	60%	3.0	4.5	0.84	50%	69.3	51.4	0.48	-26%				
Total			2.4	3.6	0.41	49%	3.2	4.6	0.82	46%	61.9	50.1	0.59	-19%				
Xianghe	39°45'N	116°58'E													136.0	234.0	0.85	72%

I

Table 2. Performance statistics for  $PM_{2.5}$  concentration and its chemical components, aerosol optical parameters at RH=10% (EXT, ABS and SSA) at the IAP site in Beijing. Mean observation (Obs), mean simulation (Sim), correlation coefficient (R) and normalized mean bias (NMB in %) are listed.

l

		FULL				NoAei	NoAerNoAFB			
Species (unit)	Samples	Obs	Sim	R	NMB	Sim	R	NMB		
PM <sub>2.5</sub> (µg m <sup>-3</sup> )	570	142.0	131.4	0.80	-7%	101.2	0.73	-29%		
$SO_4^{2-}$ (µg m <sup>-3</sup> )	33	21.0	20.3	0.92	-4%	11.9	0.88	-44%		
$NO_{3}^{-}$ (µg m <sup>-3</sup> )	33	26.0	24.3	0.88	-6%	17.6	0.87	-32%		
$NH_{4}^{+} (\mu g m^{-3})$	33	14.1	13.9	0.91	-2%	9.4	0.89	-34%		
BC (µg m <sup>-3</sup> )	33	5.2	6.7	0.92	28%	5.0	0.84	-3%		
OC (µg m <sup>-3</sup> )	33	29.1	28.3	0.88	-3%	22.3	0.78	-24%		
POC (µg m <sup>-3</sup> )	33	15.5	18.4	0.93	19%	14.1	0.87	-9%		
SOC (µg m <sup>-3</sup> )	33	13.6	9.9	0.56	-27%	8.2	0.45	-40%		
EXT (km <sup>-1</sup> )	570	0.51	0.53	0.79	4%	0.41	0.72	-19%		
ABS (km <sup>-1</sup> )	534	0.048	0.052	0.68	10%	0.043	0.59	-11%		
SSA (unitless)	534	0.85	0.88	0.65	5%	0.88	0.59	5%		

Table 3. Performance statistics for daily mean AOD at the four CARSNET sites in the BTH region. Mean observation (Obs), mean simulation (Sim), correlation coefficient (R) and normalized mean bias (NMB, in the unit of %) are listed.

				FULL			NoAerNoAFB			
_	Samples	Obs	Sim	R	NMB	Sim	R	NMB		
Nanjiao	18	1.09	0.92	0.67	-15.6%	0.82	0.74	-24.9%		
Gucheng	22	1.73	1.51	0.90	-12.8%	1.16	0.91	-33.0%		
Tianjin	22	1.37	1.29	0.86	-5.7%	1.19	0.86	-12.8%		
Shangdianzi	17	0.84	0.90	0.72	6.2%	0.89	0.85	5.0%		
Total	79	1.29	1.18	0.81	-8.6%	1.03	0.82	-20.2%		

l

Table 4. The model simulated domain and period averages of AOD, SSA and AREs from the FULL case over the BTH region.

	AOD (unitless)	SSA (unitless)	ARE <sub>surf</sub> (W m <sup>-2</sup> )	ARE <sub>TOA</sub> (W m <sup>-2</sup> )	ARE <sub>atm</sub> (W m <sup>-2</sup> )
-	Stuc	ly period (17	7 February	to 12 Mar	ch)
All day	0.78	0.91	-37	-18	19
Daytime	1.53	0.92	-79	-39	40
	Fi	rst haze epis	sode (20–20	6 February	<i>.</i> )
All day	1.59	0.93	-57	-32	25
Daytime	3.17	0.93	-123	-69	53

Table 5. The model simulated feedback-induced changes (FULL minus NoAerNoAFB) in T2, WS10, RH2, PBL height,  $PM_{2.5}$  concentration and vertical diffusion coefficient (K<sub>z</sub>) averaged over the BTH region during the entire period and the first haze episode. Inside the parentheses are percentage changes relative to the NoAerNoAFB case.

	ΔT2 (°C)	$\Delta WS10 (m s^{-1})$	ΔRH2 (%)	$\Delta PBL$ height (m)	$\Delta PM_{2.5} \ (\mu g \ m^{-3})$	$\Delta K_z (m^2 s^{-1})$
			Study period (17	February to 12 March	l)	
All day	-1.4 (-69.4%)	-0.038 (-3.1%)	+8.7 (+14.9%)	-160.0 (-18.6%)	+20.0 (+28.6%)	-3.3 (-27.0%)
Daytime	-1.8 (-42.1%)	+0.028 (+1.9%)	+9.0 (+16.9%)	-267.1 (-22.4%)	+21.1 (+35.6%)	-6.7 (-27.6%)
			First haze episode	e (20–26 February)		
All day	-1.8 (-59.7%)	-0.52 (-19.5%)	+9.8 (+12.4%)	-183.6 (-31.0%)	+45.1 (+38.7%)	-3.9 (-48.8%)
Daytime	-2.5 (-46.6%)	-0.59 (-19.8%)	+10.4 (+13.8%)	-307.3 (-37.6%)	+49.3 (+48.5%)	-8.3 (-51.9%)

в

# 12 Table 6. Same as Table 5 but for Beijing.

	ΔT2 (°C)	$\Delta$ WS10 (m s <sup>-1</sup> )	ΔRH2 (%)	$\Delta PBLH(m)$	ΔPM <sub>2.5</sub> (µg m <sup>-3</sup> )	$\Delta K_z (m^2 s^{-1})$
			Study period (17 I	February to 12 March)	-	
All day	-1.6 (-39.1%)	-0.48 (-13.9%)	+11.8 (+23.3%)	-154.0 (-18.3%)	+30.1 (+29.8%)	-4.5 (-37.5%)
Daytime	-2.3 (-33.1%)	-0.52 (-13.9%)	+12.5 (+28.1%)	-282.7 (-22.5%)	+34.0 (+43.9%)	-9.6 (-38.8%)
			First haze episode	(20–26 February)		
All day	-2.1 (-46.1%)	-0.58 (-20.4%)	+17.0 (+24.5%)	-195.6 (-35.9%)	+68.0 (+39.1%)	-5.0 (-59.5%)
Daytime	-3.4 (-44.6%)	-0.78 (-23.9%)	+17.9 (+27.2%)	-358.3 (-45.5%)	+83.2 (+59.6%)	-11.0 (-63.2%)

Table 7. The model simulated feedback-induced changes (FULL minus NoAerNoAFB) in BC, sulfate (SO<sub>4</sub><sup>2-</sup>) and nitrate (NO<sub>3</sub><sup>-</sup>) averaged over the BTH region and Beijing during the entire period and the first haze episode. Inside the parentheses are percentage changes relative to the NoAerNoAFB case.

	Beijing-Tianjin-He	Beijing			
	$\Delta BC (\mu g m^{-3}) \Delta SO_4^{2-} (\mu g m^{-3})$	ΔNO <sub>3</sub> <sup>-</sup> (µg m <sup>-3</sup> )	$\Delta BC \ (\mu g \ m^{-3})$	$\Delta SO_4^{2-} (\mu g m^{-3})$	$\Delta NO_3^-$ (µg m <sup>-3</sup> )
		Study period (17 Februa	ary to 12 March)		
All day	+0.9 (+25.1%) +5.0 (+46.4%)	+6.8 (+37.3%)	+1.6 (+27.5%)	+8.4 (+58.5%)	+8.4 (+36.9%)
Daytime	+1.0 (+39.5%) +5.4 (+60.2%)	+7.2 (+43.2%)	+1.9 (+51.5%)	+9.5 (+86.5%)	+9.5 (+48.8%)
		First haze episode (20–	26 February)		
All day	+1.9 (+32.9%) +12.6 (+66.9%)	+14.6 (+40.9%)	+3.1 (+33.6%)	+22.3 (+81.8%)	+16.7 (+34.7%)
Daytime	+2.2 (+50.1%) +13.8 (+81.4%)	+15.8 (+48.3%)	+4.1 (+62.3%)	+26.0 (+112.4%)	+20.9 (+51.5%)