

The Contributions to the Explosive Growth of PM_{2.5} Mass due to Aerosols-Radiation Feedback and Further Decrease in Turbulent Diffusion during a Red-alert Heavy Haze in Jing-Jin-Ji in China

Hong Wang^{1,2*}, Yue Peng^{1,2}, Xiaoye Zhang^{1,3*}, Hongli Liu¹, Meng Zhang⁴, Huizheng Che¹, Yanli Cheng¹, Yu Zheng^{1,2}

Author list:

Hong Wang, wangh@cma.gov.cn,

Yue Peng, 1131509950@qq.com; [nuist PY@163.com](mailto:nuist_PY@163.com)

Xiaoye Zhang, xiaoye@cma.gov.cn

Hongli Liu, liuhl@cma.gov.cn

Meng Zhang, 316398453@qq.com

Huizheng Che, chehz@cma.gov.cn

Yanli Cheng, chengyl@cma.gov.cn

Yu Zheng, hblfzhengyu@126.com

1 **The Contributions to the Explosive Growth of PM_{2.5} Mass due**
2 **to Aerosols-Radiation Feedback and Further Decrease in**
3 **Turbulent Diffusion during a Red-alert Heavy Haze in**
4 **Jing-Jin-Ji in China**

5 Hong Wang^{1,2*}, Yue Peng^{1,2}, Xiaoye Zhang^{1,3*}, Hongli Liu¹, Meng Zhang⁴, Huizheng
6 Che¹, Yanli Cheng¹, Yu Zheng^{1,2}

7 ¹ State Key Laboratory of Severe Weather (LASW), Chinese Academy of Meteorological Sciences (CAMS), CMA, Beijing 100081, China

8 ² Collaborative Innovation Center on Forecast and Evaluation of Meteorological Disasters, Nanjing University of Information Science & Technology, Nanjing
9 210044, China

10 ³ Center for Excellence in Regional Atmospheric Environment, Institute of Urban Environment, Chinese Academy of Sciences (CAS), Xiamen 361021, China

11 ⁴ Beijing Meteorological Bureau, Beijing 100089, China

12 Correspondence to: Hong Wang (wangh@cma.gov.cn), Xiaoye Zhang (xiaoye@cma.gov.cn)

13
14
15 **Abstract.** The explosive growth of PM_{2.5} mass usually results in extreme PM_{2.5} levels and severe haze
16 pollution in East China, and is generally underestimated by current atmospheric chemistry models. Based
17 on one such model, GRPAES_CUACE, three sensitivity experiments – a “background” experiment (EXP1),
18 “online aerosol feedback” experiment (EXP2), and an “80% decrease in turbulent diffusion coefficient” of
19 chemical tracers” experiment, based on EXP2 (EXP3) – were designed to study the contributions of
20 aerosol–radiation feedback (AF) and decrease in turbulent diffusion coefficient to the explosive growth of
21 PM_{2.5} during a “red-alert” heavy haze event in China’s Jing–Jin–Ji region. The results showed that the
22 turbulent diffusion coefficient calculated by EXP1 was about 60–70 m²/s on the clear day and 30–35 m²/s
23 on the haze day. This difference in diffusion coefficient was not enough to distinguish between the unstable
24 atmosphere on the clear day and extremely stable atmosphere during the PM_{2.5} explosive growth stage.
25 Also, the inversion calculated by EXP1 was obviously weaker than the actual inversion from sounding
26 observations on the haze day. This led to a 40%–51% underestimation of PM_{2.5} by EXP1; AF reduced by
27 about 43%–57% diffusion coefficient during the PM_{2.5} explosive growth stage, which strengthened the
28 local inversion obviously; plus, the local inversion indicated by EXP2 was much closer to the sounding

29 observations than that by EXP1. This resulted in a 20%–25% reduction of $PM_{2.5}$ negative errors in the
30 model, reaching as low as –16% to –11% in EXP2. However, the inversion produced by EXP2 was still
31 weaker than the actual observation, and AF could not solve all the problems of $PM_{2.5}$ underestimation.
32 Based on EXP2, the 80% decrease in turbulent diffusion coefficient of chemical tracers in EXP3 resulted in
33 near-zero turbulent diffusion, referred to as an “turbulent intermittence” atmospheric state, which resulted
34 in a further 14%–20% reduction in $PM_{2.5}$ underestimation, and the negative $PM_{2.5}$ errors were reduced to
35 –11% to 2%. The combined effects of AF and decrease in turbulent diffusion coefficient solved over 79%
36 of the underestimation of the explosive growth of $PM_{2.5}$ in this study. The results show that online
37 calculation of AF is essential for the prediction of $PM_{2.5}$ explosive growth and peaks during severe haze in
38 China’s Jing–Jin–Ji region. Besides, an improving in the planetary boundary layer scheme with respect to
39 extremely stable atmospheric stratification is also essential for a reasonable description of local “turbulent
40 intermittence” and a more accurate prediction of $PM_{2.5}$ explosive growth during severe haze in in this
41 region of China.

42 **Keywords:** aerosol–radiation feedback; turbulent diffusion; planetary boundary layer scheme; temperature
43 inversion; $PM_{2.5}$

44 **1 Introduction**

45 Since 2013, East China has been experiencing unprecedented intrusions of severe haze accompanied
46 by high levels of particulate matter (PM) of less than 2.5 microns in aerodynamic diameter (PM_{2.5}), causing
47 wide public concern (Ding et al., 2013; Wang et al. 2013; Huang et al., 2014; Wang et al., 2014; Sun et al.,
48 2014; Hua et al., 2016; Yang et al., 2015; Zhong et al., 2017, 2018a, 2018b). The instantaneous PM_{2.5}
49 concentration is usually in the hundreds of ug/m³ during severe haze episodes, occasionally exceeding one
50 thousand, in the metropolitan region of Beijing–Tianjin–Hebei, referred to here as Jing–Jin–Ji, and its
51 surroundings of East Shanxi, West Shandong, and North Henan in East China (Wang et al., 2014; Quan et
52 al., 2014; Sun et al., 2014; Yang et al., 2015; Zheng et al., 2016). Studies have shown, however, that models
53 generally underestimate the explosive growth and peak values of PM_{2.5} during severe hazes, especially in
54 Jing–Jin–Ji (Wang et al., 2013; Wang et al., 2014; Li et al., 2016).

55 The causes of PM_{2.5} explosive growth and its underestimation by atmospheric chemistry models are
56 complex and uncertain at present, but it possibly involves local emissions, regional transportation, aerosol
57 physicochemical processes, gas–particle conversion, meteorological conditions, and so on. However, the
58 actual atmospheric stability and how accurate it is described by atmospheric models is a fundamental
59 problem that cannot be ignored among others. Local or regional meteorological conditions dictate whether
60 haze occurs and what the PM_{2.5} level may be (Zhang et al., 2014; Zheng et al., 2015; Gao et al., 2016) when
61 source emissions are unchanged for a short period of time. The meteorological conditions of the planetary
62 boundary layer (PBL) are a key and direct trigger for the emergence of a haze event (Wang et al., 2014; Li
63 et al., 2016; Zhong et al., 2017). Turbulent diffusion is an important factor to characterize PBL meteorology
64 when the atmosphere is stable. Also, it is a major pathway of particle and gaseous pollutant exchange from
65 the surface to upper atmosphere; and when haze occurs, pollutant dispersal via the upper-level winds can
66 take place when haze is accompanied by calm surface winds and weak vertical motion of air in surface
67 layers and the PBL. The intensity of turbulent diffusion largely determines the severity of haze pollution.
68 Thus, a reasonable description of turbulent diffusion by PBL schemes in atmospheric chemistry models is
69 vital for the prediction of severe pollution (Hong et al., 2006; Wang et al., 2015; Hu et al., 2012, 2013a,
70 2013b; Li et al., 2016). The latest studies in this field of research show (Wang et al., 2015; Li et al., 2016)
71 that current PBL schemes may be insufficient for describing the extremely weak turbulent diffusion

72 conditions when extremely severe haze occurs in Jing–Jin–Ji, which more broadly may be one important
73 reason why PM_{2.5} peaks are underestimated by atmospheric chemistry models. More specifically, there may
74 be two independent reasons why the description of extremely weak turbulent diffusion in atmospheric
75 models is deficient. One is that aerosol–radiation feedback (AF) is not calculated online in the model run.
76 AF may restrain turbulence by cooling the surface and PBL while heating the atmosphere above it when
77 aerosols with certain absorption characteristics are concentrated in the PBL (Wang et al., 2010; Forkel et al.,
78 2012; Gao et al., 2014, 2015; Wang et al., 2015; Ding et al., 2016; Li et al., 2016; Miao et al., 2016; Petaja
79 et al., 2016; Gao et al., 2017; Qiu et al., 2017; Zhong et al., 2018). Ignoring AF is likely to lead to an
80 obvious overestimation of turbulent diffusion when the PM_{2.5} concentration exceeds a certain value, which
81 is worthy of further study. The other possible reason is that the extremely weak turbulence resulting in
82 extremely severe haze is not fully described by the atmospheric chemistry model (Li et al., 2016).

83 In the present work, a “red-alert” heavy haze event (issued by China’s Ministry of Environmental
84 Protection when the air pollution index is forecast to exceed 300 over the next three days) that occurred
85 during 15–23 December 2016 in China’s Jing–Jin–Ji region was selected to study the contributing factors to
86 PM_{2.5} explosive growth and peaks, and the possible deficiency of atmospheric models in describing
87 extremely weak turbulent diffusion.

88 **2 Model, data and methods**

89 **2.1 Model**

90 Focusing on dust and haze pollution in China and East Asia, the Chinese Unified Atmospheric
91 Chemistry Environment (CUACE) (Gong and Zhang, 2008) was online-integrated into the mesoscale
92 version of the Global/Regional Assimilation and PrEdiction System (GRAPES_meso), developed by the
93 Chinese Academy of Meteorological Sciences (Chen et al., 2008; Zhang and Shen, 2008), to build an
94 online chemical weather forecasting model, GRAPES_CUACE (Wang et al., 2009, 2010; 2015a; Zhou et
95 al., 2012). The main components of GRAPES_CUACE include: a model dynamic core; a modularized
96 physics package (Xu et al., 2008); an atmospheric chemistry module, CUCAE, with online coupling of
97 direct and indirect aerosol feedback; and an emissions inventory. The dynamic framework of
98 GRAPES_CUACE is semi-implicit, semi-Lagrangian, fully compressible, and non-hydrostatic (Yang et al.,

99 2007, 2008; Chen et al., 2008). A height-based terrain-following coordinate system is used, and there are 33
100 vertical layers from the surface to 30 km. A longitude–latitude grid is adopted in the spatial discretization of
101 the model and the horizontal resolution may vary upon request. The physics package can also be tailored by
102 the user (Xu et al., 2008), and Table 1 lists the specific physics and chemistry schemes used in this study.
103 The gas-phase chemistry of RAD II (Stockwell et al., 1990), with 63 gaseous species through 21
104 photochemical reactions and 121 gas-phase reactions, is used in this study. The aerosols include sea salt
105 (SS), sand/dust (SD), black carbon (BC), organic carbon (OC), sulfates (SFs), nitrates (NI) and ammonium
106 salts (AM), and aerosol processes involving hygroscopic growth, coagulation, nucleation, condensation,
107 dry and wet deposition, scavenging, aerosol activation, and so on. The formation of SF aerosols and
108 secondary organic aerosols from gases, NI and ammonium formed through gaseous oxidation, and
109 ISORROPIA (Fountoukis et al., 2007) calculating the thermodynamic equilibrium between NI and
110 ammonium and their gas precursors, are considered in CAUCE, which has been evaluated and introduced
111 in previous studies (Gong and Zhang et al., 2008; Zhou et al., 2008, 2012).

112 Based on the modeled aerosol concentrations, vertical profiles of temperature change, including direct
113 aerosol impacts, are calculated by the radiation model and fed back online to the model dynamic core at
114 each grid point and every time step, which reforms the model temperature field, dynamic process, regional
115 circulation and meteorological conditions, in turn ultimately impacting the aerosol concentration. The
116 external mixing of aerosols species (SS, SD, BC, OC, SF, NI, AM) and particle size bins are used in the
117 calculation of AF, as introduced and evaluated in detail in previous studies (Wang et al., 2009, 2010, 2015a,
118 2015b). With this two-way GRAPES_CUACE model, aerosol–radiation–PBL–meteorological interactions,
119 as well as aerosol–cloud–precipitation interactions and regional pollution and transportation of PM_{2.5} etc.,
120 have been successfully studied (Wang et al., 2010, 2015a, 2015b; Zhou et al., 2012, 2016; Jiang et al., 2015;
121 Zhang et al., 2018).

122 The turbulent diffusion coefficient is calculated by the YonSei University PBL scheme (Hong et al.,
123 2006), which is a revised vertical diffusion package based on the nonlocal boundary layer vertical diffusion
124 scheme in a medium-range forecast (MRF) model (Hong et al., 1996). The major ingredient of the revision
125 is the inclusion of an explicit treatment of entrainment processes at the top of the PBL, compared with the

126 MRF PBL scheme. The specific calculation method of diffusion coefficient is shown in Hong et al. (1996),
127 and has been selected as a standard option in MRF models (Caplan et al. 1997; Farfán and Zehnder, 2001;
128 Basu, et al., 2002; Bright and Mullen, 2002; Mass et al., 2002) as well as the Weather Research and
129 Forecasting model (Hong et al., 2006) in the National Centers for Environmental Prediction (NCEP) since
130 its establishment.

131 The horizontal resolution of the model adopted here was $0.15^\circ \times 0.15^\circ$, to match the resolution of the
132 emission source. Considering the impacts of the interregional transport of pollutants, East China (100° –
133 140°E , 20° – 60°N) (Figure 1a) was set as the model domain, but our discussion focuses mainly on the most
134 polluted area, Jing–Jin–Ji (red frame in Figure 1a), for which Figure 1b illustrates the geographical and
135 topographical features. There are two balloon sounding stations, Xingtai and Beijing (yellow stars in Figure
136 1b) in our study area. Xingtai, located in southern Hebei province at the eastern foot of the Taihang
137 Mountains, is influenced by descending airflow from the mountains in winter, and in recent years has
138 frequently been ranked the most polluted city in China. The topography of Xingtai and the serious haze
139 pollution it experiences are closely related to its situation on the southern plain of Jing–Jin–Ji. Beijing,
140 located next to Tianjin and surrounded by Hebei, lies in the transitional zone from the Yan Mountains to its
141 southern plain, and represents the most polluted areas in the central part of Jing–Jin–Ji.

142 **2.2 Emissions inventory**

143 Based on the Multi-resolution Emissions Inventory for China in 2012 (He et al., 2012), the changes in
144 East China of five kinds of emission sources – industrial, domestic, agricultural, natural, and traffic – were
145 obtained from national statistical data with respect to industry, energy consumption, road networks, and
146 motor vehicles, and updated to 2015 and 2016. Five reactive gases (SO_2 , NO , NO_2 , CO , NH_3), 20 volatile
147 organic compounds [VOCs (ALD, CH_4 , CSL, ETH, HC_3 , HC_5 , HC_8 , HCHO, ISOP, KET, NR, OL_2 , OLE,
148 OLI, OLT, ORA_2 , PAR, TERPB, TOL, XYL), listed in Table 2], and five aerosol species (BC, OC, SF, NI
149 and fugitive dust), were obtained via the above emissions data according to the input requirement of the
150 CUACE model. The horizontal grid resolution was $0.15^\circ \times 0.15^\circ$ and there was one emissions dataset for
151 each month at hourly intervals.

152 **2.3 Data**

153 Hourly observational $PM_{2.5}$ concentration data for more than 1440 surface observational stations (blue
154 dots in Figure 1) from the China National Environmental Monitoring Centre (<http://www.cnemc.cn>) during
155 15–23 December 2016 were used to evaluate the model results. The hourly observational meteorological
156 data, including wind speed and temperature, from 500 surface automatic observation stations of the China
157 Meteorological Administration (CMA) in the Jing–Jin–Ji region (red triangle in Figure 1b), were used for
158 model validation. Meteorological balloon sounding data from the CMA at 0000 UTC (early morning) and
159 1200 UTC (dusk, local time) in Beijing and Xingtai (yellow star in Figure 1b) during the same period were
160 also used to compare with the modeled results. There is one AERONET station (Holben et al., 1998),
161 Xianghe, and two CARSNET stations (Che et al., 2009; 2014; 2015), Beijing and Shijiazhuang, in the
162 Jing–Jin–Ji region (black crosses in Figure 1b). Observed aerosol optical depth (AOD) and single scattering
163 albedo (SSA) data from these three stations during the same period were also used for model evaluation.
164 NCEP $0.25^\circ \times 0.25^\circ$ global analysis gridded data (<https://rda.ucar.edu/datasets/ds083.3>) were used as the
165 model's initial and six-hourly lateral boundary meteorological input fields. The initial values of chemical
166 tracers were obtained according to their five-year mean climatic values. The results of the first 120 hours of
167 the model were discarded to eliminate the effects of the chemical initial fields.

168 **2.4 Experimental design**

169 Both dynamic processes of the regional atmosphere and solar radiation have important impacts on
170 turbulent diffusion and PBL processes. When severe haze occurs, it has been showed from observation
171 study (Zhong et al., 2018) that surface-level daily direct radiative exposure is reduced by around 89%
172 compared with clean days, suggesting the possibility of a huge difference in turbulent diffusion between
173 severe haze and clean days. However, it is difficult to distinguish between the two reasons for extremely
174 weak turbulent diffusion in the true atmosphere, because of the complicated relationship between
175 atmospheric dynamics and solar radiation. However, meaningful results might be possible by conducting
176 sensitivity experiments using an atmospheric chemistry model. Here, three such experiments (EXP1, EXP2,
177 and EXP3 – see Table 3 for descriptions) were designed to discuss the contributing factors to extremely
178 weak turbulence and corresponding $PM_{2.5}$ explosive growth, along with the insufficient description of
179 extremely weak turbulent diffusion by PBL schemes in atmospheric chemistry models. All other model

180 dynamic processes, physical options, and initial input data of the meteorology and chemical tracers were
181 same for the three experiments, i.e., except the differences shown in Table 3. In EXP3, a further decrease in
182 the turbulent diffusion coefficient based on EXP2 was only applied to the diffusion coefficient of chemical
183 tracers in CUACE mode; the diffusion coefficient in other physical packages and the dynamic framework
184 of GRAPES_MESO was the same as in EXP1 and EXP2.

185 **3 Results and discussion**

186 The studied haze episode began on 15 December 2016. $PM_{2.5}$ began to gather and climb slowly at this
187 time, but was below 150 ug/m^3 in most of Jing–Jin–Ji from 00:00 UTC 15 to 00:00 UTC 17 December – a
188 period we refer to as the “climbing stage” of $PM_{2.5}$. From 00:00 UTC 17 to 00:00 UTC 21 December, $PM_{2.5}$
189 increased rapidly, and reaching a peak of $400\text{--}600 \text{ ug/m}^3$ in most of the study area. We refer to this period
190 as the “explosive growth stage” of $PM_{2.5}$. In this section, we focus mainly on the contributions of AF and
191 decrease in turbulent diffusion coefficient to the $PM_{2.5}$ during this stage.

192 **3.1 Synoptic background**

193 The circulation in the upper atmosphere and the surface-level synoptic system controlling Jing–Jin–Ji
194 remained relatively stable during the maintenance of this haze episode. Figure 2 displays the geopotential
195 height, temperature, and winds in the upper (500 hPa), middle (700 hPa) and lower (850 hPa) atmosphere,
196 as well as PBL levels (900, 950, 1000 hPa), at 0000 UTC 19 December 2016, to show the meteorological
197 background. It can be seen that the geopotential height in the upper atmosphere (500 hPa) showed zonal
198 circulation in East Asia. There was a horizontal trough north of Jing–Jin–Ji (black frame) in the upper and
199 middle atmosphere (500 and 700 hPa), and the region was controlled by moderate northwesterly or
200 westerly air flow at the bottom of the trough. The temperature and wind fields at 500 and 700 hPa both
201 showed that cold air in the upper and middle atmosphere was weak. The 850-hPa geopotential height
202 showed that the subtropical high in the East Sea was strong; also, Jing–Jin–Ji was in the pressure
203 equalization field to the northwest periphery of the subtropical high and the wind was very weak at this
204 level due to the blocking of the subtropical high. The 900-, 950- and 100-hPa geopotential heights all
205 showed that Jing–Jin–Ji was located in the pressure equalization field between the “northwest land high”
206 and southeast subtropical high within the whole PBL, and the land high was weaker than the subtropical

207 high. This resulted in a small pressure gradient, weak and thin wind fields, and a stable atmospheric
208 situation within the PBL, which was conducive to the maintenance of the haze episode.

209 **3.2 Observation–model comparison**

210 Meteorological factors not only at the surface but also in the PBL are key in affecting haze processes
211 and PM_{2.5} concentrations (Wang et al., 2014a, 2014b). Unfortunately, however, most numerical models
212 struggle to simulate these aspects, which is also a key point determining the performance of atmospheric
213 chemistry models (Hu et al., 2013a, 2013b; Li et al., 2016).

214 Using hourly meteorological data from surface automatic observation stations of the CMA, the
215 surface wind speed and temperature at Beijing and Xingtai, and the average for Jing–Jin–Ji, according to
216 the results of EXP1, EXP2 and EXP3, were evaluated for the period 15–24 December 2016 (Figure 3). It
217 can be seen that, in Beijing, the modeled surface wind speed in the three experiments was in good
218 agreement with observation, in terms of the overall trend as well as the maximum and minimum values.
219 The observed and modeled wind speed was basically below 2 m/s during 17–21 December (i.e., the
220 explosive growth stage of PM_{2.5}). The modeled wind speed at Xingtai was slightly worse than that at
221 Beijing, but the overall trend of change was basically consistent with observation, and the wind speed was
222 also below 2 m/s during the explosive growth stage. The modeled wind speed was to an extent higher than
223 observed at the beginning and end in Xingtai. The trend of change in the modeled average wind speed for
224 the Jing–Jin–Ji region showed reasonable agreement with observation and was closest to the observed
225 situation in the explosive growth stage. In general, the modeled regional wind was higher than observed.
226 Comparison of the wind speed among the three experiments showed that the wind speeds in EXP2 and
227 EXP3 were basically same, but to a varying degree both were smaller than in EXP1 at Beijing and Xingtai,
228 as well as for Jing–Jin–Ji as a whole, during the explosive growth stage, showing that AF decreased the
229 surface wind speed. The trend of temperature change according to the three experiments was also consistent
230 with observation, at Beijing, Xingtai, and Jing–Jin–Ji as a whole. However, it was found that the modeled
231 temperature was obviously higher than observed, especially during the explosive growth stage. The
232 temperature in EXP2 and EXP3 was basically same, but lower than in EXP1, which was much closer to
233 observation, indicating that AF reduced the overestimation of surface temperature in Beijing, Xingtai, and

234 Jing–Jin–Ji as a whole. However, the temperature in EXP2 and EXP3 was also higher than observed during
235 the explosive growth stage, suggesting a role played by other uncertainties in the PBL scheme besides AF,
236 which is deserving of more detailed study in the future. Also shown in Figure 3 are the PBL-mean winds of
237 the three experiments for Beijing, Xingtai, and Jing–Jin–Ji as a whole. Unfortunately, no observational data
238 were available to evaluate them. However, comparison of the PBL’s wind and temperature according to the
239 three experiments showed that the PBL-mean wind was basically below 4 m/s while the temperature was
240 high in the explosive stage at Beijing, Xingtai, and in Jing–Jin–Ji as a whole. Similar to the surface-level
241 results, the PBL-mean wind speed and temperature in EXP2 and EXP3 were basically the same, but the
242 wind speed in these two experiments was obviously lower than that in EXP1. This indicated that the
243 reduction in wind speed by AF was more obvious in the PBL than at ground level. Meanwhile, comparison
244 of the surface-level and PBL temperature of the three experiments showed that the cooling effect of AF was
245 much stronger at the surface than in the PBL.

246 Aerosol optical properties, including AOD, SSA and asymmetry factor, largely determine the direct
247 radiative effects of aerosols. The observed AOD (Table 4) and SSA (Table 5) at Shijiazhuang, Beijing and
248 Xianghe were used to evaluate the modeled results for the period 15–22 December. Because the differences
249 in the modeled AOD and SSA results of EXP1, EXP2 and EXP3 were small, those of EXP1 only are
250 referred to here. The values of modeled AOD and SSA and their temporal trends of change during 15–22
251 December were basically consistent with observation at Beijing, Shijiazhuang and Xianghe, thus
252 demonstrating good model performance in terms of its description of aerosol optical properties. Both the
253 observed and modeled SSA at Shijiazhuang, Beijing and Xianghe (Table 5) showed that the SSA was
254 obviously higher during the explosive growth stage compared with that at the beginning or end of the haze
255 on 15–16 and 22 December, illustrating that the scattering characteristics of composite aerosols increase
256 obviously when high AOD and $PM_{2.5}$ occur on severe haze days in the Jing–Jin–Ji region. The accurate
257 description of AOD and SSA, especially with respect to the change in SSA from clean to haze days, is the
258 basis of the following discussion on the effects of aerosols on $PM_{2.5}$.

259 Figure 4 displays the averaged observed $PM_{2.5}$ ($PM_{2.5_OBS}$) and simulated $PM_{2.5}$ of EXP1
260 ($PM_{2.5_EXP1}$), EXP2 ($PM_{2.5_EXP2}$) and EXP3 ($PM_{2.5_EXP3}$) during the explosive growth stage. It can be

261 seen from PM_{2.5}_OBS results that the averaged PM_{2.5} values generally exceeded 100 μg/m³ in east China,
262 and Jing–Jin–Ji comprised the most polluted areas with PM_{2.5} reaching 300–400 μg/m³ in parts of Beijing,
263 Tianjin, central-south Hebei, western Shandong, and northern Henan. The most polluted area with PM_{2.5}
264 values of 500–700 μg/m³ appeared in southern Hebei and northern Henan provinces and the maximum
265 value of PM_{2.5} even exceeded 700 μg/m³ in part area in southern Hebei. Comparison of PM_{2.5}_EXP1 and
266 PM_{2.5}_OBS shows that PM_{2.5}_EXP1 was obviously lower than PM_{2.5}_OBS on the whole. Notably, EXP1
267 failed to simulate the PM_{2.5} > 300 μg/m³. PM_{2.5}_OBS was approximately 200–300 μg/m³ over most of
268 Shandong, while PM_{2.5}_bk was only 100–200 μg/m³ in this region. Compared with PM_{2.5}_EXP1, the
269 PM_{2.5}_EXP2 values were significantly improved by AF, and were much closer to PM_{2.5}_OBS. The high
270 PM_{2.5}_OBS centers of 300–400, 400–500 and 500–600 μg/m³ were almost simulated by EXP2, indicating
271 the important effects of AF in simulating such high values of PM_{2.5}. However, the simulated areas of these
272 centers were smaller than those of PM_{2.5}_OBS. EXP2 also failed to simulate the maximum PM_{2.5} values
273 over 600 μg/m³ observed in southern Hebei. PM_{2.5}_EXP3 just about made up for this shortcoming;
274 compared with PM_{2.5}_EXP1 and PM_{2.5}_EXP2, PM_{2.5}_EXP3 was undoubtedly the closest to PM_{2.5}_OBS
275 both in terms of PM_{2.5} extremes and the area of influence. These findings illustrate that both AF and
276 decrease in turbulent diffusion coefficient in atmospheric chemistry models are required for the effective
277 prediction of PM_{2.5} explosive growth during severe haze in China's Jing–Jin–Ji region.

278 **3.3 Change in downward solar radiation flux by aerosols and decrease in turbulent diffusion** 279 **coefficient**

280 PM in the atmosphere will inevitably lead to changes in surface and atmospheric solar radiation flux.
281 When severe haze occurs, most PM is concentrated in the atmosphere near the surface and within the PBL;
282 solar radiative flux reaching the ground is reduced greatly, which is a direct trigger for the subsequent
283 changes in thermodynamics, dynamics, and then atmospheric stratification. Any factor leading to a change
284 in the atmospheric PM loading might result in a change in the surface downward solar radiation flux
285 (SDSRF). We calculated the percentage changes in SDSRF (W/m²) between EXP2 and EXP1
286 $[(\text{SDSRF}_{\text{EXP2}} - \text{SDSRF}_{\text{EXP1}}) / \text{SDSRF}_{\text{EXP1}}]$, and EXP3 and EXP1 $[(\text{SDSRF}_{\text{EXP2}} - \text{SDSRF}_{\text{EXP1}}) /$
287 $\text{SDSRF}_{\text{EXP1}}]$, to study the impacts on SDSRF of aerosols and decrease in turbulent diffusion coefficient.

288 Figure 5 shows the mean percentage change in SDSRF (W/m^2) owing to aerosols (a) and aerosols plus
289 decrease in turbulent diffusion coefficient, during the explosive growth stage. It can be seen that SDSRF
290 was reduced by more than 50% by aerosols over most of the study region (60%–65% in Jing, Jin, most of Ji,
291 and northern Shandong, and even 65%–70% in Jing, Jin, and part of Ji), indicating the important influence
292 of aerosols on SDSRF. Comparison of Figures 5b and 5a shows that this reduction in SDSRF owing to
293 aerosols (Figure 5a) in EXP2 was further strengthened by the decrease in turbulent diffusion coefficient of
294 chemical tracers in EXP3 (Figure 5b) in certain regions, because decrease in turbulent diffusion coefficient
295 led to the accumulation of more $PM_{2.5}$ near the surface (Figure 3), less transport and, subsequently, an
296 increase in total $PM_{2.5}$ loading. It can also be seen that the difference between Figures 5a and 5b is
297 negligible. This is because the major impact of decrease in turbulent diffusion coefficient was to reform the
298 vertical distribution of the atmospheric loading of $PM_{2.5}$, and its impact on the total-column $PM_{2.5}$ was
299 minor. On the other hand, the reduction in SDSRF owing to aerosol radiation was already considerable, and
300 so the change in SDSRF owing to the increased total-column $PM_{2.5}$ by decrease in turbulent diffusion
301 coefficient would be secondary. This value of SDSRF reduction owing to aerosols and decrease in turbulent
302 diffusion coefficient is basically consistent with the 56%–89% difference of observational radiative
303 exposure between clear and haze days during the same period (Zhong et al., 2018).

304 **3.4 Influence of aerosols on the reforming of the local atmospheric temperature profile**

305 Offline and online studies indicate a reforming of the atmospheric temperature profile owing to the
306 direct effect of aerosol radiation (Wang et al., 2010, 2015b; Forkel et al., 2012; Gao et al., 2014, 2015;
307 Wang et al., 2014; Gao et al., 2017; Ding et al., 2016). In our previous work (Wang et al., 2015a, 2015b),
308 composite aerosol mixing of BC, OC, SF, NI, dust, ammonium, and sea salt aerosols was online coupled
309 into the GRAPES_CAUCE model. On this basis, in the present study, the changes in the mean temperature
310 profile of Jing–Jin–Ji during daytime owing to aerosol radiation were calculated for 15–20 December 2016.
311 It can be seen from Figure 6 that aerosols cooled the atmosphere below 750–800 hPa, but warmed it above
312 this height. Considering the PBL height may be as low as several hundreds to one thousand meters when
313 severe haze occurs in Jing–Jin–Ji (Wang et al., 2015a; Zhong et al., 2017), it may be concluded that the
314 whole PBL and its near upper atmosphere were cooled by aerosols to a varying extent during the different

315 stages of this haze process. The warming effects of aerosols above 750–850 hPa were very weak, and the
316 temperature differences among different days were also small. However, the cooling effects of aerosols
317 varied the most between different days from the surface to 975 hPa. For instance, surface daytime cooling
318 was about 2.2 K on 19 December, 1.5 K on 18 and 20 December, 1 K on 17 December, and 0.5–0.6 K on
319 15–16 December. This cooling effect of aerosols decreased rapidly with height. The difference in the
320 cooling rate between the surface and 850 hPa was 1.8 K on 19 December, 1.3 K on 18 and 20 December, 1
321 K on 17 December, and 0.3–0.4 K on 15 and 16 December. The difference in the cooling rate owing to
322 aerosols between the surface and the upper PBL was much bigger during the explosive growth stage than
323 the climbing stage. This may have resulted in further intensification of the temperature inversion layer that
324 already existed during the haze event, which will be discussed in the following section.

325 The meteorological data from the vertical soundings taken at Beijing and Xingtai were used to verify
326 this change in the temperature profile owing to aerosols. Figure 7 shows the vertical temperature profiles of
327 the sounding observations and the modeled temperature profiles of EXP1 and EXP2 during the climbing
328 stage (Figure 7a) and explosive growth stage (Figure 7b) at the two stations. The temperature profiles
329 (Figure 7a) show that the model results of EXP1 and EXP2 both simulated in part the observed temperature
330 inversion at Beijing and Xingtai on 15–16 December. The negligible difference between the temperature
331 profiles of EXP1 and EXP2 indicates that aerosol radiation had very little impact on the temperature
332 profiles and local inversion during the climbing stage. Nevertheless, Figure 7b shows that the observed
333 temperature inversions were obviously stronger and thicker on 18–19 December (explosive growth stage)
334 than those on 15–16 (climbing stage), both in Xingtai and Beijing. The temperature profiles of EXP2 were
335 much closer to the observational results than those of EXP1; and especially, the temperature inversions
336 were much stronger and also closer to observation than those of EXP1. This result proves that the
337 correction of local inversions by aerosols during the PM_{2.5} explosive growth stage was effective.

338 However, it can also be seen that the inversions of EXP2, which included online AF, were still
339 weaker than observed at the two stations. This suggests there must be other reasons, besides the online
340 calculation of AF, for the underestimation of the observed extremely strong inversion by the model, which
341 is worthy of further study.

342 **3.5 Contributions of AF and decrease in turbulent diffusion coefficient to PM_{2.5} explosive growth**

343 Turbulent diffusion is the main process of gas and particle exchange from surface to upper atmosphere,
344 and removal by high-altitude transport, and one of the key tasks of atmospheric chemistry models is to
345 capture this process. Firstly, the inversion and weak turbulent diffusion, which generates from atmospheric
346 dynamic processes, leads to atmospheric stabilization and determines the occurrence of haze and its
347 strength (Zheng et al., 2016). Once the haze occurs, aerosol radiation may in turn reinforce the inversion
348 when aerosols exceed a certain critical value, leading to more PM_{2.5} gathering near the ground. The relative
349 importance of these two aspects on PM_{2.5} explosive growth may vary with PM_{2.5} concentrations and
350 meteorological conditions, but they are irreplaceable for a reasonable prediction and simulation of PM_{2.5}
351 explosive growth and peaks in atmospheric models.

352 Figure 8 displays the hourly change in observed PM_{2.5} (PM_{2.5}_OBS) and the modeled PM_{2.5} of EXP1,
353 EXP2 and EXP3, together with the modeled turbulent diffusion coefficient of the three experiments, in
354 Beijing (Figure 8a) and Xingtai (Figure 8b), for the period 15–23 December. Comparison of the PM_{2.5}
355 modeled by EXP1, EXP2 and EXP3 with observation in Beijing (Figure 8a) shows that the PM_{2.5} modeled
356 by EXP3 was the closest to observation during the whole haze episode, which agreed with the results of the
357 regional distribution of the explosive growth stage illustrated in Figure 4. EXP1 underestimated the PM_{2.5}
358 obviously during 17–22 December, and this underestimation was even more obvious with increasing PM_{2.5}.
359 This difference between the modeled and observed PM_{2.5} was largest during the explosive growth stage. AF
360 reduced this difference to a considerable extent, and the PM_{2.5} of EXP2 was much closer to observation
361 than that of EXP1 during the explosive growth stage. However, there were certain differences between the
362 observed and PM_{2.5} and that modeled by EXP2, illustrating that AF cannot completely fill the sizeable gap
363 between observed and modeled PM_{2.5}. The PM_{2.5} of EXP3 reduced this gap further, showing the best
364 agreement with observation, especially during the PM_{2.5} explosive growth stage.

365 It can also be seen from Figure 8a that the diffusion coefficient of EXP1 was about 30–40 m²/s during
366 the explosive growth stage, which was about 50% of the 60–70 m²/s on clear days (15 or 22 December).
367 Obviously, this 50% diffusion coefficient difference between clear and severe haze days may be insufficient
368 to separate the difference in turbulent diffusion intensity between the extremely stable atmosphere on haze

369 days and the unstable atmosphere on clear days, which is an important reason for the underestimated $PM_{2.5}$
370 explosive growth in EXP1. Compared with EXP1, the AF in EXP2 led to a notable enhancement of the
371 temperature inversion (Figure 7b), a significant decrease in the turbulent diffusion of $PM_{2.5}$ during the
372 explosive growth stage, and a low maximum diffusion coefficient at noon (as low as $14 \text{ m}^2/\text{s}$ on 20
373 December – a reduction of 50% compared with EXP1). The maximum diffusion coefficient at noon on
374 haze days in EXP2 was only about 20% of that on clear days. The maximum diffusion coefficient at noon
375 in EXP3 was lower than $5 \text{ m}^2/\text{s}$ on 20 December and, at the same time, the $PM_{2.5}$ modeled by EXP3 was
376 further increased and was also much closer to the observed $PM_{2.5}$ than the $PM_{2.5}$ of EXP2.

377 Through comparison of the temporal change of diffusion coefficient and $PM_{2.5}$ in EXP1, EXP2 and
378 EXP3 in Beijing, it is clear that an overestimation of turbulent diffusion coefficient owing to the absence of
379 online-calculated AF, as well as a deficient description of extremely stable stratification in the PBL scheme
380 of the atmospheric model, can lead to a distinct underestimation of $PM_{2.5}$ explosive growth and peaks when
381 severe haze occurs in China's Jing-Jin-Ji region.

382 The trends of change in diffusion coefficient and $PM_{2.5}$ at Xingtai in the three experiments (Figure 8b)
383 are similar to those at Beijing. The $PM_{2.5}$ of EXP3 was also closest to observation, followed by EXP2, and
384 then EXP1 was the worst, during the whole haze episode. However, during the explosive growth stage, the
385 relative contributions of AF and decrease in turbulent diffusion coefficient to the $PM_{2.5}$ peak values showed
386 some differences to those at Beijing. The contributions of decrease in turbulent diffusion coefficient to
387 $PM_{2.5}$ peaks were more important than those of AF at Xingtai. Located in the eastern foothills of the
388 Taihang Mountains, Xingtai is usually affected by downhill airflow. Temperature inversions in this area
389 form and strengthen easily, leading to stronger inversion, weaker turbulent diffusion, and more stable
390 atmospheric stratification. However, this kind of inversion and weak turbulent diffusion, derived from the
391 local terrain, is harder for PBL schemes in atmospheric chemistry models to describe, and likely
392 underestimated.

393 Figure 9 is a diagrammatic sketch of the contributions of AF and decrease in turbulent diffusion
394 coefficient to the $PM_{2.5}$ of the explosive growth stage according to the results at Beijing and Xingtai. It can
395 be seen that the diffusion coefficient of EXP1 was $30\text{--}35 \text{ m}^2/\text{s}$, while that of EXP2 was $15\text{--}17 \text{ m}^2/\text{s}$,

396 meaning AF reduced the diffusion coefficient by about 43%–57%, which led to the rise in simulated $PM_{2.5}$
397 from 144 ug/m^3 in EXP1 to 205 ug/m^3 in EXP2 at Beijing, and from 280 ug/m^3 in EXP1 to 360 ug/m^3 in
398 EXP2 at Xingtai. This means that AF reduced the underestimation of $PM_{2.5}$ at Beijing and Xingtai by 20%
399 and 25%, respectively. The diffusion coefficient of EXP3 was as low as $4\text{--}6 \text{ m}^2/\text{s}$ during the explosive
400 growth stage, demonstrating the joint effects of AF and decrease in turbulent diffusion coefficient reduced
401 the diffusion coefficient to less than $4\text{--}6 \text{ m}^2/\text{s}$, near-zero, which we refer to as “turbulent intermittence”.
402 The direct result of this “turbulent intermittence” was a further increase in the simulated surface $PM_{2.5}$,
403 Based on EXP2, the further decrease in turbulent diffusion coefficient reduced the underestimation of
404 simulated $PM_{2.5}$ by 14% to 20%, and the $PM_{2.5}$ errors in EXP3 were reduced to as low as -11% to 2% .

405 **4. Conclusions**

406 Using an atmospheric chemistry model, GRAPES_CUACE, three experiments (EXP1, EXP2 and
407 EXP3) were designed to study the reason for the explosive growth of $PM_{2.5}$ mass during a “red-alert” heavy
408 haze event that occurred during 15–23 December 2016 in China’s Jing–Jin–Ji region. The contributions of
409 AF and decrease in turbulent diffusion coefficient to the $PM_{2.5}$, representing compensation for the deficient
410 description of extremely weak turbulent diffusion in the PBL scheme of the atmospheric model, were
411 studied by analyzing the changes in $PM_{2.5}$, SDSRF, wind speed and temperature, diffusion coefficient, and
412 the relationships among them, in the three experiments.

413 Results show that the diffusion coefficient in EXP1 was about $60\text{--}70 \text{ m}^2/\text{s}$ on clear days and $30\text{--}35$
414 m^2/s on haze days. The 50% difference between the two was considered insufficient to separate the unstable
415 atmosphere on clear days and the extreme stable atmosphere on severe haze days, compared with the
416 differences in direct downward solar radiation between clear and haze days, which was also proven
417 indirectly by the weaker inversion of EXP1 than that from sounding observations. This led to a 40%–51%
418 underestimation of the $PM_{2.5}$ peaks in EXP1 during the $PM_{2.5}$ explosive growth stage. Online calculation of
419 AF reduced the surface and PBL wind speed and cooled the surface and PBL atmosphere. The surface
420 daytime cooling due to aerosol radiation was $1.5\text{--}2.2 \text{ K}$ during the explosive growth stage and $0.5\text{--}0.6 \text{ K}$
421 during the climbing stage. The cooling effect of aerosols decreased rapidly with height, and this was a
422 major reason for the strengthening of the temperature inversion during the explosive growth stage. The

423 reduced diffusion coefficient owing to AF reached 43%–57% during the $PM_{2.5}$ explosive growth stage. The
424 local inversion simulated in EXP2 was strengthened and closer to the actual sounding observation than that
425 of EXP1. This resulted in a 20%–25% reduction in the underestimation of $PM_{2.5}$, with $PM_{2.5}$ errors in EXP2
426 being as low as –16 to –11% during the explosive growth stage. The impact on $PM_{2.5}$ owing to AF in the
427 model run was distinct during the explosive growth stage, but minor during the climbing stage, indicating a
428 critical value of $150 \mu\text{g}/\text{m}^3$ of $PM_{2.5}$ leading to an effective AF in online atmospheric chemistry models.
429 However, the local inversion simulated by EXP2 was still weaker than observed, and the $PM_{2.5}$ of EXP2
430 was still smaller than observed, illustrating AF could not solve all the $PM_{2.5}$ underestimation problems. In
431 EXP3, the decrease in turbulent diffusion coefficient of particles and gas based on EXP2 resulted in a
432 14%–20% lessening of the $PM_{2.5}$ underestimation based on EXP2, and the $PM_{2.5}$ errors of EXP3 were
433 reduced to –11% to 2%.

434 The present study illustrates that the PBL schemes in current atmospheric chemistry models are
435 probably insufficient for describing the extremely stable atmosphere resulting in explosive growth of $PM_{2.5}$
436 and severe haze in China’s Jing–Jin–Ji region. This may involve two important reasons: the absence of an
437 online calculation of AF, and/or a deficient description of extremely weak turbulent diffusion by the PBL
438 scheme in the atmospheric chemistry model. Our study suggests that an online calculation of AF and an
439 improvement in the representation of turbulent diffusion in PBL schemes, with a focus on extremely stable
440 atmospheric stratification, in atmospheric chemistry models, are indispensable for a reasonable description
441 of local “turbulent intermittence” and an accurate prediction of the explosive growth and peaks of $PM_{2.5}$ of
442 severe haze in China’s Jing–Jin–Ji region.

443

444

445 **Author Contributions:**

446 Hong Wang and Xiaoye Zhang designed the idea and experiments; Hong Wang and Yue Peng carried them
447 out; Hongli Liu prepared the emissions data and introduction; Meng Zhang performed some of the model
448 runs; Huizheng Che and Yu Zheng processed the AOD and SSA observational data; Yanli Cheng completed
449 Table 3 and the related introduction.

450 **Acknowledgements**

451 This study was supported by the National Key Project (2016YFC0203306), the National Natural Science
452 Foundation of China (41590874), and the National (Key) Basic Research and Development (973) Program
453 of China (2014CB441201).

454 **References**

- 455 Basu, S., Iyengar, G. R., and Mitra, A. K.: Impact of a nonlocal closure scheme in a simulation of a
456 monsoon system over India. *Mon. Wea. Rev.*, 130, 161–170, 2002.
- 457 Bright, D. R., and Mullen, S. L.: The sensitivity of the numerical simulation of the southwest monsoon
458 boundary layer to the choice of PBL turbulence parameterization in MM5, *Wea. Forecasting*, 17, 99–
459 114, 2002.
- 460 Caplan, P., Derber, J., Gemmill, W., Hong, S.-Y., Pan, H.-L., and Parrish, D.: Changes to the 1995 NCEP
461 operational medium-range forecast model analysis-forecast system, *Wea. Forecasting*, 12, 581–594,
462 1997.
- 463 Che, H., Zhang, X., Chen, H., Damiri, B., Goloub, P., Li, Z., Zhang, X., Wei, Y., Zhou, H., Dong, F., Li, D.,
464 and Zhou, T.: Instrument calibration and aerosol optical depth validation of the China Aerosol Remote
465 Sensing Network, *J. Geophys. Res. Atmos.*, 114, D03206, doi:10.1029/2008JD011030, 2009.
- 466 Che, H., Xia, X., Zhu, J., Li, Z., Dubovik, O., Holben, B., Goloub, P., Chen, H., Estelles, V., Cuevas-Agulló,
467 E., Blarel, L., Wang, H., Zhao, H., Zhang, X., Wang, Y., Sun, J., Tao, R., Zhang, X., and Shi, G.:
468 Column aerosol optical properties and aerosol radiative forcing during a serious haze-fog month over
469 North China Plain in 2013 based on ground-based sunphotometer measurements, *Atmos. Chem. Phys.*,
470 14, 2125–2138, doi:10.5194/acp-14-21252014, 2014.
- 471 Che, H., Zhang, X.-Y., Xia, X., Goloub, P., Holben, B., Zhao, H., Wang, Y., Zhang, X. C., Wang, H., Blarel,

472 L., Damiri, B., Zhang, R., Deng, X., Ma, Y., Wang, T., Geng, F., Qi, B., Zhu, J., Yu, J., Chen, Q., and
473 Shi, G.: Ground-based aerosol climatology of China: aerosol optical depths from the China Aerosol
474 Remote Sensing Network (CARSNET) 2002–2013, *Atmos. Chem. Phys.*, 15, 7619–7652,
475 <https://doi.org/10.5194/acp15-7619-2015>, 2015.

476 Chen, D., Xue, J., Yang, X., Zhang, H., Shen, X., Hu, J., Wang, Y., Ji, L., and Chen, J.: New generation of
477 multi-scale NWP system (GRAPES): general scientific design, *Chinese Sci. Bull.*, 53, 3433–3445,
478 doi:10.1007/s11434-008-0494-z, 2008.

479 Chou, M. D., Suarez, M. J., Ho, C. H., Yan, M. M. H., and Lee, K. T.: Parameterizations for Cloud
480 Overlapping and Shortwave Single-Scattering Properties for Use in General Circulation and Cloud
481 Ensemble Models, *J. Clim.*, 11, 202–214, 1998.

482 Chou, M. D., Suarez, M. J., Liang, X. Z., and Michael M.-H. Y.: A Thermal Infrared Radiation
483 Parameterization for Atmospheric Studies, Technical Report Series on Global Modeling and Data
484 Assimilation, NASA/TM-2001-104606, 19, America, Goddard Space Flight Center, Greenbelt,
485 Maryland, 55, 2001.

486 Ding, A. J., Fu, C. B., Yang, X. Q., Sun, J. N., Petäjä, T., Kerminen, V. M., Wang, T., Xie, Y., Herrmann, E.,
487 Zheng, L. F., Nie, W., Liu, Q., Wei, X. L., and Kulmala, M.: Intense atmospheric pollution modifies
488 weather: a case of mixed biomass burning with fossil fuel combustion pollution in eastern China,
489 *Atmos. Chem. Phys.*, 13, 10545-10554, 2013.

490 Ding, A. J., Huang, X., Nie, W., Sun, J. N., Kerminen, V. M., Petäjä, T., Su, H., Cheng, Y. F., Yang, X. Q.,
491 Wang, M. H., Chi, X. G., Wang, J. P., Virkkula, A., Guo, W. D., Yuan, J., Wang, S. Y., Zhang, R. J., Wu,
492 Y. F., Song, Y., Zhu, T., Zilitinkevich, S., Kulmala, M., and Fu, C. B.: Enhanced haze pollution by
493 black carbon in megacities in China, *Geophys. Res. Lett.*, 43, 2873-2879, 2016.

494 Farfán, L. M., and Zehnder, J. A.: An analysis of the landfall of Hurricane Nora (1997), *Mon. Wea. Rev.*,
495 129, 2073–2088, 2001.

496 Forkel, R., Werhahn, J., Hansen, A. B., McKeen, S., Peckham, S., Grell, G., and Suppan, P.: Effect of
497 aerosol-radiation feedback on regional air quality – A case study with WRF/Chem, *Atmos. Environ.*,
498 53, 202-211, 2012.

499 Fountoukis, C., and Nenes A., ISORROPIA II: a computationally efficient thermodynamic equilibrium
500 model for K^+ – Ca^{2+} – Mg^{2+} – NH_4^+ – Na^+ – SO_4^{2-} – NO_3^- – Cl^- – H_2O aerosols, *Atmos. Chem. Phys.*,
501 2007, 7, 4639–4659.

502 Gal-Chen, T., and Somerville, R. C. J.: On the use of a coordinate transformation for the solution of the
503 Navier-Stokes equations, *J. Comput. Phys.*, 17, 209–228, 1975.

504 Gao, Y., Zhao, C., Liu, X., Zhang, M., and Leung, L. R.: WRF-Chem simulations of aerosols and
505 anthropogenic aerosol radiative forcing in East Asia, *Atmos. Environ.*, 92, 250-266, 2014.

506 Gao, Y., Zhang, M., Liu, Z., Wang, L., Wang, P., Xia, X., Tao, M., and Zhu, L.: Modeling the feedback
507 between aerosol and meteorological variables in the atmospheric boundary layer during a severe fog–
508 haze event over the North China Plain, *Atmos. Chem. Phys.*, 15, 4279-4295, 2015.

509 Gao, M., Carmichael, G. R., Saide, P. E., Lu, Z., Yu, M., Streets, D. G., and Wang, Z.: Response of winter
510 fine particulate matter concentrations to emission and meteorology changes in North China, *Atmos.*
511 *Chem. Phys.*, 16, 11837-11851, 2016.

512 Gao, M., Saide, P. E., Xin, J., Wang, Y., Liu, Z., Wang, Y., Wang, Z., Pagowski, M., Guttikunda, S.K., and
513 Carmichael, G. R.: Estimates of Health Impacts and Radiative Forcing in Winter Haze in Eastern
514 China through Constraints of Surface PM_{2.5} Predictions, *Environ. Sci. Technol.*, 51, 2178-2185, 2017.

515 Gong, S., and Zhang, X.: CUACE/Dust-an integrated system for operational dust forecasting in Asia,
516 *Computers & Applied Chemistry*, 25, 1061-1067, 2008.

517 Gong, S. L., Lavoué, D., Zhao, T. L., Huang, P., and Kaminski, J. W.: GEM-AQ/EC, an on-line global
518 multi-scale chemical weather modelling system: model development and evaluation of global aerosol
519 climatology, *Atmos. Chem. Phys.*, 12, 8237–8256, doi:10.5194/acp-12-8237-2012, 2012.

520 Grell, G. A., Dudhia, J., and Stauffer, D.: A description of the fifth-generation PENN State/NCAR
521 Mesoscale Model (MM5). NCAR Tech. Note NCAR/TN-398 STR, 138, 1994.

522 He, K.: Multi-resolution Emission Inventory for China (MEIC): model framework and 1990-2010
523 anthropogenic emissions, AGU Fall Meeting, AGU Fall Meeting Abstracts, 2012.

524 Holben B. N., Eck, T .F., Slutsker, I., et al. AERONET—A Federated Instrument Network and Data
525 Archive for Aerosol Characterization, 1998.

526 Hong, S. Y. and Pan, H. L.: Nonlocal boundary layer vertical diffusion in a Medium-Range Forecast model,
527 Mon. Weather Rev., 124, 2322–2339, 1996.

528 Hong, S. Y., Noh, Y., and Dudhia, J.: A New Vertical Diffusion Package with an Explicit Treatment of
529 Entrainment Processes, Mon. Weather Rev., 134, 2318-2341, 2006.

530 Hu, X. M., Doughty, D. C., Sanchez, K. J., Joseph, E., and Fuentes, J. D.: Ozone variability in the
531 atmospheric boundary layer in Maryland and its implications for vertical transport model, Atmos.
532 Environ., 46, 354-364, 2012.

533 Hu, X. M., Klein, P. M., and Xue, M.: Evaluation of the updated YSU planetary boundary layer scheme
534 within WRF for wind resource and air quality assessments, J. Geophys. Res. Atmos., 118,
535 10490-10505, 2013a.

536 Hu, X. M., Klein, P. M., Xue, M., Zhang, F., Doughty, D. C., Forkel, R., Joseph, E., and Fuentes, J. D.:
537 Impact of the vertical mixing induced by low-level jets on boundary layer ozone concentration, Atmos.
538 Environ., 70, 123-130, 2013b.

539 Hua, Y., Wang, S., Wang, J., Jiang, J., Zhang, T., Song, Y., Kang, L., Zhou, W., Cai, R., Wu, D., Fan, S.,
540 Wang, T., Tang, X., Wei, Q., Sun, F., and Xiao, Z.: Investigating the impact of regional transport on
541 PM_{2.5} formation using vertical observation during APEC 2014 Summit in Beijing, Atmos. Chem.
542 Phys., 16, 15451–15460, 2016.

543 Huang, R. J., Zhang, Y., Bozzetti, C., Ho, K. F., Cao, J. J., Han, Y., Daellenbach, K. R., Slowik, J. G., Platt,
544 S. M., Canonaco, F., Zotter, P., Wolf, R., Pieber, S. M., Bruns, E. A., Crippa, M., Ciarelli, G.,
545 Piazzalunga, A., Schwikowski, M., Abbaszade, G., Schnelle-Kreis, J., Zimmermann, R., An, Z., Szidat,
546 S., Baltensperger, U., El Haddad, I., and Prevot, A. S.: High secondary aerosol contribution to
547 particulate pollution during haze events in China, Nature, 514, 218-222, 2014.

548 Jiang, C., Wang, H., Zhao, T., Li, T., and Che, H.: Modeling study of PM_{2.5} pollutant transport across cities
549 in China's Jing–Jin–Ji region during a severe haze episode in December 2013, Atmos. Chem. Phys., 15,
550 5803-5814, 2015.

551 Kain, J. S. and Fritsch, J. M.: Convection parameterization for mesoscale models: the Kain-Fritsch scheme,
552 Meteor. Mon., 24, 165–170, 1993.

553 Kusaka, H., Kondo, H., Kikegawa, Y., and Kimura, F.: A simple single-layer urban canopy model for
554 atmospheric models: Comparison with multi-layer and slab models. *Bound.-Layer Meteor.*, 101, 329–
555 358, 2001.

556 Li, K., Liao, H., Zhu, J., and Moch, J. M.: Implications of RCP emissions on future PM_{2.5} air quality and
557 direct radiative forcing over China, *J. Geophys. Res. Atmos.*, 121, 12985-13008, 2016.

558 Li, T., Wang, H., Zhao, T., Xue, M., Wang, Y., Che, H., and Jiang, C.: The Impacts of Different PBL
559 Schemes on the Simulation of PM_{2.5} during Severe Haze Episodes in the Jing-Jin-Ji Region and Its
560 Surroundings in China, *Adv. Meteorol.*, 1-15, 2016.

561 Lim, K. S. S., and Hong, S. Y.: Development of an effective double-moment cloud microphysics scheme
562 with prognostic cloud condensation nuclei (CCN) for weather and climate models, *Mon. Wea. Rev.*,
563 138, 1587-1612, 2010.

564 Mass, C. F., D. Ovens, K. Westrick, and B. A. Colle: Does increasing horizontal resolution produce more
565 skilful forecasts? *Bull. Amer. Meteor. Soc.*, 83, 407–430, 2002.

566 Miao, Y., Liu, S., Zheng, Y., and Wang, S.: Modeling the feedback between aerosol and boundary layer
567 processes: a case study in Beijing, China, *Environ. Sci. Pollut R.*, 23, 3342-3357, 2016.

568 Petäjä, T., Järvi, L., Kerminen, V. M., Ding, A. J., Sun, J. N., Nie, W., Kujansuu, J., Virkkula, A., Yang, X.
569 Q., Fu, C. B., Zilitinkevich, S., and Kulmala, M.: Enhanced air pollution via aerosol-boundary layer
570 feedback in China, *Sci. Rep.*, 6, 18998, 2016.

571 Pleim, J.: A combined local and non-local closure model for the atmospheric boundary layer. Part II:
572 Application and evaluation in a mesoscale meteorological model, *J. Applied Meteor. Climatology*, 46,
573 1396–1409, 2007.

574 Qiu, Y., Liao, H., Zhang, R., and Hu, J.: Simulated impacts of direct radiative effects of scattering and
575 absorbing aerosols on surface layer aerosol concentrations in China during a heavily polluted event in
576 February 2014, *J. Geophys. Res. Atmos.*, 122, 5955-5975, 2017.

577 Quan, J., Tie, X., Zhang, Q., Liu, Q., Li, X., Gao, Y., and Zhao, D.: Characteristics of heavy aerosol
578 pollution during the 2012–2013 winter in Beijing, China, *Atmos. Environ.*, 88, 83-89, 2014.

579 Stockwell, W. R., Middleton, P., Chang, J. S., and Tang, X.: The Second Generation Regional Acid

580 Deposition Model Chemical Mechanism for Regional Air Quality Modeling, *J. Geophys. Res.*, 95,
581 16343-16376, 1990.

582 Sun, Y., Jiang, Q., Wang, Z., Fu, P., Li, J., Yang, T., and Yin, Y.: Investigation of the sources and evolution
583 processes of severe haze pollution in Beijing in January 2013, *J. Geophys. Res. Atmos.*, 119,
584 4380-4398, 2014.

585 Wang, H., Gong, S., Zhang, H., Chen, Y., Shen, X., Chen, D., Xue, J., Shen, Y., Wu, X., and Jin, Z.: A
586 new-generation sand and dust storm forecasting system GRAPES_CUACE/Dust: Model development,
587 verification and numerical simulation, *Chinese Sci. Bull.*, 55, 635-649, 2009.

588 Wang, H., Zhang, X., Gong, S., Chen, Y., Shi, G., and Li, W.: Radiative feedback of dust aerosols on the
589 East Asian dust storms, *J. Geophys. Res.*, 115 , D23214, 2010.

590 Wang, H., Tan, S. C., Wang, Y., Jiang, C., Shi, G. Y., Zhang, M. X., and Che, H. Z.: A multi sources
591 observation study of the severe prolonged regional haze episode over eastern China in January 2013,
592 *Atmos. Environ.*, 89, 807-815, 2014a.

593 Wang, H., Xu, J., Zhang, M., Yang, Y., Shen, X., Wang, Y., Chen, D., and Guo, J.: A study of the
594 meteorological causes of a prolonged and severe haze episode in January 2013 over central-eastern
595 China, *Atmos. Environ.*, 98, 146-157, 2014b.

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

600 Wang, H., Shi, G. Y., Zhang, X. Y., Gong, S. L., Tan, S. C., Chen, B., Che, H. Z., and Li, T.: Mesoscale
601 modelling study of the interactions between aerosols and PBL meteorology during a haze episode in
602 China Jing-Jin-Ji and its near surrounding region - Part 2: Aerosols' radiative feedback effects, *Atmos.*
603 *Chem. Phys.*, 15, 3277-3287, 2015b.

604 Wang, J., Wang, S., Jiang, J., Ding, A., Zheng, M., Zhao, B., Wong, D. C., Zhou, W., Zheng, G., Wang, L.,
605 Pleim, J. E., and Hao, J.: Impact of aerosol-meteorology interactions on fine particle pollution during
606 China's severe haze episode in January 2013, *Environ. Res. Lett.*, 9, 094002, 2014.

607 Wang, Y., Zhang, Q. Q., He, K., Zhang, Q., and Chai, L.: Sulfate-nitrate-ammonium aerosols over China:
608 response to 2000–2015 emission changes of sulfur dioxide, nitrogen oxides, and ammonia, *Atmos.*
609 *Chem. Phys.*, 13, 2635-2652, 2013.

610 Wang, Z., Li, J., Wang, Z., Yang, W., Tang, X., Ge, B., Yan, P., Zhu, L., Chen, X., Chen, H., Wand, W., Li,
611 J., Liu, B., Wang, X., Wand, W., Zhao, Y., Lu, N., and Su, D.: Modeling study of regional severe hazes
612 over mid-eastern China in January 2013 and its implications on pollution prevention and control, *Sci.*
613 *China Earth Sci.*, 57, 3-13, 2014.

614 Xu, G., Chen, D., Xue, J., Sun, J., Shen, X., Shen, Y., Huang, L., Wu, X., Zhang, H., and Wang, S.: The
615 program structure designing and optimizing tests of GRAPES physics, *Chinese Sci. Bull.*, 53, 3470–
616 3476, doi:10.1007/s11434-008-0418-y, 2008.

617 Yang, Y., Liao, H., and Lou, S.: Increase in winter haze over eastern China in recent decades: Roles of
618 variations in meteorological parameters and anthropogenic emissions, *J. Geophys. Res. Atmos.*, 121,
619 13050-13065, 2016.

620 Yang, X., Hu, J., Chen, D., Zhang, H., Shen, X., Chen, J., and Ji, L.: Verification of GRAPES unified global
621 and regional numerical weather prediction model dynamic core, *Chinese Sci. Bull.*, 53, 3458–3464,
622 doi:10.1007/s11434-008-0417-z, 2008.

623 Yang, X. S., Chen, J. B., and Hu, J. L.: A semi-implicit semi-Lagran global nonhydrostatic model and the
624 polar discretization scheme, *Sci. China Ser D-Earth Sci.*, 50, 1885-1891, 2007.

625 Yang, Y. R., Liu, X. G., Qu, Y., An, J. L., Jiang, R., Zhang, Y. H., Sun, Y. L., Wu, Z. J., Zhang, F., Xu, W. Q.,
626 and Ma, Q. X.: Characteristics and formation mechanism of continuous hazes in China: a case study
627 during the autumn of 2014 in the North China Plain, *Atmos. Chem. Phys.*, 15, 8165–8178,
628 doi:10.5194/acp-158165-2015, 2015.

629 Zhang, M., Wang, H., Zhang, X., Peng, Y., and Che, H.: Applying the WRF double-moment six-class
630 microphysics scheme in the GRAPES_Meso model: A case study, *J. Meteor. Res.*, 32, 246, doi:
631 10.1007/s13351018-7066-1, 2018.

632 Zhang, R., and Shen, X.: On the development of the GRAPES – a new generation of the national
633 operational NWP system in China, *Chinese Sci. Bull.*, 53, 3429–3432, doi:10.1007/s11434008-0462-7,

634 2008.

635 Zhang, R. H., Li, Q., and Zhang, R. N.: Meteorological conditions for the persistent severe fog and haze
636 event over eastern China in January 2013, *Sci. China Earth Sci.*, 57, 26–35, 2014.

637 Zheng, G. J., Duan, F. K., Su, H., Ma, Y. L., Cheng, Y., Zheng, B., Zhang, Q., Huang, T., Kimoto, T., Chang,
638 D., Pöschl, U., Cheng, Y. F., and He, K. B.: Exploring the severe winter haze in Beijing: the impact of
639 synoptic weather, regional transport and heterogeneous reactions, *Atmos. Chem. Phys.*, 15, 2969–2983,
640 doi:10.5194/acp-15-2969-2015, 2015.

641 Zheng, G. J., Duan, F. K., Ma, Y. L., Zhang, Q., Huang, T., Kimoto, T. K., Cheng, Y. F., Su, H., and He, K. B.:
642 Episode-based evolution pattern analysis of haze pollution: method development and results from
643 Beijing, China, *Environ. Sci. Technol.*, 50, 4632–4641, 2016.

644 Zhong, J., Zhang, X., Dong, Y., Wang, Y., Liu, C., Wang, J., Zhang, Y., and Che, H.: Feedback effects of
645 boundary-layer meteorological factors on cumulative explosive growth of PM_{2.5} during winter heavy
646 pollution episodes in Beijing from 2013 to 2016, *Atmos. Chem. Phys.*, 18, 247–258,
647 <https://doi.org/10.5194/acp18-247-2018>, 2018a.

648 Zhong, J., Zhang, X., Wang, Y., Liu, C., and Dong, Y.: Heavy aerosol pollution episodes in winter Beijing
649 enhanced by radiative cooling effects of aerosols, *Atmos. Res.*, 209, 59-64, 2018b.

650 Zhong, J., Zhang, X., Wang, Y., Sun, J., Zhang, Y., Wang, J., Tan, K., Shen, X., Che, H., and Zhang, L.:
651 Relative contributions of boundary-layer meteorological factors to the explosive growth of PM_{2.5} during
652 the red-alert heavy pollution episodes in Beijing in December 2016, *J Meteorol. Res.*, 31, 809–819,
653 2017.

654 Zhou, C., Gong, S., Zhang, X., Liu, H., Xue, M., Cao, G., An, X., Che, H., Zhang, Y., and Niu, T.: Towards
655 the improvements of simulating the chemical and optical properties of Chinese aerosols using an
656 online coupled model CUACE/Aero, *Tellus B*, 64, 18965, doi:10.3402/tellusb.v64i0.18965, 2012.

657 Zhou, C., Zhang, X., Gong, S., Wang, Y., and Xue, M.: Improving aerosol interaction with clouds and
658 precipitation in a regional chemical weather modeling system, *Atmos. Chem. Phys.*, 16, 145-160, 2016.

659

660

661

662

Table 1. Physical and chemical processes in GRAPES_CUACE

| Process | Option | Reference |
|-------------------------|-----------------|--------------------------|
| Explicit precipitation | WDM6 | Lim and Hong (2010) |
| Cumulus cloud | KFETA scheme | Kain (2004) |
| Longwave radiation | Goddard | Chou et al. (2001) |
| Shortwave radiation | Goddard | Chou et al. (1998) |
| Surface layer | SFCLAY scheme | Pleim (2007) |
| PBL | MRF scheme | Hong et al. (1996, 2006) |
| Land surface | SLAB scheme | Kusaka et al. (2001) |
| Gas-phase chemistry | RADM II | Stockwell et al. (1990) |
| Aerosol | CUACE | Zhou et al. (2012) |
| Aerosol direct effect | External mixing | Wang et al. (2015) |
| Aerosol indirect effect | CAUCE+WDM6 | Zhou et al. (2016) |

663

664

665

666

667

668

669

670

671

672

673

674

675

676

677

678

679

680

681

Table 2. Design of sensitivity experiments

| Experiment | Description |
|------------|---|
| EXP1 | Background experiment: ignoring aerosol radiation and conventional diffusion coefficient of chemical tracers by PBL scheme in GRAPES_CUACE |
| EXP2 | Online AF online and conventional diffusion coefficient of chemical tracers by PBL scheme in GRAPES_CUACE |
| EXP3 | Online AF and DC of chemical tracers set to 20% of conventional diffusion coefficient calculated by PBL scheme, representing compensation for the deficient description of extremely weak turbulent diffusion by the PBL scheme; diffusion coefficient in physical and dynamic processes the same as EXP1 |

682

683

684
685
686
687
688
689
690
691
692
693
694
695
696
697
698
699
700
701
702
703
704
705

Table 3. VOCs in the emissions data

| VOC | Full name |
|-------|---|
| ALD | Acetaldehyde and higher aldehydes |
| CH4 | Methane |
| CSL | Cresol and other hydroxy substituted aromatics |
| ETH | Ethane |
| HC3 | Alkanes w/ $2.7 \times 10^{-13} > \text{kOH} < 3.4 \times 10^{-12}$ |
| HC5 | Alkanes w/ $3.4 \times 10^{-12} > \text{kOH} < 6.8 \times 10^{-12}$ |
| HC7 | w/kOH $> 6.8 \times 10^{-12}$ |
| HCHO | Formaldehyde |
| ISOP | Isoprene |
| KET | Ketones |
| OL2 | Ethene |
| OLI | Internal olefins |
| OLT | Terminal olefins |
| ORA2 | Acetic and higher acids |
| PAR | Paraffin carbon bond |
| TERPB | Monoterpenes |
| TOL | Toluene and less reactive aromatics |
| XYL | Xylene and more reactive aromatics |

706

707

708

Table 4. Observed and modeled daily AOD (* stands for shortage of observation)

709

710

711

712

713

714

715

716

717

718

719

| Date | Shijiazhuang | | Beijing | | Xianghe | |
|------|--------------|-------|---------|-------|---------|-------|
| | OBS | MODEL | OBS | MODEL | OBS | MODEL |
| 15 | 0.46 | 0.55 | 0.07 | 0.12 | 0.10 | 0.15 |
| 16 | 0.62 | 0.60 | 0.14 | 0.18 | 0.60 | 0.40 |
| 17 | 1.30 | 1.10 | 0.50 | 0.56 | 1.33 | 1.05 |
| 18 | 1.42 | 1.20 | 0.69 | 0.75 | 0.87 | 0.97 |
| 19 | 1.26 | 1.30 | 0.50 | 0.86 | 0.96 | 0.90 |
| 20 | * | 1.20 | 1.90 | 1.70 | * | 1.50 |
| 21 | * | 0.65 | 1.76 | 1.50 | 1.78 | 1.60 |
| 22 | 0.18 | 0.30 | 0.10 | 0.20 | 0.18 | 0.22 |

720

721

Table 5. Observed and modeled daily SSA (* stands for shortage of observation).

722

723

724

725

726

| Date | Shijiazhuang | | Beijing | | Xianghe | |
|------|--------------|-------|---------|-------|---------|-------|
| | OBS | MODEL | OBS | MODEL | OBS | MODEL |
| 15 | 0.83 | 0.85 | 0.81 | 0.83 | 0.86 | 0.84 |
| 16 | 0.83 | 0.85 | 0.88 | 0.86 | 0.92 | 0.86 |
| 17 | 0.88 | 0.89 | 0.88 | 0.90 | 0.93 | 0.90 |
| 18 | 0.87 | 0.89 | 0.91 | 0.92 | 0.90 | 0.90 |
| 19 | 0.86 | 0.91 | 0.90 | 0.93 | 0.92 | 0.91 |
| 20 | * | 0.90 | * | 0.93 | * | 0.92 |
| 21 | * | 0.88 | 0.93 | 0.93 | * | 0.90 |
| 22 | 0.82 | 0.83 | 0.84 | 0.86 | 0.88 | 0.84 |

727

728

Figure captions

729 **Fig. 1.** (a) Model domain and location of Jing–Jin–Ji. (b) Geographic location and topography of Jing–Jin–
730 Ji. Blue dots are the locations of PM_{2.5} observations; red triangles are the locations of automatic weather
731 stations; yellow stars are the two sounding stations; black crosses are the CARSNET and AEROSNET
732 stations.

733 **Fig. 2.** Geopotential height (color-shaded; gp10m), temperature (dashed black contours; K) and wind (wind
734 bars; m/s) in the (a) upper (500 hPa) and (b) middle (700 hPa) atmosphere, and geopotential height and
735 wind in the (c) lower atmosphere (850 hPa) and (d–f) PBL (900, 950, 1000 hPa), at 0000 UTC 19
736 December 2016.

737 **Fig. 3.** Observed and modeled wind speed and temperature at the surface (upper panels), and the PBL-mean
738 wind speed and temperature (lower panels), from the results of EXP1, EXP2 and EXP3 for Beijing, Xingtai,
739 and the average for Jing–Jin–Ji as a whole, during 15–24 December 2016.

740 **Fig. 4.** Mean observed (OBS_PM_{2.5}) and modeled PM_{2.5} concentration ($\mu\text{g}/\text{m}^3$) of the PM_{2.5} explosive
741 growth stage, from the results of EXP1, EXP2 and EXP3 (PM_{2.5}_EXP1, PM_{2.5}_EXP2 and PM_{2.5}_EXP3,
742 respectively).

743 **Fig. 5.** Mean percentage change in SDSRF (W/m^2) owing to (a) aerosols and (b) aerosols+ decrease in
744 turbulent diffusion coefficient during the explosive growth stage.

745 **Fig. 6.** Profiles of average temperature change in Jing–Jin–Ji owing to AF (K) during 15–20 December
746 2016.

747 **Fig. 7.** Sounding-observed and modeled temperature profiles in EXP1 and EXP2 during the (a) climbing
748 stage and (b) explosive growth stage in Beijing and Xingtai.

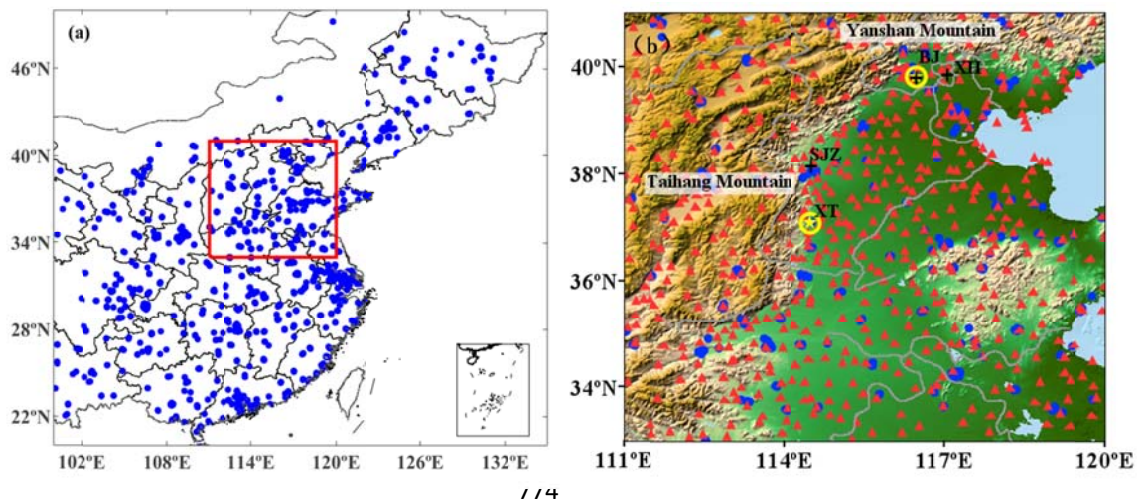
749 **Fig. 8.** Hourly change of PM_{2.5}_OBS, PM_{2.5}_EXP1, PM_{2.5}_EXP2, and PM_{2.5}_EXP3 ($\mu\text{g}/\text{m}^3$), together with
750 the diffusion coefficient (DC) at 950 hPa of the three experiments (DC_EXP1, DC_EXP2, DC_EXP3)
751 during 15–22 December 2016 in (a) Beijing and (b) Xingtai.

752 **Fig. 9.** Diagrammatic sketch of the contributions of AF and decrease in turbulent diffusion coefficient
753 (DTD) to the PM_{2.5} explosive growth.

754

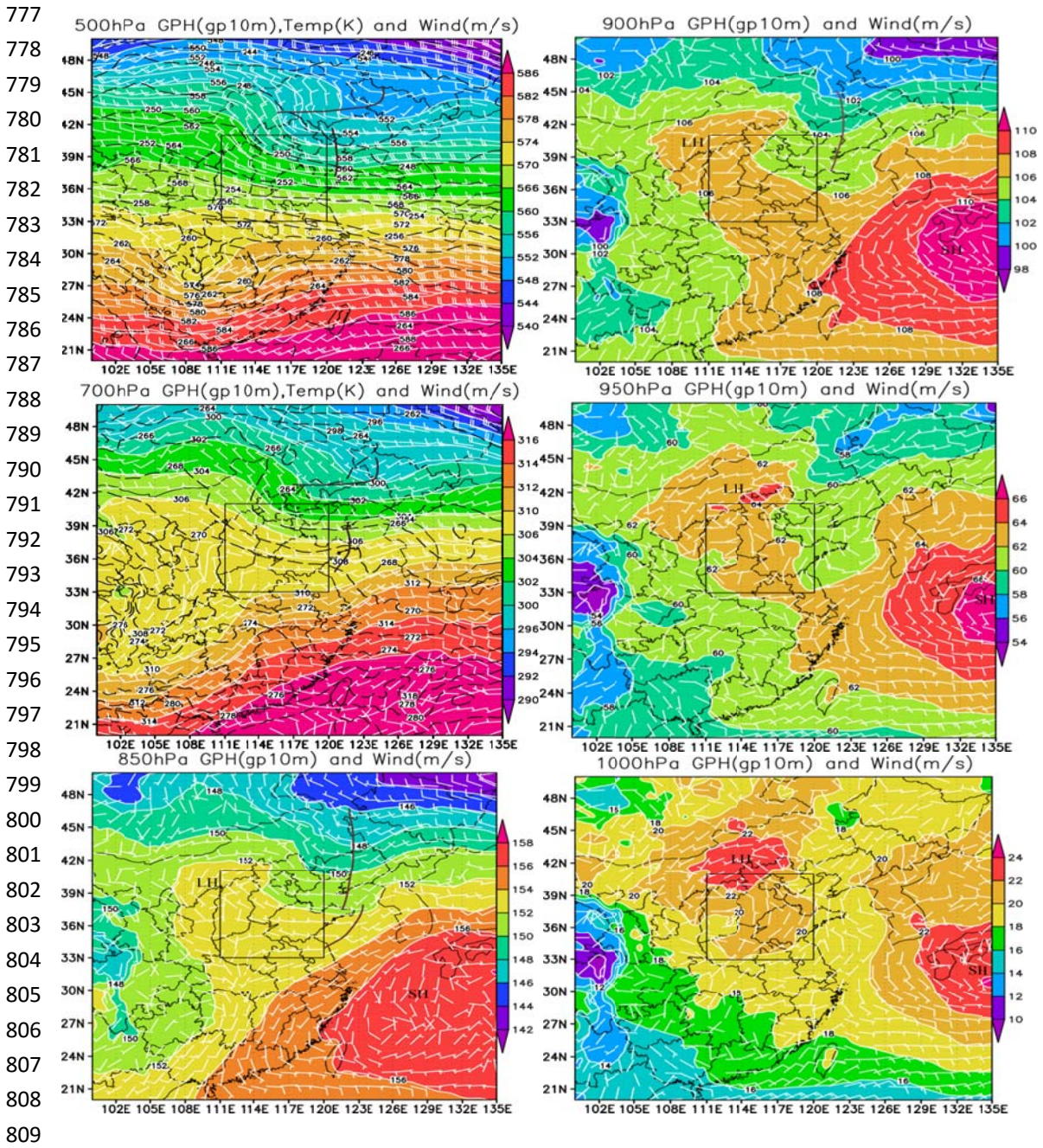
755

757
758
759
760
761



774 **Fig. 1.** (a) Model domain and location of Jing–Jin–Ji. (b) Geographic location and topography of Jing–Jin–
775 Ji. Blue dots are the locations of PM_{2.5} observations; red triangles are the locations of automatic weather
776 stations; yellow stars are the two sounding stations; black crosses are the CARSNET and AEROSNET
777 stations.

775
776

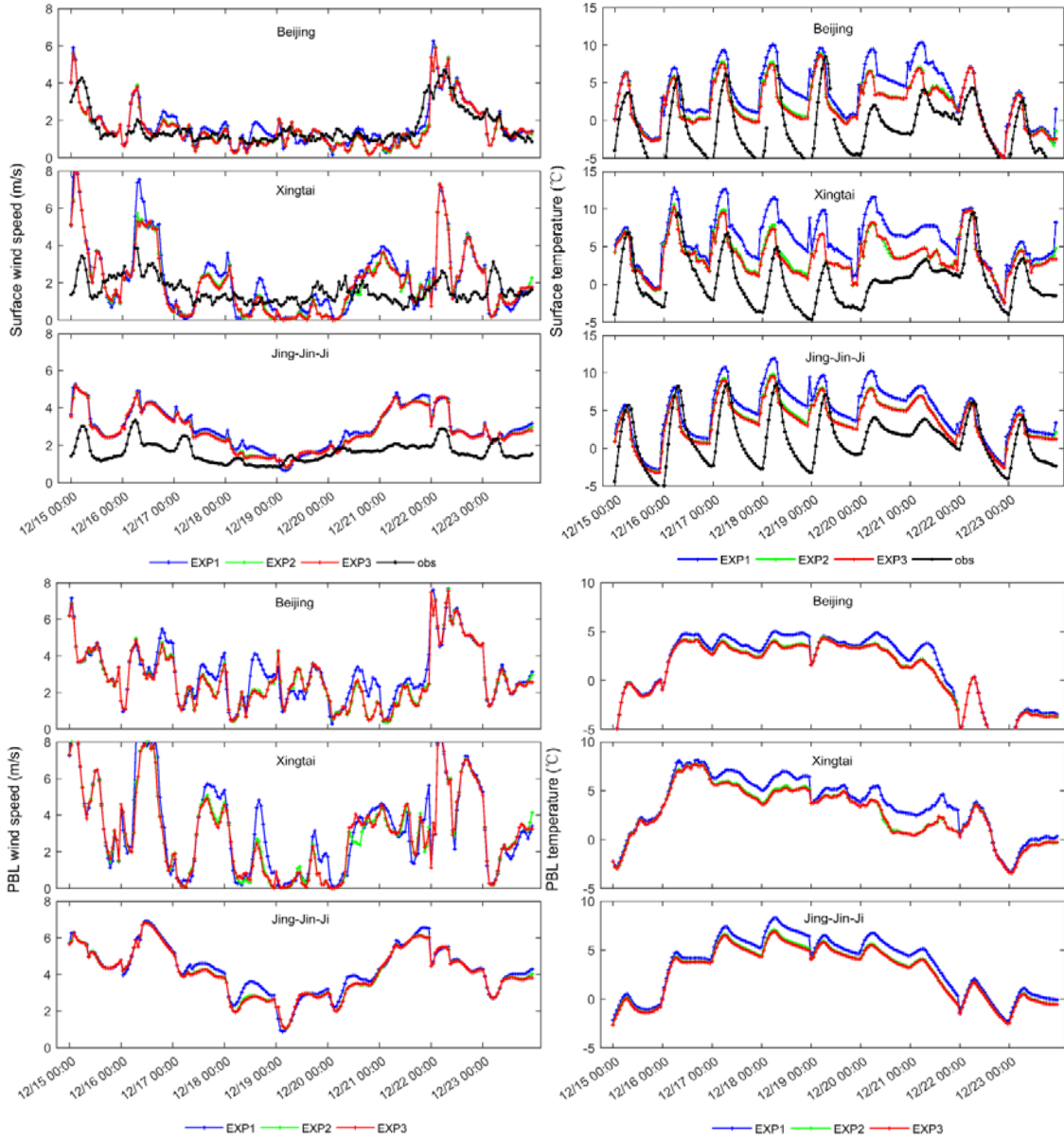


813 **Fig. 2.** Geopotential height (color-shaded; gp10m), temperature (dashed black contours; K) and wind (wind
814 bars; m/s) in the (a) upper (500 hPa) and (b) middle (700 hPa) atmosphere, and geopotential height and
815 wind in the (c) lower atmosphere (850 hPa) and (d-f) PBL (900, 950, 1000 hPa), at 0000 UTC 19
816 December 2016.

814

815

815
 816
 817
 818
 819
 820
 821
 822
 823
 824
 825
 826
 827
 828
 829
 830
 831
 832
 833
 834
 835
 836
 837
 838
 839
 840
 841
 842
 843
 844
 845
 846
 847
 848
 849



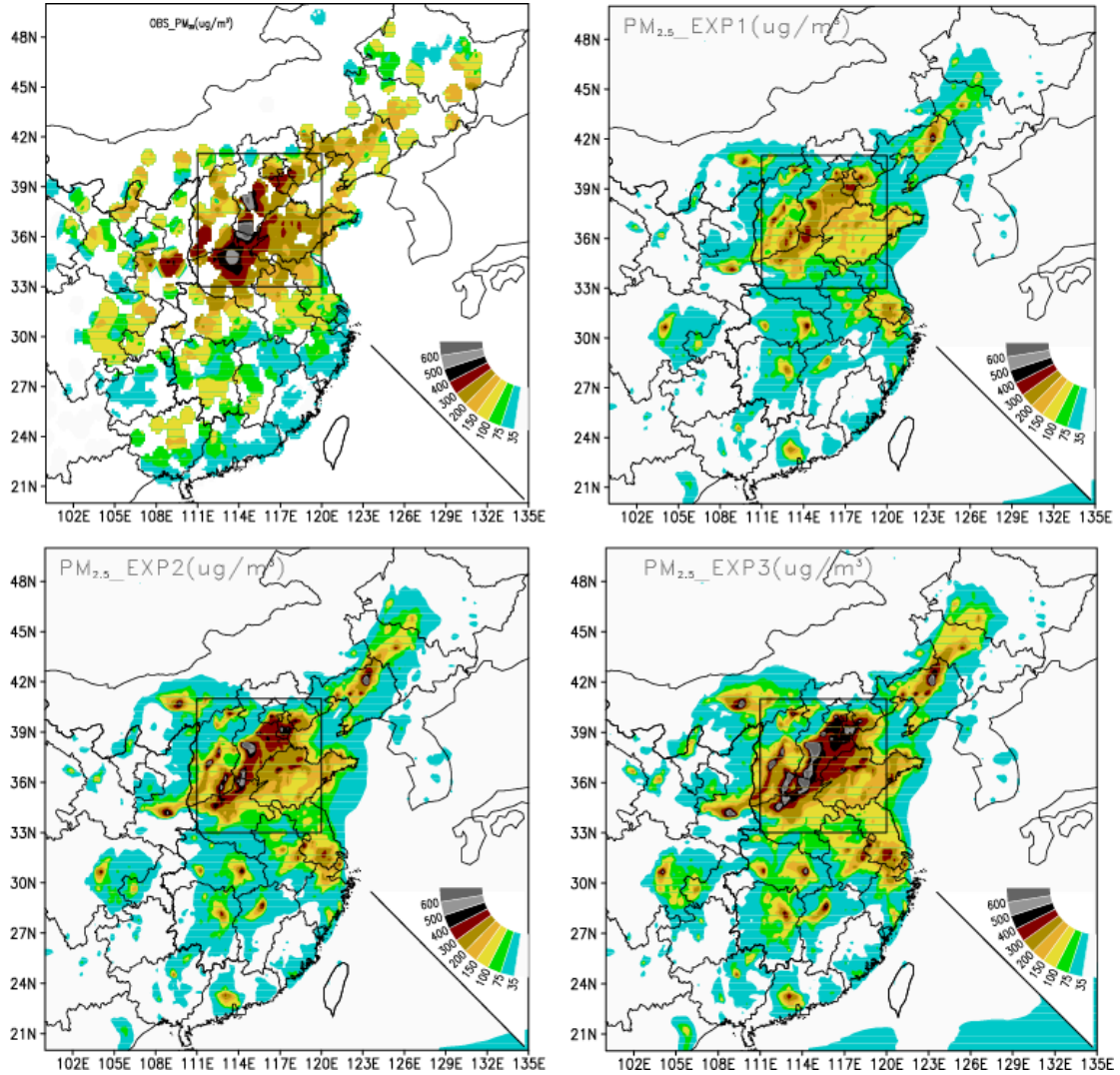
850 **Fig. 3.** Observed and modeled wind speed and temperature at the surface (upper panels), and the PBL-mean
 851 wind speed and temperature (lower panels), from the results of EXP1, EXP2 and EXP3 for Beijing, Xingtai,
 852 and the average for Jing–Jin–Ji as a whole, during 15–24 December 2016.

853
 854

855

856

857



858

859

860 **Fig. 4.** Mean observed (OBS_PM_{2.5}) and modeled PM_{2.5} concentration ($\mu\text{g}/\text{m}^3$) of the PM_{2.5} explosive

861 growth stage, from the results of EXP1, EXP2 and EXP3 (PM_{2.5}_EXP1, PM_{2.5}_EXP2 and PM_{2.5}_EXP3,

862 respectively).

863

864

865

866

867

869
870
871
872
873
874
875
876
877
878
879
880
881
882
883
884
885
887
888
888
889
890
891

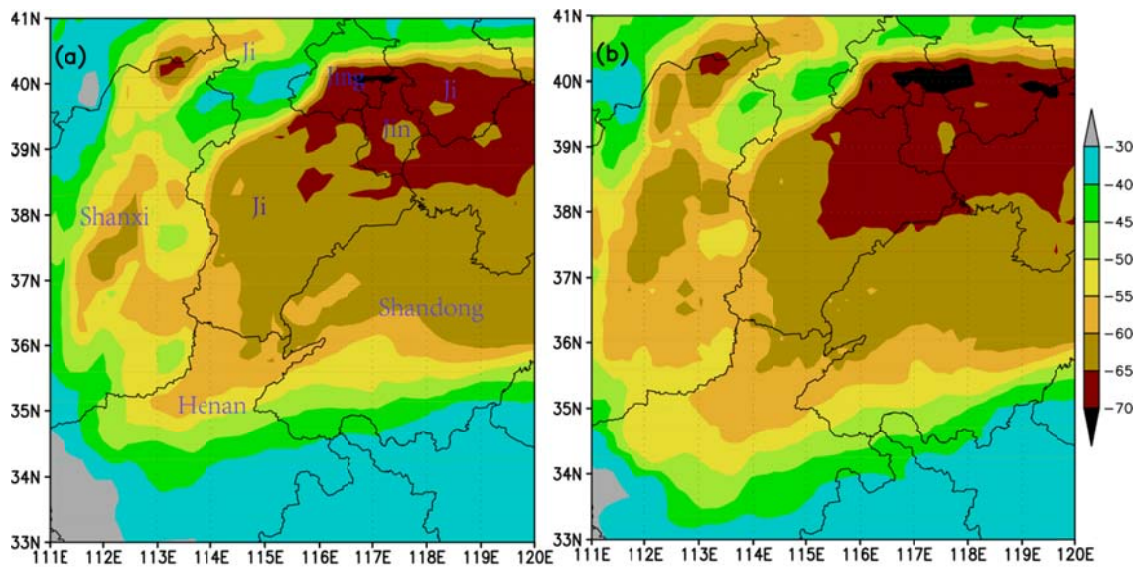


Fig. 5. Mean percentage change in SDSRF (W/m^2) owing to (a) aerosols and (b) aerosols+ decrease in turbulent diffusion coefficient during the explosive growth stage.

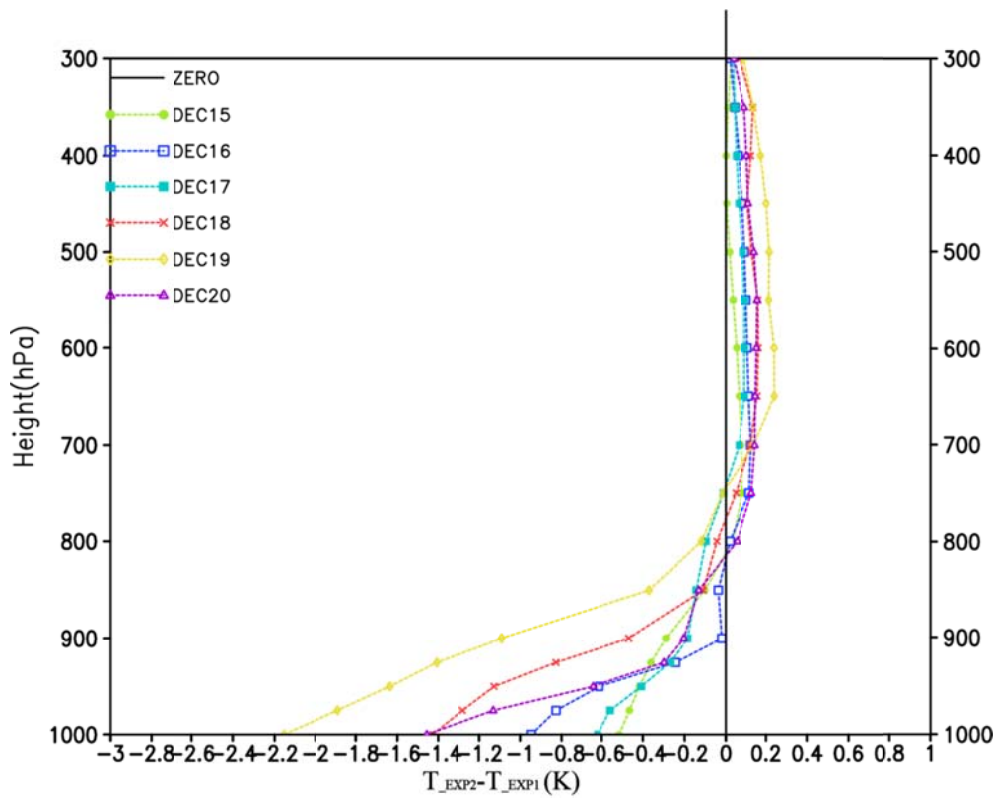
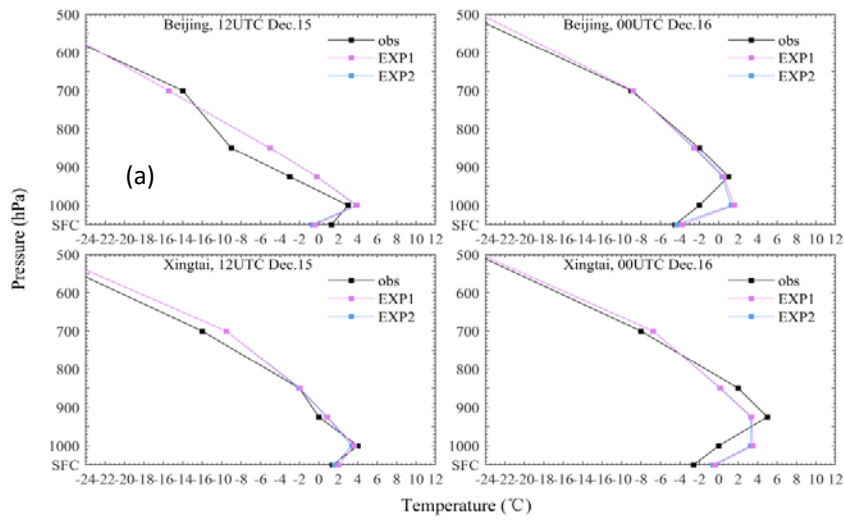
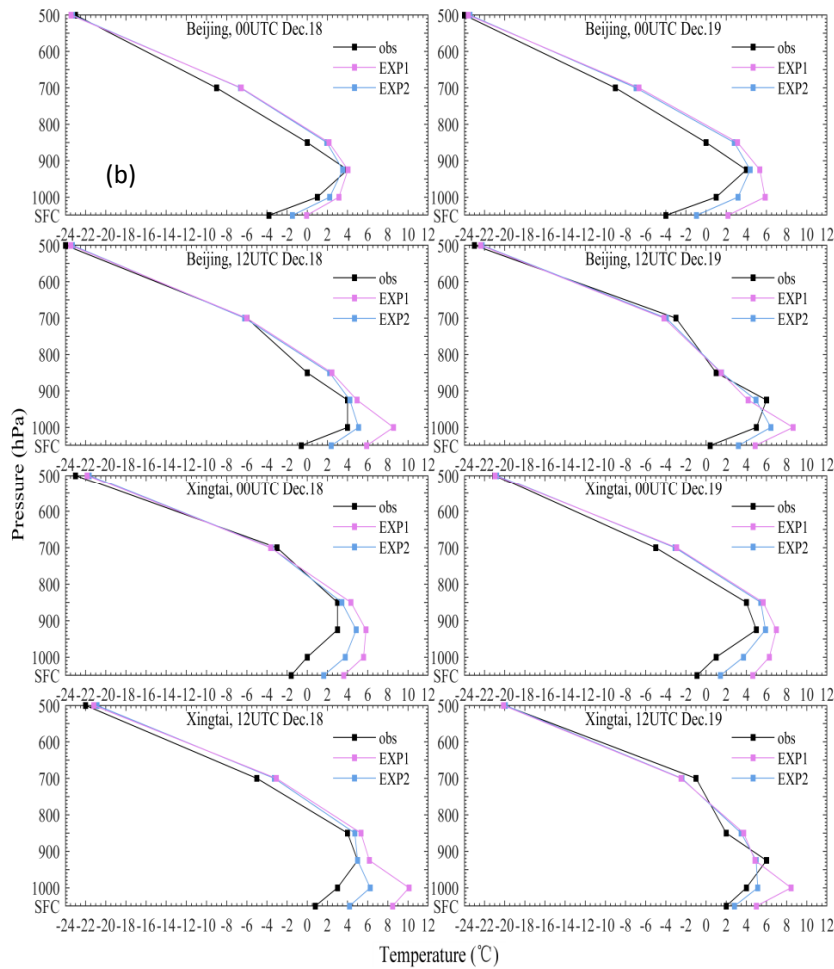


Fig. 6. Profiles of average temperature change in Jing–Jin–Ji owing to AF (K) during 15–20 December 2016.



929

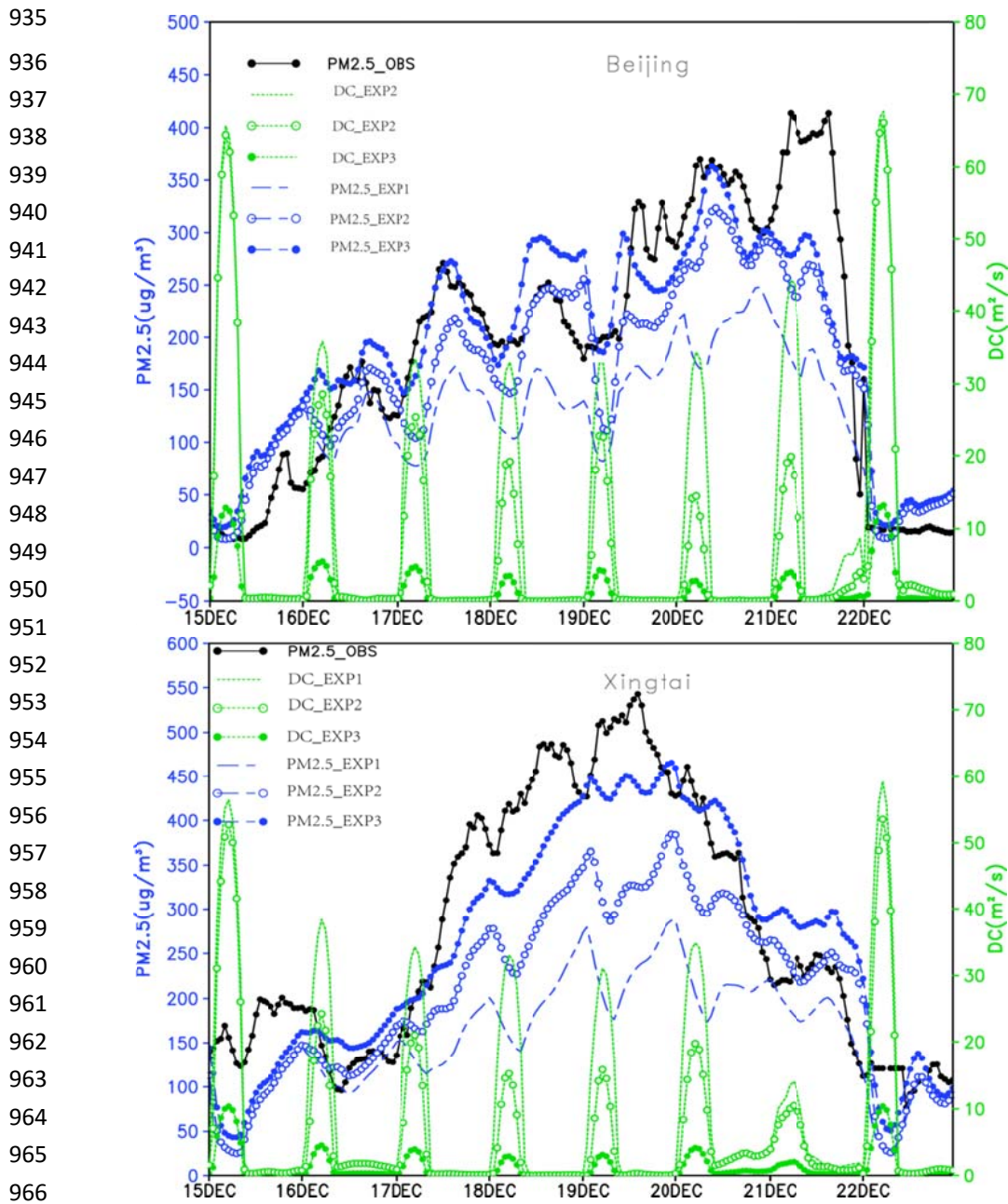


930

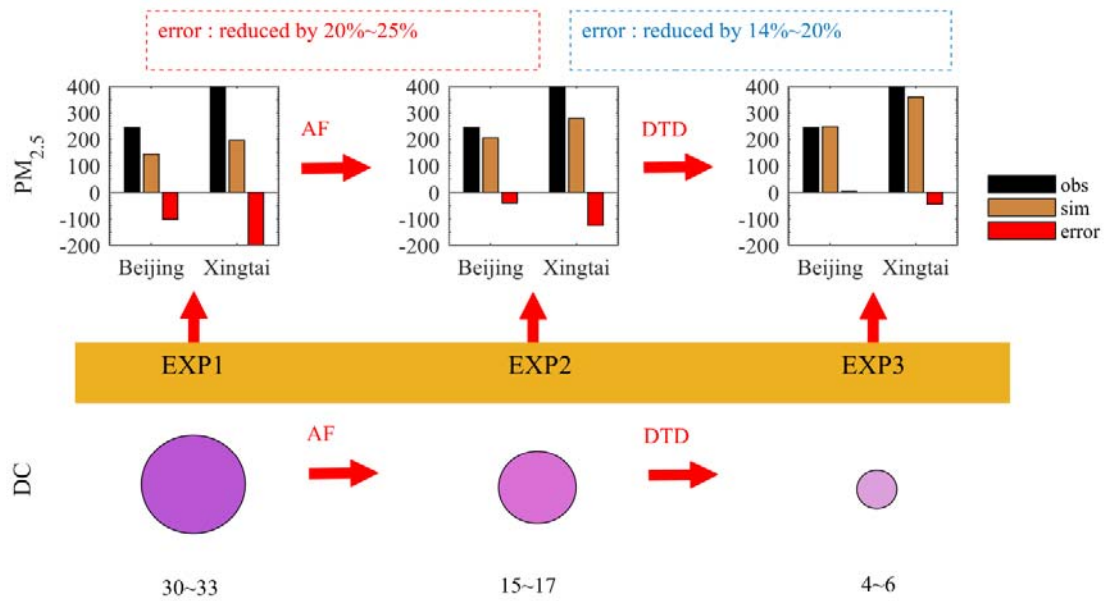
931 **Fig. 7.** Sounding-observed and modeled temperature profiles in EXP1 and EXP2 during the (a) climbing

932 stage and (b) explosive growth stage in Beijing and Xingtai.

933



969 **Fig. 8.** Hourly change of $PM_{2.5_OBS}$, $PM_{2.5_EXP1}$, $PM_{2.5_EXP2}$, and $PM_{2.5_EXP3}$ ($\mu\text{g}/\text{m}^3$), together with
 970 the diffusion coefficient (DC) at 950 hPa of the three experiments (DC_EXP1, DC_EXP2, DC_EXP3)
 971 during 15–22 December 2016 in (a) Beijing and (b) Xingtai.



969
 970
 971
 972
 973
 974
 975

Fig. 9. Diagrammatic sketch of the contributions of AF and decrease in turbulent diffusion coefficient (DTD) to the PM_{2.5} explosive growth.