# 1 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

- 5 Junting Zhong<sup>1</sup>, Xiaoye Zhang<sup>1, 2</sup>, Yunsheng Dong<sup>3</sup>, Yaqiang Wang<sup>1</sup>, Cheng Liu<sup>2, 3, 4</sup>,
- 6 Jizhi Wang<sup>1</sup>, Yangmei Zhang<sup>1</sup>, Haochi Che<sup>5</sup>
- 7 <sup>1</sup>State Key Laboratory of Severe Weather & Key Laboratory of Atmospheric Chemistry of CMA,
- 8 Chinese Academy of Meteorological Sciences, China.
- <sup>2</sup>Center for Excellence in Regional Atmospheric Environment, IUE, Chinese Academy of Sciences,
   China.
- <sup>11</sup> <sup>3</sup>Key Laboratory of Environmental Optics and Technology, Anhui Institute of Optics and Fine Mechanics,
- 12 Chinese Academy of Sciences, China.
- <sup>4</sup>School of Earth and Space Sciences, University of Science and Technology of China, Hefei 230026,
- 14 China
- 15 <sup>5</sup>Department of Physics, University of Oxford, UK.
- 16 Correspondence to: X.Y. Zhang (xiaoye@camscma.cn)

17 Abstract. In January of 2013, February of 2014, December of 2015, and December of 2016 to January 10th of 2017, 12 persistent heavy aerosol pollution episodes (HPEs) occurred in Beijing, which has 18 attracted special attention from the public. During the HPEs, the precise cause of PM<sub>2.5</sub> explosive growth 19 20 (mass concentration at least doubled in several to ten hours) is uncertain. Here, we analyzed and 21 estimated relative contributions of boundary-layer meteorological factors to such growth, using ground 22 and vertical meteorological data. Beijing HPEs are generally characterized by the transport stage (TS), 23 whose aerosol pollution formation is primarily caused by pollutants transported from the south of Beijing, 24 and the cumulative stage (CS), in which the cumulative explosive growth of PM2.5 mass is dominated by 25 stable atmospheric stratification characteristic of southerly slight or calm winds, near-ground anomalous 26 inversion, and moisture accumulation. During the CSs, observed southerly weak winds facilitate local 27 pollutant accumulation by minimizing horizontal pollutant diffusion. Established from TSs, elevated 28 PM2.5 levels scatter more solar radiation back to the space to reduce near-ground temperature, which very 29 likely causes anomalous inversion. This surface cooling by PM2.5 decreases near-ground saturation vapor 30 pressure and increases relative humidity significantly; the inversion subsequently reduces vertical

31	turbulent diffusion and boundary layer height to trap pollutants and accumulate water vapor. Appreciable
32	near-ground moisture accumulation (RH>80%) would further enhance aerosol hygroscopic growth and
33	accelerate liquid-phase and heterogeneous reactions, in which incompletely quantified chemical
34	mechanisms need more investigation. Noted positive meteorological feedback on PM2.5 mass explains
35	over 70% of cumulative explosive growth.
36	
37	Keyword: Southerly transport; anomalous inversion; moisture accumulation; meteorological feedback.

# 39 1 Introduction

40 Since a persistent heavy fog and haze event occurred in eastern China in January 2013, fine 41 particulate matter smaller than 2.5 µm in diameter (PM<sub>2.5</sub>), as a key component of pollution episodes, 42 has drawn wide attention all over China. Elevated PM2.5 leads a sharp decrease in visibility that affects 43 economic activities by causing traffic disruptions and contains toxic substances that affect respiratory 44 and circulatory system (Chen et al., 2013;Bai et al., 2007). The interaction between aerosol and radiation 45 directly and indirectly affects weather and climate (Zhang et al., 2013;Zhang et al., 2015b;Wei et al., 46 2011;Boucher et al., 2013;Wang et al., 2010). China has experienced heavy aerosol pollution episodes 47 recently, with PM2.5 reaching unprecedentedly high levels in many cities, particularly Beijing and its 48 vicinity (BIV), which is one of the nation's most polluted regions (Zhang et al., 2012).

49 To elucidate the causes of such heavy pollution episodes, a variety of explanations have been 50 proposed (Huang et al., 2014;Sun et al., 2014a;Sun et al., 2014b;Wang et al., 2014b;Wang et al., 51 2014c). Previous studies found that atmospheric conditions represented one critical parameter in 52 regulating the cycles of pollution episodes in Beijing in autumn 2013 (Guo et al., 2014;Zhang et al., 2009) 53 and in the North China Plain and other areas in China (Zhang et al., 2015c). During one pollution episode, 54 an analysis of atmospheric background fields revealed dynamic and thermodynamic effects substantially 55 affected pollution formation (Zhang et al., 2014). Specifically, aerosol pollution in Beijing was possibly 56 contributed by southerly/southwesterly surface wind (Wang et al., 2013b). This likely attribution was 57 further verified by source apportionment from the Beijing Environmental Protection Bureau in 58 2012~2013. In addition, aerosol pollution can be formed by secondary aerosol formation through 59 atmospheric chemical reactions, including liquid-phase reactions, in which aqueous SO<sub>2</sub> is oxidized by 60 NO<sub>2</sub>, H<sub>2</sub>O<sub>2</sub> and O<sub>3</sub> to form sulfate, and heterogeneous reactions, in which NO<sub>2</sub> and N<sub>2</sub>O<sub>5</sub> form nitrates 61 with water (Zheng et al., 2015a; Cheng et al., 2016).

Although these cited studies existed, the formation mechanism during different stages for heavy aerosol pollution in Beijing, especially the explosive growth stage of PM<sub>2.5</sub> mass concentration, is still not clear. Previous studies focused more on whether unfamiliar chemical mechanisms were not or inadequately considered in Beijing, a region with high concentrations of various aerosol components 66 (Wang et al., 2014b). This view was questioned by subsequent research, suggesting that such rapid 67 growth is mainly attributable to the regional transport of clean and polluted air mass, which derived from 68 the comparison between surface meteorological factors and the PM2.5 mass concentration in several cities 69 of the North China Plain (Zheng et al., 2015b). However, the attribution of such growth's drivers is 70 unreasonable occasionally, because the rapid growth may occur with weak surface winds and stable 71 stratification, which are unfavorable for transport. Then vertical meteorological variations in the 72 boundary layer (BL) in one autumnal episode have been analyzed, which significantly affect the PM<sub>2.5</sub> 73 mass concentration near the ground (Hua et al., 2016). However, in the absence of long-term observations 74 of meteorological factors and pollutant concentrations, most research concerning pollution causes 75 focuses on one or several consecutive pollution episodes in a certain time, and almost no research attempt 76 to investigate, conclude and quantify the contributions of meteorological factors to the majority of heavy 77 pollution episodes since 2013, particularly the feedback effect of meteorological factors during explosive 78 growth processes. Such investigations will definitely provide a clearer understanding of roles that various 79 vertical meteorological factors play in heavy pollution episodes. Therefore, this paper primarily uses 80 vertical measurements of meteorological factors in the BL from 2013 to 2016, investigates their 81 contributions to the explosive growth of PM<sub>2.5</sub> during the heavy pollution episodes in Beijing, and also 82 attempts to quantify the effect of meteorological factors on the explosive growth of PM<sub>2.5</sub> levels.

# 84 2 Methods

85	In this study, the following data are used. (1) Hourly PM <sub>2.5</sub> mass concentration measured by state-
86	controlled stations of the Ministry of Environmental Protection in Beijing and Baoding. PM2.5 mass
87	concentrations of urban stations were averaged to represent urban pollution conditions. (2) Atmospheric
88	vertical observations twice daily at 0800 Beijing Time (BJT) and 2000 BJT, including winds, temperature
89	and relative humidity (RH), measured using L-band radiosonde radar at the observatory (54511) in
90	southern Beijing from 1 January 2013 to 31 January 2013, 1 February 2014 to 28 February 2014, 26
91	November 2015 to 31 December 2015 and 21 December 2016 to 10 January 2017. The observatory is
92	located on the edge of urban area of Beijing, so it could be used to represent urban vertical meteorological
93	conditions to some degree due to the regional change characteristics of air masses. (3) Hourly ground-
94	level meteorological observations from automatic weather stations (AWSs) provided by the National
95	Meteorological Information Center of the China Meteorological Administration. (4) Lidar observations
96	were measured every fifteen minutes by one Mie-elastic backscatter polarization lidar emitting short
97	pulses of 20 Hz at 532nm in the Institute of Atmospheric Physics, located in the northern urban area of
98	Beijing. The optical parameters of the aerosol particles were retrieved by the backscattering signals. Then
99	the vertical profiles of the aerosol extinction coefficient and linear depolarization ratio were obtained
100	based on the assumptive lidar ratios as 50 for aerosols using the Fernald's method (Fernald, 1984;Lv et
101	al., 2017). (5) A parameterized index, PLAM (Parameter Linking Aerosol Pollution and Meteorological
102	Elements), calculated with the observations from the observatory (54511). PLAM was built as a function
103	of the following parameters:
104	PLAM (F) $\in$ f (p, t, w, rh, e, s, c',) (1)
105	where p, t, w, rh, e, s, and c' represent air pressure, air temperature, winds, relative humidity,
106	evaporability, stability, and effective parameter associated with the contribution of air pollution $\beta(c')$ ,
107	respectively. Plam is further attributed to two major separate factors: (1) initial meteorological conditions
108	$\alpha(m)$ associated with the atmospheric condensation processes and (2) a dynamic effective parameter
109	associated with the initial contribution of air pollution $\beta(c')$ :
110	$PLAM = \alpha(m) \times \beta'(c). $ (2)

- 111 It mainly indicates the regional atmospheric stability and air condensation ability. The details of its
- 112 calculation are presented in previous studies (Wang et al., 2013a;Zhang et al., 2015c;Zhang et al.,
- 113 2009;Wang et al., 2012).

# 114 **3** Results and discussion:

# 115 3.1 Characteristics of explosive growth in HPEs

A period during which the  $PM_{2.5}$  level is less than 35 µg m<sup>-3</sup> is defined as a clean period based on the  $PM_{2.5}$  daily mean mass concentration limit in the primary standard of China's national environmental quality standards, while a pollution episode is referred to as an episode during which the  $PM_{2.5}$  exceeds 80 µg m<sup>3</sup> for 3 consecutive days between two clean periods. Pollution episodes with peak  $PM_{2.5}$  values less than 300 µg m<sup>-3</sup> or more than 400 µg m<sup>-3</sup> are termed light pollution episodes (LPEs) or heavy pollution episodes (HPEs), respectively.

Based on the urban PM<sub>2.5</sub> monthly mean mass concentration in winter Beijing from 2013 to 2016, the months with highest mass concentration each year were selected to represent the severe PM<sub>2.5</sub> pollution conditions in winter, which are January in 2013, February in 2014, December in 2015 and December in 2016 respectively. These months are termed the wintertime pollution period (WPP) for the convenience of further investigation.

During the WPP, 12 HPEs occur in total (Fig. 1~4 (dark gray)), whose PM<sub>2.5</sub> mass concentration is 244.3  $\mu$ g m<sup>-3</sup> on average. The maximum mean value (307.4  $\mu$ g m<sup>-3</sup>) appears in HPE<sub>1</sub>, which has been analyzed in detail in a variety of papers (Zhang et al., 2013;Zhang et al., 2014). The concentrations of HPE<sub>6</sub> and HPE<sub>10</sub> are 304.2  $\mu$ g m<sup>-3</sup> and 294.5  $\mu$ g m<sup>-3</sup> respectively, which are slightly lower than HPE<sub>1</sub>. The minimum mean concentration of PM<sub>2.5</sub> occurs in HPE<sub>8</sub> (160.4  $\mu$ g m<sup>-3</sup>), which is nearly twice as much as the mean annual mass concentration of PM<sub>2.5</sub> in 2015, nevertheless.

Typical PM<sub>2.5</sub> rising processes (color-mark) in HPEs were selected, which appeared in 11 of the 12 HPEs. The green-mark processes are tentatively referred to as rising processes, since they generally keep rising consistently with relatively strong southerly winds compared with subsequent growth and vary sensitively and rapidly in response to wind shift from northerly to southerly in the BL. During HPEs, the 137 growth processes in which the PM<sub>2.5</sub> mass concentration is at least doubled in several or ten hours in the 138 later period of HPEs are termed explosive growth processes. The red-mark explosive growth processes 139 are tentatively termed cumulative explosive growth processes because of anomalous inversion 140 facilitating pollutant accumulation. The purple-mark explosive growth processes are tentatively known 141 as convergent explosive growth processes, for local wind convergence occurs (Fig. 6) with weak wind 142 velocity and no anomalous inversion. The early stages of HPEs during which PM<sub>2.5</sub> keeps rising are 143 defined as transport stages (TSs), while the later stages during which cumulative/convergent explosive 144 growth appears are termed cumulative stages (CSs).

# 145 3.2 Meteorological causes of the explosive growth in HPEs

# *3.2.1* PM<sub>2.5</sub> pollution formation is primarily caused by pollutants transported from the south of Beijing, which subsequently worsen weather conditions

We found that during clean periods occur mostly strong northwesterly winds whose velocity increases with height; during the HPEs, the southwesterly winds with dramatically decreased velocity were most frequent (Fig. 1-4 (a, b)). Strong northerly winds and weak southerly winds closely correspond to the clean periods and the HPEs respectively, because northwesterly winds, which are from less populated north mountainous areas, carry unpolluted air masses while southerly winds carry polluted air masses from more populated and polluted southern industrial regions (Jia et al., 2008;Liu et al., 2013;Guo et al., 2014).

155 During the TSs with southerly winds, air temperature and moisture substantially increase compared 156 with clean periods with northerly winds (Fig. 1-4 (b, c, d)), which indicates warm and humid southerly 157 airflow transport more water vapor and heat into Beijing. During 15 rising processes ((green lines)), 158 nearly no striking near-ground (<250m) moisture accumulation appears; no anomalous inversion appears 159 except brief weak inversion, which suggest that vertical variations of temperature and RH are unlike to 160 primarily cause such rising. Nevertheless, weak inversion and more near-ground moisture favor growth. 161 If we assume that the primary cause of this rising is pollution accumulation due to local emissions, the rising needs to coincide with light  $(0.3 \sim 1.5 \text{ m s}^{-1})$  or calm  $(0 \sim 0.2 \text{ m s}^{-1})$  air observed during the later 162

TSs instead of slight  $(1.6 \sim 3.3 \text{ m s}^{-1})$  or gentle  $(3.4 \sim 5.4 \text{ m s}^{-1})$  breeze observed during the early TSs, 163 because weaker winds result in a stagnant condition, which is more favorable for local accumulation. 164 165 However, the majority of later TSs with calm air do not exhibit such rising (Fig. 1-4 (a, b)), which 166 suggests local emissions under weak winds are likely conducive but not dominant with respect to rising. 167 Before rising processes during HPE<sub>1 $\sim$ 2</sub>, the urban PM<sub>2.5</sub> mass concentration of Baoding (light gray 168 lines), which is typically representative of pollution conditions in the south of Beijing, was much higher 169 than Beijing; the winds in Beijing rapidly shifted from northerly to southerly. Then the rising (green lines) 170 occurred, consistently with southerly slight or gentle breezes in the BL (green boxes). The southerly air 171 mass move more than 288 km d<sup>-1</sup> below 500 m (estimated from the measured wind speed), which are fast 172 enough to transport pollutants to Beijing. Similar conditions appeared in 8 of other 9 HPEs with rising. 173 Such processes indicate southerly pollutant transport is primarily responsible for the rising, given the 174 pollution transport pathway of the southwest wind belt determined by the unique geographic features of 175 the North China Plain, with the Tai-hang Mountains and the Yan Mountains strengthening the southwest wind belt and leading the convergence of pollutant transport in Beijing. (Su et al., 2004). Governed by 176 177 this transport pathway, PM<sub>2.5</sub> mass concentration increased by ~400  $\mu$ g m<sup>-3</sup> from less than 35  $\mu$ g m<sup>-3</sup> in 178 ten hours on 22 January 2013, when winds shifted from northerly to southerly with much higher PM<sub>2.5</sub> 179 concentrations in Baoding.

Pollutants transported from the south of Beijing primarily results in  $PM_{2.5}$  pollution formation in the urban Beijing area, to which possible weak inversion and the near-surface moisture accumulation is conducive. Warm and humid airflow from the south transports more water vapor and pollutants to the North China Plain, which creates the requisite moisture and pollution accumulation conditions for subsequently cumulative explosive growth.

185 *3*.

#### 3.2.2 Worsening meteorological conditions primarily cause cumulative explosive growth

186 Feedback of anomalous inversion on pollutant accumulation

Anomalous inversion occurs during 10 of 12 HPEs (Fig. 1~4 (a, c)). The factors that cause inversion
 in Beijing includes topography, advection and radiation. With the Tai-hang Mountains and the Yan

189 Mountains lying north of Beijing, a cold air mass flows down into the urban of Beijing from the mountain 190 peaks, which occasionally causes topography inversion; advection inversion occurs when a warm and 191 less dense air mass moves over a cold and dense air mass. However, during most cumulative stages, the 192 anomalous inversion appears with slight or calm winds, which suggests that the movement of air masses 193 is not striking, so the contribution of topography and advection to such inversion is limited. The ground 194 emits long-wave radiation at night to reduce near-ground temperature to facilitate inversion occasionally. However, almost no anomalous inversion occurs without pre-existing high PM2.5 mass concentration in 195 196 the WPP (Fig. 1~4), which suggests that the ground radiation are likely conducive to weak/normal 197 inversion, but not dominant with respect to anomalous inversion.

198 Noted anomalous inversion is preceded by existing relatively high PM2.5 levels generally established 199 by the rising processes. Before the cumulative explosive growth, existing aerosols are concentrated 200 below 500 m (Fig. 5). These low-layer aerosols back-scatter amounts of radiation to space (Wang et al., 201 2014a;Gao et al., 2015), and cause a significant reduction in radiation reaching the ground, which further 202 reduces near-ground temperature. These findings indicate that anomalous inversion is primarily due to 203 radiation cooling effect of pre-existing aerosols. Below the inversion, near-ground temperature reduction 204 cools down plumes or thermals of originally warm surface air to decrease thermal turbulence; observed 205 weakened vertical shear of horizontal winds (Fig. 1~4 (b)) produces less vorticity to reduce mechanical 206 turbulence, which further strengthens the existing inversion.

207 Anomalous inversion traps pollution-laden air beneath it due to its strong static stability (Wallace 208 and Hobbs, 2006). It facilitates pollutant accumulation by suppressing vertical air mixing and reducing 209 BL height. During the cumulative explosive growth with anomalous inversion in the HPE10, the turbulent 210 diffusion coefficient rapidly decreases from 100 m<sup>2</sup> s<sup>-1</sup> to 50 m<sup>2</sup> s<sup>-1</sup> (model output of CUACE/Chem, the 211 meso-scale China Meteorological Administration (CMA) Unified Atmospheric Chemistry modelling 212 system, personal communication with Dr. Hong Wang), and similar conditions of turbulent diffusion 213 have been modelled in another pollution episode in Beijing (Wang et al., 2015a; Wang et al., 2015b); the BL height decreases from  $\sim$ 500 m in the early morning, to  $\sim$ 350 m at noon, even to  $\sim$ 250 m at night (Fig. 214 215 5), which coincide with the increase of PM<sub>2.5</sub> from  $\sim$ 200 to  $\sim$ 450 µg m<sup>-3</sup> (Fig. 4 (a)). The striking layered 216 structure in the BL occurs at the height of ~300 m on 20 December 2016 (Fig. 5), which is consistent with the lower edge of anomalous inversion (Fig. 4), which verifies the strong inhibition of anomalous inversion. Additionally, a short cold air mass invades the northern urban area of Beijing in the early morning on 20 December 2016. The enhanced movement increases the BL height and reduces the  $PM_{2.5}$ mass concentration in part of northern urban area (Fig. 5), which slightly reduces the urban mean mass concentration of  $PM_{2.5}$  (Fig. 4 (1)). However, the anomalous inversion rapidly restores its original structure to facilitate pollutant accumulation.

The occurrence of anomalous inversion in 9 HPEs coincides with cumulative explosive growth of PM<sub>2.5</sub> levels (F1~4 (a, c)), which verifies the suppression of anomalous inversion to pollutants. Note that no cumulative explosive growth of PM<sub>2.5</sub> levels appears in HPE<sub>3</sub> despite anomalous inversion, partly because the height of the lowest inversion layer in HPE<sub>3</sub> (~750 m) is much higher than that in the 9 HPEs

- 227 (~ 250 meters), which suggests near-ground inversion is more favorable for pollutant accumulation.
- 228 Anomalous inversion results in near-surface moisture accumulation

229 During clean periods, the moisture is evenly distributed in the BL with RH less than 40%, while during the HPEs, RH is over 60% (even 80%) in the lower or upper BL (Fig. 1~4 (c, d)). During the 230 231 HPEs, in the absence of temperature inversion, moisture vertically distributes in the BL, and the RH in 232 the upper BL is occasionally higher than that of the near-ground surface; in the presence of weak 233 inversion, the lower edge of the inversion layer is in approximate agreement with the RH contour of 60%; 234 In the presence of anomalous inversion (red boxes in Fig. 1~4 (c)) in the BL, the lower edge of the strong 235 inversion layer frequently coincides with an RH contour of 80% (red boxes in Fig.  $1 \sim 4$  (d)), which is 236 observed in most cumulative explosive growth processes.

The previously noted relation of vertical temperature and RH indicates that anomalous inversion results in appreciable near-surface moisture accumulation by suppressing the vertical mixing of the water vapor (Wallace and Hobbs, 2006). The vertical diffusion of the near-surface water vapor as the anomalous inversion disappeared on 1 December 2016, 26 December 2016, and 5 January 2017 verifies the cited research outcome. Noted that mentioned near-ground temperature reduction caused by cooling effects of aerosols is also conducive to moisture accumulation by decreasing near-ground saturation vapor pressure to increase RH. 244 Moisture accumulation facilitates aerosol hygroscopic growth and additional secondary aerosol 245 formation

246 Strong absorbent aerosol particles absorb and grow, when additional water vapor appears in the air 247 (Zhang et al., 2015a). The mass concentrations of organic aerosols, sulfate, nitrate, and ammonium 248 rapidly increase with RH (Fig. S1). After moisture absorption in North China, aerosol particle size 249 increases 20%~60% (Pan et al., 2009) and aerosol direct radiative forcing increases ~50% (Zhang et al., 250 2015a). As a key component of atmospheric aerosols, aerosol water serves as a medium that enables 251 aqueous-phase reactions (Pilinis et al., 1989;Seinfeld and Pandis, 1986;Ervens et al., 2011). For example, 252 aerosol water serves as a reactor in which alkaline aerosol components trap SO<sub>2</sub>, which is then oxidized 253 by NO<sub>2</sub> to form sulfate in northern China (Cheng et al., 2016). The ratio of SO<sub>2</sub> to SO<sub>4</sub><sup>2-</sup> ranges from less 254 than 0.1 at relative humidity (RH) <20% to 1.1 at RH >90%, exhibiting an exponential increase with RH 255 (Wang et al., 2016). In addition, high RH facilitates heterogeneous chemical processes to aggravate air 256 pollution (Zhu et al., 2011). For example, the net reaction probability of HNO<sub>3</sub> uptake on CaCO<sub>3</sub> particles 257 was found to increase with relative humidity from ~0.003 at 10% to 0.21 at 80% (Liu et al., 2008).

Stable atmospheric stratification characteristic of southerly light or calm winds, anomalous inversion,
and near-ground (<250 m) moisture accumulation (RH>80%) dominates the cumulative explosive
growth of PM<sub>2.5</sub>.

261 During the HPEs, nearly all 10 cumulative explosive growth processes (Fig. 1-4) occur concurrently 262 with stable atmospheric stratification primarily characterized by southerly light or calm winds, near-263 ground anomalous inversion, and cumulative moisture (RH>80%). The weak southerly winds increased 264 with height are conducive to the growth, because relatively strong southerly winds in the upper BL 265 (~1000m) transport pollutants from the south of Beijing, while low-level (~250m) southerly light or calm 266 winds limiting the invasion of northerly cold winds facilitates local pollution accumulation by 267 minimizing horizontal pollutant diffusion. The anomalous inversion facilitates vertical pollutant 268 accumulation by suppressing convection activities. During the cumulative growth process in HPE $_{10}$ , the turbulent diffusion coefficient rapidly decreases from 100  $\frac{m^2 s^{-1}}{m^2 s^{-1}}$  to 50  $\frac{m^2 s^{-1}}{m^2 s^{-1}}$  and the BL height decreases 269

from 500 m to ~250 m, which is extremely favorable for pollutants accumulation. Additional suppression of vertical mixing of water by inversion and previously noted decreased saturation vapor pressure cause near-surface moisture accumulation (RH>80%). This accumulated moisture facilitates secondary aerosol formation in liquid-phase and heterogeneous reactions to increase PM<sub>2.5</sub> levels.

274 It's likely that merely weak southerly winds or near-ground anomalous inversion primarily can 275 cause cumulative explosive growth. However, the growth does not occur in the polluted process from 4 276 to 15 December 2015 with weak southerly winds, which indicates that weak southerly winds do not 277 suffice to cause cumulative explosive growth in the absence of anomalous inversion; even with 278 anomalous inversion, no explosive growth appeared on 14 and 24 January 2013, which suggests that 279 anomalous inversion cannot cause explosive growth without weak southerly winds. Therefore, 280 cumulative explosive growth in CSs is primarily resulted from the joint effects of southerly light or calm 281 winds, near-ground anomalous inversion and moisture accumulation.

Note that the cumulative explosive growth at 2000 BJT on 3 January was accompanied by a southerly gentle breeze (3.4~5.4 m s<sup>-1</sup>), which suggests that low-level southerly pollutant transport occasionally exerts an important impact on the growth, with anomalous inversion and near-ground moisture accumulation.

# 286 Feedback of cumulative pollutants on worsening meteorological conditions

287 Established from cumulative explosive growth, exceedingly high PM2.5 levels further decrease the 288 near-ground temperature by reflecting and scattering more solar radiation, which strengthens the existing 289 anomalous inversion and subsequently results in additional pollutant accumulation until the next synoptic 290 process occurs. The near-surface temperature decreased from 3°C at 2000 BJT on 19 December to -3°C 291 at 0800 BJT on 20 December after elevated ground PM2.5 levels (Fig. 4 (d)). Then it had remained at ~- $1^{\circ}\!C$  with PM\_{2.5} of more than 400  $\mu g$  m  $^{-3}$  over the next 2 d until northerly strong and clean winds blew 292 293 the pollution away on 22 December. Similar processes also occurred in the CSs of other HPEs, which 294 verifies the outcome.

### 295 **3.2.3** Local air convergence is favorable for convergent explosive growth.

296 The explosive growth of PM<sub>2.5</sub> appears in HPE<sub>4-5</sub> without inversion and near-ground moisture 297 accumulation (Fig. 2), which suggests previously noted stable atmospheric stratification does not 298 primarily cause the growth. Weak winds in convergent explosive growth processes, particularly the 299 process in HPE<sub>5</sub>, eliminate the likely contributions of southerly transport pollution. A comparison of 300 surface wind distributions in the North China Plain before (Fig. 6 (a, d)) and during (Fig. 6 (b, c; d, e)) 301 the convergent explosive growth processes in HPE<sub>4-5</sub> shows that the urban area of Beijing is dominated 302 by northerly winds before the growth, while is characterized by local air convergence during the 303 processes, which suggests that the persistent local convergence is conducive to the explosive growth by 304 causing pollutants to further locally accumulate. The convergent explosive growth in HPE<sub>6~7</sub> with air 305 convergence (Fig. S2) also verifies the outcome.

# 306 3.3 Quantification of meteorological contributions to PM2.5 cumulative explosive growth

307 Cooling effects of elevated PM<sub>2.5</sub> levels established from TSs worsen meteorological conditions, 308 which primarily causes cumulative explosive growth. To approximately quantify this atmospheric 309 feedback on the growth, PLAM (Parameter Linking Aerosol Pollution and Meteorological Elements) 310 was used, which was derived from the relationship of  $PM_{2.5}$  with key meteorological parameters. The 311 PLAM index, whose details of calculation have been described in Wang et al. (Wang et al., 2013a; Wang 312 et al., 2012), primarily reflects the stability of the air mass and the condensation rate of water vapor on 313 aerosol particles. It has been employed to identify the contribution of specific meteorological factors to 314 a 10 d haze-fog event in 2013 (Zhang et al., 2013) and to evaluate the contribution of meteorological 315 factors to changes in atmospheric composition and optical properties over Beijing during the 2008 316 Olympic Games (Zhang et al., 2009). During cumulative explosive growth processes, the hourly 317 variation of urban mean PM<sub>2.5</sub> mass concentration is in closely linear agreement with that of PLAM for 318 Beijing (Fig. 7 (a-d)). The squared correlation coefficients between hourly PLAM and PM<sub>2.5</sub> in 2013, 319 2015, and 2016 are 0.71, 0.69, and 0.71 respectively, exceeding the 0.05 significance level. The mean 320 value of four coefficients is over 0.70, which suggests the noted feedback of worsening meteorological

- 321 conditions on PM explains over 70% in cumulative explosive growth of PM<sub>2.5</sub>. In addition, the squared
- 322 correlation coefficients between PLAM and PM<sub>2.5</sub> in 2014 is 0.76, which indicates enhanced regional
- 323 atmospheric stability facilitate convergent explosive growth of PM<sub>2.5</sub>.

324 4 Conclusion:

We have characterized different stages of 12 HPEs during the WPP (wintertime pollution periods) in Beijing and typical explosive growth of PM<sub>2.5</sub>, including cumulative and convergent explosive growth. Meteorological causes to such growth are elucidated, based on observations of vertical meteorological factors within the BL (Fig. 8). Beijing HPEs can generally be divided into the TS, whose rising processes is primarily caused by pollutants transported from south of Beijing, and the CS, in which stable atmospheric stratification dominates the cumulative explosive growth of PM<sub>2.5</sub>.

331 Polluted and humid airflow from the south of Beijing transports water vapor and pollutants to 332 Beijing, which primarily causes rising processes and creates the requisite moisture and pollution 333 accumulation conditions for CSs. Elevated PM2.5 levels established from the TS reduce near-ground 334 temperature by back scattering short-wave solar radiation. This temperature reduction very likely results 335 in anomalous inversion, which is enhanced by the reduced mechanical turbulence that results from less 336 vorticity caused by observed weakened vertical shear of horizontal winds in the lower BL during the later 337 TSs and decreased thermal turbulence with cooling plumes or thermals of originally warm surface air 338 that result from the decreased near-ground temperature. Anomalous inversion reduces turbulent diffusion 339 and decreases the BL height to trap pollutants. The similar suppression of anomalous inversion to vertical 340 mixing of water vapor and decreased saturation water vapor pressure caused by noted temperature 341 reduction result in appreciable near-surface moisture accumulation (RH>80%). The accumulated 342 moisture facilitates pollutant accumulation by enhancing hygroscopic growth and accelerating liquid-343 phase and heterogeneous reactions. However, specific reaction mechanisms have not been fully 344 quantified and require additional investigation, particularly their contributions to the explosive growth 345 and the maintenance of PM<sub>2.5</sub> during CSs. Note that observed southerly weak winds facilitate local 346 pollutant accumulation by minimizing horizontal pollutant diffusion. The joint effects of southerly weak 347 winds, near-ground anomalous inversion, and moisture accumulation dominate cumulative explosive

- 348 growth of PM<sub>2.5</sub>. Nearly 70% of the growth is attributable to noted meteorological feedback, based on
- 349 correlation analysis between PM<sub>2.5</sub> and PLAM index during cumulative explosive growth processes.

350 Note that sporadic local air convergence also causes pollutants to further accumulate.

- 351 Established from cumulative explosive growth, exceedingly high PM<sub>2.5</sub> levels further decrease the
- 352 near-ground temperature to strengthen the existing anomalous inversion, which results in additional
- 353 pollutant accumulation until the next synoptic process occurs.

# 354 References

- Bai, N., Khazaei, M., van Eeden, S. F., and Laher, I.: The pharmacology of particulate matter air
   pollution-induced cardiovascular dysfunction, Pharmacology & Therapeutics, 113, 16, 2007.
- Boucher, O., Randall, D., Artaxo, P., Bretherton, C., Feingold, G., Forster, P., Kerminen, V. V. M.,
  Kondo, Y., Liao, H., Lohmann, U., Rasch, P., Satheesh, S. K., Sherwood, S., Stevens, B., and
  Zhang, X. Y.: Clouds and Aerosols, in: Climate Change 2013: The Physical Science Basis.
  Contribution of Working Group I to the Fifth Assessment Report of the Intergovernmental
  Panel on Climate Change, edited by: Stocker, T. F., Qin, D., Plattner, G.-K., Tignor, M., Allen,
  S. K., Boschung, J., Nauels, A., Xia, Y., Bex, V., and Midgley, P. M., Cambridge University
  Press, Cambridge, United Kingdom and New York, NY, USA, 571-658, 2013.
- Chen, Y., Ebenstein, A., Greenstone, M., and Li, H.: Evidence on the impact of sustained exposure
  to air pollution on life expectancy from China's Huai River policy, Proceedings of the National
  Academy of Sciences of the United States of America, 110, 12936-12941,
  10.1073/pnas.1300018110, 2013.
- Cheng, Y., Zheng, G., Chao, W., Mu, Q., Bo, Z., Wang, Z., Meng, G., Qiang, Z., He, K., and
  Carmichael, G.: Reactive nitrogen chemistry in aerosol water as a source of sulfate during haze
  events in China, Science Advances, 2, 2016.
- Ervens, B., Turpin, B. J., and Weber, R. J.: Secondary organic aerosol formation in cloud droplets
  and aqueous particles (aqSOA): a review of laboratory, field and model studies, Atmospheric
  Chemistry and Physics, 11, 11069-11102, 10.5194/acp-11-11069-2011, 2011.
- Fernald, F. G.: Analysis of atmospheric lidar observations: some comments, Applied Optics, 23, 652,
  1984.
- 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, Atmospheric Chemistry and
  Physics, 15, 4279-4295, 10.5194/acp-15-4279-2015, 2015.
- Guo, S., Hu, M., Zamora, M. L., Peng, J. F., Shang, D. J., Zheng, J., Du, Z. F., Wu, Z., Shao, M.,
  Zeng, L. M., Molina, M. J., and Zhang, R. Y.: Elucidating severe urban haze formation in China,
  Proceedings of the National Academy of Sciences of the United States of America, 111, 1737317378, 10.1073/pnas.1419604111, 2014.

- Hua, Y., Wang, S., Wang, J., Jiang, J., Zhang, T., Song, Y., Kang, L., Zhou, W., Cai, R., Wu, D., Fan,
  S., Wang, T., Tang, X., Wei, Q., Sun, F., and Xiao, Z.: Investigating the impact of regional
  transport on PM2.5 formation using vertical observation during APEC 2014 Summit in Beijing,
  Atmospheric Chemistry and Physics, 16, 15451-15460, 10.5194/acp-16-15451-2016, 2016.
- Huang, R.-J., Zhang, Y., Bozzetti, C., Ho, K.-F., Cao, J.-J., Han, Y., Daellenbach, K. R., Slowik, J.
  G., Platt, S. M., and Canonaco, F.: High secondary aerosol contribution to particulate pollution
  during haze events in China, Nature, 514, 218-222, 2014.
- Jia, Y., Rahn, K. A., He, K., Wen, T., and Wang, Y.: A novel technique for quantifying the regional
   component of urban aerosol solely from its sawtooth cycles, Journal of Geophysical Research
   Atmospheres, 113, 6089-6098, 2008.
- Liu, X. G., Li, J., Qu, Y., Han, T., Hou, L., Gu, J., Chen, C., Yang, Y., Liu, X., and Yang, T.:
  Formation and evolution mechanism of regional haze: a case study in the megacity Beijing,
  China, Atmospheric Chemistry & Physics, 13, 4501-4514, 2013.
- Liu, Y., Gibson, E. R., Cain, J. P., Wang, H., Grassian, V. H., and Laskin, A.: Kinetics of
  heterogeneous reaction of CaCO3 particles with gaseous HNO3 over a wide range of humidity,
  The Journal of Physical Chemistry A, 112, 1561-1571, 2008.
- Lv, L., Liu, W., Zhang, T., Chen, Z., Dong, Y., Fan, G., Xiang, Y., Yao, Y., Yang, N., and Chu, B.:
  Observations of particle extinction, PM 2.5 mass concentration profile and flux in north China
  based on mobile lidar technique, Atmospheric Environment, 2017.
- Pan, X. L., Yan, P., Tang, J., Ma, J. Z., Wang, Z. F., Gbaguidi, A., and Sun, Y. L.: Observational
  study of influence of aerosol hygroscopic growth on scattering coefficient over rural area near
  Beijing mega-city, Atmospheric Chemistry & Physics, 9, 7519-7530, 2009.
- Pilinis, C., Seinfeld, J. H., and Grosjean, D.: Water content of atmospheric aerosols, Atmospheric
   Environment, 23, 1601-1606, 1989.
- Seinfeld, J. H., and Pandis, S. N.: Atmospheric Chemistry and Physics: From Air Pollution to
  Climate Change, John Wiley & Sons, USA, 1986.
- Su, F., Gao, Q., Zhang, Z., REN, Z.-h., and YANG, X.-x.: Transport pathways of pollutants from
  outside in atmosphere boundary layer, Research of Environmental Sciences, 1, 26-29, 2004.
- 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, Journal of Geophysical
  Research: Atmospheres, 119, 4380-4398, 2014a.
- Sun, Y. L., Jiang, Q., Wang, Z. F., Fu, P. Q., 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.Atmos., 119, 4380-4398, 10.1002/2014jd021641, 2014b.
- 418 Wallace, J. M., and Hobbs, P. V.: Atmospheric science : an introductory survey, Elsevier, 2006.
- Wang, G., Zhang, R., Gomez, M. E., Yang, L., Levy Zamora, M., Hu, M., Lin, Y., Peng, J., Guo, S.,
  Meng, J., Li, J., Cheng, C., Hu, T., Ren, Y., Wang, Y., Gao, J., Cao, J., An, Z., Zhou, W., Li, G.,
  Wang, J., Tian, P., Marrero-Ortiz, W., Secrest, J., Du, Z., Zheng, J., Shang, D., Zeng, L., Shao,
  M., Wang, W., Huang, Y., Wang, Y., Zhu, Y., Li, Y., Hu, J., Pan, B., Cai, L., Cheng, Y., Ji, Y.,
  Zhang, F., Rosenfeld, D., Liss, P. S., Duce, R. A., Kolb, C. E., and Molina, M. J.: Persistent
  sulfate formation from London Fog to Chinese haze, Proc Natl Acad Sci U S A, 113, 13630–
- 425 13635, 10.1073/pnas.1616540113, 2016.

- Wang, H., Zhang, X., Gong, S., Chen, Y., Shi, G., and Li, W.: Radiative feedback of dust aerosols
  on the East Asian dust storms, Journal of Geophysical Research, 115, 6696-6705, 2010.
- Wang, H., Shi, G. Y., Zhang, X. Y., Gong, S. L., Tan, S. C., Chen, B., Che, H. Z., and Li, T.:
  Mesoscale modeling study of the interactions between aerosols and PBL meteorology during
  a haze episode in China Jing-Jin-Ji and its near surrounding region Part 2: Aerosols' radiative
  feedback effects, Atmospheric Chemistry & Physics, 15, 3277-3287, 2015a.
- Wang, H., Xue, M., Zhang, X. Y., Liu, H. L., Zhou, C. H., Tan, S. C., Che, H. Z., Chen, B., and Li,
  T.: Mesoscale modeling study of the interactions between aerosols and PBL meteorology
  during a haze episode in Jing–Jin–Ji (China) and its nearby surrounding region Part 1:
  Aerosol distributions and meteorological features, Atmospheric Chemistry and Physics, 15,
  3257-3275, 10.5194/acp-15-3257-2015, 2015b.
- Wang, J., Wang, Y., Liu, H., Yang, Y., Zhang, X., Li, Y., Zhang, Y., and Deng, G.: Diagnostic
  identification of the impact of meteorological conditions on PM2.5 concentrations in Beijing,
  Atmospheric Environment, 81, 158-165, 10.1016/j.atmosenv.2013.08.033, 2013a.
- Wang, J., Wang, S., Jiang, J., Ding, A., Zheng, M., Zhao, B., Wong, D. C., Zhou, W., Zheng, G.,
  Wang, L., Pleim, J. E., and Hao, J.: Impact of aerosol-meteorology interactions on fine particle
  pollution during China's severe haze episode in January 2013, Environmental Research Letters,
  9, 094002, 10.1088/1748-9326/9/9/094002, 2014a.
- Wang, J. Z., Gong, S., Zhang, X. Y., Yang, Y. Q., Hou, Q., Zhou, C., and Wang, Y.: A parameterized
  method for air-quality diagnosis and its applications, Advance of Meteorology,
  doi:10.1155/2012/238589, 1-10, 2012.
- Wang, X., Chen, J., Sun, J., Li, W., Yang, L., Wen, L., Wang, W., Wang, X., Collett, J. L., and Shi,
  Y.: Severe haze episodes and seriously polluted fog water in Ji'nan, China, Sci. Total Environ.,
  449 493, 133-137, 2014b.
- Wang, Z., Li, J., Wang, Z., Yang, W., Tang, X., Ge, B., Yan, P., Zhu, L., Chen, X., and Chen, H.:
  Modeling study of regional severe hazes over mid-eastern China in January 2013 and its
  implications on pollution prevention and control, Science China Earth Sciences, 57, 3-13,
  2014c.
- Wang, Z. B., Hu, M., Wu, Z. J., Yue, D. L., He, L. Y., Huang, X. F., Liu, X. G., and Wiedensohler,
  A.: Long-term measurements of particle number size distributions and the relationships with
  air mass history and source apportionment in the summer of Beijing, Atmospheric Chemistry
  and Physics, 13, 10159-10170, 10.5194/acp-13-10159-2013, 2013b.
- Wei, P., Cheng, S. Y., Li, J. B., and Su, F. Q.: Impact of boundary-layer anticyclonic weather system
  on regional air quality, Atmospheric Environment, 45, 2453-2463,
  10.1016/j.atmosenv.2011.01.045, 2011.
- Zhang, L., Sun, J. Y., Shen, X. J., Zhang, Y. M., Che, H., Ma, Q. L., Zhang, Y. W., Zhang, X. Y., and
  Ogren, J. A.: Observations of relative humidity effects on aerosol light scattering in the Yangtze
  River Delta of China, Atmospheric Chemistry & Physics, 15, 2853-2904, 2015a.
- Zhang, R. H., Li, Q., and Zhang, R. N.: Meteorological conditions for the persistent severe fog and
   haze event over eastern China in January 2013, Science China Earth Sciences, 57, 26-35, 2014.
- Zhang, R. Y., Wang, G. H., Guo, S., Zarnora, M. L., Ying, Q., Lin, Y., Wang, W. G., Hu, M., and
  Wang, Y.: Formation of Urban Fine Particulate Matter, Chemical Reviews, 115, 3803-3855,

- 468 10.1021/acs.chemrev.5b00067, 2015b.
- Zhang, X., Sun, J., Wang, Y., Li, W., Zhang, Q., Wang, W., Quan, J., Cao, G., Wang, J., Yang, Y.,
  and Zhang, Y.: Factors contributing to haze and fog in China, Chinese Science Bulletin
  (Chinese Version), 58, 1178, 10.1360/972013-150, 2013.
- Zhang, X. Y., Wang, Y. Q., Lin, W. L., Zhang, Y. M., Zhang, X. C., Gong, S., Zhao, P., Yang, Y. Q.,
  Wang, J. Z., and Hou, Q.: Changes of Atmospheric Composition and Optical Properties Over
  BEIJING—2008 Olympic Monitoring Campaign, Bulletin of the American Meteorological
  Society, 90, 1633–1651, 2009.
- 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., 11,
  26571-26615, 2012.
- Zhang, X. Y., Wang, J. Z., Wang, Y. Q., Liu, H. L., Sun, J. Y., and Zhang, Y. M.: Changes in chemical
  components of aerosol particles in different haze regions in China from 2006 to 2013 and
  contribution of meteorological factors, Atmos. Chem. Phys., 15, 12935-12952, 10.5194/acp15-12935-2015, 2015c.
- Zheng, B., Zhang, Q., Zhang, Y., He, K. B., Wang, K., Zheng, G. J., Duan, F. K., Ma, Y. L., 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,
  Atmospheric Chemistry and Physics, 15, 2031-2049, 10.5194/acp-15-2031-2015, 2015a.
- Zheng, G. J., Duan, F. K., Su, H., Ma, Y. L., Cheng, Y., Zheng, B., Zhang, Q., Huang, T., Kimoto,
  T., Chang, D., Poschl, U., Cheng, Y. F., and He, K. B.: Exploring the severe winter haze in
  Beijing: the impact of synoptic weather, regional transport and heterogeneous reactions,
  Atmospheric Chemistry and Physics, 15, 2969-2983, 10.5194/acp-15-2969-2015, 2015b.
- Zhu, T., Shang, J., and Zhao, D. F.: The roles of heterogeneous chemical processes in the formation
  of an air pollution complex and gray haze, Science China Chemistry, 54, 145-153, 2011.

495	Data Availability
496	The data that support the findings of this study are available from the corresponding author upon
497	reasonable request.
498	
499	Acknowledgements:
500	This research is supported by the National Key Project of MOST (2016YFC0203306), the Atmospheric
501	Pollution Control - the Prime Minister Fund, and the Basic Scientific Research Progress of the Chinese
502	Academy of Meteorological Sciences (2016Z001).
503	
504	Author Contributions:
505	X.Y.Z. and Y.Q.W. designed the research; X.Y.Z, J.T.Z and H.C.C carried out the analysis of observations.
506	Y.S.D provided and analyzed laser radar data. Y.M.Z provided aerosol species data. J.Z.W provided
507	PLAM data. J.T.Z. wrote the first manuscript and X.Y.Z. revised the manuscript. All authors read and
508	approved the final version.
509	
510	Additional Information:
511	Competing financial interests: The authors declare no competing financial interests.
512	

513 Figures

514



516 Figure 1. Temporal variations in urban mean PM<sub>2.5</sub> and vertical distributions of meteorological factors in

517 January 2013. (a)PM<sub>2.5</sub> mass concentration (dark gray or gray: Beijing; light gray: Baoding); (b) winds (vectors;

518 red vectors: southwesterly winds) and wind velocity (shadings; units: m/s); (c)temperature (shadings; units: °C);

- 519 (d)RH (shadings; units: %); (green boxes: rising processes; red boxes: cumulative explosive processes)
- 520



Figure 2. Temporal variations in urban mean PM<sub>2.5</sub> and vertical distributions of meteorological factors in
February 2014. (a)PM<sub>2.5</sub> mass concentration (dark gray or gray: Beijing; light gray: Baoding); (b) winds (vectors;
red vectors: southwesterly winds) and wind velocity (shadings; units: m/s); (c)temperature (shadings; units: °C);
(d)RH (shadings; units: %); (green boxes: rising processes)



Figure 3. Temporal variations in urban mean PM<sub>2.5</sub> and vertical distributions of meteorological factors in
December 2015. (a)PM<sub>2.5</sub> mass concentration (dark gray or gray: Beijing; light gray: Baoding); (b) winds (vectors;
red vectors: southwesterly winds) and wind velocity (shadings; units: m/s); (c)temperature (shadings; units: °C);
(d)RH (shadings; units: %); (green boxes: rising processes; red boxes: cumulative explosive processes)



538 December 2016. (a)PM<sub>2.5</sub> mass concentration (dark gray or gray: Beijing; light gray: Baoding); (b) winds (vectors;

- red vectors: southwesterly winds) and wind velocity (shadings; units: m/s); (c)temperature (shadings; units: °C);
- 540 (d)RH (shadings; units: %); (green boxes: rising processes; red boxes: cumulative explosive processes)
- 541



- urban area of Beijing from 19 to 20 December 2016 (The dashed lines: the approximate boundary layer height)



547 Figure 6. Surface wind distributions before (a, d) and during (b, c; d, e) two convergent explosive growth

548 processes in February 2014 on the North China Plain.



Figure 7. Correlation between PLAM and PM2.5 during the cumulative explosive growth processes in
January 2013 (a) , December 2015 (c), December 2016 (d) respectively, and the convergent explosive growth
processes in February 2014 (b).



- 557 Figure 8. A schematic figure of the formation mechanism for winter heavy pollution episodes in Beijing, which
- 558 consist of the transport stage (green background) and the cumulative stage (red background).