

1 **Mixing layer transport flux of particulate matter in Beijing, China**

2 Yusi Liu¹, Guiqian Tang^{2,3}, Libo Zhou², Bo Hu², Baoxian Liu^{4,5}, Yunting Li^{4,5}, Shu Liu⁶, Yuesi
3 Wang^{2,3,7}

4 ¹ State Key Laboratory of Severe Weather & Key Laboratory for Atmospheric Chemistry of China
5 Meteorology Administration, Chinese Academy of Meteorological Sciences, Beijing 100081, China

6 ² State Key Laboratory of Atmospheric Boundary Layer Physics and Atmospheric Chemistry,
7 Institute of Atmospheric Physics, Chinese Academy of Sciences, Beijing 100029, China

8 ³ Center for Excellence in Urban Atmospheric Environment, Institute of Urban Environment,
9 Chinese Academy of Sciences, Xiamen 361021, China

10 ⁴ Beijing Municipal Environmental Monitoring Centre, Beijing 100048, China

11 ⁵ Beijing Key Laboratory of Airborne Particulate Matter Monitoring Technology, Beijing 100048,
12 China

13 ⁶ Liaoning Provincial Environmental Monitoring & Experiment Center, Shenyang 110031, China

14 ⁷ University of Chinese Academy of Sciences, Beijing 100049, China

15

16 Correspondence: Guiqian Tang (tgq@dq.cern.ac.cn)

17

18 **Abstract**

19 Quantifying the transport flux (TF) of atmospheric pollutants plays an important role in
20 understanding the causes of air pollution and in making decisions regarding the prevention and
21 control of regional air pollution. In this study, the mixing layer height and wind profile were
22 measured by a ceilometer and Doppler wind radar, respectively, and the characteristics of the
23 atmospheric dilution capability were analyzed using these two datasets. The ventilation coefficient
24 (VC) appears to be highest in the spring ($3940 \pm 2110 \text{ m}^2 \text{ s}^{-1}$) and lower in the summer (2953 ± 1322
25 $\text{m}^2 \text{ s}^{-1}$), autumn ($2580 \pm 1601 \text{ m}^2 \text{ s}^{-1}$) and winter ($2913 \pm 3323 \text{ m}^2 \text{ s}^{-1}$). Combined with the
26 backscatters measured by the ceilometer, vertical profiles of the $\text{PM}_{2.5}$ concentration were obtained
27 and the $\text{PM}_{2.5}$ TF in the mixing layer was calculated. The TF was the highest in the spring at $4.33 \pm$
28 $0.69 \text{ mg m}^{-1}\text{s}^{-1}$ and lower in the summer, autumn and winter, when the TF values were 2.27 ± 0.42
29 $\text{mg m}^{-1}\text{s}^{-1}$, $2.39 \pm 0.45 \text{ mg m}^{-1}\text{s}^{-1}$ and $2.89 \pm 0.49 \text{ mg m}^{-1}\text{s}^{-1}$, respectively. Air pollutants transport
30 mainly occurs between 14:00 and 18:00 LT. The TF was large in the pollution transition period
31 (spring: $5.50 \pm 4.83 \text{ mg m}^{-1}\text{s}^{-1}$, summer: $3.94 \pm 2.36 \text{ mg m}^{-1}\text{s}^{-1}$, autumn: $3.72 \pm 2.86 \text{ mg m}^{-1}\text{s}^{-1}$ and
32 winter: $4.45 \pm 4.40 \text{ mg m}^{-1}\text{s}^{-1}$) and decreased during the heavy pollution period (spring: 4.69 ± 4.84
33 $\text{mg m}^{-1}\text{s}^{-1}$, summer: $3.39 \pm 1.77 \text{ mg m}^{-1}\text{s}^{-1}$, autumn: $3.01 \pm 2.40 \text{ mg m}^{-1}\text{s}^{-1}$ and winter: 3.25 ± 2.77
34 $\text{mg m}^{-1}\text{s}^{-1}$). Our results indicate that the influence of the air pollutants transport in the southern
35 regions should receive more focus in the transition period of pollution, while local emissions should
36 receive more focus in the heavy pollution period.

37 **1. Introduction**

38 With the rapid development of its economy and industry, as well as its unique local topography,
39 Beijing has become one of the cities in the world that is most seriously affected by air pollution. As
40 early as before the 2008 Olympic Games, to fulfill the promise of a “Green Olympics”, Beijing’s
41 industries were relocated to surrounding provinces and cities. After the Olympic Games, with the
42 promulgation of the “Action Plan for Prevention and Control of Air Pollution”, Beijing
43 implemented a series of measures to reduce pollutants, such as raising the emission standards of
44 motor vehicles and fuel standards for vehicles, changing coal to natural gas, coal to electricity and
45 so on. These measures have gradually improved Beijing’s air quality, with the annual average fine
46 particulate matter ($\text{PM}_{2.5}$) concentration decreasing from $90 \mu\text{g m}^{-3}$ in 2013 to $58 \mu\text{g m}^{-3}$ in 2017
47 (Cheng et al. 2018a).

48 Although the Beijing government has been committed in recent years to taking measures that could
49 ensure a steady improvement in the air quality, there is still great pressure to achieve a continuous
50 decline in the particulate matter concentration. Beijing is in the north of the North China Plain, with
51 the south side and the west side the Yanshan Mountains and the Taihang Mountains, respectively.
52 Affected by the mountains to the northwest, there are more subsiding airflows, a lower mixing layer
53 height and an extremely limited atmospheric dilution capability. In addition, pollutants tend to
54 accumulate in front of the mountains due to the influence of southerly winds and the mountain
55 obstructions. In central and northern China, the increase in $\text{PM}_{2.5}$ during the winter is closely related
56 to the adverse atmospheric dilution conditions (Wang et al. 2016). Therefore, in addition to the
57 primary emissions and secondary formation, the weak atmospheric dilution capability is also an
58 important factor leading to the frequent occurrence of serious air pollution in Beijing.

59 In recent decades, the mixing layer height (MLH) and wind speed (WS) have been two major factors

60 leading to the annual increases in the aerosol concentration and polluted days during the winter in
61 China (Yang et al. 2016). The low MLH and low WS are also important characteristics of the weak
62 atmospheric dilution capability (Huang et al. 2018; Liu et al. 2018; Song et al. 2014; Tang et al.
63 2015). The change in the MLH represents the vertical dilution capability of pollutants, and the
64 change in the WS represents the horizontal dilution capability of pollutants. The ventilation
65 coefficient (VC) is usually used to evaluate the vertical and horizontal dilution capability of the
66 atmosphere (Nair et al. 2007; Tang et al. 2015; Zhu et al. 2018). Thus, it is a good choice to use the
67 VC to evaluate the relationship between the atmospheric dilution capability and air pollution in
68 Beijing. Although previous studies have analyzed the relationship between the MLH and pollutants
69 (Geiß et al. 2017; Miao and Liu 2019; Schäfer et al. 2006; Su et al. 2018), studies on the effects of
70 the VC on the particle concentration have been extremely rare.

71 Although the problem of heavy pollution in northern China has improved in recent years, regional
72 pollution problems remain, especially in the Beijing-Tianjin-Hebei region (Shen et al. 2019). There
73 are three main transport routes affecting Beijing: the northwest path, the southwest path and the
74 southeast path (Chang et al. 2018; Li et al. 2018; Zhang et al. 2018). The occurrence of heavy
75 pollution in Beijing is closely related to the transport of pollutants in the southern regions, mainly
76 in southern Hebei, northern Henan and western Shandong, while the high-speed northwest air mass
77 is conducive to the removal of pollutants from Beijing (Li et al. 2018; Ouyang et al. 2019; Zhang et
78 al. 2018; Zhang et al. 2017). In recent years, the contribution of regional transport to Beijing has
79 been increasing annually, with a trend of 1.2% per year, which reached 31-73% in the summer and
80 27-59% in the winter (Chang et al. 2018; Cheng et al. 2018b; Wang et al. 2015). High PM_{2.5}
81 concentrations are usually accompanied by high transport flux (TF) within a day in Beijing. As
82 pollution worsens, the contribution of the surrounding areas to the PM_{2.5} in Beijing has risen from
83 52% to 65% in a month on average in 2016 (Zhang et al. 2018). However, during heavy pollution,
84 the TF decreases in Beijing (Chang et al. 2018; Tang et al. 2015; Zhu et al. 2016).

85 To solve the regional pollution problem, joint prevention and control have been recommended for a
86 long time. Many studies on regional transport have been carried out, but most observational studies
87 cannot easily quantify the TF due to the lack of particle and wind vertical profiles, and it is still
88 unclear when we need to control the emission sources and in which areas. To solve the above
89 problems, we conducted 2 years of continuous observations on MLH and wind profiles in the
90 Beijing mixing layer and analyzed the mixing layer dilution capability of the atmosphere.
91 Afterwards, using the backscattering coefficient profile, we obtained the vertical PM_{2.5}
92 concentration profiles and calculated the TF profile and mixing layer TF. Finally, using the near-
93 surface PM_{2.5} concentration as an indicator to classify the air pollution degree, we analyzed the TF
94 during the transitional and heavily polluted periods in Beijing and illuminated the main controlling
95 factors.

96 **2. Methods**

97 **2.1 Observational station**

98 To understand the dilution capability characteristics in Beijing, two years of observations were
99 conducted (2016.1.1-2017.12.31). The observational site (BJT) is at the Institute of Atmospheric
100 Physics of the Chinese Academy of Sciences, located west of the Jiande Bridge in the Haidian

101 District, Beijing (39.98° N, 116.38° W). The north and south sides of the station are the north Third
102 and north Fourth Ring Roads, respectively, and the eastern side is the Beijing-Tibet Expressway.
103 The altitude (a.s.l.) is approximately 60 m. There is no obvious emission source around the
104 observational site except for motor vehicles.

105 **2.2 Observations of MLH and wind profiles**

106 To analyze the dilution capability, the MLH was observed by a single-lens ceilometer (CL51, Vaisala,
107 Finland), and the wind profile was simultaneously observed by a Doppler wind radar (Windcube
108 100s, Leosphere, France).

109 A single-lens ceilometer measures the attenuated backscatter coefficient profile of the atmosphere
110 by pulsed diode laser lidar technology (910 nm waveband) within a 7.7 km range and determines
111 the MLH through the positions of abrupt changes in the backscattering coefficient profile. In the
112 actual measurement, the measurement interval was 16 s, and the measurement resolution was 10 m.
113 More detailed descriptions are presented in the published literature (Tang et al. 2016; Zhu et al.
114 2016). In this study, the gradient method (Steyn et al. 1999) is used to determine the MLH; that is,
115 the top of the mixing layer was determined by the maximum negative gradient value in the profile
116 of the atmosphere backscattering coefficient. Moreover, to eliminate the interference of the aerosol
117 layer structure and the detection noise, the MLH was calculated by the improved gradient method
118 after smoothly averaging the profile data (Münkel et al. 2007; Tang et al. 2015).

119 Doppler wind radar uses the remote sensing method of laser detection and ranging technology and
120 measures the Doppler frequency shift generated by the laser through the backscatter echo signal of
121 particles in the air. The Windcube 100s can provide 3D wind field data within a 3 km range from
122 the system, including u, v and w vectors. In the actual measurement, starting from 100 m, the spatial
123 resolution is 50 m, the WS accuracy is $< 0.5 \text{ m s}^{-1}$, and the radial WS range is -30 m s^{-1} to 30 m s^{-1} .

124 **2.3 Other data**

125 During the observations, the hourly $\text{PM}_{2.5}$ concentrations of the Beijing Olympic Sports Center
126 (39.99° N, 116.40° W) were obtained from the Ministry of Environmental Protection of China
127 (<http://www.zhb.gov.cn/>).

128 **2.4 Analytical method**

129 The atmospheric dilution is composed of vertical and horizontal dilution, which can be characterized
130 by the MLH and wind speed in the mixing layer (WS_{ML}), respectively. The VC ($\text{m}^2 \text{ s}^{-1}$) was obtained
131 by combining the MLH (m) and WS_{ML} (m s^{-1}) and can be used for a comprehensive evaluation of
132 the vertical and horizontal dilutions. Higher dilution-related parameters (MLH, WS_{ML} and VC)
133 indicate a stronger dilution capability, which is conducive to the transport and dilution of heavy air
134 pollution.

135 The VC calculation method is as follows:

$$136 \quad VC = H_{\text{ML}} \times \text{WS}_{\text{ML}}, \quad (1)$$

$$137 \quad \text{WS}_{\text{ML}} = \frac{1}{n} \sum_{i=1}^n \text{WS}_i, \quad (2)$$

138 $WS_i = \sqrt{u_i^2 + v_i^2}$, (3)

139 where WS_{ML} is the average WS within the mixing layer, calculated by Eq. (2); H_{ML} is the height of
 140 the mixing layer; WS_i is the WS observed at a certain height, calculated by u_i and v_i in the wind
 141 profile according to Eq. (3); and n is the number of measurement layers in the mixing layer (Nair et
 142 al. 2007).

143 The TF ($\text{mg m}^{-2}\text{s}^{-1}$) is determined by the WS and the $PM_{2.5}$ concentration in the area under analysis.
 144 The calculation method for a certain height is shown in Eq. (4):

145 $TF_{u_i} = u_i \times C_i$, (4)

146 where C_i is the concentration of $PM_{2.5}$ at a certain height. However, it is extremely difficult to
 147 observe the vertical $PM_{2.5}$ concentration in the mixing layer. To obtain the $PM_{2.5}$ concentration
 148 profile, we studied the backscattering coefficient measured by ceilometer, and found that the
 149 concentration of near-surface $PM_{2.5}$ is strongly correlated with the backscattering coefficient at 100
 150 m (Fig. S1). Thus, based on the relationship between the two, the backscattering coefficient profile
 151 can be used to invert the vertical $PM_{2.5}$ concentration profile. Then, the TFs in the mixing layer are
 152 calculated as follows:

153 $TF_u = \int_{i=1}^n (u_i \times C_i)$
 154 $TF_v = \int_{i=1}^n (v_i \times C_i)$ (5)

155 Through the above method, radial and zonal TFs can be obtained, and vector synthesis in two
 156 directions can be conducted to obtain the main transport direction to find the transport source area.

157 3. Results and discussion

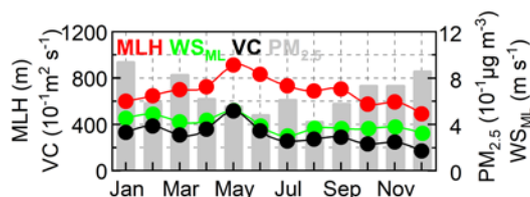
158 3.1 Boundary layer meteorology

159 3.1.1 Seasonal variation

160 To understand the variations of the atmospheric dilution capability, we carried out continuous
 161 measurements of the MLH and wind profile within the mixing layer over a 2-year period (2016.1.1-
 162 2017.12.31). The availability was verified after MLH elimination by Tang et al. (Tang et al. 2016).
 163 After the exclusion of the data of the MLH under rainy, sandstorm and windy conditions, the data
 164 availability was 95% over the 2-year period, higher than that of previous studies (Mues et al. 2017;
 165 Tang et al. 2016). The availability was the lowest in February, at 86%, and the highest in July, at
 166 99%.

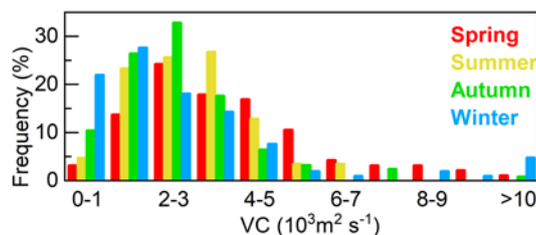
167 In terms of the seasonal variation, the average MLHs for the spring (781 ± 229 m) (value \pm standard
 168 deviation) and summer (767 ± 219 m) were higher than those of the autumn (612 ± 166 m) and
 169 winter (584 ± 221 m) (Fig. 1). However, WS_{ML} was different from the MLH in terms of the seasonal
 170 variation, with the largest value 4.6 ± 1.6 m s^{-1} in the spring, followed by the winter (4.1 ± 2.7 m s^{-1})
 171 and autumn (3.7 ± 1.6 m s^{-1}), and the smallest value 3.6 ± 1.1 m s^{-1} in the summer. The VC was
 172 calculated by the MLH and wind profile, and the results demonstrate that the dilution capability was
 173 strongest in the spring, as the VC reached as high as 3940 ± 2110 $\text{m}^2 \text{s}^{-1}$. The atmospheric dilution
 174 capabilities for the summer, winter and autumn were similar, with VC values of 2953 ± 1322 $\text{m}^2 \text{s}^{-1}$.

175 ¹, $2913 \pm 3323 \text{ m}^2 \text{ s}^{-1}$ and $2580 \pm 1601 \text{ m}^2 \text{ s}^{-1}$, respectively. A monthly analysis shows that the
 176 atmospheric dilution capability was strongest in May, when the VC was as high as $5161 \pm 2085 \text{ m}^2$
 177 s^{-1} , and worst in December, when the VC was only $1690 \pm 1072 \text{ m}^2 \text{ s}^{-1}$. The VC value in May was
 178 3.1 times that in December. To analyze the impact of the dilution capacity on $\text{PM}_{2.5}$, the seasonal
 179 variation of $\text{PM}_{2.5}$ was analyzed. The average $\text{PM}_{2.5}$ concentration for the winter ($80 \pm 87 \mu\text{g m}^{-3}$)
 180 was the highest, followed by autumn ($68 \pm 54 \mu\text{g m}^{-3}$) and spring ($67 \pm 60 \mu\text{g m}^{-3}$), and that of the
 181 summer ($51 \pm 29 \mu\text{g m}^{-3}$) was the lowest. The lowest monthly average $\text{PM}_{2.5}$ concentration was 42
 182 $\pm 26 \mu\text{g m}^{-3}$ in August. The highest monthly average was in January at $94 \pm 100 \mu\text{g m}^{-3}$, 2.2 times
 183 that in August (Fig. 1).



184
 185 Fig. 1 Monthly variations in mixing layer height (MLH), the wind speed in the mixing layer
 186 (WS_{ML}), the ventilation coefficient (VC) and $\text{PM}_{2.5}$ in Beijing.

187 Although there is little difference in the dilution capability between the summer, autumn and winter,
 188 there is serious pollution in the autumn and winter. To analyze this problem, the VC frequency
 189 distribution was studied. The results show that the VC had a high frequency in the range of 1000-
 190 $4000 \text{ m}^2 \text{ s}^{-1}$ from 2016 to 2017, but the frequency distribution was different in different seasons (Fig.
 191 2). The VC showed a strong dilution capability in the spring, mainly in the range of 2000-5000 m^2
 192 s^{-1} , with the highest frequency (24%) in the range of 2000-3000 $\text{m}^2 \text{ s}^{-1}$. In the summer, the high
 193 frequency of the VC occurred in the range of 1000-4000 $\text{m}^2 \text{ s}^{-1}$, which was slightly lower than that
 194 in the spring, and the highest frequency (27%) occurred in the range of 3000-4000 $\text{m}^2 \text{ s}^{-1}$.
 195 Additionally, the VC high frequency appeared in lower ranges in the autumn and winter. The VC
 196 occurred at a high frequency of 1000-3000 $\text{m}^2 \text{ s}^{-1}$ in the autumn, and the highest frequency occurred
 197 within the range of 2000-3000 $\text{m}^2 \text{ s}^{-1}$, accounting for 33%. In the winter, the VC appeared more
 198 frequently in the range of 0-2000 $\text{m}^2 \text{ s}^{-1}$ and was the highest in the range of 1000-2000 $\text{m}^2 \text{ s}^{-1}$, which
 199 was 28%. In the winter, when the Siberian High transits, strong northwest winds prevail in the
 200 Beijing area (Fig. 5), resulting in the higher frequency of the VC in the range of 1000-2000 $\text{m}^2 \text{ s}^{-1}$.
 201 The VC frequency of 0-1000 $\text{m}^2 \text{ s}^{-1}$ in the winter was significantly higher than that of the other
 202 seasons, up to 22%, which was 7 times that in the spring, 5 times that in the summer and 2 times
 203 that in the autumn. According to the seasonal variation in the $\text{PM}_{2.5}$ concentration, heavy pollution
 204 in the autumn and winter is related to the high frequency of poor atmospheric dilution capability.

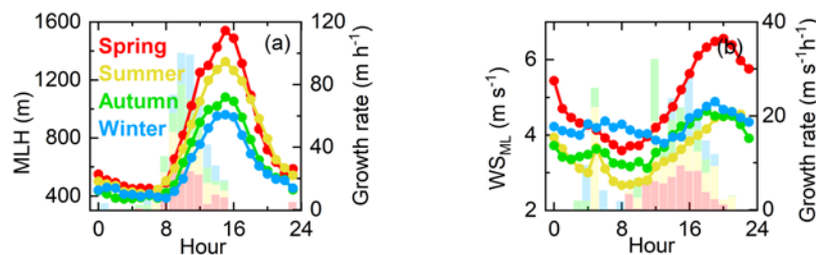


205
 206 Fig. 2 Frequency distribution of the daily VC from January 2016 to December 2017 in Beijing.

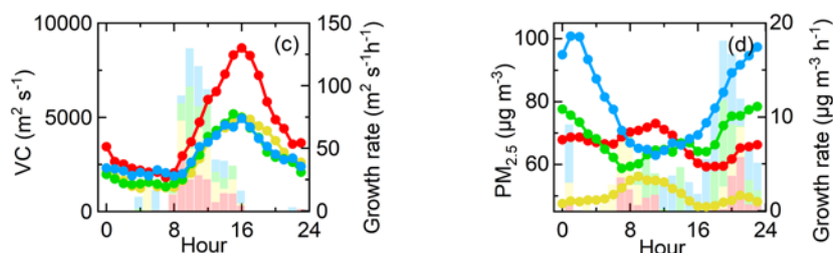
207 **3.1.2 Diurnal variation**

208 Moreover, the diurnal variations in the dilution-related parameters during different seasons were
 209 analyzed to reveal the diurnal evolution of the atmospheric dilution capability. The peak and trough
 210 values of the MLH and VC appeared simultaneously at approximately 15:30 LT and 05:30 LT,
 211 respectively. Generally, the daily variation in the MLH is characterized by a low value at night,
 212 which increases rapidly after sunrise and reaches the maximum value in the afternoon (Fig. 3a). The
 213 daily maximum value of the MLH is seasonal, where it is higher in the spring and summer and lower
 214 in the autumn and winter. The daily minimum value of the MLH generally occurs when the mixing
 215 layer is stable and is closely related to the WS. The diurnal variation in WS_{ML} is smaller, with a peak
 216 at approximately 19:30 LT and a trough at approximately 10:00 LT, which are ~4 h later than the
 217 peak and trough of the MLH (Fig. 3b). The diurnal variation in the VC is similar to that of the MLH,
 218 showing that the dilution capability is strong before sunset, gradually weakens after sunset and
 219 remains stable at night. The dilution capability in the spring was significantly stronger than that
 220 during the other seasons, and the maximum daily value reached $8678 \text{ m}^2 \text{ s}^{-1}$ (Fig. 3c). The daily
 221 maximum values of the VC in the summer, autumn and winter were close, at approximately 5000
 222 $\text{m}^2 \text{ s}^{-1}$ (Fig. 3c). The VC growth rate in the spring was significantly higher than that in the other
 223 seasons, reaching a maximum at approximately 09:00 LT. In the autumn, the VC growth rate peaked
 224 at approximately 10:00 LT, and those in the summer and winter peaked at approximately 11:00 LT.
 225 Throughout the year, the VC began to increase during the winter later than in other seasons, at
 226 approximately 09:00 LT, indicating that the weaker dilution capability remained for a longer period
 227 during the winter. The VC was weakened most rapidly in the spring; however, it was still higher
 228 than that of the other seasons after declining. In addition to the spring, the VC in the autumn and
 229 winter weakened the most rapidly, and the most slowly in the summer. In general, the vertical and
 230 horizontal dilutions are strong in the spring during both the day and night. In the winter, the vertical
 231 dilution is weak during the day, and the horizontal dilution during the night is the main component.
 232 In the summer, the vertical dilution during the day is dominant.

233 Notable differences are present when we compare the dilution-related parameters to the $PM_{2.5}$
 234 concentration. The daily maximum $PM_{2.5}$ concentrations in the spring, summer, autumn and winter
 235 were $73 \mu\text{g m}^{-3}$ (11:00 LT), $56 \mu\text{g m}^{-3}$ (09:00 LT), $78 \mu\text{g m}^{-3}$ (23:00 LT) and $101 \mu\text{g m}^{-3}$ (01:00 LT),
 236 respectively. The differences between the maximum and minimum were $14 \mu\text{g m}^{-3}$, $10 \mu\text{g m}^{-3}$, 20
 237 $\mu\text{g m}^{-3}$ and $38 \mu\text{g m}^{-3}$, respectively. Thus, the diurnal variation of $PM_{2.5}$ can be divided into two
 238 categories: (1) the highest value occurs in the midday in the spring and summer and the overall
 239 change is small, and (2) the highest value occurs during the night in the autumn and winter and
 240 differs greatly from the lowest value (Fig. 3d). The main causes of air pollution are local emissions
 241 and regional transportation. Thus, these results indicate that there are greater local contributions in
 242 the autumn and winter and higher regional transport in the spring and summer.



243



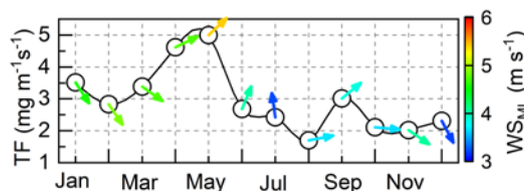
244

245 Fig. 3 Diurnal variations and growth rates of MLH (a), WS_{ML} (b), VC (c) and $PM_{2.5}$ (d) in the
 246 spring, summer, autumn and winter in Beijing. Diurnal variations are represented by lines and
 247 scatters. Growth rates are represented by columns, and only positive values are shown in the
 248 figure.

249 3.2 Mixing layer TF of $PM_{2.5}$

250 3.2.1 Temporal evolution of TF

251 To quantify the transport of $PM_{2.5}$ in Beijing, the transport direction of $PM_{2.5}$ was characterized by
 252 the average wind direction in the mixing layer. As shown in Fig. 4, the mixing layer TF in the spring
 253 was the largest, reaching $4.33 \pm 0.69 \text{ mg m}^{-1}\text{s}^{-1}$, and there was no significant difference in the
 254 summer, autumn or winter, when the TF values were $2.27 \pm 0.42 \text{ mg m}^{-1}\text{s}^{-1}$, $2.39 \pm 0.45 \text{ mg m}^{-1}\text{s}^{-1}$
 255 and $2.89 \pm 0.49 \text{ mg m}^{-1}\text{s}^{-1}$, respectively. The transport sources of the cold period in Beijing were
 256 predominantly from the northwesterly and westerly directions. With temperature warming, the
 257 transport direction gradually changed from west to south, mainly southwesterly in the spring and
 258 southerly in the summer. The monthly average maximum value of the TF occurred in May, as high
 259 as $5.00 \pm 5.21 \text{ mg m}^{-1}\text{s}^{-1}$ and mainly originated from the southwest direction, accompanied by a
 260 strong wind. The minimum value appeared in August, as low as $1.70 \pm 1.73 \text{ mg m}^{-1}\text{s}^{-1}$, which was
 261 mainly transported from western regions, with a small WS. The TF in May was 3 times that in
 262 August (Fig. 4). Therefore, the change in the transport direction leads to an obvious seasonal
 263 variation in the TF. Overall, the regional transport contributes the most to the $PM_{2.5}$ concentration
 264 in the spring, which is mainly related to increased dust activities; regional transport has a smaller
 265 contribution in the winter, but there is a high near-surface $PM_{2.5}$ concentration, which indicates that
 266 more focus should be given to local emission source control; in the summer and autumn, the
 267 southwest airflow transport influence on the Beijing should receive more focus.

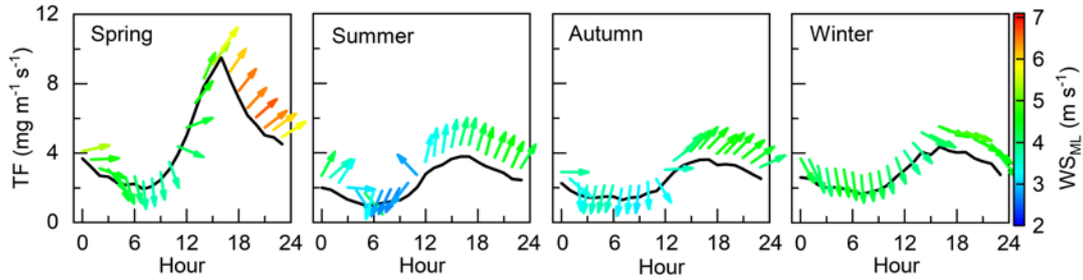


268

269 Fig. 4 Seasonal variations in the mixing layer TF of $PM_{2.5}$ and the transport direction.

270 To understand the regional transport influence on the Beijing area, the diurnal variations of the
 271 mixing layer TF were analyzed during different seasons in Beijing. The daily minimum value of the
 272 TF appeared at approximately 07:00 LT and was accompanied by a northerly wind. As the average

273 wind direction in the mixing layer gradually turned south, the daily minimum value of the TF
274 continued to rise until the daily maximum value appeared at approximately 16:00 LT (Fig. 5).
275 Transport mainly occurred between 14:00 and 18:00 LT, which was consistent with the results of a
276 previous study (Ge et al. 2018). In the spring, the WS was the highest, so the peak TF duration was
277 the shortest; it peaked at only 16:00 LT ($9.50 \text{ mg m}^{-1}\text{s}^{-1}$) and then dropped sharply to $1.94 \text{ mg m}^{-1}\text{s}^{-1}$.
278 Therefore, the diurnal variation in the TF during the spring showed the characteristics of a rapid
279 rise and rapid decline. The peak duration was approximately 3 h for a long time in the summer and
280 autumn, where the daily maximum values were $3.79 \text{ mg m}^{-1}\text{s}^{-1}$ and $3.63 \text{ mg m}^{-1}\text{s}^{-1}$ and the minimum
281 values were $1.00 \text{ mg m}^{-1}\text{s}^{-1}$ and $1.30 \text{ mg m}^{-1}\text{s}^{-1}$, respectively. The diurnal variation in the TF during
282 the summer and autumn showed the characteristics of a slow rise and slow decline. Specifically, the
283 daily variation had a strong fluctuation in the winter, peaked three times at 14:00 LT ($4.06 \text{ mg m}^{-1}\text{s}^{-1}$),
284 16:00 LT ($4.38 \text{ mg m}^{-1}\text{s}^{-1}$) and 19:00 LT ($4.07 \text{ mg m}^{-1}\text{s}^{-1}$), then dropped slowly to $1.66 \text{ mg m}^{-1}\text{s}^{-1}$.
285 Another special point is that in the spring, summer and autumn, a high TF corresponds to a
286 southerly wind, while in the winter, the southerly wind does not appear in the whole transport
287 process; instead, there is a westerly wind, which is influenced by the Siberian High.
288 Even so, the TF variation patterns can be summarized as that a high TF corresponds to a southerly
289 wind and a low TF corresponds to a northerly wind (Fig. 5). When the average wind direction in the
290 mixing layer changes from north to south, the TF gradually increases from the daily minimum to
291 the daily maximum. The TF increased by 5 times in the spring, 4 times in the summer, and 3 times
292 in the autumn and winter. The current pattern is because areas located in the south of Beijing are
293 heavily polluted and southerly winds help transport pollutants into the city, leading to high TFs in
294 the afternoons (Fig. 5). However, due to the high mixing layer in the spring, the concentration of
295 near-surface $\text{PM}_{2.5}$ did not increase. The results further confirm the conclusion that the northwest
296 wind in Beijing is a clean wind (Wang et al. 2015; Zhang et al. 2018). Thus, the northwest wind is
297 conducive to the outward transport of pollutants from Beijing, which helps to alleviate pollution. As
298 a result, there was no high TF in the winter when the northwest wind prevailed. On the other hand,
299 southerly winds are stronger than northerly winds (Fig. 5), which can also result in a high TF.
300 Therefore, the level of the TF is determined by two factors, the WS and $\text{PM}_{2.5}$ concentration. In the
301 spring, summer and autumn, a strong south wind prevails in the afternoon. As the south wind is
302 often accompanied by high $\text{PM}_{2.5}$ concentrations (Fig. S2), the TF is high. In the winter, the whole
303 day is dominated by westerly and northerly winds. Although the northerly winds are strong, the TF
304 is not high due to the low $\text{PM}_{2.5}$ concentration. Generally, a high WS means fast mixing, and the
305 corresponding MLH is also high. At this time, the TF is mainly controlled by the WS. While the WS
306 is low, the mixing speed is slow and the MLH is low. At this time, the TF is mainly controlled by
307 $\text{PM}_{2.5}$ concentration. From the above analysis, it can be inferred that if the MLH and WS gradually
308 decrease with the worsening of the pollution, the mixing layer TF is controlled by the WS first and
309 then by the $\text{PM}_{2.5}$ concentration, a maximum TF may occur at a critical moment. This moment is
310 neither the moment of the maximum WS nor the moment of the maximum $\text{PM}_{2.5}$ concentration but
311 rather should be somewhere in between. This will be discussed in more detail in section 3.3.



312

313

314

Fig. 5 Diurnal variations in the mixing layer TF of $PM_{2.5}$ and transport direction during different seasons in Beijing.

315

3.2.2 Vertical evolution of TF

316

317

318

319

320

321

322

323

324

325

326

327

328

329

330

331

332

333

334

335

336

337

338

339

340

341

342

343

344

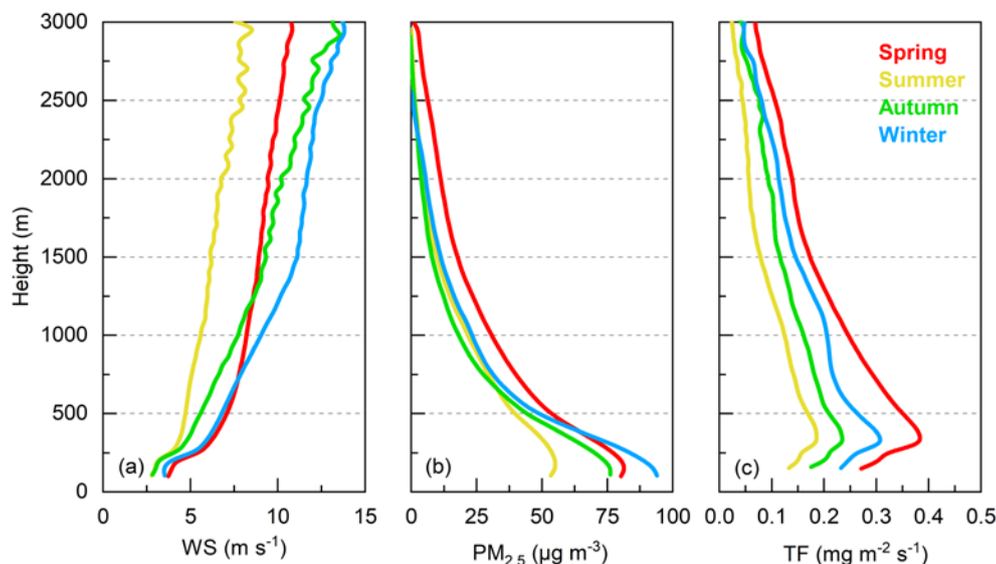
345

346

After the aforementioned analyses, the transport period is known. To further explore the height of transport, we studied the seasonal variation of the TF profile in combination with the vertical wind and $PM_{2.5}$ profiles. With the increasing altitude, the WS first increases sharply at approximately 200 m and then slowly increases, and the differences between different seasons gradually become significant. The WS is always smallest in the summer and strongest in the winter. It is the same in the spring and autumn at 1200 m. Above 1200 m, the WS in the autumn exceeds that in the spring. The $PM_{2.5}$ concentration at 100 m obtained by inversion is highest in the winter ($93.7 \text{ mg m}^{-2}\text{s}^{-1}$), similar in the spring and autumn ($80.3 \text{ mg m}^{-2}\text{s}^{-1}$ and $75.8 \text{ mg m}^{-2}\text{s}^{-1}$, respectively), and lowest in the summer ($53.5 \text{ mg m}^{-2}\text{s}^{-1}$). This finding is consistent with the near-surface results. Below 200 m, the $PM_{2.5}$ concentration is relatively uniform. As the height increases, the $PM_{2.5}$ concentration decreases gradually. Between 200-600 m, the $PM_{2.5}$ concentration begins to decrease rapidly, but the rate of decline is obviously different in different seasons. In the autumn and winter, the reduction rate of the $PM_{2.5}$ concentration is significantly higher than that in the spring and summer. As a result, the spring $PM_{2.5}$ concentration at 400 m begins to be greater than that in the winter; the summer $PM_{2.5}$ concentration at 650 m begins to be greater than that in the autumn and is at the same level as that in the winter. Over 600 m, there is no significant difference in the $PM_{2.5}$ concentration between different seasons, while the WS varies greatly. Therefore, the TF is greatly affected by the WS at high altitudes, and it is greatly influenced by the $PM_{2.5}$ concentration near the ground. The TF in the mixing layer is also affected by the MLH.

The vertical evolution of the TF is different from both the evolution of the WS and $PM_{2.5}$ concentration, and the seasonal variation remains consistent from the near-surface to the upper air. The TF for the spring is the highest, followed by the winter and autumn, and that of the summer is the lowest. The vertical variation in the TF increases first and then decreases, and a peak appears at approximately 300 m, with a value of $0.38 \text{ mg m}^{-2}\text{s}^{-1}$ in the spring, $0.19 \text{ mg m}^{-2}\text{s}^{-1}$ in the summer, $0.24 \text{ mg m}^{-2}\text{s}^{-1}$ in the autumn, and $0.31 \text{ mg m}^{-2}\text{s}^{-1}$ in the winter. In the process of the TF lowering, it has different performances in different seasons. In the spring, the decline slows down at approximately 1500 m. The changes in the summer and autumn are similar. After the peak, the TF drops rapidly in the summer and autumn. The decrease rate above 500 m becomes slow, slows down again after 1500 m, and finally the profiles become vertical. In the winter, the TF declines rapidly, followed by fluctuations at approximately 1000 m. The above results preliminarily indicate that the transport mainly occurs within 200-1500 m, which will be evaluated in Sec. 3.3. To sum up, in the

347 autumn and winter, the high concentration of $PM_{2.5}$ is concentrated near the ground, while the TF is
 348 not large, again indicating that local emissions are the main source of $PM_{2.5}$ in the autumn and winter;
 349 in the spring, affected by high-altitude transport, the $PM_{2.5}$ concentration is high; and in the summer,
 350 both the TF and $PM_{2.5}$ concentration are at their lowest levels, indicating that regional transport may
 351 play an important role in the $PM_{2.5}$ concentration in the summer.



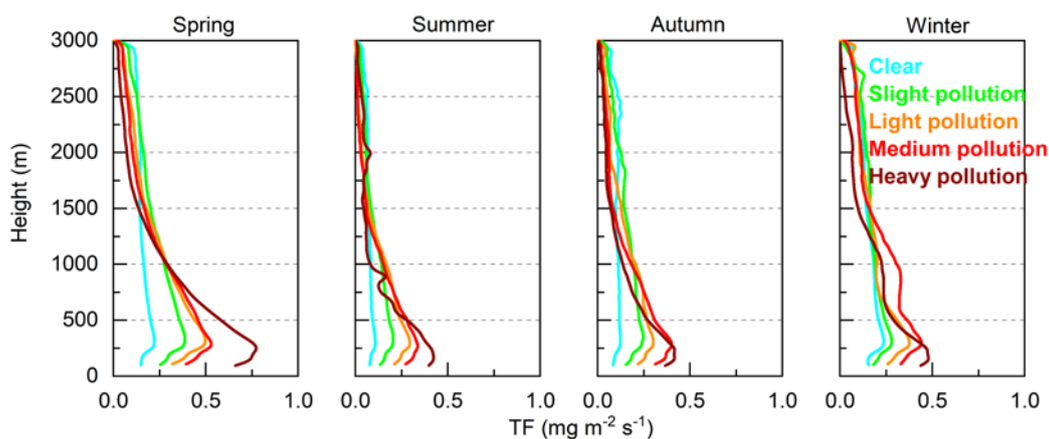
352
 353 Fig. 6 Vertical profiles of WS (a), $PM_{2.5}$ (b) and TF of $PM_{2.5}$ (c) in different seasons in Beijing.

354 3.3 TF under different degrees of air pollution

355 Previous studies have demonstrated that transport occurs only in the transition period of pollution,
 356 while it is weak at the peak of pollution (Tang et al. 2015; Zhu et al. 2016). To quantify the transport
 357 impact of different pollution levels, the $PM_{2.5}$ concentration was divided into five levels according
 358 to the “Technical Regulation on Ambient Air Quality Index (on trial)” (HJ 633-2012): $PM_{2.5} \leq$
 359 $35 \mu g m^{-3}$ (clear days), $35 < PM_{2.5} \leq 75 \mu g m^{-3}$ (slight pollution), $75 < PM_{2.5} \leq 115 \mu g m^{-3}$ (light
 360 pollution), $115 < PM_{2.5} \leq 150 \mu g m^{-3}$ (medium pollution) and $PM_{2.5} > 150 \mu g m^{-3}$ (heavy pollution).
 361 An interesting phenomenon is that with the increase in altitude, the heavier the pollution near the
 362 ground is, the greater the reduction rate of the $PM_{2.5}$ concentration is (Fig. 7). As a result, there is a
 363 reversal at 1000-1500 m. In other words, the more severe the near-surface pollution, the lower the
 364 high-altitude $PM_{2.5}$ concentration. This is particularly outstanding in the spring: from a clear to a
 365 heavy polluted day, the TF at 100 m was, in turn, $0.15 mg m^{-2} s^{-1}$, $0.26 mg m^{-2} s^{-1}$, $0.32 mg m^{-2} s^{-1}$,
 366 $0.39 mg m^{-2} s^{-1}$, $0.66 mg m^{-2} s^{-1}$, and at 2600 m, the values dropped to $0.15 mg m^{-2} s^{-1}$, $0.17 mg m^{-2} s^{-1}$,
 367 $0.13 mg m^{-2} s^{-1}$, $0.10 mg m^{-2} s^{-1}$ and $0.07 mg m^{-2} s^{-1}$, respectively. That is, the lower the pollution
 368 degree, the more vertical the TF tends to be. This is related to the MLH, because a high MLH is
 369 conducive to the diffusion of pollutants in the vertical direction. With the worsening of pollution,
 370 the MLH shows a downward trend (Fig. S3).

371 According to the previous analysis, two peaks may appear in the TF profile, indicating that the
 372 transport occurs at two different heights, approximately 200 m (low-altitude transport) and 1000 m
 373 (high-altitude transport), respectively. Due to the sudden increase in the WS at approximately 200
 374 m, the low-altitude transport at 200 m is the basic transport height, regardless of the season and the
 375 degree of pollution. In contrast, the high-altitude transport is quite special and mainly occurs in the

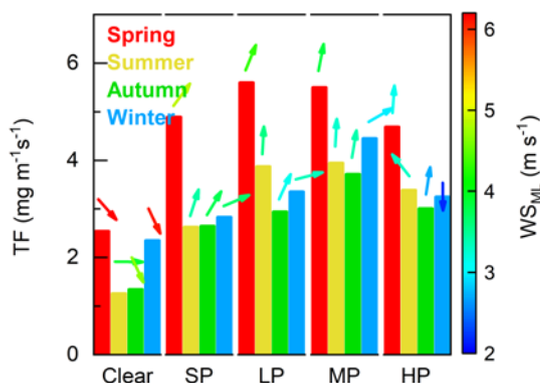
376 winter when there is significant pollution. A small peak of in the TF can also be found on heavy
 377 polluted days in the summer. Although the change in the TF profile of medium pollution in the
 378 autumn is not as obvious as that in the summer and winter, a small increase can still be seen (Fig.
 379 7). In the case of heavy pollution, the MLH is usually less than 1000 m, while in the case of clear
 380 and slight pollution, the MLH is close to the height of high-altitude transport (Fig. S3). Therefore,
 381 it can be inferred that the pollutants transported at a high altitude during heavy pollution are stored
 382 in the residual layer, and when the mixing layer becomes higher, the pollutants stored in the residual
 383 layer diffuse into the mixing layer, affecting the pollution level within the mixing layer. This may
 384 be a key contributor to the slight pollution in the summer, autumn and winter, but further research
 385 is needed.



386
 387 Fig. 7 Vertical profiles of TF of $PM_{2.5}$ under different degrees of pollution in different seasons in
 388 Beijing. Clear days: $PM_{2.5} \leq 35 \mu g m^{-3}$, slight pollution: $35 < PM_{2.5} \leq 75 \mu g m^{-3}$, light pollution: 75
 389 $< PM_{2.5} \leq 115 \mu g m^{-3}$, medium pollution: $115 < PM_{2.5} \leq 150 \mu g m^{-3}$ and heavy pollution: $PM_{2.5} >$
 390 $150 \mu g m^{-3}$.

391 According to the same division method, we further explored the seasonal variation of the TF and
 392 transport source in the mixing layer under different pollution degrees. With pollution aggravation,
 393 the mixing layer TF in Beijing increased by varying degrees during different seasons, and the
 394 transport direction gradually shifted from northwest to south (except during the winter) (Fig. 8). In
 395 particular, the mixing layer TF in the spring is significantly higher than that in the other seasons at
 396 all pollution degrees, which is 1.1-2.0 times that in the other seasons. This may be caused by the
 397 greater amount of dust during the spring. With the pollution deterioration, the TF showed an
 398 increasing trend in the initial stage of pollution and a decreasing trend during the heavy pollution
 399 period. From medium pollution to heavy pollution, the TF decreased from $5.50 \pm 4.83 mg m^{-1}s^{-1}$ to
 400 $4.69 \pm 4.84 mg m^{-1}s^{-1}$ in the spring, from $3.94 \pm 2.36 mg m^{-1}s^{-1}$ to $3.39 \pm 1.77 mg m^{-1}s^{-1}$ in the summer,
 401 from $3.72 \pm 2.86 mg m^{-1}s^{-1}$ to $3.01 \pm 2.40 mg m^{-1}s^{-1}$ in the autumn, and from $4.45 \pm 4.40 mg m^{-1}s^{-1}$
 402 to $3.25 \pm 2.77 mg m^{-1}s^{-1}$ in the winter. Among them, the largest drop was found in the winter. In the
 403 winter, with the pollution aggravation, the transport direction changed from northwest to southwest
 404 and finally to the north. In contrast to in the other seasons, the weak north wind was the main wind
 405 during heavy pollution in the winter, indicating that regional transport contributed less to the heavy
 406 pollution during the winter in Beijing. In the initial stage of pollution, the TF continued to increase,
 407 but the rate of increase gradually slowed in the spring and summer. From light pollution to medium
 408 pollution, the TF decreased by $0.1 mg m^{-1}s^{-1}$ in the spring and increased by only $0.07 mg m^{-1}s^{-1}$ in
 409 the summer. It is also not difficult to find from the changes in the TF profile (Fig. 7) that the regional

410 transport has little impact on the medium pollution in the spring and summer. These results indicate
 411 that although the region south of Beijing is the main transport source in Beijing, its contribution is
 412 significantly reduced during the severe pollution period. In general, regional transport plays an
 413 important role in the initial period of pollution, while local emissions are the main controlling factor
 414 during the period of heavy pollution. The parabolic pattern of the TF is the result of a combination
 415 of the WS and PM_{2.5} concentration. The TF reaches a threshold during medium pollution, which is
 416 the critical moment mentioned above.



417
 418 Fig. 8 The mixing layer TF of the PM_{2.5} levels and transport directions under different degrees of
 419 pollution in different seasons in Beijing. (SP denotes slight pollution, LP denotes light pollution,
 420 MP denotes medium pollution and HP denotes heavy pollution.)

421 4. Conclusions

422 To understand the characteristics of the PM_{2.5} transport flux in Beijing, the height of the atmospheric
 423 mixing layer and the wind profile within the mixing layer in Beijing were observed for a 2-year
 424 period. The main conclusions are as follows:

425 (1) By analyzing the variations in the VC, it is found that the atmospheric dilution capability in
 426 Beijing is strongest in the spring and weaker in the summer, autumn and winter. In the spring, the
 427 vertical and horizontal dilution capacities are strong; in the autumn and winter, the vertical and
 428 horizontal dilution capacities are weak; and in the summer, the vertical dilution capability is strong
 429 and the horizontal dilution capability is weak. The diurnal variation in the VC is consistent with the
 430 MLH, which shows that the dilution capability is the strongest before sunset, gradually weakens
 431 after sunset and remains stable at night. In the spring, the vertical and horizontal dilutions are strong
 432 during both day and night. In the winter, the vertical dilution is weak during the day, and the
 433 horizontal dilution during the night is the main component. In the summer, the vertical dilution
 434 during the day is dominant. Although there is little difference in the diffusivity between the summer,
 435 autumn and winter, the poor dilution capability occurs more frequently in the autumn and winter.

436 (2) The TF is the largest in the spring and smaller in the summer, autumn and winter in Beijing. The
 437 high TF mainly comes from southward transport, while the low TF is accompanied by northwest
 438 transport. The transport mainly occurred between 14:00 and 18:00 LT, and the height of the transport
 439 is at approximately 200 m and 1000 m. Using the PM_{2.5} concentration as a classification index for
 440 the air pollution, the results show that the regional transport from the southern area plays an
 441 important role in the initial period of pollution, and local emissions are the main controlling factors

442 in the heavy pollution period, especially in the winter.
443 To solve the problem of heavy pollution in northern China, joint prevention and control has been
444 suggested for a long time. Even so, there is still no concrete implementation plan. To break through
445 this embarrassing situation, this study quantifies TF to explain the time period when the transport
446 occurs and the main areas affected in Beijing. In this study, the atmospheric dilution capability
447 during different seasons and the TF during different pollution periods were also discussed. The
448 important role of transport in the initial period of pollution is emphasized, and local pollutant
449 emission control is found to be the most effective way of mitigating pollution levels. The research
450 results are of great significance to the early warning, prevention and control of atmospheric
451 particulate pollution.

452 **Data availability**

453 The data in this study are available from the corresponding author upon request (tgq@dq.cern.ac.cn).

454 **Author contribution**

455 GT and YW designed the research, LZ, BH, BL and YunL conducted the measurements. YusL and
456 GT wrote the paper. SL reviewed and commented on the paper.

457 **Competing interests**

458 The authors declare that they have no conflicts of interest to disclose.

459 **Acknowledgments**

460 This work was supported by the National Key R&D Program of China (2018YFC0213201), the
461 Young Talent Project of the Center for Excellence in Regional Atmospheric Environment CAS
462 (CERAE201802), LAC/CMA (2017A01), the National Research Program for Key Issues in Air
463 Pollution Control (DQGG0101), the National Natural Science Foundation of China (Nos. 41705113
464 and 41877312), and the Foundation of the Chinese Academy of Meteorological Sciences
465 (2019Y001).

466 **References**

- 467 Chang, X., S. Wang, B. Zhao, S. Cai, and J. Hao: Assessment of inter-city transport of particulate matter
468 in the Beijing–Tianjin–Hebei region, *Atmos. Chem. Phys.*, 18, 4843–4858, doi: 10.5194/acp-
469 18-4843-2018, 2018.
- 470 Cheng, J., J. Su, T. Cui, X. Li, X. Dong, F. Sun, Y. Yang, D. Tong, Y. Zheng, J. Li, Q. Zhang, and K. He:
471 Dominant role of emission reduction in PM_{2.5} air quality improvement in Beijing during 2013-
472 2017: a model-based decomposition analysis, *Atmos. Chem. Phys. Discussions*, 1-31, doi:
473 10.5194/acp-2018-1145, 2018a.
- 474 Cheng, N., Y. Li, B. Cheng, X. Wang, F. Meng, Q. Wang, and Q. Qiu: Comparisons of two serious air
475 pollution episodes in winter and summer in Beijing, *J. Environ. Sci-China.*, 69, 141-154, doi:

476 10.1016/j.jes.2017.10.002, 2018b.

477 Ge, B., Z. Wang, W. Lin, X. Xu, J. Li, D. Ji, and Z. Ma: Air pollution over the North China Plain and its
478 implication of regional transport: A new sight from the observed evidences, *Environ. Pollut.*,
479 234, 29-38, doi: 10.1016/j.envpol.2017.10.084, 2018.

480 Geiß, A., M. Wiegner, B. Bonn, K. Schäfer, R. Forkel, E. von Schneidemesser, C. Münkel, K. L. Chan,
481 and R. Nothard: Mixing layer height as an indicator for urban air quality?, *Atmos. Meas. Tech.*,
482 10, 2969-2988, doi: 10.5194/amt-10-2969-2017, 2017.

483 Huang, Q., X. Cai, J. Wang, Y. Song, and T. Zhu: Climatological study of the boundary-layer air
484 stagnation Index for China and its relationship with air pollution, *Atmos. Chem. Phys.*, 18, 7573-
485 7593, doi: 10.5194/acp-18-7573-2018, 2018.

486 Li, H., F. Duan, Y. Ma, K. He, L. Zhu, T. Ma, S. Ye, S. Yang, T. Huang, and T. Kimoto: Case study of
487 spring haze in Beijing: Characteristics, formation processes, secondary transition, and regional
488 transportation, *Environ. Pollut.*, 242, 544-554, doi: 10.1016/j.envpol.2018.07.001, 2018.

489 Liu, L., J. Guo, Y. Miao, L. Liu, J. Li, D. Chen, J. He, and C. Cui: Elucidating the relationship between
490 aerosol concentration and summertime boundary layer structure in central China, *Environ.*
491 *Pollut.*, 241, 646-653, doi: 10.1016/j.envpol.2018.06.008, 2018.

492 Miao, Y., and S. Liu: Linkages between aerosol pollution and planetary boundary layer structure in China,
493 *Sci. Total. Environ.*, 650, 288-296, doi: 10.1016/j.scitotenv.2018.09.032, 2019.

494 Mues, A., M. Rupakheti, C. Münkel, A. Lauer, H. Bozem, P. Hoor, T. Butler, and M. G. Lawrence:
495 Investigation of the mixing layer height derived from ceilometer measurements in the
496 Kathmandu Valley and implications for local air quality, *Atmos. Chem. Phys.*, 17, 8157-8176,
497 doi: 10.5194/acp-17-8157-2017, 2017.

498 Münkel, C., N. Eresmaa, J. Räsänen, and A. Karppinen: Retrieval of mixing height and dust
499 concentration with lidar ceilometer, *Bound-Lay. Meteorol.*, 124, 117-128, doi: 10.1007/s10546-
500 006-9103-3, 2007.

501 Nair, V. S., K. K. Moorthy, D. P. Alappattu, P. K. Kunhikrishnan, S. George, P. R. Nair, S. S. Babu, B.
502 Abish, S. K. Satheesh, S. N. Tripathi, K. Niranjana, B. L. Madhavan, V. Srikant, C. B. S. Dutt,
503 K. V. S. Badarinath, and R. R. Reddy: Wintertime aerosol characteristics over the Indo-Gangetic
504 Plain (IGP): Impacts of local boundary layer processes and long-range transport, *J. Geophys.*
505 *Res-Atmos.*, 112, D13205, doi: 10.1029/2006jd008099, 2007.

506 Ouyang, W., Y. Xu, J. Cao, X. Gao, B. Gao, Z. Hao, and C. Lin: Rainwater characteristics and interaction
507 with atmospheric particle matter transportation analyzed by remote sensing around Beijing, *Sci.*
508 *Total Environ.*, 651, 532-540, doi: 10.1016/j.scitotenv.2018.09.120, 2019.

509 Schäfer, K., S. Emeis, H. Hoffmann, and C. Jahn: Influence of mixing layer height upon air pollution in
510 urban and sub-urban areas, *Meteorol. Z.*, 15, 647-658, doi: 10.1127/0941-2948/2006/0164,
511 2006.

512 Shen, Y., L. Zhang, X. Fang, H. Ji, X. Li, and Z. Zhao: Spatiotemporal patterns of recent PM_{2.5}
513 concentrations over typical urban agglomerations in China, *Sci Total Environ*, 655, 13-26, doi:
514 10.1016/j.scitotenv.2018.11.105, 2019.

515 Song, L. C., G. Rong, L. Ying, and W. Guo-Fu: Analysis of China's haze days in the winter half-year and
516 the climatic background during 1961–2012, *Advances in Climate Change Research*, 5, 1-6, doi:
517 10.3724/sp.j.1248.2014.001, 2014.

518 Steyn, D. G., M. Baldi, and R. M. Hoff: The detection of mixed layer depth and entrainment zone
519 thickness from lidar backscatter profiles, *J. Atmos. Ocean. Technol.*, 16, 953-959, doi, 1999.

520 Su, T., Z. Li, and R. Kahn: Relationships between the planetary boundary layer height and surface
521 pollutants derived from lidar observations over China, *Atmos. Chem. Phys.*, 18, 15921-15935,
522 doi: 10.5194/acp-18-15921-2018, 2018.

523 Tang, G., J. Zhang, X. Zhu, T. Song, C. Münkel, B. Hu, K. Schäfer, Z. Liu, J. Zhang, L. Wang, J. Xin, P.
524 Suppan, and Y. Wang: Mixing layer height and its implications for air pollution over Beijing,
525 China, *Atmos. Chem. Phys.*, 16, 2459-2475, doi: 10.5194/acp-16-2459-2016, 2016.

526 Tang, G., X. Zhu, B. Hu, J. Xin, L. Wang, C. Munkel, G. Mao, and Y. Wang: Impact of emission controls
527 on air quality in Beijing during APEC 2014: lidar ceilometer observations, *Atmos. Chem. Phys.*,
528 15, 12667-12680, doi: 10.5194/acp-15-12667-2015, 2015.

529 Wang, L., Z. Liu, Y. Sun, D. Ji, and Y. Wang: Long-range transport and regional sources of PM_{2.5} in
530 Beijing based on long-term observations from 2005 to 2010, *Atmos. Res.*, 37-48, doi:
531 10.1016/j.atmosres.2014.12.003, 2015.

532 Wang, X., K. Wang, and L. Su: Contribution of Atmospheric Diffusion Conditions to the Recent
533 Improvement in Air Quality in China, *Sci. Rep-UK.*, 6, 36404, doi: 10.1038/srep36404, 2016.

534 Yang, Y., H. Liao, and S. Lou: Increase in winter haze over eastern China in recent decades: Roles of
535 variations in meteorological parameters and anthropogenic emissions, *J. Geophys. Res. Atmos.*,
536 121, 13,050-013,065, doi: 10.1002/2016jd025136, 2016.

537 Zhang, H., S. Cheng, X. Wang, S. Yao, and F. Zhu: Continuous monitoring, compositions analysis and
538 the implication of regional transport for submicron and fine aerosols in Beijing, China, *Atmos.*
539 *Environ.*, 195, 30-45, doi: 10.1016/j.atmosenv.2018.09.043, 2018.

540 Zhang, Y., J. Chen, H. Yang, R. Li, and Q. Yu: Seasonal variation and potential source regions of PM_{2.5}-
541 bound PAHs in the megacity Beijing, China: Impact of regional transport, *Environ. Pollut.*, 231,
542 329-338, doi: 10.1016/j.envpol.2017.08.025, 2017.

543 Zhu, X., G. Tang, J. Guo, B. Hu, T. Song, L. Wang, J. Xin, W. Gao, C. Münkel, K. Schäfer, X. Li, and Y.
544 Wang: Mixing layer height on the North China Plain and meteorological evidence of serious air
545 pollution in southern Hebei, *Atmos. Chem. Phys.*, 18, 4897-4910, doi: 10.5194/acp-18-4897-
546 2018, 2018.

547 Zhu, X. W., G. Q. Tang, B. Hu, L. L. Wang, J. Y. Xin, J. K. Zhang, Z. R. Liu, C. Munkel, and Y. S. Wang:
548 Regional pollution and its formation mechanism over North China Plain: A case study with
549 ceilometer observations and model simulations, *J. Geophys. Res-Atmos.*, 121, 14574-14588,
550 doi: 10.1002/2016jd025730, 2016.

551