Simultaneous measurements of particle number size distributions at ground level and 260 m on a meteorological tower in urban Beijing, China

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Abstract. Despite extensive studies into the characterization of particle number size distributions at ground level, real-time measurements above the urban canopy in the megacity of Beijing have never been performed to date. Here we conducted the first simultaneous measurements of size-resolved particle number concentrations at ground level and 260 m in urban Beijing from 22 August to 30 September. Our results showed overall similar temporal variations in number size distributions between ground level and 260 m, yet periods with significant differences were also observed. Particularly, accumulation-mode particles were highly correlated ($r^2 = 0.85$) at the two heights, while Aitken-mode particles presented more differences. Detailed analysis suggests that the vertical differences in number concentrations strongly depended on particle size, and particles with a mobility diameter between 100 and 200 nm generally showed higher concentrations at higher altitudes. Particle growth rates and condensation sinks were also calculated, which were 3.2 and 3.6 nm h$^{-1}$, and $2.8 \times 10^{-2}$ and $2.9 \times 10^{-2}$ s$^{-1}$, at ground level and 260 m, respectively. By linking particle growth with aerosol composition, we found that organics appeared to play an important role in the early stage of the growth (09:00–12:00 LT) while sulfate was also important during the later period. Positive matrix factorization of size-resolved number concentrations identified three common sources at ground level and 260 m, including a factor associated with new particle formation and growth events (NPEs), and two secondary factors that represent photochemical processing and regional transport. Cooking emission was found to have a large contribution to small particles and showed much higher concentration at ground level than 260 m in the evening. These results imply that investigation of NPEs at ground level in megacities needs to consider the influences of local cooking emissions. The impacts of regional emission controls on particle number concentrations were also illustrated. Our results showed that regional emission controls have a dominant impact on accumulation-mode particles by decreasing gas precursors and particulate matter loadings, and hence suppressing particle growth. In contrast, the influences on Aitken particles were much smaller due to the enhanced new particle formation (NPF) events.

1 Introduction

With frequent occurrence of haze episodes, the megacity of Beijing is faced with severe air pollution problems, as indicated by high concentrations of ambient aerosol particles. For example, the annual average concentration of PM$_{2.5}$...
was 80.6 µg m$^{-3}$ in 2015, which is more than twice the China National Ambient Air Quality Standard (35 µg m$^{-3}$ as an annual average) (http://www.bjepb.gov.cn/bjepb/413526/413663/413717/413719/index.html). Fine particles can significantly reduce atmospheric visibility, exert harmful effects on public health, and even have potential impacts on regional and global climate. As a result, extensive efforts have been devoted to characterize the sources, formation mechanisms, and evolution processes of aerosol particles in recent years (Ma et al., 2012; Takegawa et al., 2009; Sun et al., 2010, 2014, 2016b). Among these studies, particle number concentrations are one of the greatest concerns because particles can rapidly grow from a few nanometers to tens and even hundreds of nanometers in a short time, and hence play a significant role in haze formation (Guo et al., 2014). However, our understanding of the formation and growth of aerosol particles is not complete, particularly in highly polluted environments (Kulmala et al., 2016).

In the past decades, extensive studies have been conducted to characterize particle number size distributions in Beijing at ground level (Wehner et al., 2004; Yue et al., 2009; Wu et al., 2011; Gao et al., 2012; Wang et al., 2013b). The continuous measurements of aerosol number size distributions from 3 nm to 10 µm within the city area of Beijing in spring indicated a high variability in number concentrations, and the variations were substantially different among dust storm, clean, and polluted periods (Wehner et al., 2004). Yue et al. (2009) also found a clear shift of maximum diameter from 60 nm on clean days to 80 nm during polluted days. Most previous studies focused on new particle formation and growth events (NPEs) (Wehner et al., 2004; Wu et al., 2011; Gao et al., 2012; Zhang et al., 2011). While new particle formation (NPF) events are mostly observed under conditions with low relative humidity and clean air masses (Wu et al., 2007; Wehner et al., 2004), particle growth events are strongly associated with high relative humidity (Gao et al., 2012). The roles of chemical species in NPEs in Beijing were also explored in several studies. For example, organics were found to be the dominant species of PM$_1$ during new particle formation events in summer in Beijing (Zhang et al., 2011) and likely played a major role in NPF and growth, although sulfuric is also important as well (Yue et al., 2009). However, most of these studies were conducted at a ground site that is subject to the influences of multiple local sources, e.g., traffic and cooking emissions. Indeed, the source apportionment of particle numbers with positive matrix factorization showed significant contributions of traffic emissions and combustion sources to the total number concentration (Wang et al., 2013b; Liu et al., 2014). Therefore, measurements of size-resolved number concentrations at high altitude with less local cooking and traffic influences are essential for elucidating the NPF and growth mechanisms, and also the role of regional transport in haze formation.

During this study period, strict emission controls were implemented in Beijing and the surrounding regions, e.g., Hebei, Tianjin, and Shandong, from 20 August to 3 September to ensure good air quality during the China Victory Day (V-day) parade on 3 September 2015. The control measures, such as restricting the number of vehicles, shutting down factories and power plants, stopping construction activities, etc., were even stricter than those implemented during the Asia-Pacific Economic Cooperation (APEC) summit in 2014 (Sun et al., 2016a). Several studies have addressed the impacts of regional emission controls on aerosol composition and gaseous species (Li et al., 2016; Han et al., 2016; Zhao et al., 2017). The results are overall consistent, showing significant reductions in most aerosol and gaseous species during the control period (CP, 22 August–3 September). A recent study comparing the number size distributions with those during the same period in 2010–2013 at a rural site in Beijing illustrated the most reductions in accumulation-mode particles and condensation sink (CS) during the V-day period (Shen et al., 2016). Despite this, our understanding of the impacts of emission controls on particle number size distributions is far from complete.

Here, we conducted the first simultaneous measurements of particle number size distributions at two different heights, i.e., ground level and 260 m, within the city area of Beijing from 22 August to 30 September. This study is unique because it provides an experimental opportunity to investigate the vertical differences and processes of particle number size distributions and also the impacts of regional emission controls. The size-resolved particle number concentrations, diurnal variations, particle growth rates, and their relationships with aerosol composition at ground level and 260 m are compared in detail, and the impacts of emission controls on particle number concentrations in different sizes are elucidated. In addition, the sources of particle numbers at the two different heights are investigated with positive matrix factorization.

2 Experimental method

2.1 Sampling and data analysis

The sampling site is located at the Tower Branch of the Institute of Atmospheric Physics, Chinese Academy of Sciences, between the north third and fourth ring roads in Beijing. Two scanning mobility particle sizers (SMPSs) were deployed for simultaneous measurements of particle number size distributions at ground level and 260 m on the Beijing 325 m meteorological tower. At 260 m, the size-resolved particle number concentration (15–685 nm) was measured in situ by a condensation particle counter (TSI, 3775) equipped with a long differential mobility analyzer (TSI, 3081A). The time resolution is 5 min. Comparatively, an SMPS as part of an unattended multifunctional hygroscopic tandem differential mobility analyzer (H-TDMA) developed by the Guangzhou Institute of Tropical and Marine Meteorology, China Meteorological Administration (ITMM, CMA) was used to mea-
measure particle number concentrations (10–400 nm) at ground level. A detailed description of the H-TDMA was given in Tan et al. (2013). According to previous comparisons of particle number size distributions between different SMPSs or differential mobility particle sizers, the measurement uncertainties between 20 and 200 nm can be \( \sim 10\% \), and even larger for particles outside this range (Wiedensohler et al., 2012).

The non-refractory submicron aerosol (NR-PM\(_1\)) species, including organics (Org), sulfate (SO\(_4\)), nitrate (NO\(_3\)), ammonium (NH\(_4\)), and chloride (Chl), were measured at ground level by an Aerodyne high-resolution time-of-flight aerosol mass spectrometer (HR-AMS) and at 260 m by an aerosol chemical speciation monitor (ACSM). Co-located black carbon (BC) was measured by a seven-wavelength (AE33) and a two-wavelength Aethalometer (AE22, Magee Scientific Corp.) at 260 m and ground level, respectively. The meteorological variables, including wind speed (WS), wind direction (WD), relative humidity (RH), and temperature (T), were obtained from the measurements on the meteorological tower. The operations of the HR-AMS, ACSM, and Aethalometers and subsequent analysis details are described in Sun et al. (2015b) and Zhao et al. (2017). All the data in this study are reported in Beijing local time (UTC+8 h).

Figure S1 in the Supplement shows a comparison of the total PM\(_1\) mass (NR-PM\(_1\)+BC) with that derived from the SMPS measurements at ground level and 260 m. The particle number concentrations between 15 and 400 nm were converted to mass concentrations using chemically resolved particle density (Salcedo et al., 2006). As shown in Fig. S1, the time series of PM\(_1\) was highly correlated with that from SMPS measurements at both ground level \((r^2 = 0.94)\) and 260 m \((r^2 = 0.95)\). We also noticed some differences in the regression slopes, which are 0.44 and 0.66 at ground level and 260 m, respectively. The reasons are not very clear yet, but these differences are likely due to the different size distributions at the two different heights (Sect. 3.1).

### 2.2 Particle growth rates and condensation sink

The particle growth rates (GRs) at ground level and 260 m were calculated using Eq. (1).

\[
GR = \frac{\Delta D_m}{\Delta t},
\]

where \(D_m\) is the geometric mean diameter from the log-normal fitting of each size distribution and \(\Delta D_m\) is the increase in diameter during the growth period of \(\Delta t\).

Condensation sink (CS), indicating how rapidly vapor molecules can condense onto preexisting aerosols, is calculated using Eq. (2) (Nieminen et al., 2010).

\[
CS = 2\pi D \sum_i \beta M_i D_{p,i} N_i,
\]

where \(D\) is the diffusion coefficient of the condensing vapor, \(D_{p}\) and \(N\) are the particle diameter and the corresponding number concentration, and \(\beta M\) is the transitional regime correction factor expressed as Eq. (3).

\[
\beta M = (Kn + 1) \left[ 1 + 0.377Kn + \frac{4}{3} \alpha^{-1}Kn^2 + \frac{4}{3} \alpha^{-1}Kn^3 \right],
\]

where \(\alpha\) is assumed to be unity, and \(Kn\) is the Knudsen number. It should be noted that the CS calculated on the basis of dry particle number size distributions might be underestimated since ambient RH was not considered (Reutter et al., 2009).

### 2.3 Source apportionment of size-resolved particle number concentrations

Positive matrix factorization (PMF, 2.exe, v 4.2) was performed on the size-resolved number concentrations (Ulbrich et al., 2009; Paatero and Tapper, 1994) to resolve potential sources. In this study, the measurement uncertainties were estimated using an equation-based approach that was detailed in Ogulei et al. (2007). The required measurement errors \((\sigma_{ij})\) were first calculated using Eq. (4):

\[
\sigma_{ij} = C_1 \times (X_{ij} + \bar{X}_j),
\]
where \( C_1 \) is a constant value assumed to be 0.01, \( X_{ij} \) is the measured particle number concentration, and \( \bar{X}_j \) is the arithmetic mean value for \( j \)th size bin. The measurement uncertainties (Unc) were then calculated with Eq. (5):

\[
\text{Unc}_{ij} = \sigma_{ij} + C_2 \times X_{ij},
\]

(5)

where \( \sigma_{ij} \) is the estimated measurement error and \( C_2 \) is a constant value assumed to be 0.1. After a careful evaluation of the PMF results, five factors and four factors were chosen at ground level and 260 m, respectively. A more detailed diagnosis of PMF results is presented in Figs. S2 and S3.

3 Results and discussion

3.1 Characterization of particle number size distributions

The temporal variations in size-resolved number concentrations and aerosol species at ground level and 260 m are shown in Fig. 1. The size-resolved particle number concentrations showed overall similar evolutionary patterns between ground level and 260 m, and high number concentrations of large particles were generally associated with correspondingly higher concentrations of aerosol species, e.g., the periods of case 1 and case 2 in Fig. 1. However, periods with substantially different number size distributions were also observed. For example, we observed significantly higher particle number concentrations at ground level than 260 m in the evening on 26 August and 1 September due to the influences of local cooking emissions. On average, the particle numbers showed a broader size distribution at 260 m than ground level, peaking at approximately 85 and 45 nm, respectively (Fig. 1c). The log-normal distribution fitting further illustrated three size modes at both ground level and 260 m. While the second mode with geometric mean diameter (GMD) peaking at 41 nm accounted for the largest number fraction at ground level (52 %), the largest mode (GMD = 116 nm) dominated the total number of particles at 260 m, accounting for 62 %. Such differences were likely due to
the stronger influences of local sources (e.g., cooking) with higher emissions of smaller particles, and more influences of regional transport with large aged particles at 260 m.

Figure 2 shows the comparisons of the total number concentrations (15–400 nm, \(N_{15-400}\)) and those for three modes, including small Aitken mode (15–40 nm, \(N_{15-40}\)), large Aitken mode (40–100 nm, \(N_{40-100}\)), and accumulation mode (100–400 nm, \(N_{100-400}\)) between ground level and 260 m. The variation trends in the total number concentrations at the two heights tracked relatively well (\(r^2 = 0.40\), slope = 0.71), while the average number concentration from 15 to 400 nm at 260 m (7473 ± 4324 cm\(^{-3}\)) was 26% lower than that (10134 ± 4680 cm\(^{-3}\)) at ground level. The total particle number concentrations at ground level were generally lower than those previously observed in Beijing mainly due to the smaller size range measured in this study (Wang et al., 2013b; Yue et al., 2009). The \(N_{15-400}\) ratio of 260 m to ground (\(R_{260\text{m/ground}}\)) varied dramatically throughout the entire study, with the daily average ranging from 0.42 to 1.10. In contrast, the total volume concentrations showed much better correlations between ground level and 260 m (\(r^2 = 0.89\)) and the average ratio was close to 1. Such differences were mainly caused by the different contributions of different mode particles to the number and volume concentrations.

The correlations of particle number and volume concentrations between ground level and 260 m varied substantially for different mode particles. As shown in Fig. 2a, the small Aitken-mode particles were correlated between the two heights (\(r^2 = 0.66\)), indicating their common sources that are related to new particle formation. However, the average number concentration at 260 m (1382 ± 1281 cm\(^{-3}\)) was only approximately 40% of that at the ground level (3379 ± 2232 cm\(^{-3}\)), and the daily average ratio of 260 m to ground level for \(N_{15-40}\) varied from 0.91 to 0.51. These results illustrated additional sources for small Aitken particles at ground level. Indeed, pronounced peaks for \(N_{15-40}\) were often observed in the evening, likely indicating the influences of local emissions, e.g., cooking and traffic emissions. The large Aitken-mode particles showed the worst correlation between ground level and 260 m (\(r^2 = 0.40\), slope = 0.70), although the average number concentrations were comparable (4188 vs. 3233 cm\(^{-3}\)). These results suggested that the sources of large Aitken-mode particles were quite different between ground level and 260 m. For example, the diurnal cycle of large Aitken-mode particles at ground level was remarkably similar to that of organic cooking aerosols (Zhao...
Table 2. Summary of average number concentrations of five factors for the entire study, control period (CP), non-control period (NCP), and also the change percentages ((CP-NCP)/NCP × 100).

<table>
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<tr>
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<tr>
<td></td>
<td>260 m</td>
<td>Ground</td>
<td>260 m</td>
<td>Ground</td>
<td>260 m</td>
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<tr>
<td>Entire study (cm⁻³)</td>
<td>867</td>
<td>695</td>
<td>–</td>
<td>2567</td>
<td>2066</td>
</tr>
<tr>
<td>Control period (cm⁻³)</td>
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<td>816</td>
<td>–</td>
<td>2586</td>
<td>2271</td>
</tr>
<tr>
<td>Non-control period (cm⁻³)</td>
<td>771</td>
<td>621</td>
<td>–</td>
<td>2526</td>
<td>1967</td>
</tr>
<tr>
<td>(CP-NCP)/NCP (%)</td>
<td>38 %</td>
<td>31 %</td>
<td>–</td>
<td>2 %</td>
<td>15 %</td>
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Figure 3. Comparisons of particle volume concentrations between ground level and 260 m for different size ranges, i.e., (a) small Aitken mode (15–40 nm), (b) large Aitken mode (40–100 nm), (c) accumulation mode (100–400 nm), and (d) the total number of particles (15–400 nm). Right figure show the scatter plots of the comparisons.

et al., 2017), likely indicating a large source contribution from cooking emissions. Compared with Aitken particles, the number and volume concentrations of accumulation-mode particles were well correlated between the two heights ($r^2 = 0.85$ and 0.91, respectively). While the average number concentration at 260 m was 11 % higher than that at ground level, the volume concentration was close. Moreover, the temporal variations in accumulation-mode particles tracked well with those of secondary inorganic species that were mainly formed over a regional scale. Our results indicate that accumulation-mode particles were likely dominantly from regional transport and relatively homogeneously distributed across different heights. The different vertical ratios between number and volume concentrations suggest that the particle size distributions were slightly different between ground level and 260 m.

The regional emission control and meteorological conditions can have significant impacts on particle number size distributions. As shown in Fig. 1a and b, the GMD of number size distributions peaked at 57 nm at 260 m and 43 nm at ground level during the control period, and the average size distribution showed three similar modes between the two heights. In contrast, the size distributions had substantial changes after the control period that were characterized by much broader distributions and a clear shift from smaller to larger particles at both ground level and 260 m. For exam-
The GMD of particle number distributions was 106 nm at 260 m, which was much larger than that during the control period, and the largest mode consistently dominated the total number of particles, on average accounting for 68%. Figure 4 shows a comparison of average number and volume concentration between control and non-control periods for three-mode particles. While the average total number concentrations during the control period were lower than those during non-control periods (6139 vs. 8116 cm\(^{-3}\) at 260 m and 8708 vs. 10 824 cm\(^{-3}\) at ground level), the small and large Aitken-mode particles were comparable between control and non-control periods. As a result, the decreases in total number concentrations were mainly caused by the changes in accumulation-mode particles, which were decreased by 53 % at 260 m and 52 % at ground level during the control period. Our results illustrate that regional emission control has a large impact on accumulation-mode particles, while the influences on Aitken-mode particles were small. One of the major reasons is that emission controls substantially decrease the gas precursors (e.g., SO\(_2\) and NO\(_x\)) and PM\(_{2.5}\) mass concentrations, and hence suppress the growth of particles. This is also consistent with the large decreases in CS by 48 % at 260 m and 45 % at ground level during the control period (Fig. 4a and b). In addition to regional emission controls, we also found that the dominant northerly winds likely played an important role in decreasing the PM during the control period (Zhao et al., 2017). In contrast, the number concentrations of small particles were relatively comparable due to more frequent new particle formation events during the control period. To better evaluate the impacts of regional emission controls, cluster analysis with hourly back trajectories were performed on the entire dataset with an exclusion of precipitation days. As shown in Fig. S4, accumulation-mode particles during the control period showed the largest reductions for clusters 1 and 2 (39 and 42 %, respectively), while the large Aitken particles had small changes and the small Aitken ones even showed a large increase (43 %) for cluster 1. These results further support our conclusion above.

We also compared the particle number size distributions between polluted (PM\(_{2.5}\) > 75 µg m\(^{-3}\)) and clean days (PM\(_{2.5}\) < 75 µg m\(^{-3}\)) after the control period. As shown in Fig. S5, the average size distribution on polluted days at ground level showed a clear three-mode distribution, peaking at 36, 96, and 244 nm. The GMD of the three modes was ubiquitous larger than those (23, 41, and 106 nm) observed during clean days. While the average total number concentration increased from 10 258 (±4676) cm\(^{-3}\) during clean periods to 12 156 (±4406) cm\(^{-3}\) on polluted days (Fig. 4), we observed comparable concentrations for small and large Aitken-mode particles. Therefore, the increase in total number concentration was mainly caused by the accumulation-mode particles, which increased by 90 % during the polluted days. These results illustrate the different roles of different mode particles between clean and pol-
luted days. Similarly, the average particle number distribution showed a clear shift from a smaller size during clean periods to a larger size on polluted days at 260 m, and the total number concentration was increased by 53 \% from 7006 (±4416) to 10748 (±3615) cm\(^{-3}\). Again, the increase in total number concentration was mainly due to the increase in accumulation-mode particles by 135 \%. Compared with the number concentrations, the increases in volume concentrations for accumulation-mode particles were more significant on polluted days, which on average were 174 and 212 \% at ground level and 260 m, respectively. Indeed, the accumulation-mode particles accounted for 97 \% of total volume concentrations at both ground level and 260 m, elucidating their major roles in PM pollution. The average number ratios between 260 m and ground level increased as a function of particle sizes during both clean and polluted days. For example, the ratios increased from 0.4 to 0.9 for small and large Aitken-mode particles, and to 1.2 for accumulation-mode particles on polluted days. These results are consistent with our previous conclusion that smaller particles showed stronger vertical gradients than larger particles. We also observed ubiquitously higher \(R_{260 \text{m}/\text{ground}}\) on polluted days compared to clean periods, indicating larger vertical gradients in both number and volume concentration during polluted periods.

### 3.2 Diurnal variations

The average diurnal variations in particle number size distribution at ground level and 260 m for the entire study are shown in Fig. 5. It is clear that particle number size distributions show pronounced diurnal cycles that were characterized by the lowest values in the early morning and subsequent particle growth until midnight. During the growth period, the GMD increased from 29 to 57 nm at ground level, while it increased from 41 to 88 nm in 12 h at 260 m. After that, the GMD remained at relatively constant levels of 70 and 100 nm at ground level and 260 m, respectively. The ubiquitously lower GMD and lower growth rates at ground level were likely due to the influences of local emissions that contained a large number of small particles. Note that the changes in GMD were significant at ground level after the control period, especially on polluted days (Fig. S6), indicating that the diurnal evolution of particle number size distributions at ground level is subject to multiple influences. In contrast, the changes at 260 m were much smaller with a relatively consistent mode peaking at ~100 nm, indicating a more constant particle source at higher altitudes.

The particle number ratios between 260 m and ground level depend strongly on particle size. As shown in Fig. 5, \(R_{260 \text{m}/\text{ground}}\) increases rapidly between 15 and 100 nm as particle size increases but is typically less than 1. This is consistent with our previous conclusion that small particles are more abundant at ground level due to the influences of local emissions. \(R_{260 \text{m}/\text{ground}}\) increases continuously and reaches a maximum at \(D_p = \sim 150\) nm. One explanation is the faster condensational and coagulable growth of small particles at 260 m than at ground level. Another explanation is the enhanced regional transport of 100–200 nm particles at high altitude. This is consistent with the fact that much higher \(R_{260 \text{m}/\text{ground}}\) was observed during polluted periods than clean periods. \(R_{260 \text{m}/\text{ground}}\) decreased to less than 1 at \(D_p > 250\) nm, likely due to the deposition of large particles. Our results show that the vertical differences in particle number concentrations varied significantly as a function of size, which has important implications that the health and climate effects of aerosol particles at different heights could be substantially different.

The diurnal cycles of particle number and volume concentrations at 260 m and ground level, as well as \(R_{260 \text{m}/\text{ground}}\) during different periods, are illustrated in Figs. 6 and S7. Pronounced diurnal cycles with two clear peaks at noon and in the evening were observed at both ground level and 260 m. Further analysis highlights that these two peaks were driven by small and large Aitken-mode particles (Fig. 6b and c), likely representing two dominant sources of new particle formation and cooking emissions, respectively. In comparison, the diurnal cycles of accumulation-mode particles were relatively flat, indicating that the sources were mostly regional. Figure 6 shows that the total particle number concentration during the control period was consistently lower than that after the control period, particularly during the time period of 00:00–08:00. Such decreases were mainly caused by accumulation-mode particles that were reduced by 32–67 \% at ground level and 23–69 \% at 260 m throughout the day. In contrast, the diurnal cycle of small Aitken-mode particles was substantially different, characterized by a promi-
Figure 6. The diurnal cycles of particle number concentrations at 260 m and ground level and the ratios of 260 m to ground level for different size ranges, i.e., (a) 15–400 nm ($N_{15–400}$), (b) small Aitken mode ($N_{15–40}$), (c) large Aitken mode ($N_{40–100}$), and (d) accumulation mode ($N_{100–400}$).

The diurnal cycles of particle number concentrations at 260 m and ground level and the ratios of 260 m to ground level for different size ranges, i.e., (a) 15–400 nm ($N_{15–400}$), (b) small Aitken mode ($N_{15–40}$), (c) large Aitken mode ($N_{40–100}$), and (d) accumulation mode ($N_{100–400}$).
Figure 7. Average diurnal evolution of particle number size distributions and aerosol composition at (a, b) 260 m and (c, d) ground level for the new particle growth events. The dashed lines in panels (a) and (c) are the diurnal cycles of CS.

early morning to midnight. Also, aerosol composition had significant changes during the growth periods. As indicated in Fig. 7b and d, the contribution of organics first showed an increase during the early growth period between 08:00 and 12:00, while those of other chemical species remained small. After 12:00, both organics and sulfate showed increased contributions until 17:00. Although the increases in organics and sulfate were partly due to the decreases in nitrate and chloride because of the evaporative loss in the afternoon, our results likely indicate that organics played an important role in the early stage of particle growth, while both organics and sulfate are important in the subsequent growth.

We further calculated the particle GRs for each growth event that lasted more than 3 h (Fig. 8). The particle GR varied from 1.4 to 7.5 nm h$^{-1}$ at 260 m and from 1.5 to 6.1 nm h$^{-1}$ at ground level, which generally falls within the range that was reported previously in various environments, e.g., Beijing (Wu et al., 2007; Zhang et al., 2011), Shangdi-anzi (Shen et al., 2011), Egbert (Pierce et al., 2014), Marseille (Kulmala et al., 2005), and New Delhi (Sarangi et al., 2015). Particle growth rates strongly depend on temperature and the availability of condensable vapors. Indeed, the particle GR in the study generally correlated well with CS at both ground level and 260 m during periods with low sulfate concentrations (Fig. 8d and e). The average particle GR was 3.6 nm h$^{-1}$ at 260 m, which is slightly higher than 3.3 nm h$^{-1}$ at ground level, which is likely due to the lower temperature at high altitude. It is interesting to note that GR was correlated with the change in organic concentration ($\Delta$Org) at 260 m, and also correlated well with $\Delta$Org during periods with low sulfate concentrations (e.g., <3 µg m$^{-3}$) at ground level, likely indicating a dominant role of organics in particle growth. As shown in Fig. 8a, high sulfate concentrations were generally observed during polluted periods with high PM loadings, and correspondingly, relatively higher GR was related to higher sulfate concentration. Our results here suggest that the particle growth mechanisms could be different between clean periods with a dominance of organics and polluted periods with significantly enhanced sulfate.

3.4 Source apportionment

PMF analysis of size-resolved particle number concentrations was able to identify four factors and five factors at 260 m and ground level, respectively (Fig. 9). The five-factor solution at 260 m yielded a split factor that cannot be physically interpreted. The average number size distributions of factor 1 showed GMDs peaking at 20 and 27 nm at ground level and 260 m, respectively, and the temporal variations were characterized by frequent sharp peaks on most days (Fig. 9c). It is clear that this factor was associated with new particle events. This is further supported by the pronounced diurnal cycles showing rapid increases between 08:00 and 12:00, and a dominant source region to the west (Fig. S9a), where clean air masses were prevalent. However, we also noticed the differences in diurnal cycles between ground level and 260 m. For example, the diurnal cycle of factor 1 at the ground site showed two peaks during morning and evening traffic hours, likely indicating the influence of traffic emissions. In fact, the time series correlation between the two heights was weak ($r^2 = 0.17$), confirming that the sources of factor 1 are not the same. The average particle number concentrations of factor 1 were 816 and 1067 cm$^{-3}$ at ground level and 260 m during the control period. These values were 31 and 38% higher than those after the control period. One explanation is due to the increase in CS after the control period, which facilitated the condensation and coagulation of small particles. This result indicates that regional emission
controls could increase the number of nucleation mode particles by reducing PM loadings and decreasing CS. Note that a higher number concentration of factor 1 during the control period was also likely due to the more frequent new particle formation events associated with prevailing northerly winds (Zhao et al., 2017).

Factor 2 presented a size distribution peaking at \( \sim 32 \) nm and a distinct diurnal cycle with two comparable and pronounced peaks at noon and in the evening. The diurnal cycle of factor 2 resembled that of organic aerosol from cooking that was widely reported in Beijing (Huang et al., 2010; Sun et al., 2013; Zhang et al., 2016; Elser et al., 2016; Xu et al., 2015). On average, this factor accounted for 25% of the total particle number concentration and had only a small difference (2%) between the control and non-control periods. This factor was likely dominantly contributed by cooking emissions, although particle growth can partly explain the high concentrations during the late afternoon. Factor 3 at ground level showed a similar diurnal cycle to factor 2, yet the evening peak was much higher than the noon peak. Such a diurnal profile was remarkably similar to that of organic aerosol from cooking that was resolved from PMF analysis of organic aerosol during the same study period (Zhao et al., 2017). Also, the particle number size distribution of factor 3 was similar to that from cooking activities (Bounanno et al., 2011). These results supported the conclusion that factor 3 was mainly from cooking emissions. Similar to factor 2, there was only a small change (3%) during and after the control period, consistent with the fact that no control measures were implemented near our sampling site during the control period. Compared to the ground site, factor 3 at 260 m also showed two pronounced peaks in the diurnal pro-

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Figure 8. (a) Time series of particle growth rates and corresponding average chemical composition for selected particle growth events. Panels (b) and (c) show the correlation of particle growth rates with the changes in the concentration of organics (\( \Delta \text{Org} \)) at 260 m and ground level, respectively. Panels (d) and (e) show the correlation of particle growth rates with condensation sink at 260 m and ground level, respectively. The data points in (b–e) are color coded by the mass concentration of sulfate (\( \text{SO}_4 \)), and those with sulfate concentrations higher than 3 µg m\(^{-3}\) (ground level) and 2.5 µg m\(^{-3}\) (260 m) are marked as triangle points.
file. However, the nighttime peak was much smaller than that at ground level. This can be explained by the significantly enhanced cooking emissions at nighttime at ground level. However, the vertical mixing to high altitude was limited due to the fact that the average number concentration at ground level was 3375 cm$^{-3}$, which was 64% higher than that at 260 m. This indicates stronger influences of local cooking emissions on particle numbers at lower altitudes. This factor moderately correlated between ground level and 260 m ($r^2 = 0.37$), indicating that cooking sources could also be different at different altitudes, for example, more contributions from regional cooking emissions at higher altitudes. In addition, factor 3 at 260 m was better correlated with the sum of factor 2 and factor 3 at ground level ($r^2 = 0.40$, Fig. S8), further supporting that these three factors have similar sources. More evidence is that factors 2 and 3 have the smallest influences from regional emission control among all factors.

Factors 4 and 5 showed quite different temporal variations, but were generally explained by high concentrations during polluted periods. As shown in Fig. 9, the time series of factor 4 was highly correlated between ground level and 260 m ($r^2 = 0.74$), although the peak diameter in size distributions was slightly different (114 and 98 nm, respectively). These results suggest a similar source of factor 4 at different altitudes. The diurnal cycle of factor 4 was also similar at the two different heights, which both showed a small noon peak and high concentrations at night. Such a diurnal cycle was similar to that of less oxidized secondary organic aerosol (SOA) observed during the same study (Zhao et al., 2017). Therefore, we inferred that factor 4 is a secondary factor that was associated with photochemical processing and semi-volatile species. Compared to factor 4, factor 5 showed the best correlation between the two heights ($r^2 = 0.91$), and the time series and diurnal cycles were remarkably similar to those of highly oxidized SOA and sulfate (Zhao et al., 2017), indicating that factor 5 is an aged secondary factor and was mainly formed on a regional scale. Consistently, the bivariate polar plot of factor 5 showed a dominant source region to the south, supporting a major influence of regional transport from the south. Regional emission controls showed large yet different impacts on factor 4 and factor 5. While the average number concentrations of factor 4 showed decreases of 49 and 74% at ground level and 260 m, respectively, during the control period, those of factor 5 had the most reductions of 65 and 74%, respectively. These results are consistent with our previous conclusions that regional emission controls have the most impacts on highly aged secondary aerosols (Sun et al., 2016a; Zhao et al., 2017).

Overall, the five factors, which are associated with new particle events, local primary emissions (e.g., cooking and traffic emissions), and secondary formation with different aging process, represent the major sources of particle numbers in the megacity of Beijing. The contribution of secondary sources was dominant at 260 m throughout the day by varying from ∼50 to 80% (Fig. 10b), and the average contribution (60%) was also higher than that (34%) at ground level. In contrast, the cooking source was the largest contributor to the total particle numbers, on average accounting for 33%. Therefore, our results not only illustrated the similarities and differences of particle number concentrations and sources at different altitudes in the megacity, but also demonstrated the different responses of source factors to regional emission controls.
aerosol composition changes during the growth period, we found that organics appeared to play a more important role than sulfate during the early stage of the growth (09:00–12:00), while organics and sulfate are both important after that. The sources of particle numbers were characterized by PMF, and our results illustrated three common sources at different altitudes, i.e., new particle formation and growth, local secondary formation, and regional transport. We also observed much higher primary emissions from cooking sources at ground level than 260 m, highlighting the importance of local source emissions in the characterization of NPF and growth events at ground level. In addition, we found that regional emission controls exerted a large impact on reducing accumulation-mode particles, for example, by 65–74 % for the regional factor, while regional emission controls had minor impacts on small Aitken-mode particles, mainly due to the enhanced NPF events and the limited controls on local source emissions. These results are overall consistent with the conclusions from our previous studies during the Asia-Pacific Economic Cooperation (Sun et al., 2016a; Xu et al., 2015; Chen et al., 2015).

Data availability. The data in this study are available from the authors upon request (sunyele@mail.iap.ac.cn).

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Competing interests. The authors declare that they have no conflict of interest.

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4 Conclusions

We conducted the first simultaneous real-time measurements of particle number size distribution along with aerosol particle composition at ground level and 260 m on a meteorological tower in urban Beijing from 22 August to 30 September 2015. Our results showed that the number size distributions had significant differences between the two heights, although the particle volume and PM$_4$ mass concentrations were overall similar. The average number concentration (15–400 nm) was 7473 (±4324) cm$^{-3}$ at 260 m, which is 26 % lower than that at ground level (10 134 ± 4680 cm$^{-3}$). The number concentrations of accumulation particles (100–400 nm) at 260 m were highly correlated with those at ground level ($r^2 = 0.85$), indicating their similar sources. However, the correlations were much weaker for Aitken-mode particles, suggesting that they have more different sources at different altitudes. A more detailed analysis suggests that the vertical differences in particle number concentrations varied as functions of size. While particles in the size range of 100–200 nm showed higher concentrations at 260 m, those of smaller particles were more dominant at ground level. These results might indicate the different contributions of local emissions and regional transport to particle numbers at different altitudes. We also observed an increase in the ratio of 260 m to ground level for all particles in different size ranges during daytime, highlighting the impacts of vertical mixing.

Particle growth events were occasionally observed in this study. The average particle growth rate was 3.6 nm h$^{-1}$ at 260 m and 3.2 nm h$^{-1}$ at ground level. By comparing with
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