Highly time-resolved chemical characterization of atmospheric fine particles during 2010 Shanghai World Expo

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Abstract. Shanghai, with a population of over 20 million, is the largest mega-city in China. Rapidly increasing industrial and metropolitan emissions have deteriorated its air quality in the past decades, with fine particle pollution as one of the major issues. However, systematic characterization of atmospheric fine particles with advanced measurement techniques has been very scarce in Shanghai. During 2010 Shanghai World Expo, we deployed a high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS) and a single particle soot photometer (SP2) in urban Shanghai between 15 May and 10 June 2010 to measure fine particles with a high time resolution. The 4-min resolution PM₁ mass concentration ranged from 5.5 to 155 µg m⁻³, with an average of 29.2 µg m⁻³. On average, sulfate and organic matter (OM) were the most abundant PM₁ components, accounting for 33.3 and 28.7 % of the total mass, respectively, while the fraction of nitrate showed an increasing trend with the increasing PM₁ loading, indicating the photochemical nature of high fine particle pollution in Shanghai. Taking advantage of HR-ToF-AMS and SP2, OM was found to have an average OM/OC ratio (organic matter mass/organic carbon mass) of 1.55 and black carbon (BC) had an average number fraction of internally mixed BC of 41.2 %. Positive matrix factorization (PMF) analysis on the high resolution organic mass spectral dataset identified a hydrocarbon-like (HOA), a semi-volatile oxygenated (SV-OOA), and a low-volatility oxygenated (LV-OOA) organic aerosol component, which on average accounted for 24.0, 46.8, and 29.2 % of the total organic mass, respectively. The diurnal patterns of them with interesting time delay possibly implied a photochemical oxidizing process from HOA (and/or its concurrently emitted gaseous organic pollutants) to SV-OOA to LV-OOA. Back trajectory analysis indicated that the northwesterly continental air mass represented the most severe pollutant regional transport condition with the highest nitrate and SV-OOA fractions. In addition, the results in Shanghai were compared with similar measurements performed recently in other mega-cities in the world.

1 Introduction

Shanghai, with a population of over 20 million, is the largest commercial and industrial city in China, as well as a famous mega-city in the world. Its air quality has been deteriorated in the past decades due to rapidly increasing industrial and metropolitan emissions. For example, Ye et al. (2003) reported an annual average PM₂.₅ concentration of about 60 µg m⁻³ in Shanghai in 1999–2000, while this value reached up to about 90 µg m⁻³ in 2005–2006 (Y. Feng et al., 2009). As the adverse health impacts of air pollution increase rapidly with the population size of a city (Parrish and Zhu, 2009) and the air pollution output from mega-cities are now believed to be globally important (Crounse et al., 2009), air pollution characterization in Shanghai metropolitan areas is of great interest.

The 2010 World Expo was hosted in Shanghai during 1 May–31 October 2010, which was another major event in China after the 2008 Beijing Olympic Games. In order to improve the air quality during the World Expo, the Shanghai
government implemented a series of air pollution control measures before the event, including reducing coal burning emissions from power plants, strict regulations on vehicular emission standards, closing high-emitting factories, reducing volatile organic compound (VOC) emissions, etc. The neighboring Jiangsu and Zhejiang provinces also enacted control measures to decrease regional transport of air pollutants to Shanghai. In addition, open field biomass burning was strictly forbidden as a special temporary measure during the World Expo.

Since fine particles play a key role in air pollution in Shanghai (Chan and Yao, 2008), thorough in-situ characterization of the chemical and physical properties of atmospheric fine particles is essential for evaluation of the emission control measures during the 2010 Shanghai World Expo. As part of the project of Air Quality Monitoring Campaign for 2010 Shanghai World Expo supported by Shanghai government, we deployed a high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS) and a single particle soot photometer (SP2) in urban Shanghai to measure chemical and physical characteristics of ambient submicron particles with high time resolution. Previous aerosol studies in Shanghai were mostly based on filter sampling with a coarse time resolution like a day, which cannot match the faster variation of aerosol properties in the atmosphere. This paper summarizes and analyzes the findings based on the advanced on-line measurement techniques during the 2010 Shanghai World Expo, in order to reveal both the basic characteristics of fine particle pollution in this mega-city and some unique features related to the emission control measures during the World Expo.

2 Experimental methods

2.1 Sampling site and period

Shanghai (121.53° E, 31.23° N) is located in the east of the Yangtze River Delta region of China, facing the East China Sea. It belongs to the northern subtropical monsoon climate, with northwestern wind prevailing in the winter while southeastern wind in the summer. The sampling site was located on the roof of a 6-floor building of Shanghai Pudong Environmental Monitoring Station, which was in the eastern part of Shanghai urban areas and about 8 km far to the northeast of the World Expo Park. The surroundings of this site were mainly a residential and business area. The sampling campaign lasted from 15 May to 10 June 2010 after the opening ceremony of the World Expo, which was a transition season from spring to summer in Shanghai with a mean ambient temperature of 21 °C and a mean relative humidity of 66 %. A PM$_{2.5}$ cyclone inlet was set up on the roof to sample outdoor air into the instrument room through a copper tube with a flow rate of 101 min$^{-1}$.

2.2 HR-ToF-AMS operation and data processing

An Aerodyne high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS) was deployed in the instrument room to measure particulate matter, typically referred to as non-refractory PM$_{1}$ (NR-PM$_{1}$, Canagaratna et al., 2007), with a flow rate of 80 ml min$^{-1}$. The detailed instrumental description of HR-ToF-AMS was shown by DeCarlo et al. (2006). During the campaign, the HR-ToF-AMS operated in a cycle of 2 modes every 4 min, including: 2 min V-mode to obtain the mass concentrations of the non-refractory species, in which there was 30 s time for PToF (particle time-of-flight) mode to determine size distributions of species; 2 min W-mode to obtain high resolution mass spectral data. The HR-ToF-AMS was calibrated for inlet flow, ionization efficiency (IE), and particle sizing at the beginning, middle and end of the campaign following the standard protocols (Jayne et al., 2000; Jimenez et al., 2003; Drewnick et al., 2005). The IE calibration used size-selected pure ammonium nitrate particles of 350 and 400 nm and the size calibration used mono-disperse polystyrene latex (PSL) spheres (Duke Scientific, Palo Alto, California, USA) with nominal diameters of 100–700 nm (density = 1.05 g cm$^{-3}$).

Mass concentrations and size distributions of NR-PM$_{1}$ species measured by HR-ToF-AMS were calculated using the methods outlined by DeCarlo et al. (2006). Standard ToF-AMS data analysis software packages (SQUIRREL version 1.51H, and PIKA version 1.10H) downloaded from the ToF-AMS-Resources webpage (http://cires.colorado.edu/jimenez-group/ToFAMSResources/ToFSoftware/index.html) were used to generate unit and high-resolution mass spectra from the V-mode and W-mode data respectively. For mass concentration calculations, an empirical particle collection efficiency (CE) factor of 0.5 was used to account for the incomplete detection of species due to particle bouncing at the vaporizer (Canagaratna et al., 2007; Huang et al., 2010; He et al., 2011). The values of relative ionization efficiency (RIE) used in this campaign were 1.2 for sulfate, 1.1 for nitrate, 1.3 for chloride and 1.4 for organics (Jimenez et al., 2003; Canagaratna et al., 2007). A RIE value of 4.0 was used for ammonium based on the measurement of pure NH$_{4}$NO$_{3}$ particles. Organic elemental analysis was carried out with the methods described previously (Aiken et al., 2007, 2008).

The positive matrix factorization (PMF) analysis (Paatero and Tapper, 1994) was conducted on the high-resolution (HR) mass spectra ($m/z$ 12–150) using the PMF evaluation tool developed by Ulbrich et al. (2009). The data and noise matrices input into the model were generated with the default fragmentation waves by PIKA. The noise values were calculated as the sum of electronic and Poisson ion-counting errors for the relevant high resolution ion fragment (Allan et al., 2003; Ulbrich et al., 2009). The minimum error value for the error matrix was set as the average noise value observed for ions during low signal periods. The ions of H$_{2}$O$^{+}$, HO$^{+}$,
O⁺, and CO⁺ were not included into PMF running since they were determined according to their relationship with CO₂ and thus including them in the PMF analysis can introduce additional weight to CO₂ (Ulbrich et al., 2009). These ions were inserted back into the mass spectra of the resulted PMF components. The weak ions with signal to noise ratios of between 0.2 and 2 were downweighted by a factor of 3 while bad ions with signal to noise ratios of <0.2 were removed from the analysis (Paatero and Hopke, 2003; Ulbrich et al., 2009). More technical details of the PMF analysis can be found in our previous publications (Huang et al., 2010; He et al., 2011).

2.3 SP2 operation and data processing

For a better mass closure measurement of fine particles in the campaign, a single particle soot photometer (SP2) (Droplet Measurement Technologies Inc., Boulder, Colorado) was used for simultaneous measurement of refractory black carbon (BC) mass concentrations, which cannot be detected by HR-ToF-AMS. The sampling flow rate of SP2 was set at 30 ml min⁻¹. The technical details of SP2 have been described elsewhere (Schwarz et al., 2006, 2008b). The calibration of the SP2 was conducted with size-selected fullerene soot (Alpha Aesar, Inc., Ward HIll, MA). The volume-equivalent diameters of individual BC particles can be derived from the measured BC mass by assuming a density of 2 g cm⁻³. More details about the SP2 operation can be found in our previous publication (Huang et al., 2011a). The mixing state of BC was also detected by SP2, with the method of identifying internally mixed BC particles described in Schwarz et al. (2008b).

3 Results and discussion

3.1 Composition, size distribution and mixing state

Figure 1a shows the time series of the PM₁ mass concentration (calculated as the sum of organic matter (OM), SO₄²⁻, NO₃⁻, Cl⁻ and NH₄⁺ measured by AMS and BC measured by SP2) during the entire campaign. The PM₁ mass concentration varied largely from 5.5 to 155 µg m⁻³, with an average of 29.2 µg m⁻³. The time series of particle volume concentration, which was calculated from the number size distribution between 15 and 600 nm measured with a collocated Scanning Mobility Particle Sizer (SMPS, TSI) by assuming spherical particles, was also plotted in Fig. 1a for comparison of the variation trend. The two parameters highly correlated with each other, with a linear correlation coefficient (R²) of 0.90. The time series of the species concentrations were shown in Fig. 1b, with sulfate being the most abundant component (averagely accounting for 33.3 % of the total mass) followed by OM (28.7 %), nitrate (16.3 %), ammonium (13.4 %), BC (6.7 %), and chloride (1.6 %), as shown in Fig. 1d. Compared with other mega-cities in the world, as shown in Fig. 1e, the average PM₁ mass concentration level in Shanghai was lower than those previously measured in the other two Chinese mega-cities, i.e. Beijing in North China (Huang et al., 2010) and Shenzhen in South China (He et al., 2011), similar to in Mexico City (Aiken et al., 2009), while much higher than in New York City (Sun et al., 2011). For simplicity, the citation sources for the data of other mega-cities will not be shown in the following. Due to the limited sampling periods for the campaigns conducted in these cities, this concentration level comparison might not reflect the long-term conditions. As to the composition comparison, the most apparent feature is the higher fractions of sulfate in PM₁ in the three Chinese mega-cities. In Shanghai, the fraction of sulfate even exceeded that of OM. This indicates that high sulfur-containing fuel plays a key role in the formation of high ambient aerosol loadings in China.

In terms of size distribution, the species measured by HR-ToF-AMS showed a similar accumulation mode in mean mass size distributions, peaking at a larger size of ~450 nm in vacuum aerodynamic diameter (Dₐ), as shown in Fig. 1f, indicating large presence of aged regional origins (Allan et al., 2003; Alfarra et al., 2004; Zhang et al., 2005). However, the broader size distribution of OM at smaller sizes (around 200 nm, as a hump), similar to what observed in Beijing and Shenzhen previously, indicates significant existing of fresh combustion organic particles.

Taking advantage of HR-ToF-AMS and SP2, some unique and important properties of carbonaceous aerosols can be measured. The high-resolution organic mass spectrometry allows direct calculation of carbon, hydrogen, oxygen and nitrogen fractions of OM. On average, C, H, O and N contributed 64.7, 9.4, 24.5 and 1.3 %, to the total organic mass, respectively, corresponding to an OM/OC (organic matter mass/organic carbon mass) ratio of 1.55 ± 0.08, which can be used as an empirical factor to convert organic carbon mass to organic matter mass in future filter-based aerosol studies in Shanghai. In addition, the O/C atomic ratio is regarded as a good reference for oxidation state and photochemical age of organic aerosols (Jimenez et al., 2009; Ng et al., 2010). It varied in the range of 0.11–0.57 in the campaign, as shown in Fig. 1c, with a mean value of 0.31, which is similar to those observed in the two Chinese mega-cities, Beijing (0.33) and Shenzhen (0.30), but a little lower than those in Mexico City (0.38) and New York (0.36). This may imply that organic aerosols in Chinese urban environments are more heavily influenced by primary emissions containing low O/C ratio hydrocarbons.

Figure 1c also shows the time series of the number fraction of internally mixed BC (referred to as NIB later) detected by SP2 during the campaign. While freshly emitted BC is generally externally mixed with other aerosol materials, various processes in the ambient atmosphere especially condensation of secondary species would transform it to the internal mixing state. Although the internally mixed BC particles were ever reported to account for ~70 % in biomass.
burning plumes (Schwarz et al., 2008a), the emissions from biomass burning were strictly controlled during the World Expo and found to contribute little to fine particles by source apportionment analysis in Sect. 3.3. Therefore, the NIB can be used to roughly distinguish fresh emissions and aged particles in this study. The NIB was observed to vary in the range of 16.0–71.7 %, with an average of 41.2 %, indicating that the bulk BC was dominated by fresh and thus local combustion emissions during the campaign. This result is similar to those previously observed in another mega-city in China,
Shenzhen, with the average NIB of 40 ~ 46% (Huang et al., 2012). In addition, it is found in Fig. 1 that during the period of 19–27 May, the PM$_1$, BC, and NIB maintained at high levels, indicating that aged air mass and thus regional transport played an important role in the formation of these high pollution days. As both the O/C ratio of OM and NIB of BC can be indicative of photochemical aging of aerosols, they did show some similar variation trends during the campaign, with a correlation coefficient ($R^2$) of 0.21.

Figure 2 shows the variation of relative contributions of different species as a function of the total mass concentration during the campaign, as well as those observed by our group previously in Beijing and Shenzhen. With the PM$_1$ mass loading increasing in Shanghai, the relative contribution of sulfate showed a decreasing trend, the relative contribution of OM maintained at a stable level, while the relative contribution of nitrate showed an increasing trend. The increasing trend of nitrate suggests that active photochemical reactions were a key factor leading to high PM$_1$ pollution in Shanghai. Actually, this increasing trend of nitrate was also observed in Beijing and Shenzhen, as shown in Fig. 2, indicating that high PM$_1$ pollution in China urban environments has a common nature of photochemical smog.

### 3.2 Diurnal variation patterns

Figure 3 presents the mean diurnal variations of different species during the campaign. Sulfate showed little hourly variation, indicating that it was well mixed in the boundary layer and thus had a dominant regional origin. Both nitrate and chloride showed lower concentrations in the afternoon and higher concentrations in the early morning, suggesting that the boundary layer variation and gas-aerosol partitioning were the dominant control mechanisms for their diurnal patterns. The diurnal pattern of ammonium observed was generally the combined result of the variations of (NH$_4$)$_2$SO$_4$/NH$_4$HSO$_4$, NH$_4$NO$_3$ and NH$_4$Cl. Since OM has both large primary and secondary origins and is also influenced by its semi-volatility, the observed diurnal pattern of OM was a complicated result. However, the concurrent increase of both the OM concentration and its O/C ratio in the morning is indicative of the start of secondary organic aerosol (SOA) formation during the daytime. Identification and description of the diurnal patterns of different organic components are discussed in Sect. 3.3. BC showed two broad peaks in the morning and evening and maintained at a high level in the daytime despite of the high boundary layer, clearly reflecting the high vehicular emissions due to the World Expo traffics. The lowest hourly NIB of BC at around 8 a.m. further indicates that fresh vehicular emissions contributed most largely during the morning rush hours.

3.3 Organic source apportionment

The PMF modeling based on the high-resolution mass spectrum observed in the campaign was run for 1 to 8 factors, and it was found that the PMF solutions with factor numbers greater than 3 resulted in splitting of the existing
Fig. 3. Diurnal variation box plots of PM$_1$ species. The upper and lower boundaries of boxes indicate the 75th and 25th percentiles; the line within the box marks the median; the whiskers above and below boxes indicate the 90th and 10th percentiles; and cross symbols represent the means.

The HOA, SV-OOA and LV-OOA components have been frequently identified in previous factor analysis of AMS ambient aerosol datasets and typically attributed to fossil fuel combustion, fresh SOA and aged SOA, respectively (Aiken et al., 2009; Ulbrich et al., 2009; Jimenez et al., 2009; Ng et al., 2010; He et al., 2011). The representativeness of the three organic components identified in Shanghai was examined for three major characteristics: (1) The MS of HOA, SV-OOA and LV-OOA featured ions of C$_n$H$_{2n+1}^+$ and C$_n$H$_{2n-1}^+$, C$_x$H$_y$O$_z^+$ with a lower fraction of CO$_2^+$, C$_x$H$_y$O$_z^+$ with a higher fraction of CO$_2^+$, respectively; The O/C ratios of them well fitted in the typical ranges identified previously for HOA (0.06–0.17), SV-OOA (0.35 ± 0.14) and LV-OOA.
Fig. 4. (a) The MS profiles of the three organic components identified by PMF in this study; (b) time series of the organic components and relevant tracer species; (c) the average organic aerosol composition; (d) the comparison of HOA and OOA concentrations with other mega-cities; and (e) the mean diurnal variations of the organic components.

(0.73±0.14), respectively; The time series of HOA, SV-OOA and LV-OOA correlated well with the corresponding tracers, i.e. BC, nitrate and sulfate, respectively. More specification of this kind of judgment can be found in the literature (Ulbrich et al., 2009; He et al., 2011). On average, the HOA, SV-OOA and LV-OOA accounted for 24.0, 46.8 and 29.2% of the total organic mass, respectively, as shown in Fig. 4c. Both the average concentrations of HOA, as a surrogate of primary organic aerosol, and OOA (SV-OOA + LV-OOA), as a surrogate of SOA, in Shanghai were higher than those in New York but lower than those in Beijing, Shenzhen and Mexico City, as compared in Fig. 4d.

It should be noted that no biomass burning-related organic component was identified by PMF in this campaign, although biomass burning was ever found to be a significant source of fine particles in Shanghai in previous studies (J. Feng et al., 2006; Du et al., 2011) and the PMF analysis of HR-ToF-AMS datasets successfully identified a biomass burning organic aerosol (BBOA) component in many field campaigns (e.g. Aiken et al., 2009; Huang et al., 2011b; He et al., 2011). This was most likely a result of the strict control measures for open field biomass burning events in the rural areas around Shanghai under the air quality assurance act for the World Expo. The little presence of $m/z$ 60, a tracer ion fragment for primary biomass burning aerosols (Alfarra et al., 2007; Aiken et al., 2009), in the MS of each component in Fig. 4a also suggests that BBOA was not abundant enough to be distinguished in this campaign. A similar case of no BBOA...
A significant contribution of BBOA to urban aerosols in China can be effectively reduced by controlling the open field burning events in the surrounding rural areas. However, the changing scale of the morning peak, while the changing scale of the second peak of SV-OOA is much smaller than that of the morning peak and the changing scale of the second peak of LV-OOA is nearly negligible. This may suggest that although HOA and/or its concurrently emitted gaseous organic pollutants were also abundant in the atmosphere in the evening, the SV-OOA and LV-OOA could not be produced significantly without sunlight. The approximate occurring times of the secondary peaks of HOA, SV-OOA and LV-OOA indicate that these peaks were simultaneously caused by the changing of meteorology. Both the decreasing boundary layer and ambient temperature in the evening would favor the accumulation of semi-volatile particulate organic matter.

3.4 Back trajectory analysis

To explore the regional transport influence on PM$_1$ loading and composition in the campaign, the HYbrid Single Particle Lagrangian Integrated Trajectory (HYSPLIT4) model developed by NOAA/ARL (Draxler and Rolph, 2003) was used to produce back trajectories (BT) of air mass arriving in Shanghai. 48-h back trajectories starting at 500 m above the ground level of Shanghai (121.53° E, 31.23° N) were calculated every 6 h in the campaign, and then clustered according to their similarity in spatial distribution using the HYSPLIT4 software (Draxler et al., 2009). The five-cluster solution was found to be the optimum solution according
to the change in total spatial variance, with the mean BT of each cluster presented in Fig. 5. The five clusters were named northeastern marine (NEM) BTs, southeastern marine (SEM) BTs, southerly coastal (SCO) BTs, northwesterly continental (NWC) BTs, and northerly coastal (NCO) BTs, respectively, according to their spatial origins. In result, the NEM BTs were found to be the most frequent cluster, accounting for 51 % of all the BTs.

The average PM$_1$ compositions corresponding to the five BT clusters were then calculated and shown in Fig. 5. The NWC BTs represented the most polluted air mass origin with a mean PM$_1$ mass concentration of 72.0 µg m$^{-3}$, while the SEM BTs represented the cleanest air mass with a mean PM$_1$ mass concentration of 18.2 µg m$^{-3}$. This PM$_1$ loading distribution pattern clearly indicates that the northwestern areas to Shanghai, i.e. China Central Plains, were a much polluted region and the highest PM$_1$ loadings in Shanghai can be caused by pollutant regional transport from there. This finding is well consistent with the finding during the 2008 Beijing Olympic Games that the highest PM$_1$ loadings in Beijing can be caused by southerly pollutant transport from the Central Plains. Therefore, both the Beijing and Shanghai campaigns identified the Central Plains as a critical pollutant source region of China. In terms of composition, the polluted NWC BTs had the highest fractions of OM and nitrate among the five clusters, indicating more severe organic and NO$_x$ pollution to the northwest of Shanghai. As to the organic composition, the three continent-related BT clusters (SCO, NWC and NCO) had much higher SV-OOA fractions than the two marine-related BT clusters (NEM and SEM), which had low levels of background pollutants and were thus more representative of local pollutions of Shanghai. For the polluted NWC BTs, SV-OOA accounted for up to 20.1 % of the total PM$_1$ mass. Based on the above BT analysis, it is clearly seen that the BT type is a vital factor influencing the PM$_1$ loading and composition in Shanghai, and nitrate and SV-OOA were largely from regional transport in high fine particle pollution days of Shanghai.

4 Conclusions

Atmospheric PM$_1$ particles were on-line measured by a HR-ToF-AMS and a SP2 between 15 May and 10 June during 2010 Shanghai World Expo. The high time resolution PM$_1$ mass concentration ranged from 5.5 to 155 µg m$^{-3}$, with an average of 29.2 µg m$^{-3}$. On average, sulfate and OM were the most abundant PM$_1$ components, accounting for 33.3 and 28.7 % of the total mass, respectively, followed by nitrate (16.3 %), ammonium (13.4 %), BC (6.7 %), and chloride (1.6 %). In addition, the contribution of nitrate to total PM$_1$ mass showed an increasing trend with the increasing PM$_1$ loading, implying the photochemical nature of high fine particle pollution in Shanghai. The average size distributions of the species measured by HR-ToF-AMS were all dominated by an accumulation mode at ~450 nm in $D_{oa}$. Calculations of organic elemental composition indicated an average O/C ratio of 0.31 and an average OM/OC ratio of 1.55. The number fraction of internally mixed BC (NIB) was observed to have an average of 41.2 %, indicating that BC was dominated by fresh local emissions there. PMF analysis identified three components, including HOA, SV-OOA and LV-OOA, which on average accounted for 24.0, 46.8 and 29.2 % of the total organic mass, respectively. The average diurnal patterns of the three organic components had a common two-peak shape while with interesting time delay from HOA to SV-OOA to LV-OOA for the daytime peak, possibly implying a photochemical oxidizing process from HOA (and/or its concurrently emitted gaseous organic pollutants) to SV-OOA to LV-OOA. Back trajectory analysis indicated that the northwesterly continental air mass represented the most severe pollutant regional transport condition, with a high PM$_1$ mass concentration of 72.0 µg m$^{-3}$ and large regional transport of nitrate and SV-OOA.

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