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Aerosol composition, sources and processes during wintertime in Beijing, China

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Air pollution is a major environmental concern among all seasons in megacity Beijing, China. Here we present the results from a winter study that was conducted from 21 November 2011 to 20 January 2012 with an Aerodyne Aerosol Chemical Speciation Monitor (ACSM) and various collocated instruments. The non-refractory submicron aerosol (NR-PM₁) species vary dramatically with clean periods and pollution episodes alternating frequently. Compared to summer, wintertime submicron aerosols show much enhanced organics and chloride, which on average account for 52% and 5%, respectively of the total NR-PM₁ mass. All NR-PM₁ species show quite different diurnal behaviors between summer and winter. For example, the wintertime nitrate presents a gradual increase during daytime and correlates well with secondary organic aerosol (OA), indicating a dominant role of photochemical production over gas-particle partitioning. Positive matrix factorization was performed on ACSM OA mass spectra, and identified three primary OA (POA) factors, i.e. hydrocarbon-like OA (HOA), cooking OA (COA), and coal combustion OA (CCOA), and one secondary factor, i.e. oxygenated OA (OOA). The POA dominates OA during wintertime, contributing 69% with

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the rest of 31 % being SOA. Further, all POA components show pronounced diurnal cycles with the highest concentrations occurring at nighttime. CCOA is the largest primary source during the heating season, on average accounting for 33% of OA and 17% of NR-PM₁. CCOA also plays a significant role in chemically-resolved particulate matter (PM) pollution as its mass contribution increases linearly as a function of NR-PM₁ mass loadings. The SOA however presents a reversed trend, which might indicate the limited SOA formation during high PM pollution episodes in winter. The effects of meteorology on PM pollution and aerosol processing were also explored. In particular, the sulfate mass is largely enhanced during periods with high humidity because of fog processing of high concentration of precursor SO2. In addition, the increased traffic-related HOA emission at low temperature is also highlighted. 2078

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Introduction

Atmospheric fine particles exert highly uncertain radiative forcing impacts on climate change (Forster et al., 2007) and are detrimental to human health (Pope III et al., 2002). Aerosol particles also have series impacts on air quality and visibility reduction (Watson, 2002; Molina and Molina, 2004). In recent years, air pollution has become a major environmental concern in densely populated urban areas (Molina et al., 2010; Zhang et al., 2008b) due to large emissions of air pollutants from anthropogenic activities, e.g. traffic, industry, and power plants. However, the air pollution control remains a great challenge because of the very complex sources and evolution processes of aerosol particles. Knowledge of aerosol particle composition, sources and processes is of importance to reduce its harmful effects and improve air quality.

The particulate matter (PM) pollution in Beijing has improvement during the past decade (Chan and Yao, 2008), the visibility however is getting worse (Zhang et al., 2010). While the increasing moisture has played an important role, the consistently high concentration of fine particles (PM_{2.5}) might be another reason. Indeed, fine particles often exceed the new National Ambient Air Quality Standards of China (75 µg m⁻³ for 24-h average) among all seasons. Efforts to characterize sources and composition of fine particles thus have been extensively made in Beijing and vicinity regions. Most of the early studies rely on filter measurements (Sun et al., 2006; He et al., 2001; Yao et al., 2002; Duan et al., 2005). Carbonaceous materials, water-soluble ions, including SO_4^{2-} , NO_3^{-} , and NH_4^{+} , and mineral dust are found to be major constituents of fine particles in Beijing. However, the filter measurements with the sampling durations from hours to days are not sufficient enough to characterize the rapid evolution of atmospheric aerosols, for example, the aging of OA usually takes place in less than one day (Jimenez et al., 2009; DeCarlo et al., 2010). Further, the filter measurements might have significant loss of semi-volatile species, e.g. ammonium nitrate, due to the evaporative loss during the sampling (Dong et al., 2012).

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In recent years, the real-time measurements of aerosol particle composition have been widely conducted in China using various online instruments, e.g. Ambient Ion Monitor (AIM, URG9000B) (Wu and Wang, 2007), Gas and Aerosol Collector - Ion Chromatography (IC) systems (GAC-IC) (Dong et al., 2012), analyzer for Monitoring for AeRosols and Gases (MARGA) (Du et al., 2011), and Aerosol Mass Spectrometers (AMS) (Huang et al., 2010; Sun et al., 2010), etc. Among them, AMS is a unique instrument which can provide both chemical composition and size distributions of nonrefractory submicron aerosol species, including organics, sulfate, nitrate, ammonium, and chloride (Canagaratna et al., 2007). Due to the high sensitivity and the capability in aerosol composition detection, AMS measurements gradually become the focus of recent studies (Huang et al., 2010; Sun et al., 2010; Liu et al., 2011; Takegawa et al., 2009). Despite this, most instruments above need to be maintained routinely in order to obtain high quality data. The long-term measurements, therefore, still remain a challenge. Aerodyne Aerosol Chemical Speciation Monitor (ACSM) was recently developed based on AMS (Ng et al., 2011b). The ACSM is specially designed for routine and long-term measurements of aerosol particle composition and mass concentration, yet no size information. The first deployment of ACSM in Beijing, China shows good performance and provides rich chemical information on aerosol particles (Sun et al., 2012). With all these efforts in aerosol measurements, a number of conclusions and new findings has been obtained, which has significantly improved our understanding of aerosol sources and processes in China. In particular, organics is found to be a major component of fine particles, accounting for more than one-third of total mass. The primary organic aerosol (POA) from traffic, cooking, and coal combustion etc., and secondary OA (SOA) are distinguished and quantified using various receptor models, e.g. chemical mass balance (CMB) and positive matrix factorization (PMF). SOA dominates OA most of time in summer, and is one of the major factors leading to high PM pollution. In addition, the relationship between chemically-resolved PM pollution and meteorology has also been explored to some extent.

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Most previous AMS measurements in Beijing were conducted in summer with a major focus on photochemical processing. However, the aerosol composition, sources and processes may vary very differently between summer and winter because of different meteorology and source emissions. For example, a recent study in the winter 5 of 2010 by a High Resolution Time-of-Flight AMS (HR-ToF-AMS) found that SOA only accounts for 24% of the total OA with the rest being POA (Liu et al., 2011). The oxidation degree of OA is much reduced in winter, and the size distributions shift from a large accumulation mode peaking at ~600 nm to ~450 nm (Sun et al., 2010; Liu et al., 2011; Huang et al., 2010). In order to better characterize aerosol composition and sources during wintertime, an ACSM was deployed from 21 November 2011 to 20 January 2012 at the same site as that in summer (Sun et al., 2012). Here we have a detail characterization of submicron aerosol species including mass concentrations, chemical composition, diurnal variations, and day-of-week patterns during wintertime, and compare with those observed in summer. Then, we perform source apportionment analysis using PMF to investigate the source categories and evolution processes of OA. Finally we provide some insights into the chemically-resolved PM pollution and its association with meteorology during wintertime.

Experimental

Aerosol and gas measurements

The non-refractory submicron (NR-PM₁) aerosol species including organics, sulfate, nitrate, ammonium, and chloride were measured in-situ at the same location as Sun et al. (2012), i.e. Institute of Atmospheric Physics (IAP), Chinese Academy of Sciences located between 3rd and 4th ring road in Beijing, from 21 November 2011 to 20 January 2012 with an ACSM. The aerosol sampling setup and the ACSM operations in this study were also the same as those in Sun et al. (2012). In addition to ACSM deployment, the PM_{2.5} mass was simultaneously measured by a heated Ta**ACPD**

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pered Element Oscillating Microbalance (TEOM series1400a, Thermo Scientific), and collocated gaseous species including CO, SO₂, NO/NO_v, and O₃ were measured by various gas analyzers (Thermo Scientific). The meteorology data (wind speed, wind direction, relative humidity, temperature, pressure, precipitation and solar radiation) dur-5 ing this study was obtained from the IAP tower that is approximately 30 m away and the ground meteorology station that is $\sim 20 \, \text{m}$ away.

2.2 ACSM data analysis

The mass concentrations and mass spectra were processed using ACSM standard data analysis software (v 1.5.1.1). The detailed procedures for the data analysis have been described in Ng et al. (2011b) and Sun et al. (2012). In particular, a collection efficiency (CE) was introduced to compensate for the particle loss, mostly due to particle bounce at the vaporizer (Matthew et al., 2008). Although CE = 0.5 is appropriate for most field studies (Canagaratna et al., 2007), it also varies depending on the acidity of aerosol particles, aerosol composition, and particle phase water (Matthew et al., 2008). Given aerosol particles were dried before sampling into ACSM system and they are overall neutralized in Beijing (Huang et al., 2010; Sun et al., 2010), the particle phase water and acidity are expected to play minor roles in affecting CE values. Therefore, we use a composition-dependent CE recommended by Middlebrook et al. (2011), i.e. CE = max(0.45, 0.0833 + 0.9167 × ANMF) in this study. ANMF is mass fraction of ammonium nitrate in NR-PM₁.

The PMF with PMF2.exe algorithm (Paatero and Tapper, 1994) in robust mode was performed on ACSM mass spectra to deconvolve OA into distinct factors. Similar to our previous study (Sun et al., 2012), we limited our PMF analysis to m/z 120 due to large interferences of internal standard of naphthalene at m/z's 127-129. The PMF results were further evaluated with an Igor Pro-based PMF Evaluation Tool (PET, v2.04) (Ulbrich et al., 2009), and the number of PMF factors were determined following the procedures detailed in Zhang et al. (2011). In summer, PMF analysis only resolved two OA factors, i.e. a hydrocarbon-like OA (HOA) and an oxygenated OA (OOA). In winter,

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PMF analysis showed better deconvolution of OA factors, and was able to identify four components including three primary OA, i.e. HOA, cooking OA (COA), and coal combustion OA (CCOA), and one secondary OA, i.e. OOA.

Results and discussion

Mass concentration

Figure 1 shows the time series of NR-PM₁ and PM_{2.5} mass, submicron aerosol species, and meteorology for the entire study. Similar to the results observed in summer (Sun et al., 2012), the NR-PM₁ mass measured by the ACSM tracks well the PM_{2.5} measured by the TEOM ($r^2 = 0.77$, Fig. S1). The correlation between NR-PM₁ and PM_{2.5} yields a regression slope of 0.74, mainly because ACSM only measures PM₁ and also does not detect refractory materials, e.g. mineral dust and black carbon at vaporizer temperature of ~ 600 °C. Note that the ratio of NR-PM₁/PM_{2.5} is often close to 1 as shown in Fig. S1. One of the reasons is due to the evaporative loss of semi-volatile materials (SVM, e.g. ammonium nitrate and semi-volatile organics) for the TEOM measurements that were operated at 50°C (Eatough et al., 2003). For example, a significant loss ($\sim 50\%$) of SVM for the heated TEOM in comparison to the one with the SVM collected was reported during the SOAR-1 study (Docherty et al., 2011). Indeed, the correlation between NR-PM₁ and PM_{2.5} appears to be dependent on the mass fraction of sulfate and OOA that are less volatile components in the aerosols. Higher fraction of less volatile sulfate and OOA corresponds to lower ratio of NR-PM₁/PM_{2.5}. The higher ratio of NR-PM₁/PM_{2.5} in winter than summer (0.64) is also likely due to the evaporative loss of more semi-volatile materials in winter.

The daily average concentration of the total NR-PM₁ mass ranges from 6.6 to 192 μgm⁻³ with an average value of 66.8(±55) μgm⁻³ for the entire study, which is ~ 30 % higher than that observed in summer, 2011 (Sun et al., 2012). The daily average mass concentration often exceeds the 24-h NAAQS of US EPA of (35 µg m⁻³)

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and the second grade of NAAQS (75 µg m⁻³) released in 2012 by the Ministry of Environmental Protection (MEP) of the People's Republic of China (http://kjs.mep.gov.cn/ hjbhbz/bzwb/dqhjbh/dqhjzlbz/201203/t20120302_224165.htm). If converting the measured NR-PM₁ to PM_{2.5} using a factor of 0.74, we calculate that ~80 % and 54 % of days during wintertime exceed the NAAQS of US EPA and China MEP, respectively. This suggests that the fine particle pollution is still a major environmental concern in Beijing although it has been improved during the past decade (Chan and Yao, 2008). Considering the complexity of sources, properties, and processes of fine particles, the road to achieve the PM_{2.5} standard remains a challenge (Yuan et al., 2012).

The mass concentration of total NR-PM₁ varies very dramatically across the entire study. In particular, the clean events with NR-PM₁ $<\sim 15 \,\mu \mathrm{g}\,\mathrm{m}^{-3}$ ($\sim 24 \,\%$ of the time) and the highly polluted episodes (> 100 μ g m⁻³, ~ 25 % of the time) occur alternatively. The clean periods are often short, typically less than 2 days. In contrast, the high PM pollution events persist a longer time and sometimes over one week. The switch of clean and polluted days is strongly associated with the synoptic meteorology in Beijing. As shown in Fig. 1, the clean periods are associated with high wind speed and dominant westerly and northwesterly winds. Comparatively, the high PM pollution episodes are generally related to southerly and easterly winds with low wind speed (<~ 1.5 ms⁻¹). Such meteorology patterns influencing aerosol species in Beijing have been observed many times in previous studies (Huang et al., 2010; Sun et al., 2010, 2012; Zhang et al., 2012b). Also note wet scavenging plays a minor role in changing aerosol mass loadings given the very small precipitation during the study.

Submicron aerosol composition and diurnal variations

All submicron aerosol species show very dynamic variations day by day with the minimum and maximum of daily average concentrations being larger than 20 times, and even close to an order of 2 (e.g. chloride). In particular, organics shows an obvious sawtooth structure with the variation of mass concentration up to 100 µg m⁻³ in one day.

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PMF analysis of OA in Sect. 3.3 suggests that such structure is mainly caused by the variations of primary OA, e.g. COA and CCOA. Overall, the NR-PM₁ in winter is dominated by organics with an average contribution of 52 % (Fig. 2a). Compared to ~ 40 % observed in summer (Huang et al., 2010; Sun et al., 2010, 2012), organics is more important in PM pollution in winter. Indeed, the average mass concentration of organics is $34.4(\pm 27.9) \mu g m^{-3}$ in winter, which is almost twice higher than that in the summer of 2011 (Sun et al., 2012). The contribution of organics to NR-PM₁ is even higher during the clean periods (63%), which is consistent with our previous observations in Beijing that the air masses from the upwind areas of Beijing are characterized by high mass fraction of organics (Sun et al., 2012). As a comparison, the inorganic species sulfate and nitrate show relatively lower contribution to the total NR-PM1 mass in winter (14% and 16%, respectively) than summer although the mass concentrations between the two seasons are rather similar (Table 1). Chloride accounts for a small fraction of NR-PM₁ in winter (5%) yet with the largest seasonal difference among NR-PM₁ species. The mass concentration in winter is ~ 7 times higher than that in summer, likely due to the intensified emissions from coal combustion during the heating season.

The diurnal profiles of NR-PM₁ species, OA components, and meteorological variables for the entire study and clean periods are presented in Fig. 3. Organics shows a pronounced diurnal cycle with the highest concentration occurring at nighttime $(\sim 54 \,\mu \mathrm{g}\,\mathrm{m}^{-3})$ and the lowest value during daytime $(\sim 24 \,\mu \mathrm{g}\,\mathrm{m}^{-3})$. Our lidar measurements show that the diurnal variation of planetary boundary layer (PBL) height during the study period is relatively flat, indicating that PBL might play a minor role in driving the time-of-day variation during wintertime. Therefore, we infer that the high concentration of organics at nighttime is mainly associated with the primary organics from local emissions, in agreement with the diurnal profiles of primary OA components, e.g. HOA, CCOA, and COA (Sect. 3.3). In addition, the relatively lower wind speed and higher humidity at nighttime might also have played a role. The organics contributes the major fraction of NR-PM₁ throughout the day, ranging from 48-60%, and the contribution increases to 53-75 % during the clean periods. Note that a small peak for organics was

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observed at noon time due to the influences of cooking emissions, but such noon peak is much smaller in comparison to that observed in summer (Sun et al., 2012). The diurnal cycle of organics during the clean periods shows some differences, which is characterized by two peaks corresponding to meal times. Such diurnal pattern resembles more to that observed in summer (Sun et al., 2010, 2012; Huang et al., 2010). This implies that local cooking-related emissions play an important role in organic chemistry during the clean periods. Interestingly, a third peak between 08:00-09:00 was also observed which might be from the emissions of cooking breakfast.

Sulfate shows quite different diurnal behaviors between summer and winter. The sulfate in summer presents an enhanced noon peak associated with the photochemical processing (Sun et al., 2012). However, such noon peak is not observed in winter. This might suggest that the gas-phase photochemical production of sulfate particles is less significant in winter with low temperature and solar radiation. Instead, we found that the sulfur oxidation ratio shows a strong and positive RH dependency (Sun et al., 2013), indicating that agueous processing might have played a more important role in formation of sulfate. This is further supported by the synchronous increase of sulfate and RH from 12:00 to the late afternoon despite the rising boundary layer. In comparison, sulfate shows a very flat pattern during the clean periods demonstrating the regional characteristics of sulfate. Sulfate, overall, accounts for a relatively consistent contribution (11–15%) to the total NR-PM₁ throughout the day.

Nitrate shows the largest different diurnal cycle between summer and winter. As the gas-phase photochemical production and temperature dependent gas-particle partitioning are both important for nitrate formation in summer, the wintertime nitrate appears to be mainly driven by the photochemical production, which shows a linear increase from $\sim 7 \, \mu g \, m^{-3}$ at 08:00 to 12.5 $\mu g \, m^{-3}$ at 17:00 with a growth rate of $\sim 0.6 \,\mu \mathrm{g}\,\mathrm{m}^{-3}\,\mathrm{h}^{-1}$. Further, nitrate shows very similar diurnal pattern and tight correlation with OOA – a surrogate of SOA ($r^2 = 0.80$), confirming the importance of photochemical processing in controlling the diurnal variations of nitrate during wintertime. In addition, the ambient temperature across the entire study is generally low (< 12 °C), and the

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evaporative loss of ammonium nitrate particles due to gas-particle partitioning would not be significant. The results together suggest that the dominant formation mechanisms of nitrate in summer and winter are very different. The nitrate shows a sharp morning peak during the clean periods, which is very similar to that of NO, yet with 5 a time lag of ~ 2 h. Such diurnal pattern was also observed in Mexico City during the MCMA-2003 campaign, which is attributed to photochemical production of nitric acid from NO₂ and OH (Salcedo et al., 2006).

Chloride also shows significantly different diurnal cycles between summer and winter. In summer when the emissions of coal combustion are small, chloride shows a gradual decrease from early morning to late afternoon due to the evaporative loss and dilution effects associated with elevated boundary layer. Comparatively, the diurnal cycle of chloride in winter is guite different with the highest concentration occurring at mid-night and the lowest value at noon. Interestingly, while NH₄Cl and NH₄NO₃ share similar volatile properties and often correlate well with each other (Ulbrich et al., 2009; Sun et al., 2011), the different diurnal cycles of chloride and nitrate in this study indicate their different sources and mechanisms. As the photochemical production is the major process of nitrate formation during wintertime, the coal combustion emissions however are the major sources of chloride. Consistently, the diurnal cycles of chloride and coal combustion OA are very similar, further supporting our conclusion. The chloride during the clean periods is characterized by two pronounced peaks occurring in the early morning and evening, which is likely due to the contribution from local sources.

An even simpler comparison of diurnal variations is day-night differences. In this study, the daytime is defined as the period between sunrise and sunset which is approximately ~ 06:45-~ 17:30, and the rest of day is classified as nighttime. As shown in Table 1, the three primary OA components, i.e. HOA, COA, and CCOA, and chloride show the largest day-night differences with the nighttime concentrations being ~ twice higher than daytime concentrations. The secondary inorganic species (SO₄, NO₃, and NH_A) and OOA, however, did not show very clear day-night differences, yet with ~ 10 – 30 % higher concentration at nighttime. The day-night differences appear to be strongly

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dependent on relative humidity (Fig. 4). As the day-night RH differences are larger than 10%, most aerosol species including the secondary inorganic species often show more than twice differences in mass concentrations, highlighting the role of RH in PM pollution in winter. As mentioned in Sect. 3.2, the PBL plays a less significant role in winter than summer, thus, high concentrations of aerosol species at high RH are likely due to (1) accumulation of pollutants associated with stagnant conditions, (2) production via aqueous processing, and (3) gas-particle partitioning due to water-uptake (Hennigan et al., 2008) and low temperature (Ranjan et al., 2012). It should be noted that even when the day-night RH differences are small, POA components also show much higher concentrations at nighttime than daytime, suggesting the enhanced primary emissions from coal combustion, traffic and cooking activities at nighttime.

3.3 OA composition and sources

PMF analysis of wintertime OA spectra resolved four OA components, i.e. HOA, COA, CCOA, and OOA, of which the mass spectra and time series are presented in Fig. 5, and their diurnal profiles are shown in Fig. 3.

Hydrocarbon-like OA (HOA) 3.3.1

The mass spectral pattern of HOA is characterized by hydrocarbon ion series of m/z 27, 41, and 55, typical of cycloalkanes, and m/z 29, 43, and 57, typical of alkanes (McLafferty and Turecek, 1993). Note that the peak intensities of $C_nH_{2n-1}^+$ is about twice lower than $C_nH_{2n+1}^+$. Such spectral pattern is quite different from the HOA spectrum resolved previously by an HR-ToF-AMS in summer (Huang et al., 2010), and the standard HOA spectrum observed worldwide (Ng et al., 2011a), yet shows much similarity to that identified in winter in Beijing (Liu et al., 2011; Zhang, 2011), and the spectrum of pure diesel fuel aerosols (Canagaratna et al., 2004) and un-reacted diesel exhaust (Kroll et al., 2012). The results suggest that HOA in winter, which is emitted at low ambient temperature and low atmospheric oxidants, might be very fresh, while HOA in summer **ACPD**

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might have been oxidized to some extent already. The mass spectral difference of HOA between summer and winter is also likely due to the significant temperature differences leading to different emissions of hydrocarbons. For example, Lough et al. (2005) found an increase of lighter hydrocarbons in winter in comparison to summer based on onroad vehicle emissions measurements in two tunnels.

The time series of HOA varies dramatically day by day and correlates well with CO $(r^2 = 0.68)$, consistent with a number of previous studies that HOA is mainly associated with combustion-related emissions, e.g. diesel exhaust (Zhang et al., 2005; Canagaratna et al., 2010; Massoli et al., 2012). However, HOA does not present a morning rush hour peak as previous studies at other urban sites (Aiken et al., 2009; Sun et al., 2011), in contrast, HOA presents a pronounced diurnal cycle with the highest concentration at night $(9.4 \,\mu\text{g}\,\text{m}^{-3})$, and the lowest value during late afternoon $(2.9 \,\mu\text{g}\,\text{m}^{-3})$. Such diurnal cycle is remarkably similar to that of the number of heavy duty vehicles (HDV) and heavy duty diesel trucks (HDDT) in Beijing (Han et al., 2009) that are not allowed inside the city between 06:00 a.m.-22:00 p.m. Therefore, the diurnal cycle of HOA is more likely driven by the traffic emissions from HDV and HDDT. HOA, during clean periods, shows two visible peaks during morning and evening rush hours due to local traffic emissions and/or those transported from outside city. Overall, the average mass concentration of HOA for the entire study is 5.8 µg m⁻³ (17% of OA) with the highest contribution to OA at nighttime (22%) and the lowest during daytime (11%). The contribution of HOA to OA is close to 18% observed in summer 2008 (Huang et al., 2010), yet slightly lower than 21% in the winter of 2010 (Liu, 2012).

Cooking OA (COA) 3.3.2

The recent PMF-AMS has identified a cooking-related OA (COA) component with distinct spectral profile and diurnal cycle at various urban sites, e.g. NYC, Beijing, Fresno, Barcelona, and Manchester etc. (Sun et al., 2011; Huang et al., 2010; Ge et al., 2012; Mohr et al., 2012; Allan et al., 2010). The COA is generally characterized by prominent peaks of m/z 55 (mainly $C_3H_3O^+$, $C_4H_7^+$) and 57 (mainly $C_3H_5O^+$, $C_4H_9^+$) with the **ACPD**

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spectral pattern resembling to that of OA from fresh cooking emissions (Mohr et al., 2012; He et al., 2010). While the ratio of m/z 55/57 (= 2.3) is close to the values previously observed in Beijing (Huang et al., 2010; Liu et al., 2011), the COA spectrum in this study shows relatively higher fraction of m/z 44, yet resembles more to that observed in NYC (Sun et al., 2011). The diurnal cycle of COA is characterized by two pronounced peaks corresponding to lunch and dinner times. Note that the lunch peak in winter is less significant in comparison to the evening peak, which might suggest the significantly enhanced cooking activities at nighttime. In addition, the low temperature and high RH, which facilitates gas-particle partitioning of semi-volatile OA, could be another reason for the high concentration of COA at night. Interestingly, the COA during the clean periods shows a third peak in the morning, likely from the contribution of breakfast.

COA has been found to be an important source contribution of OA (~ 16 %-30 %) at various urban sites, and often, COA is comparable and even higher than the trafficrelated HOA. In this study, COA on average accounts for 19% of OA, with the highest contribution of 32 % during dinner time and the lowest of 11 % during late afternoon. The contribution of COA to OA is close to that reported at PKU site in Beijing in the summer of 2008 (Huang et al., 2010) and falls within the range observed at other urban sites. It should be noted that the average contribution of COA to OA increases to 36% during the clean periods, and even goes up to ~50 % at dinner time. These results suggest that COA is an important local source of OA in Beijing, particularly during clean periods.

Coal combustion OA (CCOA) 3.3.3

The wintertime PMF analysis resolved an OA factor showing overall similar spectral pattern to that of standard biomass burning OA (BBOA) (Ng et al., 2011a), yet with much lower fraction of m/z 60 and 73 – AMS marker m/z's for BB (Mohr et al., 2009; Alfarra et al., 2007). Further, such OA factor was only resolved during heating season based on PMF analysis of ACSM OA spectra from one-year measurement (Sun, 2012).

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These results suggest a different OA factor in this study from the common BBOA and/or wood combustion OA, which is very likely an OA factor representing coal combustion sources with significantly enhanced emissions during wintertime. Coal combustion in China has been found to emit a large quantity of carbonaceous aerosols, contributing ₅ ~ 70% of total emitted PM_{2.5} (Zhang et al., 2012a). While the polycyclic aromatic hydrocarbons (PAHs) and n-alkanes are the major components of OA (38% and 20%, respectively) from the coal combustion, levoglucosan only accounts for a minor fraction, up to 1.1% of OC (Zhang et al., 2008a). This is overall consistent with the low fraction of m/z 60 and the hydrocarbon ion series (e.g. $C_nH_{2n-1}^+$ and $C_nH_{2n+1}^+$) observed in this factor. Also, the diurnal profile of f60 (fraction of m/z 60 in OA) during wintertime (Fig. S2) shows a pronounced diurnal cycle with all values (0.52-0.71%) much higher than ~ 0.3 % in the absence of biomass burning (Aiken et al., 2009), confirming the significant impact of combustion source emissions on OA. In contrast, coal combustion shows minor impacts on OA in summer as suggested by the values of f60 close to ~ 0.3% although slightly higher values were observed at night. Considering above, we conclude that this factor is primarily a coal combustion related OA factor (CCOA). Not surprisingly, CCOA correlates tightly with chloride ($r^2 = 0.81$) since the coal combustion is also an important source of chloride (Zhang et al., 2012a).

CCOA shows similar and pronounced diurnal cycle to chloride. The CCOA concentration reaches a maximum of $\sim 18 \,\mu \mathrm{gm}^{-3}$ at mid-night, gradually decreases to \sim 7 $\mu g \, m^{-3}$ at 08:00–09:00, and remains a constant level till 15:00–16:00. Such diurnal cycle is consistent with the source emissions from heating activities which are intensified at nighttime. Indeed, we often can smell coal combustion at nighttime and in the early morning. Similar diurnal cycles of wood burning OA with enhanced emissions at night during wintertime were also observed at other urban sites, e.g. Fresno, CA (Ge et al., 2012) and Paris (Crippa et al., 2012). Likely, the high concentration of CCOA at night is also partially due to the partitioning of semi-volatile water-soluble organic species from gas-phase to particle-phase, which is facilitated by the higher humidity (Hennigan et al., 2009). Indeed, CCOA is the only OA component with the contribution

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to OA increasing as a function of relative humidity (Sun et al., 2012, 2013). The CCOA shows a flat diurnal pattern with very low mass concentrations during clean periods, and the average contribution of CCOA to OA drops to 10%. The high wind speeds and the clean air brought by northwesterly winds are likely the major reasons for the low CCOA during the clean periods.

CCOA is the most important primary source during wintertime, on average accounting for 33 % of OA and 17 % of NR-PM $_1$, which is almost equivalent to the sum of HOA and COA. CCOA also contributes a considerable fraction of OA and NR-PM $_1$ throughout the day, ranging from 26–37 % and 12–22 %, respectively, indicating a sustained impact of coal combustion on PM pollution across the whole day. Therefore, measures to reduce coal combustion emissions would be effective to improve air quality and visibility in Beijing during wintertime. With this, we expect an improvement of PM pollution in the next few years as the progression of boiler renovation from coal combustion to natural gas.

3.3.4 Oxygenated OA (OOA)

Similar to previously reported OOA (Zhang et al., 2005; Ng et al., 2010, 2011a), the mass spectrum of OOA in this study is characterized by the prominent peak of m/z 44 (CO_2^+) (19.4% of the total OOA signal). Although OOA shows more similar spectral pattern to that of highly oxidized LV-OOA than freshly aged SV-OOA, it doesn't necessarily indicate that OOA during wintertime is highly aged. Firstly, the m/z 44 in OOA might be overestimated due to the exclusion of m/z's larger than 120 and some m/z's (e.g. m/z 14, 39, etc.) that are not included in PMF analysis (Ng et al., 2011b). Secondly, OOA shows better correlation with NO₃ (r^2 = 0.80) than SO₄ (r^2 = 0.63), which is similar to the results from a recent HR-ToF-AMS study conducted in Fresno, CA (Ge et al., 2012). Previous studies have shown that NO₃ often correlates well with SV-OOA because of their similar volatilities. Therefore results above suggest that OOA in this study might be less photochemical processed than expected.

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The time series of OOA varies very differently from the three POA components. As POA shows dynamic variations from day to night due to influences from local emissions, OOA often remains a high concentration across several days until a change of air mass occurs. The OOA shows a pronounced diurnal cycle with the concentration being increased from $\sim 9 \, \mu g \, m^{-3}$ at 8:00 to $\sim 15 \, \mu g \, m^{-3}$ at 18:00. The contribution of OOA to the total OA also varies similarly across the day with the highest contribution of ~ 50 % occurring at late afternoon. The gradual increase of OOA during daytime is indicative of photochemical production of SOA. Comparatively, OOA shows relatively stable contributions to OA during clean periods. On average, OOA accounts for 31% of total OA in wintertime with the rest of ~ 70 % being primary OA. Even though OOA accounts for a higher fraction of OA during clean periods (40%), it's still much lower than ~ 60 % observed in summer (Sun et al., 2010, 2012; Huang et al., 2010), which illustrates the dominance of POA in winter and OOA in summer.

The OOA correlates well with secondary inorganic species NO₃ and SO₄ ($r^2 = 0.79$), consistent with the conclusions from previous AMS studies in Beijing (Sun et al., 2010, 2012; Huang et al., 2010). However, the correlations between OOA and secondary inorganic species during wintertime appear to be strongly dependent on relative humidity, in particular for sulfate. As the correlation coefficients remain significant from $r^2 = 0.80$ at RH < 20% to $r^2 = \sim 0.60$ at RH > 50%, the ratio of OOA/SO₄ decreases from 2.9 to < 1, indicating the different roles of photochemical and aqueous processing in formation of OOA and SO₄ during wintertime. Particularly, we observed a more significant enhancement of SO₄ than OOA and NO₃ during the foggy events, (e.g. 4-7 December), which might suggest that fog processing plays a more important role in sulfate production. More details can be found in Sect. 3.6.

Day-of-week variations

The day-of-week patterns for non-refractory aerosol species and OA components are shown in Fig. 6. No pronounced day-of-week patterns for both primary and secondary species, yet higher concentration at Monday and Tuesday and lower one on **ACPD**

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Wednesday were observed. The average weekday and weekend differences are less than 7% for all aerosol species except COA (> 20%). Such weekend effects are quite different from those observed at other urban sites (Lough et al., 2006; Rattigan et al., 2010). For example, Rattigan et al. (2010) observed a significant weekend effects on 5 EC, a tracer for primary emissions from traffic, at the South Bronx, New York due to the reduced traffic activities on weekends. However, HOA in our study didn't show similar behavior to EC in New York. The weak weekly cycles of POA species, e.g. HOA, indicate that the overall human activities appear not being significantly reduced during weekends. For example, while ~ 20 % of private cars are not allowed to drive in the city during weekdays, there are no restrictions during weekends. As a result, the number of cars driving on the road in Beijing during weekdays and weekends might not be significantly different. This is consistent with a previous study in Beijing by Han et al. (2009) that EC shows very similar diurnal profiles between weekdays and weekends. It should be noted that Wednesday shows the lowest concentration among the week. This is also different from a previous study in urban areas in eastern China, where the PM₁₀ presented a gradual accumulation till mid-week, then followed by a reduction with the lowest concentration appearing during weekends (Gong et al., 2007). After checking the day-of-week of meteorology, we found that the lowest concentration on Wednesday is primarily caused by the low RH and high wind speed (Fig. S3). During wintertime, the air masses often alternate rapidly between the south and the northwest, leading to a change between clean periods and pollution episodes within a few hours. Coincidently, the number of clean days that occurred on Wednesday is the highest (~ 6 days). However, the day-of-week variations of aerosol species after excluding the data of clean periods are somewhat different (Fig. S4). The weekdays show slightly higher concentrations than weekends, but the overall day-of-week variations are flat. The results suggest that meteorology is an important factor affecting the day-of-week pattern of aerosol species during wintertime.

The diurnal cycles of aerosol species and OA components between weekdays and weekends are shown in Fig. 6c. All aerosol species show overall similar diurnal cycles

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between weekdays and weekends. It's interesting to note that higher concentrations before 12:00 were observed during weekdays while it's reversed after 16:00. This appears to be consistent with the reduced human activities in the morning on weekends. After excluding the influences of clean episodes, all aerosol species except COA present larger differences between weekdays and weekends before 12:00, yet close concentrations after 16:00 (Fig. S4c). Results here indicate that the weekend effects during wintertime in Beijing mainly occur in the first half day. The minor weekend effects on COA suggest that COA remains an important OA sources throughout the whole week, contributing 17-22 % of OA.

Chemically-resolved PM pollution

Figure 7 shows the variations of mass concentrations and mass fractions of inorganic species and OA components as a function of NR-PM₁ mass loading. All aerosol species except COA and OOA show monotonic and almost linear increase as the increase of PM loading, yet the increasing rates vary for different aerosol species. For example, for an increase of 100 µg m⁻³ NR-PM₁ mass, sulfate presents the largest increasing mass (17.8 µg m⁻³) followed by nitrate (14.4 µg m⁻³), ammonium (12.3 μg m⁻³), and chloride (5.5 μg m⁻³). The OOA increases linearly below $\sim 140 \, \mu g \, m^{-3}$, and then remains relatively stable at higher mass loading, which might suggest the limited formation of secondary OA during periods with high PM pollution in winter. Although the mass concentrations of aerosol species vary significantly as a function of PM loading, the contribution of total OA and inorganic species to the total NR-PM₁ mass didn't show clear PM-mass loading dependency, which indicates that the high PM pollution in winter is a result of synergistic effects of all pollutants. As organics shows the dominant contribution (~50-60%) to PM pollution across different mass loadings, inorganic species, i.e. sulfate, nitrate, and ammonium, contribute ~ 10-15 % each to the total NR-PM₁ mass (Fig. 7b). This mass loading dependent aerosol composition is quite different from that observed in summer, particularly nitrate (Sun et al., 2012). In summer, the nitrate contribution shows a significant increase

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as a function of PM loading, highlighting the important role of nitrate in PM pollution. However, the wintertime nitrate shows a rather stable and even a slight decrease of contribution as PM mass loading increases. The different photochemical production rate, RH and temperature influences on gas-particle partitioning (Ranjan et al., 2012; Hennigan et al., 2009), as well as NH₃ ambient levels (Meng et al., 2011) are all potential reasons for such seasonal differences. The sulfate contribution shows a slight increase from ~ 10 %-19 %, indicating addition production of sulfate mass during high PM episodes. The increase of additional sulfate mass is likely from aqueous production rather than photochemical production given nitrate didn't show a correspondingly increase as sulfate. Indeed, the elevated PM levels in winter are also positively related to RH (Sun et al., 2013), which would facilitate the aqueous processing of SO₂ to form sulfate.

The OA components vary very differently as a function of PM mass loading. Despite the wide concentration range of PM, the contributions of HOA and COA to NR-PM₁ are between $\sim 5-10\%$ and $\sim 10-15\%$, respectively, and the variations are relatively small across the whole mass loadings. However, CCOA and OOA show a dramatic and reversed PM-loading dependency. The contribution of CCOA to NR-PM₁ increases significantly from $\sim 5\%$ to 25% as the NR-PM₁ mass varies from $\sim 10\,\mu g\,m^{-3}$ to more than 200 μ g m⁻³. The contribution of OOA, on the contrary, decreases from $\sim 25 \%$ to 10%. The results suggest that CCOA plays an important role in leading to high PM pollution in wintertime, while the role of OOA is less significant instead. Considering CCOA is the major component emitted from coal combustion, results here confirm that coal combustion is a major source of fine PM during wintertime.

The relationship between SOA and POA, and their roles in the PM pollution in summer in Beijing were reported in Sun et al. (2012). High PM pollution in summer is primarily associated with secondary species. The SOA vs. POA (= HOA + COA + CCOA) shows very different behaviors in winter though SOA shows similar weak correlation with POA ($r^2 = 0.28$) as summer because of their different sources. As shown in Fig. 8a, POA dominates OA most of time, and the HOA contribution is even more **ACPD**

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dominant when the OA mass fraction in NR-PM₁ is higher. For example, when the mass fraction of OA is $>\sim$ 65%, the POA concentration can reach \sim 120 $\mu g \, m^{-3}$, while SOA is generally below 10 $\mu g m^{-3}$, indicating that POA is more than 10 times higher than SOA. Even though in the periods with the highest PM loadings, POA exceeds SOA most of time, yet with a relatively low OA mass fraction, ~ 40-65 %. The diurnal cycles of POA and SOA in Fig. 8b further shows that POA exceeds SOA throughout the day. accounting for 51-76 % of OA. We also note that SOA shows an elevated contribution. from ~ 25 % to 49 % during daytime due to photochemical production. Overall, the average concentration of POA and SOA in winter is 23.7 and 10.7 µg m⁻³, accounting for 69 % and 31 %, respectively, of OA. While the SOA concentration between summer and winter is close, the POA in winter is more than 3 times higher than that in summer (Table 1). Results above demonstrate the different impacts of SOA and POA on PM pollution in different seasons. While SOA is of importance in summer, POA plays a more important role in winter. The roles of total primary and secondary species in PM pollution in winter were also investigated. The diurnal variations of secondary PM $(SPM = SO_4 + NO_3 + NH_4 + OOA)$ and primary PM (PPM = HOA + COA + CCOA)+ Chl) are shown in Fig. 8b. Chloride is classified into PPM because it's mainly from the direct emission of coal combustion in winter. While SPM and PPM both present pronounced diurnal cycles, SPM shows higher mass loading than PPM during daytime (~6:00–20:00), yet comparable to PPM at night. The contribution of SPM to the total NR-PM₁ varies from 50–72 % with the highest contribution occurring at 15:00–16:00. The average contribution of SPM and PPM for the entire study was 59 % and 41 %, respectively. Therefore, results above suggest that SPM is also important in PM pollution during wintertime although POA overwhelms SOA in OA.

3.6 Meteorological effects on PM evolution

The meteorology affects PM evolution through many ways, e.g. accumulation or dilution, gas-particle partitioning, and secondary formation, etc. Here we mainly explored the influences of wind speed, humidity and temperature on PM evolution during

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wintertime. Figure 9 shows the variations of mass concentrations of NR-PM₁ species and OA components as a function of wind speed. All aerosol species show a rapid and almost linear decrease with a reduction rate of ~ 30 % per m s⁻¹ at wind speed $< 3 \,\mathrm{m \, s^{-1}}$. When wind speed is $> 3 \,\mathrm{m \, s^{-1}}$, the total NR-PM₁ mass loading quickly drops from $\sim 100 \, \mu g \, m^{-3}$ to $< 10 \, \mu g \, m^{-3}$, at the same time, all aerosol species show very low concentrations, generally < 2 µg m⁻³. Similar wind speed dependency of EC was also observed in all seasons in Beijing (Han et al., 2009). These results suggest that winds are very efficient in cleaning up the PM pollution during wintertime. To further investigate the dilution effects of wind speed, the relationship between aerosol species and wind speed was separated into two groups with RH < 40% and > 40%. As shown in Fig. 9, the RH-separated relationship varies differently to some extent for the primary and secondary species. The secondary species, e.g. sulfate and OOA, show relatively weaker wind speed dependency in comparison to POA components at high RH. Also note the wind dependency of POA components, e.g. HOA and COA, show smaller differences between low and high RH values than secondary species, elucidating the different impacts of RH on processing aerosol species with different hygroscopicities.

The RH effects on aerosol processing were illustrated by the correlations between NO₃ and SO₄, and the ratio of NO₃/SO₄ as a function of RH (Fig. 10). Nitrate and sulfate correlates well in both summer ($r^2 = 0.61$) and winter ($r^2 = 0.64$), suggesting a common secondary nature of these two species. However, the correlations between nitrate and sulfate are strongly RH dependent, and this dependency is reversed between summer and winter. In summer, the NO₃/SO₄ ratio shows a positive correlation with RH, indicating a faster growth rate of NO₃ than SO₄. As the RH varies from ~ 20 % to > 80 %, the NO_3/SO_4 ratio jumps from ~ 0.5 to ~ 2.0, which is nearly 4 times enhancement. As we reported in our previous study (Sun et al., 2012), the high concentration of nitrate at high RH is likely due to the transformation of HNO₃ into aqueous phase, which could enhance the formation of nitrate particles, especially when there's excess gaseous ammonia in the atmosphere (Meng et al., 2011). As a comparison, the NO₃/SO₄ ratio in winter shows an opposite trend. The highest NO₃/SO₄ ratio of

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~ 2 is observed at RH <~ 20 %, and the ratio decreases linearly to ~ 0.5 when RH is above 80 %. Such differences for NO₃/SO₄ ratio between summer and winter are indications of different formation mechanisms of nitrate and sulfate. As shown in Fig. 10, the RH-binned temperature in winter is generally below 2°C and relatively constant across different RH values. Under these conditions, most of nitrate would be expected in the particle-phase, and the gas-particle partitioning would not play a significant role. Thus daytime photochemical production is the major formation mechanism for nitrate in winter. In contrast, high concentration of sulfate is mainly observed during several fog events with high RH (e.g. 2, 5-7, 13 December, and 7-10 January Fig. 1), indicating that fog processing might be the dominant pathway for the sulfate formation. To support this, we plot the correlations of ΔSO_2 versus ΔSO_4^{2-} in Fig. 11. The ΔSO_2 and ΔSO_4^{2-} is the mass concentrations after subtracting the background concentration of SO2 and $SO_4^{2-},$ which is 7 ppb and 0.33 $\mu g\,m^{-3},$ respectively, in this study. Again, the relationship of ΔSO_2 vs. ΔSO_4^{2-} shows a strong RH dependency. As RH is below 35 %, ΔSO_2 and ΔSO_4^{2-} shows tight correlation ($R^2 > 0.6$), yet the majority of SO_2 remains un-oxidized. As RH increases, the concentration of ΔSO₂ starts to decrease, and the concentration of ΔSO_4^{2-} correspondingly increases. The correlation between ΔSO_2 and ΔSO_4^{2-} is thus sharply reduced, yet with more SO₂ being oxidized to sulfate. This indicates that aqueous-processing, mostly fog processing in this study, has played a significant role in the formation of sulfate.

Figure 12 shows the variations of NR-PM₁ species and OA as a function of temperature. HOA shows similar and monotonically decreasing trend as the increase of temperature at different RH ranges, suggesting a negative influence of temperature on primary traffic emissions. Ranjan et al. (2012) found that the gas-particle partitioning of POA from diesel engines is dependent on temperature, and increasing temperature would decrease the emission factor of POA. Similarly, a study by Nam et al. (2010) also found that decrease of temperature can exponentially increase the PM emissions from light-duty vehicles. The results above are consistent with the temperature dependent POA concentration in this study. Compared to HOA, NO₃ and OOA show an increasing

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trend when temperature increases. This appears to contradict with temperature dependent gas-particle partitioning. However, as shown in Fig. 3, high temperature in one day usually appears in the afternoon when photochemistry is the most intense. Results here indicate that the photochemical production of NO_3 and OOA overcomes gas-particle partitioning under overall low ambient temperatures (< 12 °C) during wintertime. Other aerosol species appear to show different temperature dependent behaviors between low and high RH. The mass concentrations of most aerosol species peak at temperature of $\sim 0-3$ °C when RH is higher than 40 %, comparatively, most of them show a slow increasing trend as the increase of temperature when RH is below 40 %.

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We have a detail characterization of wintertime aerosol particle composition and sources in megacity Beijing, China. The NR-PM $_1$ species including organics, sulfate, nitrate, ammonium, and chloride were measured in-situ from 21 November 2011 to 20 Januar 2012 with an Aerodyne ACSM. The submicron mass varies very dynamically with an average concentration of $67\pm55\,\mu\text{g}\,\text{m}^{-3}$ for the entire study. Similar to summer, organics comprises the major fraction of NR-PM $_1$ in winter. However, the contributions of organics and chloride to NR-PM $_1$ are significantly enhanced during wintertime, which increases from 40 % to 52 % and 1 % to 5 % for organics and chloride, respectively. All NR-PM $_1$ aerosol species show quite different diurnal behaviors between summer and winter due to different source emissions and chemical processing. Of particular, the wintertime nitrate shows a gradual increase from early morning to late afternoon, and also correlates well with SOA formation, indicating that photochemical production dominates the formation mechanisms of nitrate at low ambient temperatures.

PMF analysis was performed on ACSM mass spectra to investigate the sources and processes of OA during wintertime. Four components were identified including three POA factors that are associated with traffic (HOA), cooking (COA), and coal combustion (CCOA) emissions and one secondary factor OOA. All POA components present

pronounced diurnal cycles with the highest concentrations occurring at night. OOA however presents a diurnal cycle as a result of photochemical processing. The POA dominates OA composition during heating season, on average accounting for 69%, and plays a more important role in PM pollution compared to OOA. Among POA components, CCOA is the major primary source, contributing 33 % of OA. CCOA also plays a key role in chemically-resolved PM pollution during wintertime as its contribution increases linearly as a function of to the total NR-PM₁ mass. As a comparison, the OOA concentration remains relatively stable at high PM loadings and its contribution to the total NR-PM₁ decreases as the increase of PM pollution, which might indicate the limited SOA formation during high PM pollution episodes.

Meteorology is also important in PM pollution and aerosol processing in winter. Our results show that winds can clean up the PM pollution efficiently. As the ground wind speed is >~ 3 ms⁻¹, the mass concentrations of all aerosol species quickly drop to <~ 2 μg m⁻³. Note that the cleaning effects of winds appear to be slightly different for primary and secondary species at high RH periods. The relative humidity shows very different impacts on nitrate and sulfate processes between summer and winter. In summer, high humidity leads to a faster growth of nitrate, likely due to the transformation of HNO₃ to ammonium nitrate with access NH₃ in the atmosphere. However, the humidity in winter shows a stronger effect on sulfate formation. In fact, the periods with high humidity in this study are often associated with fog events, therefore, fog processing of high concentration of precursor SO₂ to form sulfate might be the major reason. The temperature dependent aerosol composition highlights the increased PM emissions from vehicles at lower temperatures.

Supplementary material related to this article is available online at: http://www.atmos-chem-phys-discuss.net/13/2077/2013/ acpd-13-2077-2013-supplement.pdf.

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port pathways and air quality over Beijing and its surroundings, Atmos. Chem. Phys., 12,

Table 1. Summary of mass concentrations (μg m⁻³) of NR-PM₁ species and OA components.

	Winter, 2011					Summer, 2011 ^b	W/S ^c
	Entire	Clean ^a	Day	Night	Night/Day	Jul-Aug	
Org	34.4	4.8	25.1	40	1.59	20.0	1.72
HOA	5.8	0.7	3.6	7.1	1.96	7.1	3.34 ^d
COA	6.6	1.7	3.7	8.4	2.30		
CCOA	11.3	0.5	7.2	13.7	1.90		
OOA	10.7	1.9	10.6	10.7	1.01	12.7	0.85
SO_4	9.3	0.9	7.7	10.2	1.33	9.0	1.03
NO_3	10.9	8.0	10.1	11.2	1.10	12.4	0.88
NH_4	8.6	1.0	7.5	9.3	1.24	8.0	1.08
Chl	3.5	0.2	2.5	4.1	1.64	0.5	6.82
Total ^e	66.8	7.6	74.8	52.9	1.42	49.9	1.34

^a Clean periods as marked in Fig. 1.

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^b Sun et al. (2012).

^c Winter/Summer.

^d The ratio is calculated as (HOA + COA + CCOA)_{winter}/HOA_{summer}.

^e Total = Org + SO₄ + NO₃ + NH₄ + Chl.

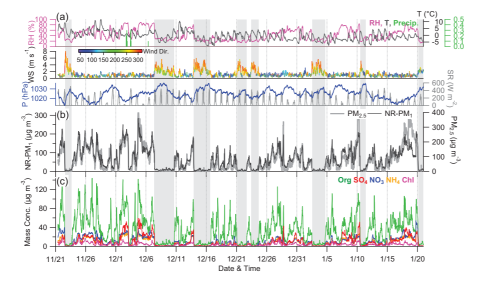


Fig. 1. Time series of **(a)** meteorological variables (relative humidity, temperature, precipitation, wind speed, wind direction, pressure, and solar radiation), **(b)** total NR-PM₁ and PM_{2.5}, and **(c)** NR-PM₁ species (Org, SO₄, NO₃, NH₄, and Chl) for the entire study. The clean periods with low aerosol mass loadings are marked as gray shaded areas.

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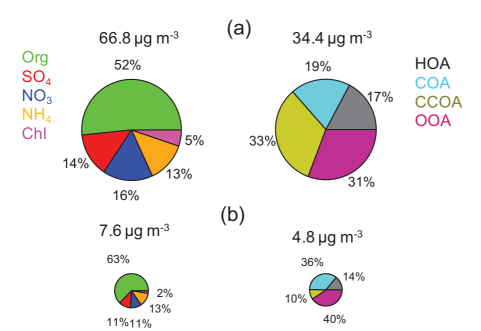


Fig. 2. Average chemical composition of NR-PM₁ and OA for the **(a)** entire study, and **(b)** clean periods marked in Fig. 1.

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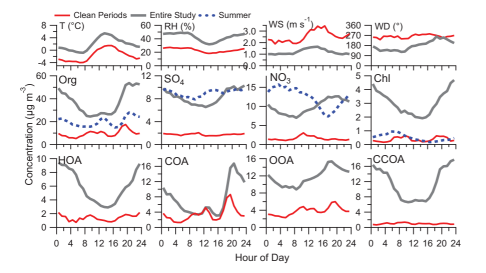


Fig. 3. Diurnal profiles of NR-PM₁ species, OA components, and meteorological variables for the entire study and clean periods marked in Fig. 1. The diurnal profiles of NR-PM₁ species in the summer of 2011 (Sun et al., 2012) are also shown for a comparison. Note, the diurnal profiles of aerosol species during the clean periods were all enhanced by a factor of 2 for clarity.

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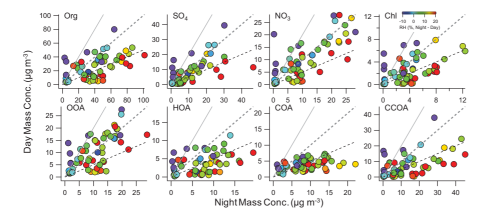


Fig. 4. Day-night comparisons of NR-PM₁ aerosol species and OA components. The data points are color-coded with night-day RH differences. The three lines in the plot refer to 2:1, 1:1, and 1:2 lines, respectively.

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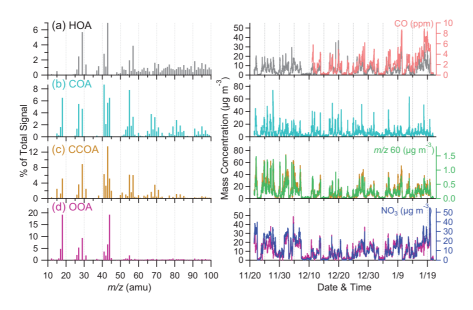


Fig. 5. Mass spectra and time series of mass concentrations of four OA components (HOA, COA, CCOA, and OOA).



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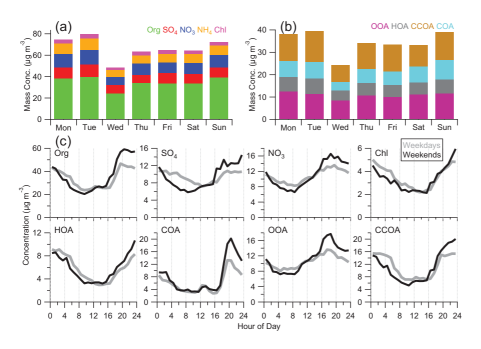


Fig. 6. Day-of-week variations of (a) NR-PM₁ chemical composition and (b) OA composition for the entire study. (c) shows a comparison of the diurnal variations of NR-PM₁ species and OA components between weekdays and weekends.



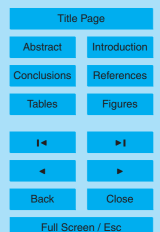
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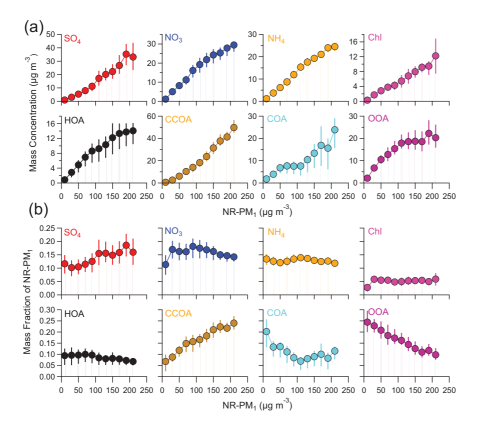


Fig. 7. Variations of (a) mass concentrations and (b) mass fractions of NR-PM₁ species and OA components as a function of NR-PM₁ mass loadings. The data was binned according to the total NR-PM₁ mass, and the vertical lines are 25th (bottom) and 75th percentiles (top).

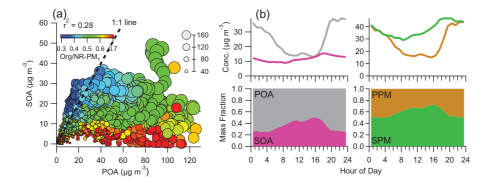


Fig. 8. (a) Relationship between SOA and POA. The data is color coded by the mass fraction of organics in the total NR-PM₁, and the maker size is proportional to the NR-PM₁ mass loading. **(b)** Diurnal variations of mass concentrations and mass fractions for SOA and POA (= HOA + COA + CCOA), and SPM (= $SO_4 + NO_3 + NH_4 + OOA$) and PPM (= POA + ChI).

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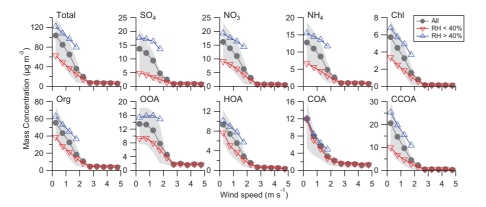


Fig. 9. Variations of NR-PM₁ species and OA components as a function of wind speed. The data was further categorized into two groups with a RH threshold of 40 %. The top and bottom of shaded areas are 25th and 75th percentiles.

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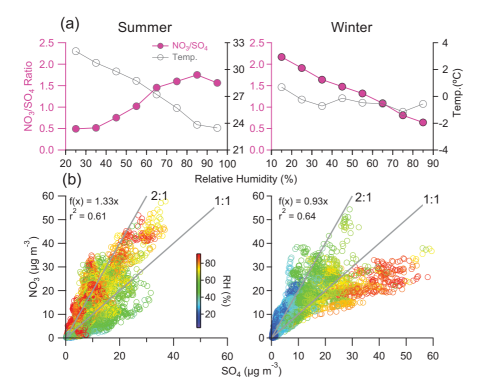


Fig. 10. (a) Variations of the ratio of NO_3/SO_4 and temperature as a function of RH, and **(b)** correlations between NO_3 and SO_4 color-coded with RH in summer and winter.

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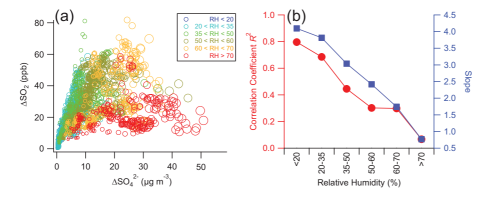


Fig. 11. (a) Scatter plots of ΔSO_2 versus ΔSO_4^{2-} at different RH ranges, and **(b)** shows the linear regression coefficient (R^2) and slope as a function of RH.

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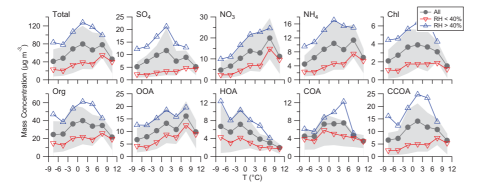


Fig. 12. Variations of NR-PM₁ species and OA components as a function of temperature. The data was further categorized into two groups with a RH threshold of 40 %. The top and bottom of shaded areas are 25th and 75th percentiles.

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