Characterization of Anthropogenic Organic Aerosols by TOF-ACSM with the New Capture Vaporizer

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Abstract. A new capture vaporizer (CV) has been developed and used recently in the Aerodyne aerosol mass spectrometers (AMS) and aerosol chemical speciation monitors (ACSM) instead of the standard vaporizer (SV) to reduce the particle bounce. It is important to characterize the CV performance in different environments. In this study, we characterized specific organic aerosols (OA) from vehicle, cooking, biomass burning, and coal burning emissions by a Time-of-Flight ACSM (TOF-ACSM) with the CV. Their corresponding marker ions that have been defined in the previous SV-based analysis are still valid in the CV mass spectra. Spectra of OA from cooking and vehicle exhaust show similarities in distinct alkyl fragments but different ratios of m/z 55 and 57. Ions related to polycyclic aromatic hydrocarbons are present in the OA spectra obtained from burning lignite and bituminous coal, but not in the spectra obtained from burning anthracite. Although the relative intensities of m/z 60 and 73 are much lower in the CV spectra than in the SV spectra for biomass burning OA, they are still relatively greater compared with the spectra for other sources. Our data suggest an atmospheric background of f_60 of below 0.03% for CV. Moreover, we deployed the CV TOF-ACSM along with a SV AMS in urban Beijing during the winter of 2017 to characterize ambient OA with strong anthropogenic influences. The CV TOF-ACSM shows a collection efficiency (CE) of about unity. The CV and SV data show consistent mass concentrations of sulfate, nitrate, ammonium, and OA. Six OA factors are identified by the positive matrix factorization (PMF) analysis for both the CV and the SV data. The SV and CV PMF factors show good correlations in mass concentrations. The SV and CV factors related to coal combustion and cooking differ significantly in loadings, explained by the PMF uncertainty and the lack of understanding of the relative ionization efficiency (RIE) and CE for primary OA. The CV factors related to secondary sources show greater loadings than the SV factors, which may be associated with the changes of signal-to-noise ratios of various ions in the PMF analysis. Our results support improved mass
quantification and useful source identification by the CV for ambient particles in the polluted urban environment. The difference in factor loadings between SV and CV should be considered when interpreting or comparing the PMF results among studies.

1 Introduction

Aerodyne AMS and ACSM have been widely used for online quantification of non-refractory (NR) aerosol components in the troposphere (Jimenez et al., 2009). These components, e.g., sulfate, nitrate, ammonium, chloride, and OA, are flash vaporized under 600 °C, ionized by electron impact, and then quantified by mass spectrometers having various mass resolution (Jimenez et al., 2003; Decarlo et al., 2006; Ng et al., 2011; Fröhlich et al., 2013). The 2-dimensional mass spectra can be further used in factor analysis to identify OA sources (Zhang et al., 2005; Lanz et al., 2007; Ulbrich et al., 2009). The commonly identified factors include hydrocarbon-like OA (HOA), cooking OA (COA), biomass burning OA (BBOA), coal combustion OA (CCOA), isoprene epoxydiols-derived secondary organic aerosol (IEPOX SOA), nitrogen-enriched OA (NOA), and various types of oxidized OA (OOA) (Jimenez et al., 2009; Zhang et al., 2011; Sun et al., 2013; Chen et al., 2015; Hu et al., 2015; Elser et al., 2016).

The majority of AMS and ACSM uses the SV that has a shape of inverted cone covered by porous tungsten. Particles may bounce on the SV, and hence the AMS and ACSM data are often corrected by CE. The CE depends on the particle composition and the sampling-line relative humidity (RH), and affects the accuracy of the mass quantification of AMS and ACSM (Crosier et al., 2007; Matthew et al., 2008). A composition-based calculation has been widely used to derive CE for the SV instruments (Middlebrook et al., 2012). In recent years, the CV with inner cavity has been developed to reduce the particle bounce (Xu et al., 2017). The CE values of ammonium nitrate (NH₄NO₃), ammonium sulfate ((NH₄)₂SO₄), sodium nitrate (NaNO₃), and ammonium chloride (NH₄Cl) for the CV are much greater than those for the SV. For example, the CE for (NH₄)₂SO₄ is 0.8-0.9 for the CV compared with ~0.4 for the SV (Hu et al., 2017a; Xu et al., 2017). Hu et al. (2017b) showed that ambient particles in forest and suburban regions have a CE of about unity for the CV AMS and quadrupole ACSM, which is twice of the typical CE of 0.5 for the SV instruments (Hu et al., 2017b).

Although the CV improves the mass quantification, the cavity of CV increases the residence time of particles in the vaporizer and leads to prolonged thermal decomposition. The fragmentation of aerosol species in the CV becomes different from that in the SV. For instance, the NO₂⁻/NO⁺ ratio of NH₄NO₃ particles is one order of magnitude lower for CV (i.e., 0.04-0.07) than that for SV (i.e., 0.3-0.7) (Fry et al., 2009; Bruns et al., 2010; Farmer et al., 2010; Xu et al., 2017; Hu et al., 2017a). Similarly, the ratios of SO₄²⁻, HSO₄⁻, and H₂SO₄⁺ to SO⁺ of (NH₄)₂SO₄ particles are much lower for CV (i.e., <0.05) than those for SV (i.e., 0.11-0.38) (Hu et al., 2017a). For OA, the CV-based mass spectra show enhanced fragmentation to small ions. Such enhancement is less pronounced for hydrocarbon-like fragments (CₓHₓ⁺) than for oxygen-containing fragments (CₓHₓOₓ⁺), and seems stronger for laboratory-generated biogenic SOA than for ambient OA (Hu et al., 2018a). Good correlations of the
relative intensities of important tracer fragments (e.g., $f_{43}$, $f_{44}$, $f_{82}$, and $f_{91}$) between the SV and the CV spectra are observed, while poor correlations are shown for fragments having low signal-to-noise ratios in the CV (e.g., $f_{60}$). Besides, prominently elevated CO$^+$ signals were observed for squalene OA in the CV spectra but not in the SV spectra, possibly caused by unknown interactions between the species and the surface with molybdenum oxides (Hu et al., 2018a).

Because of the differences in the fragmentation patterns, it is important to know how well the CV instruments characterize various types of OA in different ambient environments. Hu et al. (2018b) has tested the CV performance in the biogenic dominant and downwind regions. The results suggest reliable mass quantification and sufficient OA source apportionments for HOA, IEPOX-SOA, and OOAs. There is still lack of measurements by the CV instruments in polluted urban areas where anthropogenic OA sources such as CCOA, BBOA, COA, and NOA may contribute substantially. In this study, we investigated the CV mass spectra for specific types of anthropogenic OA produced by vehicle emissions, cooking emissions, biomass burning, coal burning and aromatic photo-oxidation. We deployed both of the CV ToF-ACSM and SV L-ToF-AMS along with other instruments to measure ambient particles in urban Beijing during the winter of 2017. Detailed comparisons for the mass quantification and the source apportionments were made between the CV and the SV data sets.

2 Experimental methods

2.1 Measurements of specific types of OA

The CV-based mass spectra of typical primary OA were obtained by using the TOF-ACSM to sample the air from the exhaust or the atomized air from water solutions of particle filter samples. For the former, the TOF-ACSM was installed in a mobile lab powered by Lithium iron phosphate battery to sample a) the vehicle exhaust in a highway tunnel (G7 highway, 40.299° N, 116.193° E) in Beijing, b) the cooking exhaust near a kitchen ventilator of a large campus cafeteria, and c) the exhaust from a common residential stove during the fierce-combustion stage when burning three types of coals (i.e., anthracite, lignite, and bituminous coal). These measurements were performed during clean days (i.e., PM$_{2.5}$ < 35 µg m$^{-3}$) to minimize the influence of the background air on the particle composition of the exhaust. As shown in Fig. S1 in the Supplement, the OA mass spectra of the three types of exhaust significantly differ from the mass spectra of background air. We did not subtract the spectra of background air from the exhaust spectra because of the lack of information about dilution ratios. Marker ions that are widely used to indicate different types of OA in the analysis of SV spectra have much greater signals in the exhaust spectra than in the spectra of background air. For example, ion peaks at $m/z$ 57 for vehicle, $m/z$ 55 for cooking, and $m/z$ 115, 128, 152, 165, 178, and 189 (i.e., fragmented polycyclic aromatic hydrocarbons (PAHs) ions) for coal combustion, are strongly present, which is in line with previous studies (Canagaratna et al., 2004; Schneider et al., 2006; Mohr et al., 2009; He et al., 2010; Bruns et al., 2015).

For offline measurements, quartz filter samples were collected while burning wheat, rice, and corn straws as well as bituminous coal and anthracite in a combustion chamber (Tian et al., 2015; Tian et al., 2018). A portion of these samples (i.e., about 0.526
cm² in each filter) were sonicated in 10 mL ultrapure water (Milli-Q, 18.2 MΩ cm) for 20 min at 30 °C, filtered through 0.45 µm nylon membrane, nebulized by an Atomizer (TSI, 3076), and dried through a Nafion dryer (Perma Pure, MD-700) before being sampled by the TOF-ACSM. The resulting mass spectra represent the water-soluble portion of BBOA (i.e., >65% of the mass) and CCOA (i.e., <20 % the mass). Moreover, anthropogenic SOA was produced in Aerodyne potential aerosol mass (PAM) oxidation flow reactor by the photo-oxidation of toluene and benzene and were sampled by the CV TOF-ACSM. The initial precursor concentrations were about 40 ppbv for toluene and 100 ppbv for benzene. The SOA mass loadings were about 1.9 to 3.3 µg m⁻³. The OH exposure was about $2.69 \times 10^{10}$ and $1.20 \times 10^{10}$ molecules cm⁻³ s and the NO:HO₂ ratio was about 1.4 and 1.9 for toluene and benzene, respectively, which are estimated by the calculations of PAMchem model on the basis of the calibration on SO₂ (Lambe et al., 2017). Details are provided in Sect. A of the Supplement.

### 2.2 Ambient measurements and data processing

Ambient measurements were carried out at Peking University Urban-Air Environment Monitoring Station (PKUERS) from 14 December 2017 to 21 January 2018. This station consists of three air-conditioned roof sites (one 8-floor and two 6-floor) in a campus building located between the fourth and the fifth north ring roads in Beijing, representing a typical urban environment influenced by anthropogenic emissions (Hu et al., 2016). Particles were detected by the CV TOF-ACSM, the SV AMS, and the scanning mobility particle sizer (SMPS, TSI, 3080), which were placed in the eight-floor site (39.99º N, 116.32º E, 80 m above sea level, and 30 m above the ground). At the same site, the concentrations of volatile organic compounds were measured by an IONICON proton transfer reaction quadrupole interface time-of-flight mass spectrometer (PTR-QiTOF). Detailed descriptions about the PTR-QiTOF measurements are provided elsewhere (Huang et al., 2019). The meteorological parameters including temperature, RH, barometric pressure, wind speed, and wind direction were acquired by a weather station at the same site (Met One Instruments Inc., 083E, 092, 010C, and 020C). Gas pollutants were measured at one of the six-floor site by the Thermo Scientific analyzers including CO (48i-TL), NO-NO₂-NOx (42i-TL), SO₂ (43i-TL), and O₃ (49i-TL). All data refers to the local time (UTC+8).

The SV AMS is LTOF-AMS equipped with regular aerodynamic lens for submicron particles (PM₁). We installed a cyclone (URG, 2000-30ED) in front of the SV AMS that had a sampling flow rate of 3 L min⁻¹ to remove coarse particles (i.e., size-cut at ~2.5 µm). The CV TOF-ACSM is equipped with an aerodynamic lens for measuring particles having aerodynamic diameter less than 2.5 µm (PM₂.5) (Xu et al., 2017). The TOF-ACSM has a different sampling line. For the purpose of comparisons, a PM₁ cyclone (URG, 2000-30EHB) was installed in front of the TOF-ACSM. The sampling flow rate was 16.7 L min⁻¹ in the first half of every hour when the TOF-ACSM samples PM₁, and then 4.8 L min⁻¹ for the other half of every hour to sample PM₂.5. For both instruments, ambient aerosols were sampled through electrolytic polished stainless steel tubes (0.95 cm outer diameter) with the residence time of less than 5 s. The inlet RH is about 12-45%, with only 15% of the values higher than 40%.
The CV TOF-ACSM data have 2-min time resolution and were processed in Tofware (Tofwerk version 2.5.13). A CE of 1 was applied. The SV AMS data have 1-min time resolution and were processed in Igor (WaveMetrics version 6.37) toolkits, Squirrel (version 1.60B) and PIKA (version 1.20B). Composition-dependent CE values for the SV AMS are calculated by the methods introduced by Middlebrook et al. (2012), showing an average CE value of 0.50 ± 0.01 (1σ) and occasionally greater values up to 0.6 (Fig. S2 in the Supplement). We therefore applied a single CE value of 0.5 to the AMS data herein, which is sufficient to achieve reasonable agreements between the AMS and the SMPS data (Fig. S3 in the Supplement). For both instruments, calibrations of ionization efficiency (IE) and relative IE (RIE) followed the standard procedures described in previous studies by using 300-350 nm pure NH$_4$NO$_3$ and (NH$_4$)$_2$SO$_4$ (Canagaratna et al., 2007; Fröhlich et al., 2013). The temperature and pressure during calibration were 293.7 K and 101.82 KPa, which represent the reference conditions of mass concentrations reported herein. Moreover, the PMF analysis was conducted on the organic mass spectra by using the Igor PMF evaluation tool (PET, version 3.00B) (Paatero and Tapper, 1994; Ulbrich et al., 2009). For the TOF-ACSM, the unit-mass-resolution (UMR) data between $m/z$ 20 and 200 are used in the PMF analysis. For the LTOF-AMS, both UMR and high-resolution (HR, a mass resolution of >5000) data are used. The UMR PMF used the spectra between $m/z$ 20 and 200, while the HR PMF analysis used the spectra between $m/z$ 20 and 130, respectively. Details of the PMF analysis are provided in Sect. B of the Supplement.

Tables S1 and S2 in the Supplement list the instrument specifications (e.g., IE, RIE, and CE) and the detection limits of each species respectively. Although our TOF-ACSM is less sensitive than the instrument deployed by Fröhlich et al. (2013) perhaps due to different detectors, the detection is sufficient given the high concentrations of fine particles in Beijing. The LTOF-AMS shows lower detection limits for ammonium and chloride and somewhat greater detection limits for sulfate, nitrate, and OA compared with the V-mode HR-TOF-AMS (Decarlo et al., 2006). Moreover, the sulfate fragmentation for the CV TOF-ACSM is modified based on the measurements of pure (NH$_4$)$_2$SO$_4$ particles (Table S3 and Fig. S4 in the Supplement). The differences of sulfate fragmentation among studies suggest instrument dependency. Each CV instrument needs its own modifications. We applied the RIE values of 1.3 and 1.4 for chloride and OA respectively, both of which are the same as the default values used for the SV instruments (Hu et al., 2017b; Xu et al., 2018). The RIE for nitrate is set to be 1.05, which is slightly lower than the value of 1.1 for the SV instruments (Xu et al., 2018). For sulfate, we applied the RIE value of 1.6 on the basis of the calibration with pure (NH$_4$)$_2$SO$_4$ particles. Previous studies reported different sulfate RIEs for the CV instruments, for example, 1.5±0.3 for a quadrupole-AMS (Xu et al., 2017), 1.7 to 2.4 for three TOF-AMS, and ~1 for a quadrupole-ACSM (Hu et al., 2017b). Such different RIEs suggest that the sulfate detection is sensitive to the vaporizer and the instrument itself. Similarly, different RIE values for sulfate may have been reported for the SV instruments (Budisulistiorini et al., 2014). Moreover, the calibration of pure (NH$_4$)$_2$SO$_4$ and NH$_4$NO$_3$ particles suggest the RIE of 3.7 for ammonium for the CV TOF-ACSM herein, which is lower than the previous finding of 5.2±0.6 (Xu et al., 2017). By contrast, the RIE values for ammonium for the SV instruments are generally 3.5~6.0 (typically an RIE of 4.0 is applied) (Canagaratna et al., 2007). In this study, the good agreement on the
ammonium concentrations between the SV AMS and the CV TOF-ACSM indicate that the RIE values of ammonium are reliable.

3 Results and Discussion

3.1 Mass spectra of anthropogenic OA

Figure 1 shows the mass spectra of five types of anthropogenic OA for CV and SV. The CV mass spectra have much higher signals at m/z 44 and lower relative intensities of many large m/z than the SV spectra have (Fig. S5 in the Supplement), which is consistent with previous findings for pure oxygenated organic compounds, chamber biogenic SOA, and ambient OA in southeast US (Hu et al., 2018a; Hu et al., 2018b). Enhanced thermal decomposition and hot surface collisions in the CV may explain such changes. The marker ions defined in the previous SV-based analysis are still distinct from their adjacent peaks in the CV spectra, suggesting that the CV instruments may work for characterizing anthropogenic OA. For example, the alkyl fragments (i.e., typically CₖH₂ₙ₊₁ and CₖH₂ₙ₋₁ series at m/z 29, 41, 43, 55, 57, 67, 69, 71 etc.) are still prominent in the CV spectra for vehicle exhaust (Fig. 1a), although their relative intensities are about half of their relative intensities in the SV spectra (Figs. 1b and S5a) (Canagaratna et al., 2004; Mohr et al., 2009). The high signal at m/z 44 is probably due to the influence of background air as well as the enhanced thermal decomposition (Fig. S1a).

High signal at m/z 55 has been used as a marker for COA (Mohr et al., 2009; He et al., 2010). As shown in Fig. 1c and Table S4, the f₅₅/f₇₅ ratio of 4.05 is much greater than that for other types of anthropogenic OA (0.86–3.24) and is close to the ratio of 4.76 reported for the CV spectra of oleic acid, a common component of cooking oils (Hu et al., 2018a; Orsavova et al., 2015). The OA spectra of cooking and vehicle exhaust show similarities in distinct alkyl fragments. Meanwhile, the OA spectra for cooking exhaust shows some PAH fragments (Fig. S1b), similar to the spectra for coal burning. This is probably because that the cafeteria often cook meat at high-temperature and lead to the production of PAHs (Phillips, 1999). The f₅₅/f₇₅ ratio in the CV spectra is much greater than the ratios of 2.34 ± 0.59 in the SV spectra (Figs. 1d and S5b), showing a stronger difference between COA and other primary OA in the CV spectra than in the SV spectra.

For coal combustion, three types of coal, including lignite, bituminous coal, and anthracite, were burned in this study (Fig. 1e and Fig. S1c). Alkyl fragments in the spectra for coal burning are less prominent than for vehicle and cooking exhaust. The PAH-related ions clearly present in the spectra for lignite and bituminous coal, which is consistent with the SV spectra for smoky coal burning (Lin et al., 2017). The relative intensities of these ions in the CV spectra are several times lower than those in the SV spectra (Figs. 1f and S5e). Anthracite has the least amounts of impurities. As expected, the signals of PAH-related fragments are insignificant in the spectrum for anthracite (Fig. 2a), which may cause difficulties in separating this source from vehicle exhaust by PMF analysis. PAH characteristics are also absent in the CV spectra for the water-soluble portion of coal-burning OA along with much weaker signals of alkyl fragments compared with the CV spectra for coal-burning exhaust (Fig.
2b). The insoluble portion of the coal-burning OA is perhaps the main contributor of the particle-phase PAH and the alkyl fragments. The CV spectra for the water-soluble portion of the three types of coal-burning OA are similar to each other.

The spectra for BBOA are typically characterized by high signal intensities of ion peaks at \( m/z \) 60 and 73, which are tracer ions for levoglucosan (Weimer et al., 2008; Schneider et al., 2006; He et al., 2010). Figure 1g shows the CV spectra for the water-soluble BBOA which consists of over 65% of the BBOA mass. The relative intensities of the two tracer ions \( f_{60} \) and \( f_{73} \) in the CV spectra are significantly lower than those in the SV spectra (Figs. 1h and S5d). Similar reduction was observed for the CV spectra of laboratory-generated levoglucosan, explained by the enhanced thermal decomposition (Hu et al., 2018a). The values of \( f_{60} \) and \( f_{73} \) depend on the straw type (Figs. 1g and 2c-d). The CV spectrum for rice straw has the greatest \( f_{60} \) (0.48%) and \( f_{73} \) (0.48%), followed by the spectra for corn \( (f_{60}=0.23\%, f_{73}=0.46\%) \) and wheat \( (f_{60}=0.19\%, f_{73}=0.41\%) \). The CV-based \( f_{60} \) values for the other four types of anthropogenic OA are listed in Table S4 in the Supplement. Most of these OAs have lower \( f_{60} \) than BBOA except for benzene SOA. The \( f_{60} \) values of several chamber biogenic SOA are also low \((0.01-0.025\%)\) in the CV spectra (Hu et al., 2017b). Therefore, \( f_{60} \) and \( f_{73} \) may be still used as the markers for BBOA in the CV spectra. However, \( f_{60} \) and \( f_{73} \) may decrease because of aging. Difficulties in separating BBOA from SOA may occur for aged air masses. The atmospheric background of \( f_{60} \) in the SV spectra is about 0.3% (Docherty et al., 2008; Cubison et al., 2011). We expect a much lower threshold in the CV spectra. Another feature in the BBOA spectra is the high \( f_{29} \) values for both CV and SV, which may originate from abundant carbohydrates in biomass (Schneider et al., 2011).

The CV and the SV spectra for aromatic SOA are dominated by the signals at \( m/z \) 29, 43, and 44 (Figs. 1i-j and 2e). The total relative intensities of ions at \( m/z \) > 50 for aromatic SOA are 3-7 times lower than those for primary OA, making aromatic SOA quite distinguishable. The SV spectra for various types of SOA produced by the ozonolysis or photooxidation of biogenic or anthropogenic precursors are similar in terms of the three major peaks and the relative contributions of ions at \( m/z \) > 50 (Kroll et al., 2009; Chhabra et al., 2010; Chen et al., 2015; Robinson et al., 2011; Li et al., 2017). Some SOA has unique features, for example, a distinct peak at \( m/z \) 82 \((C_{8}H_{8}O)\) for IEPOX SOA (Hu et al., 2015). Studies that characterize the laboratory-generated SOA by the CV instruments are however limited. Hu et al. (2018a) reported the CV spectra of the SOA from the oxidation of \( \alpha \)-pinene and \( \Delta \)-carene by nitrate radicals and the ozonolysis of \( \alpha \)-pinene. Our study shows the CV mass spectra of two anthropogenic SOA produced by the photooxidation of benzene and toluene. In comparison with the SV spectra, the CV spectra show much greater signal at \( m/z \) 44, lower signal at \( m/z \) 43, and slightly greater signal at \( m/z \) 29 (Fig. S6). Some ions at \( m/z \) > 50 have greater relative intensities in the CV spectra than in the SV spectra, but more ions have lower relative intensities which may be explained by the enhanced thermal decomposition and hot surface collisions.

3.2 Mass quantification in the polluted urban environment

Figure 3 shows the time series of the mass concentrations of NR-PM\(_1\) and its components from the ambient measurements in Beijing. The campaign-average concentrations of OA, sulfate, nitrate, ammonium, and chloride measured by the CV TOF-ACSM are 11.3±11.3, 2.2±3.6, 4.1±5.9, 2.2±3.0, and 0.56±0.71 µg m\(^{-3}\) (average ± one standard deviation), respectively. The
concentrations of these species agree well with the SV AMS results except for chloride. Because chloride only contributes to about 7% of the NR-PM$_1$ mass, the difference in chloride concentrations would not affect the comparison of the NR-PM$_1$ mass between the CV and the SV measurements significantly. As shown in Figs. 2f and S2, the NR-PM$_1$ mass concentrations measured by the CV and SV instruments are consistent with each other and also agree well with the SMPS data with assumptions of spherical particles and an effective density of 1350 kg m$^{-3}$. The results support the CE of unity and good mass quantifications of ambient aerosols under highly polluted urban conditions by the CV TOF-ACSM.

The regression slope for chloride between the CV and the SV measurements is 0.47, while the $R$ value is 0.98. Hu et al. (2017b) showed similar results (i.e., a slope of 0.52) in Boulder. Artifact chloride signal may present when the SV and CV instruments sample high concentrations of nitrate, explained by the possible reactions between the gaseous nitric acid evaporated from ammonium nitrate and the residual chloride salts on the vaporizer (Hu et al., 2017b; Drewnick et al., 2015). During the measurement period, such chloride artifact should be less than 0.16 µg m$^{-3}$ based on the empirical estimation (i.e., 0.2-0.5% of the nitrate mass) (Hu et al., 2017b). The TOF-ACSM and the AMS herein are relatively new instruments that contains less residuals (e.g., chloride salts). Our data suggest even smaller chloride artifacts (i.e., 0.13 and 0.2% of the nitrate mass for the CV TOF-ACSM and the SV AMS, respectively) (Fig. S7). Besides, the mass quantification can be affected by the product of RIE and CE. Hu et al. (2017a) showed that the CE values of ammonium chloride are 0.25 and 0.35 for the SV and CV AMS, respectively. If the chloride particles are externally mixed as indicated by the different modes in the mass size distributions of species during the non-haze periods of the measurements (Fig. S8), the lower chloride CE for the SV than for the CV instruments suggests a greater correction for the CV than for the SV chloride concentrations. Such correction may lead to a better agreement for chloride. However, information about the mixing state, the exact formula, and the RIE of chloride particles are limited to understand further.

### 3.3 OA source apportionment in the polluted urban environment

We conducted the PMF analysis on the OA spectra from the UMR and HR SV AMS data and the UMR mass spectra from the CV TOF-ACSM data. Six statistical factors were identified and labeled as CCOA, BBOA, COA, HOA, less oxidized- (LO-) OOA, and more oxidized- (MO-) OOA (Fig. 4). The mass spectra and the elemental ratios of the SV PMF factors are similar to the previous findings in winter Beijing (Fig. 4a-f and Table S5). The spectra of HOA, COA, and CCOA factors are dominated by the signals of C$_x$H$_y$ ions and hence show low atomic oxygen-to-carbon ratios (O/C). The CCOA factor is characterized by the rich contribution of alkyl ions and distinctive PAHs fragments (Fig. 4a,g). The BBOA factor shows greater relative intensities of m/z 60 (i.e., mostly C$_3$H$_4$O$_2^+$) and 73 (i.e., mostly C$_3$H$_4$O$_2^+$) compared with other primary factors (Fig. 4b,h). The O/C ratio of the BBOA factor (0.33) is greater than the ratios of freshly emitted BBOA (0.18-0.26) but lower than the ratios of two OOA factors (He et al., 2010), indicating some degree of aging during transport. The atomic nitrogen-to-carbon ratio (N/C) for the BBOA factor is also greater than for other OA factors. Similar high N/C ratios for BBOA have been reported in Beijing and Pearl River Delta regions (Sun et al., 2016; Huang et al., 2011). The COA factor is characterized by the
high f55/f57 ratio among factors (Fig. 4c,i). The f55/f57 ratio of 6.99 in the CV COA factor is greater than the ratio for the cafeteria cooking exhaust (Table S4). The HOA factor is characterized by the predominant contribution of alkyl family (i.e., typically C_{n}H_{2n+1} and C_{n}H_{2n-1}) (Fig. 4d,j). In general, the CV PMF factors show similar spectral features to the corresponding anthropogenic primary OA (Figs. S10 and S11). The average values of f60 and f73 in the CV BBOA factor is 0.084% and 0.079%, respectively, which are 2-6 times lower than in the CV spectra of water-soluble BBOA from crop burning (Figs. 1 and 2). The reduced f60 and f73 are consistent with the greater O/C ratios of the BBOA factor. Because of the reduction of tracer ion signals, the PMF identification of BBOA from the CV data may be more uncertain than from the SV data.

The spectra of LO-OOA and MO-OOA factors show more abundant oxygenated ions and thus greater O/C ratios. The mass spectra of the two OOA factors are dominated by the C_{x}H_{y}O^+ and C_{x}H_{y}O_{2-4} families (Fig. 4e-f). MO-OOA has greater O/C ratios and higher oxidation state than LO-OOA does. The ratios of f44 (CO\(_2\)) to f43 (C\(_2\)H\(_3\)O\(^+\)) are however similar for the two OOA factors in both of the SV and CV data sets, suggesting that pathways other than atmospheric aging perhaps contribute to the higher oxidation state of MO-OOA (Kroll et al., 2011). The greater O/C ratios are mainly contributed by greater f29 (CHO\(^+\)), f30 (CH\(_2\)O\(^+\)), and f31 (CH\(_3\)O\(^+\)) (Fig. S10). The CH\(_{1,3}\)O\(^+\) fragments may come from various molecules, e.g., the SOA produced by the reactive uptake of glyoxal (Ge et al., 2012;Chhabra et al., 2010). Sun et al. (2016) showed that a similar OOA factor from the PMF analysis correlated well with CH\(_2\)O\(^+\) and RH and suggest this factor may be related to aqueous processing.

Further investigations are needed to understand the origin of this factor. The relative intensities of f29, f30, and f31 are greater in the CV MO-OOA spectra than in the CV LO-OOA spectra (Fig. 4k-l), and the difference of f29, f30, and f31 between the two OOA factors are much smaller in the CV PMF results than in the SV PMF results (Fig. S9).

The diurnal patterns of the PMF factors for CV and SV are similar to each other (Fig. S12). The patterns are also consistent with the previous findings in Beijing (Hu et al., 2016;Sun et al., 2013). For example, the CCOA loading shows the strongest nighttime enhancement at 2-4 AM, which is consistent with residential coal burning in the suburban area of Beijing. The COA loading shows peak concentrations at noon and in the evening. The diurnal patterns of HOA are consistent with the morning and evening rush-hour enhancements. Another peak in the earlier morning occurs later than the CCOA nighttime enhancement (4 AM), which may be caused by emissions of heavy-duty diesel trucks that are only allowed to enter the city after 12 AM. Regional transport perhaps predominantly contributes to the BBOA in Beijing and hence the BBOA diurnal patterns show some similarity to CCOA. The diurnal variations of the OOA loadings are different from the primary OA factors. LO-OOA shows elevated mass concentrations in the afternoon (Fig. S13), indicating that the formation of LO-OOA may be photochemically driven (Hu et al., 2016;Sun et al., 2013;Sun et al., 2016). MO-OOA has a relatively flat diurnal pattern (Fig. S12).

Figure 5a-f shows the time series of the mass concentrations of each statistical PMF factor. Figure 6a-f shows the correlation coefficients of the time series of the factor loadings with those of potential tracers. The temporal variation in the CCOA loadings tracked that of the particle-phase PAH concentrations detected by the SV AMS (Figs. 5a and 6a) (Dzepina et al., 2007). The time series of BBOA tracked that of acetonitrile (R\(_{SV}=0.73\)) (Figs. 5b and 6b). The O/C ratio of BBOA suggests...
some degree of aging which may affect the correlation with acetonitrile. Consistently the scatter plot of $f_{44}$ versus $f_{60}$ shows that $f_{60}$ decreases as $f_{44}$ increases (Fig. S13). Our data suggests that the atmospheric background of $f_{60}$ in the CV spectra should be <0.03%. The COA loadings correlate well with the signals of the marker ions of C$_5$H$_3$O$^+$ and C$_6$H$_{10}$O$^+$ proposed by Sun et al. (2011) (Figs. 5c and 6c). The time series of HOA tracked the NO concentrations. Although both BBOA and HOA show good correlations with many tracers, the difference of the two factors in time series is quite clear (e.g., on December 19, 25, 27 and so on). As for OOAs, the LO-OOA loadings only show moderate correlations with the tracers in Fig. 6e, and MO-OOA shows large mass enhancements during the haze episodes (Figs. 5f). The MO-OOA loading correlates well with secondary species like sulfate, nitrate, and ammonium as well as RH (Fig. 6f). As described previously, some studies suggest this factor may be related to aqueous processing but the information is limited for further analysis.

The time series of the mass concentrations of the CV and SV PMF factors tracked each other, although the loadings of some factors differ. As shown in Fig. 5g-l, the correlation coefficients are 0.9 or greater except for BBOA ($R = 0.59$). Additionally, each of the CV PMF factors shows the best correlation with the corresponding SV factors except for BBOA (Fig. 6g-l). The difference in mass resolution between the UMR and the HR PMF analysis may affect the PMF results. Figures S12 and S14 show the comparison of the PMF factors derived from the UMR and HR SV data. The differences in the diurnal variations of the loadings of the UMR-SV vs. HR-SV PMF factors are small (Fig. S12). Good correlations with $R$ values of 0.85 to 0.98 in Fig. S14 indicate that mass resolution is perhaps not the main reason for the difference between CV BBOA and SV BBOA in Fig. 5h. The slopes between UMR and HR PMF results of the SV data range from 0.66 to 1.16, suggesting an uncertainty of about 20% for primary OA factors and about 40% for OOAs related to the mass resolution. As described previously, the average values of $f_{60}$ and $f_{73}$ in the CV BBOA factor are quite lower because of aging. The time series of $f_{60}$ and $f_{73}$ in the CV spectra are much noisier than in the SV spectra (Fig. S15). The low signal-to-noise ratio of the tracer ions may lead to insufficient separation of BBOA from other factors in the PMF analysis of the CV data. Indeed, the CV BBOA factor has weaker temporal correlation with acetonitrile than the SV BBOA factor does ($R_{CV} = 0.48$ vs. $R_{SV} = 0.73$).

The signal-to-noise ratios of the tracer ions for other primary OA factors and the OOA factors are much greater than those of the BBOA factor. Although the correlations are generally good, the regression slopes for the CV vs. SV loadings are about 1.0 for HOA, 0.4 for CCOA, 0.3 for COA, 1.15 for LO-OOA, and 1.42 for MO-OOA (Fig. 5g-l). Some difference in particle transmission efficiency exist between the two instruments, which may contribute to the loading discrepancies between the CV and SV PMF factors. Specially, the CV TOF-ACSM uses the PM$_1$ cyclone in line with PM$_{2.5}$ lens, and the SV AMS has standard PM$_1$ lens. The PM$_{2.5}$ lens have 10-20% lower transmission efficiency for 100-200 nm (vacuum aerodynamic diameter) particles (Xu et al., 2017). The PM$_1$ cyclone has 50% transmission efficiency at 1.0 µm (aerodynamic diameter), whereas the standard PM$_1$ lens have 35-40% transmission efficiency for particles having vacuum aerodynamic diameter of 0.8-1.0 µm. These differences are however small compared to the instrument and PMF uncertainties, which is consistent with the good comparisons of species loadings discussed in Sect. 3.2. Moreover, CE and RIE affect the calculation of the mass loadings. The RIE for COA (SV) may be 1.56 to 3.06, which is greater than the default value of 1.4 (Reyes-Villegas et al., 2018). Another
study showed that the CV and SV instruments may have different RIE for OA having low carbon oxidation state (e.g., less than −1.0) (Xu et al., 2018). There is lack of information about the RIE for CCOA, but the carbon oxidation state of CCOA and COA are both low (−1.26 and −1.56, respectively). A combination of externally mixing, large CE for SV (>0.5), and greater RIE in SV than CV may explain the low slopes for CCOA and COA in Fig. 5g,i. Yet related information is limited for further analysis. Another contributor is the PMF uncertainty. Enhanced thermal decomposition alters the mass spectra (Sect. 3.1). The CV spectra have much more abundant \( f_{44} \) and its related ions. The changes in the mass spectra lead to different weighting of the ions related to various factors in the PMF analysis and hence perhaps affect the PMF analysis of factor loadings. Hu et al. (2018b) also show that the slopes for MO-OOA and LO-OOA (CV vs. SV) are 1.46 and 0.59 respectively for the aircraft study in Korea, 1.20 and 0.67 respectively for the study in Southeast US. The slopes for MO-OOA and LO-OOA in our study are both greater than 1.0 (Fig. 5k-l). The greater OOA concentrations are balanced by the lower loadings of CCOA and COA, and therefore the total OA loadings are consistent with each other between the CV and SV data sets (Fig. 3a).

Figure 7 shows the fractional contribution of the campaign-average concentrations of each statistical PMF factor to the OA mass. The SV PMF results from HR (and UMR) data suggests 12% (11%) of BBOA, 12% (15%) of CCOA, 23% (27%) of COA, 13% (13%) of HOA, 15% (13%) of MO-OOA, and 25% (21%) of LO-OOA, meaning about 60% of primary OA and 40% of OOAs. The results are similar to the previous findings in winter of Beijing except for the low fraction of CCOA. Previous studies indicate 20-30% of CCOA during 2010 to 2013 (Hu et al., 2016; Sun et al., 2013; Sun et al., 2014; Sun et al., 2016). The decrease of the CCOA contribution is consistent with the stringent emission control of residential coal burning during 2016 to 2017. If the RIE of COA is underestimated, the COA loading might be overestimated and the mass fraction of COA should be lower. The CV PMF results suggest 8% of BBOA, 5% of CCOA, 10% of COA, 16% of HOA, 28% of MO-OOA, and 32% of LO-OOA, which means about 40% of primary OA factors and 60% of OOAs. BBOA is not well identified by the CV PMF analysis because of the low signal-to-noise ratio of tracer ions. The reasons for different factor loading levels between CV and SV is unclear. Changes in the mass spectra may contribute to the PMF uncertainties. Figure 7 shows that the bootstrap uncertainties of PMF are lower in the UMR-CV than in the HR-SV analysis. Hu et al. (2018b) showed lower CV bootstrap uncertainty for the study in Southeast US and greater uncertainty in the CV for the Korea aircraft and Billerica studies.

4 Conclusions

In this study, we showed the CV mass spectra of typical anthropogenic OA, including vehicle exhaust, cooking exhaust, coal combustion emission, water soluble OA from crop burning, and aromatic SOA. The marker ions for various sources defined in the previous SV-based analysis are still distinct from adjacent peaks in the CV mass spectra. Cooking and vehicle exhaust show similarities in distinct alkyl fragments but very different \( f_{55}/f_{57} \) ratios in the OA spectra. The PAH-related ions are clearly present in the spectra for lignite and bituminous coal but not for anthracite. Although \( f_{60} \) and \( f_{73} \) are significantly lower in the
CV spectra for water soluble BBOA, they are still greater than most of other OA and work as BBOA tracers. We also examined the performance of CV for mass quantification of non-refractory chemical species and source apportionment of OA in Beijing. The concentrations and temporal variations of OA, sulfate, nitrate, and ammonium derived from the CV data agree well with the SV results. Our results support the CE of about unity and good mass quantifications of ambient particles by the CV TOF-ACSM. Moreover, six statistical factors are identified, including CCOA, BBOA, COA, HOA, and two OOAs (LO-OOA and MO-OOA) by the PMF analysis on both CV and SV data. Our data suggest an atmospheric background of $f_{oo}$ of below 0.03% for CV. Despite the loading difference for some PMF factors, the good correlations of the mass concentrations of the CV and SV factors as well as the similarity between the mass spectra of ambient OA factors and the spectra of specific anthropogenic OA indicate effective PMF analysis with the CV in polluted urban environments like Beijing. Our CV-based PMF results show less fractional contribution of primary OA factors and greater contributions of OOAs than the SV. Such difference may affect the interpretation of results and the comparisons among studies. Further information about the particle mixing state, CE (SV) and RIE of primary OA may help understanding the loading difference of the CV and SV PMF factors.

Data availability. Data presented in this manuscript are available upon request to the corresponding author.

Author contribution. QC, TZ, and YZ designed the study. YZ, XC, KRL, and YWL conducted the measurements. YL, RJH, SYC, and LMZ provided data or samples. QC and YZ analyzed the data. QC and YZ prepared the manuscript with contributions from all authors.

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

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References


Figure 1. Mass spectra of OA produced from various anthropogenic sources. The CV spectra are obtained by the TOF-ACSM in this study. The SV spectra are taken from literature and the AMS spectral database (Ulbrich, I.M., Handschy, A., Lechner, M., and Jimenez, J.L., URL: http://cires.colorado.edu/jimenez-group/AMSsd/*). The intensity of \( m/z \) 28 and 18 are calculated as 100% and 22.5% of the intensity of \( m/z \) 44 for all the spectra (Aiken et al., 2008). In panels c, g, and i, the relative intensity of \( m/z \) 28 and 44 are truncated into half of the original intensities. The error bars in panels d and h show one standard deviation of the relative intensities reported in the reference spectra and only the upper sides of the bars are shown. The inserts in panels e-f show extended \( m/z \) ranges for the corresponding spectra.
Figure 2. The CV mass spectra of anthropogenic OA that are not included in Fig. 1. The intensity of m/z 28 and 18 are calculated as 100% and 22.5% of the intensity of m/z 44 for all the spectra (Aiken et al., 2008). The relative intensity of m/z 28 and 44 in all panels are truncated into half of the original intensities.
Figure 3. Time series (a-f) and scatter plots (g-l) of the mass concentrations of OA, nitrate, sulfate, ammonium, chloride, and the sum of them (i.e., total NR-PM1) measured by the CV TOF-ACSM and the SV AMS. Panel f also shows the mass concentrations of PM1 derived from the SMPS measurements for an assumption of spherical particles with an effective density of 1350 kg m\(^{-3}\). The grey lines in the scatter plots represent the orthogonal distance regressions with intercepts. The slopes and the correlation coefficients (Pearson’s R) are shown in the legend.
Figure 4. Mass spectra of the statistical OA factors, CCOA, BBOA, COA, HOA, LO-OOA, and MO-OOA that are identified by the PMF analysis. The high-resolution intensity bars in panels a-f are color-coded by the contribution of various ion families (i.e., fragments containing C, H, O, or N for subscripts of x, y, and z), as determined from the analysis of the SV AMS data. The elemental ratios of atomic hydrogen-to-carbon (H/C), oxygen-to-carbon (O/C), and nitrogen-to-carbon (N/C) ratios and the organic-mass-to-organic-carbon ratios (OM/OC) of each factor are calculated from the SV AMS data on the basis of the “Improved-Ambient” method (Canagaratna et al., 2015). The UMR bars in panels g-l are calculated from the CV TOF-ACSM data.
Figure 5. Time series (a-f) and scatter plots (g-l) of the mass concentrations of the six OA factors identified by the PMF analysis on the UMR-CV and HR-SV data sets as well as the time series of the potential tracers. The grey lines in the scatter plots represent the orthogonal distance regressions with intercepts. The slopes and the Pearson’s R values are shown in the legend. PAH refers to the particle-phase PAH concentrations detected by the SV AMS.
Figure 6. Pearson’s R values for (a-f) the correlations between the time series of the concentrations of the six OA factors that are identified by the PMF analysis on the CV and SV data sets and potential tracers and (g-l) the correlations between the time series of the concentrations of each OA factor from the CV data and the six OA factors from the SV data. PAH refers to the particle-phase PAH concentrations detected by the SV AMS.
Figure 7. The average fractional contribution to the OA mass by each of the six statistical PMF factors. The uncertainty bars are calculated from the bootstrapping analysis (Table S12).