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Revealing the sources and sinks of negative cluster ions in an urban environment through quantitative analysis

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Abstract. Atmospheric cluster ions are important constituents in the atmosphere, and their concentrations and compositions govern their role in atmospheric chemistry. However, there is currently limited quantitative research on atmospheric ion compositions, sources, and sinks, especially in the urban atmosphere where pollution levels and human populations are intense. In this study, we measured the compositions of negative cluster ions and neutral molecules using an atmospheric pressure interface high-resolution time-of-flight mass spectrometer (APi-TOF) and a chemical ionization mass spectrometer in urban Beijing. Quantitative analysis of cluster ions was performed by their comparison with condensation sink (CS), reagent ions, and neutral molecules. We demonstrate the feasibility of quantifying cluster ions with different compositions using in situ-measured ion mobility distributions from a neutral cluster and air ion spectrometer (NAIS). The median concentration of negative cluster ions was 85 (61–112 for 25 %–75 %) cm⁻³ during the measurement period, which was negatively correlated with CS. The negative cluster ions mainly consisted of inorganic nitrogen-containing ions, inorganic sulfur-containing ions, and organic ions in the form of adducts with NO_3^- or HSO_4^- . The CHON-related organic ions accounted for over 70% of the total organic ions. Although the molecules clustered with NO_3^- and $HSO_4^$ had similar compositions, we found that HSO₄⁻ clustered more efficiently with CHO and CHON_{nonNPs} species (CHON excluding nitrated phenols), while NO_3^- clustered more efficiently with nitrated phenols (CHON_{NPs}). Additionally, most organic ions were positively correlated with neutral molecules, resulting in similar diurnal cycles of organic ions and neutral molecules. However, an exception was found for CHON_{NPs}, the concentration of which is also significantly influenced by the reagent ions NO_3^- . The charge fractions are generally higher for molecules with higher molecular weight and a higher oxidation state, and the opposite diurnal variations in charging fractions between H₂SO₄ and organic species indicate a charging competition between them. Finally, we choose HSO_4^- and $C_3H_3O_4^-$ as representatives to calculate the contribution of different formation and loss pathways. We found their losses are condensational loss onto aerosol particles (73 %–75 %), ion-molecule reaction losses (19%), and ion-ion recombination losses (6%-8%).

1 Introduction

Atmospheric cluster ions are electrically charged atoms, molecules, and molecular clusters. Their mobility is usually larger than $0.5 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ (Hõrrak et al., 2000). The primary ions, such as N_2^+ , O_2^+ , NO^+ , O^- , and O_2^- , are initially formed via cosmic radiation, gamma radiation, and other near-ground local sources, such as radon decay, lightning, plant emissions, waterfall and seashore generation, combustion, and high-voltage transmission line emission (Carslaw et al., 2002; Eisele, 1989; Hirsikko et al., 2007; Tammet et al., 2006). Primary ions will subsequently undergo ion-molecule reactions, ion-ion recombination, or deposition onto particles, accompanied by the evolution of their electrical mobility, concentration, and composition (Curtius et al., 2006; Harrison, 2003; Kontkanen et al., 2013; Shuman et al., 2015). Atmospheric ions play important roles in atmospheric chemistry (Bates, 1982; Luts and Salm, 1994), ion-induced nucleation (Charlson et al., 1992; Lee et al., 2003; Lovejoy, 2004), atmospheric conductivity (Baumgaertner et al., 2013), and air quality (Jiang et al., 2018). Moreover, atmospheric ions can benefit human health at certain levels of concentration, while their low concentrations may cause headaches and insomnia (Chu et al., 2019; Malcolm et al., 2009). The effects of atmospheric cluster ions are highly related to their mobility, concentration, and composition.

There are substantial differences in the characteristics of atmospheric ions and their dynamic variations among diverse environments. Globally, atmospheric ion concentrations span a wide range from 100 to $5000 \,\mathrm{cm}^{-3}$ (Hirsikko et al., 2011; Usoskin et al., 2004), determined mainly by differences in ion production and loss rates. On one hand, ion production rates vary with geographical location, land cover type, and weather conditions (Chen et al., 2016; Ling et al., 2010; Usoskin et al., 2004). On the other hand, ion loss rates are largely related to ions carrying opposite charges, aerosol surface area concentrations, and deposition rates (Harrison, 2003; Tammet et al., 2006). For ion compositions, hundreds of new ions can be formed through ion-molecule reactions after the primary ions are initially formed. It has been found that besides the primary ions, NO_3^- , HSO_4^- , and their clusters were predominant ions due to their high acidities (Davidson et al., 1977; Viggiano et al., 1980). Oxygenated organic molecules (OOMs) were observed clustering with NO₃⁻ and HSO_4^- naturally in the atmosphere, and the composition of these organic atmospheric ions has been reported to be similar to those of neutral vapor molecules (Bianchi et al., 2017; Ehn et al., 2010; Yin et al., 2021). Specifically, ion compositions were found to be very different between new particle formation (NPF) and non-NPF events, and ion compositions during NPF imply different nucleation pathways in clean and polluted atmospheres (Bianchi et al., 2016; Ehn et al., 2010; Eisele et al., 2006; Kirkby et al., 2016; Yin et al., 2021). Thus, the measurement of ion characteristics requires high temporal measurements in various environments.

Ion characteristics at polluted urban sites, where the ion production and loss processes may be very different from the clean sites, are relatively poorly understood. At many polluted urban sites, intense pollution emissions have led to high concentrations of neutral gas species and high aerosol concentrations. For the ion production processes, the neutral gas molecules at urban sites are significantly different from those at clean sites. High NO_x concentrations were found to be accompanied by high concentrations of nitrogen-containing organic molecules at polluted urban sites (Li et al., 2022; Qiao et al., 2021). For ion loss processes, condensation sink (CS) at urban sites can reach values up to 2 orders of magnitude higher compared with clean sites (Cai et al., 2017), which could cause a significant loss of ions onto large particles in urban air. These may lead to big differences in ion concentrations and compositions between clean and urban sites. For instance, Iida et al. (2006) have shown that the ion concentration near a polluted site in Boulder might be substantially suppressed by high ion loss rates, resulting in only a minor contribution from ion-induced nucleation to NPF. Although some recent studies have revealed the characteristics of ions in clean environments (Bianchi et al., 2016; Chen et al., 2016; Ehn et al., 2010), only a few studies have focused on such characteristics in an urban atmosphere (Yin et al., 2021).

Simultaneous analysis of ion concentrations and compositions is rare in field campaigns. Various ion counters have been designed to measure ion concentrations in the last century. Among these, Gerdien counters (Gerdien, 1905), the balance scanning mobility analyzer (BSMA) (Tammet, 2006), the air ion spectrometer (AIS) (Mirme et al., 2007), and a neutral cluster and air ion spectrometer (NAIS) (Manninen et al., 2009, 2016) are the most widely used ion counters or ion mobility spectrometers. However, these electrical mobility-based instruments do not provide composition information of ions. Mass spectrometers have been widely used to measure ion compositions. Quadrupole mass spectrometers with approximate unit mass resolution have been applied in a series of ion composition measurements (Arnold et al., 1978; Arnold and Viggiano, 1982; Eisele, 1989; Eisele et al., 2006; Eisele and Tanner, 1990; Viggiano, 1993). With the development of high-resolution mass spectrometer techniques, such as an atmospheric pressure interface time-of-flight mass spectrometer (APi-TOF, Tofwerk AG), more species of atmospheric ions were identified (Ehn et al., 2010; Eisele et al., 2006; Junninen et al., 2010; Kirkby et al., 2016), but the concentration measured by the APi-TOF remains to be quantified.

Developing a robust and easy-to-use method to quantify concentrations of different ions in APi-TOFs in field measurements is needed. The detection efficiency of an APi-TOF for different ions is mainly determined by the massdependent transmission efficiency. The calibration of transmission efficiency often requires complex laboratory calibration settings, including the generation, classification, and counting of ions with different m/z ratios (Heinritzi et al., 2016; Junninen et al., 2010). Alternatively, the relative transmission efficiency of an APi-TOF compared to primary ions can be obtained using a depletion method with the help of a chemical ionization inlet (Heinritzi et al., 2016). But the settings are still complex, and only relative transmission efficiency is obtained. Ehn et al. (2011) have shown that the ion concentration measured by NAIS and the size-segregated ion signal measured by APi-TOF have good correlations in Hyytiälä. Then by converting the ion mobility into the m/z of the ions (Hõrrak et al., 2000), the in situ calibration of the mass spectrometers can be achieved using the synchronous measurements of well-calibrated ion spectrometers (e.g., NAIS). However, the feasibility of this method still needs to be examined, especially in polluted urban atmospheres.

In this study, we apply an in situ quantification method of atmospheric ion compositions measured by an APi-TOF and reveal the governing factors of atmospheric cluster ion concentration and composition at polluted urban sites. We perform field measurements of atmospheric negative cluster ions at an urban site using the APi-TOF and NAIS simultaneously. By comparing the two in situ measurements, we quantify the concentration of ions measured by the APi-TOF using an improved method. The concentration and composition of ions are compared with the clean sites, and the reasons for the observed differences are explored. The origins of and composition variations in ions are characterized and compared with the neutral clusters measured by a chemical ionization mass spectrometer with NO_3^- as the reagent ions (abbreviated as (nitrate) CI-APi-TOF). The driving factors for the variations in ion characteristics at the urban site are revealed, including CS, reagent ions, and neutral molecules. The charge fractions of various organic species were determined. To better quantify the formation and loss scheme of atmospheric ions, the ions HSO_4^- and $C_3H_3O_4^-$ are taken as representatives in the mass balance equation, and their variations are well captured.

2 Methods

2.1 Field measurement

Field measurements are conducted at the Aerosol and Haze Laboratory of Beijing University of Chemical Technology Station (AHL/BUCT station), Beijing, China (Liu et al., 2020; Yin et al., 2021). The AHL/BUCT station is a typical urban site surrounded by three traffic roads and some residential buildings. The station is ~ 18 m above the ground with no higher buildings around it within 50 m. It is equipped with many state-of-the-art instruments and routine instruments for air quality measurements. Besides, measurements at a boreal forest station SMEAR II (Station for Measuring Ecosystem-Atmospheric Relations) located at Hyytiälä, Finland, were used for comparison. Details of the instruments and their

measurement periods in these two stations are given in Table S1 in the Supplement.

The atmospheric negative cluster ion compositions were measured with an atmospheric pressure interface time-offlight high-resolution mass spectrometer (APi-HTOF) from 14 to 17 February 2018 in Beijing and from 7 April to 8 June 2013 in Hyytiälä (Yan et al., 2018). The ambient air was sampled from the window through 1.4 m long 1/4 in. stainless steel tubing. The sampling flow rate of the APi-HTOF is $\sim 0.8 \, \text{L} \, \text{min}^{-1}$, and an extra sheath flow of $3 L \min^{-1}$ was sampled to minimize the loss of ions in the sampling tubes. Thus the total sampling flow rate was $3.8 \,\mathrm{L\,min^{-1}}$. The APi-HTOF was operated in the negative ion mode to measure negative ions, and its voltages were adjusted to reduce cluster fragmentation. The mass resolution of the APi-HTOF is ~ 4500 at m/z 200. The elements C, H, O, N, and S were used for peak assignment. Peaks that cannot be assigned with these elements within 10 ppm are labeled as "unknown" or "others". The relative mass-dependent transmission efficiency in the HTOF was calibrated through the depletion method when the APi-HTOF was operated with a nitrate chemical ionization (CI) inlet in front, for which the voltages were kept the same as in the APi mode (Heinritzi et al., 2016). Detailed descriptions of the measurement and data analysis have been given in our previous study (Yin et al., 2021). In Hyytiälä, ions were sampled through a core sampling method. The inlet has a thick and long tube outside and a thin and short coaxial tube inside to minimize the sampling losses with a total flow rate of $5 \,\mathrm{L}\,\mathrm{min}^{-1}$. The details of the APi-TOF measurements in Hyytiälä can be found in a previous study (Yan et al., 2018). For further analysis in this study, we quantified the negative cluster ions in both Beijing and Hyytiälä using the in situ quantification method, as will be discussed in Sect. 2.2.

The neutral gas molecules were measured by a (nitrate) chemical ionization atmospheric pressure interface long time-of-flight mass spectrometer (CI-APi-LTOF) (Bertram et al., 2011; Jokinen et al., 2012) from 23 January to 14 April 2018 in urban Beijing. It was operated side by side with the APi-TOF to achieve simultaneous measurements of neutral molecules and cluster ions. NO_3^- and its adducts $(HNO_3)_{1-2}NO_3^-$ were used as the reagent ions to chemically ionize the gas molecules with a laminar flow ionization source mounted in front of the APi-LTOF (Eisele and Tanner, 1993). Nitric acid is volatilized and carried by a total of $20 L \text{ min}^{-1}$ sheath flow; then it is exposed to soft X-rays to produce NO_3^- and its adducts $(HNO_3)_{1-2}NO_3^-$. Mostly the species that can be ionized by NO_3^- and its adducts $(HNO_3)_{1-2}NO_3^-$ are highly oxygenated molecules, such as H₂SO₄ and OOMs. The mass resolution of the (nitrate) CI-APi-LTOF is ~ 8000 at m/z 200. The peak fitting and assignment of the molecular compositions followed our previous studies where binned positive matrix factorization (binPMF) solutions were used as assistants for the peak assignment (Nie et al., 2022). The sensitivity for H_2SO_4 was calibrated

via the controlled reaction between SO₂ and OH at the end of the campaign (Kürten et al., 2012; Li et al., 2019). The relative mass-dependent transmission efficiency of the CI-APi-LTOF was calibrated using the depletion method (Heinritzi et al., 2016). OOMs were quantified assuming they have the same collision-limit charging efficiency as H_2SO_4 with reagent ions, and the mass-dependent transmission efficiency was also used for the quantification of OOMs. The calibration results have been shown in our previous study (Yin et al., 2021).

The ion mobility distribution was measured by a NAIS (Airel Ltd.) (Mirme and Mirme, 2013) from 12 January to 31 December 2018 in urban Beijing and from 1 January to 31 December 2013 in Hyytiälä (Kontkanen et al., 2013). In both stations, the measurement periods of the NAIS covered the measurement periods of the APi-TOF. The NAIS can simultaneously measure positive and negative ions with ion mobility of $3.2-0.0013 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$, corresponding to the mobility diameter range of 0.8-40 nm. In practice, the NAIS switched between detecting naturally charged ions and total particles (including neutral and charged particles) by switching off/on a unipolar corona charger. The measurement cycle period was 2 min for the ion mode, 2 min for the particle mode, and 30s for offset. For the ion measurement, two cylindrical mobility spectrometers were operated in parallel to measure positive and negative ions simultaneously. In this study, we focus on the negative cluster ions measured in the ion mode, whose ion mobility is larger than $0.8 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ and diameter smaller than \sim 1.6 nm. A total of 109 mobility bins in the negative ion mode were measured in urban Beijing, while a total of 28 mobility bins were measured in Hyytiälä. In urban Beijing, ambient air is drawn into the system through a 1.8 m long copper tube with a diameter of 4 cm, positioned at the window. The sample flow rate is maintained at $54 L \min^{-1}$. The inversion kernel of the NAIS was calibrated based on the method described by Wagner et al. (2016), and sampling losses were further considered in the data inversion. The results of the NAIS were calibrated considering both the transfer functions and sampling losses (Mirme and Mirme, 2013; Wagner et al., 2016).

2.2 The quantification of negative cluster ions

Although we have calibrated the relative mass-dependent transmission efficiency in the APi-HTOF, there are still two problems for quantification: (1) the absolute transmission efficiency is unknown; (2) the relative transmission efficiency curve is calibrated by adding a CI inlet in front of the APi-HTOF, and the influences of the CI inlet remain unknown. For example, a previous study using a high-resolution differential mobility analyzer and electrometer to calibrate the transmission efficiencies under the APi mode and CI mode found that the transmission efficiency under the CI mode shifts to larger m/z compared to the APi mode (Heinritzi et al., 2016).

To address these two problems in quantifying the ions in the APi-HTOF, we used an improved method to obtain the absolute transmission efficiency of the APi-HTOF through comparison with the synchronous in situ measurement from the NAIS. The key to linking the measurement results of ions by the APi-HTOF with those by the NAIS is to convert the m/z of ions to ion mobilities. As the resolving power of the APi-HTOF is higher than that of the NAIS, the ion signals in the APi-HTOF in specific m/z ranges were summed up to compare them with the ion concentrations measured by the NAIS.

For some specific ions with known peak assignments and structures, like NO₃⁻ and HSO₄⁻, the ion mobility can be directly measured by ion mobility spectrometers (IMSs). The results reported in previous literature (Jen et al., 2015; Liang et al., 2013; Spangler and Collins, 1975; Stano et al., 2008) are summarized in Table S1 and were applied in this study. For the remaining ions, like the hundreds of complex organic ions, the Stokes–Millikan equation was applied to convert their m/z to mobilities (Ehn et al., 2011; Friedlander, 2000; Tammet, 1995), as in Eq. (1).

$$Z = \frac{1}{\sqrt{1 + m_g/m}} \frac{Q}{3\pi\mu} \frac{1 + Kn \left(1.257 + 0.4e^{-1.1/Kn}\right)}{d_m + d_g}$$
(1)

where Z is the electrical mobility, m_g the mass of a carrier gas molecule, *m* the mass of ions, *Q* the charge of ions, μ the viscosity of the carrier gas, *Kn* the Knudsen number, d_m the mobility diameter of ions, and d_g the diameter of the carrier gas molecule. Here, the carrier gas is ambient air, and m_g , d_g , and μ are adopted as 29.0 g mol⁻¹, 0.3 nm, and 1.73 × 10⁻⁵ kg m⁻¹ s⁻¹ at 278 K and 101.3 kPa (Ku and de la Mora, 2009). *Kn* and d_m are calculated as in Eqs. (2) and (3).

$$Kn = \frac{2\lambda}{d_{\rm m} + d_{\rm g}},\tag{2}$$

$$d_{\rm m} = \sqrt[3]{\frac{6m}{\pi\rho}},\tag{3}$$

where λ is the mean free path of carrier gas (62 nm at 278 K and 101.3 kPa) and ρ the density of ions. The appropriate density for calculating the mobility of ions is controversial. Ehn et al. (2011) used the bulk density of aconitic acid $(1.66 \,\mathrm{g}\,\mathrm{cm}^{-3})$ for the conversion, but molecular clusters may have different densities with bulk substances. To address this problem, we compared the measured mobility of an IMS for different types of OOMs with the calculated mobilities according to the Stokes-Millikan method using 1.1, 1.3, and $1.6 \,\mathrm{g}\,\mathrm{cm}^{-3}$, respectively, as the densities (Fig. S1 in the Supplement). The experimental results of the IMS are from Krechmer et al. (2016). By comparing the measured and calculated mobilities for organic ions with different compositions, densities of 1.1 and $1.3 \,\mathrm{g \, cm^{-3}}$ were ultimately used in the mass-mobility conversion of ions with m/z smaller than 150 or larger than 200, respectively, whereas between

m/z of 150 and 200, a linear curve of density is used. Though some ions with m/z larger than 400 fall into the region of 1.1 g cm⁻³ in Fig. S1, they were mostly OOM dimers that did not exist in Beijing (Qiao et al., 2021). Thus a density of 1.3 g cm⁻³ was applied for large organic ions in Beijing. Finally, the mobilities of all ions were adjusted according to the ambient temperature and pressure before being applied in the calibration of the APi-HTOF (Tammet, 1995).

Using the synchronous in situ measurement results of the NAIS as the reference, we obtained the detection efficiency of the APi-HTOF for ions with different m/z values. Here, since the ion concentrations measured by the NAIS have already considered sampling losses and represent atmospheric concentrations, the obtained detection efficiency of the APi-HTOF should represent a combined result of the transmission efficiency of the APi-HTOF and the sampling efficiency in the sampling line of the APi-HTOF. The ions used in the APi-HTOF are in the m/z range from 32 to 908, corresponding to the mobility diameters of 0.8-1.6 nm and ion mobilities of 3.16 to $0.77 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$, covering 12 mobility bins of the NAIS measurements. The number concentrations and signal intensities measured by the NAIS and APi-HTOF had good consistencies ($r \ge 0.66$) in the 12 bins, and the correlations were the best over the m/z range of 331–791 (r > 0.94) (Fig. S2). The good correlation between the APi-HTOF and NAIS indicates that both instruments captured the variation in the atmospheric negative cluster ions. Then the detection efficiency of a certain m/z in the APi-HTOF was calculated as the slope of the linear fitting line between signal intensities acquired in the APi-HTOF and number concentrations in the NAIS. Compared with the detection efficiencies calculated from the mass-dependent density, using a fixed density of $1.66 \,\mathrm{g}\,\mathrm{cm}^{-3}$ would overestimate the detection efficiency for small ions and underestimate that for large ions, with a deviation of up to 40 % (Fig. S3).

After excluding the effect of sampling losses in the APi-HTOF, the absolute transmission efficiency of the APi-HTOF was derived, and the obtained pattern agreed well with the relative transmission efficiency as shown in Fig. 1. The sampling losses of ions in the APi-TOF are assumed to be the same as those of neutral particles with the same sizes (Mahfouz and Donahue, 2021). As the mass-dependent density was applied, the absolute and relative transmission efficiency curves showed similar patterns; i.e., both peaked at $\sim m/z 400$ and the maximum absolute transmission efficiency reached \sim 4.9 %. The discrepancy between the absolute and relative transmission efficiency at small values of m/z may be caused by the uncertainty in small ion measurements in the NAIS, sampling efficiency calculation, or the influences of the CI inlet, as the voltage settings remain the same during the calibration experiment. It would result in an uncertainty of ~ 28 % for the quantification of small ions. The absolute transmission efficiency obtained by our calibration is roughly on the same order of magnitude as that in previous studies, and the shape is similar as well (Hein-



Figure 1. The detection efficiency and absolute transmission efficiency of the APi-HTOF determined through the in situ comparison with the NAIS and the relative transmission efficiency of the APi-HTOF determined through the depletion method. The relative transmission efficiency is obtained by dividing the detection efficiency by the sampling efficiency of cluster ions. The voltage settings of the APi-HTOF remain the same during the experiment.

ritzi et al., 2016; Junninen et al., 2010). Finally, the fit line of the detection efficiency was applied to quantify the ion compositions measured by the APi-HTOF, considering both the sampling efficiency and the transmission efficiency (Fig. 1).

After the quantification, the total negative cluster ion concentrations measured by the APi-HTOF were consistent with those measured by the NAIS (Fig. 2). However, when the total ion concentration was lower than $\sim 70 \,\mathrm{cm}^{-3}$, the total ion concentration measured by the NAIS was higher than that measured by the APi-HTOF. Specifically, the total cluster ion concentration measured by the NAIS was still higher than 25 cm⁻³ even when ions detected by APi-HTOF approached zero. These signals detected by the NAIS are mainly from small cluster ions $(3.16-1.28 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}, 0.8-1.27 \text{ nm})$, the fraction of which rises to > 80 % of the total counts when the total cluster ion concentration measured by the APi-HTOF is lower than $\sim 70 \,\mathrm{cm}^{-3}$. As the APi-HTOF has a much lower detection limit than the NAIS (Fig. S4), these signals could be caused by the high noise of small ions in the NAIS. The detection limit of the APi-HTOF is the lowest at a diameter of 1.4 nm, which is $\sim 0.02 \,\mathrm{cm}^{-3}$, and it increases when the detected ions move to lower or higher sizes due to the decrease in detection efficiency. Under a minimum diameter of 0.9 nm and a maximum of 1.7 nm, the detection limits of the APi-HTOF were ~ 0.4 and $\sim 4 \, \text{cm}^{-3}$, respectively. The detection limit of the NAIS decreases with an increase in ion diameter. The detection limit of the NAIS for cluster ions varies from $300 \text{ to } 30 \text{ cm}^{-3}$ at diameters ranging from 0.8 to 2 nm. Thus, the ion concentration measured by the APi-HTOF is more reliable than that measured by the NAIS when the ion concentration is extremely low. As the influence of background



Figure 2. The total concentration of negative cluster ions $(n_{sub-1.6 \text{ nm}}^{-})$ measured by APi-HTOF is well correlated with that measured by NAIS. Colors represent the ratio of negative ions smaller than 1.3 nm $(n_{sub-1.3 \text{ nm}}^{-})$ to negative cluster ions $(n_{sub-1.6 \text{ nm}}^{-})$.

noise would diminish when signals increase and the detection efficiency is calculated using data during the whole period, the background noise in the NAIS has limited influence on the calibration we have described above.

2.3 The simulation of HSO_4^- and $C_3H_3O_4^-$ concentration

To quantitatively address the sources and sinks of air ions, we simulate the concentration of ions using the mass balance equation and identify their dominant production and loss pathways. The simulated concentrations are compared with the measured concentrations. HSO_4^- and $C_3H_3O_4^-$ were chosen as they are two of the most abundant negative ions in different atmospheric environments and their reaction rates with neutral molecules are determined. The production pathways considered in this simulation are the ion-molecule reactions between NO_3^- and neutral molecules H_2SO_4 and $C_3H_4O_4$. Previously, it was reported that NO₃⁻ was formed in the earlier stage of atmospheric ion formation (Beig and Brasseur, 2000; Luts, 1995), and it continues to ionize neutral gaseous that are more acidic than HNO₃, such as H₂SO₄ and C₃H₄O₄ (Eisele, 1989; Tanner and Eisele, 1991). The loss pathways in the atmosphere mainly include condensational loss onto the particles, ion-molecule reactions, and ion-ion recombination with ions of opposite charge. The formation of H₂SO₄ dimer ions could be an important loss pathway for HSO₄⁻ (Beck et al., 2022). $C_3H_3O_4^-$ could further react with H_2SO_4 and change back to its neutral form C₃H₄O₄ (Beig and Brasseur, 2000). Thus, the following ion-molecule reactions are considered in the dynamic models to simulate the formation of HSO_4^- and $C_3H_3O_4^-$ in urban Beijing.

$$\mathrm{NO}_{3}^{-} + \mathrm{H}_{2}\mathrm{SO}_{4} \xrightarrow{k_{1}} \mathrm{HSO}_{4}^{-} + \mathrm{HNO}_{3}, \tag{R1}$$

$$NO_3^- + C_3H_4O_4 \xrightarrow{k_2} C_3H_3O_4^- + HNO_3,$$
(R2)

$$C_3H_3O_4^- + H_2SO_4 \xrightarrow{k_3} HSO_4^- + C_3H_4O_4, \tag{R3}$$

$$HSO_4^- + H_2SO_4 \xrightarrow{k_4} H_2SO_4 \cdot HSO_4^-, \tag{R4}$$

where k_1 to k_4 are the rate constants of these reactions, which are 2.32×10^{-9} , 2.5×10^{-9} , 2.0×10^{-9} , and 2.0×10^{-9} cm³ s⁻¹, respectively (Beig and Brasseur, 2000; Lovejoy and Curtius, 2001; Viggiano et al., 1997). The ionmolecule reactions between HSO₄⁻ and organic molecules are not considered in this model due to the lack of reaction rate constants.

The mass balance equations for HSO_4^- and $C_3H_3O_4^-$ can be written as Eqs. (4) and (5), respectively.

$$\frac{d[HSO_{4}^{-}]}{dt} = k_{1} [NO_{3}^{-}][H_{2}SO_{4}] - k_{4} [HSO_{4}^{-}][H_{2}SO_{4}] - CS [HSO_{4}^{-}] - \alpha n^{+} [HSO_{4}^{-}], \qquad (4)$$

$$\frac{d[C_{3}H_{3}O_{4}^{-}]}{dt} = k_{2}[NO_{3}^{-}][C_{3}H_{4}O_{4}] - k_{3}[C_{3}H_{3}O_{4}^{-}][H_{2}SO_{4}] - CS[C_{3}H_{3}O_{4}^{-}] - \alpha n^{+}[C_{3}H_{3}O_{4}^{-}],$$
(5)

where CS[HSO₄⁻] and CS[C₃H₃O₄⁻] are loss rates caused by the condensational loss onto particles, αn^+ [HSO₄⁻] and αn^+ [C₃H₃O₄⁻] are loss rates caused by the ion–ion recombination between ions of opposite charge, the value of α is assumed to be 1.6×10^{-6} cm³ s⁻¹ (Bates, 1982; Zauner-Wieczorek et al., 2022), and n^+ is the total concentration of positive ions and is measured by NAIS. Assuming a steady state, the concentrations of HSO₄⁻ and C₃H₃O₄⁻ are calculated as in Eqs. (6) and (7), respectively. HSO₄⁻ formed through the ion–molecule reaction between C₃H₃O₄⁻ and H₂SO₄ is neglected considering the low concentration of C₃H₃O₄⁻ in Beijing.

$$\left[\mathrm{HSO}_{4}^{-}\right] = \frac{k_{1}\left[\mathrm{NO}_{3}^{-}\right]\left[\mathrm{H}_{2}\mathrm{SO}_{4}\right]}{\mathrm{CS} + k_{4}\left[\mathrm{H}_{2}\mathrm{SO}_{4}\right] + \alpha n^{+}} \tag{6}$$

$$\left[C_{3}H_{3}O_{4}^{-}\right] = \frac{k_{2}\left[NO_{3}^{-}\right]\left[C_{3}H_{4}O_{4}\right]}{CS + k_{3}\left[H_{2}SO_{4}\right] + \alpha n^{+}}$$
(7)

3 Results and discussion

3.1 Negative cluster ion concentration and composition in urban Beijing

The median concentration of negative cluster ions is 85 cm^{-3} in Beijing during the measurement period, and its 25th and



Figure 3. (a) Negative cluster ions observed in global urban, rural, forest, and marine sites. The circles represent the median values, and the error bars represent the 25 %–75 % ranges. Ion mobility ranges for different sites were between sub-1.6 and sub-2 nm. (b) The decreasing trends of the negative cluster ion concentrations with the increase in CS were observed in both Beijing and Hyytiälä. The markers and error bars are median values and 25 %–75 % ranges, respectively. For comparison, NAIS data were used for both urban Beijing and the clean forest site Hyytiälä.

75th percentiles are 61 and 112 cm^{-3} , respectively. These values are significantly lower than concentrations reported at typical rural, forest, and marine sites (Fig. 3a). The total concentration of air ions ranges between 14 and 467 cm^{-3} in urban Beijing, similar to that in the Indian urban city Pune, which is also a highly polluted site (Gautam et al., 2017). Ion concentrations in these two polluted urban cities are so far the lowest, significantly lower than at other urban, rural and marine, and forest sites, which are mostly above $200 \,\mathrm{cm}^{-3}$ (Chen et al., 2017; Dos Santos et al., 2015; Hirsikko et al., 2011; Tammet, 2015a, b). It should be noted that although Paris is an urban site, its air is relatively clean in terms of aerosol concentration compared to the Chinese and Indian polluted urban sites. Similarly, cluster ion concentrations at some urban sites can also be as high as those at rural and remote sites due to the relatively clean atmosphere (Hirsikko et al., 2011).

The ion concentrations are negatively correlated with CS at both the urban site Beijing and the clean forest site Hyytiälä. In urban Beijing, the median ion concentration decreases from 181 to 51 ions per cm⁻³ when CS increases from 0.0032 to 0.12 s^{-1} (Fig. 3b). The negative correlation between the ion concentration and CS exists for all the mobility ranges and is more intensive for ions with mobilities between 1.72 and $0.90 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$, corresponding to a diameter range of about 1.03-1.44 nm (Fig. S5). A similar negative correlation is also observed in Hyytiälä, where the median ion concentration decreases from 606 to 369 ions per cm^{-3} when CS increases from 0.00017 to 0.012 s^{-1} , whereas the inverse relation between the negative cluster ion concentration and CS is much stronger in Beijing than in Hyytiälä. Combining data from these two sites, a negative correlation can still be seen over a wide CS range from 0.00017 to $0.12 \,\mathrm{s}^{-1}$. The influence of CS on the positive cluster ions also exists, as positive and negative ions are closely related to each other (Fig. S6). Thus, CS is an important influencing factor for the total cluster ion concentrations in both Beijing and Hyytiälä. This is very likely the reason for the lowest ion concentrations at the two polluted sites (Beijing and Pune, Fig. 3a) having the highest aerosol mass loadings.

In urban Beijing, the estimated lifetime of cluster ions due to the combined effect of condensation loss and ion-ion recombination of cluster ions is 0.02-0.4 min during haze periods $(PM_{2.5} > 75 \,\mu g \,m^{-3})$, average $CS = 0.091 \,s^{-1}$ and 0.1 -1.6 min during clean periods ($PM_{2.5} < 75 \,\mu g \,m^{-3}$, average $CS = 0.023 s^{-1}$) (Fig. S7). The lifetime of cluster ions in Hyytiälä is between 0.8-14 min (average CS = 0.0018 s^{-1}), much longer than that in Beijing. Such distinct CS dependence of total cluster ions in urban Beijing has rarely been reported in previous studies despite the condensational loss of ions on large particles having been widely recognized as important. On the contrary, most of the studies have found that weather conditions and ionizing radiation are the controlling factors for ion concentrations. For example, ion production rates in Hyytiälä were reported to be largely affected by variations in seasonal radiation and wind speed (Chen et al., 2016). This indicates that CS may be the driving factor of cluster ion concentration at highly polluted sites with high aerosol mass loadings and relatively constant ion production rates, while ion production rates may be the driving factor at relatively clean sites where aerosol mass loadings are low and the ion production rate varies significantly (Hirsikko et al., 2007).

The composition of negative cluster ions is mainly comprised of inorganic nitrogen-containing ions, inorganic sulfur-containing ions, and organic ions in urban Beijing, accounting for 20 %–22 %, 8 %–15 %, and 37 %–43 %, respectively (Figs. 4 and S8). NO₂⁻, NO₃⁻, and HNO₃NO₃⁻



Figure 4. The negative cluster ion compositions in Beijing and Hyytiälä. The bar plots were average mass spectra of negative cluster ions during clean and haze periods in Beijing and Hyytiälä measured by APi-HTOF. Signals for m/z > 300 were multiplied by 10 in Beijing for clearer views. Haze periods were identified based on whether PM_{2.5} concentration is higher than 75 µg m⁻³. The average NO_x concentrations for the three situations are 30.1, 15.0, and 1.5 ppb, respectively. The pie charts depict species distribution of organic ions, including the deprotonated form and adducts to NO₃⁻ and HSO₄⁻ forms of CHO and CHON species.

are the most abundant among the detected inorganic nitrogen-containing ions. They possibly exist in the form of $NO_2^- \cdot (H_2O)_n$, $NO_3^- \cdot (H_2O)_n$, and $NO_3^- \cdot HNO_3 \cdot (H_2O)_n$ in the atmosphere (Luts, 1995), with the loosely bounded water molecules being evaporated from the cluster ions when passing through the mass spectrometer. The concentrations of NO_2^- and NO_3^- were observed to be well correlated with each other in the atmosphere of Beijing, and the concentration of NO₂⁻ is ~ 20 % that of NO₃⁻ (Fig. S9a). This is consistent with the ion chemical models suggesting that $NO_2^$ and NO_3^- can be stably formed through a series of reactions between primary ions, NO, NO₂, and HNO₃ in the atmosphere (Beig and Brasseur, 2000; Kawamoto and Ogawa, 1984). HNO₃NO₃⁻ is subsequently formed by adding an HNO₃ molecule to NO₃⁻. Inorganic sulfur-containing ions are mainly in the form of HSO_4^- , SO_5^- , and $H_2SO_4HSO_4^-$. The ion HSO_4^- is mainly produced by the ion-molecule reaction between NO_3^- and H_2SO_4 , which will be discussed hereinafter. $H_2SO_4HSO_4^-$ is formed by the further addition of an H₂SO₄ molecule, which is proposed as the first step of ioninduced nucleation (Lovejoy, 2004). SO_5^- is likely to be generated through the reaction between O_2 and SO_3^- (Möhler et al., 1992), and its signal variation is similar to that of HSO_{4}^{-} (Fig. S9b). Similar compositions of and variations in inorganic sulfur-containing ions were also observed at Hyytiälä (Ehn et al., 2010).

The negative organic ions are mainly CHO-related or CHON-related organic ions in the form of the adduct with NO_3^- or HSO_4^- (Fig. 4). Only minor fractions (~4%) are

in the deprotonated form of CHO⁻ or CHON⁻. These indicate that the ionization schematic of organic ions is mainly through ion-molecular reaction with NO_3^- or HSO_4^- in Beijing. As few neutral sulfur-containing organics were observed, it is unlikely that the identified CHON-HSO₁ is a cluster of sulfur-containing organics and NO_3^- , but it is likely rather a cluster of nitrogen-containing organics and HSO_4^- . Among all negative organic ions, CHON organic ions adducted with NO_3^- (CHON $\cdot NO_3^-$) are the most abundant and account for 56 % and 69 % during clean and haze periods, respectively. Here the haze periods were identified based on whether PM_{2.5} concentration was higher than 75 μ g m⁻³. Though the fraction of $CHON \cdot NO_3^-$ does not change much, its composition is significantly different between clean and haze periods, mainly influenced by the differences in neutral gaseous molecules. During the haze periods where the average NO_x concentration is 30.1 ± 14.6 ppb, the organic ions are dominated by nitrated phenols (NPs), such as $C_6H_5NO_3 \cdot NO_3^-$ and $(C_6H_5NO_3)_2 \cdot NO_3^-$. During the clean periods when the average NO_x decreases to 15.0 ± 11.5 ppb, a series of organic ion peaks are observed for m/z above 300. The highest peaks in the series include $C_{10}H_{15}NO_{10} \cdot NO_3^-$, $C_{10}H_{14}O_{11} \cdot NO_3^-$, and $C_{10}H_{16}NO_{11} \cdot NO_3^-$, which are possibly the adducts of NO_3^- and highly oxygenated products from monoterpenes under high-NO_x conditions (Yan et al., 2020). NPs have high signals in urban Beijing, and their properties are different from most other CHON compounds, such as volatilities and charging efficiencies as will be discussed below. Thus, CHON species were divided into CHON_{NPs}

and $CHON_{nonNPs}$ groups in the following analysis (Nie et al., 2022).

The negative cluster ion spectra in urban Beijing are significantly different from those in Hyytiälä, possibly due to the high NO_x . Contrary to Beijing, CHO-related organic ions are more abundant than CHON-related in Hyytiälä, with fractions of 67 % and 33 %, respectively. The higher CHON ion fractions in urban Beijing are consistent with previous studies showing that neutral OOMs are composed of more nitrogen-containing species in urban Beijing compared to Hyytiälä due to the high NO_x concentration (Li et al., 2022; Qiao et al., 2021). In addition, the higher CHON_{NPs} ion fractions in urban Beijing are also the result of high anthropogenic emissions of aromatics and high NO_x concentrations (Cheng et al., 2021). Another difference is that the monoterpene-related OOM dimer ions are abundant in Hyytiälä (Isidorov et al., 1985) but are rarely observed in Beijing. This is also related to the neutral molecules as the formation of neutral OOM dimers would be suppressed under high-NO_x conditions (Nie et al., 2022). Moreover, a larger fraction of the organic ions are in the form of adducts with HSO_{4}^{-} in Beijing compared to in Hyytiälä. This is related to the higher HSO_4^- / NO_3^- ratio in Beijing (~ 0.36) compared with Hyytiälä (~ 0.05). The fraction of HSO₄⁻ adducted ions increased during clear days in Hyytiälä, but its fraction is still lower than that in Beijing (Bianchi et al., 2017). Despite the different compositions between urban Beijing and forest Hyytiälä, the ion mobility distributions are generally quite similar (Fig. S10). The mode mobility peaks of ions for both sites are $\sim 2 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$, corresponding to a mobility range dominated by HSO_4^- dimer, trimer, and organic ions. These indicate that similar ion mobility distributions do not necessarily indicate similar ion compositions and direct measurements should be performed.

Comparing the organic ions and neutral molecules measured by the APi-HTOF and the (nitrate) CI-APi-LTOF, we found that the composition of organic ions is similar to that of neutral molecules after subtracting the reagent ions (Fig. S11). The element number distributions of organic compounds detected in the APi-HTOF and CI-APi-LTOF are similar, in which the overlapping region has C atoms in the range of 3 to 14, O atoms in the range of 1 to 15, and N atoms in the range of 0 to 2 (Fig. S12). There are in total 293 organic species measured in both organic ions and neutral molecules, and in addition, inorganic sulfur-containing ions such as $(H_2SO_4)_{0-3}HSO_4^-$ were measured in both instruments. The neutral molecules detected by the CI-APi-TOF were identified by subtracting 1 NO_3^- . Such similar elemental distributions and abundant common ions in the APi-HTOF and CI-APi-LTOF indicate that naturally charging processes in the atmosphere are quite similar to the NO_3^- charging processes in the CI-APi-TOF, despite HSO_4^- also having a considerable concentration in the atmosphere and being able to charge neutral molecules as efficiently as NO_3^- . Here the reagent ions such as NO₃⁻ and HSO₄⁻ were already subtracted and only the neutral molecular formulas were compared. Organic compounds with carbon atom numbers fewer than 3 were not considered in our analysis. In addition, most molecules detected only by the CI-APi-TOF contain more C atoms of up to 20. This may be because the concentrations and signals of organic ions with such high carbon numbers are too low to be assigned a certain formula in the APi-HTOF.

3.2 Formation of negative cluster ions

To further characterize the formation of negative cluster ions, the influences from the reagent ions (i.e., NO_3^- and HSO_4^-) and the neutral molecules are explored in this subsection. Ions in the form of adducts with NO_3^- and HSO_4^- were compared to explore their ionization selectivity towards different compounds. The concentration variations in cluster ions were compared with that of the neutral molecules and the reagent ions to explore their driving factors. Then the charge fractions for different compositions were compared.

3.2.1 The influence of reagent ions

The two main reagent ions, NO_3^- and HSO_4^- , have different concentrations and variations in urban Beijing (Fig. S13). Firstly, NO_3^- is about 3 times higher than HSO_4^- . During the measurement period, the concentration of NO_3^- is 9.6 ± 6.0 cm⁻³ and that of HSO₄⁻ is 3.2 ± 3.4 cm⁻³ in urban Beijing. Similarly, NO_3^- is also more abundant than $HSO_4^$ in Hyytiälä, indicating that NO₃⁻ has greater chances to collide with neutral molecules than HSO_4^- in the atmosphere. Secondly, both NO_3^- and HSO_4^- increase with the increase in total negative cluster ions in urban Beijing, while this is reversed in Hyytiälä. This illustrates the significant impact of CS on the concentration of cluster ions in urban Beijing; that is, with the decrease in CS, both the concentrations of reagent ions and product ions increase to different degrees. Moreover, the variation in HSO_4^- is more significant than that of NO_3^- in urban Beijing. NO_3^- maintains a relatively constant proportion $(15\% \pm 5\%)$ in the total negative cluster ions during the whole sampling period in urban Beijing, while the fraction of HSO₄⁻ varies significantly due to the strong diurnal variation in H₂SO₄. The reason for the relatively constant NO_3^- fraction in total negative cluster ions needs further investigation, and this phenomenon might offer a chance to combine NO_3^- and organic ions attached to NO_3^- to predict the neutral molecules.

The compositions of molecules that cluster with NO₃⁻ and HSO₄⁻ are generally similar, with NO₃⁻ clustering with a few more compounds with high carbon numbers and low oxygen numbers. Figure 5a shows the carbon atom numbers and average carbon oxidation states (\overline{OS}_C) of molecules that are attached to NO₃⁻ and HSO₄⁻. For molecules with fewer than eight carbon atoms, molecules attached to NO₃⁻ and HSO₄⁻ have similar \overline{OS}_C between -3 and 3. For molecules with more than eight carbon atoms, molecules with lower \overline{OS}_C (<



Figure 5. (a) The carbon atom number and average carbon oxidation state (\overline{OS}_C) of molecules that adduct to NO_3^- and HSO_4^- , respectively. (b) The volatility distribution of CHO and CHON_{nonNPs} species in the form of adducts with both NO_3^- and HSO_4^- . Only species that attached to both NO_3^- and HSO_4^- , were considered for calculating the volatility distribution. The volatilities were calculated using the method of Qiao et al. (2021).

-0.5) are only attached to NO₃⁻. The average OS_C values of the molecules attached to NO_3^- and HSO_4^- are -0.3 and 0.22, respectively, and the former is close to the molecules detected in CI-APi-LTOF (-0.45). Similarly, the average O / C ratios for molecules adducting with NO_3^- and neutral molecules detected in CI-APi-LTOF are 0.85 and 0.82, respectively, slightly lower than the average O/C ratio of molecules adducting with HSO_4^- (1.03). These indicate that the selectivity of HSO_4^- and that of NO_3^- towards organic molecules are similar, with only slight differences for compounds with high carbon numbers and low oxygen numbers. The slight difference may be partly because the concentrations of these adducts of these compounds and HSO_4^- are too low to be accurately distinguished. Overall, the O/C ratios of compounds adducted with both NO_3^- and HSO_4^- are both significantly higher than those detected by the (iodide) CI-APi-TOF and (oxygen) CI-APi-TOF (Li et al., 2021; Riva et al., 2019), indicating their higher selectivity for oxidized compounds, which is consistent with previous theoretical simulations (Nadykto et al., 2018). In addition, when only CHO and $CHON_{nonNPs}$ molecules clustering with both NO_3^- and HSO_4^- are considered, their volatility distributions are very similar when clustering with either NO_3^- or HSO_4^- , which further indicates that the selectivity difference between $NO_3^$ and HSO_4^- towards organic molecules is very small (Figs. 5b and S14).

For the 88 species that form clusters with both NO_3^- and HSO_4^- , the ratio of the two ionization forms $(M \cdot HSO_4^- / M \cdot NO_3^-)$ is positively related to the ratio of reagent ions (HSO_4^- / NO_3^-) . In addition, HSO_4^- forms clusters with CHO and CHON_{nonNPs} more efficiently, while NO_3^- forms clusters with CHON_{NPs} more efficiently, despite the abovementioned similarity in their selectivity towards different organic molecules. Overall there are 33 CHO,

40 CHON_{nonNPs}, and 15 CHON_{NPs} species that form ions with both NO₃⁻ and HSO₄⁻. The average ratio HSO₄⁻ / NO₃⁻ is 0.36, and the relative ratios of $M \cdot HSO_4^- / M \cdot NO_3^-$ are 0.74, 0.73, and 0.23 for CHO, CHON_{nonNPs}, and CHON_{NPs} species, respectively. Generally, the $M \cdot HSO_4^- / M \cdot NO_3^-$ ratio is positively related to the HSO_4^- / NO_3^- ratio, and the dots are close to the 1:1 line as shown in Fig. 6. Specifically, the dots for CHON_{NPs} are mostly lower than the 1 : 1 line, especially under high H₂SO₄ concentration. Similar decreasing trends were also observed for the $M \cdot HSO_4^- / M \cdot NO_3^$ ratio for CHO and CHON_{nonNPs}. This is possibly due to the competition between different neutral compounds for the reagent ions, as the atmospheric concentration of ions is up to 7 orders of magnitude lower than for neutral molecules. For example, gaseous H₂SO₄ may compete with the organic molecules to react with HSO_4^- . As a highly electronegative substance, H₂SO₄ prefers to be negatively charged and form strongly bounded clusters with alkaline substances such as HSO_4^- or amines in the atmosphere; thus HSO_4^- would prefer to cluster with H₂SO₄ during the daytime, and its efficiency of clustering with organic molecules drops as a result. During the nighttime, when the competition from H₂SO₄ weakens, the formation of $M \cdot HSO_4^-$ increases.

3.2.2 The influence of neutral molecules

The diurnal cycles of negative cluster ions are mainly determined by their corresponding neutral molecules, and exceptions were found for CHON_{NPs}-related ions, the diurnal cycles of which are more related to the reagent ions (Fig. 7). The reagent ions and their inorganic clusters ((HNO₃)₀₋₂NO₃⁻ and (H₂SO₄)₀₋₂HSO₄⁻) both reach their maximum at 13:00 LT but show different diurnal patterns. (H₂SO₄)₀₋₂HSO₄⁻ has a single-peak pattern that resembles neutral H₂SO₄, while (HNO₃)₀₋₂NO₃⁻ has a three-peak pat-



Figure 6. The ratio of $CHO \cdot HSO_4^- / CHO \cdot NO_3^-$ (a), $CHON_{nonNPs} \cdot HSO_4^- / CHON_{nonNPs} \cdot NO_3^-$ (b), and $CHON_{NPs} \cdot HSO_4^- / CHON_{NPs} \cdot NO_3^-$ (c) as a function of the ratio of HSO_4^- / NO_3^- . The color represents the concentration of sulfuric acid.



Figure 7. Diurnal cycles of negative cluster ions, corresponding neutral molecules, and the charge fractions between them in urban Beijing. (a) The diurnal cycles of reagent ions and their inorganic clusters, together with those of organic ions. The organic ions in the form of adducts with NO_3^- and HSO_4^- were summed into CHO-, CHON_{nonNPs}-, and CHON_{NPs}-related ions. (b) The diurnal cycles of sulfuric acid and the corresponding neutral molecules of organic ions measured by the (nitrate) CI-APi-LTOF. The concentrations of sulfuric acid, CHO, and CHON_{nonNPs} were multiplied by a factor of 100, 20, and 20, respectively, to match that of CHON_{NPs} in the figure. (c) The diurnal cycles of charge fractions between negative cluster ions and their corresponding neutral molecules. The charge fractions for CHO, CHON_{nonNPs}, and CHON_{NPs} in the form of adducts with NO_3^- and HSO_4^- were calculated and compared with the HSO_4^- / H_2SO_4 ratio.

tern that matches the total concentration of negative cluster ions. For the organic ions, the CHO- and CHON_{nonNPs}related ions both peak in the afternoon, which resembles the diurnal pattern of corresponding neutral molecules. However, the diurnal patterns of CHON_{NPs}-related ions and neutral $\mbox{CHON}_{\mbox{NPs}}$ show opposite variations, and that of CHON_{NPs}-related ions is closer to the diurnal variation in NO_3^- . As CHON_{NPs}-related ions reach their maximum in the nighttime, neutral CHON_{NPs} reach their maximum both at noon and in the evening, indicating that the formation of CHON_{NPs}-related ions may be different from that of other organic ions. Besides, although the diurnal variations in CHOand CHON_{nonNPs}-related ions roughly follow those of neutral molecules, it was found that their peak times coincided with that of $(HNO_3)_{0-2}NO_3^-$ at 13:00 and 17:00 LT. Differently, the diurnal patterns of CHO- and CHONnonNPs-related ions tended to peak during the nighttime in Hyytiälä (Bianchi et al., 2017). This could be due to the different diurnal patterns of neutral molecules as those of the reagent ions are similar between Beijing and Hyytiälä.

Consistently, the cluster ion concentrations were found to be positively correlated with those of neutral molecules under similar CS for CHO and CHON_{nonNPs} species during the whole sampling period. As shown in Fig. 8, generally, the concentrations of all organic ions decrease with increasing CS as also indicated in Sect. 3.1. When CS is larger than 0.01 s^{-1} , the concentrations of CHO- and CHON_{nonNPs}related ions rise with their corresponding neutral gas concentration under similar CS levels, while that of CHON_{NPs}related ions remains constant or slightly declines. When CS is smaller than 0.01 s^{-1} , which is usually accompanied by low concentrations of neutral molecules, the relationships be-



Neutral molecules measured by CI-APi-TOF (cm⁻³)

Figure 8. Scatterplots of the concentrations of organic ions measured by the APi-TOF and their corresponding neutral molecules measured by the CI-APi-TOF in urban Beijing, colored by CS. The cluster ions are $\text{CHO} \cdot \text{NO}_3^-$ (**a**), $\text{CHO} \cdot \text{HSO}_4^-$ (**b**), $\text{CHON}_{\text{nonNPs}} \cdot \text{NO}_3^-$ (**c**), $\text{CHON}_{\text{nonNPs}} \cdot \text{HSO}_4^-$ (**d**), $\text{CHON}_{\text{NPs}} \cdot \text{NO}_3^-$ (**e**), and $\text{CHON}_{\text{NPs}} \cdot \text{HSO}_4^-$ (**f**). The neutral molecules are CHO (**a**, **b**), $\text{CHON}_{\text{nonNPs}}$ (**c**, **d**), and CHON_{NPs} (**e**, **f**), respectively. The data are divided into five groups based on CS in the range of 0.001–0.01, 0.01–0.02, 0.02–0.06, 0.06–0.1, and $0.1–0.2 \text{ s}^{-1}$, respectively.

tween organic ions and the corresponding neutral molecules are not obvious; this may be due to a combined effect of neutral molecules and the reagent ions.

The charge fraction between organic ions and neutral molecules reflects a charging competition between different neutral compounds. Here, we calculate the charge fraction as the proportions of the concentrations of cluster ions and their corresponding neutral molecules. As shown in Fig. 7c, different organic molecules have similar diurnal variations in charge fractions, which are highest at 05:00 LT and lowest at 15:00 LT, while the HSO_4^- / H_2SO_4 ratio has the opposite pattern and is highest during the daytime. The opposite diurnal patterns between organic ions and HSO_4^- indicate that H₂SO₄ and organic molecules compete for being charged. The median charge fractions of CHO and $CHON_{nonNPs}$ molecules with NO₃⁻ and HSO₄⁻ are close to each other and range from 2×10^{-7} to 2×10^{-6} , while the charge fractions of CHON_{NPs}-related ions are much lower than these organic ions. This suggests that CHON_{NPs} are less likely to form cluster ions with NO^-_3 and HSO^-_4 than CHO and CHONnonNPs. As the charge fraction for organic ions attached to NO_3^- and HSO_4^- has similar diurnal patterns, it is deduced that the diurnal variation in charge fractions is mainly driven by the variation in neutral molecules instead of reagent ions.

For CHO and CHON_{nonNPs}, the charge fractions between organic ions and neutral molecules generally increase with the carbon atom numbers and \overline{OS}_{C} of the molecules. As shown in Fig. 9, the average charge fractions of different organic ions vary between 3×10^{-8} and 8×10^{-6} , and species with larger carbon atom numbers and higher oxidation states tend to have higher charge fractions when clustering with either NO_3^- or HSO_4^- (Fig. S15). The charge fraction difference between molecules with different carbon atom numbers and OS_C is a reflection of the differences in reaction rates between reagent ions and the neutral molecules or the stabilities of the cluster ions. In summary, through the comprehensive quantitative analysis of atmospheric cluster ions, reagent ions, and neutral molecules, we can roughly predict the composition and concentration variations in atmospheric negative cluster ions from measured neutral molecules and CS, as indicated by Fig. 8. However, as the actual charging frac-



Figure 9. The carbon atom number and average carbon oxidation state (\overline{OS}_C) of molecules in CHO · NO₃⁻ (**a**) and CHON_{nonNPs} · NO₃⁻ (**b**). The color represents their charge fraction during the measurement periods.



Figure 10. (a) The measured and simulated concentrations of HSO_4^- and $C_3H_3O_4^-$ in Beijing. The simulation is performed according to dynamic models as described in Sect. 2.3, and four formation and loss pathways are considered. (b) The fractions of the formation or loss rates of HSO_4^- and $C_3H_3O_4^-$ contributed by different pathways. The average values of the results are shown here.

tions for different species differ in a wide range as shown in Fig. 9, the prediction of actual concentrations of atmospheric ions would require more work.

3.3 Quantification of the sources and sinks of representative negative cluster ions

We found that the ionization of H_2SO_4 and $C_3H_4O_4$ by $NO_3^$ and condensational loss to particles are the main formation and loss pathways for HSO_4^- and $C_3H_3O_4^-$, respectively. As shown in Fig. 10, the simulated concentrations of both $HSO_4^$ and $C_3H_3O_4^-$ by Eqs. (6) and (7) have a good consistency with the measured concentrations, indicating that the production and loss pathways considered can reproduce the measured ion concentration in urban Beijing. The ionization of H_2SO_4 and $C_3H_4O_4$ by NO_3^- is the main production pathway. For both HSO_4^- and $C_3H_3O_4^-$, the condensational loss is the main loss pathway, contributing to more than 70 % of the total loss rate, which explains why CS is a driving factor for the concentration of cluster ions in urban Beijing. The ion–ion recombination process only accounts for 6% and 8% of the total loss rates for HSO_4^- and $C_3H_3O_4^-$, which is due to the low ion concentrations in the polluted urban atmosphere. Notably, the ion–molecular reactions between the ions with H_2SO_4 could contribute to 19% of the loss of both HSO_4^- and $C_3H_3O_4^-$. The transformation from $C_3H_3O_4^-$ back to $C_3H_4O_4$ indicates that the conversion of ions back to neutral molecules is also significant.

4 Conclusions

We quantified the composition-resolved ion concentrations in the atmosphere by combining mass spectrometry and electrical mobility measurements. The absolute transmission efficiency in the APi-HTOF obtained with the ion mobility spectrometer (NAIS in this study) agreed well with the relative transmission efficiency obtained in the chemical ionization mode of the APi-HTOF. The calibrated ion concentrations with different m/z ranges agreed well with the concentration of ions with corresponding mobility ranges of the ion mobility spectrometer. These indicate that an APi-HTOF can be well calibrated by running an APi-HTOF and an ion mobility spectrometer side by side in ambient measurements. Furthermore, we propose that the transmission efficiency of a CI-APi-TOF can also be obtained through in situ comparison with an ion mobility spectrometer, as long as the X-ray and voltages of a chemical ionization inlet are turned off and atmospheric ions are directly measured.

The cluster ion concentrations and composition in urban Beijing are largely affected by the high condensation sink and nitrogen oxides. Firstly, ion concentration decreased significantly with an increasing CS for cluster ions in all the mobility ranges. Median cluster ion concentrations in urban Beijing were only $\sim 85 \text{ cm}^{-3}$, much lower than those reported at clean and rural sites due to the high CS in urban Beijing. Due to high concentrations of nitrogen oxides, the organic ion compositions were composed of more nitrogencontaining species in urban Beijing than those in Hyytiälä. This is consistent with the higher neutral nitrogen-containing molecular fractions in urban Beijing as previously reported.

The formation of cluster ions was analyzed by comparing their concentrations, volatility distributions, and charge fractions with CS, reagent ions, and their corresponding neutral molecules. The organic cluster ions are mainly in the form of adducts with NO_3^- or HSO_4^- . The ratio of the two ionization forms $(M \cdot HSO_4^- / M \cdot NO_3^-)$ is positively related to the ratio of reagent ions (HSO_4^-/NO_3^-) . Although the molecules clustered with HSO₄⁻ and NO₃⁻ are similar in composition, we found that M HSO₄ formed more efficiently for CHO and CHON_{nonNPs}, while $M \cdot NO_3^-$ formed more efficiently with $CHON_{NPs}$ when targeting the compounds that can be ionized by both HSO_4^- and NO_3^- . The concentrations of organic ions are positively correlated with that of neutral molecules, resulting in their similar diurnal cycles. However, an exception was found for CHON_{NPs}, the concentration of which is also significantly influenced by the reagent ions NO_3^- . The charge fractions tend to be higher for organic molecules with higher molecular weight and a higher oxidation state. We also observed a charging competition between different neutral organic and inorganic compounds, such as the competition between H₂SO₄ and organic ions. Through this quantitative analysis, it is possible to infer the variation in atmospheric negative cluster ions from the measured neutral molecules and CS in the urban atmosphere. The formation pathways of HSO_4^- and $C_3H_3O_4^-$ were well characterized by the ionization of H_2SO_4 and $C_3H_4O_4$ by NO_3^- , and their loss processes are dominated by condensation loss, with minor contributions from ion-molecular reactions and ionion recombination.

Inspired by the high dependence of the charge fraction on the molecule species, we can take the ambient atmosphere as a natural ion-molecule reaction chamber, and the charge fraction observed for different neutral molecules may provide some insights into the charging efficiency of organic species measured with the reagent ions of NO_3^- and $HSO_4^$ in the chemical ionization mass spectrometers and help better quantify them.

Data availability. The detection efficiency of the APi-TOF, the time series of total negative cluster ions and CS in urban Beijing, and the average spectra of negative clusters ions measured by APi-TOF during haze and clean periods in urban Beijing are available from https://doi.org/10.5281/zenodo.7791785 (Yin, 2023).

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