

Molecular composition and processing of aqueous secondary organic aerosol in clouds at a mountain site in southeastern China

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Abstract. Aqueous secondary organic aerosol (aqSOA) contributes substantially to organic aerosol (OA), affecting air quality, human health, and climate. However, the molecular composition and processing of aqSOA in clouds remain unclear due to limited online field measurements. We measured molecular composition of OA online (time resolution 20 s) and tracked its processing at a mountain site in southeastern China, using an Extractive ElectroSpray Ionization inlet coupled with a Time-of-Flight Mass Spectrometer (EESI-ToF-MS). We identified 2084 molecular formulas and compared OA composition from three sample types: cloud droplet residuals (CDR), interstitial aerosol particles (INT), and cloud-free aerosol particles (CF) in representative cloud episodes. CHO class was the dominant constituent, followed by CHON class. In most cloud episodes, the fraction of CHO was lower in CDR than that in INT and CF, while the fraction of CHON was higher, which may result from the uptake of organonitrates or nitration in cloud water. Compounds in CDR had more carbon number and higher molecular weight than CF, which is attributed to accretion reactions in cloud water. We identified 39 significantly enriched compounds in CDR compared with CF, which could be potentially used as aqSOA tracers formed via cloud processing. This study also reveals rapid changes in aqSOA composition, which highlight the necessity for high time resolution measurements to capture the processing of aqSOA in clouds. Overall, this study provides clear information on processing of aqSOA in clouds and highlights the importance of accretion reactions, which have implications on the composition and physicochemical properties of SOA.

1 Introduction

Secondary organic aerosol (SOA) is a major component of organic aerosol (OA) with diverse emission sources, gaseous precursors, and composition, exerting significant impacts on air quality, climate, and human health (Jimenez et al., 2009; Nault et al., 2021). SOA is primarily produced through the oxidation of volatile organic compounds (VOCs), while the atmospheric aging of primary organic aerosol (POA) may also contribute. Numerous previous studies have investigated the formation mechanisms of SOA, with particular emphasis on gas-phase pathways (Odum et al., 1996; Ervens et al., 2011). However, SOA formed solely through gas-phase reactions (gasSOA) cannot fully account for the observed SOA concentrations (de Gouw et al., 2005; Volkamer et al., 2007; Volkamer et al., 2006). In addition to the traditional gas-phase processing, aqueous-phase pathways have been recognized as an important source of SOA (Sehested et al., 1975; Galloway et al., 1976; Graedel and Weschler, 1981; Fu et al., 2008; Tan et al., 2009; Zhang et al., 2010; Ervens et al., 2011; Lamkaddam et al., 2021).

Mounting evidence for aqueous secondary organic aerosol (aqSOA) has been reported in field observations in various atmospheric aqueous systems, i.e., aerosol liquid water (ALW), fog water, and cloud water. For example, several studies on source apportionment in different sites showed that aqSOA formed in ALW is an important contributor to SOA, with its fraction particularly elevated (up to 44 %) under high relative humidity (RH) conditions (Wang et al., 2021; Zhao et al., 2019; Tong et al., 2021; Gilardoni et al., 2016; Duan et al., 2022; Xu et al., 2019; Sun et al., 2016). Relative to ALW, fog water and cloud water are diluted aqueous systems where aqSOA can also be formed (Herckes et al., 2013). For fog water, the ratio of aqSOA to OA during fog-rain days is enhanced compared with non-fog-rain days (Duan et al., 2021). Additionally, OA composition of fog water is more oxidized (Brege et al., 2018), has more N-containing compounds (Mattsson et al., 2025; Sun et al., 2024a; Kim et al., 2019) compared with aerosol particles, and shows signs of oligomerization based on fragments in the mass spectrum (Gilardoni et al., 2016; Mandariya et al., 2019). In contrast to fog, the cloud is more common, ubiquitously presents in the atmosphere, and consists of a large quantity of droplets generated by aerosol activation, providing an aqueous medium for physical processes and chemical reactions (McNeill et al., 2012; McNeill, 2015). Within clouds, aerosol particles may undergo repeated hydration-dehydration cycles, including hygroscopic growth, activation, and subsequent evaporation. Such cloud processing could influence the concentration of OA composition (Wang et al., 2024b; Gao et al., 2023; Liu et al., 2023b), thereby influencing aerosol size distribution, hygroscopicity, volatility, and cloud condensation nuclei (CCN) activity (Jimenez et al., 2009; Sun et al., 2025). Additionally, cloud processing may facilitate the formation of brown carbon, including N-containing heterocyclic compounds, which could affect atmospheric radiative forcing (Liu et al., 2023b).

A number of field campaigns have been conducted to measure the chemical composition of OA in cloud droplet residuals. Several previous field campaigns found that more highly oxygenated OA is present in cloud droplet residuals compared to cloud-free aerosol particles using online techniques, Aerosol Mass Spectrometer (AMS) or Aerodyne Aerosol Chemical

Speciation Monitor (ACSM), which provide information on fragment ions of compounds, such as the fraction of m/z 44 (CO_2^+) in the mass spectra (Dadashazar et al., 2022; Lance et al., 2020; Gao et al., 2023). Although these studies provide valuable information on the chemical composition of aqSOA, the use of AMS or ACSM leads to molecular fragmentation and thus cannot provide molecular formulas for the components of aqSOA. As a result, the molecular composition of aqSOA and mechanisms of its formation and transformation remain incompletely understood. This gap hinders the analysis of sources, evolution, health effects, and climate impacts with respect to specific OA compounds.

Molecular formulas of OA in cloud samples can be assigned and classified into several groups, including CHO, CHON, CHOS, and CHONS, with CHO and CHON accounting for the largest fractions (Liu et al., 2023b; Cook et al., 2017; Pailler et al., 2024; Zhao et al., 2013; Bianco et al., 2019; Sun et al., 2021; Gramlich et al., 2023). Oligomers (Cook et al., 2017; Zhao et al., 2013), organosulfates (Sun et al., 2021; Bianco et al., 2019), and N-containing compounds such as nitroaromatics (Sun et al., 2021; Cook et al., 2017; Bianco et al., 2019) have been observed in cloud droplets. Although the molecular composition of OA in cloud droplets has been characterized using offline techniques such as Fourier Transform Ion Cyclotron Resonance Mass Spectrometry (FT-ICR-MS), the formation mechanisms of many compounds in clouds remain uncertain. For example, it is not clear whether the oligomers originate from cloud processing or from aqueous aerosol due to a lack of concomitant aerosol measurements and limited temporal variation analyses (Cook et al., 2017; Zhao et al., 2013). The coarse time resolution of filter-based sampling (several hours to one day), together with limited sample numbers, prevents these studies from resolving cloud-processing reaction processes that occur on minute-to-hour timescales and are subjected to the influence of rapid variability in meteorological conditions within clouds. The chemical characteristics of aqSOA obtained from comparisons between cloud droplets and cloud-free aerosol particles are subject to large uncertainties, because the composition of both cloud droplets and aerosols may change over long-time sampling. Therefore, it is necessary to obtain online molecular information on OA in clouds by comparing OA composition of cloud droplets, interstitial aerosol particles, and cloud-free aerosol particles, to provide new insights into the detailed chemical composition, evolution variation, and the mechanism of cloud processing.

To get a detailed understanding of cloud processing of aqSOA, we measured the real-time molecular composition of aqSOA in clouds using an Extractive ElectroSpray Ionization inlet coupled with a Time-of-Flight Mass Spectrometer (EESI-ToF-MS) in a mountain site in southeastern China. In this study, we identify molecular formulas of OA in cloud processing and compare differences in OA characteristics between cloud droplet residuals (CDR), interstitial aerosol particles (INT), and cloud-free aerosol particles (CF). We explore new compounds formed in cloud processing and explain their potential formation mechanisms. We also aim to track the temporal evolution of compounds in aqSOA during cloud processing.

2 Methods

We conducted this field campaign from May 1st to May 29th in 2024 at Shanghuang Eco-Environmental Observatory of Chinese Academy of Sciences at the summit of the Damaojian mountain (119.51° E and 28.58° N, 1128 m above sea level) located in Jinhua city, Zhejiang province, China. The site is a background monitoring station surrounded by coniferous and broad-leaved forests away from megacities, as shown in Fig. 1. In addition to biogenic emissions, this site may be affected by anthropogenic activities originating from the surrounding small counties, as mentioned in Zhang et al. (2024).

Cloud droplets were collected using a Ground-based Counterflow Virtual Impactor (GCVI, Brechtel Manufacturing Inc., Model 1205). The GCVI collected cloud droplets with diameters larger than 8.5 μm (Shingler et al., 2012) under conditions of visibility < 3 km, RH > 95 %, and absence of precipitation. After separation from INT (non-activated aerosol in clouds), the cloud droplets were dried by mild heating (40 °C) within the GCVI (Lin et al., 2017) and further by a Nafion dryer downstream, and CDR were subsequently measured. Because the focus is on the relative compositional change of OA in CDR and CF, the GCVI enhancement factor was not applied. A PM_{2.5} (particulate matter smaller than 2.5 μm) cyclone inlet (URG, USA) was used to collect INT and CF. A switching system alternated between the GCVI and the URG inlet: PM_{2.5} was sampled when GCVI detected no cloud, whereas CDR sampling was triggered automatically once cloud presence was detected by GCVI. During cloud episodes, the switch was also configured to alternate between CDR and INT sampling. It should be noted that the terms “cloudy days” and “cloudless days” in this study specifically refer to periods with and without low clouds.

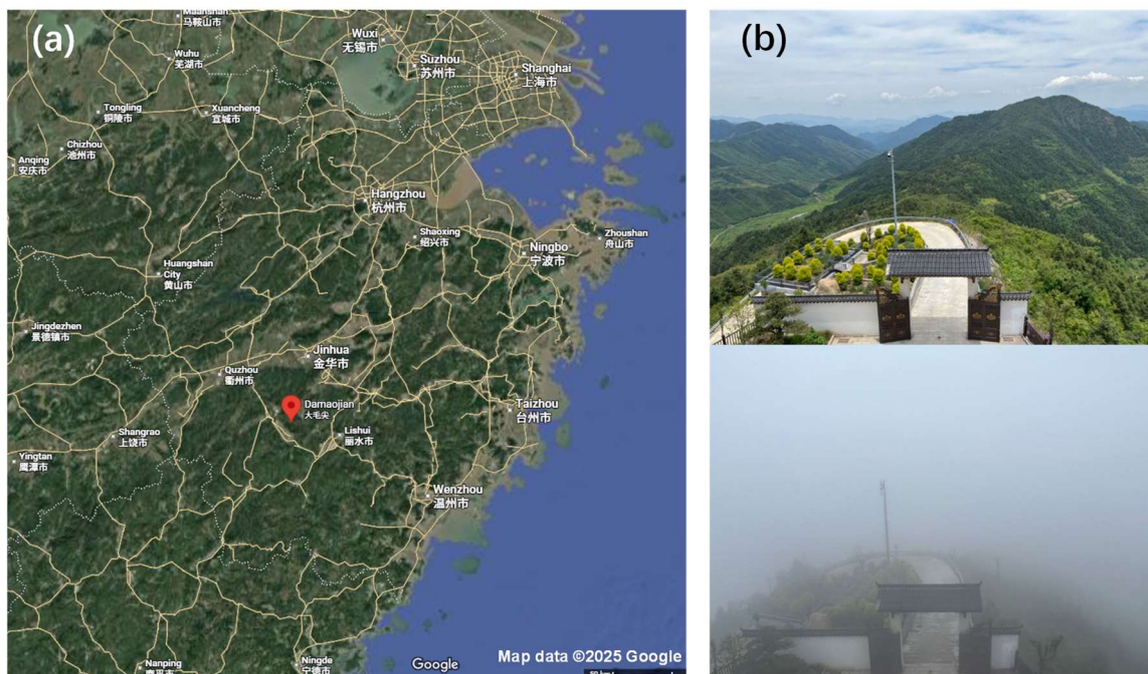


Figure 1. Location of the Shanghuang site. (a) a map (from Google Maps) and (b) two photos of the sampling site, one with cloud and another without cloud.

Measurements of CDR, INT, and CF were conducted through a manifold positioned downstream of both the GCVI and URG inlets. The concentrations of OA composition were measured online using EESI-ToF-MS (Aerodyne Institute) with a

time resolution of 20 s. This mass spectrometer achieves soft ionization while preserving the structure of compounds, measuring molecular formulas with high mass resolution (8000–10000) and low detection limit (Wang et al., 2024a). Detailed information regarding EESI-ToF-MS has been reported previously (Lopez-Hilfiker et al., 2019; Stefenelli et al., 2019; Brown et al., 2021; Kumar et al., 2022; Luo et al., 2024; Xue et al., 2025). Aerosol was sampled after gaseous compounds were removed by entering a charcoal denuder, and subsequently intersected with an electrospray generated from a working solution containing 100 ppm NaI in a 1:1 (v/v) water and acetonitrile mixture, allowing aerosol compounds to be detected as $[M+Na]^+$ in positive ion mode. Background measurements were obtained by switching the inlet to a filter. The durations of sample and background collection can be adjusted to ensure aerosol signal levels return to baseline within the time of background (Qi et al., 2019). In this campaign, the sample and background were set in combinations of 10 min and 5 min typically. The sampling volume of EESI-ToF-MS was 0.9 L m⁻³. Weekly calibration was performed using levoglucosan, and the sensitivity was assumed identical for all compounds. All organic compound signals are shown as relative intensities normalized to (NaI)Na⁺ to avoid interference from the ion source fluctuations in EESI-ToF-MS. Mass spectral data were processed using Tofware 3.2.5 in Igor Pro 8. For data screening, the signal-to-background ratio (s/b) was calculated as the median value of (sample signal–background)/background, thereby excluding compounds showing insignificant differences between sample and background. Only compounds with the s/b ratio greater than 0.1 were included (Tong et al., 2021). After this screening, 79, 148, 604, and 126 compounds were retained from cloud episodes one to four, respectively. All results presented in Sect. 3.2 are based on these screened data.

The OA size distribution was characterized using a scanning mobility particle sizer (SMPS, TSI 3936), which, together with an atomizer (TSI, 3076), was used to facilitate EESI-ToF-MS calibration. PM_{2.5} concentration was monitored using a Thermo Scientific instrument (Thermo Scientific. Model 5014i), while CO was measured by a Picarro greenhouse gas analyzer (Picarro Inc., G2401). Meteorological parameters including RH, Temperature (T), wind speed (WS), and wind direction (WD) were monitored by an automatic weather station.

The 72 h backward trajectories of air masses arriving at the Shanghuang site were calculated by the Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPPLIT) model, with Global Data Assimilation System (GDAS) meteorological data at 1°×1° spatial resolution (Stein et al., 2015; Rolph et al., 2017). These trajectories were clustered into several appropriate groups for selected cloud episodes. The clustering was based on the total spatial variance (TSV) method (Song et al., 2023; Roland et al., 2025) in MeteInfo software (Wang, 2014).

145 **3 Results and discussion**

3.1 Characteristics of cloud episodes

During the entire campaign, $\text{PM}_{2.5}$ concentration was $13.4 \pm 10.8 \mu\text{g m}^{-3}$ (mean value \pm standard deviation), with the highest concentration of $72.2 \mu\text{g m}^{-3}$ observed on cloudless days, as shown in Fig. 2. The $\text{PM}_{2.5}$ concentration is typical in rural areas of China and is substantially lower than that observed in the same season in metropolitan cities such as Beijing ($\sim 40 \mu\text{g m}^{-3}$) and Shanghai ($\sim 30 \mu\text{g m}^{-3}$) (Liu et al., 2023a; Yin et al., 2023).

Cloud episodes accounted for 27.1 % of the one-month campaign, with the sample types of CDR and INT representing 13.1 % and 14.0 %, respectively. Of the 16 recorded cloud episodes, we selected those without precipitation to avoid the influence of wet deposition and with a cloud-free period within < 2 h from the cloud episode, and for which CDR, INT, and CF samples were all available. Six out of 16 episodes meet these criteria, and four cloud episodes (CEs) are further selected. CE1, CE2, CE3, and CE4 differ in $\text{PM}_{2.5}$ and CO concentration, meteorological conditions, origin of air mass, and duration time, as shown in Table S1. The duration of each CE ranged from several minutes to three days. Meteorological conditions and origin of air masses are discussed in Text S1, and backward trajectories from HYSPLIT are shown in Fig. S2.

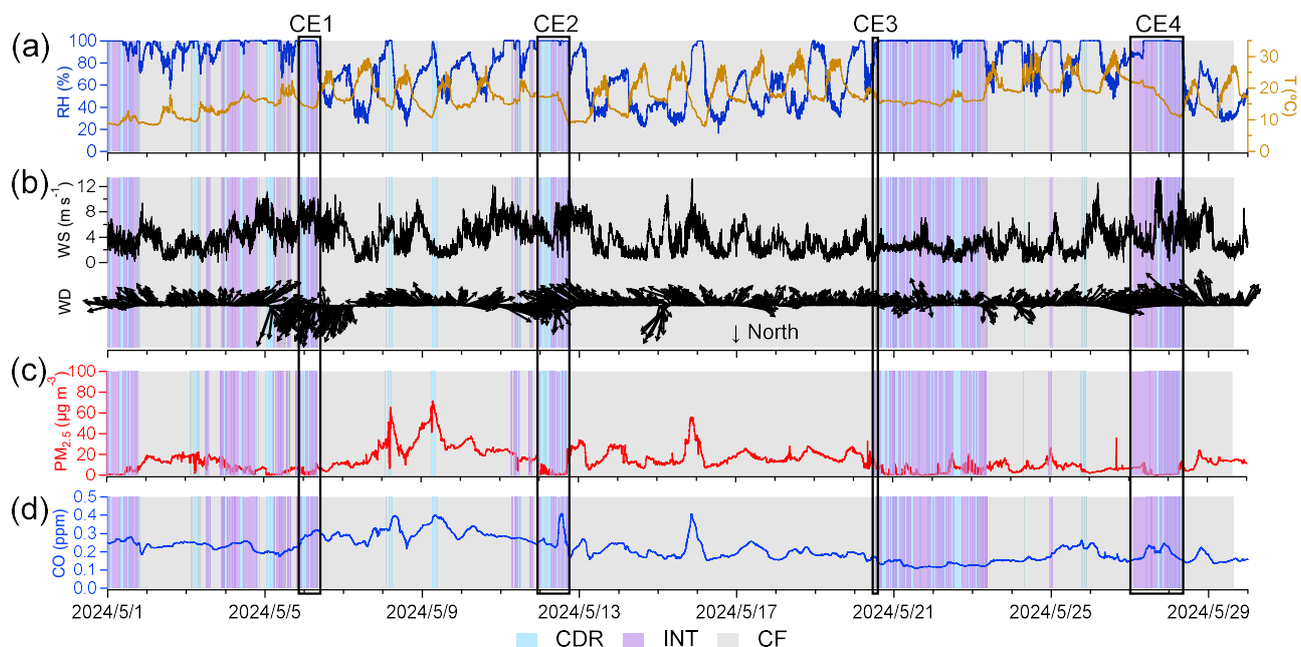


Figure 2. Time series of (a) RH and T, (b) WS and WD, (c) $\text{PM}_{2.5}$, and (d) CO. Sample types of cloud droplet residuals (CDR), interstitial aerosol particles (INT), and cloud-free aerosol particles (CF) are shaded as blue, purple, and gray, respectively.

Each CE's sampling period is divided into three stages (pre-cloud, in-cloud, and post-cloud) to compare OA characteristics. The in-cloud stage corresponds to the sample types of CDR and INT, whereas both the pre-cloud and post-cloud stages correspond to the sample type of CF ($\text{PM}_{2.5}$). Detailed characteristics of sample types in four CEs, such as mean chemical formula, H/C, O/C, N/C, and OSc (carbon oxidation states, $2 \times \text{O/C} - \text{H/C}$), are shown in Table 1, and division of stages is shown in Fig. S1.

A total of 2084 molecular formulas of OA were identified in the campaign. Mean formula of CDR was $C_{9.95-12.92}H_{14.53-21.78}O_{5.15-6.02}N_{0.32-0.42}S_{0-0.01}Si_{0-1.29}$ for CE1–CE4. Compared with pre-cloud aerosols with formula $C_{8.17-10.57}H_{12.99-16.14}O_{4.98-5.64}N_{0.12-0.30}S_{0-0.01}Si_{0-0.24}$, CDR exhibited increased numbers of carbon, hydrogen, oxygen, and nitrogen atoms, with the differences being statistically significant ($p < 0.05$) (Table S2). These molecular formulas were classified into eight classes, that is, CHO (only C, H, O atoms are contained in the chemical formula, hereafter), CHON, CHONS, CHOS, CHN, CHS, CHNS, and CHOSi. Since the composition of OA varied in different CEs, the fractions of these OA classes are discussed for each CE in Sect. 3.2. The O/C ratio was generally lower in CDR (0.45–0.66) than in pre-cloud aerosols, INT, and post-cloud aerosols in 4 CEs. The O/C ratio in CDR is comparable to those reported for fog water (0.52–0.68), aqSOA (0.61–0.84), and oxygenated OA (0.44–0.83) by Gilardoni et al. (2016). In general, O/C of CDR in this study is comparable to that of fog (0.58–0.8) in the Po Valley in Brege et al. (2018), while H/C of CDR (1.50–1.87) is higher than that of fog (1.29–1.37) in that study. Furthermore, CDR showed elevated N/C (0.033–0.045) relative to other sample types, while its OSc value (–0.83 to –0.25) is generally lower than in other sample types.

Table 1. Mean molecular formulas and elemental parameters (H/C, O/C, N/C, and OSc; mean \pm standard deviation) of OA for CE1–CE4 in pre-cloud aerosols, CDR, INT, and post-cloud aerosols.

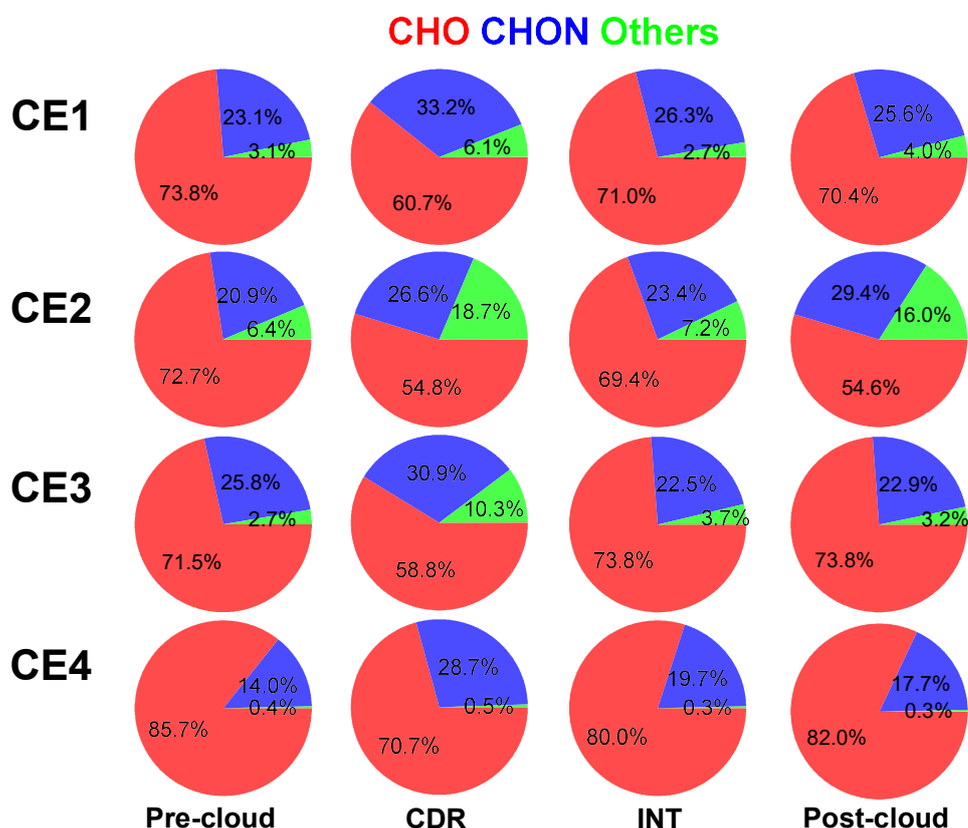
| | Pre-cloud aerosols | CDR | INT | Post-cloud aerosols |
|-------------|---|---|--|---|
| CE1 Formula | $C_{8.72}H_{13.81}O_{5.64}N_{0.20}S_{0}Si_{0.24}$ | $C_{10.45}H_{17.25}O_{6.02}N_{0.32}S_{0}Si_{0.69}$ | $C_{8.99}H_{14.21}O_{5.68}N_{0.26}S_{0}Si_{0.23}$ | $C_{8.86}H_{14.45}O_{5.71}N_{0.22}S_{0}Si_{0.40}$ |
| H/C | 1.61 \pm 0.04 | 1.64 \pm 0.11 | 1.61 \pm 0.04 | 1.61 \pm 0.04 |
| O/C | 0.71 \pm 0.02 | 0.66 \pm 0.03 | 0.69 \pm 0.02 | 0.70 \pm 0.02 |
| N/C | 0.022 \pm 0.004 | 0.033 \pm 0.008 | 0.028 \pm 0.005 | 0.025 \pm 0.006 |
| OSc | -0.19 \pm 0.06 | -0.33 \pm 0.14 | -0.22 \pm 0.06 | -0.20 \pm 0.06 |
| CE2 Formula | $C_{8.33}H_{13.72}O_{4.98}N_{0.28}S_{0.01}Si_{0.16}$ | $C_{10.13}H_{19.51}O_{5.44}N_{0.39}S_{0.01}Si_{1.29}$ | $C_{8.52}H_{14.19}O_{4.92}N_{0.33}S_{0.01}Si_{0.20}$ | $C_{11.12}H_{21.10}O_{6.26}N_{0.33}S_{0.01}Si_{1.52}$ |
| H/C | 1.70 \pm 0.03 | 1.87 \pm 0.15 | 1.72 \pm 0.04 | 1.78 \pm 0.13 |
| O/C | 0.63 \pm 0.01 | 0.58 \pm 0.03 | 0.61 \pm 0.02 | 0.61 \pm 0.02 |
| N/C | 0.037 \pm 0.004 | 0.045 \pm 0.01 | 0.042 \pm 0.009 | 0.031 \pm 0.006 |
| OSc | -0.43 \pm 0.04 | -0.71 \pm 0.19 | -0.50 \pm 0.08 | -0.56 \pm 0.15 |
| CE3 Formula | $C_{10.57}H_{16.14}O_{5.08}N_{0.30}S_{0.01}Si_{0.02}$ | $C_{12.92}H_{21.78}O_{5.15}N_{0.42}S_{0.01}Si_{0.87}$ | $C_{10.42}H_{16.21}O_{5.09}N_{0.27}S_{0.004}Si_{0.11}$ | $C_{10.51}H_{16.18}O_{5.11}N_{0.27}S_{0.01}Si_{0.07}$ |
| H/C | 1.57 \pm 0.01 | 1.73 \pm 0.08 | 1.59 \pm 0.01 | 1.58 \pm 0.01 |
| O/C | 0.55 \pm 0.008 | 0.45 \pm 0.03 | 0.55 \pm 0.005 | 0.55 \pm 0.005 |
| N/C | 0.033 \pm 0.002 | 0.040 \pm 0.007 | 0.028 \pm 0.002 | 0.029 \pm 0.002 |
| OSc | -0.48 \pm 0.02 | -0.83 \pm 0.12 | -0.49 \pm 0.02 | -0.47 \pm 0.02 |
| CE4 Formula | $C_{8.17}H_{12.99}O_{5.29}N_{0.12}S_{0.004}Si_{0}$ | $C_{9.95}H_{14.53}O_{5.68}N_{0.32}S_{0.01}Si_{0}$ | $C_{8.94}H_{13.36}O_{5.52}N_{0.20}S_{0.004}Si_{0}$ | $C_{8.77}H_{13.20}O_{5.52}N_{0.18}S_{0.004}Si_{0}$ |
| H/C | 1.65 \pm 0.02 | 1.50 \pm 0.04 | 1.52 \pm 0.02 | 1.52 \pm 0.01 |
| O/C | 0.69 \pm 0.01 | 0.62 \pm 0.02 | 0.66 \pm 0.008 | 0.67 \pm 0.01 |
| N/C | 0.014 \pm 0.005 | 0.034 \pm 0.01 | 0.021 \pm 0.005 | 0.019 \pm 0.003 |
| OSc | -0.28 \pm 0.04 | -0.25 \pm 0.07 | -0.20 \pm 0.02 | -0.18 \pm 0.02 |

3.2 Comparison of cloud episodes

The fractions of three OA classes which are CHO, CHON, and Others (including CHONS, CHOS, CHN, CHS, CHNS, and CHOSi) exhibited general similarities across the four CEs, as shown in Fig. 3. In all sample types (pre-cloud aerosols, CDR, INT, and post-cloud aerosols) of the four CEs, CHO dominated OA composition, accounting for > 50 % of signal intensity of OA (54.6 %–85.7 % from CE1 to CE4), followed by CHON (14.0 %–33.2 %) and Others (lower than 18.7 %). The Others class, predominantly CHOSi, accounted for 0.5 %–18.7 % in CDR, exceeding the fractions in other sample types. In most cloud episodes, CDR showed the lowest CHO fraction (54.8 %–70.7 %) and the highest CHON fraction (26.6 %–33.2 %) among the four sample types. In CE2, the higher CHON (29.4 %) and lower CHO (54.6 %) in post-cloud aerosols than in other sample types are attributable to air mass changes during the long-time interval between post-cloud and others (shown in Fig. 2 and Fig. S1), as indicated by elevated CO concentration. CHON compounds have been detected in cloud and fog water in numerous studies (LeClair et al., 2012; Sun et al., 2024a; Sun et al., 2021; Sun et al., 2024b). Higher CHON fraction (59.0 %–63.5 %) in fog water than aerosol particles (51.2 %–51.5 %) was reported previously in Sun et al. (2024a), which is in agreement with the results in this study. In addition, the greater number of CHON compounds in CDR compared with CF underscores the role of cloud processing in enhancing CHON, as reflected in the number fraction rather than the intensity fraction (Boone et al., 2015; Liu et al., 2023b). The fraction of CHON (19.7 %–26.3 %) in INT was lower than in CDR (26.6 %–33.2 %) and higher than in pre-cloud aerosols (14.0 %–23.1 %) in CE1, CE2, and CE4. However, in CE3, the slightly lower CHON fraction in INT compared to pre-cloud aerosols may be due to turbulence in clouds resulting from the short duration time of the cloud (several minutes). A higher relative abundance of CHON in CDR (43.6 %–65.3 %) compared to INT (31.8 %–51.0 %) has been observed at Tianjing Mountain in southern China (Sun et al., 2021), consistent with our results. The higher CHON fraction in CDR than in pre-cloud aerosols suggests that cloud processing promoted CHON formation. Higher CHON in INT compared to pre-cloud aerosols indicates that although not activated into cloud droplets, high RH experienced by INT (close to 100 %) and corresponding high aerosol water content could still promote CHON formation in INT, consistent with the elevated N/C ratio of aqSOA of aerosol particles under high RH conditions (Zhao et al., 2019). We would like to note that it is assumed that different classes of compounds have similar sensitivity in EESI-ToF-MS.

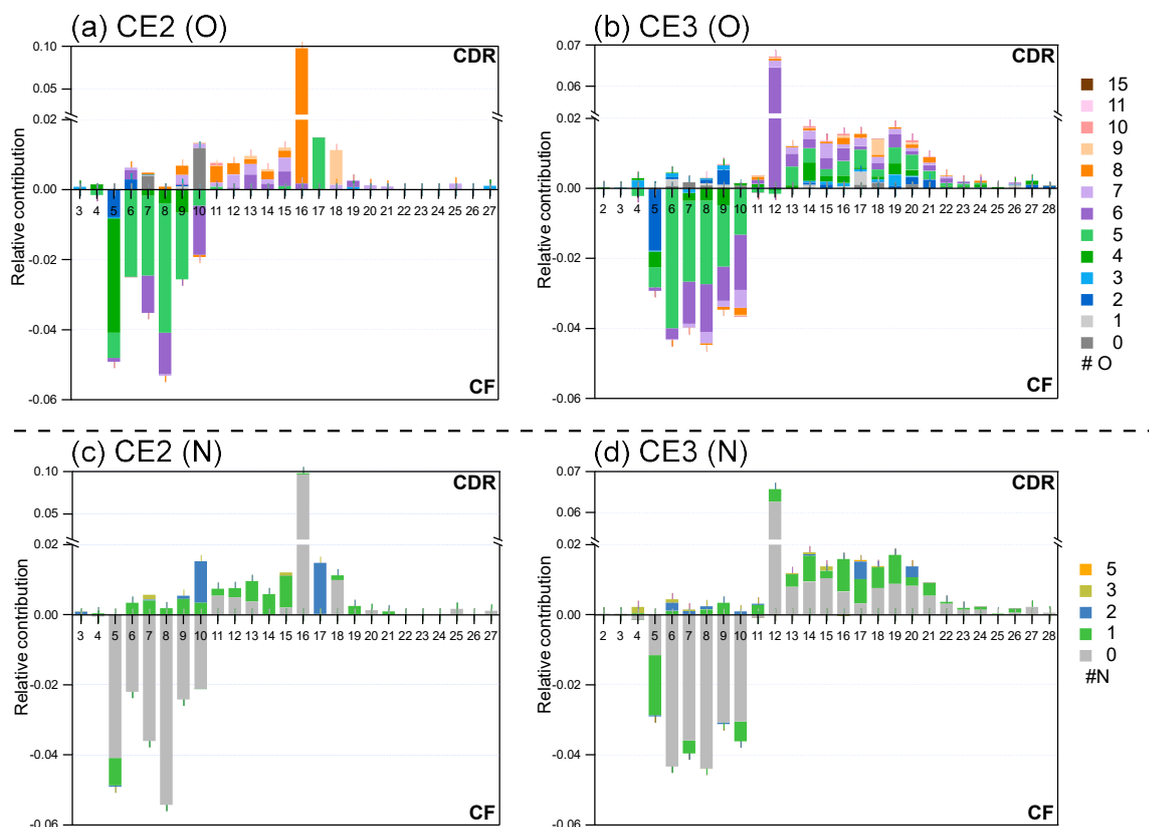
Among the CHON class, the compounds enriched in CDR, such as $C_{8-12}H_{11-19}NO_{5-8}$ and $C_{14-16}H_{21-27}NO_{4-9}$, with an O/N ratio of ≥ 3 (71 %–88 % for CE1, CE3, and CE4, and 17% for CE2), suggesting that they are likely organonitrates, amino acids, or N-containing heterocyclic compounds. At the Shanghuang site, emissions of monoterpenes and sesquiterpenes are abundant (Zhang et al., 2024). Consequently, $C_{10}H_{15}NO_x$ and $C_{10}H_{17}NO_x$ may be formed via hydroxyl oxidation of monoterpene in the presence of NO (Shen et al., 2022) or NO_3 oxidation (Shen et al., 2021; Guo et al., 2022) and dissolve in the aqueous phase, whereas $C_{15}H_{23}NO_x$ and $C_{15}H_{25}NO_x$ may originate from similar reactions involving sesquiterpenes. Additionally, precursors could form organonitrates through aqueous reactions, e.g., with NO_3 radicals (Ng et al., 2017), or involving NO_3^- (Sun et al.,

2024b; Huang et al., 2023; Barber et al., 2024). These reactions can occur at night or even during the day under reduced light conditions in clouds. This finding contrasts with the observation at Mt. Tai, where, despite the higher number of CHON compounds in CDR relative to CF, a larger fraction contained reduced nitrogen groups ($O/N < 3$) (Liu et al., 2023b). Such disparity may arise from differences in precursors between the two sampling sites. Additional information, such as the gas-phase CHON composition and concentration, is required to further elucidate the formation mechanisms of these compounds.



220 **Figure 3. Fractions of three classes of OA, namely CHO, CHON, and Others (including CHONS, CHOS, CHN, CHS, CHNS, and CHOSi) in pre-cloud aerosols, CDR, INT, and post-cloud aerosols in CE1, CE2, CE3, and CE4.**

The molecular composition characteristics of OA in four CEs exhibit similar patterns, presented as carbon number distribution colored according to the numbers of oxygen and nitrogen; therefore, only CE2 and CE3 are shown in Fig. 4 (CE1 and CE4 in Fig. S3). In each CE, comparison is carried out between CDR and CF which was the closest to CDR temporally: for CE1–3, CDR is compared with pre-cloud aerosols, while CE4 is compared with post-cloud aerosols, as shown in Fig. S1. In CDR, the carbon number of OA ranged from 2 to 28 (CE2: 3–27; CE3: 2–28), and the oxygen number ranged from 0 to 10 in CE2 and 0 to 15 in CE3. Comparing with OA in CF, OA in CDR contained a higher fraction of compounds with $n_C > 10$ as well as elevated n_O (CE2: $n_O=7-10$; CE3: $n_O=6-15$).



230 **Figure 4. Detailed relative contribution of OA. The average carbon number distribution of differences between CDR and CF are colored by oxygen number of (a) CE2, (b) CE3; and nitrogen number of (c) CE2, (d) CE3. Positive value stands for significant molecular characteristics of CDR, and negative value stands for that of CF. Fractions of compounds are normalized to sum of signals of all organics in CDR and CF, respectively.**

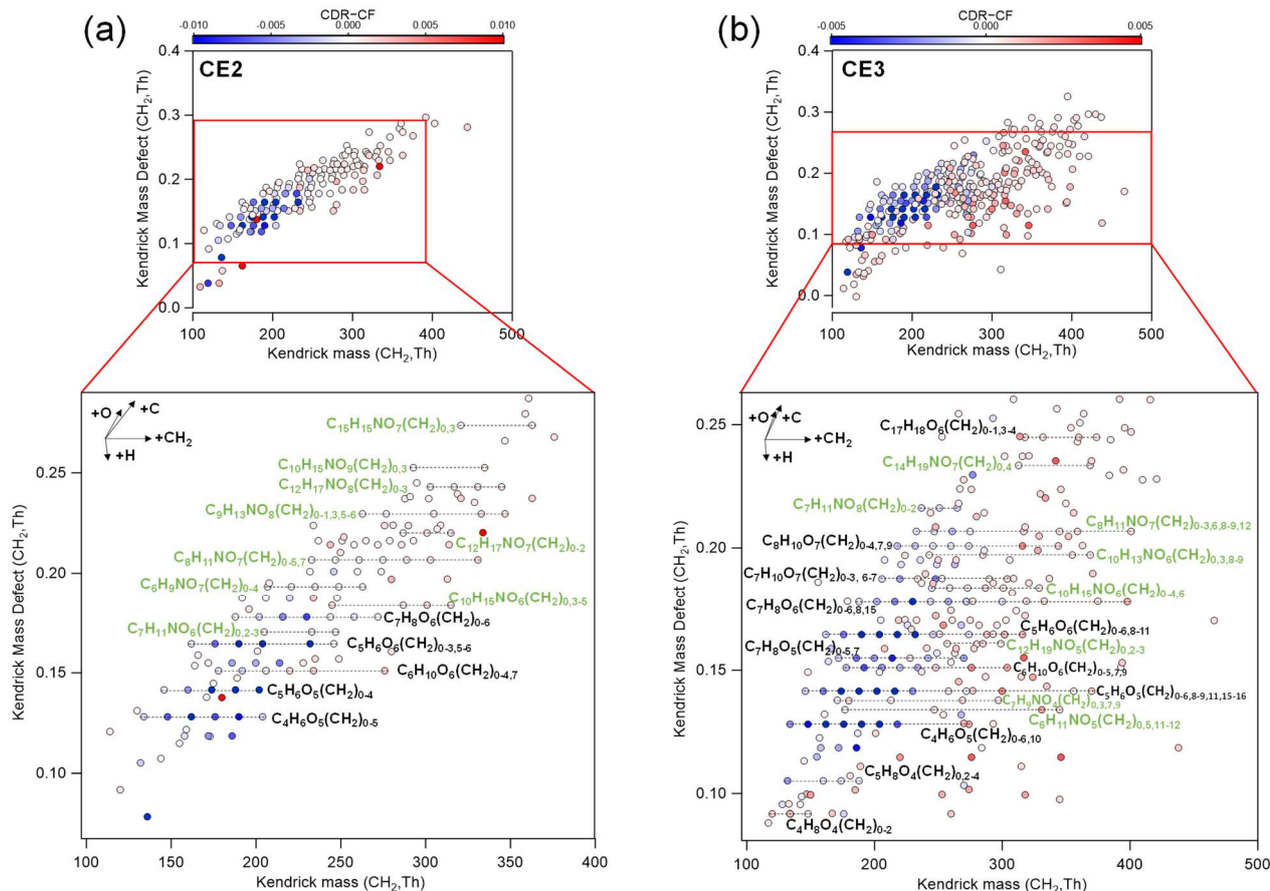
The nitrogen number (n_N) distributions relative to n_C exhibit similar patterns in all CEs. In CE2, the n_N of OA is distributed
 235 from 0 to 3, and from 0 to 5 in CE3. The n_C of N-containing OA ranged from 3 to 21 in CE2 and 4 to 27 in CE3. Compared with CF, CDR contained a larger fraction of N-containing OA, especially those with $n_N=1-3$ and higher n_C . Collectively, compounds in CDR had more n_C , n_O , and n_N than those in CF. The molecular characteristic of higher n_C is likely attributed to accretion reactions such as oligomerization (Yu et al., 2016; Fenselau et al., 2025). This finding is consistent with several laboratory studies of aqSOA formation. For instance, enriched high-molecular-weight compounds (HMWC) in aqSOA were
 240 reported in the bulk phase experiments of methylglyoxal and glyoxal under cloud-relevant conditions (Tan et al., 2009; Altieri et al., 2008). And aqSOA from in-cloud simulation using a wetted-wall flow reactor has more highly oxygenated and carbon-containing compounds than gasSOA simulated by an oxidation flow reactor (OFR) from the same biomass burning samples (Wang et al., 2024b). Experiments in the bulk phase and the wetted-wall flow reactor which better represents atmospheric aqueous conditions, indicate that accretion reactions could be prevalent in cloud droplets. Field observations in the Arctic also
 245 show potential evidence of accretion reactions, with compounds of longer carbon chains enriched in CDR relative to CF

(Pasquier et al., 2022), hinting at the possible importance of accretion reactions. Notably, this study provides direct molecular-level evidence for the contribution of accretion reactions during cloud processing of OA.

To investigate OA processing when the cloud episode changed from CF to CDR, CH₂-based Kendrick mass defect (KMD) plots for CE2 and CE3 are analyzed (Fig. 5). The chemical formulas of compounds with a larger fraction in CDR than CF in four CEs are listed in Table S3. Several series of compounds in CE2 and CE3 exhibit sequential increase in CH₂ groups, such as C₁₀H₁₅NO₆(CH₂)_n, C₈H₁₁NO₇(CH₂)_n, C₄H₆O₅(CH₂)_n, C₅H₆O₅(CH₂)_n, C₅H₆O₆(CH₂)_n, C₇H₈O₆(CH₂)_n. Specifically, numerous CHON compounds were present at higher fractions in CDR, with some labeled by formulas such as series of C₆H₉NO₇(CH₂)₀₋₄, C₇H₁₁NO₆(CH₂)_{0,2,3}, C₈H₁₁NO₇(CH₂)_{0-5,7}, and C₁₂H₁₇NO₈(CH₂)₀₋₃ in CE2 and series of C₇H₉NO₄(CH₂)_{0,3,7,9}, C₈H₁₁NO₇(CH₂)_{0-3,6,8-9,12}, C₁₀H₁₅NO₆(CH₂)_{0-4,6}, C₁₂H₁₉NO₅(CH₂)_{0,2-3} in CE3. This result is in agreement with the higher fraction of total CHON compounds in CDR compared with CF, as discussed above. The observed CH₂-based homologous series likely reflects carbon-chain growth through aqueous accretion reactions. Possible formation pathways include peroxy radical (RO₂) addition, aldol condensation, hydroxyl-carbonyl addition (hemiacetal/acetal formation), and esterification involving precursors (Tilgner et al., 2021; Mayhew et al., 2025), which warrant further investigation. For most homologues, CDR contained higher fractions of larger compounds (with more CH₂ groups) than CF, while lower fractions of smaller compounds. As detailed above, it is likely that cloud processing enhanced accretion reactions by extending the length of the carbon chain, which further highlights the importance of accretion reactions of organics in cloud droplets. In CF, CHO had a larger fraction than CHON; for example, CHO compounds such as C₅H₆O₆(CH₂)_n, C₆H₁₀O₆(CH₂)_n, and C₅H₆O₅(CH₂)_n were more abundant. The pattern of adding CH₂ groups in cloud processing is similar in all CEs. However, the KMD plots based on O show that compounds in CE2 and CE3 did not exhibit a clear pattern with a sequential increase in O (Fig. S4). The dominant pattern of CH₂ addition, rather than O addition, suggests that sequential OH addition or auto-oxidation was not prevalent in cloud processing. In terms of the increments of CH₂ and O, CH₂ displays a wider growth trend (0–7) among all series, whereas O shows a narrower increase, confined to a range of 0 to 3. Consequently, results of KMD plots suggest that as cloud processing proceeded, n_C of OA increases, while the increase in n_O is lower than the n_C, agreeing with the lower O/C ratio in CDR than that in CF. The possible reason is that aqueous processing is more significant in accretion (enhancing n_C) than oxygenation (enhancing n_O).

Furthermore, although oligomer formation involving subunits such as C₂H₂O₃ (Lim et al., 2010) and C₃H₄O₂ (Cook et al., 2017; Altieri et al., 2008; Tan et al., 2009) has been reported, compounds in CE of this study such as CE2 and CE3 did not seem to show clear sequential increases in these subunits (Fig. S5). This may be attributed to differences in precursors and formation mechanisms during cloud processing between the Shanghuang site and other observations and laboratory studies.

275 In addition, some siloxane compounds showed higher fractions in CDR than in CF. The reason for the higher fraction of siloxane warrants further study.



280 **Figure 5.** Kendrick mass defect plots based on CH_2 of compounds in (a) CE2 and (b) CE3. Data points are color-coded by differences in fractions of compounds between CDR and CF. Fractions of compounds are normalized to the sum of signals of all organics in CDR and CF, respectively. Note that, for conciseness, data points in CE3 with normalized signal difference between -0.0003 and 0.0003 (appearing nearly white) are not shown here. Siloxane compounds are not shown here for clarity.

3.3 Characteristic compounds in cloud processing and formation mechanisms

We identified enriched OA compounds in cloud droplet residuals by comparing the intensity fractions of all compounds between CDR and CF using a *t*-test at a significant level of 0.05. A total of 144, 421, 274, and 537 organic compounds in CE1, CE2, CE3, and CE4, respectively, passed the *t*-test. Among these compounds, 39 organic compounds in CDR were significantly enriched in three or four CEs, as shown in Table 2. Two were consistently significant in CDR across all four CEs: $\text{C}_{14}\text{H}_{42}\text{O}_7\text{Si}_7$ and $\text{C}_9\text{H}_{22}\text{N}_2\text{O}_4$. Furthermore, sulfate compounds were enriched in CDR compared with CF in three CEs, of which time series is shown in Fig. S6. Sulfate is a well-established tracer for aqueous-phase processing, and its elevated concentration in cloud droplets and fog has been widely reported (Dadashazar et al., 2022; Brege et al., 2018; Kim et al., 2019), which further enhances the potential of identifying the enriched OA compounds as aqSOA tracers formed via cloud processing. The number

290

of CHO, CHON, CHN, and CHOSi compounds is 15, 19, 2, and 3, respectively. The majority of the enriched OA compounds exhibit carbon numbers greater than nine, which is also an indication of accretion reactions in cloud droplets. Most of these enriched OA compounds have not been reported in previous literature (Cook et al., 2017; Bianco et al., 2019; Tong et al., 2021; Sun et al., 2024b).

Table 2. Thirty-nine enriched OA compounds observed in CDR of three or four CEs. These OA compounds are classified into four classes: CHO, CHON, CHN, and CHOSi.

| CHO | CHON | CHN | CHOSi |
|---|---|---|--|
| C ₆ H ₁₂ O ₆ | C ₄ H ₇ NO ₄ | C ₉ H ₁₈ N ₂ | C ₁₂ H ₃₆ O ₆ Si ₆ |
| C ₁₀ H ₁₆ O ₂ | C ₅ H ₁₁ N ₂ O | C ₁₂ H ₂₃ N | C ₁₄ H ₄₂ O ₇ Si ₇ |
| C ₁₀ H ₂₂ O ₄ | C ₉ H ₂₂ N ₂ O ₄ | | C ₁₆ H ₄₈ O ₈ Si ₈ |
| C ₁₂ H ₂₆ O ₅ | C ₉ H ₁₃ NO ₂ | | |
| C ₁₃ H ₂₂ O | C ₁₀ H ₁₉ NO | | |
| C ₁₃ H ₂₆ O ₅ | C ₁₀ H ₁₉ NO ₃ | | |
| C ₁₅ H ₂₄ O ₁₄ | C ₁₁ H ₁₉ NO ₄ | | |
| C ₁₅ H ₂₆ O ₈ | C ₁₃ H ₃₀ N ₂ O ₄ | | |
| C ₁₅ H ₃₂ O ₆ | C ₁₃ H ₂₃ NO ₃ | | |
| C ₁₆ H ₃₀ O ₄ | C ₁₄ H ₂₉ N ₃ O ₃ | | |
| C ₂₀ H ₂₂ O ₅ | C ₁₄ H ₂₉ NO ₄ | | |
| C ₂₁ H ₃₆ O ₈ | C ₁₈ H ₃₃ NO ₅ | | |
| C ₂₃ H ₄₄ O ₃ | C ₁₈ H ₂₉ NO ₅ | | |
| C ₂₄ H ₄₀ O ₃ | C ₁₉ H ₃₇ NO ₃ | | |
| C ₃₀ H ₅₆ O ₂ | C ₂₀ H ₂₉ NO ₅ | | |
| | C ₂₁ H ₄₁ NO ₂ | | |
| | C ₂₂ H ₃₄ N ₂ O ₆ | | |
| | C ₂₉ H ₅₁ NO ₂ | | |
| | C ₂₉ H ₅₁ NO ₆ | | |

Furthermore, 236 OA compounds were significantly enriched in two of four CEs, including the common aqSOA tracer, oxalic acid (C₂H₂O₄), previously reported in field observations and laboratory studies (Rogers et al., 2025; Ervens et al., 2011). The compound C₂O₄Na₃⁺ is identified as oxalic acid, of which the hydrogen atoms in the carboxylic functional group (-COOH) are substituted by Na⁺ (Surdu et al., 2024). The oxalic acid signal was exclusively observed during CDR, whereas it remained weak and noisy in CF and INT, as shown in Fig. 6. The oxalic acid signal was significantly enhanced only in CE2 and CE4, rather than in all CEs, which may be related to larger inhomogeneity within clouds due to strong turbulence. Meanwhile, the C₆H₁₂O₆ signal was as low as the detection limit in CF; however, it increased gradually when CDR began. C₆H₁₂O₆ in aqueous formation was reported in a laboratory study and may be produced from the aqueous reaction of formaldehyde or acetaldehyde (Li et al., 2011). Therefore, it is reasonable to classify C₆H₁₂O₆ as a tracer of aqSOA.

Notably, N-containing compounds were significant in CDR, such as C₁₂H₂₃N and C₇H₁₂N₂ (shown in Fig. 6). C₁₂H₂₃N was reported to be emitted from primary sources including vehicle emissions (Thomas et al., 2025) and agricultural residue burning (Lin et al., 2012), whereas C₇H₁₂N₂ (enriched in CE2) has been observed in emissions from traditional biomass fuel burning and agricultural residue burning (Fleming et al., 2018; Wang et al., 2017; Lin et al., 2012; Hao et al., 2025). C₁₂H₂₃N

and $C_7H_{12}N_2$ may be formed in aerosol phase and undergo uptake into cloud droplets. These compounds could also be heterocyclic compounds containing imine or amine functional groups, potentially resulting from secondary formation in the aqueous phase (Zhao et al., 2015; Li et al., 2023). In addition, $C_{12}H_{23}N$ may be a compound with a pyrrole structure. Pyrrole-derived SOA may contribute to brown carbon chromophore and influence radiative forcing (Chen et al., 2024). $C_7H_{12}N_2$ is likely 1-butyylimidazole, a derivative of imidazole, reported in reactions of methylglyoxal and amines in cloud simulation in De Haan et al. (2011). Moreover, imidazole has been reported as a type of brown carbon influencing regional radiative forcing (Kim et al., 2019; Lian et al., 2020; Gan et al., 2024) and may contribute to reactive oxygenated species, potentially relating to adverse health effects (Dou et al., 2015). The enhanced concentration of N-containing compounds in CDR could therefore have significant atmospheric implications and warrants further investigation.

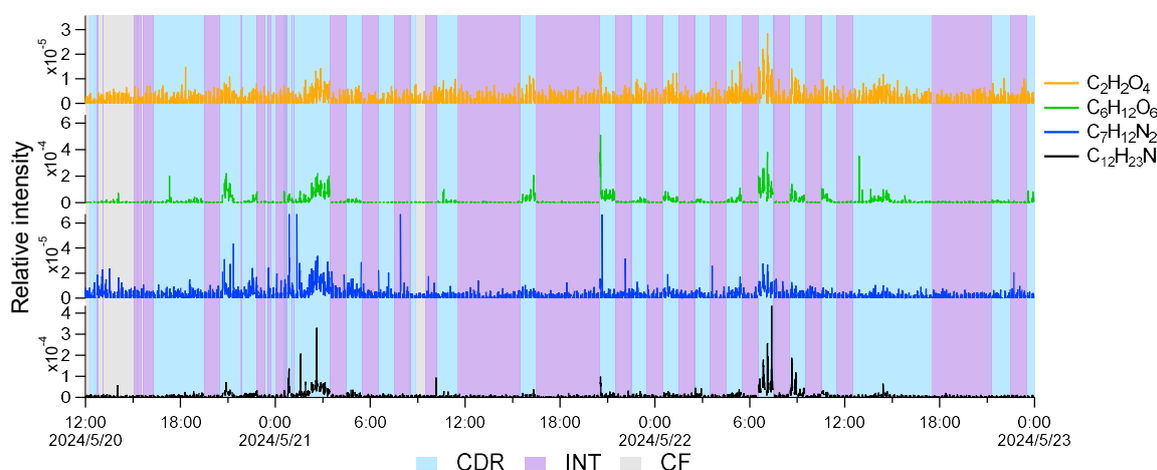


Figure 6. Typical time series of compounds: carboxylic acid ($C_2H_2O_4$), $C_6H_{12}O_6$, N-containing substances including $C_7H_{12}N_2$ and $C_{12}H_{23}N$, in a cloud episode. CDR, INT, and CF are shaded as blue, purple, and gray, respectively. Gaps in the time series represent EESI-ToF-MS background measurement periods, which are not shown.

3.4 Dynamic variation of OA in clouds

Relatively stable T, WS, and CO concentration in a typical 3-day cloud indicate that the cloud was stable and primary emission sources remained largely constant throughout the cloud episode. The time series of CHO, CHON, and Others are shown in Fig. 7. CHO and CHON were the major constituents for most of the episode, whereas Others was the lowest. The O/N ratio was generally lower in CDR than in INT, while the ratio of O/C and N/C varied irregularly in CDR and INT. Although the time resolution of our measurement (~ 20 s) is enough to capture the evolution of a compound in clouds, either in CDR or INT, there is no clear trend in the time series of the compounds, either from the fractions of OA classes or elemental ratios during the sample types of CDR or INT. This phenomenon is likely due to the dynamic characteristics of clouds, in

which turbulence and chemical processes continuously induce rapid changes in organic compounds, resulting in no gradual trends in their concentrations.

From the perspective of the molecular composition, the relative intensities of representative compounds in cloud episodes exhibited frequent and pronounced fluctuations during individual CDR periods, as shown in Fig. 6. Even within a 1-h CDR, the signal of compounds increased and decreased irregularly, likely due to turbulence. Consequently, it is difficult to track and capture information on the chemical transformation of OA in clouds. Most previous comparisons of the chemical composition of cloud droplets with cloud-free aerosol particles or interstitial aerosol particles are based on long sampling (hours to a day) and offline analysis (Sun et al., 2021; Liu et al., 2023b). Based on the findings in this study, the results obtained using methods with low time resolution may be subject to uncertainties due to the dynamic nature of clouds.

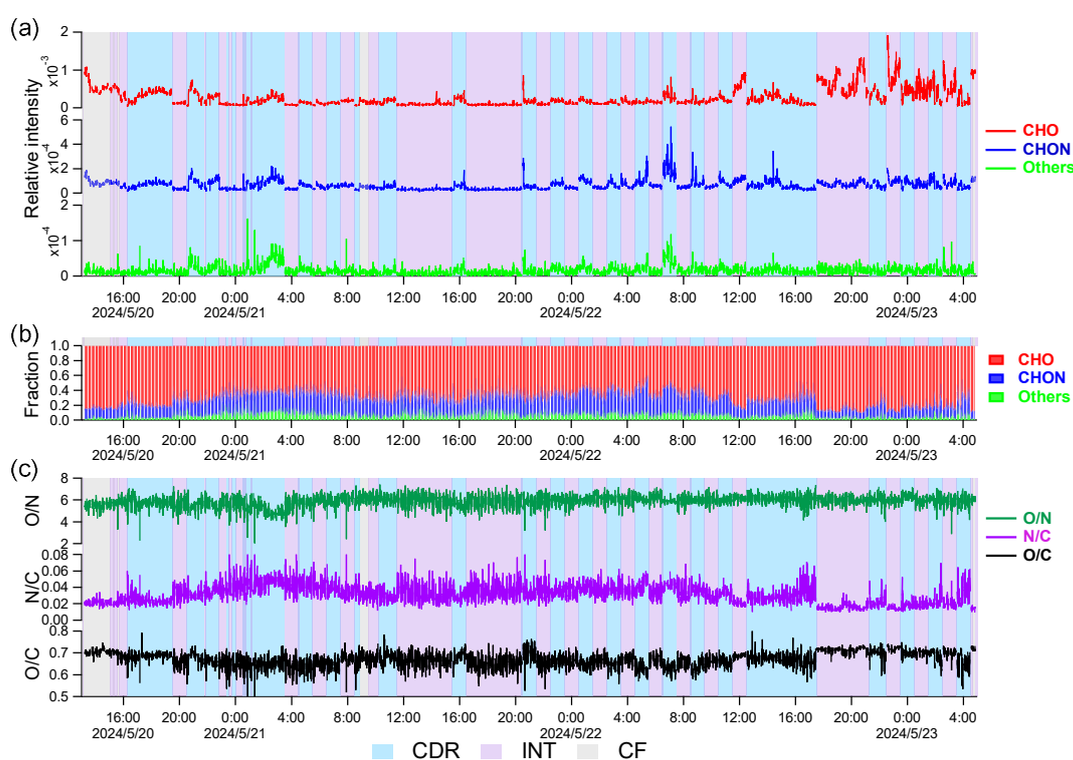


Figure 7. Time series of three classes of organic compounds (CHO, CHON, and Others) in a long cloud. (a) relative intensity, (b) fraction in OA, (c) O/N, N/C, and O/C ratio of OA. CDR, INT, and CF are shaded in blue, purple, and gray, respectively. Gaps in the time series represent EESI-ToF-MS background measurement periods, which are not shown.

350 4 Conclusions and implications

AqSOA molecular composition and processing in cloud episodes were studied using online molecular information obtained by EESI-ToF-MS at a high-mountain site in China. Cloud processing substantially influences OA composition, resulting in large differences among distinct cloud episodes. Organics in cloud droplet residuals had an average molecular formula $C_{9.95-12.92}H_{14.53-21.78}O_{5.15-6.02}N_{0.32-0.42}S_{0-0.01}Si_{0-1.29}$ for the selected four cloud episodes. CHO compounds contributed

355 predominantly to OA in cloud droplet residuals. CHON was enhanced markedly in cloud droplet residuals compared with cloud-free aerosol particles and interstitial aerosol particles in most cloud episodes. The majority of CHON compounds were likely organonitrates, highlighting the enrichment of organonitrates compounds in cloud processing. OA in cloud droplet residuals contained higher numbers of C, O, and N atoms, exhibited a CH₂-based homologous series, and showed an enrichment of higher-molecular-weight compounds compared with aerosol particles sampled under the temporally closest
360 cloud-free conditions, collectively highlighting the importance of accretion reactions in cloud processing of OA at the molecular level. We identified several compounds significantly enriched in cloud droplet residuals, including typical aqSOA tracers such as oxalic acid. The new aqSOA tracers, such as C₆H₁₂O₆ and C₉H₂₂N₂O₄, could help future studies identify cloud processing aqSOA.

This study provides direct molecular-level evidence for the contribution of accretion reactions during cloud processing
365 of OA. Although previous cloud observations using FT-ICR-MS reported the presence of oligomers in cloud samples, these studies could not distinguish whether such compounds originated from cloud processing or aqueous aerosols, as no concomitant aerosol samples were collected for comparison (Zhao et al., 2013; Cook et al., 2017). By directly comparing OA composition in cloud droplet residuals with that in cloud-free aerosol particles, our results clearly demonstrate that accretion reactions occur within cloud droplets. It has been assumed that HMWC are predominantly formed in aerosol liquid water
370 rather than cloud water, owing to the lower reaction rates of accretion reactions in the more dilute cloud-water environment (Ervens et al., 2011). In contrast, our study provides direct molecular-level evidence that such compounds can also be formed in cloud water, extending earlier observations by Cook et al. (2017). These findings highlight that accretion reactions should be considered when modeling aqSOA formation in clouds.

The HMWC formed via accretion reaction may have implications for the environment and climate. Due to the increase
375 in the HMWC, accretion reactions likely reduce the volatility of organics and could potentially enhance OA mass concentration and alter the aerosol size distribution after cloud evaporation. The formation of HMWC can also modify physicochemical properties, such as lifetime, oxidation state, viscosity, and hygroscopic properties, which may further influence the cloud activation of these aerosols. In addition, the formation of N-containing compounds in cloud droplet residuals, such as organonitrates, pyrrole, and imidazole, may also affect the physicochemical properties of aqSOA, e.g., contributing to brown
380 carbon and thus affecting regional radiative forcing.

Based on the measurement of high time resolution (~20 s), we find that the concentrations of individual organic compounds were highly dynamic in clouds, which is likely due to the turbulence in clouds. Such a highly dynamic nature in clouds poses difficulties in extracting the influence of chemical processes on individual compounds for instrumentation with low temporal resolution. Therefore, our results highlight the necessity of high time resolution measurements (< 1 h), especially

385 online systems achieving minute-level resolution to investigate the chemical processes in clouds, considering dynamic variations of compounds in clouds due to turbulence in clouds and alterations in air masses.

It should be noted that this study provides molecular formulas only, while detailed structural information is warranted to better constrain the sources, formation mechanisms, and climate impacts of aqSOA in clouds. In addition, sources of compounds enriched in cloud droplet residuals will be investigated in future studies.

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Data availability. The data used in this study are available from the corresponding authors upon request: Defeng Zhao (dfzhao@fudan.edu.cn).

Supplement.

Author contributions. DZ conceptualized the research. YJ conducted the measurements with the aid of HL, DZ, ST, SX, and CN. XiaocP and GZ conducted GCVI measurements. XiaolP conducted the meteorological measurements. WX, YZ, YS, QC and LL provided support for sampling and operation of the Shanghuang site. YJ processed data and wrote the manuscript. YJ and DZ edited the manuscript with the inputs of all authors.

Competing interests. Qi Chen is a member of the editorial board of ACP.

Financial support. This work is supported by the National Natural Science Foundation of China (No. 42575109), Shanghai Pilot Program for Basic Research-Fudan University (No. 21TQ1400100 (22TQ010)), the National Natural Science Foundation of China (No. 42330605), and “Island Atmosphere and Ecology” Category IV Peak Discipline.

Acknowledgement. The authors gratefully acknowledge the NOAA Air Resources Laboratory (ARL) for the provision of the HYSPLIT transport and dispersion model and/or READY website (<https://www.ready.noaa.gov>) used in this publication.

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