Molecular characterization of organic aerosol in the Himalayas: insight from ultra-high-resolution mass spectrometry

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Received: 7 July 2018 – Discussion started: 20 August 2018
Revised: 28 November 2018 – Accepted: 8 January 2019 – Published: 29 January 2019

Abstract. An increased trend in aerosol concentration has been observed in the Himalayas in recent years, but the understanding of the chemical composition and sources of aerosol remains poorly understood. In this study, molecular chemical composition of water-soluble organic matter (WSOM) from two filter samples collected during two high aerosol loading periods (denoted as P1 and P2) at a high-altitude station (Qomolangma Station, QOMS; 4276 m a.s.l.) in the northern Himalayas was identified using electrospray ionization Fourier transform ion cyclotron resonance mass spectrometry (ESI-FTICR MS). More than 4000 molecular formulas were identified in each filter sample which were classified into two compound groups (CHO and CHON) based on their elemental composition, with both accounting for nearly equal contributions in number (45–55%). The relative abundance weighted mole ratio of O/Cw for P1 and P2 was 0.43 and 0.39, respectively, and the weighted double bond equivalents (DBEw), an index for the saturation of organic molecules, were 7.12 and 7.87, respectively. Although the O/Cw mole ratio was comparable for CHO and CHON compounds, the DBEw was significantly higher in CHON compounds than CHO compounds. More than 50% molecular formulas in the Van Krevelen (VK) diagram (H/C vs. O/C) were located in 1–1.5 (H/C) and 0.2–0.6 (O/C) regions, suggesting potential lignin-like compounds. The distributions of CHO and CHON compounds in the VK diagram, DBE vs. number of C atoms, and other diagnostic diagrams showed high similarities among each other, suggesting their similar source and/or atmospheric processes. Many formulas formed from biogenic volatile organic compounds (e.g., ozonolysis of α-pinene products) and biomass-burning-emitted compounds (e.g., phenolic compounds) were found in the WSOM, suggesting the important contribution of these two sources in the Himalayas. The high DBE and high fraction of nitrogen-containing aerosol can potentially impact aerosol light absorption in this remote region. Further comprehensive study is needed due to the complexity of organic aerosol and limited molecular number identified in this study.

1 Introduction

Relatively high aerosol concentration events have been frequently observed over the Himalayas during the premonsoon period (March to June) (Bonasoni et al., 2010). The aerosol plumes are originated from the southern regions of the Himalayas such as northwestern India and/or the Indo-Gangetic Plain based on air mass back trajectory analysis and satellite observation (Liu et al., 2008; Lu et al., 2012; Lüthi et al., 2015). Due to increased consumption on fuels (including biofuels and fossil fuels) by industry and residents in recent decades, air pollution has been a serious issue in south Asia (Gustafsson et al., 2009). Under favorable atmospheric circulation, air pollutants emitted or formed in these regions can be transported fast to the Himalayas and Tibetan Plateau (HTP; Xia et al., 2011).
Enhanced aerosol concentration for the remote region of the HTP is thought to have many negative climate and environmental effects. For example, the transported aerosol could heat the air in the higher layer of the troposphere over the HTP and impact the monsoon system of south Asia and accelerate the melting of glaciers in the Himalayas (Lau et al., 2006; Ramanathan et al., 2007). This heating effect is predominantly from light-absorbing particulate aerosol (LAPA) such as black carbon (BC) and brown carbon which are types of organic aerosol (OA; Ram et al., 2010; Zhang et al., 2015, 2017). BC is from incomplete combustion and dominates the absorption of LAPA; brown carbon can originate from primary emission and/or secondary processes, and has made an increasing contribution (up to ~20%) to light absorption in recent years (Laskin et al., 2015, and reference therein). Due to the light absorption of brown carbon being strongly dependent on their molecular structure, light-absorbing compounds at a molecular level have been explored during recent years, and it was found that nitrogen-containing compositions are important brown carbon compounds (Lin et al., 2016, 2017). Many studies show that biomass burning is an important source of brown carbon (e.g., Saleh et al., 2014; Washenfelder et al., 2015), which is very common in developing regions in the southern Himalayas. In comparison with other regions, high elevation and mixed biomass fuels in the southern Himalayas could make the evolution and chemical composition of OA from biomass burning emission more complicated (Stockwell et al., 2016; Fleming et al., 2018; Jayaratne et al., 2018).

The details on the molecular composition of OA are important for understanding the sources and chemical evolution of OA (Laskin et al., 2018). Previous studies conducted in the HTP have focused on a limited number of molecular markers such as organic acids, which are closely related to biomass burning emission (Cong et al., 2015), and some toxicology species such as polycyclic aromatic hydrocarbons (PAHs) and persistent organic pollutants (POPs), which are related to anthropogenic activities (Wang et al., 2015, 2016). In addition, online measurement using the Aerodyne high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS) has provided more details on the chemistry and sources of OA with a high time resolution (Xu et al., 2018). However, different instruments have limitations on OA detection, and an ultra-high mass resolution of mass spectrometry which can identify many molecular formulas is lacking.

Fourier transform-ion cyclotron resonance mass spectrometry (FTICR MS) coupled with a soft ionization source, such as electrospray ionization (ESI), can be used to identify the individual molecular formula of extremely complex mixtures because of its ultra-high resolution and mass accuracy (Mazzoleni et al., 2010). Similar methods have been used for the identification of components in aqueous secondary OA (SOA) and in ambient samples, and allow the identification and separation of thousands of compounds in a sample (e.g., Mazzoleni et al., 2010; Altieri et al., 2012; Mead et al., 2013). Types of methods such as double bond equivalents (DBEs), elemental ratios, and Kendrick mass defects (KMDs) can be applied to deduce the chemical characterization of obtained molecular formulas. In this study, we focus on the molecular composition of water-soluble organic compound in fine particle aerosol in the Himalayas using ESI-FTICR MS and evaluate the sources, chemical processing, and potential impact of aerosol in this region.

2 Methodology

2.1 Aerosol sampling

A field study was conducted at the Qomolangma Station (QOMS; 28.36° N, 86.95° E; 4276 m a.s.l.) located at the toe of Mt. Qomolangma from 12 April to 12 May 2016 using a suite of online instruments (X. Zhang et al., 2018), including a HR-ToF-AMS (Aerodyne Research Inc., Billerica, MA, USA) for 5 min size-resolved chemical compositions (organics, sulfate, nitrate, ammonium, and chloride) of non-refractory submicron particulate matter (NR-PM$_1$) and a photoacoustic extinction meter (PAX; DMT Inc., Boulder, CO, USA) for BC mass concentration. In addition, a low-volume (16.7 L min$^{-1}$) particulate matter (PM) sampler (BGI, USA, model PQ 200) with an Aerodyne diameter cutoff of 2.5 μm at the inlet was used to collect PM$_{2.5}$ filter samples on pre-baked quartz fiber filters (47 mm, Pall Life Science, NY, USA). Due to the low aerosol loading at this remote region, a 2-day sampling strategy was adopted for each filter collection from 08:00 to 07:45 the following day (local time). A total of 18 filter samples were collected during the field study with three procedure blanks which were used to assess potential contamination during sampling and transportation. The sampling air volume ranged from 35.1 to 48.1 m$^3$ at ambient conditions. Two samples collected during 25–27 April (P1) and 29 April–1 May (P2), respectively, were used in this study due to the relatively higher aerosol loading based on HR-ToF-AMS results (Sect. 3.1) and distinct particulate matter on the filter. One procedure blank was also adopted in this study like that of aerosol samples to subtract the potential background.

2.2 Chemical analysis

For FTICR MS analysis, filter samples were extracted in 20 mL Milli-Q water in an ultrasonic bath for 30 min and filtered using 0.45 μm pore-size Acrodisc syringe filters to remove water-insoluble matter (Pall Science, USA). The sample tubes were immersed in the mixture of ice–water during ultrasonic extraction to prevent potential chemical reaction. Prior to FTICR MS analysis, the extraction was concentrated and purified using PPL (Agilent Bond Elut-PPL cartridges, 500 mg, 6 mL) solid-phase extraction (SPE) cartridges for water-soluble organic matter (WSOM) to avoid possible ESI artifacts. PPL cartridges generally have the best
properties for WSOM enrichment for subsequent FTICR MS analysis (Raeke et al., 2016). In addition, we control the concentration of SPE effluent to be ~0.2 mg mL⁻¹, which was not too concentrated for artifact adducts. Note that through SPE cartridges, the most hydrophilic compounds, such as inorganic ions, and low-molecular-weight organic molecules, such as organic acids and sugars, were removed, whereas the relatively hydrophobic fraction was retained. Details on the SPE method using PPL cartridges and analysis using FT-ICR MS can be found in our previous paper (Feng et al., 2016).

Briefly, the mass spectrometry analyses of these samples were performed using a Solarix XR FTICR MS (Bruker Daltonik GmbH, Bremen, Germany) equipped with a 9.4 T refrigerated actively shielded superconducting magnet (Bruker Biospin, Wissembourg, France) and a Paracell analyzer cell. The samples were ionized in positive ion modes using the ESI ion source (Bruker Daltonik GmbH, Bremen, Germany).

A typical mass-resolving power of > 400000 was achieved at m/z 400, with an absolute mass error of < 0.5 ppm. The ions detected in filter blank were subtracted and molecular formulas in the samples were assigned to all ions with signal-to-noise ratios of greater than 10, with a mass tolerance of ±1.5 ppm, using custom software. Molecular formulas with their maximum numbers of atoms were defined as 30 12C, 60 1H, 20 16O, 3 14N, 1 32S, 1 13C, 1 18O, and 1 34S. Identified formulas with H/C, O/C, N/C, S/C, and DBE/C, ranging 0.5–3.0, 0.3–3.0, 0–0.5, 0–0.2, and 0–1, were selected, and formulas containing isotopomers (i.e., 13C, 18O, or 34S) were not considered. Compounds were detected as either sodium adducts, [M + Na]⁺, or protonated species, [M + H]⁺. Although ammonium could also readily be adduct to assist in the formation of positive ions, this possibility in our sample was because of the low concentration of ammonium (Fig. 1). We report all detected compounds as neutral species, unless stated otherwise. Note that the ESI+ mode more easily detects basic functional group compounds, and the reported organic molecules here are only part of organic aerosol which are biased ionized in ESI+ mode. In addition, highly functionalized compounds could be detected in both positive and negative modes (Lin et al., 2012).

2.3 Data processing

The assigned molecular formulas were examined using the Van Krevelen diagram (Wu et al., 2004), DBEs, KMD series, and aromatic indices (AI mod). The O/C and H/C ratios were calculated by dividing the number of O and H atoms, respectively, by the number of C atoms in a formula. DBE analysis was used to determine the number of rings and double bonds in a molecule. The DBE was calculated using Eq. (1):

\[
DBE = 1 + c - h/2 + n/2.
\]

where c, h, and n are the numbers of C, H, and N atoms, respectively, in the formula.

Figure 1. The average mass concentration (mass avg.) and chemical composition of PM1 during P1 (a) and P2 (b) periods, respectively, measured by the HR-ToF-AMS and PAX. Note that the compounds of PM1 include sulfate, nitrate, ammonium, chloride, BC (black carbon), organics, BBOA (biomass-burning-emitted OA), NOA (nitrogen-containing OA), and OOA (oxidized oxygenated OA).

The weighted DBE (DBEw), O/C (O/Cw), and H/C (H/Cw) were calculated using Eq. (2):

\[
X_w = \frac{\sum (w_i \times X_i)}{\sum w_i},
\]

where \(X_i\) and \(w_i\) are the parameters above and the relative intensity (RI) is for each individual formula, \(i\).

The Kendrick mass (KM) and KMD for CH2 series, used to search for potential oligomeric units (Hughey et al., 2001), were calculated using Eqs. (3) and (4):

\[
KM = \text{observed mass} \times \frac{14}{14.01565},
\]

\[
KMD = \text{NM} - \text{KM},
\]

where 14 is the nominal mass (NM) of CH2, 14.01565 is the exact mass of CH2, and NM is KM rounded to the nearest integer.

Furthermore, a second-order mass defect analysis using the base units of CH2 and H2 was applied following the method described in Roach et al. (2011), which greatly simplified the visualization of complex mass spectra.

\[AI_{mod} = \frac{1 + c - 0.5a - 0.5h - 0.5n}{(c - 0.5a - n)},\]

where c, a, and h are the numbers of C, O, H, and N atoms in the formula. \(AI_{mod}\) ranges from 0 for a purely aliphatic compound to higher values for compounds with more double bonds and that are more aromatic.

3 Results and discussions

3.1 Chemical characterization of PM1 during P1 and P2 measured by the HR-ToF-AMS

The average mass concentration and chemical composition measured by the HR-ToF-AMS during P1 and P2 periods are

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The air mass back trajectory analysis using the HYSPLIT model (Draxler and Hess, 1998) during Figure 2. The air mass trajectories were recovered back to 72 at a 1 h interval from the sampling site (QOMS) at 1000 m above ground level using a 1° resolution Global Data Assimilation System (GDAS) dataset (https://ready.arl.noaa.gov/gdas1.php; last access: 10 September 2017). The cluster analysis for these trajectories was completed based on the directions of the trajectories (angle distance) and colored according to air pressure (vertical profile). The firespot observed from MODIS (https://firms.modaps.eosdis.nasa.gov; last access: 10 September 2017) and the average wind rose plot colored by wind speed (WS) for during each filter sampling period are also shown. The firespot is sized by fire radiative power.

shown in Fig. 1. The mass concentrations of PM$_1$ were 9.2 and 10.6 µg m$^{-3}$, respectively, which were at the high range of all filters (1.3–10.6 µg m$^{-3}$) because of a continuous long-range transport event at the QOMS (X. Zhang et al., 2018). Due to our sample processing error, the mass concentration of filter measured gravimetrically could not be used, and thus the fractions of PM$_1$ to PM$_{2.5}$ are not available. However, most of WSOM in PM$_{2.5}$ is in the accumulation size mode (less than 1 µm) which could be detected by the HR-ToF-AMS (Zhang et al., 2005). The chemical composition of PM$_1$ during P1 and P2 was all dominated by OA (55% and 57%), followed by BC (26% and 22%), sulfate (7% and 8%), nitrate (5% and 6%), and ammonium (5% and 6%). The OA was comprised of biomass-burning-emitted OA (BBOA), nitrogen-containing OA (NOA), and more oxidized oxygenated OA (MO-OOA) decomposed using positive matrix factorization (PMF) analysis (Fig. 1). Details on PMF analysis can be found in X. Zhang et al. (2018). The mass contribution of BBOA was higher during P2 than P1 (32% vs. 22%), whereas the contribution of MO-OOA was higher during P1 (24% vs. 16%). The mass spectra of OA for these two filter periods were closely similar to a Poisson correlation efficiency ($r$) of 0.9. The elemental ratios of oxygen (O) to carbon (C) of OA were 1.04 and 0.97 for P1 and P2 periods (Improved Ambient method; Canagaratna et al., 2015), respectively, and the ratios of hydrogen (H) to carbon (C) of OA (BBOA), nitrogen-containing OA (NOA), and more oxidized oxygenated OA (MO-OOA) decomposed using positive matrix factorization (PMF) analysis (Fig. 1). Details on PMF analysis can be found in X. Zhang et al. (2018). The mass contribution of BBOA was higher during P2 than P1 (32% vs. 22%), whereas the contribution of MO-OOA was higher during P1 (24% vs. 16%). The mass spectra of OA for these two filter periods were closely similar to a Poisson correlation efficiency ($r$) of 0.9. The elemental ratios of oxygen (O) to carbon (C) of OA were 1.04 and 0.97 for P1 and P2 periods (Improved Ambient method; Canagaratna et al., 2015), respectively, and the ratios of hydrogen (H) to carbon (C) of OA were 1.26 and 1.32 accordingly. These suggest that the OA during P2 was relatively less oxidized than that during P1 ($t$ test, $p < 0.05$). The six category ions ($C_xH_y^+_z$, $C_xH_yO_z^+$, $C_xH_yO_2^+$, $C_xH_yN_z^+$, $C_xH_yO_2N_z^+$, and HO$^+$) detected by the HR-ToF-AMS for these two filter periods were all dominated by $C_xH_yO_2^+$, followed by $C_xH_y^+$, $C_xH_yO_2^+$, $C_xH_yN_z^+$, $C_xH_4O_2N_z^+$, and HO$^+$. The air mass trajectory analyses using the hybrid single particle Lagrangian integrated trajectory (HYSPLIT) model for P1 and P2 periods show air mass mainly originated from west and southwest of the QOMS across north and northwest India where there were many firespots during these two periods (Fig. 2). The air mass during P2 was partly (13%) transported with a low wind speed and across a short distance (less than 100 km), and could contain some fresh OA as illustrated by the higher fraction of BBOA.

3.2 The chemical characteristics of WSOM from ESI-FTICR MS

A total of 4295 and 4770 molecular formulas were identified by ESI-FTICR MS over the mass range of 100–700 Da for P1 and P2, respectively. The identified molecular formulas were grouped into two subgroups based on their elemental composition, i.e., CHO and CHON, all of which had equally important contributions (45%–55%) in number (Fig. 3). Note that individual species in the ESI-FTICR MS mass spectra could have many different isomeric structures, in which case the percentages only reflect the number of unique molecular formulas in each category. The mass spectra of these two samples were highly similar in the distributions of molecular formulas (Fig. 3). The average weighted element ratios of P1 and P2 were 0.43 vs. 0.39 for O/C$_w$, 1.36 vs. 1.31 for H/C$_w$, and 1.72 vs. 1.68 for OM/OC$_w$ (Table 1), suggesting a relatively higher oxidation and saturation degree for P1 than P2. These trends are consistent with the results of the HR-ToF-AMS, although the elemental ratios are different between them, which is due to the difference in the detection

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Table 1. Chemical characterization of all molecular assignments detected in WSOM for P1, P2, and common ions. Relative intensity weighted ("w") for each data subset (O/C, H/C, OM/OC, DBE, and DBE/C) is given.

|               | All   | CHO   | CHON |
|---------------|-------|-------|------|
| P1            |       |       |      |
| O/C_w         | 0.43  | 0.44  | 0.42 |
| H/C_w         | 1.36  | 1.37  | 1.34 |
| OM/OC_w       | 1.72  | 1.70  | 1.76 |
| DBE_w         | 7.12  | 6.74  | 7.79 |
| DBE/C_w       | 0.39  | 0.37  | 0.43 |
| Percentage (%) of aliphatic (AI_{mod} = 0) | | | |
| Olefinic (0.5 > AI_{mod} > 0) | | | |
| Aromatic (AI_{mod} ≥ 0.5) | | | |
| P2            |       |       |      |
| O/C_w         | 0.39  | 0.39  | 0.39 |
| H/C_w         | 1.31  | 1.32  | 1.31 |
| OM/OC_w       | 1.68  | 1.63  | 1.73 |
| DBE_w         | 7.87  | 7.38  | 8.32 |
| DBE/C_w       | 0.42  | 0.40  | 0.44 |
| Percentage (%) of aliphatic (AI_{mod} = 0) | | | |
| Olefinic (0.5 > AI_{mod} > 0) | | | |
| Aromatic (AI_{mod} ≥ 0.5) | | | |
| Common ions   |       |       |      |
| O/C_w         | 0.42  | 0.43  | 0.42 |
| H/C_w         | 1.35  | 1.36  | 1.33 |
| OM/OC_w       | 1.72  | 1.69  | 1.76 |
| DBE_w         | 7.18  | 6.75  | 7.79 |
| DBE/C_w       | 0.40  | 0.38  | 0.43 |
| Percentage (%) of aliphatic (AI_{mod} = 0) | | | |
| Olefinic (0.5 > AI_{mod} > 0) | | | |
| Aromatic (AI_{mod} ≥ 0.5) | | | |

Table 2. Chemical characterization of the molecular assignments detected in aerosol samples from selected studies (adapted and modified from Table 3 in Dzepina et al., 2015). Note that all values are presented as arithmetic means, which are given for convenience of comparison. The data for our study are the arithmetic means for P1 and P2.

| Sample type       | Measurement site | Instrument               | O/C     | H/C     | OM/OC   | DBE     | DBE/C   | Reference            |
|-------------------|------------------|--------------------------|---------|---------|---------|---------|---------|-----------------------|
| Aerosol           | Remote           | ESI(+)-FTICR MS          | 0.39–0.42 | 1.30–1.34 | 1.68–1.72 | 7.71–8.38 | 0.41–0.42 | This study            |
| Aerosol           | Free troposphere | ESI(−)-FTICR MS          | 0.42–0.46 | 1.17–1.28 | 1.67–1.73 | 9.4–10.7  | 0.42–0.47 | Dzepina et al. (2015)  |
| Aerosol           | Remote           | ESI(−)-FTICR MS          | 0.53 ± 0.2 | 1.48 ± 0.3 | 1.91 ± 0.3 | 6.18 ± 3.0 | NA       | Mazzoleni et al. (2012) |
| Aerosol           | Rural            | ESI(−)-FTICR MS          | 0.28–0.32 | 1.37–1.46 | 1.54–1.60 | 6.30–7.45 | 0.33–0.38 | Wozniak et al. (2008)  |
| Aerosol           | Suburban         | ESI(−)-FTICR MS          | 0.46     | 1.34     | 1.85     | 5.3      | 0.45     | Lin et al. (2012)      |
| Aerosol           | Urban            | ESI(+)-FTICR MS          | 0.31     | 1.34     | NA       | 8.68     | 0.41     | Choi et al. (2017)     |
| Aerosol           | Marin boundary layer | ESI(−)-FTICR MS    | 0.35     | 1.59     | 1.67     | 4.37     | 0.28     | Schmitt-Kopplin et al. (2012) |
| Aerosol           | Marine boundary layer | ESI(−)-FTICR MS    | 0.36–0.42 | 1.49–1.56 | 1.70–1.74 | 5.88–6.76 | 0.28–0.32 | Wozniak et al. (2014)  |
| Cloud water       | Remote           | ESI(−)-FTICR MS          | 0.61–0.62 | 1.46     | 2.06–2.08 | 6.29–6.30 | 0.38     | Zhao et al. (2013)     |
| Cloud water       | Rural            | ESI(−)-FTICR MS          | 0.51     | 1.47     | NA       | 6.03     | NA       | Cook et al. (2017)     |
| Fog water         | Rural            | ESI(−)-FTICR MS          | 0.43     | 1.39     | 1.77     | 5.6      | 0.40     | Mazzoleni et al. (2010) |

NA: not available

The elemental ratios of WSOM from ESI-FTICR MS in our study are similar to those results observed in aerosol samples in remote sites using ESI-FTICR MS (e.g., 0.35–0.53 for O/C) (Table 2). The O/C and H/C in Van Krevelen diagrams (Wu et al., 2004) for these two filters and the subgroup molecular formulas show similar distributions, and are all concentrated in ranges of 1.2–1.8 for H/C and 0.3–0.7 for O/C (Fig. 3), suggesting their similar aerosol sources and atmospheric processes. The similar distributions for these two filters are also observed in plots of KMD vs. KM and DBE vs. C (Fig. 3).

Structural information for the assigned molecular formulas is inferred from the DBE_w value which was higher for P2 than for P1 (7.87 vs. 7.12) (Table 1). Compared with other studies, the DBE_w values in our filter are relatively close to the results from biomass burning aerosol and aerosol samples from remote sites (Table 2) (Dzepina et al., 2015). The DBE_w
values for each molecular subgroup were higher for CHON than for CHO (Table 1), especially for P2 (8.32 vs. 7.38), suggesting more rings and double bonds in CHON molecular formulas. The AI\text{mod}, reflecting the minimum number of carbon–carbon double bonds and rings (Koch and Dittmar, 2006), was correspondingly higher in P2 as illustrated by its higher contribution of olefinic (75.0 % vs. 73.9 % for P2 and P1) and aromatic compounds (10.3 % vs. 7.7 % for P2 and P1) (Table 1). For aromatic compounds (AI\text{mod} ≥ 0.5) in P2, 52 % of them were CHON formulas (45 % for P1). Higher DBE and AI\text{mod} values in CHON compounds suggest more unsaturated compounds in them which could contain a certain number of chromophores. The distribution of DBE vs. the carbon number of two filters showed a systematic increase in a concentrated region and a high similarity with each other. This similarity further suggests the consistent source and chemical processes for the aerosol of these two filters.

There were 3955 common molecular formulas between these two filters with the number contribution of CHO of 50.7 % and CHON of 49.3 %. These common molecular formulas accounted for 92.1 % (P1) and 82.9 % (P2) of two filters, respectively. There were 340 unique molecular formulas in P1, with 73 % being CHO compounds, whereas there were 815 unique molecular formulas in P2, with 65.3 % being CHON compounds. For more confidence in molecule assignment, we focus on the common molecular formulas detected in these two samples in the section below. Note that the mass spectrum of common ions was calculated from the average RI from two mass spectra and normalized to the highest peak.

3.3 The potential sources and formation processes

3.3.1 CHO compounds

CHO compounds have been frequently detected in ambient aerosol samples (Altieri et al., 2009b; Mazzoleni et al., 2010; Lin et al., 2012; Fleming et al., 2018), which could comprise high molecular weight humic-like substances (HULIS) or oligomers, and in primary emission or secondary formation of different aerosol sources (Mazzoleni et al., 2012; Wozniak et al., 2014; Lin et al., 2016; Cook et al., 2017). In our samples, the weighted molecular weight of CHO compounds was 361.9, with an average C atom of 19.3 ± 5.3 per molecule; the most abundant O atoms were located in the range 5–10, with an average value of 7.8 ± 2.9 per molecule (Fig. 4a and b). The oxygen distribution is also evidenced by the longest homologous series in the second-order mass defect analysis (CH\textsubscript{2}-H\textsubscript{2}), which were all O\textsubscript{3}-O\textsubscript{10} compounds (Fig. 4d). The DBE of CHO increased with carbon number, with a DBE\textsubscript{w} value of 6.75 (Fig. 3); the carbon-normalized DBE (DBE/C) was 0.39 ± 0.14. These two values were close to the results from biomass burning aerosol samples in other studies and are in a high range of published data (Table 2) (Lin et al., 2012; Mazzoleni et al., 2012), suggesting relatively high aromaticity in our samples. The carbon oxidation state (OS\textsubscript{C}) values (Kroll et al., 2011), a useful metric for the degree of oxidation of organic species in the atmosphere, exhibited between −1 and 0 for high number density formulas, with 25 or fewer carbon atoms, suggesting that they are semi- and low-volatile organic compounds, corresponding to “fresh” (BBOA) and “aged” (LV-OOA) SOA by multistep oxidation reactions (Fig. 4c).

There are several possible sources and chemical formation pathways for high-oxygen-containing CHO compounds. Ozonolysis of α-pinene has been found to form highly oxygenated molecules with some important products such as C\textsubscript{17}H\textsubscript{26}O\textsubscript{8} (m/z 358) and C\textsubscript{19}H\textsubscript{28}O\textsubscript{7} (m/z 368), which
are possible esterification products of *cis*-pinic (C_{9}H_{14}O_{4}; m/z 186) and diaterpenylic acid (C_{9}H_{14}O_{5}; m/z 190) (Kristensen et al., 2013). The first three formulas were all found in our common CHO molecules with high relative abundance (Table 3). The appearance of these formulas together with high relative abundance gave us more confidence in the products of ozonolysis of α-pinene. Ozone concentration in the Himalayas during the pre-monsoon period was the highest based on online measurement at the Nepal Climate Observatory at Pyramid (NCO-P) during 2006–2008 (61 ± 9 ppbv) (Cristofanelli et al., 2010). Biogenic volatile organic compounds could be transported from the low elevation regions in subtropical India and biogenic secondary organic aerosol has been found to be an important source in the Himalayas (Stone et al., 2012). A number of other previously reported monoterpene oxidation product formulas were also observed in our study (Table 3) (Claeys et al., 2007; Kleindienst et al., 2007; H. Zhang et al., 2018). In addition, some biomass burning aerosol markers were also found in CHO compounds. Sun et al. (2010) and Yu et al. (2014, 2016) observed that aqueous-phase oxidation of lignin produces phenol (C_{6}H_{6}O), guaiacol (C_{7}H_{8}O_{2}), and syringol (C_{8}H_{10}O_{3}), yielding a substantial fraction of dimers and higher oligomers, with key dimer markers identified as C_{16}H_{18}O_{6} and C_{14}H_{14}O_{4}. The dimer markers C_{16}H_{18}O_{6} and C_{14}H_{14}O_{4} were also present in our sample with high RI (5.5 % and 27.5 %). The high relative intensity of these compounds indicates that fog and cloud processing of phenolic species (biomass burning aerosol) could be an important mechanism for the production of low-volatility SOA in the Himalayas. Compounds observed in biomass burning emission (cow dung and brush wood) during residential cooking in Nepal were also found in our samples (Fleming et al., 2018).

3.3.2 CHON compounds

The frequency distribution for \( n_\text{O} \) and \( n_\text{C} \) in CHON formulas was shown in Fig. 5a, which show peaks between 6 and 10 and 15 and 20, respectively. The DBE of CHON formulas ranged from 4 to 10, with DBE\(_\text{w}\) being 7.79 (Fig. 5b and Table 1). In the CHON class, compounds contained one or two nitrogen (1N or 2N) atoms, with 1N compounds accounting for 70.5 % and 2N for 29.5 %, respectively. Most (93.6 %) 1N compounds contained more than 3 oxygen atoms and up to 13 oxygen atoms, whereas about 62.5 % of 2N compounds contained more than 6 oxygen atoms (Fig. 6a). The average O atoms contained in each molecular formula were therefore higher for 1N compounds than 2N compounds (8.1 ± 2.9 vs. 6.3 ± 2.3). The ratios of O/C\(_w\) and O/SOC\(_w\) for 1N compounds were accordingly higher than those for 2N compounds (0.42 vs. 0.37 for O/C\(_w\) and −0.48 vs. −0.54 for O/SOC\(_w\), respectively), suggesting a higher oxidation state for 1N compounds (Fig. 5c). In contrast, the DBE\(_w\) and AI\(_\text{mod}\) values for 2N compounds were higher than for 1N compounds (Table 1). The higher H/C\(_w\) for 2N compounds (Fig. 5d) suggests that 2N compounds could contain many aromatic N-heterocyclic compounds. For 1N compounds, longer and higher relative intensity CH\(_2\) homologous series compounds were found based on the Kendrick mass defect
Many CHON compounds could be identified in ESI+ mode, such as reduced N functional compounds and nitroaromatic compounds (Altieri et al., 2009a; Laskin et al., 2009; Lin et al., 2012; O’Brien et al., 2013; Wang et al., 2017). Laskin et al. (2009) identified the amount of N-heterocyclic alkaloid compounds from types of fresh biomass burning aerosol. Lin et al. (2012) and Wang et al. (2017) also identified many CHON compounds in fresh and aged biomass burning aerosol. Oxygenated organic nitrogen compounds in ambient aerosol (Dzepina et al., 2015), rainwater (Altieri et al., 2009a), and fog water (Mazzoleni et al., 2010) from biomass-burning-emission-influenced regions were also observed. Lin et al. (2017) found aged biomass burning aerosol in the present urban oxidants (such as NOx) could result in a higher fraction of CHON compounds compared to the fresh biomass burning aerosol. Considering the high influence of biomass burning emission in the Himalayas (X. Zhang et al., 2018), the CHON compounds in our study were probably related to biomass burning emissions. Recent studies have proven that burning of mixed biomass fuels in Nepal could emit amounts of nitrogen species such as NH3, NOx, HCN, benzene, and organics, and the emission factors for these species are higher than for wood (Stockwell et al., 2016; Jayarathne et al., 2018). In addition, it is likely that smoldering burning of bio-fuels in high elevation areas is also re-

### Table 3. List of identified BVOC products and biomass-burning-emission-related compounds in this study.

| Mass (m/z) | Molecular formula | Relative intensity | Compound type | References |
|-----------|-------------------|--------------------|---------------|------------|
| 358.1622  | C17H26O8          | 33.1%              | α-Pinene products | Kristensen et al., 2013 |
| 368.1829  | C19H28O7          | 11.5%              |               |            |
| 186.0887  | C9H14O4           | 26.0%              |               |            |
| 231.0737  | C9H13NO6          | 12.7%              |               |            |
| 233.0894  | C9H15NO6          | 14.0%              |               |            |
| 215.1152  | C10H17NO4         | 10.6%              |               |            |
| 229.0945  | C10H15NO5         | 16.5%              |               |            |
| 231.1101  | C10H17NO5         | 16.8%              |               |            |
| 233.1257  | C10H19NO5         | 9.5%               |               |            |
| 245.0894  | C10H15NO6         | 26.4%              |               |            |
| 247.1050  | C10H17NO6         | 16.5%              |               |            |
| 249.1207  | C10H19NO6         | 9.6%               |               |            |
| 306.1099  | C16H18O6          | 5.5%               | Aqueous-phase oxidation of biomass burning aerosol | Yu et al. (2014, 2016), Sun et al. (2010) |
| 246.0886  | C14H14O4          | 27.5%              |               |            |
| 166.0624  | C9H10O3           | 4.9%               |               |            |
| 178.0624  | C10H10O3          | 5.7%               |               |            |
| 190.0624  | C11H10O3          | 7.7%               |               |            |
| 192.0781  | C11H12O3          | 7.0%               |               |            |
| 188.0832  | C12H12O2          | 4.3%               | Cow dung and brush wood burning | Fleming et al., 2018 |
| 216.0781  | C13H12O3          | 12.5%              |               |            |
| 218.0937  | C13H14O3          | 16.7%              |               |            |
| 198.0866  | C13H14O4          | 30.5%              |               |            |
| 248.1043  | C14H16O4          | 32.2%              |               |            |
| 232.1094  | C14H16O3          | 21.6%              |               |            |
| 338.0745  | C14H14N2O8        | 5.7%               |               |            |
| 341.1105  | C15H19N1O8        | 27.2%              | Biomass burning aerosol | Song et al. (2018) |
| 355.1261  | C16H21N1O8        | 29.4%              |               |            |
| 369.1418  | C17H23N1O8        | 24.0%              |               |            |
| 243.0890  | C14H13N1O3        | 9.3%               |               |            |
| 257.1046  | C15H15N1O3        | 9.5%               |               |            |
| 235.1203  | C13H17N1O3        | 17.0%              |               |            |
Figure 5. The molecular information for common CHON compounds. (a) The number frequency distribution of carbon ($n_c$) and oxygen ($n_o$). (b) The 3-D plot for $n_o$, $n_c$, and double bond equivalents (DBEs) colored by their relative intensity. (c) Scatter plot of carbon-based oxidation state (OSc) vs. $n_c$ colored by the distribution of number of molecules. (d) The Van Krevelen diagram by H/C vs. N/C colored by number of oxygen ($n_o$). The different sizes of dot marker in (c) and (d) represent the 1N and 2N compounds.

Figure 6. (a) The number frequency distribution of $n_o$ for 1N and 2N compounds and (b) the longest 10 CH$_2$ homologous series compounds in 1N compounds.

Sonsible for the presence of many nitrogen-containing compounds in BBOA (Chen et al., 2010). Nitroaromatic compounds such as methyl nitro catechols (C$_7$H$_7$NO$_4$) are introduced to be tracers for biomass burning secondary organic aerosols (Iinuma et al., 2010). Although the C$_7$H$_7$NO$_4$ formula is not found in our measurement, C$_{14}$H$_{14}$N$_2$O$_8$ was found in our measurement, which was probably its dimer formula. In addition, the homologous series compounds in which C$_7$H$_7$NO$_4$ serves as the core molecule were also found in our samples. Some high relative abundance CHON molecular formulas identified in a recent paper from biomass burning aerosol were also found in our measurement (Table 3) (Song et al., 2018).

Besides primary emission and/or secondary formation from biomass burning emission, nitrogen-containing OA could also be formed through other chemical processes. For example, biogenic volatile organic compounds (BVOCs) can react with NO$_3$ radicals or RO$_2$ + NO into organic nitrate (Ng et al., 2017). Although organic nitrate is not favored to be ionized in positive ESI-MS (Wan and Yu, 2006), organic nitrate formed from BVOC could be highly functionalized (Lee et al., 2016) and ionized in positive MS through other alkaline functional groups. Recent studies have shown that BVOCs, including isoprene (C$_{5}$H$_{8}$) and monoterpenes (C$_{10}$H$_{16}$), dominate the organic nitrate formation in the southeastern United States under the condition of the mixed anthropogenic NO$_x$ and BVOCs (Xu et al., 2014; Lee et al., 2016; H. Zhang et al., 2018). Several molecular formulas formed from monoterpene and NO$_3$ radicals were found in our study (Table 3) (Lee et al., 2016; H. Zhang et al., 2018).
4 Implications and limitations

This study analyzed the WSOM using ESI(+) FTICR MS in fine particulate aerosol from the Himalayas and found that the molecular compositions of WSOM were mainly comprised of two group compounds (CHO and CHON) with equally important contributions. The two compound groups could be originated from biomass burning emission and BVOC oxidation products because many markers for these two sources were found in these molecular compounds. All our compounds had relatively high DBE values, suggesting a potential high light absorption feature. Due to the relative higher mass concentration and higher contribution of nitrogen-containing compounds (9% of PM$_1$) during these two periods based on HR-ToF-AMS results (X. Zhang et al., 2018), it is believed that aerosol transported to the Himalayas plays an important role in atmospheric radiative forcing.

Ramanathan and Carmichael (2008) found a distinct warming effect of light-absorbing aerosol over the Himalayas through estimating aerosol radiative forcing by BC. However, brown carbon had not been considered before, which could also be an important light-absorbing aerosol due to its high mass loading (Laskin et al., 2015). Zhang et al. (2017) estimated the light absorption contribution of brown carbon from the inland Tibetan Plateau, which was up to $\sim$13% of that of BC. The high DBE and nitrogen-containing OA in our study suggested aerosol in the Himalayas could also contain amounts of light-absorbing organic matter because the light absorption properties of organic molecules are closely related to the number of double bonds and rings in the molecule and nitrogen atoms. Many studies have found that the dominating chemical molecules in brown carbon were related to nitrogen-containing aerosol (e.g., Lin et al., 2016). This kind of aerosol combined with BC could have a higher radiative forcing than before in this area.

More comprehensive methods are needed in the future for identifying BrC in the Himalayas due to the chemical complexity of BrC. For example, BrC extraction is highly dependent on the solvent used, and water-insoluble OA can contribute higher light absorption than water-soluble OA (Chen and Bond, 2010). In addition, Budisulistiorini et al. (2017) found that a number of compounds can dominate the light absorption of BrC, although they make a minor contribution to the aerosol mass. Therefore, it is important to know the exact chromophores of BrC which can be obtained by combining light absorption measurement with a photodiode (PDA) detector and chemical composition with a high-resolution mass spectrometer with high performance liquid chromatography (HPLC-PDA-HRMS system) (Lin et al., 2016). For mass spectrometry analysis, different ionization sources which are also favorable for different compounds, such as ESI, only detect a part of polar compounds; non-polar compounds which could dominate the contribution of BrC can be measured using atmospheric pressure photo ionization (Lin et al., 2018). A recent study indicated that over 40% of the solvent-extractable BrC light absorption is attributed to water-insoluble, non-polar to semi-polar compounds such as PAHs and their derivatives (Lin et al., 2018). In contrast, the polar, water-soluble BrC compounds, which are detected in ESI, account for less than 30% of light absorption by BrC (Lin et al., 2018).

Data availability. The processed FTICR-MS data, AMS data, and meteorological data in this study are available upon request from the corresponding author (jzxu@lzb.ac.cn).

Author contributions. YA analyzed the data and wrote the manuscript. JX and SK organized the campaign, analyzed the data, and wrote the manuscript. LF, BJ, and YL organized the mass spectrometry analysis. XZ analyzed AMS data and YL collected the filter.

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

Special issue statement. This article is part of the special issue “Study of ozone, aerosols and radiation over the Tibetan Plateau (SOAR-TP) (ACP/AMT inter-journal SI)”. It is not associated with a conference.

Acknowledgements. This research was supported by grants from the National Natural Science Foundation of China (41771079, 41421061), the Key Laboratory of Cryospheric Sciences Scientific Research Foundation (SKLCS-ZZ-2018), and the Chinese Academy of Sciences Hundred Talents Program. The authors thank the QOMS for logistic support.

Edited by: Wenshou Tian
Reviewed by: three anonymous referees

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