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Atmospheric organic matter in clouds: exact masses and molecular formula identification using ultrahigh-resolution FT-ICR mass spectrometry

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Abstract. Clouds alter the composition of atmospheric aerosol by acting as a medium for interactions between gas- and particulate-phase substances. To determine the cloud water atmospheric organic matter (AOM) composition and study the cloud processing of aerosols, two samples of supercooled clouds were collected at the Storm Peak Laboratory near Steamboat Springs, Colorado (3220 m a.s.l.). Approximately 3000 molecular formulas were assigned to ultrahigh-resolution mass spectra of the samples after using a reversed-phase extraction procedure to isolate the AOM components from the cloud water. Nitrogen-containing compounds (CHNO compounds), sulfur-containing compounds (CHOS and CHNOS compounds) and other oxygen-containing compounds (CHO compounds) with molecular weights up to 700 Da were observed. Average oxygen-to-carbon ratios of ∼0.6 indicate a slightly more oxidized composition than most water-soluble organic carbon identified in aerosol studies, which may result from aqueous oxidation in the clouds. The AOM composition indicates significant influences from biogenic secondary organic aerosol (SOA) and residential wood combustion. We observed 60% of the cloud water CHO molecular formulas to be identical to SOA samples of α-pinene, β-pinene, d-limonene, and β-caryophyllene ozonolysis. CHNO compounds had the highest number frequency and relative abundances and are associated with residential wood combustion and NOx oxidation. Multiple nitrogen atoms in the assigned molecular formulas for the nighttime cloud sample composite were observed, indicating the significance of nitrate radical reactions on the AOM composition. Several CHOS and CHNOS compounds with reduced sulfur (in addition to the commonly observed oxidized sulfur-containing compounds) were also observed; however further investigation is needed to determine the origin of the reduced sulfur-containing compounds. Overall, the molecular composition determined using ultrahigh-resolution Fourier-transform ion cyclotron resonance (FT-ICR) mass spectrometry provides an unambiguous identification of the cloud water organic anion composition in the Rocky Mountain area that could help to improve the understanding of aqueous-phase processes.

1 Introduction

Cloud and fog droplets play a major role in the processing of atmospheric organic matter (AOM) (Blando and Turpin, 2000; Collett et al., 2008; Mazzoleni et al., 2010; Ervens et al., 2011; Herckes et al., 2013). Under conditions of high relative humidity, aerosol particles may act as cloud condensation nuclei (CCN), where water vapor condenses onto them to form droplets. Suspended droplets are subject to gas–liquid equilibrium of volatile compounds and solid–liquid equilibrium of non-volatile compounds (Graedel and Weschler, 1981), and thus new transport and transformation pathways affecting the composition of AOM may occur (Collett et al., 2008; Lim et al., 2010; Ervens et al., 2011). Volatile water-soluble compounds like glyoxal and methylglyoxal can readily transfer into the aqueous phase.
of droplets due to their high effective Henry’s law constant (e.g., $H_{\text{eff}} = 3.710 \times 10^3 \text{M atm}^{-1}$ at 25°C for methylglyoxal (Betterton and Hoffmann, 1988)). Similarly, organic compounds in CCN aerosol can partition into the aqueous phase. Since water-soluble organic carbon (WSOC) may comprise up to 90% of the total organic carbon mass in aerosols (Samburova et al., 2013) and the particle activation process usually favors particles with a large fraction of water-soluble components (Facchini et al., 1999), a large fraction of the WSOC may partition to the aqueous phase. Thus a considerable number of species, either from the gas phase or the particle phase, can be found in the liquid phase of the droplets.

The multiphase environment of the droplets facilitates aqueous-phase reactions of the water-soluble species. A number of laboratory experiments mimicking wet aerosols showed that precursors like glyoxal or methylglyoxal react with inorganic species like ammonium nitrate or ammonium sulfate to produce a complex mixture of light-absorbing organic compounds (Carlton et al., 2007; De Haan et al., 2009; Galloway et al., 2009; Perri et al., 2009; Shapiro et al., 2009; Volkamer et al., 2009; Sareen et al., 2010; Sun et al., 2010). Likewise, a number of laboratory experiments have demonstrated the production of secondary organic compounds under cloud-relevant conditions (Tan et al., 2010; Yasmeen et al., 2010; Bateman et al., 2011), indicating cloud processing of AOM. In fact, in-cloud production of oxalic acid (Crahan et al., 2004), organic acids (Sorooshian et al., 2010; Sun et al., 2009; Völkmann et al., 2009; Laskin et al., 2009; Schmitt-Kopplin et al., 2010; Lin et al., 2010) and HULIS found in aerosols (Havers et al., 1998; Graber and Rudich, 2006) were also recently observed in clouds (Feng and Möller, 2004; Deecesari et al., 2005; Reyes-Rodríguez et al., 2009). For example, Reyes-Rodríguez et al. (2009) observed the high-molecular-weight organic compounds in cloud samples to be mostly aliphatic and oxygenated, with a small amount of aromatic compounds based on study of the functional groups. Overall, the previous studies either observed several individual compounds or observed functional groups rather than the complex mixture of the organic compounds on a molecular level.

Electrospray ionization (ESI) coupled with Fourier-transform ion cyclotron resonance (FT-ICR) mass spectrometry provides detailed molecular characterization of organic matter due to its extremely high resolution and mass accuracy (Marshall et al., 1998; Kujawinski, 2002; Sleight and Hatcher, 2007). ESI is a soft ionization method that minimizes the fragmentation of analytes, allowing for detection of intact molecules, while the ultrahigh resolution and high accuracy of FT-ICR allows for assignment of a unique chemical formula to each peak detected. FT-ICR mass spectrometry has been successfully applied to ambient fog water samples (Mazzezoni et al., 2010; LeClair et al., 2012), rainwater samples (Altieri et al., 2009a, b, 2012), and aerosols (Reemtsma et al., 2006; Wozniak et al., 2008; Laskin et al., 2009; Schmitt-Kopplin et al., 2010; Lin et al., 2012b; Mazzezoni et al., 2012). This powerful analytical approach was applied in this study to characterize the complex mixture of higher molecular weight water-soluble organic compounds in ambient cloud water samples collected at the Storm Peak Laboratory (SPL). Here we present the results of ultrahigh-resolution FT-ICR mass spectrometry of two supercooled cloud samples collected in February 2010.

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The comprehensive characterization of the cloud water AOM composition after negative ion ESI FT-ICR mass spectrometry will be discussed.

2 Methods

2.1 Cloud water sample collection

To investigate the composition of cloud water AOM, samples of ambient clouds consisting of supercooled droplets were collected at the SPL. The remote continental site is located on the western summit of Mt. Werner (3220 m a.s.l.) in the Park Range Colorado near Steamboat Springs. The mountain-top laboratory facilitates observations of free-tropospheric air and in-cloud conditions. SPL is situated at the tree line on a 70 km ridge oriented perpendicular to the prevailing westerly winds (Hallar et al., 2013). SPL experiences transport from distant sources including urban areas, power plants, and wildfires (Obrist et al., 2008). There are also abundant monoterpene emissions from the pine trees at SPL during summer months (Amin et al., 2012). Clouds that surround the SPL with supercooled droplets form in the wintertime (Lowenthal et al., 2002). A custom cloud sieve with stainless steel strands of 0.5 mm diameters was used to collect supercooled droplets by impaction (Fig. S1). As described by Hindman et al. (1992), cloud droplets with diameter > 8 µm will be collected with a 50 % collection efficiency on the 0.5 mm strand diameter with an average wind speed of 2 m s$^{-1}$ (average wind speed during sample collection time periods). Cloud sieves were mounted on the SPL rooftop deck railing on the west side during cloud events. After ~20 min, the cloud sieves were carried into a cold laboratory space to remove the frozen cloud water (collected as rime) from the strands using a stainless steel blade and collection tray. The collected rime was stored in a freezer at $-5 \degree C$. Four cloud water samples were collected during short separate cloud events. The first sample (CW1) is a composite of two samples collected between 20:30 and 21:00 LT on 24 February 2010 and 07:00 and 08:00 LT on 25 February 2010. The second composite sample (CW2) is a composite of two samples, collected between 08:00 and 11:00 LT on 26 February 2010. The 24 h air back trajectory analyses showed the air coming from the west and northwest of SPL (Figs. S2 and S3). The particle concentrations (with diameter > 3 nm) were around 1000 cm$^{-3}$ during the sampling time determined by the condensation particle counter (TSI, model #3025). One field blank sample was prepared by leaving the cloud sieve in the air in the absence of a cloud event and then rinsed with high-purity water.

2.2 Sample preparation

Cloud water samples were prepared for analysis using a reversed-phase solid-phase extraction (SPE) method to concentrate the analytes and remove inorganic ions (Mazzoleni et al., 2010). The Strata-X (Phenomenex) cartridges, with both hydrophilic and hydrophobic functional groups, were found to obtain a diverse spectrum of analytes. Strata-X SPE cartridges were conditioned with consecutive application of the following: 1 mL of isopropyl alcohol, 2 mL of acetonitrile, 2 mL of acidified methanol-containing 0.1 % formic acid, and 2 mL of aqueous 0.1 % formic acid. Then 100 mL of cloud water without pH adjustment was applied at a rate of 1–2 mL min$^{-1}$ to the cartridge to allow the AOM components to partition to the SPE stationary phase. pH values of the cloud water samples were not determined in this study. Hindman et al. (2006) measured the pH values of wintertime clouds collected at Storm Peak Laboratory from 1983 to 2004 and reported a pH range of 3.7 to 5.7; thus the cloud samples in this study are expected to be acidic. The cartridges were rinsed with high-purity acidified water to remove salts and subsequently dried. Some of the low-molecular-weight analytes (< 100 Da) are expected to be lost in this step (Sambugrova et al., 2013). Analytes were eluted with 1.5 mL of acetonitrile. Extraction recoveries were not determined in this study, however a large fraction of the high-molecular-weight cloud water organic carbon is expected to be present due to the both hydrophilic and hydrophobic characteristics of the Strata-X cartridges. The field blank and lab blank were prepared using the same method except that 100 mL of high-purity water without pH adjustment was applied to the SPE cartridge for the lab blank. All solvents were HPLC grade or higher. Prepared samples were stored at $-5 \degree C$ until further analysis.

2.3 Instrumental parameters

Ultrahigh-resolution mass spectrometry analysis was performed using a hybrid linear ion trap FT-ICR mass spectrometer (LTQ FT Ultra, Thermo Scientific) equipped with an ESI source. Cloud water samples were infused directly into the ESI interface, and the ESI parameters were adjusted to obtain a stable ion current with minimum ion injection time into the mass analyzer. After optimization, the infusion flow rate was 4 µL min$^{-1}$, the ESI needle voltage was $-3.7 \, kV$, and the capillary temperature was 265 \degree C. Besides the instrumental parameters, the electrospray efficiencies of the analytes depend on their chemical structures. Analytes with hydroxyl or carboxyl functional groups favor ionization in the negative ion electrospray mode. Many water-soluble organic species in clouds contain these hydrophilic functional groups, and thus most of the cloud water organic carbon species are expected to be observed in the negative ion mass spectra, though not all the organic carbon species are ionized. The sample delivery apparatus was flushed in between each analysis with a minimum of 500 µL of acetonitrile and water (50 : 50) and 500 µL of 100 % acetonitrile until background noise levels were reached. Negative ion mass spectra were collected using full-scan analysis for the range of 100 < m/z < 1000. Mass resolving power, m / Δm$^{50\%}$, in which Δm$^{50\%}$
is peak full width at half-maximum peak height, was set at 200 000 (at $m/z$ 400) for all spectra. Automatic gain control was used to consistently fill the LTQ with the same number of ions ($n = 10^6$) for each acquisition and to avoid space charge effects from overfilling the mass analyzer. The instrument was externally calibrated in negative ion mode with a standard solution of sodium dodecyl sulfate and taurocholic acid. The resulting mass accuracy was better than 2 ppm. Over 200 individual mass spectra recorded in the time domain were collected and stored as transients for each sample using Thermo Xcalibur software.

2.4 Data processing and assignment of molecular formula composition

Ultrahigh-resolution FT-ICR mass spectra were post-processed using Composer software (Sierra Analytics, version 1.0.5) as previously described in Putman et al. (2012) and Mazzoleni et al. (2012). Briefly, approximately 200 transients were co-added to improve analysis detection (Kujawinski et al., 2002; Stenson et al., 2003). Internal recalibration of the mass spectra (Sleighter and Hatcher, 2008) was performed to improve the molecular formula assignment accuracy. A set of naturally occurring cloud water analytes from the two cloud water samples was selected to represent internal recalibrants over the mass range of 100–650 Da. The match tolerances for the internal recalibrants were set at 3 ppm with a minimum relative abundance (RA) of 18 times the root mean square (RMS) of the signal-to-noise ratio between the range $m/z$ 900–1000, (RMS of signal-to-noise ratio was 0.018 % for CW1 and 0.038 % for CW2). The list of internal recalibrants is given in Table S1 in the Supplement.

The molecular formula calculator was set to allow up to 70 carbon (C), 140 hydrogen (H), 25 oxygen (O), 4 nitrogen (N), and 1 sulfur (S) per molecular formula composition. The calculator is based on the PREDATOR algorithm (Blakney et al., 2011) and uses a Kendrick mass defect (KMD) analysis (Hughes et al., 2001) to sort ions into CH2 homologous series (species with a specific number of double bond equivalents (DBE) and heteroatom content) and then assigns the “de novo” molecular formulas ≤ 500 Da. The KMD analysis is determined from Eqs. (1) and (2) (Hughes et al., 2001):

Kendrick mass (KM) = (measured mass) $\times (14.00000)/(14.01565).$

KMD = nominal Kendrick mass (NKM) − KM. (2)

Using these equations, compounds of the same CH2 homologous series will have the same KMD. An assignment threshold of 6 times the RMS of the signal-to-noise ratio was applied to the detected anions. The molecular formulas resulting in measurement errors > 3 ppm were discarded. Additional data filtering of the assigned formulas was done by applying the rules and assumptions as described by Koch et al. (2005). The procedures applicable to atmospheric samples are described in more detail in Putman et al. (2012). DBE is determined by Eq. (3) (McLafferty and Tureek, 1993):  

$$DBE = c - h/2 + n/2 + 1$$

for each elemental composition $C_hN_nO_oS_s$. S and O are divalent and N is trivalent in Eq. (3). Thus, the additional double bonds formed by tetravalent and hexavalent S or pentavalent N are not represented in the DBE calculations (Mazzoleni et al., 2012).

Molecular formulas were assigned to the field and lab blank samples using the same method. The blank mass spectra contained several low-molecular-weight negative ions ($m/z$ < 400), most of which were assigned unambiguous molecular formulas as described above. Approximately 800 of the assigned monoisotopic formulas were in common with the cloud water samples CW1 and CW2. The common formulas represent approximately 20 % of the assigned molecular formulas in cloud water samples. Although some carryover of the sample components to blanks may have occurred (Mazzoleni et al., 2012), other sources of contamination could not be ruled out. Thus, all of the common molecular formulas assigned to negative ions in either of the blank samples were removed from the presented cloud water composition.

3 Results and discussion

3.1 Mass spectra and molecular formula assignments

Molecular formulas of the form $C_hN_nO_oS_s$ were assigned to the negative ion ultrahigh-resolution FT-ICR mass spectra of AOM extracted from cloud water samples. Overall, 82–83 % of the total ion current of the CW1 and CW2 sample mass spectra were assigned. After blank subtraction and quality assurance filtering, approximately 3000 monoisotopic molecular formulas were unambiguously identified for CW1 and CW2 (Table 1). Assignments of corresponding polyisotopic molecular formulas containing $^{13}$C were observed for 71 % and 69 % of the formulas assigned to CW1 and CW2, respectively. Likewise, polyisotopic molecular formulas with $^{34}$S were observed for 54 % and 50 % of the S-containing molecular formulas assigned to CW1 and CW2. Detection of polyisotopic anions is more limited than detection of monoisotopic anions due to the lower RA, which reflects the natural abundances of $^{13}$C at 1 % and $^{34}$S at 4 %. The high number frequency of corresponding polyisotopic molecular formulas with $^{13}$C at exactly 1.0034 Da higher than the monoisotopic molecular formulas confirms the molecular formula assignment and the singly charged state of the detected anions. No evidence of multiply charged species was observed.
Reconstructed mass spectra of the monoisotopic molecular formulas assigned to the cloud water samples after blank subtraction are shown in Fig. 1. In both of the cloud water mass spectra, the anions occupy a mass range of 100–700 Da, covering both the range of previously identified cloud water carbonyl species (Limbeck and Puxbaum, 2000; Marinoni et al., 2004; van Pinxteren et al., 2005; Samy et al., 2010; Charbouillot et al., 2012) and the range of water-soluble macromolecular compounds also known as “HULIS” (Feng and Möller, 2004). Macromolecular compounds were also observed in fog water by several studies (Krivacsy et al., 2000; Herckes et al., 2002a, b; Cappiello et al., 2003). Similar to the ambient aerosol collected at this site, the highest RA ions were observed between m/z 200–400 (Mazzoleni et al., 2012). Consistent with the ultrahigh-resolution mass spectra of other atmospheric samples like aerosols (Wozniak et al., 2008; Schmitt-Kopplin et al., 2010, fog water (Mazzoleni et al., 2010; LeClair et al., 2012), and rainwater (Altieri et al., 2009b, 2012), the mass spectra for cloud water samples are quite complex with a high number of isobaric anions. An example of the isobaric complexity with 14 molecular formulas within 0.25 Da is shown in an excerpt of the reconstructed CW1 mass spectrum (Fig. 1b).

Four subgroups of molecular formulas were determined based on their elemental composition; they include compounds containing only C, H, and O (CHO); compounds containing C, H, N, and O (CHNO); compounds containing C, H, O, and S (CHOS); and compounds containing C, H, N, O, and S (CHNOS). The CHO compounds (N = 840 and 712) represented ~26% of the total number of assigned molecular formulas from the CW1 and CW2 sample mass spectra (Fig. 1c, e). In general, a clear pattern of mass differences of the CHO compounds can be seen in both of the mass spectra (blue peaks in Fig. 1) with spacing of the high-intensity anions of 14 Da. This has been commonly observed in natural organic matter samples (Stenson et al., 2003; Koch et al., 2007; Mazzoleni et al., 2010). This pattern is a likely consequence of the naturally occurring CH$_2$ “homologous series” or formula extensions in organic matter (Hughes et al., 2001). All of the identified CHO molecular formulas belong to a CH$_2$ homologous series (Fig. 2). The series range from 2 to 26 molecular formulas in length, with an average length of ~10. However, it should be noted that the addition of the – CH$_2$ to a molecular formula does not necessarily lengthen the compounds’ carbon backbone, but may appear in any aspect of the compounds’ molecular structure. Moreover, each molecular formula identified may represent several organic compounds with different chemical structures (Hertkorn et al., 2008). Therefore, the clouds have extremely complex organic composition considering the ~ 3000 molecular formulas identified in each sample. In previous studies, the CHO compounds typically represented the highest number fraction of the overall compounds (Wozniak et al., 2008; Schmitt-Kopplin et al., 2010; Altieri et al., 2012; Mazzoleni et al., 2012). However in this study, CHNO compounds (N = 1638 and 1431) are the most frequently observed compounds in both cloud water samples and represented >50% of the total number of molecular formulas (Fig. 1c and 1e). From the mass spectra, the CHNO compounds (red peaks in Fig. 1) with the highest relative abundances are in the mass range of m/z 200–400 and the highest density is in the mass range of m/z 300–600. Consistent with the CHO compounds, a wide number of CH$_2$ homologous series were observed in the cloud water CHNO compounds (Fig. 2). The S-containing compounds (CHOS and CHNOS) represented ~21% of the total number of molecular formulas (Fig. 1c, e). In CW1, 281 CHOS and 381 CHNOS molecular formulas were identified, and in CW2, 221 CHOS and 360 CHNOS formulas were identified. Typically the S-containing compounds (green peaks in Fig. 1) are in the mass range of 200 < m/z < 600. Often the relative abundances of these compounds are lower than the compounds in other groups. The CH$_2$ homologous series of the S-containing compounds are less clear in the reconstructed mass spectra. However, they are clearly shown in the KMD plots (Fig. 2). The composition and molecular properties for each of the subgroups will be discussed further in the following sections.

### 3.2 Compounds containing only C, H, and O

Oxygen-to-carbon ratios (O/C) and hydrogen-to-carbon ratios (H/C) are commonly used to describe the chemical properties of aerosol (Aiken et al., 2007, 2008; Jimenez et al., 2009; Bateman et al., 2010; Kroll et al., 2011). In general, O/C represents the degree of oxidation and H/C reflects the degree of saturation of the composition studied. The O/C and H/C elemental ratios of the individual CHO compounds of the cloud water samples are plotted in Fig. 3a (for CW1) and Fig. S4a (for CW2). The van Krevelen diagram (O/C vs. H/C) indicates the oxidation, hydration, hydrogenation, and alkylation relationships between the observed molecular formulas (Kim et al., 2003; Wu et al., 2004; Heald et al., 2010). In Fig. 3a and S4a, we plotted simultaneously the analyte isobundance and individual points for each of the CHO molecular formulas. This permits visualization of the CH$_2$ homologous series of molecular formulas in lines pointing to the upper left corner (H/C = 2.0) of the plot as well as their significance in terms of isobundance. In both samples, the majority of the CHO compounds have O/C ratios < 1.0 and H/C ratio from 0.5 to 2.0. The highest intensity molecular formulas are in the vicinity of O/C = 0.5 and H/C = 1.5. Interestingly, there are several compounds with O/C ratios > 1.0 in both samples and many of these have 18–22 oxygen atoms and molecular weights of 400–500 Da. These highly oxidized high-molecular-weight compounds may be the products of aqueous-phase reactions (Lee et al., 2011). In addition, a few CHO compounds are located in the lower left corner of the van Krevelen diagram (O/C < 0.3, H/C < 1). These appear to be aromatic compounds with high DBE values. Examples of the molecular formulas in this
region include the homologous series of C_{20}H_{20}O_{4}(CH_{2})_{1−7} and C_{10}H_{18}O_{5}(CH_{2})_{1−6}. The average O/C ratios for the CW1 and CW2 CHO compounds are 0.54 (±0.35) and 0.51 (±0.28) (Table 1). Likewise, the average H/C ratios are 1.42 (±0.36) and 1.41 (±0.34). Similarly, Altieri et al. (2009b) observed an average O/C ratio of 0.7 (±0.5) and an average H/C ratio of 1.5 (±0.4) for the CHO compounds in ambient rainwater. Thus, the CHO compounds reported here are
less oxidized and somewhat less saturated, likely reflecting seasonal differences and cloud processing time differences. Other reported elemental ratios for CHO compounds in ambient aerosol water-soluble organic compounds include 0.47 (±0.2) and 1.42 (±0.3) (Mazzoleni et al., 2012), 0.6 and 1.69 (Rincon et al., 2012), and 0.40 (±0.21) and 1.29 (±0.35) (Lin et al., 2012a). Note that the solid-phase extraction step used in this study may not retain well some of the low-molecular-weight CHO compounds like formic acid, acetic acid, or oxalic acid, which usually have high O/C ratios (> 1). Thus, the O/C ratios of the clouds may be somewhat higher than reported here. Overall, the O/C and H/C ratios of CHO compounds in the cloud water samples are within the wide range of the previously reported values, but are closer to those of aerosol water-soluble organic carbon than rainwater. This is probably related to the different extents of aqueous processing and temperature in supercooled and precipitating clouds.

Another measure of molecular unsaturation is the DBE, which indicates the number of double bonds and rings in a molecular structure (Eq. 3). A wide range of DBE values (0–19) was observed among the cloud water CHO compounds with a clear trend of increasing DBE values with increasing carbon content (C_3–C_36) (Fig. 3b). In both samples, the majority (> 90%) of the compounds have DBE values ≤ 10 and the high-relative-abundance compounds have DBE values within the range of 2–7. Overall, the average DBE values for CW1 and CW2 CHO compounds are 6.4 (±3.6) and 6.4 (±3.2), similar to that of CHO compounds from other studies. Average DBE values for CHO compounds of 5.6 and 6.4 were reported by Lin et al. (2012a) and Mazzoleni et al. (2012) for ambient aerosol; Mazzoleni et al. (2010) reported an average DBE value for CHO compounds of 5.3 for low-molecular-weight fog water AOM; and Putman et al. (2012) reported an average DBE value for CHO compounds of 7.0 for chamber-generated α-pinene ozonolysis SOA. Interestingly, several CHO compounds with high DBE values (i.e., 14–19) were observed and are located in the aromatic region (O/C < 0.3 and H/C < 1.0) of the van Krevelen diagram (Fig. 3a). Generally, these aromatic compounds are within the mass range of 300–400 Da and have < 6 O atoms.

The relative abundances of the compounds with respect to carbon are also illustrated in Fig. 3b. The higher intensity compounds are compounds with C_5–C_25. Among these compounds, the highest relative abundances are ~C_10, followed by ~C_15. There is also a relatively high-intensity spot ~C_20 in the CW1 sample. These high-relative-abundance compounds at C_10, C_15, and C_20 indicate terpene characteristics in the cloud water CHO compounds (Claeyts et al., 2013).

To evaluate the oxygen content, the cloud water CHO compounds were separated into subclasses based on the number of oxygen atoms (O) in the molecular formulas, labeled as O_1, O_2, and so on. The total relative abundances for each of the subclasses are shown in Fig. 3c. Overall, cloud water CHO compounds range from O_2 to O_22, with the majority of compounds in the range of O_2 to O_15. These subclasses with variable oxygen numbers may reflect multiple and/or various functional groups like hydroxyl, ether, peroxyl, carbonyl, carboxyl, or ester. Interestingly, the highest relative abundances were observed for the O_5, O_7, and O_10 subclasses. The O_10 subclass contains the highest number of CHO compounds (~ 130 molecular formulas), while the highest relative abundance CHO compounds were observed in the O_5 and O_7 subclasses. Examples of the high-relative-abundance CHO compounds detected in both samples include C_{10}H_{18}O_{5}, C_{11}H_{18}O_{5}, C_{10}H_{14}O_{7}, C_{11}H_{16}O_{7}, C_{12}H_{18}O_{7}, C_{13}H_{20}O_{7}, and C_{14}H_{22}O_{7}. These high-relative-abundance compounds have DBE values of 2, 3, and 4. Interestingly, no formulas in the O_16 and O_17 classes were observed, but there are several compounds in the higher subclasses (O_18–O_22). These high oxygen numbers correspond to compounds with O/C ratios > 1.0 in the van Krevelen diagram.

The reconstructed mass spectra of the cloud water CHO compounds have similar characteristics to laboratory-generated SOA samples. Three groups of high abundance anions were observed; they include m/z 200–350, m/z 350–500, and m/z 500–650 (Fig. 4). Similar clusters of high-relative-abundance anions (sometimes referred to as “oligomers” or monoterpane accretion products in SOA.
samples) have been observed in SOA samples and have decreasing ion intensities with increasing molecular weight (Reinhardt et al., 2007; Bateman et al., 2011; Kundu et al., 2012; Putman et al., 2012). Also the average O/C and H/C ratios of the cloud water CHO compounds are consistent with those determined for biogenic SOA samples (Bateman et al., 2009, 2010; Putman et al., 2012). In addition, there are several “hot spots” at C_{10}, C_{15}, and C_{20} in the isobundance DBE plots (Fig. 3b) representing the high-relative-abundance compounds. The DBE values and the carbon numbers of the high-relative-abundance compounds reflect monoterpene and sesquiterpene characteristics (Putman et al., 2012). To further explore the SOA composition similarities, the cloud water CHO molecular formulas were compared to the CHO molecular formulas assigned to several laboratory-generated biogenic SOA mass spectra. The biogenic SOA samples used in the comparison include α-pinene ozonolysis SOA, β-pinene ozonolysis SOA, d-limonene ozonolysis SOA, and β-caryophyllene ozonolysis SOA, all of which were generated under dark conditions without an OH scavenger (e.g., Kundu et al., 2012; Dalbec et al., 2013). Of the CW1 and CW2 CHO molecular formulas, 58 and 64 %, were identical to those in the biogenic SOA samples (Fig. 4). Furthermore, most of these common molecular formulas are high-intensity anions in the cloud water samples. Overall, the three groups of anions assigned as CHO compounds in the reconstructed mass spectra are less distinct than those in the SOA mass spectra. This reflects the more complex conditions for ambient aerosols and clouds than laboratory experimental conditions. Some of the differences include the presence of biogenic SOA from unevaluated precursors, aqueous-phase reactions that alter the composition, and compounds from other sources, etc. Despite the differences between ambient conditions and those of laboratory experiments, we observed highly similar characteristics in the CHO molecular formulas of cloud water and biogenic SOA. This suggests that a high percentage of the cloud condensation nuclei in the Rocky Mountain region during the cloud sampling study contain biogenic SOA components. Similarly, Sun et al. (2011), using aerosol mass spectrometry, found ~90 % of the aerosol water-soluble organic carbon at a rural site (Centreville, Alabama) in the summer are biogenic related. The fraction decreased in the winter, but still comprised ~50 % of the total water-soluble organic carbon.

In addition to biogenic SOA, the CHO composition of the clouds appears to be affected by residential wood combustion. Guaiacols and syringols (also known as methoxyphenols) are commonly found in biomass combustion emissions and are from the pyrolysis of wood lignin (Hawthorne et al., 1992; Mazzoleni et al., 2007). Methoxyphenols have been shown by Sun et al. (2010) to contribute to aqueous SOA and have been observed in ambient foggy atmospheres (Sagebiel and Seiber, 1993). Two of the molecular formulas considered to be markers of syringol and guaiacol aqueous-phase reactions in the Sun et al. (2010) experiments were observed in the cloud water samples. They are C_{16}H_{18}O_{6} (RA = 7 %

![Isoabundance van Krevelen diagram for CW1 CHO compounds](image-url)
in both samples) and C_{14}H_{14}O_{4} (RA = 2.7 % in CW1 and 4 % in CW2). As described, molecular formulas assigned to exact mass measurements using ultrahigh-resolution FT-ICR mass spectrometry provide valuable insights regarding the chemical structure (e.g., elemental ratios and DBE). A large fraction of the high-relative-abundance O_{5} (50 out of 79 formulas in CW1, 39 out of 63 formulas in CW2) and O_{7} (68 out of 74 formulas in CW1, 74 out of 77 formulas in CW2) CHO compounds have DBE values ≥ 4. This combination is consistent with methoxyphenol-derived structures since there is both enough oxygen for the functional groups of -OCH_{3} and -OH and enough DBE for the benzene ring. Samy et al. (2010) analyzed aerosol and supercooled cloud water samples collected at the SPL. They found abundant concentrations of methoxyphenols in both the aerosols and the cloud samples, which suggested a strong biomass burning chemical signature. The emissions likely arise from nearby residential wood combustion in Yampa Valley. Similarly, Collett et al. (2008) also reported fog processing of carbonaceous particles from wood smoke. Residential wood combustion may contribute to the observed biogenic SOA in the cloud samples. It is known that terpenoid emissions from living plants are highly correlated to ambient temperature (Kesselmeier and Staudt, 1999), and thus biogenic SOA is generally not expected to be significant during the winter time periods. However, trees have a large storage reservoir of the monoterpenes and low DBE molecules assigned. For example, the molecular formulas of the homologous series C_{5}H_{10}N_{2}O_{6} (CH_{2})_{1−3} with high relative abundances show these characteristics. Also, there are a large number (~50 %) of CHNO molecular formulas with O/C < 0.1, many of the CHNO compounds have O/C > 1.0; they also have some of the highest relative abundances of the CHNO group. Most of these CHNO compounds with high O/C ratios also have high H/C ratios (~1.2), suggesting that the compounds are both highly saturated and oxygenated. For example, the molecular formulas of the homologous series C_{5}H_{10}N_{2}O_{6} (CH_{2})_{1−3} with high relative abundances show these characteristics. Also, there are a large number (~50 %) of CHNO molecular formulas with O/C < 0.7. In general, these compounds have a relatively low relative abundance and have a large range of H/C ratios (0.3 to 2.2). Overall, the average O/C ratios of the CHNO compounds in CW1 and CW2 are 0.72 (±0.34) and 0.73 (±0.31), respectively. The average H/C ratios for the CHNO compounds in CW1 and CW2 are 1.56 (±0.29) and 1.56 (±0.30) (Table 1).

3.3 Compounds containing only C, H, N, and O

A wide range of elemental ratios was observed for the CHNO compounds (Fig. 5a). In contrast to the elemental ratios of the CHO compounds, where the majority of the compounds have O/C < 1.0, many of the CHNO compounds have O/C > 1.0; they also have some of the highest relative abundances of the CHNO group. Most of these CHNO compounds with high O/C ratios also have high H/C ratios (~1.2), suggesting that the compounds are both highly saturated and oxygenated. For example, the molecular formulas of the homologous series C_{5}H_{10}N_{2}O_{6} (CH_{2})_{1−3} with high relative abundances show these characteristics. Also, there are a large number (~50 %) of CHNO molecular formulas with O/C < 0.7. In general, these compounds have a relatively low relative abundance and have a large range of H/C ratios (0.3 to 2.2). Overall, the average O/C ratios of the CHNO compounds in CW1 and CW2 are 0.72 (±0.34) and 0.73 (±0.31), respectively. The average H/C ratios for the CHNO compounds in CW1 and CW2 are 1.56 (±0.29) and 1.56 (±0.30) (Table 1). Previous studies of aerosol WSOC and fog AOM observed overall less oxygenated CHNO compounds, with average O/C ratios typically < 0.5 (Wozniak et al., 2008; Mazzoleni et al., 2010; Schmitt-Kopplin et al., 2010). However, in the study of rainwater AOM by Altieri et al. (2009b), several CHNO compounds were identified with high O/C ratios (some even higher than 2.0). They reported an average O/C ratio of 1.6 (±1.4) for the CHNO compounds. The presence of highly oxygenated CHNO compounds (O/C > 1.0) appears to be related to aqueous-phase reactions in cloud and rainwater droplets.

Consistent with the H/C elemental ratios, the majority of the CHNO compounds have DBE values ≤ 10 (Fig. 5b). Similar to the CHO compounds, the number of carbon atoms in the formulas range from C_{2} to C_{35} and the DBE values increase with the number of carbon atoms. The high-intensity CHNO compounds with C_{2}–C_{20} have DBE values ≤ 6. The most abundant compounds are those with C_{11} and low DBE values (2–4). These values indicate that most of the CHNO compounds are relatively saturated, especially those with higher relative abundances. Only a small fraction (~3 %) of the CHNO compounds have high DBE values from 11 to 19. As illustrated in Fig. 5b, the high-DBE CHNO compounds have low relative abundances. Overall, the average DBE values are 5.6 (±2.6) and 5.6 (±2.8) for CW1 and CW2 CHNO compounds. Therefore the cloud water CHNO compounds are mostly saturated, with a few exceptions.

Cloud water CHNO compounds contain 1–4 nitrogen atoms (N_{1}–N_{4}) per molecular formula. The compounds with N_{1} represented ~43 % of the CHNO compounds in both samples, followed by compounds containing N_{2} (~29 %), N_{3} (~21 %), and N_{4} (~7 %) in both samples. To further examine the N in the molecular formulas, CHNO subclasses were defined by the numbers of nitrogen and oxygen in the molecular formulas; they include NO_{3}–23, N_{2}O_{8}–18, N_{3}O_{12}–16, and N_{4}O_{14}–19. The total relative abundance for the compounds contained within the subclasses is shown in Fig. 5c. Interestingly, the plot clearly indicates that the high-relative-abundance CHNO compounds are in the subclasses of NO_{6}–14, N_{2}O_{8}–16, and N_{3}O_{11}–16. The compounds in high-relative-abundance subclasses correspond to the hot spots in the upper right region of the isoabundance van Krevelen diagram (O/C > 0.7, H/C > 1.5, Fig. 5a). All the high-relative-abundance CHNO subclasses are rich in oxygen with respect to nitrogen. Almost all of these classes show an oxygen-to-nitrogen ratio (O/N) ≥ 3. Thus, the CHNO compounds appear to contain a large number of organonitrate (with the functional group of −NO_{3}); however CHNO compounds are not exclusively organonitrate since multifunctional compounds are anticipated. For example a study using an ultrahigh-resolution “tandem mass spectrometry” technique reported only 63 % HNO_{3} losses from CHNO and CHNOS compounds in fog water samples (LeClair et al., 2012).

Organonitrates or nitrooxy organosulfates have been observed in several AOM samples (Allen et al., 1994; Nielsen
Fig. 4. Reconstructed mass spectra of the CHO compounds in CW1 (a) and CW2 (b). Blue peaks show the common formulas between the cloud water samples and biogenic SOA samples (combined data of ozonolysis of α-pinene, β-pinene, d-limonene and β-caryophyllene); red peaks show the formulas unique to cloud water samples.

Fig. 5. Isoabundance van Krevelen diagram for CW1 CHNO compounds (a). Isoabundance plot of DBE vs. number of carbon atoms in molecular formulas for CW1 CHNO compounds (b). Total relative abundance (RA) of each subclass for CW1 CHNO compounds; the dashed lines separate the subclasses by number of nitrogen atoms contained in the formulas (c). The corresponding figures for CW2 CHNO compounds are shown in Fig. S5.

et al., 1998; Zhang and Anastasio, 2001; Garnes and Allen, 2002; Herckes et al., 2007; Reemtsma et al., 2006; Surratt et al., 2006; Altiere et al., 2008; Bruns et al., 2010; Fry et al., 2013). The organonitrates or nitrooxy organosulfates observed in the clouds may be from dissolution of CCN aerosol, gas organic compounds containing nitro- or nitrate groups partitioned to the droplets (Lüttke and Levsen, 1997; Lüttke et al., 1997), or aqueous formation within the cloud droplets. Alkyl nitrates have relatively high vapor pressures, so they are not expected to readily partition to the particle phase. In a study by Nielson et al. (1998), only alkyl nitrates with > 18 carbon atoms can contribute to the particulate organic nitrogen mass at 20 °C. However hydroxyl or carboxylic acid functional groups lower the vapor pressure (Arp and Goss, 2009), allowing the multifunctional organonitrates with less carbon atoms to partition to the particle phase. This coupled with the low temperatures during the sampling period (∼ −10 °C) enhances the significance of organonitrates in the
condensed phase. Therefore it is highly possible that a considerable number of organonitrates are present in the super-cooled cloud droplet samples. Similarly, nitrate esters were also observed in fog water (Herckes et al., 2007).

As mentioned ∼50% of the CHNO compounds have O/C ratios < 0.7 (Fig. 5a). These low O/C ratios are similar to those reported by Schmitt-Kopplin et al. (2010) for a biomass burning aerosol sample. In that study, almost all the identified nitrogen-containing species had O/C ratios less than 0.7. The lower oxygen content suggests the presence of reduced nitrogen compounds. Wood combustion emissions are known to emit reduced nitrogen compounds. In a study by Laskin et al. (2009) a substantial fraction of N-heterocyclic alkaloid compounds were observed in biomass burning aerosols. In another study by Bateman et al. (2010), an O/C ratio of 0.19 was reported for combustion of dried pine needles and sticks. Thus, the residential wood combustion in the vicinity of the SPL is a likely contributor to the reduced nitrogen-containing species observed in the cloud water samples. However this does not exclude the possibility of aqueous-phase reactions resulting in nitrogen-containing species like those formed by aqueous-phase reactions of glyoxal (Galloway et al., 2009; Shapiro et al., 2009) or methylglyoxal (De Haan et al., 2010; Sareen et al., 2010) with (NH₄)₂SO₄/NH₄NO₃. Reaction products such as amine, imine, or imidazole provide another likely source of the reduced nitrogen compounds in the cloud water samples. Alkyl amines have also been detected in ambient aerosols (Gilardoni et al., 2009) and fog water (Zhang and Anastasio, 2001; Herckes et al., 2007). Regardless of their origin, reduced nitrogen functional groups can be oxidized in the cloud droplets to form oxidized functional groups as indicated by the high O/C ratios of products from aqueous-phase reactions (Lim et al., 2010; Ervens et al., 2011).

### 3.4 Compounds containing only C, H, O, and S

A wide range of elemental ratios is shown in Fig. 6a for the CHOS compounds and Fig. 7a for the CHNOS compounds. Similar to the CHO (Fig. 3a) and CHNO compounds (Fig. 5a), the elemental ratios align into several straight lines indicating CH₂ homologous series. However instead of one large group in the isoabundance van Krevelen diagrams, subgroups of the CHOS and CHNOS compounds in the isoabundance van Krevelen diagrams are depicted. Some obvious differences in the isoabundance van Krevelen diagrams of the CHOS and CHNOS compounds compared to those of the CHO and CHNO compounds were observed (Figs. 6a and 7a). First, the presence of a high number of highly unsaturated CHOS and CHNOS compounds was observed. Second, we observed a wider overall range of elemental ratios with variable relative abundances. These compound properties are explored in more detail in the following paragraphs.

Four subgroups of CHOS molecular formulas with varied unsaturations or DBE values were observed (Fig. 6). The CHOS compounds in subgroup 1 with H/C > 2 are highly saturated with DBE values of 0 or 1 (Fig. 6b). These compounds belong to the subclasses of O₄S, O₃S, and O₂S (Fig. 6c). Subgroup 2 CHOS compounds are less saturated and contain the subclasses of O₇−₁₃S with relatively low DBE values (mostly 3 or 4). The CHOS compounds in subgroup 3 are highly oxygenated with the subclasses of O₁₅−₁₈S. They are less saturated than those in subgroup 2 with DBE values ranging from 7 to 9. Most of the subgroup 3 molecular formulas have relatively high molecular weight, typically > 400 Da. Thus from subgroups 1 to 3, the DBE values increase (from 0 to 9) with the increase of oxygen content in the molecular formulas (O₄ to O₁₈). The CHOS compounds in subgroup 4 are the most unsaturated of the CHOS compounds. The DBE values of these molecular formulas range from 8 to 19, suggesting aromatic moieties. Unlike the compounds in the previous subgroups, these have low O/C ratios (< 0.4). The subclasses of O₁−₇S in this group contain too little oxygen to contain organic sulfate functional groups. Overall around 1/3 of the detected CHOS compounds are in subgroup 4 which appears to be reduced sulfur species.

Similarly, the CHNOS compounds have similar elemental ranges (Fig. 7a). The number of CHNOS compounds in subgroups 1–3 are less than those of the CHOS compounds and the subgroup 4 CHNOS compounds are clearly dominant among the CHNOS compounds. Overall, a total of 75% of the CHNOS compounds are in subgroup 4 and 25% in subgroups 1–3. The CHNOS compounds in subgroups 1–3 have DBE values ranging from 1 to 8 (Fig. 7b). These compounds contain 1 or 2 nitrogen atoms in the molecular formulas with a high number of oxygen atoms (> 7, subclasses NO₇−₁₂,₁₈S and N₂O₁₂−₁₆S; Fig. 7c). Based on the oxygen numbers and DBE values, some of the molecular formulas may represent organonitrates, organosulfates, or nitroxy organosulfates. In contrast, the subgroup 4 CHNOS compounds are highly aromatic, with DBE values from 6 to 19. Most of the CHNOS compounds with 3 or 4 nitrogen atoms in their molecular formulas are in this group, including: N₃O₁−₁₁S and N₄O₁−₅S. Furthermore, CHNOS compounds with 1 or 2 nitrogen atoms and a low number of oxygen atoms are in this group, including: NO₂−₄S and N₂O₂−₉S. Therefore, consistent with the cloud water CHOS compounds, the CHNOS compounds also include characteristics that are either highly oxygenated or highly aromatic. The aromatic S-containing compounds are dominant in number and thus greatly influence the average DBE value for these elemental classes.

Considering the adequate oxygen content and low DBE values (DBE for a sulfate functional group is 0 using Eq. (3)), the S-containing compounds detected in subgroups 1–3 can be organosulfates or nitroxy-organosulfates. Examples include the several high-relative-abundance, low-molecular-weight CHOS compounds detected in both samples (200–300 Da): C₁₂H₂₆O₄S, C₉H₁₈O₈S, C₁₄H₃₀O₄S.
and C₅H₁₆O₆S. LeClair et al. (2012) also detected these molecular formulas in fog water and used MS/MS to confirm the sulfate group in the structures. Organosulfates have been detected commonly in atmospheric samples (Surratt et al., 2006; Gómez-González et al., 2008; Hatch et al., 2011). In fact, most of the atmospheric S-containing organic compounds reported in the literature are organosulfates or nitrooxy organosulfates. Observations from both chamber simulations (Inumma et al., 2007; Liggio et al., 2007; Surratt et al., 2007) and field measurements (Surratt et al., 2008; Stone et al., 2012; Pratt et al., 2013) indicate that organosulfates and nitrooxy organosulfates can form in biogenic secondary organic aerosols. The identified organosulfates in biogenic SOA usually have carbon atom numbers similar to their biogenic precursors, e.g., ~5 or 10 (Surratt et al., 2008). In the cloud water samples, most of the CHOS and CHNOS compounds with less than 12 carbon atoms contain more than 4 oxygen atoms. Considering the cloud water biogenic SOA characterization identified from the cloud water CHO compounds, these organosulfates and nitrooxy organosulfates compounds (S-containing compounds in subgroups 1–3) may have originated from biogenic precursors. Several high-molecular-weight organosulfate candidates (> 300 Da) with more carbon atoms (16 to 33) were also detected in the cloud water, including: C₁₇H₁₈O₁₆S, C₁₈H₃₈O₆S, and C₁₇H₃₀O₁₆S. The high-molecular-weight organosulfates have been observed previously in atmospheric aerosols (Romero and Oehme, 2005; Reemtsma et al., 2006). Mostly AOM organosulfates are secondary reaction products (Hatch et al., 2011). Two formation mechanisms were proposed, including: esterification of hydroxyl groups with sulfuric acid (Liggio et al., 2005; Surratt et al., 2007) and acid-catalyzed reactions of epoxides (Minerath and Elrod, 2009). Both of the pathways require sulfuric acid anions (Surratt et al., 2007; Minerath and Elrod, 2009). Considering the coal-fired power plants located to the west of the SPL area, which are emission sources of SO₂, and the aqueous-phase processing that may take place within cloud droplets, formation of organosulfates is likely.

Although organosulfates and nitrooxy organosulfates were observed, a dominant number of S-containing compounds contain reduced S (e.g., aromatic S) due to insufficient oxygen content in the molecular formulas (subgroup 4 in Figs. 6 and 7). AOM with reduced S is rarely reported.
Fig. 7. Isoabundance van Krevelen diagram for CW1 CHNOS compounds (a). Isoabundance plot of DBE vs. number of carbon atoms in molecular formulas for CW1 CHNOS compounds (b). Number of formulas identified in each of CW1 CHNOS subclasses (c). The red circles indicate the approximate groupings (1 to 4) of the CW1 CHNOS formulas. The bars not boxed in (c) belong to group 4. The corresponding figures for CW2 CHNOS compounds are shown in Fig. S7.

(Saranjampour, 2012; Kundu et al., 2013). Most of the high-DBE, S-containing compounds observed in cloud water have high carbon numbers (> 10, Figs. 6b and 7b). Instead of being secondary products like the organosulfates, they may be from primary emissions. The diesel fuel used to power snow cats used to groom the ski area at night near the SPL may be a source of reduced S compounds. Further investigation is needed to determine the origin of the unambiguously identified reduced S compounds.

3.5 Cloud water AOM properties

Overall, the bulk properties of the two cloud water samples are very similar (Table 1). The average O/C ratios of the CW1 and CW2 samples are 0.62 (±0.37) and 0.61 (±0.34) and the average H/C ratios are 1.46 (±0.36) and 1.46 (±0.35). The average elemental ratios of each compound group with and without relative-abundance-weighting (Eqs. S1 and S2) are summarized in Table 1. The relative-abundance-weighted elemental ratios are very similar in most cases to the unweighted values discussed here. As shown in Table 1, the O/C ratios of the CHO, CHOS, and CHNOS compounds are all approximately 0.5, but the O/C of the CHNO compounds is ~ 0.7. Thus the overall elemental ratios are highly influenced by the CHNO compounds given their high number frequency and high relative abundance. Low-molecular-weight organic compounds like formate, acetate, and formaldehyde observed previously in ambient samples (van Pinxteren et al., 2005; Herckes et al., 2007; Collett et al., 2008; Samy et al., 2010) but not here have relatively high O/C ratios (> 1). In general the overall O/C ratios observed here are consistent with the O/C ratios observed for other aerosol water-soluble organic carbon and AOM samples, but they are at the higher end of the reported range. For example, Aiken et al. (2008) reported O/C ratios for various ambient aerosols from 0.16 to 0.76 using aerosol mass spectrometry. Similarly, Lee et al. (2012) estimated O/C ratios of low-volatility cloud water organics to be between 0.52 and 0.59. The observed cloud water O/C values are higher than Mazzoleni et al. (2010) observed in fog water collected in Fresno, California (0.43), but lower than Altieri et al. (2009b) observed in rainwater collected in New Jersey (0.96). Similar to the O/C ratios, the H/C ratios of CHNO compounds are higher than those of the CHO, CHOS, and CHNOS compounds, so the overall H/C values are mainly driven by the high number and relative abundances of CHNO compounds. The observed H/C ratios for the cloud water are also within the range of reported values for aerosol water-soluble organic carbon and AOM samples. Aiken et al. (2008) reported H/C ratios for ambient aerosols from 1.41 to 1.89. A similar range of H/C values was observed in fog and rainwater samples.
Average DBE values and relative-abundance-weighted average DBE values (Eq. S3) of each compound group are listed in Table 1. The average DBE values of the CW1 and CW2 samples are 6.30 (±3.37) and 6.29 (±3.29). Again despite the aromatic compounds identified in the CHO, CHOS, and CHNOS groups, the high number of saturated CHNO compounds lowers the overall average DBE values. The relative-abundance-weighted average values of the samples are even lower than the unweighted average DBE values due to the influence of high-relative-abundance saturated CHNO compounds. Similarly, the relative-abundance-weighted DBE value for the summer aerosol water-soluble organic carbon at the SPL was observed to be 5.35 (±0.05) (Mazzoleni et al., 2012). In general, the DBE values for the cloud water AOM are consistent with a strong biogenic influence (e.g., oxidation products of terpenes and methoxylphenols) and are comparable to other aerosol water-soluble organic carbon and AOM samples.

The average and relative-abundance-weighted organic mass-to-organic carbon ratios (OM/OC) for the samples and each of the compound groups were also evaluated (Table 1). Consistent with previous studies (Altieri et al., 2009b; Mazzoleni et al., 2012), we define OM/OC as the measured mass divided by the calculated mass of C in the assigned formulas. Overall, the OM/OC ratios for the clouds were ~2.1. As expected, the compound groups with heteroatoms (N and S) have higher OM/OC ratios than the CHO compound group. The observed OM/OC ratios for the clouds are slightly higher than those for the ambient aerosols (Aiken et al., 2008; Mazzoleni et al., 2012), reflecting a higher oxidized characteristic and the importance of N- and S-containing compounds identified in clouds.

Although the overall bulk properties for the two cloud water samples are quite similar, detectable compositional differences between the two samples were observed within the N-containing compounds. The reconstructed difference mass spectrum between the CW1 and CW2 CHNO compounds (Fig. 8) shows several unique peaks with high relative abundance in CW1 in the range of m/z 250–350, while those in CW2 are within the range of m/z 350–450. The unique high-relative-abundance CHNO compounds in CW1 are mostly saturated compounds with low DBE values (2 and 4). Some examples of these compounds are the homologs of N₂O₁₀ with DBE values of 2 (C₄H₈(CH₂)₁₋₅N₂O₁₀), and homologs of N₂O₈ with DBE values of 4 (C₅H₁₂(CH₂)₁₋₃N₂O₈). However, the unique high-relative-abundance CHNO compounds in CW2 contain one nitrogen atom instead of two. Some example compounds are homologs of NO₁₀ with DBE values of 3, 5, or 7, and homologs of NO₁₁ with DBE values of 4, 5, 6, or 8. The average N/C ratios for the unique CHNO compounds are 0.15 in CW1 and 0.13 in CW2. Similarly, the average N/C for the unique CHNOS compounds in CW1 is higher than in CW2 (0.12 versus 0.10). These observations may be the result of nighttime NO₅ chemistry (Iinuma et al., 2007; Fry et al., 2013). At night, nitrate radical (NO₃⁻) is produced from the reaction of O₃ and NO₂, which can react with organics (RO₂) to produce organonitrates RONO₂ (Kroll and Seinfeld, 2008). As discussed in Section 3.3, organonitrates observed in the clouds may be from the CCN particulate phase, the gas phase, or formed within the aqueous phase. The NO₅ chemistry can be involved in all these pathways at night. For example, more organonitrates would be produced in the gas phase and partitioned to the aerosols. Also, it has been shown that nitrate radical can transfer from the atmospheric gas phase to the aqueous phase and is the main source of nitrate radical in aqueous phase (Herrmann et al., 2010). Oxidation of reduced N-containing compounds or nitrate radical oxidation of CHO or CHOS compounds in the aqueous phase may occur (Perraud et al., 2010). Since part of the CW1 composite was collected at night, it is reasonable to observe more N-containing compounds with higher nitrogen content in the molecular formulas from nighttime NO₅ chemical pathways in CW1 than in CW2.

4 Conclusions

Approximately 3000 monoisotopic molecular formulas were assigned to each of the two supercooled cloud water samples collected at Storm Peak Laboratory. An overall complex organic composition with N, O, and S, as well as a wide range of DBE values similar to other ambient aerosol water-soluble organic carbon and aqueous AOM samples, was observed. CHO, CHNO, CHOS, and CHNOS compounds with molecular weights up to 700 Da were observed in the cloud samples. Approximately 70% of the identified molecular formulas were observed in the mass range of 300 to 500 Da. Unique to this observation, the CHNO compounds had the highest number frequency and relative abundances. The average elemental ratios (O/C = 0.6 and H/C = 1.5) and DBE values (6.3) were largely influenced by the highly oxygenated CHNO compounds in the clouds and were slightly higher than the aerosol water-soluble organic carbon collected in the summer at this site. Consistent with the summer aerosol water-soluble organic carbon composition, the majority of the cloud water CHO compounds matched CHO compounds assigned to chamber-generated biogenic SOA. Furthermore, the organosulfates observed in the clouds suggest biogenic contributions to the cloud water AOM. Biomass stored terpenes are likely released during residential wood combustion, yielding significant biogenic SOA-like composition. Combustion products also influence the cloud water AOM composition as evidenced by the high number (~1500 unique formulas) of nitrogen-containing compounds. Approximately one-third of the CHOS and approximately three-quarters of the CHNOS compounds were identified as reduced S-containing compounds. They might be from diesel fuel combustion or power plant emissions, but further investigation is needed to determine their origin. More nitrogen atoms were associated with the high
RA molecular formulas assigned to the cloud sample with a nighttime composite than the daytime sample (2 versus 1 nitrogen atom), likely reflecting the nighttime NO\(_x\) chemistry. The molecular composition determined from ultrahigh-resolution FT-ICR mass spectrometry provides insights into the organic composition of cloud water AOM in the Rocky Mountain area and indicates significant biogenic SOA and residential wood combustion contributions to the overall composition.

Supplementary material related to this article is available online at http://www.atmos-chem-phys.net/13/12343/2013/acp-13-12343-2013-supplement.pdf.

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