Observation of playa salts as nuclei in orographic wave clouds

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[1] During the Ice in Clouds Experiment–Layer Clouds (ICE–L), dry lakebed, or playa, salts from the Great Basin region of the United States were observed as cloud nuclei in orographic wave clouds over Wyoming. Using a counterflow virtual impactor in series with a single-particle mass spectrometer, sodium-potassium-magnesium-calcium-chloride salts were identified as residues of cloud droplets. Importantly, these salts produced similar mass spectral signatures to playa salts with elevated cloud condensation nuclei (CCN) efficiencies close to sea salt. Using a suite of chemical characterization instrumentation, the playa salts were observed to be internally mixed with oxidized organics, presumably produced by cloud processing, as well as carbonate. These salt particles were enriched as residues of large droplets (>19 μm) compared to smaller droplets (>7 μm). In addition, a small fraction of silicate-containing playa salts were hypothesized to be important in the observed heterogeneous ice nucleation processes. While the high CCN activity of sea salt has been demonstrated to play an important role in cloud formation in marine environments, this study provides direct evidence of the importance of playa salts in cloud formation in continental North America has not been shown previously. Studies are needed to model and quantify the impact of playas on climate globally, particularly because of the abundance of playas and expected increases in the frequency and intensity of dust storms in the future due to climate and land use changes.


1. Introduction

[2] The effects of aerosol particles on cloud formation and properties represent the largest source of uncertainty in climate prediction [Intergovernmental Panel on Climate Change, 2007]. Certain aerosol particles can serve as cloud condensation nuclei (CCN) or ice nuclei (IN), indirectly affecting climate by changing cloud reflectivity, cloud lifetime, and precipitation patterns [Andreae and Rosenfeld, 2008]. To model current and future climate change, the aerosol population and corresponding cloud nucleating ability must be understood, particularly with regard to spatial and temporal variability. The critical supersaturation ($S_c$) at which a particle activates to become a cloud droplet is dependent on particle diameter and chemistry [Andreae and Rosenfeld, 2008]. While chemical composition can be less important than size, it is predicted to have a significant role in the activation of aerosols containing less than approximately 10–15% soluble material [Junge and McLaren, 1971; Roberts et al., 2002]. Thus, Dusek et al. [2006] suggested that regional and global climate models can assume a “typical” size distribution to estimate CCN abundance and the effects of aerosols on cloud physics. However,
Hudson [2007] examined the relationship between particle size and S\(_4\) for a greater range of ambient conditions, showing greater variability than observed by Dusek et al. [2006], particularly for “clean” air masses. Due to variability between different air masses, Hudson [2007] concluded that particle chemistry must be considered to avoid inaccurate estimates of CCN. Recently, Twohy and Anderson [2008] observed salts (chloride and sulfate salts of Na, K, Ca, and Mg) as preferred cloud droplet nuclei for particles <0.2 \(\mu\)m in diameter. However, most cloud studies have focused on marine locations with few studies investigating the chemical composition of clouds in remote continental locations.

[3] Uncoated mineral dust particles are generally assumed to be insoluble, not acting as CCN until becoming coated with sulfate, nitrate, or chloride, for example [Andreae and Rosenfeld, 2008]. However, Twohy et al. [2009] found submicron African dust near the source acted as CCN, and recent work has highlighted the role of dust mineralogy on cloud activation properties [Kelly et al., 2007]. In particular, dust from saline dry lakebeds, or playas, is generally composed of \(\sim 10-44\%\) soluble salts [Abuduwaili et al., 2008; Blank et al., 1999; Reheis, 1997; Singer et al., 2003]. Salt crusts form on playas following rapid evaporation of rainwater or groundwater fluctuations [Reynolds et al., 2007], which concentrates soluble salts at the top surface layer, producing a fluffly, wind-erodible crystalline surface [Bluckwelder, 1931; Lowenstein and Hardie, 1985; Russell, 1883]. Common playa evaporite minerals include halite, gypsum, mirabilite, thendardite, epsomite, trona, sylvite, carnallite, calcite/aragonite, dolomite, hanksite, northupite, and aphthitalite [Bryant et al., 1994; Droste, 1961; G"{u}ven and Kerr, 1966; Lowenstein and Hardie, 1985]. Chemical analysis of playa dust has shown increased mass fractions of potassium, chloride, sulfate, and sodium with respect to the bulk (deeper) playa soil [Blank et al., 1999].

[4] Playas are present globally [Jones and Deocampo, 2004; Prospero et al., 2002] and contribute to the high global dust burden [Andreae and Rosenfeld, 2008]. Formenti et al. [2003] identified the presence of playa salts within Saharan dust plumes over the Atlantic Ocean. Recent laboratory CCN measurements demonstrated that a more hygroscopic fraction of playa dust showed CCN activities similar to sodium chloride; whereas, a less hygroscopic fraction heterogeneously nucleated ice at lower relative humidities than required for homogeneous ice nucleation at temperatures below \(-40^\circ\)C [Koehler et al., 2007]. Studies in the western U.S. hypothesized that playa dust may be incorporated into clouds and rained out when air masses encounter mountain ranges [Reheis, 2006; Reheis and Kihl, 1995]. Recent satellite-based studies of Aral Sea dust storms suggest that playa dust particles act as giant CCN and promote precipitation [Rudich et al., 2002].

[5] Results derived herein from the Ice in Clouds Experiment-Layer Clouds (ICE-L) provide the first in situ chemical composition measurements of playa salts within cloud droplets and ice crystals. A highly detailed perspective of the individual cloud residue chemistry is obtained through a unique combination of mass spectrometry and microscopy techniques. CCN measurements are shown for a playa crust sample with similar chemistry to that of the cloud residues sampled during ICE-L. The incorporation of the playa salts within the orographic wave cloud is discussed.

2. Experimental Methods

2.1. Overview and Cloud Sampling

[6] Measurements in orographic wave clouds were made aboard the National Science Foundation/National Center for Atmospheric Research (NSF/NCAR) C-130 aircraft during ICE-L. In this paper, we primarily focus on measurements made on November 16, 2007 during research flight #3 (RF03) over Wyoming (41.6–42.8°N, 105.8–108.5°W) at 5.9–7.0 km above mean sea level (MSL). The National Oceanic and Atmospheric Administration (NOAA) Air Resources Laboratory HYSPLIT (Hybrid Single-Particle Lagrangian Integrated Trajectory) model was utilized for air mass trajectory analysis (R. R. Draxler and G. D. Rolph, HYSPLIT (HYbrid Single-Particle Lagrangian Integrated Trajectory) Model, 2003, access via NOAA ARL READY Web site, http://www.arl.noaa.gov/ready/hysplit4.html) (hereinafter Draxler and Rolph, NOAA Web site, 2003). In clear air, an Ultra High Sensitivity Aerosol Spectrometer (UHSAS, Particle Metrics Inc., Boulder, CO) provided size-resolved aerosol number concentrations (0.1–1.0 \(\mu\)m in diameter). Clear air total condensation nuclei (CN) concentrations were measured using a condensation particle counter (CPC, TSI Model 3760), detecting particles >13 nm in diameter.

[7] Cloud particle size distributions, habits and phase were measured with a suite of instruments. Cloud droplet size distributions from 2 to 50 \(\mu\)m were measured with a cloud droplet probe (CDP, Droplet Measurement Technologies, Boulder, CO). A two dimensional optical array cloud probe (2D-C, Particle Measuring Systems Inc., Boulder, CO), updated with fast electronics, was used as the primary measurement of number concentrations of cloud ice crystals >50 \(\mu\)m in diameter. The presence of supercooled droplets was detected using a Rosemount icing detector (Model 871, Rosemount Engineering Co., Minneapolis, MN). A cloud particle imager (CPI, Stratton Park Engineering Company Inc., Boulder, CO) was used to identify the presence of small droplets, as well as particle habits for particles above \(\sim 40–50 \mu\)m.

[8] Cloud residual particles were isolated using a counterflow virtual impactor (CVI) [Noone et al., 1988; Twohy et al., 1997]. The CVI separates individual cloud droplets and ice crystals from interstitial aerosol and gases, collecting and concentrating them within a small flow of high-purity dry nitrogen gas heated to 50°C. Cloud particles (droplets and ice crystals) with aerodynamic diameters \(>7 \mu\)m were selected using the CVI. Condensed water was evaporated providing individual residual particles for in situ analysis, described below. While cloud particles larger than \(\sim 50 \mu\)m are collected, they may shatter due to inertial forces or upon contact with internal inlet surfaces [Twohy et al., 1997]. The break-up of a large ice crystal may result in more than one residual particle if the initial ice crystal had previously collected liquid droplets or scavenged aerosol particles during its lifetime. Size distributions of the CVI residual particles were measured using an optical particle counter (LAS-Air 1001, Particle Measuring Systems Inc., Boulder, CO), which groups particles in eight size ranges between...
0.1 and 4.0 μm diameter with six second time resolution. Ambient particle impactor samples, described below, were also collected through the CVI inlet by turning off the counterflow, allowing it to function as a simple subisokinetic aerosol inlet.

[9] Aerosol and cloud residue chemical composition were measured using real-time mass spectrometry and microscopy, described below. For CVI data analysis, time periods were divided into three categories: liquid, ice, and mixed phase (both liquid droplets and ice crystals present simultaneously). Generally, cloud particles with aerodynamic diameters >7 μm were selected using the CVI; however, during one cloud period consisting of primarily liquid droplets, >19 μm cloud particles were selected by increasing the counterflow for a higher cut-point. CVI sampling occurred during periods with equivalent potential temperatures ($\theta_e$) of 318.3–323.8 K. Thus, for the mass spectrometry analysis, clear air analysis time periods with $\theta_e$ values within this range were chosen for comparison with the in-cloud periods.

### 2.2. Aircraft Aerosol Time-of-Flight Mass Spectrometer

[10] In situ measurements of the size-resolved chemical composition of individual submicron particles were made using the recently developed aircraft aerosol time-of-flight mass spectrometer (A-ATOFMS) [Pratt et al., 2009]. The A-ATOFMS measures, in real-time, the vacuum aerodynamic diameter ($d_{wv}$) and dual-polarity mass spectra of individual particles from ~70–1200 nm ($d_{w}$). Following a $^{210}$Po neutralizer and pressure-controlled inlet [Bahreini et al., 2008], particles are focused in an aerodynamic lens system. Particles are optically detected by two continuous wave 532 nm lasers spaced 6.0 cm apart, providing particle velocity and, thus, $d_{wv}$. Polystyrene latex spheres of known physical diameter from 95 to 1500 nm were used for the particle size calibration. During ICE-L, particles were described and ionized using 266 nm radiation from a Q-switched Nd:YAG laser operating at ~0.4–0.6 mJ. Positive and negative ions resulting from individual particles are detected within the time-of-flight mass spectrometer. For the analysis discussed herein, 283 individual particles were chemically analyzed: 35 particles during clear air sampling, 23 during liquid CVI sampling, 124 during mixed phase CVI sampling, 80 during ice CVI sampling, and 21 during the 19 μm cut-point CVI sampling. Standard errors of particle type number percentages were calculated based on Poisson statistics.

[11] Mass spectral peak identifications correspond to the most probable ions for a given $m/z$ ratio based on previous lab and field studies. Based on dual-polarity mass spectra, individual particles were divided into 6 categories: salt (Na–K–Mg–Ca–Cl, described in Section 3.1), biomass, organic carbon (OC), soot, sulfate, and other. The mass spectra of the biomass particles, from wildfires and wood combustion, are dominated by potassium ($m/z$ 39(K$^+$)) with less intense carbonate positive ions [Silva et al., 1999]; most of the biomass particles contained sulfate ($m/z$ ~97 (HSO$_4^-$)), and to a lesser extent, nitrate ($m/z$ ~62(NO$_3^-$)). The OC particles were characterized by carbonaceous marker ions at $m/z$ 12(C$^+)$, 27(C$_2$H$_3$/CHN$^+$), 36(C$_6$), and 37(C$_3$H$^+$); ammonium ($m/z$ 18(NH$_4^-$)), amines, and sulfate were typically observed in these particles. Soot particles, dominated by carbon cluster ions (C$_n^-$), were often mixed with potassium and sulfate; a fraction of these particles contained organic carbon markers as well. Sulfate particles were characterized by having only negative ion mass spectra with intense peaks at $m/z$ ~97(HSO$_4^-$) and ~195(H$_3$SO$_2$HSO$_4^-$). The “other” particle category contains less abundant particles, such as biological particles and particles producing only negative ion mass spectra containing both nitrate and sulfate.

### 2.3. Scanning Transmission Electron Microscopy: Energy Dispersive X-Ray Analysis

[12] Downstream of the CVI inlet, a two-stage round-jet impactor [Rader and Marple, 1985] allocated separation of particles by size. At typical sampling pressures of 500 mb, the small particle impactor plate collected 0.11 to 0.59 μm diameter unit-density particles, or 0.08 to 0.42 μm diameter 1.7 g cm$^{-3}$ density particles. The large particle plate collected larger particles up to several microns in size. Particles were impacted onto transmission electron microscope grids (nickel base coated with formvar and carbon) and were analyzed individually via scanning transmission electron microscopy (STEM) with energy dispersive X-ray analysis (EDX). Note that some organics and nitrates volatilize under the vacuum environment of the microscope. Approximately 50–100 particles were randomly selected and analyzed per stage per sample. The clear air sample was taken at $\theta_e$ values of 315.9–317.2 K, which is lower than the $\theta_e$ values during cloud sampling; thus, the air probably did not flow into the cloud from this level. The mixed-phase CVI sample was collected at $\theta_e$ of 321.7–322.3 K.

[13] For STEM-EDX analysis, particles were classified into eight primary types as given in Table 1. “Mixed” types were particles described by multiple categories; for example, crustal dust with sulfate or salts as these may be important for their role as both IN and CCN. Particles were classified as “unknown” if they did not fit into any of the specific categories or did not produce detectable X-ray signatures above background (for example, volatile organics or nitrates). This is the same classification scheme as used by Twohy and Anderson [2008], with the addition of the biomass category which would have been included in the “salts” category in previous work. Standard errors of particle type number percentages were calculated based on Poisson statistics.

### Table 1. STEM-EDX Particle Classification Scheme

<table>
<thead>
<tr>
<th>Category</th>
<th>STEM-EDX Compositional Signature</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crustal dust</td>
<td>Silicates (rich in Si, variable Na, Mg, Al, K, Ca, and Fe), carbonates (Mg and/or Ca with large C peak), phosphates (rare).</td>
</tr>
<tr>
<td>Soot</td>
<td>Carbon only. Characteristic morphology (chain aggregates).</td>
</tr>
<tr>
<td>Organic</td>
<td>Amorphous or irregular and may contain minor S and K, as well as C.</td>
</tr>
<tr>
<td>Sulfates</td>
<td>Ammonium sulfate/sulfate and sulfuric acid.</td>
</tr>
<tr>
<td>Sulfates</td>
<td>S, O only.</td>
</tr>
<tr>
<td>Industrial Metals</td>
<td>Metals such as Al, Fe, Cr, Ti, Mn, Co, Zn, Cu.</td>
</tr>
<tr>
<td>Biomass Salts</td>
<td>Sulfates with potassium, may contain C as well.</td>
</tr>
<tr>
<td>Mixed</td>
<td>Combinations of two or more of above categories</td>
</tr>
</tbody>
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2.4. Compact Time-of-Flight Aerosol Mass Spectrometer

[14] The Aerodyne compact time-of-flight aerosol mass spectrometer (C-ToF-AMS) [Drewnick et al., 2005] measured the mass concentrations of non-refractory species (organics, sulfate, nitrate, ammonium, and a fraction of chloride) in real-time. Following a pressure-controlled inlet [Bahreini et al., 2008], particles of \( d_{wa} \sim 50-800 \) nm are focused by an aerodynamic lens system, sent through a 3.5% chopper, and then impacted onto a tungsten vaporizer operated at 550–560°C. The chopper can be operated in three modes to gather either background mass spectra, ensemble average mass spectra over all particle sizes, or size-resolved mass spectra. Measurements of chloride and other semi-refractory species are reported herein, but concentrations of these species are significantly underestimated by the C-ToF-AMS because they are not efficiently volatilized at the vaporizer temperature (550–560°C) used. Once vaporized, molecules undergo electron impact ionization and are extracted at a rate of \( \sim 53 \) kHz into a time-of-flight mass analyzer.

[15] Detection limits for out of cloud aerosol were calculated as the standard deviation of the noise when measuring filtered air: organics (0.06 \( \mu g/m^3 \)), sulfate (0.02 \( \mu g/m^3 \)), nitrate (0.01 \( \mu g/m^3 \)), ammonium (0.07 \( \mu g/m^3 \)), and chloride (0.01 \( \mu g/m^3 \)). Detection limits for measurements made in cloud were calculated as the standard deviation of measurements made on the CVI inlet during cloud-free periods: organics (0.003 \( \mu g/m^3 \)), sulfate (0.0006 \( \mu g/m^3 \)), nitrate (0.0007 \( \mu g/m^3 \)), ammonium (0.0007 \( \mu g/m^3 \)), and chloride (0.0005 \( \mu g/m^3 \)). In practice, detection is often limited by counting statistics at low aerosol loadings. Lower detection limits during CVI sampling were due to the enrichment of particles by the CVI and entrainment in \( N_2 \) (rather than air) which reduces the noise at certain organic \( m/z \)'s. Corrections for CVI enhancements (typically a factor of \( \sim 20–30 \)) were completed for reported detection limits and mass concentrations. One important issue concerning C-ToF-AMS data is the collection efficiency (CE) of particles within the instrument [Huffman et al., 2005]; in this study, C-ToF-AMS mass loadings were not corrected for CE due to the lack of another mass measurement on the plane and because of possible changes in particle phase [Matthew et al., 2008] between clear air and CVI dry residue sampling. By not adjusting the mass loadings for CE, we are effectively assuming a collection efficiency of 1, although we note that the CE likely varied during the flight. Standard deviations are noted as uncertainties for average species mass fractions. Oxygen/carbon (O/C) ratios were calculated from unit resolution mass spectra following the estimation method developed by Aiken et al. [2008]:

\[
O/C = (0.0382 \pm 0.0005)(x) + (0.0794 \pm 0.0070)
\]

where \( x \) is the percentage of organic aerosol mass from \( m/z \) 44 (CO\(_2\)).


[16] Three large stage impactor samples, collected as described in Section 2.3, were analyzed with scanning transmission X-ray microscopy (STXM) with near edge X-ray absorption fine structure spectroscopy (NEXAFS) at Lawrence Berkeley National Laboratory’s Advanced Light Source (ALS) [Kilcoyne et al., 2003]. The clear air sample was taken at \( \theta_e \) values of 320.2–322.1 K, which is within the \( \theta_e \) values of cloud sampling. The ice phase CVI sample was collected at \( \theta_e \) of 319.0–323.1 K; similarly, the 19 \( \mu m \) cut-point CVI sample was collected in primarily liquid phase at \( \theta_e \) of 319.3–322.8 K. For each sample region of interest, \( \sim 100 \) images were acquired over the carbon K edge from 278 to 320 eV. Peaks arising from the potassium L2 and L3 edges occur at \( \sim 299 \) and 297 eV respectively. For some of the carbonate samples, the presence of calcium was confirmed at the L2 and L3 edges occurring at 353 and 349 eV respectively [Henke et al., 1993].

[17] Maps of elemental carbon (EC), organic carbon, potassium, and regions rich in non-carbon elements (inorganic, In) were used to classify the different particle types seen in the different samples (R. C. Moffet et al., Automated assay of internally mixed individual particles using X-ray spectromicroscopy maps, submitted to Analytical Chemistry, 2010). EC regions were identified as those regions containing greater than 35% carbon-carbon sp\(^2\) hybridized double bonds [Hopkins et al., 2007]. The pre-edge region occurring around 278 eV is proportional to the number of non-carbon atoms, and the difference between the post-edge and pre-edge regions is proportional to the number of carbon atoms. In regions were identified by the ratio of the pre-edge height to post edge height (\( h_{pre}/h_{post} \)); if \( h_{pre}/h_{post} \) was greater than 0.4, the area is defined as being dominated by inorganic material on a thickness basis. Finally, if an area contained absorbance due to COOH groups at 288 eV (near the carbon edge), the region was identified as containing OC. Particle class labels (OC, OCED, OCEDIn, and OCIn) were used to identify particles that contain one or more of these three components (In, OC and/or EC).

2.6. Continuous Flow Diffusion Chamber

[18] To measure the number concentrations of ice-nucleating aerosol particles in real-time, a continuous flow diffusion chamber (CFDC) [Eidhammer et al., 2010; Rogers et al., 2001] re-processed cloud residues near the cloud observation temperature and at a relative humidity (with respect to water) exceeding 100% to simulate ice formation conditions at expected wave cloud parcel entry conditions. This means of operating the CFDC emphasized condensation/immersion freezing nucleation as is expected for the cloud entry regions of orographic wave clouds on the basis of previous studies [Cooper and Vali, 1981]. Use of an impactor limited sampling to primarily submicron particles (<1.5 \( \mu m \) in diameter) prior to CFDC processing in order to distinguish ice crystals nucleated in the CFDC from non-activated aerosol particles using optical methods.

2.7. Laboratory Measurements

[19] For comparison to ICE-L results, dry lakebed crust particles collected from the surface of Owens (dry) Lake, CA were aerosolized by dry generation using \( N_2 \), as detailed previously by Sullivan et al. [2009]. The dust potential and chemistry of Owens Lake has been studied in detail [e.g., Cahill et al., 1996; Gill et al., 2002], making it a good comparison source; however, this does not suggest it as the exact source of the ICE-L particles. The A-ATOFMS,
described above, was used to measure particle chemistry in tandem with measurements of CCN activity. For CCN analysis, aerosolized particles were passed through a 210Po neutralizer and then size-selected (80, 90, and 100 nm) using a differential mobility analyzer (DMA, Model 3081, TSI Inc.) to obtain a monodisperse aerosol distribution. Total particle concentrations were measured by a condensation particle counter (CPC, Model 3010, TSI Inc.) and compared to the number concentrations of particles activated within a miniature cloud condensation nuclei counter (CCNc) at supersaturations of ~0.1–1.1% [Roberts and Nenes, 2005]. The supersaturation of the CCNc is a function of the column temperature gradient (dT), which was calibrated using (NH4)2SO4 (Aldrich, 99.999% purity). Activation curves of CCN/CN ratios, providing the fraction of activated particles, were generated by scanning through supersaturations (column dT) at a fixed dry diameter; the S∞ is defined as CCN/CN = 0.5. The contribution of multiply charged particles was corrected using the method of Rose et al. [2008]. Herein, CCN activity is presented as the single hygroscopicity parameter kappa (κ) [Petters and Kreidenweis, 2007], which was determined by plotting S∞ for each dry diameter (Dd) on a log-log plot with the different S∞-Dd pairs for each sample falling on κ isolines. Uncertainty in κ is from errors associated with the CCNc supersaturation calibration and from selecting a monodisperse aerosol using a DMA.

3. Results and Discussion

3.1. Single-Particle Mixing State and Origin of Playa Salts

While flying through mixed-phase orographic wave clouds over Wyoming during ICE-L, the A-ATOFMS identified playa salts among the residues of cloud droplets and ice crystals. Figure 1 shows example A-ATOFMS positive and negative ion mass spectra of an individual salt residue from a RF03 evaporated cloud droplet. These salt particles were characterized by intense sodium (m/z 23, Na+), magnesium (m/z 24, Mg2+), potassium (m/z 39, K+), and calcium (m/z 40, Ca2+) ion peaks with less intense positive ions at m/z 55(KO2), 56(KOH+/CaO2), 81(Na2ClO4), and 97(Na2ClO3). The negative ion mass spectra featured distinguishing peaks at m/z −16(O−), −17(OH−), −26(CN−), −35(Cl−), −42(CNO−), −58(NaCl), and −93(NaCl2). The high relative intensities of the magnesium, calcium, and negative ion organic nitrogen markers, in particular, distinguish these playa salts from sea salt [Gard et al., 1998]. A fraction of the negative ion mass spectra also contain carbon cluster ions (Cn), silicates, phosphate, nitrate, and/or sulfate. The mass spectral signatures of these salts are expected for the typical evaporite minerals associated with playas, including halite (NaCl), gypsum (CaSO4 · 2H2O), mirabilite (Na2SO4 · 10H2O), thenardite (Na2SO4), epsomite (MgSO4 · 7H2O), trona (NaHCO3 · Na2CO3 · 2H2O), sylvite (KCl), carnallite (KCl · MgCl2 · 6H2O), calcite/aragonite (CaCO3), dolomite (CaMg(CO3)2), hanksite (Na32K(SO4)9(CO3)2Cl), northupite (Na32Mg(CO3)2Cl), and aphthitalite (K2SO4) [Bryant et al., 1994; Drosie, 1961; Güven and Kerr, 1966; Lowenstein and Hardie, 1985].

[21] To further confirm the A-ATOFMS salt signature as from playas and dry lakebed crust particles from Owens Lake, CA were compared using the A-ATOFMS analysis method of Toner et al. [2008]. The average mass spectral signature of the Owens Lake crust was compared to the average mass spectrum of the cloud residue salts using calculated dot products for the ion peaks in the spectra. Lower dot products indicate less similarity between the two sets of particles; whereas high dot products indicate greater similarity. In this case, the mass spectral comparison showed a strong similarity (dot product = 0.7) between the Owens Lake crust and cloud residue salt particles, supporting the ICE-L signature as playa dust, as opposed to sea salt (dot product = 0.58); it is important to note that a higher
dot product would likely be obtained if the exact playa source of the cloud residues could have been measured, particularly at the time of the dust lofting.

[22] STEM-EDX analysis of cloud residues also showed the presence of salts, defined as containing Na, K, Ca, and Mg with S or Cl (Table 1). An example STEM image and corresponding EDX spectrum are shown in Figure 2 for a single RF03 cloud residue salt, likely from a liquid droplet. In addition to the salt-defining elements, this particle contains C, O, F, Zn, and P above background levels. This signature is in excellent agreement with that determined by A-ATOFMS, as discussed above, as well as previous proton-induced X-ray emission (PIXE) analysis of playa sediments, where samples contained Na, Ca, Si, and trace metals [Gill et al., 2002]. By comparison, for Owens Lake (CA) playa sediments, silicon and aluminum contents ranged from ∼16–22% and ∼3–5% by mass, respectively; Zn content was found to be ∼46–100 ppm [Gill et al., 2002]. It should also be noted that a “halo” surrounds the solid residue salt shown in Figure 2. During previous aircraft-based aerosol sampling over Montana, NaCl particles from the Great Salt Lake area were detected simultaneously with soil particles [Hobbs et al., 1985]. Using scanning electron microscopy (SEM)-EDX, “haloes” containing Si and Cl with trace S were found surrounding soil particles containing Si and Al with trace Fe and K [Hobbs et al., 1985].

[23] To further characterize the cloud residue salts with particular focus on the organic component, STXM/NEXAFS was utilized to examine the EC, OC, K, and regions rich in In elements within individual particles. Figure 3 shows chemical component maps for a single mixed organic/inorganic (OCIn) RF03 cloud residue collected during the 19 μm CVI cut-point sampling period with liquid droplets, as discussed below. For this particular particle, Ca, K, COOH

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Figure 2. STEM image of cloud residue salt with corresponding EDX spectrum.

Figure 3. (a) STXM/NEXAFS maps of inorganic, potassium, sp² bonds, COOH and CO₃ for a single residue of a cloud droplet (>19 μm). (b) STXM/NEXAFS spectrum over the area of the individual cloud droplet residue shown in Figure 3a.
groups, and CO$_3$ are all internally mixed. As discussed above, Ca, K, and carbonate are common components of playa evaporite minerals. Based upon STXM/NEXAFS analysis, the “haloes” observed by STEM-EDX were likely from an oxidized organic coating, presumably from cloud processing [Chen et al., 2007]. This is further confirmed by an elevated C-ToF-AMS O/C ratio (0.84 ± 0.36), indicative of oxygenated organic species [Aiken et al., 2008] present during cloud droplet residue sampling. Indeed, the A-ToFMS observed ion peaks of possible organic acids, including $m/z$ –43(CH$_3$COH/HCNO$^-$), –73(C$_2$H$_5$O$_3^-$), glyoxylic acid), –159(C$_3$H$_7$O$_4^-$, pimelic acid), and –173 (C$_4$H$_7$O$_4^-$, suberic acid), within the mass spectral signatures of the playa salt cloud droplet residues.

[24] Overall, the ICE-L chemical signatures observed for the individual playa salt particles are consistent with previous measurements of playa dust. For dust from Lake Ebinur (China) and Owens Lake (CA), playa dust was found to be ∼10–30% soluble salts, predominantly sodium, calcium, and magnesiu chloride and sulfate [Abudawalli et al., 2008; Reheis, 1997; Reheis and Kuhl, 1995]. Daily carbonate and total salt fluxes were found to be correlated with typical carbonate mass fractions of up to ∼25% [Reheis, 1997]. Compared to playa sediments, increased fractions of soluble salts and carbonate have been observed in the playa crustal dust from Owens Lake [Reheis, 1997]. Further, playa dust has been found to be up to ∼30% organic matter by mass [Reheis, 1997].

[25] While our focus is on RF03, sodium chloride salts enriched in Mg, Ca, and K were observed during three of seven ICE-L orographic wave cloud flights (Figure 4). Examination of HYSPLIT (Draxler and Rolph, NOAA Web site, 2003) air mass back trajectories for the Mg-Ca-K-enriched salt flights (RF02, RF03, RF04) show transport over the northern edge of the Great Basin region of Nevada and Utah, the largest dust source region in the U.S. [Prospero et al., 2002; Washington et al., 2003]. The Great Basin region, once the sites of Lake Lahontan and Lake Bonneville during the Quaternary period, contains many playas [Güven and Kerr, 1966], as shown in Figure 4 [Soller and Reheis, 2004]. During flights in which these enriched salts were not observed (RF01, RF06, RF09, RF12), the air masses followed more northerly trajectories, over land where playas are not present in abundance. For the “salt” flights, air mass back trajectories show transport times of less than 18 h from northern Utah and Nevada. While the vertical profiles of these HYSPLIT back trajectories do not show transport within the boundary layer (transport ∼2–6 km above ground level (AGL)), previous measurements of playa dust storms have shown high wind speeds [Pelletier, 2006], rapid lofting to high altitudes (>2 km AGL) [Cahill et al., 1996; Reid et al., 1994], as well as associated long-range transport [Formenti et al., 2003]. Indeed, for RF03, HYSPLIT forward trajectories from both the Black Rock Desert in Nevada and the Bonneville Salt Flats in Utah show lofting to >2 km AGL with transport to Wyoming, in agreement with instability above ∼2 km AMSL based on radiosonde data taken over Salt Lake City. Further, primarily southwesterly wind gusts of up to 18 m/s were observed over northwestern Nevada with westerly wind gusts of up to 12 m/s measured over northwestern Utah. For RF03, elevated mass concentrations of particulate matter less than 10 µm or 2.5 µm (PM$_{10}$ and PM$_{2.5}$) were observed in Salt Lake City and Logan City, UT on November 16, 2007. For example, at the Utah State Division of Air Quality North Salt Lake monitoring site in Salt Lake City, the 24 h PM$_{10}$ mass concentration was 86 µg/m$^3$ on Nov. 16, compared to the month average of 44 µg/m$^3$ (standard deviation (σ) 25 µg/m$^3$). Likewise, at the Rose Park monitoring site in Salt Lake City, the 24 h PM$_{2.5}$ mass concentration was 30.3 µg/m$^3$ on Nov. 16, compared to the month average of 13.5 µg/m$^3$ (σ 7.3 µg/m$^3$). While the exact playa source(s) could not be identified, together these trajectories and PM observations further support the detection of playa salts from the Great Basin region.

![Figure 4](image-url). HYSPLIT air mass back trajectories (up to 48 h, with dots shown every 6 h) for ICE-L orographic wave cloud flights, classified as either having Na-Mg-Ca-K-Cl salts present (green) or not (black). For reference, a map of playa sediments [Soller and Reheis, 2004] is shown.

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3.2. Supercooled Liquid Droplets

[26] To evaluate the ability of playa salts to act as CCN, the chemistry and CCN ability of Owens (dry) Lake crust was evaluated in the laboratory. Having a similar A-TOFMS mass spectral signature to the cloud residue salts, the Owens (dry) Lake crust had an elevated CCN activity (high hygroscopicity) \( (\kappa = 0.84 \pm 0.10) \) compared to that measured previously by Koehler et al. [2009] for most mineral dust \( (\kappa \leq 0.08). \) Previous CCN measurements of dry-generated 200 nm Owens Lake dust found 35% of the particles to have a \( \kappa \) of 0.69 with 65% having a \( \kappa \) of 0.05; particles <100 nm were more CCN-active with \( \kappa \) values up to 1.07 [Koehler et al., 2007]. For comparison, highly CCN-active particles are considered to be those with \( \kappa \) values between 0.5 and 1.4 [Petters and Kreidenweis, 2007]. NaCl has a \( \kappa \) of 1.12–1.28 [Petters and Kreidenweis, 2007]; as a pure salt, it is slightly more hygroscopic than sea salt [Niedermeier et al., 2008], showing that playa salts can have CCN activities similar to those of sea salt. Magnesium chloride and calcium chloride, enriched in playa salts compared to sea salt, have significantly lower deliquescence relative humidities (DRHs) \( (\text{MgCl}_2 \cdot 6\text{H}_2\text{O}, 33\% ; \text{CaCl}_2 \cdot 6\text{H}_2\text{O}, 28.3\%) \) compared to sodium chloride \( (\text{NaCl}, 75.3\%) \) [Tang et al., 1997] with mixtures of these salts having even lower mutual DRHs [Kelly and Wexler, 2006]. In addition, the internal mixing of oxidized organics with salts is suggested to enhance water uptake at low relative humidities [Hersey et al., 2009], as well as to reduce or eliminate clear deliquescent phase transitions compared to those expected for inorganics only [Carrico et al., 2008]. Thus, it is likely that hygroscopic playa salts, observed as dry cloud residues, retained water under clear air conditions (average relative humidity of 42\% (\( \sigma \) 29\%)).

[27] To further understand the activation of the supercooled liquid droplets observed within the orographic wave clouds during RF03, the maximum cloud supersaturation was modeled using the 1-D parcel model described by Heymsfield et al. [1991]. Using the average temperature \( (\sim 21.8^\circ \text{C}) \), pressure \( (\sim 333 \text{ Torr}) \), and altitude \( (6573 \text{ m}) \) with a peak updraft velocity of 2.6 m/s of the first (and lowest altitude) liquid phase cloud pass, the maximum supersaturation was estimated to be up to \( \sim 8\% \), similar to previous observations of supercooled liquid wave clouds [Heymsfield and Miloshevich, 1993]. While 8\% supersaturation may be an overestimate, supersaturations reach very high levels because the droplet growth rates, which deplete the buildup of supersaturation, are relatively low at a temperature of \( \sim 20^\circ \text{C} \) [Heymsfield and Miloshevich, 1993]. Collision-coalescence was not observed to occur in the model for the given microphysical constraints and timescale of the cloud parcels; similarly, CPI, radar, and lidar data showed no evidence of drizzle formation. Thus, in contrast to the warm clouds observed by Rudich et al. [2002], the playa salts were not able to or observed to initiate a precipitation mode during RF03. To examine the particles activated as cloud droplets during RF03, measured clear air particle and cloud droplet number concentrations were compared. During the clear air measurement periods \( (\theta_e = 318.3–323.8 \text{ K}) \), number concentrations of particles 0.1–1.0 \( \mu \text{m} \) in diameter, measured by an UHSAS, averaged 25 \( \text{ cm}^{-3} \) (maximum 164 \( \text{ cm}^{-3} \) with average CN concentrations of 217 \( \text{ cm}^{-3} \) (maximum 704 \( \text{ cm}^{-3} \)). During liquid-only sampling, average cloud droplet number concentrations, as measured by the CDP, were 64 \( \text{ cm}^{-3} \) (maximum 125 \( \text{ cm}^{-3} \)) with an average of 88\% (\( \sigma \) 40\%) of droplets larger than 7 \( \mu \text{m} \).

[28] To examine the variation of aerosol chemistry with cloud droplet diameter, CVI sampling was completed at two different CVI cut-points \( (7 \mu \text{m} \text{ and } 19 \mu \text{m}) \). Sampling of cloud droplets \( >7 \mu \text{m} \) was completed at altitudes of 6.6–7.7 km and ambient temperatures of \( \sim 21 \) to \( \sim 32^\circ \text{C} \); sampling of cloud droplets \( >19 \mu \text{m} \) was completed at 7.2 km and \( \sim 26 \) to \( \sim 27^\circ \text{C} \). Average cloud droplet size distributions during these two periods are shown in Figure 5; residue chemistry, as measured by the A-TOFMS and C-ToF-AMS are also shown. During the \( >19 \mu \text{m} \) period, the average cloud droplet number concentration, measured by the CDP, was 66 \( \text{ cm}^{-3} \) (maximum 125 \( \text{ cm}^{-3} \)) with a small contribution of cloud ice particles \( (>50 \mu \text{m}) \) of 2 \( \text{ L}^{-1} \) on average (maximum of 18 \( \text{ L}^{-1} \)) measured by the 2DC probe; 35\% (\( \sigma \) 18\%) of droplets were larger than 19 \( \mu \text{m} \). As shown in Figures 5b and 5e, the playa salts were preferentially found in larger cloud droplets with the number fraction increasing from 57 \( \pm 10\% \) to 90 \( \pm 6\% \) with the increase in CVI cut-point. For residues of the larger cloud droplets, STXM/NEXAFS analysis classified all measured particles as internally mixed organic carbon and inorganic components, with \( \sim 71\% \) by number mixed with carbonate and \( \sim 29\% \) mixed with potassium as well (Table 2). Concurrent with the increase in the fraction of playa salts, the C-ToF-AMS non-refractory mass fractions showed an increase in organics \( (37 \pm 17\% \text{ to } 80 \pm 4\%) \) and decrease in sulfate and ammonium (Figure 5); as shown by the A-TOFMS mass spectra and confirmed with STXM/NEXAFS, the majority of the organics measured by the C-ToF-AMS were internally mixed with the playa salts. Although within uncertainties, the C-ToF-AMS observed a higher O/C ratio [Aiken et al., 2008] for the organic species observed during the \( >19 \mu \text{m} \) cloud droplet residue sampling than during the \( >7 \mu \text{m} \) sampling \( (1.19 \pm 0.15 \text{ versus } 0.84 \pm 0.36, \text{ respectively}) \), suggesting that the larger cloud droplets may have undergone additional cloud processing during transport. In addition to the probable organic acids (glyoxylic, pimelic and suberic acid) observed by the A-TOFMS to be internally mixed with the \( >7 \mu \text{m} \) cloud droplet playa salt residues, additional ion peaks of potential organic acids were observed for the \( >19 \mu \text{m} \) cloud droplet playa salt residues: \( \text{m/z } 45 \) (HCOO, formic acid), \( \sim 87 \) (\( \text{C}_3\text{H}_4\text{O}_3 \), pyruvic acid), and \( \sim 103 \) (\( \text{C}_3\text{H}_4\text{O}_4 \), malonic acid) [Sullivan and Prather, 2007]. Considering the oxidation pathways of organic gas phase species, oxalic, glutaric, adipic, and pyruvic acids have previously been suggested as the primary contributors to organic mass production during aqueous phase processing [Ervens et al., 2004]. The O/C ratio observed by the C-ToF-AMS during \( >19 \mu \text{m} \) sampling is among the highest reported, significantly above reported values for laboratory secondary organic aerosol [e.g., Aiken et al., 2008; Hildebrandt et al., 2009], highlighting the need to further understand cloud processing of organics. Previously, Sun et al. [2009] observed an O/C ratio of 1.08 for organic aerosol, hypothesized to have undergone aqueous phase processing during transport over the Pacific Ocean to Whistler Mountain (Canada).

[29] While organics are generally considered to be less efficient CCN than most inorganic species, small amounts of
salt (<5% of dry mass for NaCl) have been found to greatly decrease the $S_c$ of even slightly soluble organic compounds [Bilde and Svenningsson, 2004]. At the vaporization temperature utilized by the C-ToF-AMS during ICE-L (550–560°C), the C-ToF-AMS is unable to efficiently detect refractory species present in sea salt (e.g., NaCl); however, due to the low ionization potentials of Na and K, a small fraction of the salts may undergo surface ionization [Allan et al., 2004]. Through comparison of the C-ToF-AMS non-refractory mass concentrations with residue volume concentrations measured by the CVI OPC (and assuming a density of 1.7 g cm$^{-3}$), it is estimated that a significant fraction of the particle mass was likely refractory during both the >7 and >19 μm periods, respectively; this refractory mass was not detected by the C-ToF-AMS. However,

<table>
<thead>
<tr>
<th>Type</th>
<th>&gt;19 μm Liquid</th>
<th>Ice (&gt;7 μm)</th>
<th>Clear Air</th>
</tr>
</thead>
<tbody>
<tr>
<td>OC</td>
<td>0</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>OCEC</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>OCECIn</td>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>OClO$_x$</td>
<td>17, (12 CO$_2$), (5 K)</td>
<td>6, (6 CO$_2$), (3 K)</td>
<td>7, (2 CO$_2$), (1 K)</td>
</tr>
</tbody>
</table>

*Numbers in the table represent the number of particles in each sample that contained specified particle type: organic carbon (OC), organic carbon-elemental carbon (OCEC), organic carbon-elemental carbon-inorganic (OCECIn), and organic carbon-inorganic (OClO$_x$). The numbers of particles found to have CO$_2$ and/or K are shown in parentheses. For the three sample periods (>19 μm liquid, ice, and clear air), the total numbers of particles analyzed were 9, 9, and 17, respectively.
with the largest drops forming on

113 L

(a) Number fractions of particle types measured by the A-ATOFMS for residues of cloud ice >7 μm; 80 particles were chemically analyzed; 78% of the salts were observed to be internally mixed with organics. (b) Non-refractory mass fractions measured by the C-ToF-AMS for residues of cloud ice >7 μm.

Figure 6. (a) Number fractions of particle types measured by the A-ATOFMS for residues of cloud ice >7 μm; 80 particles were chemically analyzed; 78% of the salts were observed to be internally mixed with organics. (b) Non-refractory mass fractions measured by the C-ToF-AMS for residues of cloud ice >7 μm.

qualitatively, the C-ToF-AMS observed higher Na, K, and Cl signals in the larger cloud droplets, in agreement with the enrichment of playa salts in the larger cloud droplets, as measured by the A-ATOFMS.

[30] The observed enrichment of playa salts in the larger cloud droplets is consistent with the relationship between cloud droplet size and S, with the largest drops forming on the largest, most hygroscopic CCN [Hudson and Rogers, 1986]. Twyoh et al. [1989] used a parcel model to predict size-dependent droplet chemistry for an ammonium sulfate small aerosol mode and a sea salt large aerosol mode; similar to that observed herein, sea salt was calculated to be within the largest droplets, showing that bulk droplet composition does not represent the composition of individual cloud droplets. For coastal stratus clouds heavily influenced by sea salt, Munger et al. [1989] observed higher Na⁺, Ca²⁺, and Mg²⁺ concentrations in larger droplets with higher concentrations of SO₄²⁻, NO₃, NH₄⁺, and H⁺ in smaller droplets, similar to that observed for the RF03 residues. It is predicted that cloud drops will retain the size-resolved chemistry of the original CCN until coalescence produces drizzle-sized drops [Ogren and Charlson, 1992]. Previously, giant aerosol particles primarily consisting of NaCl, observed over the eastern Great Lakes region of the United States and Canada, were found to contribute to the presence of supercooled large drops [Lasher-Trapp et al., 2008]. In addition, satellite measurements showed that playa dust from the Aral Sea region increased cloud drop size and promoted precipitation [Rudich et al., 2002]. For particle sizes similar to large mineral dust [Patterson and Gillette, 1977; Reid et al., 1994], playa salts can serve as giant CCN [Andreae and Rosenfeld, 2008], increasing cloud droplet size similar to large hygroscopic salts used in cloud seeding studies [Drofa, 2006; Ghate et al., 2007].

3.3. Cloud Ice Residues

[31] During sampling in cloud ice regions, the 2DC probe concentrations averaged 57 L⁻¹ (± 113 L⁻¹) with sampling at 6.5–7.7 km. Although flight level temperatures ranged from −21 to −31°C, the wave cloud tops reached below −38°C, resulting in influence from homogeneous freezing (P. R. Field et al., Contrasting ice nucleation in two lee wave clouds observed during the ICE-L campaign, manuscript in preparation, 2010), discussed below. Similar to the >19 μm droplet residues, the ice residues were dominated by playa salts internally mixed with organics, as shown in Figure 6. This was further confirmed by STXM/NEXAFS analysis, which showed that 6 out of the 9 analyzed ice residues were internally mixed organics and inorganics (Table 2). A-ATOFMS results showed that approximately half as many ice salt residues were internally mixed with sulfate and/or nitrate (28 ± 5%) compared to the cloud droplet salt residues (69 ± 13%) (Figure 7). This is consistent with lower mass fractions of sulfate and nitrate measured by the C-ToF-AMS for the ice residues compared to the cloud droplet residues (Figures 5 and 6). Similarly, Cziczo et al. [2004] observed large, unreacted sea salt particles as residues of cirrus ice produced by homogeneous freezing. It was hypothesized that the preferential freezing of sea salt was due to the extensive water uptake of NaCl, resulting in larger, more dilute, sea salt particles which were more likely to freeze compared to the smaller, more numerous sulfate/K/organic/nitrate particles observed in the clear air [Cziczo et al., 2004]. Similarly, organic carbon/sulfate particles were not observed by the A-ATOFMS for the RF03 ice residues. Similar to the >19 μm cloud droplet residues, the O/C ratio observed by the C-ToF-AMS during ice residue sampling was elevated (1.15 ± 0.25), indicative of highly oxygenated organic aerosol [Aiken et al., 2008]. In addition to the probable organic acid ion peaks observed during cloud droplet residue sampling, oxalic acid (m/z = 89(C₂H₂O₄) and m/z = 179((C₂H₂O₄)H⁺)) [Sullivan and Prather, 2007] was observed to be preferentially mixed with the playa salt ice residues.

[32] Despite sampling at temperatures above −31°C, the wave cloud tops reached below −38°C as confirmed by the radar signature and a kinematic 1-D model (Field et al., manuscript in preparation, 2010). This temperature would allow homogeneous freezing to occur in dilute solution droplets and activated cloud droplets [DeMott and Rogers, 1990], as confirmed by the radar signature and a kinematic 1-D model (Field et al., manuscript in preparation, 2010). During RF03, homogeneously nucleated ice particles were observed to fall from above the sampling altitude, as well as follow the wave cloud streamlines to a lower altitude; this ice greatly influenced the sampled ice residues (Field et al., manuscript in preparation, 2010), suggesting that the majority of the ice playa salt residues were homogeneously frozen at higher altitudes, similar to the ice sea salt residues observed by Cziczo et al. [2004] and Twyoh and Poellot [2005]. In warmer mixed-phase cloud regions, riming, as indicated in CPI images (not shown), would have
contributed additional soluble salts to the ice. However, in addition to homogeneous freezing, the Field et al. (manuscript in preparation, 2010) model also indicated a small contribution (∼6 L⁻¹ to the total ice particle concentration of ∼50 L⁻¹) of heterogeneous condensation/immersion freezing. This is consistent with a maximum ice nuclei concentration of 3 L⁻¹ measured by the CFDC operating at −31°C and a supersaturation (with respect to water) of up to 1%, for the ice residues. For previous marine anvil cirrus ice measurements by the CFDC, a small fraction of the salts were found to act as ice nuclei, freezing below water saturation [Prenni et al., 2007]. Less hygroscopic playa dust has been shown to nucleate ice heterogeneously at lower relative humidities than required for homogeneous freezing of aqueous particles at temperatures less than −40°C; smaller contributions to heterogeneous ice formation from about −37 to −40°C were also observed [Koehler et al., 2007]. However, the study by Koehler et al. [2007] did not report ice formation for activated fractions of <1% or examine playa dust greater than 400 nm in size, so potential heterogeneous ice formation by small fractions of particles at temperatures above about −37°C are not resolved. For the RF03 ice salt residues, 21 ± 5% were found to contain silicates (20 ± 4% of the total ice residues), while none of the cloud droplet salt residues contained silicates. We can assume that ∼2–3% of these “dust-like” playa salts may activate as ice nuclei [Field et al., 2006; Prenni et al., 2009]. Considering the concentration of residues measured by the CVI OPC, we estimate that an average of 2.9–4.4 L⁻¹ (range of 1.1–8.0 L⁻¹ given uncertainties) playa dust may have been IN active. This estimate is in excellent agreement with the modeled (∼6 L⁻¹) and measured IN concentrations (maximum 3 L⁻¹, average 1.0 L⁻¹, σ 0.7 L⁻¹), suggesting that the silicate-containing playa salts may have served as heterogeneous ice nuclei. Further, during sampling regions containing only cloud droplets, IN concentrations were below the detection limit (0.1 L⁻¹) of the CFDC, consistent with the relatively low abundance of silicate-containing particles present.

[33] It should be noted that during cloud ice sampling, a significant fraction (35 ± 5%) of the total ice residue A-ATOFMS mass spectra also contained markers of inlet metals, such as chromium (m/z 52, Cr⁺), manganese (m/z 55, Mn⁺), iron (m/z 56, Fe⁺), and molybdenum (m/z 98, Mo⁺), indicating that a fraction of the ice particles impacted the stainless steel CVI aircraft inlet during sampling, as previously observed in even higher concentrations by Murphy et al. [2004], likely due to higher aircraft speeds. Indeed, a greater concentration (∼50 times) of ice residues >110 nm, was measured by the CVI OPC compared to ice crystal concentrations, as measured by the 2DC, further indicating the occurrence of ice crystal shattering within the CVI inlet, as well as possible residue multiplication due to dissolved salts within homogeneously frozen ice crystals. However, while a fraction of the positive ion mass spectra were influenced by these metals, the distinguishing positive ion mass spectral signature of the playa salt (i.e., Na, K, Mg, Ca) was still present, allowing the residues to be classified as salts. Further, the negative ion mass spectra of these particles were identical to salt particles that had not collided with the inlet (i.e., did not contain Cr, Mn, Fe, Mo), showing how the dual-polarity mass spectrometer of the A-ATOFMS can help distinguish the source signature of the residue despite interferences from the inlet metals. Although inlet metals were only observed during ice-containing sampling periods, no correlation was observed between ice particle size (or riming) and the presence of inlet metals in the A-ATOFMS mass spectra. Further, ice nuclei concentrations measured within clouds behind the CVI during ICE-L did not generally seem enhanced relative to IN concentrations measured.

Figure 7. Number fractions of cloud residues identified as salts, measured by the A-ATOFMS, found to be internally mixed with silicates or nitrate and/or sulfate; 75, 104, and 13 salt particles, respectively, were chemically analyzed for the three cloud periods.
3.4. Mixed Phase Cloud Residues

Concurrent CVI sampling by the A-ATOFMS, C-ToF-AMS, and impactor for STEM-EDX analysis was conducted for mixed phase (coexisting supercooled liquid droplets and ice particles) cloud residues, allowing a comparison of the three aerosol chemical analysis techniques (Figure 8). During mixed phase sampling (−20 to −32°C, 6.4−7.9 km), supercooled droplets, measured by the CDP, were present at an average of 44 cm$^{-3}$ (maximum 144 cm$^{-3}$); cloud ice particles, measured by the 2DC probe, were present at an average of 10 L$^{-1}$ (14 L$^{-1}$). The relative contribution of the ice particles to the mixed phase residue number is expected to be enhanced since residue multiplication can occur from ice particles larger than ∼75 μm (aerodynamic diameter) that may break up within the CVI.
inlet [Twohy et al., 2003], although residue mass will be unaffected. The number fraction of mixed-phase residues containing inlet metals, as measured by the A-ATOFMS, was minor (4 ± 2%). For the mixed-phase STEM-EDX sample, 3% of the small impactor stage residues and 7% of the large impactor stage residues contained stainless steel markers (not shown), in agreement with that observed by the A-ATOFMS. As expected for these mixed phase residues, the number fraction of playa salts (84 ± 3%), as measured by the A-ATOFMS, was between that of the droplet residues (57 ± 10%) and ice residues (94 ± 3%) (Figures 5, 6, and 8). Similarly, the organic mass fraction, as measured by the C-ToF-AMS, increased from droplets (37 ± 17%) to mixed phase (59 ± 20%) to ice (82 ± 7%), with a corresponding increase in the O/C ratio (0.84 ± 0.36 to 1.00 ± 0.31 to 1.15 ± 0.25, respectively). While the number fraction of salts containing sulfate and/or nitrate was similar for the mixed phase (70 ± 4%) and droplets (69 ± 13%) (Figure 7), the mass fraction of sulfate in all residues decreased from droplets (57 ± 10%) to ice (48 ± 5%), in agreement with the presence of large playa salts with a size distribution similar to that of mineral dust [Patterson and Gillette, 1977]. STEM-EDX showed a greater fraction of sulfate particles for the small stage (13 ± 3%) compared to the large stage (5 ± 2%, not shown).

[35] For comparison between STEM-EDX, A-ATOFMS, and C-ToF-AMS, one must first consider the relationship between particle geometric diameter \(d_g\), measured by STEM, and vacuum aerodynamic diameter \(d_{va}\), measured by the A-ATOFMS and C-ToF-AMS. For spherical particles with no voids, \(d_g\) and \(d_{va}\) are related by the following equation:

\[
d_{va} = d_g \frac{\rho_p}{\rho_0}
\]

(2)

where \(\rho_0\) is the standard density (1 g cm\(^{-3}\)) and \(\rho_p\) is the average particle density for a multiple component particle [DeCarlo et al., 2004]. Thus, the size ranges of the A-ATOFMS and C-ToF-AMS are most similar to the small impactor stage, which collected approximately 0.08 to 0.42 \(\mu m (d_g)\) or 0.136 to 0.714 \(\mu m (d_{va})\) particles, compared to the large impactor stage (>0.42 \(\mu m (d_g)\) or >0.714 \(\mu m (d_{va})\)), given an assumed effective density of \(\sim 1.7\) g m\(^{-3}\) for the salt particles, as observed previously for ambient sea-salt particles at low relative humidity [Moffet et al., 2008b]. It also should be noted that the STEM-EDX sample was collected for a shorter time period than the A-ATOFMS and C-ToF-AMS mixed phase residue samples, which may yield minor differences in the samples.

[36] Comparison of the number fractions of different particle types measured by the A-ATOFMS and STEM-EDX small impactor stage (Figures 8a and 8b) shows a smaller fraction of salts detected by STEM-EDX compared to the A-ATOFMS, although some of the salts are included in the STEM-EDX “mixed” particle category. In a previous comparison ATOFMS and SEM-EDX of K- and Na-containing particles in Mexico City, the ATOFMS observed twice as many particles containing Cl\(^-\) compared to SEM-EDX due to the increased sensitivity of the ATOFMS to this species, as well as possible Cl\(^-\) loss reactions following filter collection [Moffet et al., 2008a]. In agreement, the A-ATOFMS observed 6 ± 2% of the mixed phase residues to be silicate-containing salts, whereas STEM-EDX categorized 6 ± 2% of the residues as crustal dust. The large fraction of industrial metals observed by STEM-EDX may have been above the \(d_{va}\) range of the A-ATOFMS due to high densities, accounting for this discrepancy between the two results. Last, STEM-EDX identified a greater fraction of sulfate particles compared to the A-ATOFMS (13 ± 3% versus 2 ± 1%, respectively); however, it is likely that a fraction of these particles were internally mixed with organic species and/or soot and, thus, were classified by the A-ATOFMS as such. Overall, A-ATOFMS and STEM-EDX were found to be in good agreement.

3.5. Comparison to Clear Air

[37] In the ambient air surrounding the orographic wave clouds, the A-ATOFMS observed the majority of submicron particles to be either externally mixed sulfate or an internal carbonaceous/sulfate mixture (biomass, organic carbon, and soot particle types), which is consistent with the majority of the non-refractory submicron mass being composed of organics and sulfate, as measured by the C-ToF-AMS (Figure 8). Given the uncertainties associated with the observed low mass loadings, the C-ToF-AMS did not observe a significant difference in the mass fractions of organics, sulfate, nitrate, and ammonium between the ambient air and cloud droplet residues; similar O/C ratios (0.77 ± 0.28 versus 0.84 ± 0.36) were observed for the organic aerosol as well. Relative to cloud sampling, externally mixed sulfate particles, measured by A-ATOFMS and STEM-EDX, were present in higher concentrations in the clear air. In addition, organics were enriched compared to sulfate (C-ToF-AMS) in the ice residues compared to the clear air, possibly due to the internal mixing of a fraction of the organics with playa salts, as confirmed by STXM/NEXAFS analysis. A-ATOFMS and STEM-EDX results both show an enrichment of the playa salts by number in the cloud residues (>80 nm \(d_g\)) compared to the clear air. Similar enrichments were observed by Cziczo et al. [2004], wherein 26% of homogeneously frozen ice residues were sea salt compared to <1% outside of the cirrus; however, these anvil cirrus clouds were associated with convective systems which were hypothesized to transport sea salt from the surface. Twohy and Anderson [2008] observed similar enrichments of salts among <0.2 \(\mu m\) cloud droplet residues collected within eastern Pacific stratocumulus, Caribbean Sea small cumulus, and Indian Ocean small cumulus compared to ambient particles.

[38] As discussed in Section 3.2, average CN clear air concentrations were 217 cm\(^{-3}\) (maximum 704 cm\(^{-3}\)) with average submicron (0.1–1.0 \(\mu m\)) number concentrations of 25 cm\(^{-3}\) (maximum 164 cm\(^{-3}\)); in comparison, cloud droplet number concentrations averaged 64 cm\(^{-3}\) (maximum 125 cm\(^{-3}\)). This comparison suggests that most particles greater than 100 nm, as well as a fraction of 13–100 nm particles, would be expected to serve as nuclei for cloud droplet formation. However, CVI cloud droplet residues >110 nm in diameter had average number concentrations of only 3.3 cm\(^{-3}\) (maximum 6.9 cm\(^{-3}\)), despite an average of
88% of droplets possessing diameters larger than 7 μm, as measured by the CDP. CVI concentrations were much greater for residues <110 nm (average 21 cm⁻², maximum 47 cm⁻²). Also, during liquid droplet (>7 μm) sampling, the C-ToF-AMS measured an average non-refractory PM₁ mass concentration of 0.035 ± 0.006 μg m⁻³, which was only 13 ± 5% of that observed for the clear air (0.27 ± 0.09 μg m⁻³). However, this particularly low fraction may be partially due to preferential activation of larger, more refractory salts (which are not efficiently detected by C-ToF-AMS) compared to smaller organic-sulfate particles. Thus, it appears that all particles >110 nm did not result in cloud droplets >7 μm, suggesting that playa salts are preferentially activated over other particle types in this size range, as measured by A-ATOFMS and STEM-EDX.

4. Conclusion

[39] Playa salts, comprised of Na–K–Mg-Ca–Cl salts, internally mixed with oxidized organics, presumably produced by cloud processing, and carbonate have been shown to act as cloud nuclei in orographic wave clouds over Wyoming. Similar to previous observations of sea salt in marine environments, these playa salts were enhanced in larger droplets (>19 μm) compared to smaller droplets (>7 μm). Enhancement of the playa salts in ice, compared to the clear air and droplets, was also observed, due to influences of homogeneous freezing and riming. A small fraction of silicate-containing playa salts are hypothesized to be important in the heterogeneous ice nucleation observed in the wave clouds. The unique ability of the A-ATOFMS to distinguish between sea salt and playa salts was essential for the major findings of this study. Excellent agreement was achieved between the aerosol chemical analysis techniques used in this study (A-ATOFMS, C-ToF-AMS, STEM-EDX, STXM/NEXAFS). The techniques are complementary, combining together to provide unprecedented detail on single-particle mixing state with quantification of non-refractory particle components.

[40] Although the generation of wind-blown dust from playas is relatively well-known, the relative role of playa dust globally in cloud formation is highly uncertain. Previous laboratory studies of the CCN and IN abilities of playa dust have suggested their importance in cloud formation [Koehler et al., 2007]. Further, satellite-based studies of the dust storms from the Aral Sea suggested that playa salts promote precipitation and act as giant CCN [Rudich et al., 2002]; however, cloud residue analysis was not completed to confirm the chemistry and associated CCN activity of the observed dust. Thus, these ICE-L results represent the first direct detection of playa salts serving as cloud nuclei within clouds. Due to the global abundance of playas [Jones and Deocampo, 2004; Prospero et al., 2002] and increasing potential for playa dust storms due to land use changes [Gill, 1996], it is particularly important to understand the impact of playa dust on clouds. Based on the results shown herein, we suggest that playa salts impact cloud droplet and ice formation and cloud properties, particularly in remote continental locations. In-depth studies of the salt-producing potentials of playas globally are needed to further evaluate the transport, mixing, and cloud interactions of these salts in the global atmosphere. The implications of this study have ramifications for land use decisions and climate predictions, particularly of the indirect effect and regional weather patterns.

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