Hygroscopicity of Fresh and Aged Salt Mixtures from Saline Lakes

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Abstract: The high hygroscopicity of salt aerosol particles makes the particles active in aerosol and cloud formations. Inland saline lakes are an important and dynamic source of salt aerosol. The salt particles can be mixed with mineral dust and transported over long distances. During transportation, these particles participate in atmospheric heterogeneous chemistry and further impact the climate and air quality on a global scale. Despite their importance and potential, relatively little research has been done on saline lake salt mixtures from atmospheric perspectives. In this study, we use experimental and model methods to evaluate the hygroscopic properties of saline lake brines, fresh salt aerosol particles, and aged salt aerosol particles. Both original samples and literature data are investigated. The original brine samples are collected from six salt lakes in Shanxi and Qinghai provinces in China. The ionic compositions of the brines are determined and the hygroscopicity measurements are performed on crystallized brines. The experimental results agree well with theoretical deliquescence relative humidity (DRH) values estimated by a thermodynamic model. The correlations between DRHs of different salt components and the correlations between DRHs and ionic concentrations are presented and discussed. Positive matrix factorization (PMF) analysis is performed on the ionic concentrations data and the hygroscopicity results, and the solutions are interpreted and discussed. The fresh and aged salt aerosol particles are analyzed in the same way as the brines, and the comparison shows that the aged salt aerosol particles completely alter their hygroscopic property, i.e., transferring from MgCl$_2$ governed to NH$_4$NO$_3$ governed.

Keywords: hygroscopicity; aerosol; saline lake; playa; thermodynamic model; PMF; ageing

1. Introduction

Salt aerosol plays an important role in many processes related to atmospheric chemistry and climate systems [1], especially as active parts in aerosol and cloud formations [2,3] due to their high hygroscopicity [4]. The formation of salt particles has several pathways, including sea spray [5], wind erosion [6], anthropogenic processes [7], etc. An important and dynamic source is saline lakes and playas [8,9], which produce inland salt aerosol particles externally and internally mixed with mineral dust that can be transported over
long distances [6,10–17]. These long-range transported aerosols can alter the radiative balance [18], influence precipitation [19], participate in atmospheric heterogeneous chemistry, and form secondary aerosol to deteriorate air quality [16,20].

The salt aerosol particles from saline lakes and playas impact the climate on a global scale [11,14–16,21]. Pratt and coworkers studied the cloud droplet residues over the Great Basin, where Na-K-Mg-Ca-Cl playa salt fingerprint components were identified, showing that these salt components act as active cloud condensation nuclei (CCN) [15]. Gaston et al. used thermodynamic models to describe the hygroscopicity of playa dust based on their chemical compositions [14]. Tang and coworkers studied the hygroscopicity of saline mineral dust from playa regions, and the experimental results are well described by thermodynamic models [22]. Kong and coworkers revealed that most of the playa surface salts are dominated by NaCl, regardless of the chemical compositions of the nearby saline lakes [23]. In the same work, the hygroscopicity of various sampled salts is well described by thermodynamic models, based on the ionic compositions [23].

The inland salt aerosol particles are also found to associate with haze formation in more populated regions. For example, the formation of secondary inorganic aerosol (SIA) in Shanghai is promoted by the salt aerosol after long-distance transportation from East Asian desert regions [16]. The formed haze is more harmful to human health compared to the primarily emitted dust and salt aerosol. Moreover, during transportation the salt-dust mixture can undergo heterogeneous reactions and be transformed to materials of higher hygroscopicity [24,25]. Mitroo et al. reported that the Cl-rich dust from playas can react with N₂O₅ and produce atmospheric reactive chlorine that facilitates tropospheric ozone formation [21]. Thus, facilitated by the long-range transportation and heterogeneous pathways, the inland salt aerosol particles impact the air quality and human health [26].

The origins of inland salt aerosol particles, i.e., saline lakes and playas, are under continuous and dynamic developments, especially in the context of climate change [27] and increased land and water use [28,29]. This makes saline lakes and playas more important due to their growing spatial extent. Despite the active and important roles of inland salt aerosol particles play in the climate system, the modern climate models have not yet taken them into account [30]. The main reasons are the relatively low abundance compared to other components (e.g., sea salt), as well as the poorly understood characteristics and mechanisms (e.g., salt-dust mixing ratio and state). In this study, we use experimental and model methods to evaluate the hygroscopic properties of saline lake brines, fresh and aged salt aerosol particles. Ionic compositions reported in the literature are extensively collected and analyzed by the model. The comparison between fresh and aged salt particles shows the change of hygroscopicity-governing components before and after ageing.

2. Methods

2.1. Sampling Sites

The original brine samples are collected from six salt lakes, located in Shanxi and Qinghai provinces, as shown in Figure 1.

2.1.1. Two Lakes (XC and YC) in Shanxi Province

The Yuncheng Salt Lake is in topographic depressions formed by local faults and subsidence which is situated in the southwest of Yuncheng city, Shanxi Province [31,32]. As a unique low-altitude inland salt lake in China [33], Yuncheng Salt Lake is one of the largest sodium sulfate type inland salt lakes in the world, covering an area of 132 km², and the catchment area of the lake exceeds 120 km². The lake is located in a monsoonal climatic region with an average temperature of 12.8 °C, its average annual precipitation is approximately 550 mm, and evapotranspiration is about 1240 mm. The lake basin has a high salinity center surrounded by low salinity and freshwater areas, and abundant salt mine is distributed at the bottom of the lake. Mirabilite, halite, and astrakhanite are the main mineral phases of the salt lake, with sulfate, chloride, and sodium as the dominant ions [33].
Two brine samples were collected from the Yuncheng Salt Lake region, Xiao Chi (XC) and Yan Chi (YC), on 2nd September 2020. The XC is the original saline lake and the YC is an evaporation pond next to the XC.

2.1.2. Four Lakes (CK, KK, QH and MA) in Qaidam Basin

The Qaidam Basin is located on the northern side of the Qinghai-Tibet Plateau and is surrounded by Qilian Mountains, Kunlun Mountains and Altun Mountains [34]. The Qaidam Basin which has a continental climate, mainly controlled by the westerly circulation, as well as influenced by India, East Asian winter, and East Asian summer monsoon is windy all year round [35]. The climate is dry and there is little vegetation in this area, mostly playas, Gobi and deserts. The total annual average precipitation is 42 mm, the average annual temperature is 5.1 °C, the average annual evaporation capacity which far exceeds the rainfall is 2597.6 mm [35]. Despite the low temperature latitude, the Qaidam Basin is far away from the ocean and the climate is extremely dry which has led to the formation of many salt lakes and playas in the basin. Basin lakes migrate from west to east, and the salt formation stage in the west is earlier than that in the east.

Chaka (CK) Salt Lake, Keke (KK) Salt Lake, Qarhan (QH) Salt Lake and Mang’ai (MA) Salt Pond are the sampling lakes in the Qaidam Basin, where the sampling was done during 11–14 September 2020. Detailed information about sampling locations can be found in a previous publication [23]. The brines from the six different salt lakes were taken 2–5 cm below the lake surface with a glass beaker and then all the samples were filtered by qualitative filter papers (pore size <20 μm) within 2 days after sampling. The samples were stored in polyethylene bottles sealed with Parafilm membrane and refrigerated at 4 °C.

2.2. Ion Chromatography

Prior to the ion chromatography measurements, the brine samples were diluted by a factor of 1000 to fulfill the detection range. The diluted brines were then filtered by 0.22 μm water filter membranes. The Dionex AQUION IC system (Thermo Fisher Scientific, MA, USA) was used to measure cations (Na+, K+, Mg2+ and Ca2+) and anions (Cl− and SO42−) simultaneously. CS12A with methanesulfonic acid eluent (30 mmol L−1) and AS11-HC with KOH eluent (20 mmol L−1) were used to measure cations and anions, respectively. All the samples were measured in duplicate with good reproducibility.

2.3. Brine Crystallization

Filtered brines were heated to evaporate water to yield crystalized brines. During heating, the brines were continuously stirred with a glass rod to prevent salt splashing especially at the late stage of boiling. When the brines completely dried up, the crystallized brines were bottled and stored in a refrigerator at 4 °C.
2.4. Hygroscopicity Measurements

The hygroscopic properties of the crystalized brines were measured by a vapor sorption analyzer (Q5000SA; TA Instruments). During experiments (at a room temperature of 25 ± 0.1 °C), the RH was elevated stepwise. The precise masses of the samples along the RH ramp were measured by a highly sensitive balance. The balance has a dynamic range of 0–100 mg and a measurement sensitivity is <0.1 μg. The detailed instrumental and operational information can be found in previous studies [36–38]. The masses of used samples were about 0.5–1 mg. The mass change of the empty sample holder during the RH ramp was measured as baseline, which was removed during data processing. The apparatus had been calibrated by NaCl, KCl and (NH₄)₂SO₄ and the differences from theoretical deliquescence RH are within 1% [36]. Prior to each experiment, the sample was first dried at <1% RH, and then the RH was elevated to 90% RH with steps of 5–10% RH. At each RH step, 30 min was given for the samples to equilibrate with the environment, i.e., mass change ratio <0.05%. Three repeated samples were prepared and measured, and the average values are reported.

2.5. Thermodynamic Model

The AIOMFAC (aerosol inorganic-organic mixtures functional groups activity coefficients) model was used to simulate the deliquescence relative humidity (DRH) of the crystalized brine samples mixtures based on their activity coefficients [39,40]. The cations and anions are balanced by scaling. From the result output, the molar ion activity products (IAP) are calculated based on the calculated activities of ions. Taking MgCl₂ as an example, the IAP is calculated as

\[ IAP = \left[ a_m(\text{Mg}^{2+}) \right] \times \left[ a_m(\text{Cl}^-) \right]^2 \]  

(1)

where \( a_m \) is the activity of relevant ions. The IAP is a function of RH, and when the IAP reaches the IAP of saturated solutions (IAP_{sat}), the corresponding salt deliquesces is the DRH. The IAP_{sat} is calculated based on the ionic concentrations of saturated solution from literature. Figure 2 shows two example plots with the IAP versus RH. The horizontal dashed lines indicate the IAP_{sat} of each corresponding salt, and the vertical dashed lines point to the DRH on the x-axis. The complete set of IAP figures for all samples is included in Figure S1.

![Figure 2. IAP as a function of RH, including four salt components in two brine samples.](image)

2.6. Positive Matrix Factorization Analysis

The positive matrix factorization (PMF) receptor model was used to analyze the results, both the ion chromatography and the DRH results determined from the thermodynamic model. An open-source program EPA PMF 5.0 was employed for the analysis [41].

3. Results and Discussions

The ionic concentrations of brine samples (including six brines measured in this study and 90 values from literature) are presented. With the thermodynamic model, AIOMFAC,
DRHs are calculated based on the ionic compositions. Correlation analysis and PMF analysis are applied on the ionic concentrations and the DRH values. In a similar way, the ionic compositions of fresh and aged salt aerosol particles, from the literature, are analyzed. The chemical and hygroscopic evolutions from brine, fresh aerosol to aged aerosol are discussed.

3.1. Saline Lake Brines
3.1.1. Ionic Concentrations

The complete collection of ionic concentrations from literature can be found in Table S1 [6,16,42–48], and representative values are plotted in Figure 3. Overall, the anion Cl− is most abundant in brines, followed by the cations Na+ and Mg2+ (Figure 3a). The concentrations of the other three ions (K+, Ca2+, SO42−) are relatively low, and their relative abundances are more visible in the Na-normalized log-scale Figure 3b. The concentrations of the two major cations Na+ and Mg2+ show an anti-phase oscillation, while the Mg2+ concentrations follow the Cl− concentrations (Figure 3b).

![Figure 3](imageurl)

**Figure 3.** (a) Ionic concentrations of brine samples. (b) Na-normalized ionic concentrations of brine samples. The letters after the numbers indicate the literature source, and the detailed literature information can be found in Table S1. c from [43,44], d from [45], e from [46], g from [47,48].

The correlations between the ions are quantified and the results are shown in Table 1. Between the major ions, Na+ and Mg2+ are competing with a negative correlation coefficient of −0.55. Mg2+ and Cl− are positively correlated with a coefficient of 0.86. As for the minor ions, K+ shows weak associations with Mg2+ (0.24) and Cl− (0.29), and a negative correlation with Na+ (−0.06) and Ca2+ (−0.21). Another noticeable negative correlation is between Ca2+ and SO42− (−0.33), as the two ions form CaSO4 that easily precipitates, leading to the low coexistence of the two ions in the aqueous phase.

**Table 1.** Correlations coefficient of ionic concentrations of all brine samples.

|                | Na+  | K+  | Mg2+ | Ca2+ | Cl−  | SO42− |
|----------------|------|-----|------|------|------|-------|
| Na+            | 1.00 |     |      |      |      |       |
| K+             | −0.06| 1.00|      |      |      |       |
| Mg2+           | −0.55| 0.24| 1.00 |      |      |       |
| Ca2+           | −0.07| −0.21| 0.16 | 1.00 |      |       |
| Cl−            | −0.13| 0.29| 0.86 | 0.35 | 1.00 |       |
| SO42−          | 0.25 | −0.02| 0.11 | −0.33| 0.05 | 1.00  |
3.1.2. PMF Analysis of Ionic Concentrations

PMF analysis is performed to further investigate the relationship of the ions. The PMF analysis returns a 3-factor solution, and the factor profile is presented in Figure 4a. The three factors are led by (1) Factor 1: Ca$^{2+}$, (2) Factor 2: K$^+$, Mg$^{2+}$, Cl$^-$, and (3) Factor 3: SO$_4^{2-}$, respectively. The major ion Na$^+$ is evenly split into Factor 1 and 3, implying the anti-correlation between Na$^+$ and the cations in Factor 2 (Mg$^{2+}$ and K$^+$).

![Figure 4](image)

Figure 4. (a) Factor profiles of PMF 3-factor results on ionic concentrations. (b-1,b-2) PMF factor distributions of all brine samples. (c-1,c-3) Three factor-representing samples: (c-1) 41c for Factor 1, (c-2) QH for Factor 2, (c-3) KK for Factor 3. b from [42], c from [43,44], d from [45], e from [46], g from [47,48].

The factor contributions of the PMF results are shown in Figure 4b, including three representative samples whose ionic concentrations are presented in Figure 4c. The representative sample of Factor 1 (41c) shows a high level of Cl$^-$ and Na$^+$. Although the Ca$^{2+}$ concentration (leader ion of Factor 1) is relatively low compared to the major ions (Cl$^-$, Na$^+$ and Mg$^{2+}$) as shown in Figure 4c-1, the absolute Ca$^{2+}$ concentration reaches 0.265 mol/L, which is significantly higher than the Ca$^{2+}$ levels of most other samples. The QH case, representing Factor 2, shows outstandingly high level of SO$_4^{2-}$, matching the factor profile of Factor 3, though it is still much lower than the main anion Cl$^-$. The three-factor PMF result shows that the saline lake brines can be classified into three categories, featured with significant Ca$^{2+}$, Mg$^{2+}$, and SO$_4^{2-}$, respectively.
3.1.3. Deliquescence RH

The complex chemical compositions of brines lead to complicated hygroscopic properties. Nevertheless, compared to other natural substances, saline lake brines are relatively simple due to their relatively low bioactivities and organic contents. Here, a thermodynamic model, AIOMFAC, is applied to describe the hygroscopic properties of salt mixtures with brine chemical textures. In total, there are six original brine samples from this study and 27 sets of representative literature data (22 sets of brine samples and 5 sets of aerosol samples) are analyzed. All the calculated DRHs can be found in Table S2.

Figure 5 shows calculated DRHs of four common deliquescing salts (NaCl, MgCl₂, KCl, MgSO₄). Among the four components, MgCl₂ in general has the lowest DRHs, i.e., the MgCl₂ component most easily deliquesces. NaCl is a major component in most brines, thus its hygroscopic property strongly influences the behavior of the mixtures. MgSO₄ has the highest DRH, but it is not an effective component as MgCl₂ (easiest to deliquesce) and NaCl (very abundant). As for KCl, its hygroscopicity is influential in K-rich brines. Moreover, the abundance of K affects the hygroscopicity of other components. For example, in the QH sample, the high K content leads to a depressed DRH_{NaCl}.

The red and blue squares in Figure 5 indicate the RH onsets for initial and significant water uptake that are experimentally observed, respectively. The experimental results of the six original samples are presented in Figure 6. Figure 6a–c show the growth factors as a function of RH, and Figure 6d shows the RH of water uptake onsets. In Figure 6a, the dotted lines indicate the theoretically predicted DRH_{NaCl} in the six samples. The dashed lines in Figure 6b are the calculated DRHs_{MgCl₂}. To visualize the initial mass change due to water update, Figure 6c zooms in on the y-axis. Identified initial water uptake onsets are indicated as red squares in Figure 6d (same as the red squares in Figure 5). Since mixtures have complicated components that result in complex water update behaviors, most of the samples show continuous mass increases along with the increasing RH. Nevertheless, the XC and QH samples present identifiable major water uptake onsets, which are indicated as blue squares in Figure 6d, the same as the blue squares in Figure 5.
As seen in Figure 6d, in most samples the initial water uptakes are comparable to the MgCl₂ deliquescence onsets. However, the QH sample shows a large discrepancy between the experimental and theoretical results, where the initial water uptake is observed at 20%, which is considerably lower than the lowest calculated DRH_{MgCl₂} at 37%. Note that the QH sample has an exceptionally high K level, which significantly lowers the DRH_{NaCl} to 42% (DRH of pure NaCl ≈ 75%). The particularly complicated composition of the QH sample makes it very sensitive to ambient water vapor. Further, the heterogeneity of samples may be another potential reason for the derivation. Moreover, the major water uptake of the XC sample is also well described, which is largely determined by the NaCl component.

The DRHs of mixtures are complicated due to ionic interactions. To understand the relationships between ions and DRHs, Figure 7 plots the ionic concentrations against the predicted DRHs. Figure 7a shows that the DRH_{NaCl} are positively correlated with Na⁺ but negatively correlated with Mg^{2+} and Cl⁻, i.e., the existence of Mg^{2+} lowers the DRH_{NaCl}. The Cl⁻ follows the Mg^{2+} trend as its counter-ion. In contrast, Figure 7b shows the opposite trends for MgCl₂, which is negatively correlated with Na⁺ and positively associated with Mg^{2+} and Cl⁻. In another word, both DRH_{NaCl} and DRH_{MgCl₂} are lowered by the impurity cations (Mg^{2+} for NaCl; Na⁺ for MgCl₂). As for KCl, the K⁺ concentrations are relatively low, and the DRH_{KCl} is associated with Mg^{2+} and Cl⁻ (Figure 7c), showing good coherence with DRH_{MgCl₂}. A complete correlation table between the DRHs and ion concentrations can be found in Table S3.

Figure 7d–f shows the correlations between the DRHs of three major salt components (NaCl, MgCl₂, and MgSO₄). The full correlation parameters can be found in Table S4. In Figure 7d, the DRH_{NaCl} is widely distributed (between 40% to 75%), compared to MgCl₂ that fluctuates between 20% to 40%. In addition, the DRH of these two components are negatively correlated, i.e., high DRH_{NaCl} are accompanied by low DRH_{MgCl₂}. The KCl shows a generally positive correlation with MgCl₂ (Figure 7f) and is negatively correlated with NaCl (Figure 7e).
Correlations of DRHs and ionic concentration (mol/L) of four ions. (d–f) Correlations of the DRH value of three main salt components in brine samples.

The DRH results are also analyzed by the PMF, where a 2-factor solution is converged, and the results are shown in Figure 8. Factor 1 is dominated by NaCl and MgSO$_4$ and Factor 2 is characterized by MgCl$_2$ and KCl. In Figure 8b, most samples show the co-existence of both factors, and only five samples are dominated by Factor 1. Comparing with the PMF analysis on ionic concentrations (previously presented in Figure 4), the current Factor 1 cases are the typical Factor 3 cases in Figure 4, i.e., of high SO$_4^{2-}$ and Na$^+$ concentrations.

3.1.4. Aged Salt Aerosol Particles

Salt aerosol particles undergo ageing along with the atmospheric transportation, which alters the chemical and hygroscopic properties of the salt particles. For example, the ageing of NaCl (fresh salt particles) can occur by reacting with nitrogen oxides and sulfur oxides in atmosphere to form Cl-depleted aged salt particles [49]. To shed light on the chemical
evolution during ageing, we analyze two previously reported ionic concentrations of fresh and aged salt aerosol particles. Figure 9a shows the Na-normalized ionic concentrations of the two types of salt aerosol particles, i.e., long-range transported and aged aerosol (1a, 2a, 3a) [16], and fresh salt aerosol (74f, 84f, 87f) [6]. Compared to Na\(^+\), the aged salt aerosol particles contain high levels of NH\(_4\)\(^+\), NO\(_3\)\(^-\), and SO\(_4\)\(^2-\), which are typical secondary inorganic aerosol components. On the other hand, the fresh salt aerosol has high Na\(^+\), Cl\(^-\), and Ca\(^2+\) fractions, which are common aerosol components found near playas and deserts.

**Figure 9.** (a) Normalized concentration of ions. (b) Modeled DRHs of size salts in five aerosol samples. The 1a–3a are aged salt aerosol particles [16]; 74f, 84f, 87f are fresh salt aerosol particles [6].

Based on the ionic concentrations, the DRHs of the aged and fresh salt samples are calculated and shown in Figure 9b. The hygroscopicity of aged aerosol is dominated by the NH\(_4\)NO\(_3\) component, while the fresh salt particles are regulated by MgCl\(_2\), like the brine samples. Ageing strongly alters the hygroscopic property of salt particles. Note that not all the salt components deliquesce in the aged cases, i.e., IAP never reaches IAP\(_{sat}\) below 100\% RH, e.g., MgCl\(_2\), NaCl, and KCl in the aged samples.

The PMF method is applied to the ionic concentrations and calculated DRHs of aerosol samples. Both datasets are resolved with 2-factor solutions, i.e., fresh and aged factors. Figure 10a-1 shows the factor profiles of ionic concentrations, where Mg\(^{2+}\), Ca\(^{2+}\), SO\(_4\)\(^{2-}\), and NO\(_3\)\(^-\) are the dominating ions of Factor 1 (aged); Na\(^+\) and Cl\(^-\) are the fingerprint ions for Factor 2 (fresh). Figure 10a-2 shows the factor contributions, which confirms that Factor 1 dominates the aged aerosol, especially 1a and 2a, and Factor 2 are important in the fresh aerosol (74f, 84f, 87f). Factor 10b shows the PMF results of calculated DRHs. Factor 1 is governed by secondary inorganic aerosol components and therefore dominates the aged samples. Factor 2 reflects the deliquescing components for fresh salt aerosol. Figure 10c shows the ionic compositions of 1a and 74f, which represent the aged and fresh aerosol factors, respectively.
Atmosphere 2021, 12, 1203

4. Conclusions

In this study, we use experimental and model methods to evaluate the hygroscopic properties of saline lake brines, namely fresh and aged salt aerosol particles. The ionic compositions of the brines are measured by ion chromatography in this study. The comparison of ions measured in this study and taken from the literature shows that Na\(^+\) and Mg\(^{2+}\) are negatively correlated, and Mg\(^{2+}\) and Cl\(^-\) are positively correlated. K\(^+\) shows significant associations with Mg\(^{2+}\) and Cl\(^-\), and a slight negative correlation with Na\(^+\). Hygroscopicity measurements were performed on the crystallized brines, and the experimental results are compared with theoretical DRH values from a thermodynamic model. The DRHs of four common deliquescing salts (NaCl, MgCl\(_2\), KCl, MgSO\(_4\)) are presented. Among the four components, MgCl\(_2\) in general has the lowest DRHS, which determines the initial water uptake. NaCl is a major component in most brines, thus its hygroscopic property strongly influences the behavior of the mixtures. In addition, the presence of abundant K affects the hygroscopicity of other components, e.g., the high K content leads to a depressed DRH\(_{\text{NaCl}}\) in the QH sample. The correlations between the DRHS and the correlation between DRHs and ionic concentrations are presented and discussed. Positive matrix factorization analysis is performed on the ionic concentrations data and the hygroscopicity results, and the solutions are interpreted and discussed. The fresh and aged salt aerosol particles are analyzed in the same way as the brines, and the comparison shows that the aged salt aerosol particles completely alter their hygroscopic property, i.e., transferring from MgCl\(_2\) governed to NH\(_4\)NO\(_3\) governed.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/atmos12091203/s1, Figure S1a: AIOMFAC modelled IAPs as a function of RH (brine samples from Shan’xi and Qinghai). The horizontal dashed lines are DRH for individual salts in the corresponding salt mixture. The vertical dashed lines are the DRH found by the AIOMFAC model; Figure S1b: AIOMFAC modelled IAP as a function of RH (aerosol samples). The horizontal dashed
lines are DRH for individual salts in the corresponding salt mixture. The vertical dashed lines are the DRH found by the AIOMFAC model; Figure S1c: AIOMFAC modelled IAP as a function of RH (brine samples). The horizontal dashed lines are DRH for individual salts in the corresponding salt mixture. The vertical dashed lines are the DRH found by the AIOMFAC model; Figure S1d: AIOMFAC modelled IAP as a function of RH (brine samples). The horizontal dashed lines are DRH for individual salts in the corresponding salt mixture. The vertical dashed lines are the DRH found by the AIOMFAC model; Table S1: Molar concentration (mol/l) of all samples; Table S2: DRH predicted by the AIOMFAC model; Table S3: Correlations of the DRH and ions in brine samples; Table S4: Correlations of the DRH value of four kinds of salts in brine samples.

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