Hygroscopic behavior and chemical composition evolution of internally mixed aerosols composed of oxalic acid and ammonium sulfate

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Abstract. Although water uptake of aerosol particles plays an important role in the atmospheric environment, the effects of interactions between components on chemical composition and hygroscopicity of particles are still not well constrained. The hygroscopic properties and phase transformation of oxalic acid (OA) and mixed particles composed of ammonium sulfate (AS) and OA with different organic to inorganic molar ratios (OIRs) have been investigated by using confocal Raman spectroscopy. It is found that OA droplets first crystallize to form OA dihydrate at 71 % relative humidity (RH), and further lose crystalline water to convert into anhydrous OA around 5 % RH during the dehydration process. The deliquescence and efflorescence point for AS is determined to be 80.1 ± 1.5 % RH and 44.3 ± 2.5 % RH, respectively. The observed efflorescence relative humidity (ERH) for mixed OA / AS droplets with OIRs of 1 : 3, 1 : 1 and 3 : 1 is 34.4 ± 2.0, 44.3 ± 2.5 and 64.4 ± 3.0 % RH, respectively, indicating the elevated OA content appears to favor the crystallization of mixed systems at higher RH. However, the deliquescence relative humidity (DRH) of AS in mixed OA / AS particles with OIRs of 1 : 3 and 1 : 1 is observed to occur at 81.1 ± 1.5 and 77 ± 1.0 % RH, respectively. The Raman spectra of mixed OA / AS droplets indicate the formation of ammonium hydrogen oxalate (NH₄HC₂O₄) and ammonium hydrogen sulfate (NH₄HSO₄) from interactions between OA and AS in aerosols during the dehydration process on the time scale of hours, which considerably influence the subsequent deliquescence behavior of internally mixed particles with different OIRs. The mixed OA / AS particles with an OIR of 3 : 1 exhibit no deliquescence transition over the RH range studied due to the considerable transformation of (NH₄)₂SO₄ into NH₄HC₂O₄ with a high DRH. Although the hygroscopic growth of mixed OA / AS droplets is comparable to that of AS or OA at high RH during the dehydration process, Raman growth factors of mixed particles after deliquescence are substantially lower than those of mixed OA / AS droplets during the efflorescence process and further decrease with elevated OA content. The discrepancies for Raman growth factors of mixed OA / AS particles between the dehydration and hydration process at high RH can be attributed to the significant formation of NH₄HC₂O₄ and residual OA, which remain solid at high RH and thus result in less water uptake of mixed particles. These findings improve the understanding of the role of reactions between dicarboxylic acid and inorganic salt in the chemical and physical properties of aerosol particles, and might have important implications for atmospheric chemistry.

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1 Introduction

Atmospheric aerosols have vital impacts on the Earth’s climate directly by scattering, reflecting and absorbing solar radiation, and indirectly by influencing formation of clouds and precipitation (Tang and Munkelwitz, 1994b; Jacobson et al., 2000; Penner et al., 2001; Pöschl, 2005; Von Schneidemesser et al., 2015). Direct and indirect effects depend on the chemical and physical properties of atmospheric aerosols, including size, structure, hygroscopicity and chemical composition. Field observations indicate that aerosol particles are generally internal mixtures of inorganic and organic compounds in the atmosphere (Saxena et al., 1995; Murphy et al., 1998, 2006; Pratt and Prather, 2010). Ammonium sulfate (AS) is one of the most abundant inorganic constituents in the atmosphere, the hygroscopicity of which has been widely investigated (Liu et al., 2008; Cziczo et al., 1997; Laskina et al., 2015).

Oxalic acid (OA) is ubiquitous and has been identified as the dominant dicarboxylic acid in urban and remote atmospheric aerosols (Chebbi and Carlier, 1996; Kanakidou et al., 2004; Yang and Yu, 2008; Wang et al., 2012; Kawamura and Bikkina, 2016). Previous studies have focused on deliquescence behavior of pure OA (Peng et al., 2001; Braban et al., 2003; Miñambres et al., 2013; Ma et al., 2013a; Jing et al., 2016). It was found that due to its high deliquescence point, OA exhibited no deliquescence transition or hygroscopic growth within the relative humidity (RH) range studied by an electrodynamic balance (EDB) (Peng et al., 2001), vapor sorption analyzer (Ma et al., 2013) or hygroscopicity tandem differential mobility analyzer (HTDMA) (Jing et al., 2016). Braban et al. (2003) reported that OA could deliquesce at 98 % RH using aerosol flow tube Fourier transform infrared spectroscopy (AFT-FTIR). However, the study on the efflorescence behavior of OA during the dehydration process remains limited (Peng et al., 2001; Mikhailov et al., 2009). Peng et al. (2001) observed the efflorescence transition of OA using EDB while Mikhailov et al. (2009) reported continuous hygroscopic growth of OA during both hydration and dehydration processes using the HTDMA.

The dicarboxylic acids can affect properties of internally mixed aerosol particles such as hygroscopicity, phase transition, solubility and chemical reactivity (Lightstone et al., 2000; Brooks et al., 2002; Jørgensen et al., 2007; Pradeep Kumar et al., 2003; Treuel et al., 2011; Laskin et al., 2012; Drozd et al., 2014; Peng et al., 2016; Jing et al., 2016, 2017; Li et al., 2017). Field measurements have observed the formation of low-volatility organic salts in atmospheric particles due to the reactions of organic acids with mineral salts, chloride salts, nitrate salts, ammonium and amines (Sullivan and Prather, 2007; Laskin et al., 2012; Wang and Laskin, 2014; Smith et al., 2010). The organic salts formed typically have varying hygroscopicity compared to the corresponding organic acids. Thus, these drastic changes in aerosol composition have potential effects on the water uptake and related physicochemical properties of particles. The effects of OA on deliquescence behaviors of AS have been extensively investigated (Brooks et al., 2002; Prenni et al., 2003; Wise et al., 2003; Miñambres et al., 2013; Jing et al., 2016). The majority of studies found that the presence of OA had no obvious effect on the deliquescence process of OA / AS mixtures with minor OA content (Brooks et al., 2002; Prenni et al., 2003; Wise et al., 2003). To our knowledge, there is still a lack of studies on the efflorescence process of OA / AS mixed systems. In fact, the efflorescence behavior is a critical hygroscopic characteristic of atmospheric aerosols, which may favor specific chemical interactions between components within the supersaturated droplets. For example, previous studies have found that chloride depletion could occur in the NaCl / dicarboxylic acid mixed aerosols during the dehydration or efflorescence process, which led to the formation of organic salts and in turn affected subsequent deliquescence behaviors of aerosols (Laskin et al., 2012; Ghorai et al., 2014). OA has been found to react with both mono- and divalent cations to form low-volatility and low solubility compounds (Drozd et al., 2014). Miñambres et al. (2013) proposed that OA might react with AS to form ammonium hydrogen oxalate and ammonium hydrogen sulfate within OA / AS solution. Due to the lack of available thermodynamic data, the aerosol thermodynamic models typically assume that upon dehydration dicarboxylic acid could only form an organic solid without the organic salt in the inorganic electrolyte / dicarboxylic acid system (Clegg and Seinfeld, 2006; Amundson et al., 2007). Thus, the incorporation of organic salts formed from interactions between inorganic salts and organic acids is crucial to the modeling of hygroscopic properties of mixed organic / inorganic particles. It merits further investigation on the interactions between OA and AS and the related influence on the water uptake behaviors of aerosols during the dehydration and hydration processes.

Raman spectroscopy is a powerful technique to characterize aerosol compositions, water contents, molecular interactions and particle phases especially for the efflorescence process (Ma and He, 2012; Laskina et al., 2013; Zhou et al., 2014; Wang et al., 2015). In this study, the phase transformations and hygroscopic properties of OA and mixed OA / AS droplets with varying OA content were studied by confocal Raman spectroscopy in conjunction with optical microscopy. Furthermore, we explored the effects of reactions between OA and AS on the chemical compositions and hygroscopic properties of mixed OA / AS droplets.

2 Experimental section

2.1 Sample preparation

AS and OA dihydrate were purchased from Sinopharm Chemical Reagent Co. Ltd. (99.0 % purity) and used with-
out further purification. The 0.5 mol L$^{-1}$ pure component AS and OA solutions were prepared by dissolving AS and OA dihydrate in ultrapure water (18.2 MΩ cm, Barnstead Easy-pure II), respectively. The mixed OA / AS solutions with different organic to inorganic molar ratios (OIRs) of 1 : 3, 1 : 1 and 3 : 1 were obtained by dissolving a designated amount of OA into AS solutions. The sample solution was discharged from a syringe. Then, residual solution in the syringe was pushed rapidly to generate aerosol droplets spraying onto a polytetrafluoroethylene (PTFE) substrate fixed to the bottom of the sample cell. Then, the sample cell was promptly sealed by a transparent polyethylene film. The RH in the sample cell was regulated by nitrogen streams consisting of a mixture of water-saturated N$_2$ and dry N$_2$ at controlled flow rates. At $\sim$ 95 % RH, the droplets with a diameter of 30–40 µm detected by an optical microscope (50× objective, 0.75 numerical aperture) were selected to acquire the Raman spectra. The dry size of these particles after efflorescence ranged from 10 to 20 µm. The RH and temperature of the outflow from the sample cell was measured by a humidity/temperature meter (Centertek Center 313) with an accuracy of $\pm$ 2.5 % below 90 % RH and $\pm$ 0.7 K placed near the exit of the sample cell. The temperature accuracy of 0.7 K could result in an uncertainty of 4 % at a RH of 95 %. The temperature of the sample was maintained at 297 ± 0.5 K by using an automatic thermostat.

2.2 Apparatus and conditions for the measurements

The experimental setup used in this study was described in detail in a previous work (Wang et al., 2008; Dong et al., 2009; Zhou et al., 2014). Briefly, the Renishaw InVia confocal Raman spectrometer equipped with a Leica DMLM microscope was used to acquire the Raman spectra. An argon-ion laser (wavelength 514.5 nm, model Stellar-REN, ModuLaser) was used as an excitation source with an output power of 20 mW, and a 514.5 nm notch filter was adopted to remove the strong Rayleigh scattering. A 1800 g mm$^{-1}$ (grooves per millimeter) grating was used to obtain the spectra in the range of 200–4000 cm$^{-1}$ with a resolution of about 1 cm$^{-1}$. Spectral calibration was made using the 520 ± 0.05 cm$^{-1}$ Stokes shift of the silicon band before performing measurements. Then, spectroscopic measurements were made on droplets observed by using the Leica DMLM microscope with a 50× objective lens (0.75 numerical aperture). The spectra were obtained with three spectral scans, and each time with an accumulation time of 10 s. The sample droplets were injected onto the substrate at high RH ($\sim$ 95 % RH). Subsequently, the RH was decreased stepwise for a slow dehydration process, and then increased stepwise from RH < 3 % to high RH for a hydration process. The decrease rate was typically 5–6 % RH every 40 min and the rate remained 2–3 RH every 40 min near the phase transition. The RH was decreased continuously in a few minutes for a rapid dehydration process. The particles were equilibrated with water vapor at a given RH for about 40 min, during which the intensity ratio of the water peak (3430 cm$^{-1}$) to the sulfate peak (980 cm$^{-1}$) remained constant. The spectra of AS, OA and mixed OA / AS droplets were monitored and measured through a full humidity cycle. Multiple particles (three or four) were selected to acquire the Raman spectra through each humidity cycle. Each humidity cycle experiment was repeated at least three times. All the measurements were taken at ambient temperature of about 297 K.

Raman growth factor, $g$(RH), is defined as the ratio of the integrated area of the OH stretching mode of water (3350–3700 cm$^{-1}$) at each RH ($A_{RH}$) normalized to that of a dry particle ($A_{RH0}$) according to Eq. (1) (Laskina et al., 2015).

$$g(RH) = \frac{A_{RH}}{A_{RH0}},$$

where $A_{RH}$ is the integrated area of the OH stretching mode from water (3350–3700 cm$^{-1}$) at a specific RH and $A_{RH0}$ is that of a dry particle. Hygroscopic growth curves are acquired by plotting the average Raman growth factor of duplicate particles as a function of RH.
and hydrate remains unchanged for 40 min. Once RH decreases to 6.6 % RH, at which the Raman spectrum of dihydrate indicates that OA droplets crystallize to form OA dihydrate. The observation of no spectral change until 94 % RH suggests that OA dihydrate shows no deliquescence transition in the 0–94 % RH range studied, consistent with previous studies (Peng et al., 2001; Braban et al., 2003; Ma et al., 2013a; Jing et al., 2016). The transition point of anhydrous OA to OA dihydrate upon hydration is 17.9–19.6 % (Fig. 2b), in agreement with the results reported by Braban et al. (2003) and Ma et al. (2013a).

### 3 Results and discussion

#### 3.1 Raman spectra of pure AS and OA droplets

The Raman spectra of AS droplets during the dehydration and hydration processes can be found in Fig. 1a and b, respectively. AS droplets effloresce at 44.3 ± 2.5 % RH, as indicated by the disappearance of the water peak centered at 3437 cm⁻¹ and a red-shift in the ν₃(SO₄²⁻) peak position from 979 to 974 cm⁻¹ during the dehydration process. For the hydration process, the deliquescence of AS particles is observed to occur at 80.1 ± 1.5 % RH, resulting in an abrupt increase in the absorbance of the water peak centered at 3437 cm⁻¹ and a blue-shift in the ν₄(SO₄²⁻) peak position from 974 to 979 cm⁻¹.

The Raman spectra of OA droplets with varying RH during the dehydration and hydration processes are shown in Fig. 2, and the assignments of the peaks for OA are presented in Table 1 according to previous studies (Hibben, 1935; Ebisuzaki and Angel, 1981; Mohaček-Grošev et al., 2009). As seen in Fig. 2a, the feature bands for OA droplets are observed at 1460, 1750 and 3433 cm⁻¹ at 92.5 % RH. At lower RH around 71 % (Fig. 2a; magenta line), these bands shift to 1490, 1737, 3433 and 3474 cm⁻¹, and a new band at 1689 cm⁻¹ occurs, which is entirely consistent with the spectrum of OA dihydrate (Fig. 2a; black dashed line). It indicates that OA droplets crystallize to form OA dihydrate. OA particles after efflorescence exist in the form of dihydrate until 6.6 % RH, at which the Raman spectrum of dihydrate remains unchanged for 40 min. Once RH decreases to ~ 5.0 %, the peaks promptly shift to 1477, 1710, 2587, 2760 and 2909 cm⁻¹, and peaks at 3433 and 3474 cm⁻¹ assigned to ν(OH) vanish, which is the spectral feature of anhydrous OA. This result implies that OA dihydrate is converted to anhydrous OA in the RH range around 5.0 %. The Raman spectra of anhydrous OA particles during the hydration process as a function of RH are shown in Fig. 2b. It can be found that the Raman spectra feature for anhydrous OA particles occurs at RH < 19.6 %. At 19.6 % RH, the peaks observed at 1490, 1737, 3433 and 3474 cm⁻¹ are identical to that of OA dihydrate (Fig. 2a; black dashed line), indicating the formation of OA dihydrate. The observation of no spectral change until 94 % RH suggests that OA dihydrate shows no deliquescence transition in the 0–94 % RH range studied, consistent with previous studies (Peng et al., 2001; Braban et al., 2003; Ma et al., 2013a; Jing et al., 2016). The transition point of anhydrous OA to OA dihydrate upon hydration is 17.9–19.6 % (Fig. 2b), in agreement with the results reported by Braban et al. (2003) and Ma et al. (2013a).

#### 3.2 Raman spectra of OA / AS mixtures

The Raman spectra of mixed OA / AS droplets with OIRs of 1 : 3, 1 : 1 and 3 : 1 at various RHs during the dehydration and hydration processes are depicted in Figs. 3 and 4, respectively. Since spectral features upon hydration are identical to the dehydration process, here we only analyzed spectral evolution of the efflorescence process in detail. The detailed assignments are summarized in Table 2. For the mixed OA / AS droplets (OIR = 1 : 3) at 96.2 % RH (seen in Fig. 3a), the bands at 979 and 1049 cm⁻¹ are characteristic peaks of aqueous SO₄²⁻ and HSO₄⁻ (ν₃(SO₄⁻)), respectively. In addition, the peak at 1741 and 1446 cm⁻¹ can be assigned to the vibrating mode of aqueous OA and H₂C₂O₄, respectively. With decreasing RH, only small changes are observed in the spectra until the RH reaches 34.4 %. At 34.4 % RH, the shift of the ν₃(SO₄²⁻) peak from 979 to 974 cm⁻¹ indi-

### Table 1. Molecular vibration assignments of pure OA and AS droplets.

<table>
<thead>
<tr>
<th>Solid H₂C₂O₄</th>
<th>H₂C₂O₄ droplets (92.5 % RH)</th>
<th>(NH₄)₂SO₄ droplets (94.8 % RH)</th>
<th>References</th>
<th>Assignments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anhydrous</td>
<td>Dihydrate</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>482</td>
<td>477</td>
<td>457</td>
<td>Spinner (2003)</td>
<td>δ₄(SO₄²⁻)</td>
</tr>
<tr>
<td>828</td>
<td>855</td>
<td>845</td>
<td>Ebisuzaki and Angel (1981)</td>
<td>r(CO)</td>
</tr>
<tr>
<td>1477</td>
<td>1490</td>
<td>1460</td>
<td>Ebisuzaki and Angel (1981)</td>
<td>ν(C-C)</td>
</tr>
<tr>
<td>1627</td>
<td>1636</td>
<td></td>
<td>Ebisuzaki and Angel (1981)</td>
<td>ν(C=O)</td>
</tr>
<tr>
<td>1689</td>
<td>1737</td>
<td>1750</td>
<td>Hibben (1935)</td>
<td>ν(C=O)</td>
</tr>
<tr>
<td>2587, 2760</td>
<td>2909</td>
<td></td>
<td>Mohaček-Grošev et al. (2009)</td>
<td>Combinations</td>
</tr>
<tr>
<td>3433, 3474</td>
<td>3433</td>
<td></td>
<td>Spinner (2003)</td>
<td>ν(OH)</td>
</tr>
</tbody>
</table>

ν: stretching, δ: bending, r: rocking and s: symmetric mode.
Table 2. Molecular vibration assignments of mixed OA / AS systems.

<table>
<thead>
<tr>
<th>H$_2$C$_2$O$_4$-(NH$_4$)$_2$SO$_4$ (1 : 3), RH = 96.2 %</th>
<th>H$_2$C$_2$O$_4$-(NH$_4$)$_2$SO$_4$ (1 : 1), RH = 96.1 %</th>
<th>H$_2$C$_2$O$_4$-(NH$_4$)$_2$SO$_4$ (3 : 1), RH = 95.9 %</th>
<th>References</th>
<th>Assignments</th>
</tr>
</thead>
<tbody>
<tr>
<td>450 450</td>
<td>852 852</td>
<td>461 Spinner (2003)</td>
<td>$\delta_1$($SO_4^{2-}$)</td>
<td></td>
</tr>
<tr>
<td>979 979</td>
<td>1050 Dawson et al. (1986)</td>
<td>$\nu_5$($SO_4^{2-}$)</td>
<td>$\nu(C-C)$</td>
<td></td>
</tr>
<tr>
<td>1049 1051</td>
<td>1382 Chang and Huang (1997)</td>
<td>$\nu_2$($SO_4^{2-}$)</td>
<td>$\nu(C-O)$</td>
<td></td>
</tr>
<tr>
<td>1446 1448</td>
<td>1460 Ebisuzaki and Angel (1981)</td>
<td>$\nu_6$($COO$)</td>
<td>$\nu(C-O)$</td>
<td></td>
</tr>
<tr>
<td>1694</td>
<td>1752 Ebisuzaki and Angel (1981)</td>
<td>$\nu_7$($C=O$)</td>
<td>$\nu(OH)$</td>
<td></td>
</tr>
<tr>
<td>1741</td>
<td>1751</td>
<td>$\nu_8$($C=O$)</td>
<td>$\nu(OH)$</td>
<td></td>
</tr>
<tr>
<td>3430 3427</td>
<td>3426 Spinner (2003)</td>
<td>$\nu_9$($C=O$)</td>
<td>$\nu(OH)$</td>
<td></td>
</tr>
</tbody>
</table>

$v$: stretching, $\delta$: bending, $\omega$: wagging and $s$: symmetric mode.

cates the crystallization of AS, as also seen in Fig. 10b. A new band centered at 874 cm$^{-1}$ corresponds to combination bands of the vibrational mode ($\delta(S-OH)$) of the HSO$_4^-$ ion from NH$_4$HSO$_4$ (Dawson et al., 1986) and the HC$_2$O$_4^-$ ion vibrating (Villepin and Novak, 1971), suggesting the formation of crystalline NH$_4$HC$_2$O$_4$. Moreover, the several new peaks at 1416, 1469 and 1660 cm$^{-1}$ can be attributed to the HC$_2$O$_4^-$ ion vibration of crystalline NH$_4$HC$_2$O$_4$ (Villepin and Novak, 1971). Therefore, the evolution of Raman spectra of the mixed OA / AS droplets (OIR = 1 : 3) during the dehydration process confirms that OA could react with AS to form NH$_4$HSO$_4$ and NH$_4$HC$_2$O$_4$, which supports previous speculation for the reaction between OA and AS (Miñambres et al., 2013). The reaction of OA with AS occurs via the following pathway:

$$(\text{NH}_4)_2\text{SO}_4(aq) + \text{H}_2\text{C}_2\text{O}_4(aq) \rightarrow \text{NH}_4\text{HSO}_4(aq) + \text{NH}_4\text{HC}_2\text{O}_4(aq).$$

For the mixed OA / AS droplets (OIR = 1 : 1; Fig. 3b), the evolution of spectra shows a resemblance to that of mixed droplets (OIR = 1 : 3). At 96.1 % RH, the peaks at 979, 1751, 1051 and 1448 cm$^{-1}$ can be assigned to the vibrating mode of SO$_4^{2-}$, OA, HSO$_4^-$ ($\nu_6$($SO_4^-$)) and HC$_2$O$_4^-$, respectively. At 75.0 % RH, a new peak at 874 cm$^{-1}$ corresponding to the vibrational mode ($\delta(S-OH)$) of HSO$_4^-$ and the HC$_2$O$_4^-$ ion vibrating as well as the new peaks at 494, 1469 and 1677 cm$^{-1}$ due to the HC$_2$O$_4^-$ vibrating mode, indicates that crystalline NH$_4$HC$_2$O$_4$ is generated from the reaction of OA with AS. As the RH further decreases to 44.3 %, the $\nu_5$($SO_4^{2-}$) band shifts from 979 to 974 cm$^{-1}$, indicating the formation of crystallized AS particles.

For the mixed OA / AS droplets (OIR = 3 : 1; Fig. 3c) at 95.9 % RH, the bands at 970, 1752 and 1050 cm$^{-1}$ are characteristic of the SO$_4^{2-}$ ion, OA and the HSO$_4^-$ ion ($\nu_6$($SO_4^-$)), respectively. And the peak at 1460 cm$^{-1}$ can be attributed to the vibrating mode of the HC$_2$O$_4^-$ ion. When the RH decreases to 74.4 %, a new band at 874 cm$^{-1}$ is contributed by the vibrational mode of both HSO$_4^-$ ($\delta(S-OH)$) and HC$_2$O$_4^-$. 

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Figure 2. Raman spectra of oxalic acid (OA) droplets during the (a) dehydration process and (b) hydration process. In panel (a), the black dashed line indicates the spectrum of pure H$_2$C$_2$O$_4$·2H$_2$O particles with the peak height of $\nu$(OH) located at 3433 cm$^{-1}$ scaled by a factor of 1/6.
Meanwhile, the bands at 494, 1471 and 1654 cm\(^{-1}\) can be attributed to the HC\(_2\)O\(_4^-\) vibrating mode, suggesting OA reacts with AS to yield crystalline NH\(_4\)HC\(_2\)O\(_4\) during the dehydration process. At 64.4 % RH, the peaks at 494, 874, 1471, 1654, 1718 cm\(^{-1}\), and the peak at 3426 cm\(^{-1}\) from OA dihydrate become sharp and narrow, indicating that the OA / AS droplets (OIR = 3 : 1) completely crystallize to form NH\(_4\)HC\(_2\)O\(_4\) and H\(_2\)C\(_2\)O\(_4\)·2H\(_2\)O. No obvious change in spectral features of the major bands is observed with RH decreasing from 64.4 to 1.1 %.

3.3 Hygroscopicity of pure AS, OA and OA / AS mixtures

3.3.1 Phase transitions and chemical transformation of AS in mixed systems

Considering that the peak position is sensitive to the chemical environment in the aerosols, the position of the \(\nu_3(SO_4^{2-})\) mode can be used to determine the phase transitions of AS. The previous studies have also applied the abrupt shift in characteristic peak position to indicate a phase transition of AS during the hygroscopic process (Braban and Abbatt,

Figure 3. Raman spectra of mixed oxalic acid / ammonium sulfate (OA / AS) droplets with OIRs of (a) 1 : 3, (b) 1 : 1 and (c) 3 : 1 at various RH values during the dehydration process.

Figure 4. Raman spectra of mixed oxalic acid / ammonium sulfate (OA / AS) droplets with OIRs of (a) 1 : 3, (b) 1 : 1 and (c) 3 : 1 at various RH values during the hydration process.
Figure 5. The peak position of the $\nu_5$-SO$_{4}^{2-}$ peak of mixed OA / AS particles and pure AS particles at various RHs during the (a) dehydration and (b) hydration process. The red and blue dashed lines indicate the ERH and DRH, respectively.

2004; Ling and Chan, 2008; Yeung et al., 2009). Figure 5 presents the peak position of the $\nu_5$(SO$_{4}^{2-}$) for AS droplets and mixed OA / AS droplets during the dehydration and hydration processes, respectively. During the dehydration process, a red shift from 979 to 974 cm$^{-1}$ can be observed for AS and OA / AS mixed particles with OIRs of 1 : 3 and 1 : 1, indicating crystallization of AS from droplets. During the hydration process, the observations of blue shift from 974 to 979 cm$^{-1}$ for AS and OA / AS mixed particles with OIRs of 1 : 3 and 1 : 1 suggest the deliquescence transition of AS from crystal phase to aqueous solution. For OA / AS mixed particles with an OIR of 3 : 1, the peak shift between $\sim$ 966 and $\sim$ 979 cm$^{-1}$ is determined during the whole RH cycle. The shift of the $\nu_5$(SO$_{4}^{2-}$) mode to 966 cm$^{-1}$ suggests the formation of letovicite (NH$_4$)$_3$H(SO$_4$)$_2$(s) (Damak et al., 1985). The DRH and ERH for pure and mixed systems are shown in Fig. 5 and a detailed discussion is given in the following section.

The peaks at $\sim$ 1049 and $\sim$ 979 cm$^{-1}$ for mixed OA / AS droplets (OIRs = 1 : 3, 1 : 1 and 3 : 1) can be attributed to the HSO$_4^-$ and SO$_4^{2-}$ stretching mode, respectively. The area ratio of Raman peaks assigned to the HSO$_4^-$ and SO$_4^{2-}$ is used to indicate the degree of conversion of SO$_4^{2-}$ into HSO$_4^-$ ($\alpha_{\text{HSO}_4^-}$) in mixtures, which can be expressed as the following:

$$\alpha_{\text{HSO}_4^-} = \frac{A_{1049}}{A_{1049} + A_{979}},$$

where $A_{1049}$ and $A_{979}$ are the peak areas of the HSO$_4^-$ and SO$_4^{2-}$ signals, respectively. The peak at $\sim$ 1049 cm$^{-1}$ for HSO$_4^-$ is not obvious after crystallization of the droplets. Thus, the calculations are based on the bands at RH approaching the full efflorescence point. The estimated $\alpha_{\text{HSO}_4^-}$ value for OIR = 1 : 3 (36.1 % RH), OIR = 1 : 1 (46.2 % RH) and OIR = 3 : 1 (66.2 % RH) is 0.048, 0.368 and 0.644, respectively, indicating the enhanced conversion of SO$_4^{2-}$ into HSO$_4^-$ with increasing OA content in the mixed systems. Due to the effects of the Raman cross section, $\alpha_{\text{HSO}_4^-}$ could not represent the actual degree of conversion. In fact, here $\alpha_{\text{HSO}_4^-}$ is only used for comparisons of degree of conversion of SO$_4^{2-}$ into HSO$_4^-$ between mixed particles with varying OIRs.

Figure 6. Hygroscopicity of (a) AS and (b) OA as a function of RH. The red and blue dashed lines indicate the ERH and DRH, respectively. The black lines show the phase transition points for the transformation between OA dihydrate and anhydrous OA.
3.3.2 Hygroscopic growth of pure and mixed components

Hygroscopicity curves of AS and OA particles are shown in Fig. 6. The optical images of the AS particle at the phase change points can be seen in Fig. 7. The ERH of AS is determined to be 44.3 ± 2.5 % RH, which generally falls into the range from 33 to 52 % RH reported in the literature (Tang and Munkelwitz, 1994a; Cziczo et al., 1997; Dougle et al., 1998; Laskina et al., 2015). The DRH of AS particles is observed to occur at 80.1 ± 1.5 % RH, which agrees well with reported values of 80 % RH by EDB (Tang and Munkelwitz, 1994a) and 82.3 ± 2.5 % RH by micro-Raman spectroscopy (Laskina et al., 2015). As shown in Figs. 6b and 8, the measured ERH of OA is 71 ± 2.5 % RH, which deviates from the reported value of 51.8–56.7 % RH by Peng et al. (2001) using the EDB technology. It is worthwhile to point out that the conversion of OA droplets to OA dihydrate at 71 % RH is inconsistent with the observation of Peng et al. (2001). They observed that OA droplets crystallized to form anhydrous OA rather than OA dihydrate at 51.8–56.7 % RH. The discrepancy on the ERH of OA compared to that reported by Peng et al. (2001) is likely due to the effects of substrate and sample purity. The size of dry particles ranging from 10 to 20 µm in our experiment is consistent with observation using EDB by Peng et al. (2001), which eliminates the influence of particle size. The substrate supporting droplets may promote the heterogeneous nucleation of OA while the levitated droplets in the EDB study can avoid induced nucleation by the substrate. Ghorai et al. (2014) also reported the potential effects of substrate on the efflorescence transition of NaCl/dicarboxylic acid mixed particles. In addition, the OA purity in our study is 99.0 % lower than that of 99.5 % in the study by Peng et al. (2001). Thus, trace amounts of impurities in OA droplets acting as a heterogeneous nucleus could contribute to crystallization and result in a higher ERH of OA. Due to the effects of substrate and sample purity, the heterogeneous nucleation should be responsible for the discrepancy on the observed ERH of OA. The water content of the supersaturated droplet at the onset of crystallization determines the form of OA crystal generated, i.e., anhydrous OA or OA dihydrate. Due to a higher ERH, OA droplets with more water content favor the formation of a dihydrate after crystallization. It should be noted that our experiment appears to be favored in the atmospheric environment, considering that insoluble material such as mineral dust mixed with OA may play the role of substrate thus facilitating the heterogeneous nucleation of OA aerosols. The Raman growth factor of OA shows no obvious change between ∼71 and 6.6 % RH upon dehydration. At RHs lower than 5 %, the Raman growth factors drop abruptly due to the transformation of crystalline H$_2$C$_2$O$_4$·2H$_2$O into anhydrous OA, as also indicated by the Raman spectrum. It seems that the structure of the anhydrous OA particle is not as compact as that of the dihydrate, seen in Fig. 8. Thus, the loss of crystal water results in no obvious change in particle size. During the hydration process, the Raman growth factor of OA shows a slight increase at 19.6 % RH, which can be attributed to the conversion of anhydrous OA to dihydrate. The transition point of anhydrous OA to OA dihydrate agrees with previous studies (Braban et al., 2003; Ma et al., 2013b; Miñambres et al., 2013). No deliquesce behavior is observed for OA dihydrate even at 94 % RH, consistent with earlier observations (Ma et al., 2013b; Miñambres et al., 2013; Jing et al., 2016).
Figure 9. Hygroscopicity of OA/AS mixtures with OIRs of (a) 1:3, (b) 1:1 and (c) 3:1 as a function of RH. The red and blue dashed lines indicate the ERH and DRH, respectively. In panels (a) and (b), Raman growth factors of pure AS and OA above 80 % RH in the dehydration process are also included for comparisons. In the panel (c), Raman growth factors of pure AS and OA dihydrate above 80 % RH during the dehydration process are also given for comparisons.

Figure 10. Optical micrographs of the mixed oxalic acid/ammonium sulfate (OA/AS) particle (OIR = 1:3) at phase change points. Dehydration: (a) 36.1 % RH and (b) 34.4 % RH. Hydration: (c) 79.4 % RH and (d) 81.1 % RH. In the panel (d), the visual solid in aqueous phase is marked with a red dashed circle.

Figure 9 presents hygroscopic growth of OA/AS mixtures with OIRs of 1:3, 1:1 and 3:1 as a function of RH. The red and blue dashed lines indicate the ERH and DRH, respectively. In panels (a) and (b), Raman growth factors of pure AS and OA above 80 % RH in the dehydration process are also included for comparisons. In the panel (c), Raman growth factors of pure AS and OA dihydrate above 80 % RH during the dehydration process are also given for comparisons.

The mixed OA/AS droplets with an OIR = 1:1 first partially effloresce at 75.0 ± 1.6 % due to the crystallization of NH₄HC₂O₄, as indicated by Raman spectra. Then, the full efflorescence occurs at 44.3 ± 2.5 % RH with the crystallization of AS. The full ERH of 1:1 OA/AS mixed droplets is highly consistent with that of pure AS. During the hydration process, the Raman growth factor of 1:1 mixed particles increases slightly at 35.5 % RH, and then remains al-
In contrast to the surface, the obvious features of 980 and 1050 cm$^{-1}$ of a mixed OA dihydrate and other components in the outer phase reveals that the major component on the surface at 1471 cm$^{-1}$ is assigned to SO$_4^{2-}$, HSO$_4^-$ and HC$_2$O$_4^-$, respectively. The sharp absorption at 874 cm$^{-1}$ is assigned to SO$_4^{2-}$, HSO$_4^-$ and HC$_2$O$_4^-$, respectively. The water contents of mixed droplets after deliquescence are significantly lower than those upon dehydration. The Raman features at 494 and 874 cm$^{-1}$ have confirmed the presence of solid NH$_4$HC$_2$O$_4$ upon hydration across all RHs studied (seen in Fig. 4), which should be responsible for the decreasing water uptake of the mixed particles at high RH.

For mixed OA / AS droplets with an OIR = 3 : 1, the partial and full efflorescence transition could be observed at 74.4 ± 1.0 % RH and 64.4 ± 3.0 % RH, respectively (seen in Figs. 9 and 11). As seen in Fig. 3c, the bands at 494, 1471 and 1654 cm$^{-1}$ suggest the formation of crystalline NH$_4$HC$_2$O$_4$ at 74.4 ± 1.0 % RH. Figure 12 presents the spatial distribution of chemicals within mixed OA / AS (OIR = 3 : 1) particles at 74.4 % RH. The characteristic peak of 980, 1050 and 1471 cm$^{-1}$ is assigned to SO$_4^{2-}$, HSO$_4^-$ and HC$_2$O$_4^-$, respectively. The sharp absorption at 874 cm$^{-1}$ and obvious peak at 1471 cm$^{-1}$ indicate the abundant content of NH$_4$HC$_2$O$_4$. The comparison of characteristic peaks between inner and outer phase reveals that the major component on the surface of a mixed OA / AS (OIR = 3 : 1) particle is NH$_4$HC$_2$O$_4$. In contrast to the surface, the obvious features of 980 and 1050 cm$^{-1}$ at the core of the particle suggest that (NH$_4$)$_2$SO$_4$ and NH$_4$HSO$_4$ mainly exist in the inner aqueous phase. During the dehydration process, crystalline NH$_4$HC$_2$O$_4$ in the outer phase acts as the heterogeneous nucleus, leading to the crystallization of OA dihydrate and other components in the inner phase. Thus, the full ERH of 3 : 1 OA / AS mixed droplets is higher than that of pure AS (44.3 ± 2.5 % RH) and NH$_4$HSO$_4$ (22–0.05 % RH). During the hydration process, Raman growth factors of mixed particles slightly increase at 34.5 % RH. No deliquescence transition or significant water uptake is observed over the RH range studied. This phenomenon can be explained by the fact that most of the AS in the mixtures has been converted into NH$_4$HC$_2$O$_4$ and NH$_4$HSO$_4$ or letovicite. Although NH$_4$HSO$_4$ with a low DRH may contribute to water uptake of mixed particles, the minor NH$_4$HSO$_4$ or letovicite formed in the mixtures is likely to be coated by NH$_4$HC$_2$O$_4$ and OA with a high DRH. Thus, the mixed OA / AS particles with an OIR = 3 : 1 show no obvious hygroscopic growth upon hydration due to the change in aerosol composition and morphological effects. The effects of morphology on the hygroscopic growth of aerosols have been reported for AS particles containing adipic acid (Sjogren et al., 2007). The water uptake of AS particles containing relatively high content of adipic acid could be suppressed due to AS enclosed by the crust of solid adipic acid with a high DRH.

The observed ERH for mixed droplets was dependent on the molar ratio of OA to AS. The mixed OA / AS droplets with an OIR of 1 : 3 are observed to effloresce completely at 34.4 ± 2.0 % RH relative to ERH of pure AS (44.3 ± 2.5 %) or OA (71 ± 2.5 %). It can be seen that AS as a major fraction of the particle does not promote the heterogeneous nucleation of OA. Meanwhile, the crystallization of AS is also influenced due to the presence of OA. A similar phenomenon was also observed for malonic acid / AS mixtures with minor organic content (Braban and Abbatt, 2004; Parsons et al., 2004). Braban and Abbatt (2004) found that the ERH of malonic acid / AS mixed particles was considerably decreased compared to that of pure AS for mass fractions of malonic acid less than 0.3. They concluded that the presence of AS in the supersaturated droplet could exert the extra barrier to nucleation of malonic acid crystals rather than play the role of a heterogeneous nucleation site. As for 1 : 3 OA / AS mixed droplets, OA may also inhibit the nucleation of OA at relatively high RH. With decreasing RH, aqueous OA could enhance the viscosity of the droplet due to hydrogen bond interactions (Mikhailov et al., 2009), thus limiting the nucleation of AS and resulting in a lower ERH with respect to the value of pure AS (Parsons et al., 2004). In the case of mixed OA / AS droplets with an OIR of 1 : 1 and 3 : 1, the NH$_4$HC$_2$O$_4$ formed at ~75 % RH upon dehydration likely acts as a heterogeneous nucleus for crystallization of other components, which increases the full efflorescence point of mixed particles. One study indicated that humic acid sodium salt (NaHA, Aldrich) could also promote the ERH of AS (Badger et al., 2006). Similar to OA, succinic acid and adipic acid have a high deliquescence point and low solubility. However, it has been found that the efflorescence point of AS in mixed particles is not elevated even when the content of succinic acid or adipic acid is more than 50 % by mass or mole fraction (Ling and Chan, 2008; Yeung et al., 2009;
Laskina et al., 2015). The chemical nature of a solid determines its ability to act as a heterogeneous nucleus (Braban and Abbatt, 2004). In contrast to AS particles containing succinic acid or adipic acid, our results suggest that the addition of OA into AS droplets may trigger partial and full crystallization of aerosols at relatively higher RH upon dehydration due to the NH$_4$HC$_2$O$_4$ product acting as an effective nucleus.

During the deliquescence process, the OA / AS mixed particles with an OIR of 1:3 and 1:1 exhibit a slightly lower deliquescence point than that of pure AS, consistent with previous observations of effects of crystalline OA on deliquescence transition of AS (Brooks et al., 2002; Wise et al., 2003; Jing et al., 2016). It should be noted that prior literature results also showed that continuous or smooth water uptake from low RH was observed for particles composed of AS and OA with a mass ratio of 1.5:1 due to the fact that after drying OA existed in an amorphous or liquid-like state that prevented nucleation of AS even under dry conditions (Prenni et al., 2003). In the present study, water uptake by the OA / AS mixed particles at high RH upon hydration is dramatically lower than that upon dehydration and significantly decreased with elevated OA content. This phenomenon distinguishes itself from hygroscopic characteristics of typical water-soluble mixtures in the literature. It has been found that hydration growth curves and dehydration growth curves are typically merged above the deliquescence point for mixed systems containing inorganic salts and water-soluble organic compounds (Choi and Chan, 2002; Chan and Chan, 2003; Gysel et al., 2004; Clegg and Seinfeld, 2006; Sjogren et al., 2007; Pope et al., 2010; Ghorai et al., 2014; Estilllore et al., 2016). In this study, Raman spectra and the micrographs suggest the presence of solid NH$_4$HC$_2$O$_4$ and residual solid OA at high RH should be responsible for the decreased water uptake during the hydration process. In contrast, Prenni et al. (2003) reported that the hygroscopic growth of OA / AS mixed particles remained unchanged at 90% RH with OA mass fraction ranging from 0.01 to 0.4. In addition, they also found that water uptake after deliquescence was well described by the model method assuming complete dissolution of OA in aqueous phase as well as no interactions between OA and AS, which was also observed by Jing et al. (2016) using the HTDMA. The previous HTDMA studies for OA / AS mixed particles indicate no composition change and no specific interactions existing between OA and AS (Prenni et al., 2003; Jing et al., 2016). However, it should be noted that the HTDMA studies did not perform measurements for the dehydration process such that aerosols underwent rapid drying on the time scale of seconds, i.e., the total residence time for transformation of droplets into dry particles in the drying section of the HTDMA is typically tens of seconds (Prenni et al., 2003; Jing et al., 2016), much shorter than (10–12 h) in our study. In the HTDMA experiments, the combination of faster drying and smaller particles with submicron size implies that the aqueous phase obtained higher supersaturations than in our present study (Rosenoern et al., 2008), leading to less dissociation of OA and thus less HC$_2$O$_4^−$ formed in the droplets as well as the inhibited formation of NH$_4$HC$_2$O$_4$. The fast evaporation of water from the surface of an aqueous droplet upon rapid drying could result in a higher surface concentration of solutes than the slow drying process (Treuel et al., 2011). The higher surface concentration of OA corresponds to less formation and hence decreased supersaturation of HC$_2$O$_4^−$. Due to the dependence of the nucleation rate on the extent of supersaturation, it can be expected that the nucleation of NH$_4$HC$_2$O$_4$ is suppressed within OA / AS mixed droplets undergoing rapid drying.

Considering the potential effects of drying time on the reactions between OA and AS, we explored the hygroscopicity of OA / AS particles with an OIR of 1:1 after the rapid drying process. The mixed OA / AS droplets undergo dehydration to form dry particles in 3–5 min. We observed one-step efflorescence of rapidly dried particles (1:1, molar ratio) occurring at 47 ± 2.5% RH, compared to the two-step efflorescence of slowly dried particles occurring at 75 and 44.3% RH. The Raman spectra and hygroscopic curve upon hydration for OA / AS particles with an OIR of 1:1 are presented in Fig. 13. The obvious discrepancies can be observed for spectra at ~2% RH between...
of OA dihydrate, as indicated by the Raman feature. The deliquescent growth factors of mixed particles with an OIR of 1 : 1 and 3 : 1, OA droplets effloresce to form OA dihydrate at 71 ± 2.5 % RH, and then OA dihydrate further loses crystalline water to form anhydrous OA at ∼ 5.0 % RH during the dehydration process. The Raman spectra of mixed OA / AS droplets reveal the formation of NH₄HSO₄ and NH₄H₂SO₄ from the reaction of OA with AS in aerosols during the dehydration process. The deliquescent and efflorescence point for AS is observed to occur at 80.1 ± 1.5 and 44.3 ± 2.5 % RH, respectively. The ERH of the mixed OA / AS droplets with 1 : 3, 1 : 1 and 3 : 1 ratios is determined to be 34.4 ± 2.0, 44.3 ± 2.5 and 64.4 ± 3.0 % RH, respectively, indicating significant effects of OA content on the efflorescence transition of AS. The mixed OA / AS particles with 1 : 3 and 1 : 1 ratios show deliquescent transitions at 81.1 ± 1.5 and 77 ± 1.0 % RH, respectively, which is close to the DEH of AS. The mixed OA / AS particles with 3 : 1 ratio exhibit no deliquescent transition over the RH range studied due to the transformation of (NH₄)₂SO₄ into high-DRH NH₄H₂SO₄. The hygroscopic growth of mixed particles at high RH upon hydration is substantially lower than that of the corresponding dehydration process and further decreases with increasing OA content. The discrepancies for water content of mixed particles between the two processes at high RH can be explained by the significant formation of low hygroscopic NH₄H₂SO₄ and residual OA, which still remain solid and thus result in less water uptake of mixed particles.

The prior hygroscopic studies suggest that crystallization of internally mixed AS / dicarboxylic acid particles may lead to the formation of trace organic salt. Lightstone et al. (2000) estimated that approximately 2 % of the initial succinic acid may form ammoniated succinate within mixed ammonium nitrate / succinic acid particles during the efflorescence process. Ling and Chan (2008) inferred that crystallization of mixed OA / succinic acid droplets likely generated metastable organic salt based on a change in the Raman peak form of succinic acid. Braban and Abbatt (2004) reported that NH₄H₂SO₄ and ammoniated malonate were likely generated upon crystallization of mixed AS / malonic acid particles. However, due to the trace amount of organic salt below the

Figure 13. (a) Raman spectra of equal molar mixed OA / AS particles after the rapid drying process at various RH values upon hydration. The Raman spectrum (black short dash) at 2.5 % RH obtained from the slow drying process is also given for comparison. (b) Deliquescence curve of OA / AS mixtures with an OIR of 1 : 1. The hygroscopic curve (olive symbols) of particles after the slow drying process is also included for comparison. The blue dashed lines indicate the DRH.

4 Conclusions and atmospheric implications

In this work, confocal Raman spectroscopy is used to investigate the hygroscopic properties and phase transformations of OA and internally mixed OA / AS droplets (OIRs = 1 : 3, 1 : 1 and 3 : 1). OA droplets effloresce to form OA dihydrate at 71 ± 2.5 % RH, and then OA dihydrate further loses crystalline water to form anhydrous OA at ∼ 5.0 % RH during the dehydration process. The Raman spectra of mixed OA / AS droplets reveal the formation of NH₄H₂SO₄ and NH₄H₂SO₄ from the reaction of OA with AS in aerosols during the dehydration process. The deliquescent and efflorescence point for AS is observed to occur at 80.1 ± 1.5 and 44.3 ± 2.5 % RH, respectively. The ERH of the mixed OA / AS droplets with 1 : 3, 1 : 1 and 3 : 1 ratios is determined to be 34.4 ± 2.0, 44.3 ± 2.5 and 64.4 ± 3.0 % RH, respectively, indicating significant effects of OA content on the efflorescence transition of AS. The mixed OA / AS particles with 1 : 3 and 1 : 1 ratios show deliquescent transitions at 81.1 ± 1.5 and 77 ± 1.0 % RH, respectively, which is close to the DEH of AS. The mixed OA / AS particles with 3 : 1 ratio exhibit no deliquescent transition over the RH range studied due to the transformation of (NH₄)₂SO₄ into high-DRH NH₄H₂SO₄. The hygroscopic growth of mixed particles at high RH upon hydration is substantially lower than that of the corresponding dehydration process and further decreases with increasing OA content. The discrepancies for water content of mixed particles between the two processes at high RH can be explained by the significant formation of low hygroscopic NH₄H₂SO₄ and residual OA, which still remain solid and thus result in less water uptake of mixed particles.

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Raman or infrared detection limit, they found no apparent influence of organic salt formed upon dehydration on the water uptake or phase change of mixed particles. In contrast, our results indicate that the chemical processing upon drying of droplets containing OA and AS influences the efflorescence transition and water uptake of mixed aerosols during the humidity cycle by modifying particulate components.

Our results highlight the atmospheric importance of dicarboxylic acid–AS interactions in aerosol aqueous chemistry. Such chemical processing upon drying of aerosols comprised of organic acid / (NH₄)₂SO₄ mixtures may enhance the acidity of the aqueous phase in the intermediate RH due to the transformation of (NH₄)₂SO₄ into NH₄HSO₄. These experiments also imply that the chemical reaction between aqueous (NH₄)₂SO₄ and OA upon slow dehydration is a possible formation pathway for the low-volatility oxalate in ambient particles, which could enhance partitioning of dicarboxylic acids to aqueous particles with the presence of AS (Yli-Juuti et al., 2013; Hakkinen et al., 2014). It has been reported that the aerosol aqueous processing within organic acid / AS mixtures partly contributes to enhanced loadings of secondary organic aerosol from biogenic precursors (Hoyle et al., 2011). Compared to aqueous processing such as condensed-phase acid-catalyzed reactions relevant to formation of organosulfates, the contribution of other aerosol processes containing organic salt formation to secondary organic aerosol burden likely becomes important under less acidic conditions. Formation of low-solubility organic salts from aqueous processing within aerosols alters the particle-phase component and thus modifies aerosol’s hygroscopicity, optical properties and chemical reactivity. Our findings provide fundamental insights into the effects of drying conditions (drying rate or time) on formation of organic salts from reactions of organic acids with inorganic salts in the particle phase under ambient RH conditions. Overall, a better understanding of the chemical interactions between species in a multicomponent system during the humidity cycle is critical for the accurate modeling efforts of aerosol phase behavior in thermodynamic models.

Data availability. All data are available upon request from the corresponding authors.

Author contributions. YZ, MG and BJ designed the experiments and XW carried them out. XW and BJ performed the data analysis and prepared the manuscript with contributions from all co-authors.

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

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