Liquid-liquid phase separation and morphology of internally mixed dicarboxylic acids/ammonium sulfate/water particles

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Abstract. Knowledge of the physical state and morphology of internally mixed organic/inorganic aerosol particles is still largely uncertain. To obtain more detailed information on liquid-liquid phase separation (LLPS) and morphology of the particles, we investigated complex mixtures of atmospherically relevant dicarboxylic acids containing 5, 6, and 7 carbon atoms (C5, C6 and C7) having oxygen-to-carbon atomic ratios (O:C) of 0.80, 0.67, and 0.57, respectively, mixed with ammonium sulfate (AS). With micrometer-sized particles of C5/AS/H2O, C6/AS/H2O and C7/AS/H2O as model systems deposited on a hydrophobically coated substrate, laboratory experiments were conducted for various organic-to-inorganic dry mass ratios (OIR) using optical microscopy and Raman spectroscopy. When exposed to cycles of relative humidity (RH), each system showed significantly different phase transitions. While the C5/AS/H2O particles showed no LLPS with OIR = 2:1, 1:1 and 1:4 down to 20 % RH, the C6/AS/H2O and C7/AS/H2O particles exhibit LLPS upon drying at RH 50 to 85 % and ∼90 %, respectively, via spinodal decomposition, growth of a second phase from the particle surface or nucleation-and-growth mechanisms depending on the OIR. This suggests that LLPS commonly occurs within the range of O:C < 0.7 in tropospheric organic/inorganic aerosols. Both core-shell structures and partially engulfed structures were observed for the investigated particles, suggesting that such morphologies might also exist in tropospheric aerosols.

1 Introduction

Aerosol particles affect the Earth’s energy budget directly by scattering and/or absorbing solar radiation and indirectly by acting as cloud condensation and ice nuclei modifying cloud properties (Yu et al., 2006; IPCC, 2007). These effects are influenced by the aerosol size, physical state, chemical composition and morphology, which also affect the particle hygroscopicity (Hanel, 1976; Martin, 2000). Therefore, the interrelation between chemical composition and aerosol phase might be critical to attain a correct description of aerosol effects on climate.

The hygroscopic behavior of inorganic salts such as ammonium sulfate (AS), sodium chloride and ammonium nitrate have been well characterized (Tang and Munkelwitz, 1984, 1994; Cziczo et al., 1997; Martin, 2000). When crystalline particles of these inorganic salts are exposed to increasing relative humidity (RH), they take up water from the gas phase and become solution droplets at their deliquescence relative humidity (DRH). In turn, when the RH is decreased, the smaller solution droplets may reach high supersaturation before they crystallize and lose their water to the gas phase at the efflorescence relative humidity (ERH). While the inorganic aerosol constituents are relatively small in number and well known, the chemical composition of the organic matter is highly complex with a large fraction still

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Liquid-liquid phase separation and morphology of multicomponent systems. They use as a basis the group-contribution model called Aerosol Inorganic-Organic Mixtures Functional groups Activity Coefficients (AIOMFAC) (Zuend et al., 2008). Calculations that they performed for ternary and multicomponent alcohol/polyol-water-salt mixtures suggest that liquid-liquid equilibria are indeed a prevalent feature of organic-inorganic aerosol systems. LLPS in smog chamber experiments is supported by a recent study of phase transitions and growth factors of mixed particles consisting of AS and secondary organic material prepared from α-pinene ozonolysis that are consistent with a model of phase separation into an aqueous electrolyte phase and an organic-rich phase in individual particles (Smith et al., 2011).

Particles consisting of two liquid phases can adopt different morphologies, e.g. fully engulfed core-shell structures or asymmetric partial engulfed structures. Knowledge of the aerosol morphology is important for determining the nature of the gas-particle interface, and thus the rate of heterogeneous chemical reactions, and the rate of water uptake (Cai et al., 1996; Fuzzi et al., 2006; Kwamena et al., 2010, Reid et al., 2011). Formation of organic coatings by condensation of monoterpene oxidation products onto aqueous sulfate particles is suspected to significantly reduce the heterogeneous hydrolysis rates of N$_2$O$_5$ (Folkers et al., 2003; Amtiaux et al., 2006, 2007). Model simulations for Europe suggest that this process could decrease particulate nitrate concentrations by up to 90 % (Riemer et al., 2009). Strong reduction of N$_2$O$_5$ hydrolysis rates have also been observed for sulfuric acid and sea salt aerosols that were coated with organic surfactants (Thornton and Abbatt, 2005; McNeill et al., 2006; Cosman and Bertram, 2008; Cosman et al., 2008).

Ciobanu et al. (2009) used optical microscopy and micro-Raman spectroscopy to investigate LLPS of micrometer-sized droplets consisting of poly(ethylene glycol)-400 (PEG-400)/AS/water. The resulting morphology was an aqueous AS inner phase surrounded by a mainly PEG-400 containing outer phase. Core-shell morphologies have also been observed by Bertram et al. (2011) for mixed organic/AS particles deposited on a hydrophobically coated substrate. However, Kwamena et al. (2010) and Reid et al. (2011) have shown in model calculations and experiments that the adoption of a partially engulfed structure should predominate with the organic component forming a surface lens when the solubility of the organic component and the salt-out of the organic component to the surface by the presence of concentrated inorganic solutes in the aqueous phase are considered.

Previous studies on phase transitions of single aerosol particles have mostly been performed with one organic substance in the presence of AS (Parsons et al., 2006; Ling and Chan, 2008; Ciobanu et al., 2009; Yeung et al., 2009; Bertram et al., 2011). Studies on more complex organic mixtures are needed to obtain insight into the phase transitions of organic/inorganic aerosols considering atmospheric aerosol composition. In this study, we investigate single particles containing different dicarboxylic acids and AS.
During humidity cycles. Dicarboxylic acids are one of the most abundant organic compound classes in the troposphere (De Cesari et al., 2006; Gilardoni et al., 2009; Hawkins et al., 2010). For our model systems, we mix together three isomeric acids to better mimic the complexity of real aerosol compositions. Three mixtures with C5, C6, and C7 dicarboxylic acids with oxygen-to-carbon ratios (O:C) of 0.80, 0.67, and 0.57, respectively were investigated. AS has been chosen as the model inorganic salt due to its abundance in the atmosphere and its well-characterized thermodynamic and spectroscopic properties (Braban and Abbatt, 2004; Ling and Chan, 2008; Yeung et al., 2009). To explore the phase diagram, we mixed the components in different organic-to-inorganic dry mass ratios (OIR). We use optical microscopy and micro-Raman spectroscopy to elucidate the phase transitions of these model systems when exposed to RH cycles. A main goal is to investigate the RH conditions and mixture compositions for which the model systems exhibit LLPS. The study of single particles allows for the investigations of miscibility gaps in solutions that are supersaturated with respect to a crystalline species and, hence, not directly accessible to bulk measurements. It also allows studying particle morphologies. Laboratory experiments of single aerosol particles are thus essential to improve our understanding of aerosol processes.

2 Experimental and Modelling Methods

2.1 Materials

Aqueous solutions containing different OIR of the three model systems were prepared by dissolving the pure substances in purified water (resistivity \( \geq 18.0 \text{ M} \Omega \text{ cm} \)). The mixture “C5” consists of three C5-dicarboxylic acids (glutaric, methylsuccinic, and dimethylmalonic acid), “C6” of three C6-dicarboxylic acids (2-methylglutaric, 3-methylglutaric, and 2,2-dimethylsuccinic acid), and “C7” of three C7-dicarboxylic acids (3-methyladipic acid, 3,3-dimethylglutaric acid and diethylmalonic acid), mixed together in equal mass (and molar) ratios. All components were purchased from Sigma-Aldrich with purities \( \geq 98\% \) and were used without further purification.

2.2 Raman Microscopy of Single Droplets

Figure 1 shows schematically the experimental setup. Single droplets in a new custom-made cell are investigated with a Raman microscope (Jobin Yvon, model: Labram). Using a droplet generator (Knopf, 2003), micrometer-sized aqueous droplets were deposited on a glass cover slip (\( \approx 15 \text{ mm diameter} \)) that was hydrophobically coated by a silanization solution (Sigma-Aldrich) containing 5% dimethyldichlorosilane in heptane to avoid heterogeneous nucleation due to surface contact. The coating procedure is described in detail in Knopf (2003). The cover slip was mounted in the cell with inner dimension of 30 mm in diameter and 0.1–0.2 mm in height. The cell is kept at constant temperature (20°C) by water cooling/heating of the cell housing. RH is regulated by a \( \text{N}_2/\text{H}_2\text{O} \) flow that is brought to cell temperature before entering the cell. A single droplet on the cover slip was chosen and monitored with video microscopy during a humidity cycle. To change RH, the water vapor mixing ratio of a constant total \( \text{N}_2/\text{H}_2\text{O} \) flow (180 sccm) was varied by mixing dry and water-saturated \( \text{N}_2 \) flows using automatic mass flow controllers. Temperature and RH of the gas flow inside the cell were measured by a G-TUCN.34 sensor (U.P.S.I., France; capacity probe for RH, NTC for \( T \)) which is positioned about 5 mm downstream from the gas entrance. During the experiments, the gas flow temperature was maintained at 20 ± 0.2°C. The cell temperature was also measured directly on the substrate by a Pt100 temperature sensor (Minco, USA). The deviation between the two sensors was less than 0.3°C. Adjustment of the flows through the mass flow controllers, readout of the temperature and humidity and the synchronization with the video signal was controlled by a PC using custom software implemented in VEE (Agilent, USA). During humidity cycles, morphological changes of the droplet are monitored optically with the microscope equipped with a long working distance objective (Olympus BX-40, magnification 50, aperture 0.7). A black and white CCIR video camera with an acquisition frequency of 25 frames s\(^{-1}\) was used to acquire images and movies of
the droplet morphology. At the beginning of the experiment, the droplet was equilibrated at high RH (−95%). Humidity cycles comprise a RH reduction from 95 to 20% and subsequent increase from 20 to 95% at a rate of 0.1−0.3% RH min⁻¹.

To identify the composition of the droplet phases, Raman spectra were acquired at constant RH and temperature conditions with the Raman system coupled to the optical microscope. A Nd:YAG (532 nm wavelength) laser operated at 75−100 mW power was used as the excitation source. An 1800 grooves mm⁻¹ grating and a CCD camera were used to analyze the backscattered light from the sample. We obtained Raman spectra at different positions on the particles in the range of 150−4000 cm⁻¹ with a spectral resolution of about 2−4 cm⁻¹ and an integration time of 200 s.

2.3 RH and temperature sensor calibrations

The specified accuracy of the G-TUCN.34 RH sensor by the manufacturer is ±2% in the RH range from 2−98%. However, we observed that this type of sensor might show offsets compared to the calibration curve provided by the company. Therefore, by measuring the DRH of various pure inorganic salts at room temperature, such as KNO₃ (93.5%), (NH₄)₂SO₄ (80.0%), NaCl (75.0%), NH₄NO₃ (65.5%), NaI (38.0%) and MgCl₂·6H₂O (33.0%), which cover a wide range of RH values, we recalibrated the G-TUCN.34 RH sensor. To do this, the RH in the cell was continuously increased at a rate of about 0.05% RH min⁻¹ and the RH value at which onset of deliquescence was observed was registered as the DRH and compared with literature values. Each DRH value was determined using at least two independent particles in the diameter range of 10−35 μm. With this calibration, we obtained an accuracy of the humidity sensor of ±1.5% RH. In order to detect sudden offsets of RH, the calibration was checked by a DRH experiment with a pure AS crystal every two weeks when experiments have been performed. We calibrated the temperature sensors by determining the melting point of ice particles using the same procedure as described in Knopf and Lepori (2009). Water droplets of 15−60 μm in diameter froze at about −35°C in the cell. Subsequently, the ice melting point was determined using a heating ramp of 0.03°C min⁻¹. The temperature sensors proved to work accurately with deviations from the melting temperature of about ±0.1°C.

2.4 Saturation solubility and water activity measurements

To determine the physical properties of the dicarboxylic acids and AS, we measured their individual solubility in water and the water activity (a_w) of the saturated solutions. By stepwise adding small portions of water until the solute was completely dissolved, saturated aqueous solutions were prepared. The solutions were allowed to equilibrate in a thermostated water bath at 25.0±0.1°C in between the additions. The estimated uncertainty of this procedure is less than 1.0 wt%. The a_w of the saturated aqueous solutions were measured at 25°C, using a water activity meter (AquaLab Model 3TE, Decagon Devices, USA). In order to correct for instrument offset and drift, the performance of the sample block was frequently controlled and readjusted with reference samples (Marcoll and Krieger, 2006). The experimental errors for the measurements with standard samples such as KNO₃ (a_w = 0.925) and NaCl (a_w = 0.753) were ±0.002 a_w and ±0.009 a_w, respectively.

2.5 Thermodynamic modeling

We use the method developed by Zuend et al. (2010) for the computation of LLPS in mixed organic/inorganic systems. This method is based on the global minimization of the Gibbs energy of a system to find the number of phases and corresponding compositions at thermodynamic equilibrium. In order to calculate the Gibbs energy at a given composition, activity coefficients of all components are required to account for non-ideal mixing. Following Zuend et al. (2010), we use the group-contribution model AIOMFAC developed by Zuend et al. (2008) with the new and extended set of organic/inorganic interaction parameters determined by Zuend et al. (2011). This recent implementation of the new AIOMFAC model parameters to describe interactions between various important organic functional groups and inorganic ions enables the computation of activity coefficients in multicomponent mixtures such as those investigated in this study.

Phase separation computations were performed at 298 K for the three 5-component systems C5/AS/H₂O, C6/AS/H₂O, and C7/AS/H₂O with the components as given in Table 1. In addition, phase diagrams for three corresponding 3-component systems were computed, each containing only one of the dicarboxylic acids (one of the C5, C6, or C7 diacids used in the 5-component systems), AS, and water. In contrast to a laboratory experiment, in a model calculation the potential crystallization of a single dicarboxylic acid in a mixture can be suppressed. Hence, the ternary systems allow a direct comparison with the results for the corresponding 5-component systems at the same dry (water-free) mass fractions of AS (mf(a,AS), mf(AS) = mass(AS)/mass(organics + AS)). Such comparisons of the 5-component with 3-component systems allow an evaluation of the influence of using a mixture of three structural isomers for the organic fraction versus only one corresponding organic compound in AIOMFAC group-contribution model calculations. Furthermore, the ternary systems are of use as they allow a mathematically simpler computation of metastable and unstable regions of the phase diagram employing the algorithm of Zuend et al. (2010), as detailed in Sect. 3.5. In the calculations of the miscibility gap, the formation of crystalline AS at concentrations above its solubility limit was ignored, instead
Table 1. Molecular weights (MW), aqueous solubility in weight percent at 25°C, water activities ($a_w$) of saturated solutions at 25°C, oxygen-to-carbon atomic ratios (O:C) and hydrogen-to-carbon atomic ratios (H:C) of investigated dicarboxylic acids and AS.

<table>
<thead>
<tr>
<th>System components</th>
<th>MW (g mol$^{-1}$)</th>
<th>Solubility (wt %)</th>
<th>$a_w$ of saturated solution</th>
<th>O:C</th>
<th>H:C</th>
</tr>
</thead>
<tbody>
<tr>
<td>C5</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Glutaric acid</td>
<td>132.12</td>
<td>58.8</td>
<td>0.883</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methylsuccinic acid</td>
<td>132.12</td>
<td>32.0</td>
<td>0.943</td>
<td>0.80</td>
<td>1.60</td>
</tr>
<tr>
<td>Dimethylmalonic acid</td>
<td>132.12</td>
<td>10.5</td>
<td>0.979</td>
<td></td>
<td></td>
</tr>
<tr>
<td>C6</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2-methylglutaric acid</td>
<td>146.14</td>
<td>52.7</td>
<td>0.929</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3-methylglutaric acid</td>
<td>146.14</td>
<td>45.3</td>
<td>0.935</td>
<td>0.67</td>
<td>1.67</td>
</tr>
<tr>
<td>2,2-dimethylsuccinic acid</td>
<td>146.14</td>
<td>8.3</td>
<td>0.982</td>
<td></td>
<td></td>
</tr>
<tr>
<td>C7</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3-methyladipic acid</td>
<td>160.17</td>
<td>17.1</td>
<td>0.979</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3,3-dimethylglutaric acid</td>
<td>160.17</td>
<td>14.2</td>
<td>0.982</td>
<td>0.57</td>
<td>1.71</td>
</tr>
<tr>
<td>Diethylmalonic acid</td>
<td>160.17</td>
<td>19.5</td>
<td>0.979</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AS</td>
<td>132.14</td>
<td>43.2</td>
<td>0.802</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

For the following experiments, a C5 mixture consisting of the three C5 components, a C6 and a C7 mixture were prepared. In a first step, single droplets of these purely organic mixtures were injected into the cell and investigated in the microscope. They remained as one liquid phase droplets down to dry RH conditions and did neither show LLPS nor efflorescence. In a second step, these three-component organic mixtures were mixed together with AS and water in different OIR and investigated as single droplets. The following section shows examples of time sequences for OIR = 1:1. In Sect. 3.3 the chemical compositions of the observed phases are discussed. In Sects. 3.4 and 3.5, experimental and calculated phase diagrams are presented and discussed. Considerations of particle morphologies are given in Sect. 3.6.

3.2 LLPS features of C5/AS/H$_2$O, C6/AS/H$_2$O and C7/AS/H$_2$O droplets

Humidity cycles were performed with 4 droplets of C5/AS/H$_2$O, 9 droplets of C6/AS/H$_2$O, and 5 droplets of C7/AS/H$_2$O all with OIR = 1:1 and dry diameters in the range of 20–60 µm. Figure 2 exemplifies the phase and morphological changes that were observed optically. At the beginning of the humidity cycles at high RH (set to time $t = 0$), the droplets displayed only one liquid phase as shown in the first frames of Fig. 2a, b and c, taken at RH of 87 %, 78 % and 90 %, respectively.

The C5/AS/H$_2$O droplet effloresced at 39.2 % RH ($t = 138$ min, Fig. 2a) upon drying without showing any evidence of precedent LLPS. This value is within the range of ERH of pure AS particles (e.g. Ciobanu et al., 2010). This together
with the finding that pure C5/H₂O (AS-free) droplets did not effloresce during humidity cycles down to 20 % RH indicates that nucleation of AS initiated the efflorescence process. Upon subsequent moistening, the effloresced particle gradually absorbed water, followed by sudden dissolution of most solid material at ≈77.5 % RH (t = 251 min). Some crystals remained up to ≈85.2 % RH, when discernible dissolution again set in (t = 275 min). This high DRH cannot be ascribed to the deliquescence of AS and shows that at least a part of the C5 dicarboxylic acids crystallized, probably by heterogeneous nucleation on AS crystals. To ascertain the dissolution order of the substances, we also conducted a bulk measurement at 20 °C by adding water stepwise to the crystalline C5/AS mixture until a clear solution was obtained.

We observed by eyes that the AS crystals were completely dissolved at \( \alpha_w = 0.79 \pm 0.004 \) with the glutaric and methylsuccinic acids. Therefore, it indicates that AS in C5/AS/H₂O deliquesced at between 77.5 % and 79.0 % RH.

LLPS was observed for the C6/AS/H₂O droplet by the abrupt appearance of schlieren at 73.4 % RH (t = 30 min, Fig. 2b) upon water evaporation. Such schlieren (Kosztorz, 2001), which appear as small separated regions all over a droplet are involved in spinodal decomposition. The schlieren developed into small droplets that coalesced to form an inner phase within the particle. Ciobanu et al. (2009) have also observed spinodal decomposition for LLPS of PEG-400/AS/H₂O particles. At 71.9 % RH, inner and outer phases together with satellite inclusions can be clearly discriminated in the liquid droplet. Interestingly, after LLPS had occurred, the inner phase that was confirmed to consist of aqueous AS by Raman spectroscopy (see Sect. 3.3) tended to move from the core toward the edge of the particle as more water was released. This partially engulfed configuration can be seen at 67.2 % RH in Fig. 2b. Such particle morphology has recently been postulated by Kwamen et al. (2010) and Reid et al. (2011). A further decrease in RH leads to efflorescence and crystal growth in both phases of the particle.
at 46.3 % ($t = 238$ min). This value is above the typical range observed for ERH of pure AS droplets (e.g. Ciobanu et al., 2010) and will be discussed in more detail in Sect. 3.4. For subsequently increasing RH, the crystals in the outer phase deliquesced first, followed by fast dissolution of the solid material in the inner phase at 79.1 % RH. When AS had deliquesced at 79.6 % RH, a remnant of crystallized organic material, most likely 2,2-dimethylsuccinic acid (considering results of bulk measurement using the same method as for the C5/AS) remains (marked by an arrow on Fig. 2b, $t = 782$ min). The particle fully deliquesced as one liquid phase at 87.4 % RH.

In the C7/AS/H$_2$O droplet shown in Fig. 2c, LLPS was induced by spinodal decomposition at RH as high as 90 % ($t = 3$ min) when the humidity was decreased. Schlieren formation and coalescence can be seen at almost the same RH ($t = 3$ min and 3.1 min in Fig. 2c) leading to a core-shell morphology without satellite inclusions in the outer phase at $t = 3.6$ min. It is interesting to note that the core-shell or completely engulfed morphology is the main configuration in the C7/AS/H$_2$O particle for OIR = 1:1 while for C6/AS/H$_2$O droplets the partially engulfed morphology prevailed. A gradual loss of water resulted in a shrinkage of the particle until the inner phase effloresced at 41.4 % RH. The outer phase that was confirmed to consist mainly of organics (see Sect. 3.3) did not seem to crystallize, probably because the very low AS concentration or a high viscosity in this phase derates crystal growth. Rather, it was sucked into cavities of the crystalline inner phase within a few seconds ($t = 139.3$ min) most probably because of capillary forces (Sjogren et al., 2007). Undergoing water uptake, the particle deliquesced and returned into a two-liquid-phases state at 79 % RH. In contrast to the C6/AS/H$_2$O particle (Fig. 2b, $t = 782$ min), the C7/AS/H$_2$O particle shows no remaining solid above the DRH of AS. The two liquid phases merged into one liquid phase at 89.1 % RH.

### 3.3 Chemical compositions of the different phases

To identify the chemical composition of the different phases present in C6/AS/H$_2$O and C7/AS/H$_2$O particles which have shown LLPS, we collected Raman spectra at constant RH on particles with OIR = 1:1. Figure 3 depicts representative Raman spectra collected at different positions as indicated in the optical images of the particles. Spectra of an AS crystal (Fig. 3, c1) and a liquid droplet of purely organic 3-components C6 and C7 mixtures at RH = 0 % are also included as references (Fig. 3, c2). The most prominent feature

![Fig. 3. Raman spectra and microscopic images of (a) C6/AS/H$_2$O and (b) C7/AS/H$_2$O particles with OIR = 1:1 at different RH. Reference spectra are included for AS (c1) and for the pure C6 and C7 organic mixtures (c2). Size bars in the microscopic images are 20 µm.](http://www.atmos-chem-phys.net/12/2691/2012/)

www.atmos-chem-phys.net/12/2691/2012/
in the reference spectrum of crystalline AS is the symmetric stretching vibration of \( \text{SO}_4^{2-} \) at \( \sim 975 \text{ cm}^{-1} \) (Zhang and Chan, 2002; Ling and Chan, 2008; Ciobanu et al., 2009). The band of the stretching vibration of \( \text{NH}_3^+ \) at around \( 3130 \text{ cm}^{-1} \) (Ciobanu et al., 2009; Yeung et al., 2009) is also distinguished. The C6 and C7 references reveal C-H stretching vibrations at 2880–2984 and 2875–2990 \text{ cm}^{-1}, respectively, and several peaks between 700 and 1670 \text{ cm}^{-1}. The C=O stretching vibrations appear as relatively sharp peaks at \( \sim 1656 \text{ cm}^{-1} \) for the C6 and at \( \sim 1658 \text{ cm}^{-1} \) for C7 mixtures.

In the one-liquid-phase state of C6/AS/H\(_2\)O and C7/AS/H\(_2\)O droplets, the sharp peak at \( \sim 980 \text{ cm}^{-1} \) and the broad band between 2870 and 3000 \text{ cm}^{-1}, originating from AS and dicarboxylic acids, respectively (a1 and b1 of Fig. 3), are clearly visible. In addition, at this high RH, symmetric and antisymmetric O-H stretching modes of water molecules can be detected as an additional intense broad band in the range of 3030–3700 \text{ cm}^{-1} (Yeung et al., 2009). This is clear evidence for the presence of both AS and organics in one aqueous phase.

At 64 % RH, when the C6/AS/H\(_2\)O droplet is in the two-liquid-phases state, Raman spectra were collected from the satellite inclusions, the inner phase and the outer phase. As shown by the dominance of the AS peak at 979 \text{ cm}^{-1}, the low intensity of the water band and the hardly visible spectral features of the dicarboxylic acids in spectra (a2) and (a3) of Fig. 3a, the inner phase and the satellite inclusions consist of a concentrated aqueous AS solution containing only small amounts of organics. On the other hand, spectrum (a4) acquired from the outer phase of C6/AS/H\(_2\)O shows well discernable signals of the dicarboxylic acids and the AS and water bands appear only as weak features, establishing the outer phase as highly concentrated aqueous dicarboxylic acid solution with AS as a minor component (see also Ciobanu et al., 2009; Smith et al., 2011).

Raman spectra of the outer and the inner phases of the C7/AS/H\(_2\)O droplet measured at 71 % RH give evidence of an almost complete phase separation: the dicarboxylic acid peaks are hardly visible in the Raman spectrum of the inner aqueous AS phase (b2 of Fig. 3b) and the main AS peak at 979 \text{ cm}^{-1} is very weak in the Raman spectrum of the outer organic-rich phase (b3). Also, the spectral signature of the water band can be hardly detected in the organic-rich phase indicating its low water content. More complete phase separation in the C7/AS/H\(_2\)O system than in the C6/AS/H\(_2\)O system is expected because of the more hydrophobic nature (lower O:C) of the C7 compared with the C6 dicarboxylic acids.

Spectra of the effloresced particles were collected at 35 % and 13 % RH for the C6/AS/H\(_2\)O and the C7/AS/H\(_2\)O droplets, respectively. For both systems efflorescence of the AS component is confirmed by the increased sharpness of the peaks at 451 and 614 \text{ cm}^{-1} and the shift of the main peak from \( \sim 979 \) to \( \sim 975 \text{ cm}^{-1} \). Such effects due to the crystallization of AS have been described by Zhang and Chan (2002) and Ciobanu et al. (2009). The Raman spectra do not provide any clear evidence of the phase state of the dicarboxylic acids. The distribution of the two liquid phases within the droplet is mostly conserved during efflorescence as can be seen optically and from comparison of the Raman spectra (c1) and (c2) of Fig. 3c: the effloresced inner phase is enriched in crystalline AS compared with the outer phase. Nevertheless, a certain remixing of the phases takes place after efflorescence, as can be clearly seen from the increased intensity of the dicarboxylic acid spectral features in the inner phase of the C6/AS/H\(_2\)O droplet after efflorescence (spectrum (a5) compared with (a2)) and in the inner phase of the C7/AS/H\(_2\)O droplet (spectrum (b4) compared with (b2)). We assume that this remixing is driven by capillary forces that act in veins of polycrystalline AS and cavities between AS needles.

### 3.4 Phase diagrams of C6/AS/H\(_2\)O and C7/AS/H\(_2\)O systems

To fully assess the dependence of LLPS on mixing ratio and RH and to establish corresponding phase diagrams, C5/AS/H\(_2\)O, C6/AS/H\(_2\)O and C7/AS/H\(_2\)O particles with different OIR were investigated at 20 °C. We performed small humidity cycles (from 95 to 60 % RH) to determine whether there is a hysteresis between the onset of LLPS upon drying and the merging of the two liquid phases upon moistening, and large humidity cycles (from 95 to 20 % RH) to observe efflorescence and deliquescence processes. Typically, a small humidity cycle was followed by a large one. For some particles, we performed one to several large humidity cycles. The sizes of the investigated droplets were in the range from 24–75 µm diameters.

In the C5/AS/H\(_2\)O particles LLPS did not occur for the investigated OIR = 2:1, 1:1 and 1:4, instead, efflorescence occurred at 32.4 %, 37.9 %, and 40.8 % RH, for OIR = 2:1, 1:1, and 1:4, respectively. If LLPS occurred in this system, it would be at RH < ERH. However, LLPS appeared in both C6/AS/H\(_2\)O and C7/AS/H\(_2\)O systems over a broad composition range. Figure 4 presents phase diagrams of (a) C6/AS/H\(_2\)O and (b) C7/AS/H\(_2\)O systems as a function of RH in terms of \( \text{m}_{\text{eff}}(\text{AS}) \) (lower x-axis) and OIR (upper x-axis).

In C6/AS/H\(_2\)O particles LLPS occurred upon drying for \( \text{m}_{\text{eff}}(\text{AS}) \) of 0.07–0.89. Humidity cycles performed between 95 % and 60 % RH show no discernable hysteresis between onset and end of LLPS, i.e. the open and filled black circles in both Figs. 4a and b coincide within experimental uncertainties. The experiment carried out at \( \text{m}_{\text{eff}}(\text{AS}) = 0.92 \) did not show any discernable LLPS, marking the end of the miscibility gap to high AS concentrations, though we cannot exclude the possibility that separation occurred but could not be detected because of lack of contrast. For \( \text{m}_{\text{eff}}(\text{AS}) = 0.30–0.89 \) the onset and end of LLPS is in the narrow range from 72–74 % RH. On the other hand, in C7/AS/H\(_2\)O particles...
we observed LLPS over the whole investigated composition range with onset values between 84–92 % RH upon drying and end values between 86–90 % RH during moistening, i.e. with no discernable hysteresis (Fig. 4b). The larger miscibility gap of the C7/AS/H2O compared with the C6/AS/H2O system in terms of composition and RH range is consistent with the more hydrophobic character of the C7 compared with the C6 dicarboxylic acids. A further decrease in RH leads to efflorescence of the particles. In Fig. 4a and b the average ERH of all experiments carried out with the same composition are shown as green filled circles, ERH of individual crystallization events as red filled circles. Open red circles show cases when crystal growth was very slow and mark the state with full needle coverage (this RH value depends on the rate at which particles were dried). The ERH scatters between individual experiments because nucleation is a stochastic process (Martin, 2000) while the instrumental uncertainty is smaller than this scatter by factors 3 to 4. In general, ERH decreases as AS concentrations decrease. For mf_d(AS) ≥ 0.5, ERH values range from 35–48 % RH for the C6/AS/H2O system and between 36–48 % RH for the C7/AS/H2O system as well. These ranges are at the high end or above ERH observed for other dicarboxylic acid/AS/H2O systems: e.g. Parsons et al. (2004) and Zardini et al. (2008) report ERH for malonic acid/AS, glutaric acid/AS, and citric acid/AS particles consistently below 40 %. A possible explanation for the high ERH values observed in this study might be an increase of the ion activity coefficients due to the energetically disadvantageous interactions between the ions of AS and the C6 and C7 dicarboxylic acids, enhancing the probability for the formation of stable AS crystal nuclei at higher RH as
compared to mixtures with the more hydrophilic malonic (C3 dicarboxylic acid), glutaric (C5 dicarboxylic acid) and citric acids.

Figure 5 illustrates nucleation and crystal growth of AS for a C6/AS/H2O particle with OIR = 11:1. The main phase of this particle is organic-rich. Aqueous AS inclusions are marked by arrows. Crystallization started from the rim of the particle at 35 % RH and slowly proceeded until after 6.5 min the particle was fully covered with needles. Because our experiments were carried out at constant drying rate, this translates to a crystal growth that occurs over several percent in RH until a droplet is fully covered by needles as indicated in Fig. 4 (open red circles). The needles mostly consist of AS because they deliquesce at around the DRH of AS (open blue circles in Fig. 4). The aqueous AS inclusions crystallized when the AS needle had reached them. We attribute the slow crystal growth to the high viscosity that is reached in the organic-rich phase.

Upon increasing RH, deliquescence of AS occurred between 70–80 %RH for both systems over the whole composition range. This is in good agreement with previous work (Braban and Abbatt, 2004; Parsons et al., 2004; Sjogren et al., 2007; Ling and Chan 2008; Smith et al., 2011). Upon moistening, the C6/AS/H2O and C7/AS/H2O particles showed noticeably different behavior. Whereas C6/AS/H2O particles returned into a one-liquid phase state upon deliquescence of AS, a two-liquid-phases state was restored when AS deliquesces in C7/AS/H2O particles. The reason for this different behavior is that for C7/AS/H2O the miscibility gap extends to RH > AS DRH while the miscibility gap of C6/AS/H2O is restricted to compositions that are supersaturated with respect to AS and therefore ends at RH < AS DRH.

In the C6 system, the situation is intricate at low AS concentration. LLPS was clearly visible for mf_d(AS) ≈ 0.1. In this concentration range, the highest observed onset of LLPS occurred (at RH ≈ 85 %RH for mf_d(AS) = 0.095, OIR = 9.5:1). For mf_d(AS) = 0.08, LLPS drops to 50 %RH marking the end of the miscibility gap to low AS concentrations. Surprisingly, for mf_d(AS) = 0.19 and 0.21 (hatched area in Fig. 4a) LLPS was only detectable in 7 out of 12 and 7 out of 9 experiments, respectively. In cases with detectable LLPS, the onset RH scattered over quite a large RH range of 65–78 %. Interestingly, this large range cannot be explained by kinetically limited nucleation of the minor phase because no phase separation hysteresis could be observed when exposing the same particle to humidity cycles. Some particles showed reproducible LLPS during several humidity cycles while LLPS was reproducibly absent in others.

In order to clarify LLPS at high OIR in the C6/AS/H2O system, we collected Raman spectra of the C6/AS/H2O and the C7/AS/H2O systems in the one-liquid-phase state and in the organic-rich phase of the phase separated system for different mf_d(AS), see Fig. 6. From theory, we expect that mixtures with OIR within the miscibility gap partition into an organic-rich and an aqueous AS phase with constant compositions corresponding to the ones of the limits of the miscibility gap, irrespective of their OIR. This should be strictly true for systems measured at the same RH. The organic-rich phase (given by the red circles in Fig. 6) should therefore exhibit a constant dicarboxylic acid/AS Raman peak ratio irrespective of the OIR. By comparing the Raman peak height ratios of the organic-rich phase with the ones in the one-liquid-phase state, it is furthermore possible to estimate the AS concentration of the organic-rich phase. To do this, we use the C6 (2931 cm−1)/AS (979 cm−1) and the C7 (2937 cm−1)/AS (979 cm−1) Raman peak height ratios of the one-liquid-phase states (at 85 %RH and 95 %RH for the C6/AS/H2O and C7/AS/H2O systems, respectively) as a measure of AS concentration given as mf_d(AS) on the x-axis of Fig. 6. From this, we deduce an estimate of the AS concentration in the organic-rich phase in the phase separated system at RH = 65 %. Figure 6b shows a constant C7/AS ratio of ~15 corresponding to mf_d(AS) well below 0.1 for the organic-rich phase of the C7/AS/H2O system – as expected from theory. The organic-rich phase of the C6/AS/H2O system exhibits a C6/AS ratio of ~1 that corresponds to mf_d(AS) ~ 0.3 for overall compositions of mf_d(AS) = 0.50 and 0.67 (Fig. 6a). This implies that the miscibility gap of this system is less complete than the one of the C7/AS/H2O system and should end at mf_d(AS) ~ 0.3. However, the observed phase separation persists to lower mf_d(AS), and, in addition, the AS content of the organic-rich phase is no longer a constant but decreases with decreasing mf_d(AS) of the particle (corresponding to an increase of the C6/AS ratio up to ~8 for an overall mf_d(AS) of 0.08). This implicates
that C6/AS/H2O systems with \( m_f(AS) \) below 0.3 do not exhibit a typical LLPS. This issue will be discussed in detail in Sect. 3.5.

Ciobanu et al. (2009) found that LLPS may occur by spinodal decomposition, nucleation-and-growth, and growth of a second phase at the surface of the particle depending on OIR within internally mixed PEG-400/AS/H2O particles. Nucleation-and-growth typically occurs in the metastable region of the phase diagram and is associated with overcoming an energy barrier to form stable nuclei of the second phase while spinodal decomposition occurs without a nucleation energy barrier at the border to the unstable region of the phase diagram (Shelby, 1995; Papon et al., 1999). To determine the different LLPS mechanisms from the microscopic images, we used the same methods as described by Ciobanu et al. (2009), which rely on the analysis of droplet morphology at the onset of phase separation and the evaluation of the number of AS inclusions as a function of time. Such assignment of phase separation mechanism is always prone to uncertainties because the very first stages of LLPS are not visible in optical microscopes due to limitations in spatial resolution and optical contrast. Given these limitations, in the C6/AS/H2O systems, LLPS occurring at 0.07 < \( m_f(AS) \) < 0.40 matched best with a nucleation-and-growth mechanism whereas spinodal decomposition seemed to occur for 0.40 < \( m_f(AS) \) < 0.70 (Fig. 4a). However, we were not able to determine the separation mechanism for LLPS occurring at 0.70 < \( m_f(AS) \) < 0.89 in the C6/AS/H2O systems (Fig. 4a) because the very first instances of phase separation are not clearly discernable due to lack of optical contrast. Figure 7a shows as an example for this case a time sequence of a droplet with OIR = 1:4. This droplet remained very dynamic with continuous appearance and coalescence of inclusions for a time period over 30 min (corresponding to an RH change of 20 %). Here, the organic-rich phase does not seem to spread on the aqueous AS phase but to form droplets or lenses on the surface which might be moved by the \( N_2/H_2O \) flow passing over the particle.

On the other hand, the C7/AS/H2O particles are showing nucleation-and-growth mechanism (example in Fig. 7b), spinodal decomposition, and growth of a second phase at the surface of the particle (example in Fig. 7c) at \( m_f(AS) \) of <0.30, 0.30 to 0.60 and 0.6 to 1.0, respectively. These different phase separation mechanisms are indicated as different shadings in Fig. 4b.
3.5 Calculated phase diagrams

Thermodynamic model calculations for the three systems have been conducted to support the comparison and interpretation of the experimentally determined phase diagrams. While the measurements establish the phase diagrams in the RH vs. dry (water-free) composition space, a thermodynamic calculation reveals the compositions (including water content) of the two phases in liquid-liquid equilibrium (LLE).

On the basis of a mathematically much simpler computation of phase stabilities in case of only three thermodynamically independent components, the limit of stability between metastable and unstable one-phase states, the spinodal curves, were calculated only in case of the ternary systems (see Zuend et al., 2010). However, for reasons of similarity, further discussed below, the computed spinodal curves also support the interpretation of the phase stabilities in case of the C5/AS/H2O, C6/AS/H2O, and C7/AS/H2O systems.

Figure 8 shows the computed phase diagrams for the different systems at room temperature. The model predicts a miscibility gap for all systems, the extent of which however differs in composition and RH range. In agreement with the experimental phase diagrams, the miscibility gap is largest in case of the C7/AS/H2O system and becomes smaller with decreasing size of the dicarboxylic acids and increasing O:C, until it disappears completely. The mass fraction compositions of coexisting phases, as shown in the upper panels of Fig. 8, reveal that water is present in both phases in similar fractions, the difference being largest in case of the C7/AS/H2O system, as seen from the slopes of the tie-lines. However, the difference in water contents is less pronounced when considered in terms of mole fractions since the difference in molar mass of the C5, C6, and C7 dicarboxylic acids explains a part of the apparent difference between panels (a), (b), and (c). The difference in terms of AS and organic fractions in coexisting phases is much more distinct. This implies that both the dicarboxylic acids and AS (dissociated into NH4+ and SO42− ions in solution) are quite hydrophilic, but that interactions between the ions and the organics lead to a LLPS at higher AS mass fractions, when, at lower RH, the present fraction of water is not sufficient to moderate those interactions. Since mostly the interactions between non-polar organic molecule segments, here the CH2 and CH3 groups, and ions are causing the phase separation, it is not surprising that the miscibility gap increases with a decrease in the O:C. The model also predicts that all three salt-free systems are miscible with water in all proportions.

While many features of the calculated phase diagrams agree with the droplet experiments at least qualitatively, there are also a number of differences, discussed in the following. Most strikingly, in contrast to the measurements, the model predicts a phase separation region in the C5/AS/H2O system. In principle, there could also be a phase separation region in the C5/AS/H2O droplets in reality, but only present at RH below the ERH of the droplets and therefore outside of the experimentally accessible range. However, the model predicts the maximum onset RH of LLPS already at ~60 % (a0 = 0.6), see Fig. 8b. It is therefore obvious, that the predicted onset RH of LLPS is too high, in case of C5/AS/H2O by at least 20 % RH. For an analogous ternary C4/AS/H2O system containing succinic acid, the model still predicts a hypothetical LLPS (not shown), but only below the ERH of AS (<40 % RH) and only for mf0(AS) above 0.6. For a ternary C3/AS/H2O system containing malonic acid, no LLPS is predicted in the entire composition space. Comparisons of predictions and measurements for the C6/AS/H2O and C7/AS/H2O systems, show that the model also predicts higher LLPS onset RH for these systems, with a difference (model – measurement) of ~8 % RH in case of C6/AS/H2O and ~2 % RH in case of C7/AS/H2O at OIR = 1:1. These differences are considerably smaller than in the case of the C5/AS/H2O system. Regarding the group-contribution concept used within AIOMFAC to represent organic molecules and their interactions with other species/functional groups, it is not surprising that there are some differences between model predictions and measurements. As described in Zuend et al. (2011), the required AIOMFAC interaction parameters between ions and organic functional groups were determined based on a large, but unevenly distributed amount of experimental data, using a sophisticated parameter optimization procedure to obtain best simultaneous agreement between calculated and measured thermodynamic data. As it turns out, the parameterization database contains mainly water activity data for the optimization of interaction parameters between NH4+ and SO42− ions of AS and the carboxyl group of dicarboxylic acids. Only three solid-liquid equilibria datasets and not a single LLE dataset have been available to better constrain these interactions with potential consequences for the prediction of LLPS. In view of this, the model predictions are in remarkably good agreement with the measured C7/AS/H2O data. The larger deviations between model and measurements in case of the C5/AS/H2O mixtures can also be understood as a result of less constrained AIOMFAC interaction coefficients towards high organic concentrations (and low water mass fractions).

The calculated and measured coexistence curves of the C6/AS/H2O systems agree within ~8 % RH for mf0(AS), from 0.3 to 0.9. Comparing the coexistence curve of the C6/AS/H2O system at mf0(AS) < 0.25 from Fig. 4a (measurements) with panel (e) of Fig. 8 shows that there is disagreement at those lower salt concentrations. While the model suggests a single mixed phase, the experimental data show that a phase separation is present, however, with rather unusual variations in the LLPS onset RH. For some experiments at compositions 0.19 < mf0(AS) < 0.21 no LLPS was observed, which would be in agreement with the model results. Moreover, the mf0(AS) measured by Raman spectroscopy in the organic-rich phase of C6/AS/H2O phase separated systems agree with a miscibility gap that only extends to mf0(AS) of ~0.3 (Fig. 6a). Furthermore, the measured
variable AS concentration of the organic-rich phase for OIR = 4:1 and higher, suggests that we do not have a regular LLPS in this composition range. The reproducibly high onset and end of LLPS at RH = 85% for mfd(AS) = 0.095 might be considered as an indication that the minor phase has an almost fixed stoichiometry and might be highly structured. Also, additional ionic species might be important for the formation of this phase because at those lower AS mass fractions and higher water mass fractions, the dicarboxylic acids dissociate to a significant degree, yet concentration-dependent and additional ionic species such as bisulfate ions (HSO$_4^-$) form, leading to a different mixture composition (than used in the model calculations). From Fig. 4b, it looks like a similar but less pronounced phase behavior could be present in the C7/AS/H$_2$O droplets at mfd(AS) < 0.15.
The lower panels (c), (f), and (i) of Fig. 8 show model calculations for ternary mixtures containing glutaric acid, 2-methylglutaric acid, and 3,3-dimethylglutaric acid, respectively, corresponding to the 5-component systems of the panels above them. The predicted phase separation regions of the ternary systems are highly similar to those of the corresponding 5-component systems. This result is expected, since the main difference between the 5-component systems and the ternary systems is the mixture of three similar dicarboxylic acids of equal mass, O:C and hydrogen-to-carbon atomic ratios (H:C), and the same kinds of functional groups, instead of only one dicarboxylic acid. Hence, the applied group-contribution approach of AIOMFAC can only lead to very small differences among the activity coefficients of structural isomers of dicarboxylic acids in aqueous AS solutions. Based on these phase diagram similarities and the underlying modeling reasons, it is also clear that the calculated spinodal curves of the ternary diagrams apply as well to the 5-component systems. The position and shape of the unstable one-phase region enclosed by the spinodal curve in panels (f) and (i) of Fig. 8, relative to the position of the coexistence curve (binodal), suggests that spinodal decomposition may occur in the C6/AS/H2O system in the composition range of \(\sim 0.4 < mf_d(\text{AS}) < \sim 0.7\), which is in good agreement with the observed mechanism in the experiments. At higher AS mass fractions, \(mf_d(\text{AS}) > \sim 0.7\), a spinodal decomposition could still take place, but the \(aw\)-difference between binodal and spinodal curves becomes larger, so that nucleation of a second phase already at a higher RH, before the spinodal curve is reached, becomes more likely. In case of the C7/AS/H2O droplets, the model suggests that spinodal decomposition may occur in the mass fraction range \(\sim 0.11 < mf_d(\text{AS}) < \sim 0.55\). As compared to the experiments, the predicted range is extended to lower mass fractions of AS; the experiments suggest a lower range limit of \(mf_d(\text{AS}) = \sim 0.3\). The measured onset RH (89.7 %) for spinodal decomposition in C7/AS/H2O droplets at OIR = 1:1 agrees well with the predicted spinodal curve (90 % RH) at this composition.

Figure 9 shows the computed Gibbs energy difference between a forced one-liquid-phase state of the mixture and the equilibrium state (two-liquid-phases state in the LLPS region) for the C6/AS/H2O and C7/AS/H2O systems at 298 K in the RH range from 100 to 30 %. Close to the coexistence curve, the Gibbs energy difference \(\Delta G\) is rather shallow but increases steeply at lower RH within the phase separation region. This indicates that metastable solutions can exist at water activities slightly below the coexistence curve so that the spinodal limit can be reached in experiments before a nucleation-and-growth event happens. The curvature of the ammonium sulfate solubility limit (the white dashed-dotted curve in Fig. 9) in both systems indicates AS deliquescence RH close to the value in the binary aqueous ammonium sulfate solution for the dry mass fraction range \(\sim 0.2 < mf_d(\text{AS}) \leq 1.0\). At higher organic contents, \(mf_d(\text{AS}) < \sim 0.2\), the deliquescence RH drops sharply to lower values due to the reduced amount of AS in the mixture. These DRH predictions agree quite well with the measured AS DRH values shown in Fig. 4.

### 3.6 Possible morphologies of droplets on a hydrophobically coated substrate

The optical microscope gives only the top view of the droplets on the substrate. It is therefore not possible to directly observe whether inclusions of the minor phase have contact to the droplet/air or the droplet/substrate interface or whether they are floating within the droplet volume.
To resolve this issue, we performed confocal Raman scans through the droplets by changing the laser focus from the bottom to the top of the droplets with step sizes of 6–7 µm. Figure 10 shows an example of such a vertical scan. Panel (a) gives the optical image of the droplet. The scan was performed through an inclusion with a diameter of ∼19 µm (marked by a white arrow). For comparison, a Raman spectrum of the outer phase (marked by a black arrow) was also collected. Evaluation of the Raman spectra relied on the ratio of the peak height of the main peaks of AS (979 cm$^{-1}$) and C6 (2931 cm$^{-1}$) and C6 (2931 cm$^{-1}$). This AS/C6 peak ratio is shown in Fig. 10b as a function of laser focus within the droplet. We determined the bottom of the droplet (droplet height = 0 µm) optically as the point where the rim of the droplet appeared sharp and the diameter of the droplet was largest. Moving the focus upwards, the observed diameter decreased (gray arrows in c) and the droplet edge became increasingly blurred making the diameter measurement inaccurate. To determine the top of the droplet, we in addition reconstructed the droplet shape assuming a spherical cap with a contact angle of 81.6 ± 2° (determined for 2 µl C6/AS/H$_2$O droplets with OIR = 1:1 (∼40 wt % (C6 + AS)) deposited on the hydrophobically coated substrate (see panel c) leading to a droplet height of 25.4 µm for the investigated droplet with diameter of 58.9 µm. This droplet height is given as dashed gray line in (b). The highest AS/C6 Raman peak ratios were measured at droplet heights of 6 and 12 µm. Above 20 µm the AS/C6 peak ratio remained constant at a value slightly above the one of the outer phase (given as a black triangle in b) while the absolute Raman intensities decreased because the laser focus was moving out of the droplet. This clearly shows that the inclusion was at the bottom of the droplet, probably with a high contact angle to the substrate (>90°) or forming almost a sphere. We performed similar experiments for two other droplets with smaller inclusions (∼10 µm diameter), which clearly showed that these inclusions were also at the bottom of the droplet. This morphology is also supported by the calculation of the settling velocity of aqueous AS inclusions in the organic-rich phase: settling of 10 and 20 µm diameter AS-rich phase inclusions should occur at a rate of 0.5–3.3 and 2–13 µm min$^{-1}$, respectively. We therefore expect that the AS-rich inclusions settle to the bottom of the droplet driven by gravity within the timescale of the experiment (see Appendix A for details of the calculation).

In all investigated C6/AS/H$_2$O droplets with OIR = 1:1, the main AS inclusion moved from the middle to the edge of the droplet between ∼70–66 % RH and remained in this partially engulfed configuration down to ERH. A similar behavior was observed for C6/AS/H$_2$O droplets between ∼63–50 % RH with OIR = 1:2. Partial engulfed configurations were also observed for C7/AS/H$_2$O droplets ∼88 % RH with OIR = 2:1.

The morphology of droplets with sizes of several tens of micrometers is influenced by surface and interfacial forces and by gravity. The spreading coefficient (Torza and Mason, 1970; Kwamena et al., 2010; Reid et al., 2011) can be used to predict the equilibrium morphology of two immiscible phases when in contact with each other. The spreading coefficient ($S_i$) of a liquid component (i) spreading over another component (j) can be determined when the surface ($\sigma_{jk}$, $\sigma_{ik}$) and interfacial tensions ($\sigma_{ij}$) are known:

$$S_i = \sigma_{jk} - (\sigma_{ij} + \sigma_{ik}).$$

For a partial engulfed configuration, the spreading coefficient of the organic-rich phase on the aqueous phase needs to be smaller than zero, which implies an interfacial tension
between the two liquid phases of $\sigma_{\text{org/AS}} > \sigma_{\text{AS}} - \sigma_{\text{org}}$. Assuming an upper limit for the surface tension of the C6-rich phase of $\sigma_{\text{org}} = 52 \text{ mN m}^{-1}$ and a lower limit for the AS-rich phase of $\sigma_{\text{AS}} = 67 \text{ mN m}^{-1}$ (see Appendix B for estimates of surface and interfacial tensions), an interfacial tension of at least $15 \text{ mN m}^{-1}$ is required for a partial engulfed configuration, which is in the typical range of interfacial tensions of aqueous/organic two-liquid-phases systems. It can be assumed that the interfacial tension increases with decreasing RH because water as mediating component between the phases becomes more and more scarce. Between 70–66 % RH the interfacial tension seems to reach a critical value where the spreading coefficient of the C6-rich phase changes sign from a positive to a negative value making a partially engulfed configuration energetically more favorable.

Using Young’s equation, the contact angle of the aqueous AS inclusion with the substrate can be estimated (see Appendix B), leading to values in the range of 105–140°. Droplets with these lower and upper values are sketched in blue in Fig. 10c, the best estimation of the droplet’s morphology is given in red.

3.7 Atmospheric implications

Recently, O:C has been stressed as crucial parameter to describe hygroscopicity (Jimenez et al., 2009), oxidation (Heald et al., 2010; Kroll et al., 2011), volatility and mixing thermodynamics of organic aerosol (Donahue et al., 2011). O:C and H:C of ambient and laboratory aerosol particles can be derived from analysis of high resolution mass spectral data (Aiken et al., 2007, 2008). Heald et al. (2010) established that atmospheric organic aerosols, obtained from various field measurements and laboratory studies, cluster along a narrow line with a slope of $-1$ in Van Krevelen space (H:C vs. O:C). O:C can also be an indicator of phase separation of mixed particles. Figure 11 gives the position of the organic components of hydroxyl and PEG-400/AS/H$_2$O (blue) and aromatic or aliphatic carboxylic acid/AS/H$_2$O systems (red) from literature (see Table 2) in the Van Krevelen diagram. The C5/AS/H$_2$O, C6/AS/H$_2$O and C7/AS/H$_2$O systems from this study are also included (orange symbols). It can be seen that systems with O:C < 0.7 show LLPS (circles) while systems with O:C > 0.7 do not (crosses). This is in perfect agreement with Bertram et al. (2011) who reported in a recent study that O:C is a critical factor for LLPS of aerosol particles and showed that there is no phase separation for O:C > 0.7 for various single organic compounds in the presence of AS. Therefore, we expect that in the troposphere LLPS occurs in mixtures of AS and organic components with an O:C < 0.7. Analysis of ambient data indicates O:C up to 0.6 for semivolatile oxygenated organic aerosols (SV-OOA) and from 0.5–1.1 for low volatility OOA (LV-OOA) (Ng et al., 2011). All of the SV-OOA and a part of the LV-OOA, therefore, fall into the O:C range for which we expect LLPS. The degree of oxidation that chamber organic aerosol reach strongly depend on the volatile organic precursor molecules: secondary organic aerosol (SOA) formed by oxidation of α-pinene, isoprene, m-xylene and toluene mostly exhibit O:C < 0.7 while e.g. acrolein and glyoxal SOA cover O:C > 0.7 (Chhabra et al., 2011). In addition, LLPS might promote the formation of particles with highly viscous and glassy coatings because the more viscous organic components are spatially separated from the less viscous inorganic salt solution. In this study, partially engulfed configuration was observed in the C6/AS/H$_2$O system for OIR = 1:1 and 1:2 while both partial (OIR = 2:1) and complete engulfed (OIR = 1:1) configurations occurred in the C7/AS/H$_2$O system. No size dependence of the morphology was observed for particles within the investigated size ranges. Due to the Kelvin effect, more complex morphologies are thermodynamically not favored in accumulation mode particles. Based on this and other recent studies (Ciobanu et al., 2009; Kwamena et al., 2010; Bertram et al., 2011; Reid et al., 2011), core-shell and partially engulfed configurations seem to occur. A core-shell configuration will have more drastic consequences for heterogeneous chemistry and hygroscopicity than a partially engulfed configuration because the aqueous inorganic-rich phase will be totally enclosed by a probably highly viscous organic coating with low diffusivity for reactants and water. E.g., atmospheric chemistry related to the N$_2$O$_5$ heterogeneous hydrolysis onto
Table 2. List of “no LLPS” and “LLPS” exhibiting mixtures consisting of aqueous AS with organic components containing hydroxyl (OH), carboxyl (COOH), ether (CHnO), ester (CCOO), and aromatic (ACHn) functional groups. Range of experimental conditions: 30 % < RH < 100 % and mfd(AS) > 0.1. Data from the literature and this work.

<table>
<thead>
<tr>
<th>Compounds</th>
<th>Formula</th>
<th>Functional Groups</th>
<th>O:C</th>
<th>H:C</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>glycerol</td>
<td>C3H8O3</td>
<td>OH</td>
<td>1.00</td>
<td>2.67</td>
<td>Marcolli and Krieger (2006)</td>
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<tr>
<td>1,2,4-butane triol</td>
<td>C4H10O3</td>
<td>OH</td>
<td>0.75</td>
<td>2.50</td>
<td>Bertram et al. (2011)</td>
</tr>
<tr>
<td>1,2,6-dioxabicyclo[3.2.1]octane-2,3,4-triol</td>
<td>C5H8O4</td>
<td>COOH</td>
<td>0.80</td>
<td>1.60</td>
<td>Bertram et al. (2011)</td>
</tr>
<tr>
<td>citric acid</td>
<td>C6H8O7</td>
<td>COOH, OH</td>
<td>1.17</td>
<td>1.33</td>
<td>Bertram et al. (2011)</td>
</tr>
<tr>
<td>propanedioic acid</td>
<td>C3H4O4</td>
<td>COOH</td>
<td>1.33</td>
<td>1.33</td>
<td>Bertram et al. (2011)</td>
</tr>
<tr>
<td>C5: glutaric acid + dimethylmalonic acid + methylsuccinic acid</td>
<td>C5H8O4</td>
<td>COOH</td>
<td>0.80</td>
<td>1.60</td>
<td>This work</td>
</tr>
<tr>
<td>1,2,4-hexanediol</td>
<td>C6H14O2</td>
<td>OH</td>
<td>0.33</td>
<td>2.33</td>
<td>Marcolli and Krieger (2006)</td>
</tr>
<tr>
<td>1,4-butanediol</td>
<td>C4H10O2</td>
<td>OH</td>
<td>0.50</td>
<td>2.50</td>
<td>Marcolli and Krieger (2006)</td>
</tr>
<tr>
<td>1,2,6-hexane-triol</td>
<td>C6H14O3</td>
<td>OH</td>
<td>0.50</td>
<td>2.33</td>
<td>Bertram et al. (2011)</td>
</tr>
<tr>
<td>2,4-pentanediol</td>
<td>C5H12O2</td>
<td>OH</td>
<td>0.40</td>
<td>2.40</td>
<td>Zuend et al. (2008)</td>
</tr>
<tr>
<td>2,5-hexanediol</td>
<td>C6H14O3</td>
<td>OH</td>
<td>0.33</td>
<td>2.33</td>
<td>Zuend et al. (2008)</td>
</tr>
<tr>
<td>1,7-heptanediol</td>
<td>C7H16O2</td>
<td>OH</td>
<td>0.29</td>
<td>2.29</td>
<td>Zuend et al. (2008)</td>
</tr>
<tr>
<td>monomethyloctane-1,8-dioate</td>
<td>C10H16O4</td>
<td>COOH, CCOO</td>
<td>0.44</td>
<td>1.78</td>
<td>Bertram et al. (2011)</td>
</tr>
<tr>
<td>4-dihydroxy-3-methoxybenzeneacetic acid</td>
<td>C10H16O5</td>
<td>COOH, OH, CHnO, ACHn</td>
<td>0.56</td>
<td>1.11</td>
<td>Bertram et al. (2011)</td>
</tr>
<tr>
<td>2,5-dihydroxybenzoic acid</td>
<td>C7H6O4</td>
<td>COOH, OH, ACHn</td>
<td>0.57</td>
<td>1.50</td>
<td>Bertram et al. (2011)</td>
</tr>
<tr>
<td>2,2-dimethylbutanedioc acid</td>
<td>C6H10O4</td>
<td>COOH, OH, ACHn</td>
<td>0.67</td>
<td>1.67</td>
<td>Bertram et al. (2011)</td>
</tr>
<tr>
<td>PEG-400</td>
<td>C2nH4n+2O4n+1,</td>
<td>(n = 8–9)</td>
<td></td>
<td></td>
<td>Ciobanu et al. (2009)</td>
</tr>
<tr>
<td>C6: 2-methylglutaric acid + 3-methylglutaric acid + 2,2-dimethylsuccinic acid</td>
<td>C6H10O4</td>
<td>COOH</td>
<td>0.67</td>
<td>1.67</td>
<td>This work</td>
</tr>
<tr>
<td>C7: 3-methyladipic acid + 3,3-dimethylglutaric acid + diethylmalonic acid</td>
<td>C7H12O4</td>
<td>COOH</td>
<td>0.57</td>
<td>1.71</td>
<td>This work</td>
</tr>
</tbody>
</table>

organic coated aerosol particles is strongly influenced by particle morphology (Thornton and Abbatt, 2005; Anttila et al., 2006; Cosman and Bertram, 2008; Cosman et al., 2008). Significant reduction in reactive uptake coefficient with a factor of 2–42 has been observed with dependence on organic components on aqueous sulfuric acid solutions. Assuming that the aqueous AS phase is surrounded by the organic-rich phase, the N2O5 hydrolysis may be suppressed effectively.

4 Conclusions

We investigated phase transitions of micrometer-sized particles consisting of multicomponent mixtures of AS and C5, C6, and C7 dicarboxylic acids having O:C of 0.80, 0.67, and 0.57, respectively, during humidity cycles. We used optical microscopy to follow phase changes and Raman spectroscopy to infer chemical composition. Micrometer-sized particles of C5/AS/H2O, C6/AS/H2O, and C7/AS/H2O showed significantly different phase transition behavior during RH cycles. While the C5/AS/H2O particles showed no LLPS with OIR = 2:1, 1:1, and 1:4, the C6/AS/H2O and C7/AS/H2O particles exhibit LLPS upon drying at RH between 50 and 85 % and at ∼ 90 %, respectively, depending on the OIR. Phase equilibria calculations using AIOMFAC are for the most part in good agreement with the experimentally determined phase diagrams. From the investigated systems and literature data, we suggest that LLPS commonly occurs
in atmospheric aerosol particles with O:C < 0.7. This is in excellent agreement with a very recent study by Bertram et al. (2011). Further investigations have to show how sensitive this limit is to the specific chemical composition of the organic and inorganic aerosol fractions, temperature, and the complexity of the organic fraction in aerosols.

Core-shell structure and partially engulfed configurations were observed in the particles on a hydrophobically coated substrate and therefore such morphologies might very likely be present in the troposphere. A core-shell configuration will have consequences for heterogeneous chemistry and hygroscopicity because the aqueous inorganic-rich phase will be totally enclosed by a probably highly viscous organic coating with low diffusivity for reactants and water. Definitive conclusions whether a core-shell or a partially engulfed morphology is prevalent in airborne particles is not possible from the present study because all experiments have been performed on a hydrophobically coated substrate, which might influence surface forces and the preferred morphology. Supplementary experiments should therefore be carried out on levitated particles in an electrodynamic balance or optical tweezers. Since also levitated particles are subject to additional forces that are not present in atmospheric aerosols, these measurements should be complemented by calculations of the preferred morphology using accurate surface and interfacial tension data of the investigated mixtures.

Appendix A

Settling velocity

The settling velocity (\(v\)) of an AS-rich inclusion within a C6-rich fluid is given by Stokes’ law:

\[
v = \frac{2(\rho_i - \rho_f)gr^2}{9\mu}
\]  
(A1)

where \(\rho_i\) and \(\rho_f\) are the densities of the inclusion and the fluid, respectively, \(g\) the gravitational acceleration, \(r\) the radius of the inclusion and \(\mu\) the viscosity of the fluid. At \(a_w = 0.7\), AS solution density is calculated as 1.30 g cm\(^{-3}\) at 20°C using the parameterization by Tang and Munkelwitz (1994). We estimate the density of the C6-rich phase at 20°C as 1.20–1.23 g cm\(^{-3}\) by extrapolating the densities of solutions of the structurally similar glutaric acid (Ben-Hamo et al., 2007) to estimated solute mass fractions of 0.75–0.80 at \(a_w = 0.7\). Hence, we expect the aqueous AS phase to be of higher density than the organic-rich one.

The viscosity of aqueous adipic acid, the straight-chain C6 dicarboxylic acid, has only been measured for dilute solutions (Chmielewska and Bald, 2008). In the range where data is available, densities of aqueous solutions of adipic, glutaric, and citric acid show all similar concentration dependences (Laguerie et al., 1976; Chmielewska and Bald, 2008). To estimate the viscosity of the C6-rich phase, we therefore extrapolate the citric acid values to concentrations corresponding to solute mass fractions of 0.75–0.80, yielding viscosities of 100–500 mPa s. Applying these values, settling of 10 and 20 µm diameter AS-rich phase inclusions should occur at a rate of 0.5–3.3 and 2–13 µm min\(^{-1}\), respectively. We therefore expect that the AS-rich inclusions settle to the bottom of the droplet driven by gravity within the timescale of the experiment, which is in accordance with the observed morphology of the Raman scans.

Appendix B

Estimation of surface and interfacial tensions

Surface tensions of aqueous solutions of the straight chain dicarboxylic acids have been measured by several groups (e.g. Shulman et al., 1996; Varga et al., 2007). A decrease of surface tension is observed with increasing solute concentration. For 1–2 wt % aqueous adipic acid (C6) solutions surface tensions of ~68 mN m\(^{-1}\) (pure water case: ~72 mN m\(^{-1}\)) and for 30–40 % aqueous glutaric acid (C5) solutions a reduction of 10–20 % has been reported (Shulman et al., 1996; Varga et al., 2007; Booth et al., 2009). Topping et al. (2007) give values for adipic acid surface tension in the supercooled liquid state of 45.5 mN m\(^{-1}\) and 52.4 mN m\(^{-1}\) depending on estimation method. The organic-rich phase in C6/AS/H\(_2\)O droplets is highly concentrated and we expect it to exhibit similar surface tensions as the supercooled liquid state of adipic acid, i.e. 45–52 mN m\(^{-1}\). Surface tension in AS solutions is increased by 20 % at AS saturation (\(a_w = 0.8\) at 25°C) compared with pure water and is expected to increase even more in the supersaturated regime (Lewis, 2006). However, in the presence of small amounts of adipic acid (~0.5 wt %), this trend is reversed and surface tension decreases with increasing AS concentration (Booth et al., 2009). A solution containing 0.0007 mole fraction adipic acid and 0.009 mole fraction AS exhibited a surface tension of 67.1 mN m\(^{-1}\). This value might quite well represent the surface tension of the AS-rich phase in the C6/AS/H\(_2\)O droplets.

Interfacial tensions of binary aqueous/organic two-phase systems typically vary between 6–52 mN m\(^{-1}\) (Backes et al., 1990). On the other hand, aqueous two-phase systems consisting of immiscible aqueous phases of incompatible polymers, surfactants or salts exhibit very low interfacial tensions (<1 mN m\(^{-1}\)) (Giraldo-Zuniga et al., 2006; Nan et al., 2006; Spyropoulos et al., 2008).

Young’s equation relates the interfacial forces (\(\sigma\)) to the contact angle (\(\theta\)) that a droplet (d) in a fluid (f) makes with the surface (s):

\[
\sigma_{s/d} + \sigma_{d/f} \cos \theta = \sigma_{s/f}.
\]  
(B1)
The fluid (f) is usually, but not necessarily, the vapor phase. Using this relation, the contact angle of the minor-phase inclusions on the substrate can be roughly estimated. In a first step, the interfacial tensions between the substrate and the C6-rich phase $\sigma_{\text{org}/s}$ and the substrate and the AS-rich phase $\sigma_{\text{AS}/s}$ can be estimated using the following values for the surface tensions of the substrate $\sigma_s = 25.3 \text{ mN m}^{-1}$ (Frank and Belfort, 2001), the C6-rich phase $\sigma_{\text{org}} = 45–52 \text{ mN m}^{-1}$, and the AS-rich phase $\sigma_{\text{AS}} = 67.1 \text{ mN m}^{-1}$, and contact angles for saturated C6 and 40 wt% AS droplets of 76.7 $^\circ$ and 94.0 $^\circ$, respectively. This yields $\sigma_{\text{org}/s} = 17.7–18.7^\circ$ and $\sigma_{\text{AS}/s} = 25.3^\circ$. Using these numbers and lower and upper limits of interfacial tension $\sigma_{\text{org}/\text{AS}}$ of 15 and 44 mN m$^{-1}$, respectively, yields contact angles of 105–140 $^\circ$.

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