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Hygroscopic properties of aminium sulfate aerosols

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Received: 27 October 2016 – Discussion started: 1 November 2016
Revised: 10 February 2017 – Accepted: 9 March 2017 – Published: 31 March 2017

Abstract. Alkylaminium sulfates originate from the neutralisation reaction between short-chained amines and sulfuric acid and have been detected in atmospheric aerosol particles. Their physicochemical behaviour is less well characterised than their inorganic equivalent, ammonium sulfate, even though they play a role in atmospheric processes such as the nucleation and growth of new particles and cloud droplet formation. In this work, a comparative evaporation kinetics experimental technique using a cylindrical electrodynamic balance is applied to determine the hygroscopic properties of six short-chained alkylaminium sulfates, specifically mono-, di-, and tri-methylaminium sulfate and mono-, di-, and tri-ethyl aminium sulfate. This approach allows for the retrieval of a water-activity-dependent growth curve in less than 10 s, avoiding the uncertainties that can arise from the volatilisation of semi-volatile components. Measurements are made on particles > 5 µm in radius, avoiding the need to correct equilibrium measurements for droplet-surface curvature with assumed values of the droplet-surface tension. Variations in equilibrium solution droplet composition with varying water activity are reported over the range 0.5 to > 0.98, along with accurate parameterisations of solution density and refractive index. The uncertainties in water activities associated with the hygroscopicity measurements are typically < ±0.2 % at water activities > 0.9 and ~ ±1 % below 0.9, with maximum uncertainties in diameter growth factors of ±0.7 %. Comparison with previously reported measurements show deviation across the entire water activity range.

1 Introduction

Quantifying the response of aerosol particles to variations in relative humidity (RH) in the atmosphere is important for predicting the impact of aerosols on climate through both direct and indirect radiative forcings (Albrecht, 1989; Kanakidou et al., 2005; Kolb et al., 2010), for understanding the mechanisms of heterogeneous chemistry (Dennis-Smither et al., 2014; Lee et al., 2012), and the partitioning of products in the condensed phase (Dusek et al., 2006; Topping et al., 2013a), and for inferring the deposition and loss mechanism of aerosols on surfaces and on inhalation into the respiratory tract (Broday and Georgopoulos, 2001; Haddrell et al., 2015). With an increase in RH, solution droplets must absorb water to maintain an equilibrium balance of water between the gas and condensed phases, leading to the dilution of solutes, increase in mass of condensed phase water and hygroscopic growth in particle size. The hygroscopic response of a wide range of inorganic and organic solutes of varying solubility, often present as complex mixtures, must be quantified. Thus, the influence of interactions between multiple solutes of varying charge, acidity, and hydrophilicity must be understood through predictions of activity coefficients that reflect the departure of solution composition from ideality (Dutcher et al., 2013; Shiraiwa et al., 2013).

Measurements of the hygroscopic growth of ambient aerosol are routinely made; the extent of growth at high RH or the critical supersaturation for activation as cloud condensation nuclei are often represented by a single parameter, such as the κ parameter in κ-Köhler theory (Petters and Kreidenweis, 2007). Resorting to such a reduced parameter model is a consequence of the complex chemical composition of ambient aerosol and the intractability of providing ac-
Accurate predictions based on a detailed chemical speciation. However, laboratory measurements are essential for exploring the complex details often obscured in measurements of ambient aerosol. These include the roles of pH, liquid–liquid phase separation, and mixing state (Topping et al., 2013c; You et al., 2014) and the co-condensation of semi-volatile organic species with varying RH (Topping et al., 2013b).

When considered alongside measurements of hygroscopic response and change in particle composition with RH for aerosol of well-known chemical composition, detailed predictive tools can be benchmarked and improved, providing a crucial framework for interpreting field measurements. We present here measurements of the hygroscopic response of a class of compounds, aminium sulfates, with the objective of providing accurate data to refine predictive tools of their equilibrium hygroscopic response.

Low molecular weight amines are mostly emitted into the atmosphere as gaseous compounds by a number of different sources, both anthropogenic (animal husbandry, food processing and cooking, combustion, pesticides) and natural (oceans, biomass burning, vegetation; Ge et al., 2011a). These chemical species are basic (with $pK_a$ values from 9.8 to 10.84; Lide, 2005), highly water soluble, and have high pure liquid vapour pressures (up to hundreds of kPa) and partitioning due to a variety of chemical processes (Ge et al., 2011b): direct solubilisation, oxidation reactions that lead to the formation of secondary organic aerosols, acid–base reactions similar to those of ammonia (Seinfeld and Pandis, 2006) with both inorganic (sulfuric, nitric and hydrochloric acids) and organic acids (Lavi et al., 2015; Liu et al., 2012), and displacement reactions of ammonium cations with aminium cations (Bzdek et al., 2010; Qiu and Zhang, 2013). The product of these last two types of chemical reactions and the salts formed by protonated amines following new particle formation are aminium sulfates, which result from the neutralisation of sulfuric acid and short-chained alkylamines:

$$H_2SO_4 + 2R_3N(R_3NH)_2SO_4,$$

where $R$ can be either $-CH_3$, $-CH_2CH_3$, or $-H$. The formation of aminium sulfates and other similar aminium salts increases the solubility of short-chained alkylamines from 7–45 wt % for the unprotonated form (calculation from data in Ge et al., 2011b) to 84–91 wt % for five aminium sulfates in Clegg et al. (2013). This, correspondingly, increases their partitioning from the gas to the condensed phase (Barsanti et al., 2009; Yli-Juuti et al., 2013). However, the physicochemical properties of aminium sulfates are much less well characterised than their inorganic counterpart, $(NH_4)_2SO_4$, even though they can play a fundamental role in the nucleation and growth of new particles (DePalma et al., 2012; Loukonen et al., 2010; Wang et al., 2010b) and in cloud formation (Lavi et al., 2013). Some recent work has attempted to fill this gap, reporting the densities (Clegg et al., 2013; Lavi et al., 2013; Qiu and Zhang, 2012), CCN activity, optical properties (Lavi et al., 2013), hygroscopicity and phase transitions of aminium sulfates (Chu et al., 2015; Clegg et al., 2013; Lavi et al., 2013; Qiu and Zhang, 2012; Sauerwein et al., 2015), specifically for the methyl and ethylaminium sulfates (or subsets of these six compounds). In this work, we report measurements of the hygroscopic behaviour of the three methylaminium sulfates (methylaminium sulfate, MMAS; dimethylaminium sulfate, DMAS; trimethylaminium sulfate, TMAS) and the three ethylaminium sulfates (ethylaminium sulfate, MEAS; diethylaminium sulfate,
DEAS; triethylaminium sulfate, TEAS) from water evaporation experiments made using a comparative kinetic electrodynamic balance (CK-EDB; Rovelli et al., 2016). In Sect. 2, we review the experimental details and procedures for performing hygroscopic growth measurements using the CK-EDB. In Sect. 3, we present measurements of the hygroscopic growth of the six ammonium sulfate salts, comparing our measurements with literature values and considering the accuracy and reproducibility of measurements with the CK-EDB.

2 Experimental

We first describe the CK-EDB technique and the data analysis before describing the method for preparing aqueous starting solutions of the ammonium sulfates.

2.1 Hygroscopic properties from comparative kinetics measurements in a CK-EDB

Electrodynamic trapping of single charged droplets has been used to investigate various properties of confined particles, including optical properties (Barnes et al., 1997), vapour pressures of low-volatility compounds (Pope et al., 2010), hygroscopic properties of atmospherically relevant aqueous solutions (Choi and Chan, 2002; Chu et al., 2015; Peng et al., 2001; Rickards et al., 2013) and of pharmaceutical aerosols (Haddrell et al., 2013; Peng et al., 2000), and evaporation dynamics of aqueous droplets (Heinisich et al., 2009; Shulman et al., 1997; Zobrist et al., 2011). The experimental set-up used in this work has been presented in previous publications (Davies, 2014; Davies et al., 2012a, b; Haddrell et al., 2012; Miles et al., 2012) and a schematic of it can be found in the Supplement (Fig. S1). In particular, the approach used to quantify the hygroscopic properties of single confined particles from comparative kinetics measurements using a CK-EDB has also been previously discussed (Davies et al., 2013) and its application over a wide-solution water-activity range from 0.5 to > 0.99 has been extensively validated (Rovelli et al., 2016).

In a CK-EDB single charged droplets from two solutions with known chemical composition and concentration can be sequentially dispensed on demand by means of two alternatively operated microdispensers and trapped within the electrodynamic field generated by a set of concentric cylindrical electrodes. This electrode geometry guarantees a stable and tight trapping of single droplets within 100 ms from their generation. Single droplets are confined in the trapping chamber within a nitrogen stream with controlled mass flow, temperature, and relative humidity. The gas flow RH is modified by mixing different ratios of a humidified and a dry nitrogen flow and is inferred from the evaporation kinetics of probe droplets, as described below. The temperature within the trapping chamber is controlled by a circulating 1 : 1 volume mixture of water and ethylene glycol, through the lid and the bottom of the chamber. Temperature and gas phase RH ranges that are accessible with this experimental set-up are −25 to 50 °C and 0 to 99 %, respectively. All the comparative evaporation kinetics measurements presented here were performed at 20 °C and at gas phase RH values between ∼50–90 %. Trapped droplets are illuminated by laser light (532 nm) and the resulting elastic scattering light pattern is collected every 0.01 s by means of a CCD camera and used to keep track of changes in size of a single levitated droplet using the geometrical optics approximation (Glantschnig and Chen, 1981). Note that the variations in refractive index due to water evaporation from each droplet are taken into account for an accurate determination of droplet size, as described in a previous publication (Davies et al., 2012b).

The evaporation rate of water from a droplet containing one or more solutes at a certain RH and temperature depends on the hygroscopic properties of the solution. In comparative kinetics experiments, the evaporation rate of single probe droplets with known composition and known evaporation kinetics (either pure water or a NaCl solution) is compared to that of single sample droplets in order to determine the hygroscopic properties of the solution droplet being studied (Davies et al., 2013; Rovelli et al., 2016). To do so, typical CK-EDB experiment are composed of a series of at least 10 alternating single probe and sample droplets (Fig. 1a).

The retrieval of the hygroscopic properties from the radius (a) vs. time evaporation profiles of a sequence of singly trapped probe and sample droplets is shown schematically in Fig. 1. Different analyses are carried out for probe (Fig. 1b) and sample droplets (Fig. 1c). Either pure water (as shown in Fig. 1a) or NaCl solution droplets with known initial salt concentration can be used as a probe. The evaporation profiles of the probe droplets are compared with simulations obtained from the evaporation/condensation kinetics model by Kulmala et al. (1993). By doing so, the gas phase RH can be inferred, either from fitting the a² vs. time evaporation profile of pure water droplets or from the equilibrated radius of the NaCl solution droplets. Both methods, together with their associated uncertainties and their RH ranges of applicability, have been discussed previously (Davies et al., 2013; Rovelli et al., 2016). RH fluctuations over the run of ten pairs or more of probe and sample droplets are very slight, typically of the order of 0.2 % RH and never exceeding 0.5 % RH. However, it should be noted that slight RH fluctuations are taken into account in our approach: the gas phase RH is monitored before every sample droplet by injecting a probe droplet and data from this probe droplet are directly used in the sample droplet evaporation analysis.

For the analysis of the radius (a) vs. time data of sample droplets, the radius data (Fig. 1c, inset panel i) are converted to droplet mass (inset ii) by using a third-order polynomial parameterisation of density as a function of the solute mass fraction, which is discussed below. The mass flux of water leaving the droplet during evaporation (dm/dt) is then
Figure 1. Schematics representing the retrieval of hygroscopic growth curves from comparative kinetics experiments in a CK-EDB. An experimental sequence of alternating single evaporating probe and sample droplets is collected (a). For each pair of probe and sample droplets, the gas phase RH is inferred from the evaporation kinetics of the probe (b) and this information is used to analyse the corresponding sample droplet hygroscopic properties as indicated in (c).

calculated and, knowing the gas phase RH from the probe droplet analysis, Kulmala’s equations for evaporation kinetics are applied to calculate the temporal variation of water activity ($a_w$) in the droplet (inset iii). In addition, knowing the initial concentration and size of the droplet at generation, the radius of the dry particle ($a_{dry}$) can be estimated and a radial growth factor ($GF_r = a/a_{dry}$) can be calculated for each of the measured radii (inset iv). Results from inset panels (iii) and (iv) are then combined and typical $GF_r$ vs. $a_w$ growth curves are obtained. The key thermodynamic quantities that describe the hygroscopic properties of the tested solution (moles of water per mole of solute in solution, $n_{water}/n_{solute}$; osmotic coefficients, $\phi_{st}$) can be calculated if the densities of the aqueous solutions are also known (see below). Details of the treatment of the experimental uncertainties and their influence on each of the computed quantities are described in the Supplement (Table S1). When error bars are not shown in the figures in the following sections, the reader can assume that they are smaller than the size of the corresponding data point.

A previous study (Cai et al., 2016) showed that the molar refraction mixing rule, together with a third-order polynomial parameterisation of density as a function of the square-rooted mass fraction of solute (mfs), represent the best approach to predicting refractive indices ($m$) and densities ($\rho$) of solutions of organic compounds for which bulk data of such quantities are available for solute mass fractions up to at least 0.4. In this work the densities of at least 10 solutions with different concentrations for each aminium sulfate were measured with a density metre (Densito 30PX, Mettler Toledo; accuracy of ±0.001 g cm$^{-3}$, calibrated with pure water before each use). Densities were measured at ambient temperature, which varies in the laboratory between 293 and 295 K; temperatures were always registered together with the measured density values. In addition, refractive indices of the same solutions were measured at 589 nm by means of a refractometer (Palm Abbe II, Misco, precision of ±0.0001, calibration with pure water before each use). The measured density and $m$ values for each aminium sulfate solution are provided in the Supplement (Table S2), together with the third-order polynomial and the molar refraction mixing rule fittings for each compound (Table S3 and Fig. S2). A brief description of the molar refraction mixing rule application is also provided in the Supplement. These data have been presented previously and discussed by Cai et al. (2016) along with measurements from a large number of organic aqueous solutions.

2.2 Preparation of the solutions

Aminium sulfate stock solutions were prepared by the neutralisation of solutions of each of the six amines with aqueous sulfuric acid. The commercial amines stock solutions (Sigma Aldrich, MMA, ~ 40 wt %; DMA, ~ 40 wt %; TMA,
ice bath (0 °C) solutions with H2SO4 measurements. During both the titration of the amine stock concentration for the CK-EDB comparative kinetics measurement was monitored for the whole duration of the reaction both solutions around 40 wt %. The pH of the solution mixture was determined to be 81.6 wt % (0.52 wt % standard deviation) by the neutralisation reaction and to avoid any possible slow and dropwise, in order to dissipate the heat generated by the neutralisation reaction and to minimise volatilisation of the amine. In addition, the HCl (and later the H2SO4) used for the standardisation of the commercial amine solutions were titrated with Na2CO3 (≥ 99.5 %, Alfa Aesar), which was first dried at 225 °C for 3 h before weighing to make sure that no water was adsorbed on it. Three repetitions were performed in these cases.

For the preparation of the amonium sulfates stock solutions, stoichiometric amounts of the standardised H2SO4 and amine solutions were mixed with an initial concentration of both solutions around 40 wt %. The pH of the solution mixture was monitored for the whole duration of the reaction to ensure that all of the amine in the solution had reacted. The concentrated stock solutions of the salts that result from this procedure were subsequently diluted down to a weight percentage of ≈ 5 %, in order to obtain a suitable starting concentration for the CK-EDB comparative kinetics measurements. During both the titration of the amine stock solution with HCl and the preparation of the ammonium sulfates solutions with H2SO4, the amine solution was kept in an ice bath (0 °C), and the addition of the acid was performed slowly and dropwise, in order to dissipate the heat generated by the neutralisation reaction and to avoid any possible amine volatilisation. Phase separation was observed when titrating the TEA commercial solution and its concentration was determined to be 81.6 wt % (0.52 wt % standard deviation over three repetitions), which is considerably lower than the ≥ 99 wt % concentration value given by the manufacturer. This is possibly due to the much lower solubility of TEA (0.7 mol kg⁻¹) than the other amines (Ge et al., 2011b), leading to incomplete solvation of the amine in water and an inaccurate measurement of pH during the titration. Thus, we instead assumed that the commercial TEA solution was a 99 wt % concentration and note that the results for the TEA system should be interpreted with some caution. This experimental procedure ensured that the concentrations of the reagents were well known and, consequently, that the concentrations of the stock solutions prepared for amonium sulfates were similarly well known. The uncertainties in the commercial solution concentrations of the reagents, as determined from the repeated titrations, were taken into account for the calculation of the overall experimental uncertainties indicated in the Supplement (Table S1).

To validate the procedure for solution preparation described above, the steps were carried out for the preparation of (NH4)2SO4 from the reaction of ammonia and sulfuric acid. The resulting salt solution was then used in comparative kinetics measurements and the hygroscopicity of the ammonium sulfate from reaction was compared with calculations from the Extended Aerosol Inorganics Model (E-AIM; Wexler and Clegg, 2002). Figure 2 shows the hygroscopic properties of (NH4)2SO4 droplets prepared in this way from the direct reaction of ammonia and sulfuric acid, reporting values of nwaterr/n sulfate, vs. am (panel a) and osmotic coefficients (φst) plotted against the square root of the sulfate molality (m(SO42⁻)0.5, panel b). Osmotic coefficients are useful parameters to represent the deviation of a solution from an ideal behaviour and they are defined as indicated in Eq. (2):

\[
\phi_{\text{st}} = -\frac{\ln(a_w)}{M_w 3m/1000},
\]

where \(M_w\) is the molecular weight of water, \(m\) is the molality of the solute and 3 is the stoichiometric number of ions in the salt. The value of \(\phi_{\text{st}}\) tends to 1.0 in the limit of an infinitely dilute solution, in accordance with the Debye–Hückel limiting law (Robinson and Stokes, 1970).

In a previous publication (Rovelli et al., 2016) we showed that it is possible to achieve very good agreement with predictions from the E-AIM model for well-characterised inorganic compounds with the CK-EDB experimental technique within an uncertainty in \(a_w\) of ±0.002. The plots in Fig. 2 show the averaged data obtained from two datasets of 10 droplets of aqueous (NH4)2SO4 (black and open circles). In Fig. 2b the effect of a ±0.002 error on \(a_w\) on the modelled osmotic coefficients values is shown with dashed lines. The osmotic coefficients agree well with calculations from the E-AIM model and lie within the envelope associated with this previous estimate of typical experimental uncertainty for the CK-EDB technique. This demonstrates that the volatility of ammonia is not a significant problem when the neutralisation reaction with H2SO4 is performed. In addition, since the vapour pressure of ammonia (1956 kPa at 298.15 K; Lide, 2005) is even higher than the vapour pressure of the most volatile of the six considered amines (methylamine, 336 kPa at 298.15 K; Ge et al., 2011b), it is likely that there is also no evaporative loss of amines from solution during the preparation of the amonium sulfate solutions. This result confirms that the estimated concentrations of the (NH4)2SO4 solutions prepared from NH3 and H2SO4 are accurate, and that the preparation method is reliable for both (NH4)2SO4 and all the six amonium sulfates.

3 Results and discussion

We first report our measurements of the hygroscopic response of the sequence of six aminium salts before comparing our results with previous studies and assessing the accuracy and reproducibility of our data.
3.1 Hygroscopic properties of aminium sulfate droplets

The hygroscopic properties of the series of six aminium sulfates were characterised by means of comparative kinetics measurements and using the density and refractive index parameterisations discussed for the CK-EDB data treatment, as described in Sect. 2.1. First, the radial growth curves are shown in Fig. 3a. Considering the compound with the lowest molecular weight first, the hygroscopic behaviour of MMAS is the most similar to ammonium sulfate, in terms of GF<sub>r</sub>. Continuing in the methylaminium sulfates series, a slight decrease in GF<sub>r</sub> is observed in the high water activity region (a<sub>ω</sub> > 0.8) for DMAS and TMAS, whereas at lower water activities, the hygroscopic properties of these compounds converge to (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> within the uncertainties of the measurements as the amount of water in the particles decreases. With respect to the ethylaminium sulfates series, a more evident decreasing trend in the radial growth curve is observed with an increasing number of C atoms in the cation (MEAS > DEAS > TEAS), once again especially in the upper part of the curves. If the mono-, di-, and tri- pairs within the two different series are compared, the methyl compound always presents higher values of radial growth factor than its equivalent in the ethylaminium sulfates series. As indicated in Sect. 2.2, it is worth reiterating that the results presented here and below for TEAS need to be interpreted cautiously because of the uncertainty of the TEA stock solution. However, the trends that have been observed for TEAS when compared to the other five aminium sulfate systems seem to be completely plausible, and this may indicate that the assumed initial TEA concentration is reasonable.

For the calculation of the dry radius reference state in the denominator of GF<sub>r</sub>, the pure melt density is used (Sect. 2.1); if the pure solid density values were known and used, one could expect that the calculated radial growth curve would be slightly higher, a consequence of the solid densities having higher values than the melt density (Clegg et al., 2013). A comparison of the pure melt densities (ρ<sub>melt</sub>) from our work and from Clegg et al. (2013) and solid densities (ρ<sub>solid</sub>) estimated by Qiu and Zhang (2012) is provided in Table S4, with our data previously published in Cai et al. (2016). As an example, if a hypothetical increase of 5 % from ρ<sub>melt</sub> to ρ<sub>solid</sub> as a rough estimate is considered for DMAS, the obtained GF<sub>r</sub> curve would increase by less than 1 % (less than 0.01 in GF<sub>r</sub>); this would not affect the trends shown in Fig. 3a, although it would marginally change the relative position of the aminium sulfates curves to that of ammonium sulfate.

The same datasets shown in Fig. 3a have been converted to n<sub>water</sub>/n<sub>solute</sub> vs. a<sub>ω</sub> and are shown in Fig. 3b. Note that the y axis is displayed on a logarithmic scale for clarity. Different trends can be recognised: the primary factor influencing how many moles of water are absorbed per mole of solute is the number of alkyl groups in the cation and not the length of the carbon chains (i.e. mono- vs. di- vs. tri-methyl or ethyl). In fact, when the n<sub>water</sub>/n<sub>solute</sub> curves (Fig. 3b) are compared with the hygroscopic behaviour plotted as GF<sub>r</sub> (Fig. 3a), more significant differences are apparent in Fig. 3b within each series (e.g. among MMAS, DMAS, and TMAS) rather than between analogous compounds belonging to the two series (e.g. between TMAS and TEAS), even though the ethylaminium sulfate in a pair always has slightly higher values of n<sub>water</sub>/n<sub>solute</sub> (i.e. MMAS ≈ MEAS < DMAS ≈ DEAS < TMAS ≈ TEAS).

The seemingly different trends found for the hygroscopic properties of aminium sulfates when reported either as GF<sub>r</sub> or n<sub>water</sub>/n<sub>solute</sub> as a function of a<sub>ω</sub> originate in the different physicochemical properties that each quantity is dependent on. When GF<sub>r</sub> is calculated, the density and molecular

Figure 2. Measured n<sub>water</sub>/n<sub>solute</sub> vs. a<sub>ω</sub> (a) and osmotic coefficients (φ<sub>s</sub>) vs. the square root of sulfate molality (m(SO<sub>4</sub>)<sup>0.5</sup>) (b) of ammonium sulfate solution obtained from the reaction between NH<sub>3</sub> and H<sub>2</sub>SO<sub>4</sub>. Symbols: black dots – 0.03 mfs of (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> in the initial solution, 88.5 % RH in the gas phase; open circles – 0.004 mfs of (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> in the initial solution, 90% RH in the gas phase; solid lines – calculations from E-AIM model; dashed lines – uncertainty on the osmotic coefficients corresponding to an error in a<sub>ω</sub> of ±0.002.

Figure 3. Aminium sulfate GF\(_r\) vs. \(a_w\) (a) and \(n_{\text{water}}/n_{\text{solute}}\) vs. \(a_w\) (b) hygroscopic growth plots from CK-EDB experiments. Symbols: • – MMAS; ■ – DMAS; ♦ – TMAS; ○ – MEAS; □ – DEAS; ◆ – TEAS; line – E-AIM model calculation for (NH\(_4\))\(_2\)SO\(_4\). Note that only E-AIM predictions for (NH\(_4\))\(_2\)SO\(_4\) are reported in this and subsequent figures with the CK-EDB measurements and the E-AIM model compared in Fig. 2.

Figure 4. Moles of salt \((n_{\text{salt}})\) in a 1 µm radius solid particle as a function of the molecular weight \((M_w)\) of each aminium sulfate.
means of an HTDMA (Hygroscopicity Tandem Differential Mobility Analyser). They inferred hygroscopic growth factors from the ratio of the mobility diameters measured at a variable RH (up to 90 %) and at an RH of ~ 12 %. A monotonic increase in the particle size with RH for each compound was observed and no deliquesence/efflorescence behaviour was observed. If their growth curves are compared with the GF values shown in Fig. 3a (Fig. S3), a systematically smaller growth factor is reported from the HTDMA measurements. The largest deviations are for TEAS (− 16 % on average in the $a_w$ range where the two datasets overlap, from ~ 0.5 to 0.9) and for TMAS (− 13 %); a deviation of − 10 % is observed for DEAS and DMAS and of − 7 % for MMAS. These discrepancies are most likely due to the presence of some residual water at the conditions at which the reference diameter was measured in the HTDMA experiments (RH ~ 12 %), which would result in an overestimated reference “dry” size and in an underestimated growth factor values. This explanation is supported by the studies of Chan and Chan (2012), who reported the presence of water for some aminium sulfates even at an RH of ~ 3 %. In addition, the possible volatilisation of the amine during the drying step in the HTDMA would alter the chemical composition of the particles (Chan and Chan, 2012, 2013), not only artificially reducing the apparent dry size recorded but also shifting the dry particle towards an aminium bisulfate composition (1 : 1 molar ratio of sulfuric acid to amine), which has been shown to be less hygroscopic than its sulfate counterpart (Sauerwein et al., 2015).

Consistent with the above discussion, the two compounds with the biggest deviation between the GF values reported here and by Qiu and Zhang (TMAS and TEAS) are those that were found by Chan and Chan (2012) to be affected by the largest evaporative losses of amine. They reported studies of ammonium displacement reactions by alkylamines by levitating single droplets in an EDB and evaluating the changes in the recorded Raman spectra during an experiment. After TEAS solution droplets were levitated at RH < 3 % for more than 5 h, TEAS was found to have converted to TEA bisulfate almost completely, indicating that half of the amine in the initial solution droplet had evaporated. Similar behaviour was observed for TMAS, while DMAS and DEAS showed a smaller degree of evaporation of the amine (~ 25 and ~ 5 %, respectively); MMAS and MEAS did not show any relevant evaporation over the experimental timescales. These results support the hypothesis of possible evaporation of the amine from the drying solution droplets during HTDMA experiments, and at the same time represent an interesting comparison to evaluate the timescales over which the evaporative loss of amine is significant. The hygroscopic growth measurement of each droplet of aminium sulfate solution in CK-EDB studies extends over 30 s at the longest; this allows the effective decoupling of the fast water evaporation and the slow amine evaporation, since these two processes occur over different timescales. In addition, the hygroscopicity measurements in the present work were carried out at RHs in excess of 50 % at all times; for this reason, the evaporation of the amines is necessarily smaller than those measured by Chan and Chan (2012).

Clegg et al. (2013) have converted the size growth curves measured by Qiu and Zhang to $n_{\text{water}}/n_{\text{solute}}$ vs. $a_w$. The growth curves that resulted did not show any discernible trends in the hygroscopicity of aminium sulfates. Indeed, the results were essentially similar to the water uptake of ammonium sulfate in the lower $a_w$ range. At higher $a_w$ above 0.7, the scatter in the data limited the identification of any apparent trend apart from the suggestion that the methylaminium sulfates absorb fewer moles of water per moles of solute than the compounds in the ethyl series. Figure 5 shows the large differences between the water uptake determined from the results of Qiu and Zhang, and that from the work in this study. Because of the uncertainties associated with the HTDMA GF values, it is hard to draw conclusions with comparison to our results.

Sauerwein et al. (2015) have recently reported bulk water activity measurements with an activity metre (Aqua lab Series 3TE) for various amine-to-sulfate ratios and over a concentration range of dissolved electrolyte up to 9 mol kg$^{-1}$ at 25 °C. (Note that TEAS was not considered in their study.) Using a bulk measurement technique for the determination of hygroscopic properties of aminium sulfates has the advantage of limiting possible evaporative losses of the amine during the experiments, especially compared to accumulation-mode aerosol measurements with an HTDMA for which timescales of evaporative loss/equilibration are assumed very short. As a first comparison, the hygroscopic growth curves previously shown in Fig. 3b in terms of $n_{\text{water}}/n_{\text{solute}}$ vs. $a_w$ are compared with results from Sauerwein et al. (2015; Fig. 6a in their paper) in Fig. 5. The hygroscopic growth curves determined from the two different experimental methods differ marginally, but these discrepancies are considerably smaller than the differences between the results of both studies – CK-EDB and bulk water activity measurements – and HTDMA data from Clegg et al. (2013).

A comparison of the two datasets represented in terms of osmotic coefficients (Eq. 2) can provide further insight; see Fig. 6. Notably, the measurements reported here extend both more dilute and more concentrated solutions and include measurements for TEAS. If the errors associated with each dataset are considered, the two sets of measurements overlap not only over some of the range in $m(SO_4^{2-})^{0.5}$, especially in the low molality limit (high water activity region) for MMAS, TMAS, MEAS, and DEAS ($m(SO_4^{2-})^{0.5} < 1$, approximately), but also in the region $m(SO_4^{2-})^{0.5} > 2$ for DMAS and TMAS. A comparison with the uncertainty in the osmotic coefficients that arises from an uncertainty of ±0.002 in water activity (same as in Fig. 2b) indicates that the observed discrepancies are larger in magnitude than this typical experimental uncertainty of our technique, except for
very low sulfate molality values (below \(~0.8\)). However, despite the discrepancies that do exist, results from the two different experimental methods show the same qualitative hygroscopicity trends over the methyl and ethylaminium sulfates series. For further comparison, the results of Clegg et al. (2013; from the HTDMA data) were converted to stochiometric osmotic coefficients and are plotted in Fig. 6. These values are broadly comparable for all of the compounds, and close to those for ammonium sulfate as noted earlier. They do not agree with either the results of the experiments reported here, or the bulk measurements of Sauerwein et al. (2015).

We now consider the differences that exist between the data of Sauerwein et al. (2015) and our measurements. It is worth stating here that the evaporation kinetics measurements at the core of the CK-EDB approach have been validated in a previous publication (Rovelli et al., 2016) by means of the determination of the hygroscopic properties of well-characterised inorganic compounds and their mixtures. In addition, we have performed sensitivity tests to evaluate possible effects of random experimental errors associated with the proposed experimental method. The reproducibility of our measurements and the uncertainties associated with the treatment of densities are discussed below in Sect. 3.3 and 3.4, where they are shown not to affect significantly the determined hygroscopic properties of a compound.

If the differences between our measurements and the results of Sauerwein et al. (2015) were caused by partial volatilisation of the amine from the droplets evaporating in the CK-EDB, the observed bias between the two datasets would be reversed; i.e. an underestimation of \(n_{\text{water}}/n_{\text{solute}}\) and of osmotic coefficients would be expected, similar to what has been discussed in the case of the HTDMA size-based measurements of Qiu and Zhang (2012) when converted to a molar basis by Clegg et al. (2013). Another potentially significant source of error could be inaccuracies in the starting concentrations of the amminium sulfate solutions. In this respect, the solution preparation and measurement method described in Sect. 2.2 has been validated through the reaction of ammonia and sulfuric acid and the measurements of the hygroscopic properties of the obtained \((\text{NH}_4)_2\text{SO}_4\). If the volatilisation of \(\text{NH}_3\) during the neutralisation reaction was not an issue in preparing the solution of \((\text{NH}_4)_2\text{SO}_4\), it is unlikely to present a problem for the alkylamine solutions, with all alkylamines characterised by lower vapour pressures than ammonia (Ge et al., 2011b). Furthermore, in order to achieve complete agreement with the data from Sauerwein et al. (2015), the amminium sulfates solutions should be 5–10 % more concentrated than calculated from the titration of the commercial stock solutions and from the amount of the reagents. Not only does this percentage seem unrealistically high, but also the solute concentrations would be overestimated and not underestimated if any volatilisation of the amine occurred during the preparation of the solutions.

Sauerwein et al. (2015) provided a framework for the estimation of the water content in mixtures (in this case ternary mixtures of \(\text{H}_2\text{O–H}_2\text{SO}_4–\text{amine}\)) at any molar ratio of solutes. This allows for a calculation of the water content (in terms of kilograms of water per mole of solute, or molality) and the stochiometric osmotic coefficients (Eq. 2) for the relevant amminium:sulfate ratio of 1:1 (bisulfate) and 2:1 (sulfate). However, it should be noted that unlike the study presented here, measurements were not explicitly made for the 2:1 molar ratio aqueous amminium sulfate solution. Instead, Sauerwein et al. used a modified Zdanovskii–Stokes–Robinson expression (ZSR; Eq. 3 of Sauerwein et al.) to represent the water uptake of the different mixtures of aqueous amminium sulfates and \(\text{H}_2\text{SO}_4\); these fits are shown in Fig. 7 along with the actual measured points. \(x_{\text{salt}}\) can be considered as the degree of neutralisation of sulfuric acid. \(x_{\text{salt}}=0\)

\[
\begin{align*}
\text{solute} & = \text{degree of neutralisation of sulfuric acid,} \\
x_{\text{salt}} & = 0
\end{align*}
\]

Figure 5. \(n_{\text{water}}/n_{\text{solute}}\) vs. \(a_w\) plots for the six amminium sulfates. Symbols: black circles – CK-EDB comparative kinetics measurements; open circles – Sauerwein et al. (2015); grey dots – Clegg et al. (2013) (based on measurements from Qiu and Zhang, 2012); line – \((\text{NH}_4)_2\text{SO}_4\) calculation from E-AIM model.
Figure 6. Osmotic coefficients ($\phi_{st}$) vs. square root of sulfate mass fraction ($m(SO_2^{2-})^{0.5}$) for the six aminium sulfates. Symbols: black circles – CK-EDB comparative kinetics measurements; open circles – Sauerwein et al. (2015); grey circles – Clegg et al. (2013) (based on measurements from Qiu and Zhang, 2012); solid line – ($NH_4)_2SO_4$ calculation from E-AIM model; dashed lines – uncertainty of the osmotic coefficients for ($NH_4)_2SO_4$ corresponding to an error in $a_w$ of ±0.002, included to provide a guide as to the level of expected error in the osmotic coefficient with varying molality.

corresponds to pure H$_2$SO$_4$ and the water content in this case is well known (Clegg and Brimblecombe, 1995); the ZSR fit is therefore constrained to this value. A value at $x_{salt} = 1$ corresponds to the completely neutralised aminium sulfate. The results of the fit were used to estimate the water uptake of each solution of aqueous aminium sulfate (i.e. the 2 : 1 molar ratio at $x_{salt} = 1$) and it is these values that are shown in Fig. 6. In order to compare our data with that of Sauerwein et al., values at the same water activities must be compared (0.8 and 0.925 are chosen here) and a linear interpolation between actual measurement points is required to achieve this. Normally, the data from CK-EDB measurement are calculated with $a_w$ steps of 0.01 for $a_w > 0.8$ where hygroscopic growth curves are the steepest, with a spacing of 0.02 for the water activity range ~0.65–0.8 and of 0.03 for $a_w < 0.65$. As in Sauerwein et al. (2015), the uncertainty of the interpolated points is set as the largest uncertainty among the experimental points used for the interpolation. The discrepancies between the estimations from bulk and CK-EDB measurements shown in Fig. 5 are also apparent in Fig. 7 at $x_{salt} = 1$; the amount of water associated with each mole of aminium sulfate is systematically higher from the CK-EDB measurements when compared with the bulk measurements at both $a_w$ with the discrepancy higher at the higher water activity.

In view of the differences between our measurements at $x_{salt} = 1.0$ and the extrapolations from the fits of Sauerwein et al., we include a second fit to the ZSR model used previously but including our own measurements of the water uptake. The fit equations and parameters are provided in the Supplement (Table S6). As well as the fitted values more closely reflecting our measured values for the pure aminium sulfates at $x_{salt}$ equal to unity, the revised fit does not significantly worsen the fit to the measurements of Sauerwein et al. for the acidified mixtures when $x_{salt} < 0.6$. We therefore suggest that the discrepancies seen in terms of $n_{water}/n_{solute}$ (Fig. 5), osmotic coefficients (Fig. 6), and kilograms of water per mole of solute in Fig. 7 can be attributed, in part, to the uncertainties associated with the ZSR fits and interpolation.
necessary to estimate the water uptake of the pure aqueous aminium sulfates from the results of Sauerwein et al. (2015).

Certainly there is no physical reason why a simple relation such as the modified ZSR equation (Eq. 3 of Sauerwein et al.) should exactly represent the relationship between water activity and chemical composition in such systems, especially considering the large and unquantified influence of the sulfate–bisulfate equilibrium on the thermodynamic properties of the solutions.

A possible constant error in measured water activity could be the origin of the differences, but this seems unlikely. Such an offset in water activity results in a large change in the osmotic coefficient at the lowest concentrations for which measurements were made, but a much smaller change at the highest concentrations. This is not consistent with the differences in $\phi_{st}$ shown in Fig. 6. As a final consideration, we now explore the reproducibility of the CK-EDB measurements and the sensitivity of the retrieved hygroscopicity to the chosen treatment for solution density.

3.3 Reproducibility of measurements

A full validation of the retrieval of the hygroscopic properties of single trapped solution droplets from CK-EDB experiments has already been presented in a previous publication (Rovelli et al. 2016), where we demonstrated the accuracy of the approach by reporting hygroscopicity measurements for well-characterised inorganic components. In this section, we evaluate the reproducibility of the solution preparation method (Sect. 2.2) with the aim of demonstrating that the results presented in the previous sections are not affected by any random error associated with our approach for making the sample solutions. Data obtained from three different datasets of evaporating DMAS solution droplets are compared in Fig. 8. The variation in $n_{\text{water}}/n_{\text{solute}}$ with $a_w$ (panel a) and the change in osmotic coefficient with square root of sulfate mass fraction (panel b) are calculated from three different datasets, each arising from measurements with 10 droplets and prepared from different starting stock solution of diethylaminium sulfate, obtained separately one from the other by mixing DEA and $\text{H}_2\text{SO}_4$ on different days. Data from Sauerwein et al. (2015; open circles) and calculations for ammonium sulfate (E-AIM model, line) are shown for comparison. Measurements were made with droplet evaporation into a gas phase RH of $\sim 80\%$. The reproducibility of the data is very satisfactory, giving further evidence that the applied solution preparation procedure coupled to the retrieval of the hygroscopic properties with CK-EDB experiments is reliable. In addition, this consistency in experimental reproducibility strongly suggests that the discrepancies with the data of Sauerwein et al. (2015) do not originate from random errors associated with the CK-EDB experiments.

3.4 Sensitivity to parameterisation of solution density

A knowledge of the solution density as a function of solute concentration is needed to process CK-EDB evaporation radius profiles (Sect. 2.1). The effect of uncertainties in the density parameterisation used is here evaluated for DMAS, in order to estimate how potential errors in the density measurements and uncertainties in the extrapolation of $\rho_{\text{melt}}$ at mfs equal to unity could affect the hygroscopicity data retrieved from comparative kinetics measurements. In Fig. 9, the densities of bulk DMAS solutions measured in this work are shown together with their third-order polynomial fit, which is the parameterisation used in all the calcula-
Table 1. Parameters of the third-order polynomial parameterisation for density and refractive index (R.I.) of the melt calculated with molar refraction mixing rule (Sect. 2.1). DMAS is the original fitting of experimental data (black circles in Fig. 9), while “Error+” and “Error−” are calculated supposing a ±2% error on the extrapolated ρmelt, as described in the text.

<table>
<thead>
<tr>
<th></th>
<th>Third-order polynomial density coefficients</th>
<th>Melt</th>
<th>Melt ρ</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>a</td>
<td>b</td>
<td>c</td>
</tr>
<tr>
<td>DMAS</td>
<td>0.99847</td>
<td>−2.55 × 10⁻³</td>
<td>0.34103</td>
</tr>
<tr>
<td>Error+</td>
<td>0.99847</td>
<td>+0.017256</td>
<td>0.342053</td>
</tr>
<tr>
<td>Error−</td>
<td>0.99847</td>
<td>−0.02231</td>
<td>0.339909</td>
</tr>
</tbody>
</table>

Figure 9. DMAS density ρ dependence on mass fraction (mfs⁰.⁵) of solute. Symbols: black squares – measured densities, this work; solid line – third-order polynomial fit of measured ρ values; grey shaded area – evaluated uncertainty of the density parameterisation (calculated as discussed in the main text); open circles – measured densities from Clegg et al. (2013); dashed line – densities from the apparent molar volumes fitting by Clegg et al. (2013).

Figure S4 shows a further comparison of density data from this work and from Clegg et al. (2013) converted in terms of apparent molar volumes; both measured and fitted apparent molar values are provided and the coefficient of the fitted equations can be found in Table S5. These discrepancies may be attributable to the different preparation procedures of the aminium sulfates solutions and to the different experimental techniques for the measurement of densities.

The hygroscopic properties of DMAS obtained from the treatment of the evaporation kinetics data with the “original” density parameterisation and with the upper and lower bounds on the density treatment are shown in Fig. 10. Differences in the hygroscopic properties of DMAS are represented either in terms of the dependence of either n_water/n_solute or GF on water activity (panels a and b), the three curves deriving from the three different density treatments are virtually indistinguishable; thus, the uncertainty in the applied density parameterisation does not significantly alter the analysis. In the case of the osmotic coefficients plot (panel c) some very slight deviations between the three treatments can be distinguished at the two extremes of the plot. With respect to the low sulfate molality region, these small differences are due to the fact that a small variation in the sulfate molality results in more significant variations in the osmotic coefficients, because m appears in the denominator in the osmotic coefficient expression (Eq. 2). If the high sulfate molality region is considered, the variations among the three curves are more significant because the simulated error on the density parameterisation is larger for more concentrated solutions (i.e. the size of the grey envelope in Fig. 9).

It is clear that the hygroscopic properties of aminium sulfates determined from CK-EDB measurements are relatively insensitive to reasonable variations in the extrapolated ρ_melt value (±2%) and to the applied density parameterisation. In addition, the variations introduced by different density parameterisations are very small compared to the differences from the results of Sauerwein et al., who obtain lower values of the hygroscopicity and osmotic coefficient. We conclude...
that these differences cannot be caused by inaccuracies in the approaches for treating variations of the density of evaporating droplets.

4 Atmospheric importance and conclusions

Quantifying the hygroscopic properties of aminium sulfates is important for understanding and modelling the atmospheric processes in which they are involved. In particular, the role of short-chained alkylamines in the formation of new particles has been investigated in recent literature and found to be significant (Sect. 1). Aminium sulfate-rich nanoparticles that derive from new particles formation events can potentially act as CCN, and their hygroscopic properties must be well-characterised with the aim of reducing the overall uncertainties that currently affect our understanding of the indirect effects of atmospheric aerosols on climate. Robust and accurate data are essential for improving microphysical models of aerosol hygroscopicity; this study presents an extensive dataset for an homologous series of six compounds, compared to ammonium sulfate, extending over a wide range in RH. In addition, it represents the most comprehensive characterisation of the hygroscopic response of aminium sulfate aerosol so far, complementing previous bulk phase measurements (comparable in accuracy but limited to higher water activity) and aerosol measurements at lower RH (with lower accuracy than achieved here). Previously, the bulk and aerosol measurements reported in the literature were in disagreement. Here, we report aerosol measurements that are in good agreement with the previously most accurate bulk phase data, resolving this discrepancy.

In order to measure their hygroscopic properties with a CK-EDB, the dependence of aminium sulfates solution densities and refractive indices on mass fraction of solute are reported. The experimental technique together with the application of the molar refractive mixing rule and a third-order polynomial parameterisation of density for the representation of the refractive indices and densities of solutions with variable solute mass fractions were presented and validated in previous works (Cai et al., 2016; Davies et al., 2013; Rovelli et al., 2016). The procedure for preparation of the aminium sulfates stock solutions was validated by using it for the preparation of ammonium sulfate solutions from the direct reaction of ammonia and sulfuric acid. The hygroscopic properties of the obtained (NH$_4$)$_2$SO$_4$ solutions retrieved from CK-EDB measurements were in very good agreement with calculations from E-AIM model and we demonstrated that the uncertainty of such measured values was comparable with a typical uncertainty of ±0.002 on $\omega_w$, which was previously shown to be typical for CK-EDB measurements (Rovelli et al., 2016). This result demonstrated that the used preparation procedure is robust and reliable.

The experimental results for the aminium sulfates were compared with the few studies available in the literature and the observed discrepancies were discussed in the light of the different experimental approaches. Measurements from our new approach provide a level of accuracy that reveals clearly the fine variations in hygroscopic growth (down to discernible difference is GF of the order of 0.01–0.02) that occur with molecular structure and substitution, and avoids the additional complexity of volatilisation of semi-volatile components during hygroscopic growth with measurements complete in a matter of a few seconds. The largest discrepancies with previous data were found when comparing the CK-EDB results with H-TDMA measurements (Clegg et al., 2013; Qiu and Zhang, 2012). These differences can be attributed to a possible overestimation of the dry size of particles due to residual water in the reference dry state in HTDMA measurements and/or to a shift of the chemical composition of particles towards the bisulfate composition because...
of partial volatilisation of the amines from solution. Closer comparability was found with the bulk water activity measurements by Sauerwein et al. (2015). The main differences in approaches are that we perform aerosol measurements that cover a wider range in water activity as compared with the bulk measurements of Sauerwein et al. (2015), and we provide direct measurement at amine-to-sulfate ratios of exactly 2 : 1, whereas Sauerwein et al. (2015) performed a ZSR fitting on data from solutions with variable amine-to-sulfates ratios and extrapolated water content for the exact 2 : 1 ratio.

These new CK-EDB measurements suggest a higher level of hygroscopic growth for the aminium sulfates than previously reported by Sauerwein et al. (2015) when inferred from measurements over a range of amine-to-sulfates ratios; we have provided a refined parameterisation for all compositions.

It is worth noting that the characterisation of the hygroscopic properties of aminium sulfates up to $a_w$ of 0.99 was possible with the CK-EDB technique. The other literature approaches that were discussed in this section were able to cover larger $a_w$ ranges (down to 0.1) but none of them could be applied to obtain any data for $a_w > 0.9$. Thus, the comparative kinetics measurements in a CK-EDB provide a powerful tool for investigating a water activity region that is otherwise hard to characterise with such accuracy, but which is of great importance for the understanding of the activity of aerosol particles as cloud condensation nuclei (Wex et al., 2009).

As a final remark, aminium sulfates are the first class of mixed inorganic–organic aerosol systems to be investigated by means of the CK-EDB comparative kinetics technique, described in Rovelli et al. (2016), over a wide range of water activities. Therefore, besides the atmospheric relevance of these compounds, this study also provides a deeper understanding of the possible effects caused by random errors in the experimental procedure, and by uncertainties on the representation of the density of a compound. We demonstrated that the CK-EDB measurements are characterised by a very good level of reproducibility and that a typical ±2 % uncertainty on the value of the extrapolated melt density only marginally affects the measured hygroscopic properties.

**Data availability.** The experimental data presented in this paper are provided through the University of Bristol data repository at Reid et al. (2017).

The Supplement related to this article is available online at doi:10.5194/acp-17-4369-2017-supplement.

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

**Acknowledgements.** Rachael E. H. Miles, Jonathan P. Reid, and Simon L. Clegg acknowledge support from the Natural Environment Research Council through grant NE/N006801/1. Grazia Rovelli acknowledges the Italian Ministry of Education for the award of a PhD studentship.

Edited by: D. Topping
Reviewed by: three anonymous referees

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Aminium sulfate aerosols

G. Rovelli et al.: Aminium sulfate aerosols

Atmos. Chem. Phys., 17, 4369–4385, 2017

www.atmos-chem-phys.net/17/4369/2017/


