The effect of atmospherically relevant aminium salts on water uptake

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Abstract. Atmospheric new particle formation is initiated by clustering of gaseous precursors, such as small acids and bases. The hygroscopic properties of those precursors therefore affect the hygroscopic properties of aerosol particles. In this work, the water uptake of different salts consisting of atmospheric small acids and amines was studied computationally using the conductor-like screening model for real solvents (COSMO-RS). This method allows for the prediction of water activities in atmospherically relevant salts that have not been included in other thermodynamics models. Water activities are reported here for binary aqueous salt solutions, as well as ternary solutions containing proxies for organic aerosol constituents. The order of the studied cation species regarding water activities is similar in sulfate, iodate, and methylsulfonate, as well as in bisulfate and nitrate. Predicted water uptake strengths (in mole fraction) conform to the following orders: tertiary > secondary > primary amines and guanidinos > amino acids. The addition of water-soluble organic to the studied salts generally leads to weaker water uptake compared to pure salts. On the other hand, water-insoluble organic likely phase separates with aqueous salt solutions, leading to minimal effects on water uptake.

1 Introduction

Water uptake of atmospheric aerosol has a large effect on the radiative forcing through, e.g., cloud formation. Theoretical and experimental evidence show that even a small amount of salt has a dominant effect on cloud condensation nucleus (CCN) activity (Bilde and Svenningsson, 2004; King et al., 2012). These salts may originate from sea spray or from the nucleation and condensation of acidic and basic compounds, such as sulfuric acid and amines.

The nucleation of gas-phase acids and bases has been investigated computationally during recent years by computing formation energies of clusters that contain various small molecules (Elm, 2021). Computational studies have shown that various amines can form molecular clusters with acidic species, such as sulfuric acid (Elm et al., 2016; Myllys et al., 2018; Kubečka et al., 2023). Liu et al. (2022) found that guanidino-containing compounds (guanidine, agmatine) enhance methylsulfonic-acid-driven nucleation. Chen et al. (2022) simulated the nucleation of nitric acid with various alkylamines, alkanolamines, and polyamines. They found that alkanolamines and polyamines can nucleate with nitric acid at room temperature, while ammonia and alkylamines require lower temperatures.

The activity of water in aerosol particles describes the response of the particles to variations in RH. Some experimental studies have investigated water activities of ammonium and aminium sulfate as well as nitrate salts (Bonner, 1981; Clegg et al., 2013; Sauerwein et al., 2015; Rovelli et al., 2017). For example, Rovelli et al. (2017) found that water activity decreases with the increasing alkyl group length and number of alkyl groups. However, water activity measurements of salt solutions comprising larger amines relevant to salt solutions are still lacking.

This study investigates the effect of different amine and acid species involved in the atmospheric nucleation on water activity computationally. Many thermodynamics models commonly used for activity coefficient calculations, such as AIOMFAC (Zuend et al., 2008, 2011), are parameterized for only a few atmospheric ions, such as $\text{NH}_4^+$, $\text{SO}_4^{2-}$, $\text{HSO}_4^-$, $\text{NO}_3^-$, and $\text{IO}_3^-$. On the contrary, the quantum-
chemistry-based conductor-like screening model for real solvents (COSMO-RS; Klamt, 1995; Klamt et al., 1998; Eckert and Klamt, 2002) can be used to compute thermodynamic properties of any atmospherically relevant species. COSMO-RS and AIOMFAC predict similar water activities in (NH₄)₂SO₄, NH₄HSO₄, NH₄NO₃ and NH₄IO₃, within a factor of 1.08 and 1.30 in aqueous solutions with 0.1 and 0.2 mole fraction of the salt, respectively (see Fig. S1 of the Supplement). Though COSMO-RS calculations are much slower (several hours) than group contribution methods (∼1 s), mainly due to the quantum chemistry input needed for the COSMO-RS calculations, COSMO-RS is currently the only method capable of estimating water activities in aminium salts. For this reason, COSMO-RS was used here to predict water activities of the salt solutions of interest.

The acids selected for this study are abundant in the atmosphere. Sulfuric acid (H₂SO₄) is derived from anthropogenic emissions of SO₂, nitric acid (HNO₃) is produced from both anthropogenic and natural processes, and iodic acid (HIO₃) and methylsulfonic acid (CH₃SO₃H) are more abundant in marine environments. In aqueous solutions, these strong acids are deprotonated to form sulfate (SO₄²⁻) and bisulfate (HSO₄⁻), iodate (IO₃⁻), and methylsulfonate (CH₃SO₃⁻), respectively. The following atmospherically abundant amines have many biogenic (vegetation, biomass burning) and anthropogenic (pesticides, combustion, livestock) sources (Ge et al., 2011). Additionally, the effect of the salts on water uptake was investigated in the presence of a hydrophilic and a hydrophobic organic.

2 Computational methods

The COSMO-RS model (Klamt, 1995; Klamt et al., 1998; Eckert and Klamt, 2002), implemented in the BIOVIA COSMOtherm program (BIOVIA COSMOtherm, 2021a, abbreviated COSMOtherm), was used for activity coefficient and liquid–liquid equilibrium (LLE) calculations. Activity coefficient γ of compound i is computed in COSMOtherm using the following equation:

\[
\ln \gamma_i(x) = \frac{\mu_i^\circ(x) - \mu_i^\circ(x^\circ)}{RT},
\]

where \(x\) is the composition of the mixture, \(T\) is the temperature (here 298.15 K), \(R\) the gas constant (in kJ K⁻¹ mol⁻¹), \(\mu^\circ\) is given in kJ mol⁻¹, and \(P = 10^5\) Pa reference pressure. Pseudo-chemical potential (Ben-Naim, 1987) \(\mu^\circ\) is an auxiliary quantity defined using the chemical potential at the reference state \(\mu^\circ\) :

\[
\mu_i^\circ(x) = \mu_i^\circ(x^\circ, T, P) + RT \ln \gamma_i(x).
\]

The pure compound \((x_i = 1)\) is used as the reference state composition \(x^\circ\). The use of pseudo-chemical potential allows for the calculation of activity coefficients in infinitely diluted states \((x_i \to 0)\).

The occurrence of liquid–liquid or solid–liquid phase separation (LLPS or SLPS, respectively) is not considered in COSMOtherm activity coefficient calculations. Instead, the mixtures are assumed to be homogeneous. Small aerosol particles are less likely to phase separate than larger droplets (Freedman, 2020). One-phase states are therefore possible in small particles even if the bulk solution would phase separate.

The solubility of the hydrophobic organic in the salt solutions was found using the LLE condition:

\[
\gamma_i(x^\alpha) x_i^\alpha = \gamma_i(x^\beta) x_i^\beta.
\]

Here, \(\alpha\) and \(\beta\) denote the aqueous phase and the organic-rich phase. In the LLE calculation, the salt is assumed to be fully dissolved into the liquid phases.

2.1 Electrolytes in COSMOtherm

The framework of COSMOtherm calculations treats electrolytes (i.e., solvent and one salt) as ternary solutions (solvent, cation, anion). This affects the mole fractions (input) and activity coefficients (output) of the program. All mole fractions presented with the results were converted to the laboratory-binary framework (solvent, salt) for easier comparison with experiments.

The highest level of theory parameterization in COSMOtherm (BP_TZVPD_FINE_21, abbreviated FINE, using BP/TZVPD/BP/TZVP level cosmo files) works poorly with the strong charges of small ions, such as atomic ions and strong small semi-spherical ions (such as SO₄²⁻, NH₄⁺) (BIOVIA COSMOtherm, 2021b). The use of an electrolyte parameterization (BP_TZVPD_ELYTE_21; abbreviated ELYTE; using BP/TZVP level cosmo files) is recommended for up to 6 mol kg⁻¹ solutions, corresponding to around 0.1 mole fraction of salt in water (BIOVIA COSMOtherm, 2021b). However, the use of ELYTE is not recommended for large ions (BIOVIA COSMOtherm, 2021b). Based on these recommendations, the use of ELYTE should be limited.
to systems for which it was developed unless the calculated values can be corroborated by experiments.

The different COSMOtherm parameterizations are compared in Sect. S1 of the Supplement. Overall, the experimental water activities agree equally well with both ELYTE and FINE parameterizations with the exception of ammonium sulfate, with which FINE significantly overestimates the experiments (see Figs. S2–S4). For some cations (e.g., ARG), ELYTE results are highly inconsistent with the results of FINE and the BP_TZVP_21 parameterization (abbreviated TZVP), which for most salts give fairly similar estimates. Additionally, the current electrolyte parameterization is only available for the lower level of theory BP/TZVP cosmo files and not for the higher level of theory BP/TZVPD cosmo files, corresponding to the FINE parameterization. In case there are systematic errors within the different parameterizations or levels of theory, the comparison of the different salts is more reliable when the same parameterization is used in all calculations. The FINE parameterization is therefore used for all of the studied salts, with the exception of ammonium sulfate, which is calculated using the ELYTE parameterization.

Using the selected parameterizations, COSMOtherm agrees with the experimental water activities of ammonium sulfate, bisulfate, and nitrate solutions (x_{salt} < 0.2) within factors of 1.39, 1.18, and 1.23, respectively. Similar uncertainties can be expected for other amonium salts. Additionally, as can be seen from Figs. S2–S4, the disagreement between calculations and experiments increases with the increasing salt mole fraction in the solution.

### 2.2 Input files for COSMOtherm calculations

The input files for COSMOtherm calculations (cosmo-files) were obtained through a series of density functional theory calculations with increasing levels of theory. The process has been discussed in more detail in a previous publication (Hyttinen et al., 2020). In short, all conformers were found using the systematic conformer search algorithm in the Spartan20 program (Wavefunction Inc., 2020). The geometries of all conformers were optimized, and duplicate conformers were removed using the BIOVIA COSMOfree program (BIOVIA COSMOfree, 2021), which uses the Turbomole program (TURBOMOLE, 2020) for the quantum chemistry calculations. The final cosmo-files were computed at the BP/def2-TZVPD-FINE//BP/def-TZVP level of theory (BP/def-TZVP for ELYTE and TZVP calculations).

Many of the studied cations have multiple conformers. At most the 10 lowest chemical potential conformers were selected as input for the COSMOtherm calculations. However, only conformers with chemical potentials within 2 kcal mol$^{-1}$ of the lowest chemical potential were used in order to avoid including conformers with low COSMO energies but high chemical potentials (Hyttinen, 2023b). More specifically, COSMOtherm gives high weights to conformers containing intramolecular H bonds (Hyttinen and Prisle, 2020) because intramolecular H bonds are favored in the COSMO energies (Kurtén et al., 2018). There is an excellent correlation between chemical potentials in water and in electrolyte solutions (see Fig. S7). This indicates that conformer distributions are similar in pure water and electrolytes. The conformer sets for electrolyte calculations can therefore be selected using the chemical potentials of the ions in water.

### 3 Results and discussion

#### 3.1 Salt–water solutions

Water activities were calculated in solutions ranging from pure water to 0.2 mole fraction of salt. It should be noted that the water solubility of some salts at 298.15 K may be below 0.2 mole fraction. However, all salt was assumed to be dissolved in the solvent water as ions. Additionally, all cations in the calculations are singly charged for simpler comparison, even if some contain multiple amine groups. Figure 1 shows COSMOtherm-derived water activities in selected salt solutions at 298.15 K. The values are given in Tables S1–S16 in the Supplement and in Fig. S8. Unfortunately, determining reliable error estimates of water activities in these salts is not possible due to the lack of experimental data for similar salts. However, the errors are likely to be systematic for similar compounds, such as amonium ions. The order of the cations may therefore be more reliable than the order of the anions.

Figure 1 shows that both the anion and cation affect the water activity of the solutions. Assuming that the ions investigated here follow the Hofmeister series, the order of cations should be similar regardless of counter ion. Based on COSMOtherm estimates, the order of the cations is fairly similar in sulfate (Fig. 1d), iodate and methylsulfonate (Fig. 1f) salts. Similarly, bisulfate and nitrate (Fig. 1e) salts have a similar order for the cations. This indicates that the effect of the cation on water uptake also partially depends on the anion. Additionally, the order of the studied anions is not the same for all of the cations.

Some patterns can be seen between the different functional groups of the cations. For example, water uptake is enhanced in the following order: tertiary amine > secondary amine > primary amine (Fig. 1a). This order follows the hygroscopicity of ammonium sulfate salts (Rovelli et al., 2017) and is the same as Dawson et al. (2014) found for water uptake of amonium methylsulfonate salts (TMA > DMA > MMA). Additionally, two amine groups in the cation lead to lower water activity than one amine group, even though only one of the amine groups is protonated. This is seen in the comparison between TMA and TMEDA, as well as MMA and EDA (Fig. 1a).

Carboxylic acid groups have inconsistent effects on water activity. Amino acids have lower water activities than MMA in some salts (bisulfate, nitrate; Fig. 1f) and higher in others (sulfate; Fig. 1d, iodate, methylsulfonate; Fig. 1e). Addition-
ally, there is no clear trend between the studied amino acids with one (ALA, GLY, SER) or two (GLU, ASP) acid groups. On the other hand, water activity is lower in all five ARG (one carboxylic acid group) salts than in the corresponding AGM (no carboxylic acid group) salts (Fig. 1d–f).

In sulfate solutions, the water activities have the largest dependence on the cation compared to the other studied anions. Additionally, GLY and ASP sulfates are predicted to inhibit water uptake at low salt concentrations (water activity coefficients above one, see Tables S12 and S14). It should be noted that the ionic strength of the sulfate solutions is 3 times the ionic strength of the other studied solutions with the same salt mole fraction.

In their experiments, Sauerwein et al. (2015) observed similar osmotic coefficients of aqueous AM, MMA, DMA, TMA, and DEA bisulfate solutions, indicating similar water activities independent of the cation. Similarly, the COSMOtherm-derived water activities in the bisulfate solutions are similar (within a factor of 1.1 in $x_{\text{salt}} = 0.1$ solutions) for all studied cations. Due to uncertainties in the calculated values, it is possible that there are no real differences between the water activities of the different aminium bisulfate solutions. Conversely, MMA, DMA, TMA, and DEA sulfate solutions had lower experimental water activities than the corresponding bisulfate solutions, while water activities in AM bisulfate solutions were similar to those in AM sulfate solutions (Sauerwein et al., 2015). A similar trend is seen in the COSMOtherm-derived water activities.

Myllys (2023) studied the effect of water on the growth of acid–guanidine clusters in the gas phase computationally. At the molecular level, water was seen to enhance particle formation at 298 K in systems containing nitric acid and methylsulfonic acid at relatively low acid concentrations. No significant effect was seen in sulfuric acid systems. The same order in these three acids is seen in COSMOtherm bulk-phase calculations. Nitrate enhances water uptake to the aqueous phase more than the other two anions, methylsulfonate enhances water uptake slightly less than nitrate, and the bisulfate has the weakest enhancing effect. However, COSMOtherm predicts the lowest water activity in solutions containing sulfate ions, indicating the strongest water uptake. However, since the concentration of guanidine was lower than the sulfuric acid concentration in the simulations of Myllys (2023), the sulfuric acid is expected to form bisulfate ions and not sulfate ions. If the correlation between COSMOtherm-estimated water activities and the enhancement on cluster formation at high RH holds for other acid–base pairs, the relative effect of RH on particle forma-
tion could be estimated using the computationally simple water activity calculations instead of the more time-consuming molecular cluster calculations.

Higher equilibrium water content in aerosol particles indicates a higher hygroscopic growth factor. Because the salts have different molar masses, mass fraction water content provides a more direct comparison for the hygroscopic growth factors of the salts. Table 1 shows the mass fraction water content of each of the studied salts at 70 % RH as an example. Comparing the anions, the highest water content is found in either the sulfate or the nitrate salt for all studied cations. For most of the studied cations (all except TMEDA), the lowest water content in terms of mass fraction is in the iodate salt. COSMOtherm predicted water mass fractions at 70 % RH vary between 0.19 (ASP-IO₃⁻) and 0.64 (TMA₂-SO₄³⁻) for the studied salts.

### 3.2 Addition of water soluble organic

Next, an organic compound was added to the aqueous salt solutions to investigate the effect of the salts on water activity in the presence of an organic component. An isoprene-derived epoxydiol (trans-β-IEPOX, C₅H₁₀O₃) was selected as the organic compound because of its high estimated gas-phase production rate in the atmosphere (100 Tg C per year; St. Clair et al., 2016). Additionally, COSMOtherm predicts miscibility between water and IEPOX and only slight deviation from ideality in all mixing ratios (Hyttinen et al., 2020). This is important, as limited solubility between water and the chosen organic would lead to strong deviation from ideality for water by the organic. In such cases, liquid–liquid phase separation should be taken into account in the activity coefficient calculations.

#### 3.2.1 Water content

Figure 2 shows the water activities in ternary salt–IEPOX–water solutions of three different methylsulfonate salts and IEPOX. The values were calculated with 0.1 intervals of salt mole fraction in the dry mixture (x_{salt,dry}).

![Figure 2](https://doi.org/10.5194/acp-23-13809-2023)
The order of the studied cations varies depending on both of the other anions, likely due to the higher ionic strength. The salting out effect of sulfate is much stronger than that of the other anions, indicating that organosulfates are less likely to form in solutions containing equal mole fractions of sulfuric acid and amines (Aoki et al., 2020). Reactive uptake of IEPOX into aqueous sulfate aerosols may counteract the strong salting out effect of sulfate salts for organic compounds that can form ionic species by reacting with the sulfate ion.

Since organosulfates are likely to form from the selected organic (IEPOX), additional water activities were computed for mixtures containing equal mole fractions of ammonium organosulfate salt and neutral amine. This mixture may arise if 1 : 1 mole ratio of sulfate salt and IEPOX react, neutralizing half of the aminium ions in the process:

$$\text{epoxide} + \text{SO}_4^{2-} + 2\text{RNH}_3^+ \rightarrow \text{organosulfate}^- + \text{RNH}_3^+ + \text{RNH}_2.$$  \hspace{1cm} (R1)

While the elemental content of the mixture remains the same, organosulfate formation reaction reduces the ionic strength of the mixture by neutralizing half of the amines and converting the doubly charged sulfate into singly charged organosulfate.

Water activities were additionally computed for organosulfate solutions that do not contain neutral amines. These mixtures correspond to either organosulfate formation from an epoxide reaction with bisulfate ion, or evaporation of the neutral amines after the organosulfate formation from a reaction with sulfate ion.

Figure 4 shows the COSMOtherm-derived water mass fraction in the bisulfate, sulfate (x axis), and organosulfate (y axis) mixtures at 70 % RH. Similar water content is predicted for aminium bisulfate–IEPOX and aminium organosulfate mixtures (magenta circles in Fig. 4). On the other hand, larger differences are seen between aminium sulfate–IEPOX and the corresponding aminium organosulfate–amine mixtures. For most of the studied cations, water uptake is predicted to be stronger for the aminium sulfate–IEPOX mixture than for the corresponding aminium organosulfate–amine mixture. One reason may be the difference in ionic strength between sulfate and organosulfate. Additionally, the COSMOtherm estimates of sulfate salts may be less accurate than the of salts comprising singly charged non-spherical ions, such as bisulfate and organosulfate.

While amines can also react with epoxides in their neutral forms, they are likely to exist in protonated forms in acidic secondary organic aerosol. The nucleophile strengths of protonated amines are much weaker than those of neutral amines (Stropoli and Elrod, 2015). This makes reactions between the studied aminiums and IEPOX unlikely. The nitrate ion can also act as a nucleophile and react with the epoxy group.
However, the product tertiary organonitrate is not as stable as the corresponding organosulfate, leading to water substituting the nitrate group to form a methyltetrol (Darer et al., 2011). The formation of an organonitrate species was therefore not considered here.

3.3 Addition of water-insoluble organic

A hydrophobic water-insoluble organic matter (WIOM; CC(=O)C\textsubscript{1}O\textsubscript{2}C\textsubscript{2}C\textsubscript{3}O\textsubscript{4}C\textsubscript{1}C\textsubscript{1}=CC(C)=CC(C)=C\textsubscript{1}) proposed by Kalberer et al. (2004) has often been used as a proxy in COSMO-RS calculations of atmospheric aerosol properties (Wania et al., 2014; Wang et al., 2017; Kurtén et al., 2016; Hyttinen et al., 2021; Lumiaro et al., 2021). COSMO\textit{therm}-predicted solubility of water in WIOM is low (0.07 mole fraction; Hyttinen, 2023b), indicating that relatively high RH is needed to increase the water content in pure WIOM particles. For example, at equilibrium in 70 % and 90 % RH, the predicted water mole fraction in WIOM is only 0.047 and 0.065, respectively. Adding a salt to the mixture increases equilibrium water content compared to pure WIOM. However, high mole fractions of salt lead to the salting out of WIOM from the aqueous salt solution.

When the salt–WIOM–water mixtures phase separate into an aqueous phase and a organic-rich phase, the aqueous phase will contain mostly water and salt, while the organic-rich phase is almost pure WIOM. The COSMO\textit{therm}-derived solubility of WIOM in water at 298.15 K is very low, only \(2.5 \times 10^{-5}\) mole fraction. The solubility of WIOM further decreases with the addition of salt (salting out of WIOM). The predicted water activity in the aqueous phase is very close to that of the binary salt–water mixtures (see Fig. S13). The effect of very hydrophobic organics on water activity of salts is therefore expected to be negligible, as water activity in both phases at equilibrium is equal according to the LLE condition of Eq. (3).

4 Conclusions

Salts are expected to have a larger effect on water activity and equilibrium water content than neutral organic compounds. Additionally, the critical supersaturation is lowered through decreased water activity in salt-containing particles. The water content of atmospheric aerosol also affects the aerosol radiative forcing, which is a crucial factor in climate models. It is therefore important to know how each ionic species in atmospheric aerosol affects water activity.

Water activities can be used to determine the hygroscopic growth and CCN activation of atmospheric aerosol particles. Knowledge of how different atmospheric ions affect water uptake and CCN activation of aerosol particles will help improve climate models. COSMO\textit{therm} calculations show how the different atmospheric anions and cations affect water activity. Most of the salts of this study enhance water uptake relative to ideal solutions (e.g., pure water).

Organosulfate and bisulfate–organic mixtures have similar effects on the equilibrium water content. However, water uptake may be slower in organosulfate than in sulfate–organic mixtures. This indicates that the formation of organosulfate may reduce the water uptake ability compared to particles that do not contain organic compounds. This should be considered in models that use experimental hygroscopicities of pure sulfate salts to describe the hygroscopicities of mixed salt–organic particles. Future studies are needed to further investigate other ionic species and highly oxygenated multifunctional organics.

Data availability. The research data (cosmo files) can be accessed through the Jyväskylä University Digital Repository (JYX) at https://doi.org/10.17011/jyx/dataset/89238 (Hyttinen, 2023a).

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