Analysis of the hygroscopic and volatile properties of ammonium sulphate seeded and unseeded SOA particles


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Abstract. The volatile and hygroscopic properties of ammonium sulphate seeded and unseeded secondary organic aerosol (SOA) derived from the photo-oxidation of atmospherically relevant concentrations of \( \alpha \)-pinene were studied. The seed particles were electrospray generated ammonium sulphate \((\text{NH}_4)_2\text{SO}_4\) having diameters of approximately 33 nm with a quasi-mono-disperse size distribution (geometric standard deviation \(\sigma_g=1.3\)). The volatile and hygroscopic properties of both seeded and unseeded SOA were simultaneously measured with a VH-TDMA (volatility – hygroscopicity tandem differential mobility analyzer). VH-TDMA measurements of unseeded SOA show a decrease in the hygroscopic growth (HGF) factor for increased volatilisation temperatures such that the more volatile compounds appear to be more hygroscopic. This is opposite to the expected preferential evaporation of more volatile but less hygroscopic material, but could also be due to enhanced oligomerisation occurring at the higher temperature in the thermodenuder. In addition, HGF measurements of seeded SOA were measured as a function of time at two relative humidities, below (RH 75%) and above (RH 85%) the deliquescence relative humidity (DRH) of the pure ammonium sulphate seeds. As these measurements were conducted during the onset phase of photo-oxidation, during particle growth, they enabled us to find the dependence of the HGF as a function of the volume fraction of the SOA coating. HGF’s measured at RH of 85% showed a continuous decrease as the SOA coating thickness increased. The measured growth factors show good agreements with ZSR predictions indicating that, at these RH values, there are only minor solute-solute interactions. At 75% RH, as the SOA fraction increased, a rapid increase in the HGF was observed indicating that an increasing fraction of the \((\text{NH}_4)_2\text{SO}_4\) is subject to a phase transition, going into solution, with an increasing volume fraction of SOA. To our knowledge this is the first time that SOA derived from photo-oxidised \( \alpha \)-pinene has been shown to affect the equilibrium water content of inorganic aerosols below their DRH. For SOA volume fractions above \( \sim 0.3 \) the measured growth factor followed roughly parallel to the ZSR prediction based on fully dissolved \((\text{NH}_4)_2\text{SO}_4\) although with a small difference that was just larger than the error estimate. Both incomplete dissolution and negative solute-solute interactions could be responsible for the lower HGF observed compared to the ZSR predictions.

1 Introduction

The atmospheric aerosol is the assembly of liquid or solid particles which are suspended in the gaseous phase of our atmosphere. This aerosol consists of a mixture of organic and inorganic substances. Organic matter contributes 20% to 60% of the mass of fine particles, depending on location (Chow et al., 1994; Duce et al., 1983; Krivacsy et al., 2001a, b). Particulate organic carbon may be either directly emitted as primary organic aerosol, or produced in the atmosphere as secondary organic aerosol (SOA) from chemical degradation of gas-phase precursors such as natural hydrocarbons (e.g. terpenes) and anthropogenic hydrocarbons.
Inorganic particles typically experience pronounced phase transitions with varying RH, with sudden changes from dry crystals to solution droplets and vice versa that are characterized by sharply defined points of deliquescence relative humidity (DRH) or efflorescence relative humidity (ERH). DRH and ERH are both dependent on composition and between these points the particles undergo a hysteresis in water uptake, i.e. for a specific composition and at a RH between DRH and ERH the particle can be either liquid or solid. The knowledge of the values of DRH and ERH is of great importance since, given the RH history of an air parcel they define which fraction of the atmospheric aerosol is present as purely liquid (possibly supersaturated) droplets, which as mixed-phase solid/liquid particles, and which as purely solid particles. The role of organics in the deliquescence and efflorescence behavior of inorganic aerosols is not fully understood (Cruz and Pandis, 2000; Marcolli et al., 2004) despite the importance of the physical state of such aerosols.

Conventionally, the water uptake of atmospheric particles has been associated with their inorganic fraction, while the contribution of organic aerosols has generally been neglected. However, Saxena et al. (1995) have pointed out the importance of atmospheric water-soluble organic carbon (WSOC) for the hygroscopicity of atmospheric aerosols. In addition, WSOC has been postulated to have roles similar to that of inorganics in visibility degradation and the formation of cloud condensation nuclei (Cruz and Pandis, 1998; Novakov and Penner, 1993). Although there have been measurements of the hygroscopic properties of organic salts and carboxylic acids of atmospheric relevance (Peng et al., 2001; Peng and Chan, 2001; Sjogren et al., 2007b; Gysel et al., 2004), and a number of organic species have been incorporated into a liquid phase thermodynamic model (Clegg and Seinfeld, 2006; Topping et al., 2005; Ming and Russell, 2002) there is still a need to expand our understanding using a combined approach to describe the behaviour of multi-component mixtures of inorganic and organic compounds including potential phase separation such as through the use of LIFAC (Kiepe et al., 2006) or X-UNIFAC (Erdakos et al., 2006).

The literature on the effect of organics on the hygroscopic growth factor (HGF) of inorganic aerosols is, in general, limited and offers varying examples of their influence: some organic compounds affect the hygroscopic growth of inorganic aerosols positively and others negatively. Prenni et al. (2001) found that some dicarboxylic acids retain water to very low RH and never exhibit efflorescence. Thus, organics have, if they are sufficiently water soluble, an increasing effect on HGF below the DRH of the inorganics, and a diminishing effect above DRH (Sjogren et al., 2007b). In addition, if the organic compounds are present as surface films on aqueous droplets the mass transfer of gaseous molecules into and out of the droplet can be impeded and the hygroscopic growth factor (HGF) is reduced (Andrews and Larson, 1993; Edney et al., 2005; Hansson et al., 1998; Lightstone et al., 2000; Xiong et al., 1998). Saxena et al. (1995) found that organic compounds appeared to diminish the water uptake of inorganic compounds at urban locations, while a positive contribution was reported for a continental nonurban location. Cruz and Pandis (2000) found that glutaric acid and pinonic acid, in general, enhance water sorption of ammonium sulphate ((NH4)2SO4, hereinafter referred to as AS) but reduce that of NaCl relative to that of the pure inorganic salts. Thus, the effects of organics on inorganic aerosols appear to be species-specific. The measurements of Marcolli et al. (2004) for a range of organics and inorganics showed, in general, little influence of the organic/inorganic interactions on the water activity in mixed particles, whereas the solubility of the inorganics was clearly influenced. Also, Dick et al. (2000) found that the organic-associated water content was less than that of sulphate compounds at high relative humidity (RH) but comparable or greater at low RH.

The measured positive effect on growth due to organics is likely due to water absorption by the organics (and/or the chemical effect with the inorganics, i.e., chemical interactions between the inorganic and organic species) while observed negative effects can be either due to kinetic limitations caused by organic coatings (Xiong et al., 1998), mass transfer limitations due to inhomogeneous morphology inside the particle (Sjogren et al., 2007b), or a thermodynamic effect caused by persistent chemical interactions. The distinction between these three effects is important because organic films may not necessarily affect the water absorption of atmospheric particles at sub-saturation because of longer time scales for RH changes compared to just a few seconds applied in the traditional H-TDMA technique (Rader and McMurry, 1986; McMurry and Stolzenburg, 1989).

Choi and Chan (2002) investigated the effect of malonic and citric acid on the hygroscopic properties (water content, DRH and ERH) of NaCl and ammonium sulphate using an electrodynamic balance (EDB). They found that at low RH (<30%) the inorganic components in the mixed particles have crystallized, but the mixtures often contain water due to the presence of the organic species. The mixtures can then absorb water even though the RH is much lower than the DRH of the pure inorganic species. Again, hysteresis effects exist, and therefore prediction of the water content requires knowing the RH history of a particle. However, in the presence of organics deliquescence may occur at low RH and may easily be missed in laboratory and field hygroscopic measurements. At high RH (>80%), particles become completely deliquesced. A big range of ambient RH falls between the DRH and the ERH of NaCl and ammonium sulphate, i.e. RH=45–80%, where the effects of organics on deliquescence and efflorescence are very complex and require special attention.
A useful technique to study the water uptake of mixed organic inorganic species is the combined Volatility and Hygroscopicity Tandem Differential Mobility Analyzer (VH-TDMA). Briefly, the VH-TDMA system operates on the same principles as a typical TDMMA (Rader and McMurry, 1986) with the additional feature that the sample aerosol is passed through a humidified region after exiting the thermodenuder. The particle hygroscopic growth factor can then be determined as a function of the thermodenuder temperature. By increasing the temperature of the thermodenuder the more volatile species will start to evaporate from the particle (i.e. organic species are usually more volatile than inorganic sulphates) and the change in the HGF will be observed. The VH-TDMA technique was first applied in the investigation of internally mixed species and common atmospheric species (Johnson et al., 2004). Since then it has been used to study marine aerosols (Fletcher et al., 2007; Johnson et al., 2005), atmospheric iodine formation (Ristovski et al., 2006), or diesel nanoparticles (Meyer and Ristovski, 2007). The benefits of using this system are highlighted in Meyer and Ristovski (2007) where it has been shown that removal of outer layers from multi-component aerosols enables the study of internal fractions.

This paper presents an analysis of the hygroscopic properties of ammonium sulphate seeded and unseeded SOA as measured with the Hygroscopicity Tandem Differential Mobility Analyser (H-TDMA) technique. Growth factors of seeded particles were observed above and below the DRH of AS during the onset phase of the photo-oxidation experiments. The aim of this study was to characterise the relative contributions of the inorganic and organic aerosol components to the overall water uptake above and below the DRH of AS. Additional to this, combined volatility and hygroscopicity studies were conducted on seeded and unseeded particles using the VH-TDMA technique.

2 Experimental methods

SOA was generated via photo-oxidation of α-pinene in a 27-m³ Teflon chamber at 20°C and 50% relative humidity (Paulsen et al., 2005). The chamber was first humidified to 50% RH before introducing NOₓ. A known concentration of around 10 ppb of α-pinene (Fluka, 99.8%) was evaporated in a heated glass sampling bulb and continuously flushed into the chamber with pure air. Four xenon arc lamps were used to simulate the solar light spectrum and initiate the photochemical reaction. Two types of chamber experiments were conducted with and without ammonium sulphate seeds. During seeded experiments ammonium sulphate was generated using an electrospray aerosol generator (TSI-3480). Ammonium sulphate was used because of its tendency to form approximately spherical seed particles (Gysel et al., 2002). This was considered important when determining total water uptake based on an increase in mobility diameter. The ability of the electrospray to generate high concentrations of nearly mono-disperse (geom. std. dev. σg=1.3) aerosol particles differentiates this study from, for example, the study by (Varutbangkul et al., 2006) where poly-disperse (σg~1.75) seed samples were used. The general approach with poly-disperse seed aerosol leads to mixed sizes of seed species and gives greater uncertainty in determining SOA coating thicknesses with respect to seed size. Size distribution data and number concentrations were measured using a scanning mobility particle sizer (SMPS, TSI 3080) and a condensation particle counter (CPC, TSI 3022). The SMPS performed a scan every 3 minutes while the CPC continuously measured with a 1s time resolution. Chamber temperature and relative humidity were monitored using Rotronic Hygro Clip SC05 probes with an accuracy of 0.3°C and 1.5% respectively. Broad chemical speciation of the aerosol components in the chamber was monitored with an Aerodyne quadrupole aerosol mass spectrometer (AMS) (Canagaratna et al., 2007; Alfarra et al., 2006). Volatile and hygroscopic properties were measured with a VH-TDMA (Johnson et al., 2004).

3 Experimental results

A representative time series of the number size distribution measured during a seeded SOA experiment is shown as a contour plot in Fig. 1. During the hour prior to lights on, the injection of the seed particles can be noted as an increase in concentration centered around a mean diameter of 33 nm. At time t=0 h, the chamber is illuminated, initiating the photochemical oxidation process. As the reactions proceed, gas-to-particle conversion via condensation to the seed surface occurs as the volatility of the reaction products is reduced (in the case of non-seeded experiments, this condensation phase is preceded by homogeneous nucleation of low volatility organic compounds). In all experiments, the condensation process becomes equal to the wall loss after 5 h. From this

![Fig. 1. Time series of size distributions measured inside the smog chamber starting with (NH₄)₂SO₄ seed aerosol (seed diameter of approximately 33 nm and σg=1.3) followed by the growth due to photo-oxidation of α-pinene.](image-url)
time onwards the total aerosol mass is constant, and after 6 h the total mass is decreasing (more losses than condensation). Although there was some growth after this period, our interest was in the period leading up to 6 h after lights on (Fig. 1). Chemical ageing processes, however, continue and become increasingly apparent when observing volatile and hygroscopic properties.

3.1 Particle composition

The top graph in Fig. 2 shows the temporal evolution of the AMS signal. During the onset phase of the reaction both the organic and sulphate concentrations increase. This is attributed to the condensation of the organic species on the seed surface leading to an increase in particle diameter thereby improving the transmission efficiency of the aerosol particles to the AMS (Hinz und Sprengler, 2007). After three hours however a decline in the sulphate begins due to wall losses such that by 6 h after lights on the sulphate mass concentration has been reduced to approximately 50% of its peak value. Within measurement uncertainties, ammonium follows the sulphate evolution. For the same period the measured organic concentration continually increases until ~5 h after lights on. After this period our calculations show that the wall losses offset the condensational growth keeping the total mass roughly constant. Given wall losses, condensation and subsequently increasing particle diameters within the chamber it was thought that the particle detection efficiency of the AMS would increase rather than decrease. In principle, a reduction in sulphate mass could also be explained by formation of organic sulphates. Surratt et al. (2007) have recently shown the formation of ester sulphates in similar chamber measurements conducted using ammonium sulphate seeds and photo-oxidized α-pinene and isoprene. However the formation of ester sulphates was observed to a much greater extent when the (NH$_4$)$_2$SO$_4$ seed aerosol was acidified by addition of sulphuric acid or an acidic seed aerosol was produced by photo-oxidation of SO$_2$. Our experiments did not utilise acidic seed aerosol or SO$_2$.

Initially, this does suggest a limited likelihood of any possible reaction occurring between the organic and the sulphate and therefore in our case further investigation is required to clarify the reduction in sulphate mass. The bottom graph in Fig. 2 displays the evolution over time of the ratio of the organic to inorganic mass (solid red circles) and of the organic to sulphate mass (solid squares). In the case of the former, the inorganic fraction incorporates the sulphate and ammonium mass fractions.

3.2 Hygroscopic properties of seeded particles

Analysis of the hygroscopic properties of the seeded species was conducted during the onset phase of the photo-oxidation experiments. During two consecutive seeded chamber experiments the VH-TDMA was operated with the thermodenuder bypassed, therefore operating in the H-TDMA mode, at fixed relative humidities of 75% and 85%, respectively. These RH’s were chosen to be below and above the deliquescence relative humidity (DRH) of the pure (NH$_4$)$_2$SO$_4$ seed material. The evolution of the hygroscopic growth factor (HGF), for a 33 nm dry diameter particle, was measured continuously from the moment the lights in the chamber were turned on every 2 min for around 1.5 h. The results for the two experiments conducted at 75% and 85% RH are presented in Fig. 3.

The growth factor of the pure seed was equal to 1 at 75 % RH as expected for any RH below the deliquescence point of (NH$_4$)$_2$SO$_4$. A slow increase of the HGF was initially observed after turning the lights on, followed by a strong increase to a HGF of ~1.25 approximately 30 minutes later, when the (NH$_4$)$_2$SO$_4$ to SOA volume ratio dropped below 0.7. Then the HGF decreased continuously until a HGF of about 1.07 was recorded after about 1.7 h. For pure SOA particles of equal dry diameter measured in unseeded experiments after similar light exposure time growth factors of ~1.05 were exhibited (green line on the bottom graph of
Fig. 3. Change in the hygroscopic growth factor of seeded SOA as a function of elapsed time from the moment the lights were turned on. Top graph at RH=85% (full circles) and bottom graph at RH=75% (open circles). In addition the theoretical HGF of ammonium sulphate (magenta) and the measured HGF of unseeded SOA (green) at the two different RH are also presented.

Fig. 4. Change in the hygroscopic growth factor (full circles, left y-axis) and volume fraction remaining (VFR) (squares, right y-axis) as a function of the thermodenuder temperature. These measurements were conducted after 5 h of measurement time had elapsed and condensation of organic species was considered to have ceased. Top graph is of 100-nm seeded particles and the bottom graph is of pure 100-nm SOA particles. In addition the HGF* (open circles) corrected for the Kelvin effect (at constant $a_w$) is also presented. For comparison, on the top graph, the VFR of pure AS particles is presented with a dashed blue line.

3.3 VH-TDMA spectra of seeded and unseeded particles

The volatile and hygroscopic properties were measured simultaneously with the VH-TDMA both for the unseeded (pure SOA) and seeded particles. The particles were heated in the thermodenuder to a temperature that was ramped from room temperature to 200°C in steps of 5 or 10°C. For each thermodenuder temperature the hygroscopic growth factor was measured. The bottom graph in Fig. 4 shows the HGF at RH=85%, for 100 nm pure SOA particles with respect to the thermodenuder temperature as well as the volume fraction remaining (VFR). In addition the HGF corrected for the Kelvin effect at $a_w=0.85$ (HGF*; assuming surface tension of pure water) is also presented. These measurements were conducted after 5 h since the lights were turned on and have taken around 1 h for a temperature scan from room to 200°C. During this time although there was some condensation of organic species it was not considered to be significant. Further any aging effects during the VH-TDMA scan, of 1 h, were also negligible.

For pure SOA particles a reduction in particle diameter in response to heating results in a decrease of the HGF at constant RH due to the increasing influence of the Kelvin effect. In the absence of chemical transformation the Kelvin corrected HGF* would remain constant, while for mixed particles the relative fractions of more and less hygroscopic compounds may also be altered resulting in a change of both HGF and HGF*. Here a decreasing trend of both HGF and HGF* is seen for temperatures from 70 to 115°C. This would indicate that the volatilized compounds are more hygroscopic than the remaining material. A further increase in the thermodenuder temperatures up to 180°C results in a more pronounced reduction of the HGF than the Kelvin corrected one, HGF*, indicating that the reduction in the growth factor...
is more influenced by the reduction in particle size than a change in the composition. One would expect that preferential evaporation of more volatile (less hygroscopic) material occurs with increasing temperature, resulting in an increased HGF. However, the opposite is true, suggesting that the remaining aerosol has an enhanced degree of oligomerisation, resulting in larger molecules with lower HGF. It should be noted that enhanced oligomerisation could occur as a result of the increased temperature. This would mean that the V-TDMA introduces an artefact, by rendering the particles less volatile than without the heating.

The top graph in Fig. 4 shows the change in HGF and HGF* of 100-nm seeded particles measured at 75% RH with respect to thermodenuder temperature. Similar to the measurements of pure SOA these measurements were also conducted 5 h since the lights were turned on and condensation of organic species was considered to have ceased. In contrast to the previous case we observe an increase in the HGF as the thermodenuder temperature is increased. This increase is due to the influence of the ammonium sulphate seed. As has been previously shown (see bottom graph in Fig. 3) SOA enables (NH₄)₂SO₄ seeds to take up water at RH of 75%, which is below the DRH of pure AS. As the temperature is increased more of the volatile, but less hygroscopic, SOA is removed from the particles and the volume fraction of (NH₄)₂SO₄ increases therefore resulting in an increase of the HGF. For temperatures above 140°C (NH₄)₂SO₄ starts to evaporate. In the temperature range from 140–165°C the HGF* of pure SOA particles remains constant therefore the reduction in HGF of seeded particles is due to the evaporation of (NH₄)₂SO₄. After 170°C most of the (NH₄)₂SO₄ has evaporated and the decrease of the HGF is due to the Kelvin effect as can be seen from the constant value of the HGF*.

4 Discussion

The HGF of a mixed particle (HGFₘ) can be estimated from the growth factors of the individual components of the particle and their respective volume fractions, applying the ZSR relation (Stokes and Robinson, 1966) which is equivalent to:

\[ HGF_m = \left( \sum_k \varepsilon_k HGF_k^3 \right)^{1/3} \]  

where the summation is performed over all of the compounds that are present in the particle. \( \varepsilon_k \) is the volume fraction and \( HGF_k \) is the hygroscopic growth factor of each of the components present in the particle. The major simplifying assumption of the ZSR relationship compared to full thermodynamic models is that there are no solute-solute interactions, i.e. that the water uptake of the mixed particle is equal to the sum of the individual water uptakes of each component. Correct treatment of the Kelvin effect within the ZSR relationship means to take the \( HGF_k \) at equal water activity as \( HGF_m \), or in good approximation at equal RH and dry size as the mixed particle. The ZSR relation in the form of Eq. (1) contains additional assumptions: sphericity of the particles and no change in the volume upon mixing (ideal mixing behaviour).

In the case of a seeded SOA particle a two component system, with ammonium sulphate as one component and SOA as the other is assumed. Applying the ZSR relationship the HGF of such a mixed particle becomes:

\[ HGF_m = \left( \epsilon \times HGF_{SOA}^3 + (1 - \epsilon) \times HGF_{AS}^3 \right)^{1/3} \]  

where \( \epsilon \) is the volume fraction of SOA; \( HGF_{SOA} \) and \( HGF_{AS} \) are the hygroscopic growth factors of pure SOA and ammonium sulphate. \( HGF_{AS} \) is taken from ADDEM (Topping et al., 2005), whereas \( HGF_{SOA} \) is taken from the measurements conducted during unseeded experiments during this campaign and previous studies (Duplissy et al., 2008). It should be noted that \( HGF_{SOA} \) increases slightly with ongoing photooxidation (Duplissy et al., 2008). The measurements of pure SOA were usually not available at exactly equal dry size, RH and time after lights on. For this reason the semi-empirical k-Köhler theory (Petters and Kreidenweis, 2007) was used to extrapolate the original measurement to the RH and dry size required as input for the ZSR model. Associated uncertainties are taken into account for the error estimates of the ZSR predictions.

The volume fraction of the SOA (\( \epsilon \)) in a particle of a given dry size at a certain time was determined from the evolution of the particle number size distribution measured by the SMPS (see Fig. 1). The diameter of the seed core, \( D_{seed} \), corresponding to coated particles with diameter \( D_{coated} \) at a certain time \( t \) is obtained from the following implicit equation:

\[ \frac{1}{N_{tot}(t_0)} \int_0^{D_{seed}} dN(D_p, t_0) \frac{d \log D_p}{d \log D_p} d \log D_p = \frac{1}{N_{tot}(t)} \int_0^{D_{coated}} dN(D_p, t) \frac{d \log D_p}{d \log D_p} d \log D_p, \]

where \( N_{tot} \) denotes the total number of particles. \( t_0 \) is the time at the beginning of the experiment after introducing the pure seed aerosol into the reaction chamber and before starting the photochemical reaction. The volume fraction of the SOA, \( \epsilon \), becomes then:

\[ \epsilon = 1 - \frac{D_{seed}^3}{D_{coated}^3} \]  

Calculating \( \epsilon \) with equations 3 and 4 contains the assumption that the percentile values of \( D_{seed} \) and \( D_{coated} \) in the number size distributions at time \( t_0 \) and \( t \), respectively, are equal. This approach assumes that the particles don’t change
their relative order in particle diameter upon condensational growth, i.e. coagulation is ignored. The uncertainty of seed diameters determined in this way is estimated to be about 5% and is propagated through the ZSR predictions. Comparisons of the ZSR predictions with the H-TDMA measurements of the seeded SOA particles at 85% and 75% are shown in Fig. 5 and Fig. 6, respectively.

Comparison of the measured HGF at 85% relative humidity and calculated applying the ZSR relationship Eq. (2) is shown on Fig. 5 (top graph). The bottom graph in Fig. 5 shows the change of the volume fraction of SOA (ε) derived from the SMPS measurements as described above, which was used for the ZSR calculations. Two types of ZSR predictions are shown: the first one assuming that the (NH$_4$)$_2$SO$_4$ seed is fully dissolved (cyan line; $HGF_{AS}=1.49$) and the second under assumption that the (NH$_4$)$_2$SO$_4$ seed is not dissolved (dark blue line; $HGF_{AS}=1$). The shaded band around the calculated curves illustrates the high and low estimates for a 5% uncertainty in the SMPS-derived seed diameter and a 3% uncertainty in the HGF of the pure SOA. The error bars for the measured HGF data reflect an RH accuracy of $\Delta RH=\pm 1.2\%$. A good agreement between the calculated growth factor, assuming a fully dissolved seed, and measured growth factors is observed. This indicates that at RH of 85% the ZSR relation is accurate for this mixture, which means that there is little interaction between the organic and inorganic solutes and that they independently absorb water. This further confirms that the (NH$_4$)$_2$SO$_4$ is fully dissolved at the RH of 85%. Similar observations of good agreement between the ZSR theory and measured HGF for mixtures of organics and inorganics at high RH, when the components have completely deliquesced, was observed previously (Choi and Chan, 2002; Cruz and Pandis, 2000; Marcolli et al., 2004).
close to the “undissolved” ZSR prediction and as the volume fraction of SOA (ε) increases the HGF_{in} moves towards the “fully dissolved” ZSR prediction. The observed increase in the growth factor, with an increase in ε is larger than what the “undissolved” ZSR prediction implies. This indicates that an increasing fraction of the (NH_{4})_{2}SO_{4} is in solution with an increase of the volume fraction of SOA. This is in agreement with the previous observation of Marcolli et al. (2004) where the addition of a small amount of organic species (dicarboxylic acids) has increased the solubility of (NH_{4})_{2}SO_{4} in the mixed particle by up to 40%.

The maximum HGF_{in} was observed for ε ~ 0.3 after 0.5 h. A similar maximum water uptake at RH of 75% was also observed by Marcolli et al. (2004) for a mixture of 5 dicarboxylic acids and (NH_{4})_{2}SO_{4} (see Fig. 7 in Marcolli et al., 2004). The main difference is that they observed a maximum water uptake at a larger fraction of organics of 50 wt% while we observed a maximum water uptake at around 20 wt% (under the assumption that the SOA density is 1.3 g/cm^{3} (Varutbangkul et al., 2006; Alfarra et al., 2006) and the AS density is 1.77 g/cm^{3}).

After this time the measured growth factor follows roughly parallel to the ZSR prediction based on fully dissolved (NH_{4})_{2}SO_{4}. It is important to note that there is a difference between the calculated and measured growth factors which is just larger than the error estimate.

There are several possible reasons for over prediction of HGF_{in} with the ZSR approach assuming fully dissolved (NH_{4})_{2}SO_{4}. The first to consider would be that there are kinetic limitations such that the equilibrium RH is not reached in the instrument for the time the particles are exposed to high humidity (which is around 2 s in the RH conditioner and an additional 3 s in the H-DMA). The mass transfer analysis for submicron particles (Kerminen, 1997) shows that the required time to achieve equilibrium is less than the times experienced in our system. On the other side if organic coatings, such as in our case, are present on the particles it has been suggested that they may reduce the rate of water transport across the surface by acting as a physical barrier (Chuang, 2003; Xiong et al., 1998; Johnson et al., 2005). If insufficient time is allowed for particles to achieve their final size a lower HGF will be measured. However it is not likely that there were kinetic limitations in the measurements at RH of 75% as we would have observed similar effects at RH of 85%. As discussed previously measurements at RH of 85% show good agreement with the ZSR prediction.

A second possible reason could be then that the (NH_{4})_{2}SO_{4} seed was not completely dissolved. Let us now calculate the apparent dissolved fraction of (NH_{4})_{2}SO_{4}. SOA produced by photo-oxidation of α-pinene exhibits a continuous water uptake even at very low RH’s (Duplissy et al., 2008; Varutbangkul et al., 2006) therefore we can assume that at RH=75% SOA is fully dissolved. If SOA is fully dissolved and AS is partially dissolved the hygroscopic growth factor can then be calculated as:
$HGF_m = \left( \varepsilon_{SOA} \times HGF_{SOA}^3 + \varepsilon_{DAS} (1 - \varepsilon_{SOA}) \times \right) \frac{1}{(1 - \varepsilon_{DAS}) (1 - \varepsilon_{SOA}) \times 1} \right)^{1/3}$

Where $HGF_{SOA}$ is the growth factor of fully dissolved SOA with a volume fraction of $\varepsilon_{SOA}$; $HGF_{AS}=1.38$ is the growth factor of pure AS (at RH=75%) and $\varepsilon_{DAS}$ the apparent dissolved volume fraction of AS. From the measured $HGF_m$ and knowing the volume fraction of SOA ($\varepsilon_{SOA}$) as well as the $HGF_{SOA}$ it is possible to calculate $\varepsilon_{DAS}$ from Eq. (5). Note that $\varepsilon_{DAS}$ used in Eq. (5) is the ratio of the volume of dissolved AS to the total volume of AS present in the particle. The calculated $\varepsilon_{DAS}$ is presented on the bottom graph (right axis) of Fig. 6. The coloured band around the line representing the apparent dissolved volume fraction presents the overall uncertainty arising from the uncertainties of $\varepsilon_{SOA}$, $HGF_{SOA}$, and $HGF_m$, as already introduced above.

As the volume fraction of SOA increases so does the $\varepsilon_{DAS}$ of AS until it reaches a maximum of around 80% for $\varepsilon_{SOA} \approx 0.3$. For larger values of $\varepsilon_{SOA}$ the $\varepsilon_{DAS}$ decreases though this trend is not significant regarding the large uncertainty. The initial increase in $\varepsilon_{DAS}$ is similar to what would be expected from binary salt mixed solutions (see discussion on p. 517, Seinfeld and Pandis, 1998). Seinfeld and Pandis (1998) discuss the changes in a particle of a given composition as RH is increased. Let us discuss a slightly different case, which corresponds to our situation, where the composition of a binary mixture of two electrolytes at a constant RH is changed. Marcolli et al. (2004) estimated that the DRH of the eutonic composition of five dicarboxylic acids and ammonium sulphate was below 40%. Hence we can assume that at RH=75% we are between the DRH of the component with a higher DRH (AS) and the DRH of the eutonic composition. Initially we have pure AS and the particle is solid. Addition of SOA causes the particle to consist of two phases: a solid phase consisting of pure AS and an aqueous solution of SOA and AS. The composition of the aqueous solution corresponds to the composition of a mixed particle that will deliquesce at RH of 75%. All of the SOA that is added to the particle goes into solution. As the composition of the aqueous solution has to remain constant at a constant RH this will cause more AS to be dissolved. Once the particle composition becomes equal to the composition of the particle that will deliquesce at RH of 75% all of the AS should have been dissolved and the particle should have only one phase—liquid.

As mentioned previously our calculations show that AS is never fully dissolved and reaches a maximum value of 80%. A possible reason for the apparent dissolved fraction to be smaller than unity could be due to negative solute-solute interactions between SOA and AS. Topping et al. (2005) applied the ADDEM model to the results of Marcolli et al. (2004) where they studied the water activity of a multi-component organic mixture (5 components) with a single inorganic salt. They showed that at higher relative humidity, when the solutions are more dilute, the additive approximation works very well. This means that the interaction between the organic and inorganic fractions is not as strong as in more concentrated solutions. In the case of the AS-organic system that they analysed, the deviations increase with decreasing relative humidity, i.e., more concentrated solutions. The model tends to over predict the observed water content which indicates that the organic—inorganic interactions lead to negative contributions to the total water content for AS. This agrees well with our experimental observation where the measurements at RH of 85% agree well with the additive approximation (ZSR theory – no interactions between AS and SOA) while for the measurements at RH of 75%, the interaction is negative leading to a smaller HGF than predicted by the ZSR theory.

We also compared the calculated HGF’s using the ZSR relationship for the VH-TDMA measurements conducted on seeded 100-nm particles (previously presented on the top graph of Fig. 4). This comparison is now presented on the top graph of Fig. 7. Same as before two calculations are shown with a fully dissolved AS seed and under assumption that the AS seed is not dissolved. As the particle diameter reduces with increased thermodenuder temperature a variable $HGF_{AS}$ of pure ammonium sulphate at RH=75%, reflecting the Kelvin effect, had to be used as input for the ZSR calculations. To calculate the $HGF_{SOA}$ of SOA for different thermodenuder temperatures, measurements conducted with pure (unseeded) SOA particles were used (see bottom graph on Fig. 4) and extrapolated to correct RH and dry size using the k-Köhler theory (Petters and Kreidenweis, 2007; Sjogren et al., 2007a).

A similar situation is observed for the VH-TDMA measurement as in the H-TDMA measurements conducted at RH of 75% shown on Fig. 6. There, the SOA coating thickness increased with time, while for the case presented in Fig. 7 the SOA coating thickness was reduced by passing it through a thermodenuder at an elevated temperature (note that the SOA volume fraction is being reduced as the temperature is increased from left to right which is opposite to the situation shown on Fig. 6, where the SOA volume fraction is increased as time passes going from left to right). We have also calculated the apparent dissolved volume fraction of AS ($\varepsilon_{DAS}$) from Eq. (5). This is presented on the bottom graph (right axis) of Fig. 7. In this case the maximum apparent dissolved fraction of AS was around 60% although the uncertainties involved in these calculations were much larger than in the previous case. This also points out to incomplete dissolution of AS or negative solute-solute interactions due to the more concentrated solutions at 75% RH.
5 Conclusions

The hygroscopic and volatile properties of SOA formed by photo-oxidation of \(\alpha\)-pinene were studied. Two types of chamber experiments were conducted with and without ammonium sulphate seeds. The combined measurements of the volatile and hygroscopic properties of pure, unseeded, SOA were measured with a VH-TDMA. They show a decrease in the HGF for thermodenuder temperatures from 70 to 115°C that is larger than the expected decrease due to the Kelvin effect. This indicates that the compounds that are volatilised in this temperature range are more hygroscopic than the remaining material. This is opposite to the expected preferential evaporation of more volatile but less hygroscopic material that should have resulted in the increase in the HGF. However, this result could also be explained by enhanced oligomerisation occurring at the higher temperature in the thermodenuder.

Hygroscopic growth factors of SOA seeded with AS were measured below (RH 75%) and above (RH 85%) the deliquescence relative humidity of AS. The nearly monodisperse AS seeds used in this study enabled us to calculate the SOA thickness with smaller uncertainty compared to the previous studies. This enabled measurements of the dependence of the HGF at two different humidities as a function of the volume fraction of the SOA coating. As has been seen previously the HGF at RH of 85% shows a continuous decrease as the SOA coating thickness increases. The measured growth factors show good agreement with the ZSR relationship indicating that the additive approach could be used and that solute-solute interactions have a minor influence on the HGF of mixed SOA-AS particles at these humidities.

At 75% RH the observed change was more complex and showed a strong dependence on the SOA volume fraction. As the SOA fraction increased a rapid increase in the HGF was observed indicating an increasing fraction of AS in solution. This shows that the SOA coating, produced from the photo-oxidation of \(\alpha\)-pinene, enables at least partial dissolution of ammonium sulphate at RH below its deliquescence point in pure form. A maximum HGF was observed for a 30% volume fraction of SOA. At this SOA fraction the calculated apparent dissolved fraction of AS did not reach unity but was at a maximum value of 0.8. A further increase in the SOA fraction had a small influence on the apparent dissolved fraction of AS, with any observed change within the measurement error. Both incomplete dissolution of AS or negative solute-solute interactions could be responsible for an apparent dissolved fraction smaller than unity.

The implication of our findings on the atmospheric aerosols is that the presence of SOA coating will enhance the water uptake of the inorganic seeds at RH’s lower than the DRH of the inorganic component. This also indicates that at the RH relevant to the atmospheric conditions SOA coatings could broaden the range over which particle-bound water influences the physicochemical and optical properties of organic/inorganic mixed particles.

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