Hygroscopic properties of the Paris urban aerosol in relation to its chemical composition

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Abstract

Aerosol hygroscopic growth factors and chemical properties were measured as part of the MEGAPOLI “Megacities Plume Case Study” at the urban site LHVP in the city center of Paris from June to August 2009, and from January to February 2010. Descriptive hygroscopic growth factors (DGF) were derived in the diameter range from 25 to 350 nm at relative humidities of 30, 55, 75, and 90 % by applying the summation method on humidified and dry aerosol size distributions measured simultaneously with a humidified differential mobility particle sizer (HDMPS) and a twin differential mobility particle sizer (TDMPS). For 90 % relative humidity, the DGF varied from 1.06 to 1.46 in summer, and from 1.06 to 1.66 in winter. Temporal variations in the observed mean DGF could be well explained with a simple growth model based on the aerosol chemical composition measured by aerosol mass spectrometry (AMS) and black carbon photometry (MAAP). In particular, good agreement was observed when sulfate was the predominant inorganic factor. A clear overestimation of the predicted growth factor was found when the nitrate mass concentration exceeded values of 10 µg m\(^{-3}\), e.g. during winter.

1 Introduction

Aerosol particles are ubiquitous in the atmosphere, and play an important role in a variety of fields such as the terrestrial radiation budget, atmospheric chemistry, climate, and human health. Most aerosol-related effects are controlled by particle size, including aerosol dynamical reaction rates, particle deposition, cloud condensation nuclei activation, and light scattering. In the atmosphere, particles take up water well below deliquescence depending on particle hygroscopicity and ambient relative humidity (Swietlicki et al., 2008). Because the degree of particle hygroscopicity influences ambient particle size and therefore light scattering, hygroscopicity has a direct impact on direct radiative forcing (Haywood and Boucher, 2000). Emphasizing the fact that
aerosol water uptake affects both the direct and indirect light scattering properties of aerosols, an incomplete understanding of aerosol hygroscopicity has been identified as a major limiting factor in estimating the aerosol climate forcing (Solomon et al., 2007).

The scientific understanding of the hygroscopic properties of atmospheric particles has improved considerably during the recent years. While the hygroscopic growth of pure soluble ionic species is known quite accurately (Tang and Munkelwitz, 1994; Brechtel and Kreidenweis, 2000), the growth of atmospheric organic matter (Saxena and Hildemann, 1996; Gysel et al., 2004) or complex mixtures of various substances that are typical for the atmosphere is more uncertain.

In the atmosphere, a few inorganic salts constitute the major part of the inorganic aerosol fraction (Heintzenberg, 1989; Putaud et al., 2010). While the impact of the organic fraction on aerosol hygroscopicity has been neglected for a long time, some recent studies stressed the influence of organic compounds on overall particle hygroscopicity (e.g. Gysel et al., 2007). Organic matter may contribute 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). Consequently, there has been a significant shift in focus toward organic hygroscopicity in the last decade. A number of theoretical (e.g. Clegg and Seinfeld, 2006; Topping et al., 2005a, b), laboratory (e.g. Peng et al., 2001; Sjogren et al., 2007; Cruz and Pandis, 2000; Massling et al., 2007) and chamber (e.g. Ansari and Pandis, 2000; Cocker et al., 2001a, b; Duplissy et al., 2008) studies investigated how the presence of organics affects the hygroscopic properties of the atmospheric aerosol. Despite recent improvements in understanding the water uptake characteristics of organic-containing particles, in situ measurements are sparse. Combining measurements of particle composition and their hygroscopic behavior, laboratory studies and hygroscopic growth models can be compared with field measurements.

The standard method for calculating hygroscopic growth from the aerosol chemical composition is based on volume-weighted water uptake by the individual chemical constituents, the so-called Zdanovskii–Stokes–Robinson mixing rule (ZSR; Stokes and Robinson, 1966). The prediction of water uptake for the inorganic fraction of at-
mospheric aerosols is usually reliable. In contrast, the vast number of different organic species in the atmosphere (e.g. Saxena and Hildemann, 1996) combined with a limited understanding of organic aerosol hygroscopicity has led to the practice of assigning the water uptake of organics as the particle growth not explained by inorganic constituents (e.g., Malm et al., 2005). This study compares measured and calculated hygroscopic growth factors from field measurements of the aerosol chemical composition and particle number size distributions in the city center of Paris, France.

A limited number of studies (Eichler et al., 2008; Meier et al., 2009; Achtert et al., 2009; Birmili et al., 2009) with the HDMPS system are available. Whereas Meier et al. (2009), Birmili et al. (2009) and Eichler et al. (2008) obtained chemical particle information by impactor measurements, Achtert et al. (2009) were the only with combined HDMPS and AMS data, but at a rural/suburban site in the North China Plain. Our study is therefore the first one investigating HDMPS size resolved particle growth (cf. HDMPS vs. HTDMA differences in Sect. 3.1) in combination with chemical measurements with high time resolution in a European megacity.

2 Experimental

2.1 The MEGAPOLI field study

The measurements analyzed in this study were performed under the umbrella of the European FP7 project “Megacities: Emissions, urban, regional and global atmospheric pollution and climate effects, and integrated tools for assessment and mitigation” (MEGAPOLI) investigating interactions of megacities, air quality and climate. MEGAPOLI includes both basic and applied research, and bridges spatial and temporal scales connecting local emissions, air quality and weather with global atmospheric chemistry and climate (Baklanov et al., 2008). Two intensive field campaigns were conducted in the summer of 2009 and the winter of 2010 in Paris, a European megacity of more than 10 million inhabitants. Quantifying sources of primary and secondary
Hygroscopicity of the Paris urban aerosol

K. A. Kamilli et al.

Aerosol in and around a megacity was a major objective within MEGAPOLI (Baklanov et al., 2008). Our measurements correspond to the atmospheric aerosol obtained under the impact of various diffuse sources found in such a large conurbation. In practice, detailed aerosol and precursor gas measurements were carried out at several urban and suburban sites, and using several mobile laboratories. MEGAPOLI also included airborne observations of the evolution of the megacity plume.

2.2 Sampling site in Paris

The measurements were carried out in the backyard of Le laboratoire d’hygiène de la Ville de Paris (LHVP; 48.828° N, 2.359° E) situated near Place d’Italie in the 13th quarter of Paris, a residential and business district in the south-eastern part of the city of Paris.

The suite of specialized measurements to characterize the hygroscopic growth properties and the chemical composition included a twin differential mobility particle sizer (TDMPS), a humidifying differential mobility particle sizer (H-DMPS), an aerodynamic particle sizer (APS), a high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS), and a multiangle absorption photometer (MAAP). All instruments were placed inside a laboratory container. Air conditioning was used to stabilize room temperatures, which are crucial for reliable measurements of humidified particle number size distributions. A PM$_{10}$ inlet on the container rooftop served as the main inlet for all instruments. A custom-built regenerative adsorption drier (Tuch et al., 2009) which was installed on the rooftop kept the relative humidity of the sample flow well below 30%.

2.3 Instrumentation

2.3.1 Twin differential mobility particle sizer

The particle number size distribution (PNSD) under dry conditions was measured with a twin differential mobility particle sizer (TDMPS) at a time resolution of 10 min. The
particle diameter range was 3–700 nm during the summer campaign and 3–800 nm in winter. The TDMPS follows the principles outlined in Birmili et al. (1999). Briefly, two Vienna-type differential mobility analyzers (Vienna DMA, Winklmayr et al., 1991) and two corresponding condensation particle counters (UCPC 3025 and CPC 3010; TSI Inc., St. Paul, MN, USA) are used in the TDMPS. With the aid of additional regenerative diffusion dryers inside the TDMPS, typical conditions were between 5 and 26 % relative humidity (RH) of the aerosol inlet flow. The measured raw mobility concentration was inverted taking individual CPC collection efficiencies and DMA transfer functions into account.

2.3.2 The humidifying differential mobility particle sizer

Measurements of humidified particle number size distributions were carried out using a humidifying differential mobility particle sizer (HDMPS, Nowak, 2006; Birmili et al., 2009), which was successfully deployed in several atmospheric field experiments (Eichler et al., 2008; Meier et al., 2009; Achtert et al., 2009; Birmili et al., 2009). The measurement principle is similar to the TDMPS except that the PNSD is recorded in a diameter range from 25 to 800 nm under controlled relative humidities of 30, 55, 75 and 90 %.

In a first step, the sample aerosol is humidified to approximately 90 % RH in a Nafion pre-humidifier (Model MH-110-12-S, ANSYCO Inc., Karlsruhe, Germany) to ensure that the deliquescence point is exceeded. Because of pre-humidification to 90 % RH, one can assume spherical solution droplets from hygroscopic particles or particles consisting of mixtures of hygroscopic and hydrophobic compounds such as mineral dust or crystalline organics (Sjogren et al., 2007).

Subsequently, the aerosol is conditioned to the target humidity by passing the flow through a membrane diffusion dryer (Model MD-110-12F, ANSYCO Inc., Karlsruhe, Germany). It is important to bring the aerosol to the target humidity before entering the neutralizer to guarantee constant RH conditions when reaching the charge equilibrium. The sheath air is also conditioned to reach the target RH before entering the DMA.
nally, the aerosol particles are classified according to their electrical mobility in the DMA
and counted with a condensation particle counter (CPC 3010, TSI Inc., St. Paul, MN,
USA). To ensure reliable measurements, at least three complete PNSD scans were
performed at each humidity step. Thus, a complete cycle through all four target RHs
and the selected particle size ranges lasts 2.5 to 3 h, e.g. the data for a chosen diam-
5
eter at the target RH = 90 % are available every 150 min. After 3 data points follows
a gap of approximately 125 min, where the other humidity conditions are recorded. The
humidity, temperature and flow conditions inside the HDMPS were constantly mon-
tored using temperature/humidity sensors (HMP230, Vaisala Inc., Helsinki, Finland)
and differential pressure transducers. The humidity sensors were calibrated before the
campaign with a dew point mirror.

2.3.3 Chemical composition and black carbon

In order to measure the chemical composition, we used a high-resolution time-of-flight
aerosol mass spectrometer (AMS, Aerodyne Research Inc., Billerica, MA, USA), which
uses flash thermal vaporization under high vacuum followed by electron impact ioniza-
tion and time-of-flight mass analysis (Jayne et al., 2000; Jimenez et al., 2003; Cana-
garatna et al., 2007). The AMS provides quantitative information of all non-refractory
species that are vaporized in a few seconds at a temperature of 600 °C including organ-
ics, sulfate, nitrate, ammonium and non sea-salt chloride. The non-refractory fraction
includes the majority of atmospheric components, with the notable exception of black
5

carbon, crustal oxides (dust) and sea salt which require vaporization at higher tempera-
tures. The AMS mass spectra were analyzed using the data analysis software Squirrel
v1.51. (D. Sueper, University of Colorado-Boulder, CO, USA).

Black carbon (BC) was measured using a multiangle absorption photometer (MAAP,
type 5012, Thermo Scientific Inc., Franklin, USA) which measures the amount of light
that is transmitted through and scattered back from a particle-laden glass fiber filter at
a wavelength of 637 nm (Petzold and Schönlinner, 2004). The wavelength of 637 nm
corresponds with the region of the solar spectrum where BC is the primary absorber, thus interferences with “brown carbon” and mineral dust are minimized.

In order to obtain a more complete description of the aerosol chemical composition, the AMS and MAAP measurements were combined and averaged to match the time resolution of the hygroscopicity measurements. Particle density was calculated by comparison of the time series of dry particle volume derived from the PNSD measurements to the total AMS plus MAAP mass concentration.

3 Data processing

3.1 Determination of hygroscopic growth factors

A special data reduction method, the so-called summation method (SM, Birmili et al., 2009; Eichler et al., 2008; Achtert et al., 2009) is employed on combined HDMPS/TDMPS data to extract hygroscopic growth factors with high time resolution. The summation method is different from the procedure used to derive hygroscopic growth factors from humidified tandem differential mobility analyzers (HTDMA, e.g. Rader and McMurry, 1986). While in an HTDMA measurement, one or more hygroscopic growth factors representative of different particle fractions are derived directly for a particular preselected particle diameter, the SM yields a number of descriptive hygroscopic growth factors (DGF) for different particle diameters, because of the combined HDMPS/TDMPS method. By these coupled instruments a dry and a humidified size distribution are obtained, which are divided into the same number of segments. The SM assigns every dry a humidified segment, beginning with the uppermost segment in every size distribution. So a large number of DGFs are obtained, depending on the number of segments. With this method the DGF value represents an average of all hygroscopic particle fractions at the corresponding particle diameter.

One notable requirement for successful application of the SM is that the total number under the complete PNSD remains constant with humidification. This implies that no
particles are lost or formed during the humidification process. If this is the case, the particle number concentrations measured by the TDMPS and the HDMPS are the same, and the humidification may be considered as a size distribution rearrangement (Birmili et al., 2009). In this study, the total particle number concentration ratio TDMPS/HDMPS was 1.12 on average in summer, and 1.18 in winter for 90% RH. Due to the departing TDMPS/HDMPS ratio from unity, the computed GFs had to pass a quantile filter. Data were eliminated when the DGF value exceeded the 10% or 90% quantiles because one can assume that those extreme DGF values result from extreme values in the TDMPS/HDMPS ratio.

The summation method (Nowak, 2006; Birmili et al., 2009) may be considered as a tool that relates dry and humidified PNSDs. In this method, the dry and humidified cumulative number size distributions are split into 500 disjunct segments with an equivalent number of particles. Note that the segments may be different in diameter width. The SM now allocates the lower boundary diameter of the first segment of the dry PNSD to the lower boundary diameter of the corresponding humidified segment, and continues with the next segments. This process starts from the largest particle diameter and steps towards smaller diameters until the lower end of the PNSD is reached. For each segment, the descriptive hygroscopic growth factor is computed as $DGF(D_{p_{(dry)}}) = D_{p_{(wet)}}/D_{p_{(dry)}}$. This procedure yields 500 DGF values representative of 500 particle diameter bins distributed over the PNSD size range. As the DGF bins are usually variable, the DGF curve was re-binned onto a fixed grid from 25–350 nm by interpolation in this study.

One might argue that the segmentation of particle number distributions is not appropriate, because it assumes internally mixed particles. H-TDMA studies usually identify an external particle mixture with regard to their hygroscopicity (Swietlicki et al., 2008). However, as long as no stepwise changes with particle size are observed in the average hygroscopic growth factors, the SM remains valid. In this study stepwise changes were not observed.
Using the summation method, the growth factors were computed for conditions of 55, 75 and 90 % RH based on the dry PNSD. The 30 % RH measurement of the HDMPS can be considered a dry reference measurement providing results comparable to the TDMPS.

### 3.2 Hygroscopic growth model

The determination of the growth factor based on the Zdanovskii–Stokes–Robinson relation (ZSR) requires knowledge of the volume fractions and the growth factors of the pure substances of mixed aerosol particles. The growth factor is estimated from

\[
GF_{\text{mix}}(a_w) \approx \left( \sum_i \varepsilon_i g_i(a_w)^3 \right)^{\frac{1}{3}},
\]

where \(GF_{\text{mix}}\) is the growth factor of the mixed aerosol particle, \(a_w\) is the water activity, \(i\) is the number of compounds, \(g_i\) is the growth factor of compound \(i\) in pure form, and \(\varepsilon_i\) is the volume fraction of compound \(i\) in the dry particle. In good approximation, the water activity \(a_w\) can be replaced by RH (Gysel et al., 2007) to obtain a functional dependence of \(GF_{\text{mix}}\) on RH, \(GF_{\text{mix}}(RH)\). The ZSR growth model implies internally mixed particles and the possibility to calculate the water uptake of mixed particles as the sum of the water uptake of each compound, independently.

The AMS measurements provide mass concentrations of ammonium (NH\(_4\)), nitrate (NO\(_3\)), sulfate (SO\(_4\)) and chloride (Cl). Assuming exclusive formation of ammonium salts, ammonium nitrate, ammonium chloride, ammonium sulfate, or ammonium bisulfate may be formed. If the main ammonium salts are NH\(_4\)NO\(_3\), NH\(_4\)Cl and (NH\(_4\))\(_2\)SO\(_4\), the ion charge balance requires

\[
\frac{m(\text{NH}_4^+)}{18} = \frac{m(\text{Cl}^-)}{35} + \frac{m(\text{NO}_3^-)}{62} + 2\frac{m(\text{SO}_4^{2-})}{96}
\]
where $m$ is the mass concentration of the ions, respectively.

In analogy, if the main ammonium salts are $\text{NH}_4\text{NO}_3$, $\text{NH}_4\text{Cl}$ and $\text{NH}_4\text{HSO}_4$, the ion charge balance requires

$$\frac{m(\text{NH}_4^+)}{18} = \frac{m(\text{Cl}^-)}{35} + \frac{m(\text{NO}_3^-)}{62} + \frac{m(\text{SO}_4^{2-})}{96}. \quad (3)$$

In the case of the inorganic salts, the growth factors of the pure substances were obtained from the modified Köhler theory according to the model by Pitzer (1973). A detailed description can be found in Nowak (2006). For elemental carbon (EC), we assume a growth factor of unity. The hygroscopic growth curve of organics has been determined by a commonly used empirical $\gamma$ model (e.g. Massling et al., 2003; Swietlicki et al., 2000),

$$\text{GF} = (1 - \text{RH}/100\%)^{-\gamma}, \quad (4)$$

where $\gamma$ is the model parameter.

4 Results

4.1 Campaign overview

The measurements in the city center of Paris started in the evening of 29 June and ended at 1 August 2009 in summer and lasted from 12 January to 15 February 2010 in winter. Figure 1 gives an overview of the TDMPS/HDMPS and AMS measurements. The time series of the measured DGF shows a clear difference between summer and winter; the values in winter are higher with a maximum value of 1.66 in winter compared to a maximum value of 1.46 in summer. This is consistent with the differences of the aerosol chemical composition. The mass fraction of organic carbon, which reduces the hygroscopic growth factor, is higher in summer than in winter, whereas the fraction of
inorganic salts is higher in winter than in summer. Also, one can notice a shift in the mass fractions of nitrate and sulfate. Whereas sulfate is the dominant inorganic salt in summer, the influence of nitrate increases in winter.

4.2 Correlation between DGF, organic and inorganic mass fractions

As obvious by the temporal patterns of the DGF and the organic mass fraction in Fig. 2a, and the negative regression between the two variables (Fig. 2b), the DGF and the organic mass fraction are clearly anti-correlated. With an increasing mass fraction of OC, the DGF decreases. OC was therefore classified as less hydrophilic, and a value of $\gamma = 0.075$ (Gysel et al., 2004) was chosen in Eq. (4), which leads to a GF for the organics of 1.19 at 90% RH. If we evaluate the linear regression presented in Fig. 2b with the equation $y = -0.33x + 1.43$, the growth factor for the OC mass fraction can be determined by setting $x = 1$ (purely organic sample). By this computation we arrive at a slightly lower value of DGF $= 1.10$. For the winter campaign the linear regression yields a value of DGF $= 1.11$. Figure 2c, d shows the correlation of the mass fraction of the inorganic compounds with DGF. In contrast to the organic fraction, obviously the inorganic compounds have a promoting effect on hygroscopic growth. The time series of the inorganic mass fraction and DGF follow the same pattern, but with differing factors between both. The correlation coefficient ($R$) is $R = 0.79$ indicating a reasonable correlation. For the winter data we obtained a similar correlation, however, the correlation coefficient ($R = 0.57$) was lower than in summer.

4.3 Ion balance

To initialize the hygroscopic growth model on the basis of chemical composition measurements, knowledge of the ion balance is required. Crippa et al. (2013) calculated the ion charge balance for three MEGAPOLI sites during winter assuming the presence of ammonium nitrate, ammonium sulfate, and ammonium chloride. In this study, the observed mass concentration of $\text{NH}_4^+$ and the mass concentration of $\text{NH}_4^+$ in ion
charge equilibrium calculated from Eqs. (2) and (3) is compared in Fig. 3 to determine, whether the ion charges are balanced with ammonium nitrate, ammonium chloride, and ammonium sulfate or ammonium bisulfate. The color of the symbols indicates the measurement date.

Apparently, in summer the relation between the observed and calculated \( \text{NH}_4^+ \) concentrations is split up into two branches for the calculation with ammonium bisulfate (Fig. 3a). It seems that this calculation shows better correlation at the beginning of the campaign until 5 July, and worse correlation for the rest of campaign shown by a steeper slope of the correlation line deviating from unity. On average, the slope of the regression line between the observed mass concentration of \( \text{NH}_4^+ \) and the calculated concentration of \( \text{NH}_4^+ \) in ion charge equilibrium has the value of 1.21. This means the measured values exceed the calculated ones. For the whole campaign the calculation with ammonium sulfate (Fig. 3b) fits the observations best. The correlation line has a slope of \( n = 0.92 \), which means that on average there is only an 8% deviation from perfect charge equilibrium.

In winter (Fig. 3c, d), there is no evident separation of certain time periods, neither for the calculation with ammonium bisulfate nor ammonium sulfate. The computation with ammonium bisulfate yields the best results for the whole campaign. For ammonium bisulfate, a slope of 0.98 indicates that ion charge equilibrium is reached, while for ammonium sulfate, the slope is 0.77. Overall, chloride can be neglected because of its vanishingly low amount.

Using the corresponding densities (Table 1), the volume fractions \( \varepsilon \) of the particular compounds were determined from the observed mass concentrations of inorganic as well as organic compounds. After the preceding considerations concerning the ion balance, all computations were made with ammonium sulfate for summer and with ammonium bisulfate for winter data.
4.4 Case studies of measured and calculated DGFs

We measured and evaluated the DGF at three different RH conditions, i.e. at 90 %, 75 % and 55 % RH. All evaluations of growth factors at different relative humidities were made at a diameter of 285 nm. The reason for this is a concentration maximum within the measurement range of the HDMPS directly below a diameter of 300 nm. This maximum becomes obvious in Fig. 5. The chemical data measured by the AMS represents the PM$_1$ range. Therefore, the computed GF is representative of the submicron aerosol, and a diameter of 285 nm is the best point of comparison within the diameter range of the HDMPS from 25–350 nm. The averaged DGF at 285 nm particle diameter are summarized in Table 2 both for the summer and winter campaigns. While the average DGF at 90 % RH is clearly higher in winter than in summer, the average DGF values at 75 % and 55 % RH are similar in winter and summer.

In order to investigate the dependence of hygroscopic growth on particle composition in more detail, several periods of constant meteorological conditions were examined. In summer, we determined one period with a strong diurnal variation of nitrate in the beginning of the campaign from 29 June to 3 July, shown in the time series of the AMS measurements in Fig. 1a. In this period, the air masses were advected slowly to Paris, as can be seen in the trajectory plot (Fig. 4a) calculated with the HYSPLIT model provided online by the NOAA Air Resources Laboratory (Draxler and Rolph, 2012; Rolph, 2012). The two backward trajectories arriving at 300 m and 750 m a.g.l. travelled over eastern Germany and northern France 96 h before arriving in Paris. The upper air layer arriving at 1850 m a.g.l. travelled a longer distance above the Atlantic Ocean, however, it did not mix with the other layers and, therefore, did not affect Paris in the lower boundary layer. Due to the slow horizontal transport where the wind speed did not exceed 3 m s$^{-1}$, high total particle as well as black carbon concentrations, and a diurnal variation in nitrate developed, which is clearly visible in Fig. 1a. The volume fractions of the aerosol compounds are depicted as pie charts in Fig. 5. Organic carbon is the predominant compound in this period (Fig. 5a) with a volume fraction of 48 %. Ammo-
nium sulfate has a volume fraction of 28%, ammonium nitrate of 13%, EC of 10%, and ammonium chloride a negligible fraction of 0.27%. The size dependence of the corresponding DGF is also represented in Fig. 5a. Obviously, the DGF increases with increasing particle diameter. In general, due to aging processes one can expect more hydrophilic material in older, larger particles than in fresh, smaller particles. Because the volume fractions of the aerosol compounds are derived from AMS measurements roughly representing PM$_1$, the upper size range of the DGF distribution is crucial for an intercomparison of measured and calculated growth factors.

The second time period of interest lasted from 5 to 12 July 2009, when air masses were brought from over the Atlantic Ocean to Paris (Fig. 4b). The particle mass concentration, which varied from 3 to 18 µg m$^{-3}$, and the concentration of the chemical compounds were lower than in the first period. Also, the change in air mass origin led to a shift in the particle composition (Fig. 5b). In the maritime air masses, precursor gases formed by combustion are less abundant than in the urban aerosol. Therefore, less ammonium nitrate (3.8%) was formed, whereas the organic volume fraction (58%) increased. All other volume fractions remained roughly the same. One would expect a lower DGF than in the first period, as less hydrophilic material (OC) is more dominant and the volume fraction of hygroscopic inorganic salts is reduced. However, the maximum DGF is slightly higher in the second period than in the first period. One can speculate that the organic fraction of the second period is more hydrophilic than in the first period.

A third interesting period was found during the winter campaign from 26 to 27 January. Air masses from the east, which crossed densely populated industrial areas in Poland and Germany (Fig. 4c), brought continental aerosol to Paris. Due to the long residence times over these areas, high particle mass concentrations of aged particles were advected. This is also reflected in the aerosol composition with a high volume fraction of ammonium nitrate of 40%, and ammonium bisulfate of 20%. Ammonium chloride volume fractions are slightly higher with 1.2% but still negligible. The OC vol-
volume fraction of 31 % is distinctly lower than in summer. Therefore, the maximum DGF of 1.4 is higher than in summer.

During a fourth period from 2 to 5 February 2010, the air masses were transported over southern France coming from the Atlantic (Fig. 4d). Similar to the second period in summer (Fig. 4b), this period is influenced by maritime air masses. The size dependence of the DGF is approximately the same in these two periods. As in the summer period, the OC volume fraction of more than 50 % is dominant, however, the inorganic salts are mainly ammonium nitrate (21 %) and less than 10 % of ammonium bisulfate.

Overall, the absolute values and the size dependence of the DGF do not show pronounced variations even if the chemical composition and the air mass origin are quite variable. Part of the reason for the small variability of the DFG may by the averaging over extended time periods which may reduce the differences between the four selected periods.

4.5 Overall statistics of measured and calculated DGFs

As already mentioned in Sect. 4.4, the first few days of the summer campaign are different from the rest of the campaign. This is also evident in the DGF measurements of 285 nm particles which show the strongest variation in this time interval (Fig. 6a). Furthermore, the deviation of the measured and the calculated DGF is strongest during this period, whereas the rest of the time series fits well if one considers the use of a simplified growth model. The correlation coefficient of the two time series of measured and calculated DGF at 90 % RH reaches a value of 0.66 for the total of the summer campaign (Fig. 6a). The diurnal nitrate variation and the associated expected variation in the hygroscopic growth factor is indeed well reproduced by the calculated growth factor, but not by the measured DGF. Possible reasons for this discrepancy are either the increased nitrate concentrations or inaccurate assignment of the corresponding ions in the calculation of the growth factor. In the scatter plot of measured DGF and predicted GF (Fig. 6b), the measurements from 29 June to 3 July 2009, are indicated in red. Removing these data points, the correlation coefficient is $R = 0.76$. Overall, the
pattern of the two time series is similar, but the absolute values differ. It is remarkable
that the measured time series of the DGF shows a much higher variability in winter
than in summer. This decreases the correlation coefficient of the wintertime data to $R =
0.52$ (Fig. 6c). During two periods, the calculated GF is distinctly higher than the DGF.
During these periods, 23 to 29 January 2010 and 6 to 15 February 2010, the nitrate
mass fraction is increased. Just like in summer, the correlation between measured and
calculated growth factor improves if these periods (indicated in red in Fig. 6d) are not
taken into account.

Previously, McFiggans et al. (2007) obtained reasonable agreement of measured
and predicted GF if sulfate was the predominant inorganic compound. Worse agree-
ment was found under nitrate enriched conditions. In these cases the computations
overestimated the hygroscopic growth. It was speculated that either the ZSR failed for
this case, or that nitrate measured in the AMS was not from ammonium nitrate but from
organic nitrates. Similar observations were made by Swietlicki et al. (1999) und Gysel
et al. (2002, 2007). These authors reached hygroscopic closure only in the absence
of significant nitrate masses. They all concluded that the ZSR could predict the hy-
groscopic growth with sufficient accuracy, however, nitrate loss based on evaporation
within the HTDMA could induce a discrepancy. For a short-term measurement during
the MEGAPOLI campaign, the AMS was directly connected to the HDMPS inlet and
showed a sudden drop in the nitrate concentration. After connecting the AMS to the
standard inlet again, the nitrate concentration was back at the higher level. Therefore,
we cannot exclude the possibility of nitrate losses due to evaporation in our system.

The possibility of inaccurate assignment of the ions in the GF calculations was eval-
uated using the Extended Aerosol Inorganics Model (AIM, Aerosol Thermodynamics
Model, Model 2; Clegg et al., 1998). The ion pairing scheme was evaluated under
the constraint that only ammonium sulfate, ammonium bisulfate and ammonium nitrate
may form. To run the model, the mole fractions of the different ions at a specific time
were determined at the beginning and end of the campaign when the particle mass
exceeded $25 \mu g m^{-3}$. Due to the fact that we only obtained one organic mass fraction
containing many different organic compounds from the AMS measurements, for the model calculation an artificial organic model compound was created. As a representative model compound, we used the thermodynamic properties of l-malic acid, and a molar mass of 250 g/mol.

The result was not in full agreement with the previous assignments. Ammonium sulfate and ammonium bisulfate occurred in parallel. In summer, at the beginning of the campaign the computed mass of ammonium sulfate was two to three times higher than the mass of ammonium bisulfate. This is similar to our original assignment, where charge equilibrium is reached with ammonium sulfate, ammonium nitrate and ammonium chloride for the summer period. However, for the end of the campaign, the AIM model calculation reached ion balance with ammonium bisulfate in absence of ammonium sulfate. This is in contrast to our original assignment. For the winter period, the AIM model results were in better agreement with our original assignment. Although ammonium sulfate and ammonium bisulfate existed in parallel, ammonium bisulfate concentrations were one to two orders of magnitude higher than ammonium sulfate concentrations, which is consistent with ammonium bisulfate as the dominant sulfate compound.

The water uptake not explained by inorganic constituents (Malm et al., 2005) is often assigned to organics. Then, it is possible to compare the growth factor of the unexplained water uptake with the growth factor of organic matter estimated from the empirical γ model in Eq. (4) (i.e. 1.19, cf. Table 1), thus evaluating the accuracy of the selected growth factor. By rearranging the ZSR-relations to yield the growth factor for the organic fraction, and applying all the known values for the inorganic constituents, a growth factor for the organic fraction can be determined, for which the DGF and the calculated GF coincide. The computations for this study yielded growth factors even below unity for the organic fraction which corresponds to shrinkage with humidification. Thus, the discrepancies between measured DGF and calculated GF could not be equated by a lower growth factor for organic compounds. The median growth factor of organics was 1.06 in summer and 0.87 in winter, respectively. In summer, values below
unity only occurred in the beginning when the fluctuations in the GF values were strong. In winter the time series for the measured GF fluctuated steadily, so the computed GF for organics reached values below unity more often.

5 Conclusions

Aerosol hygroscopicity has been investigated for the inner city of Paris using growth factor measurements by HDMPS/TDMPS. In addition, the ZSR mixing rule was used to predict hygroscopic growth factors based on the aerosol chemical composition measured by AMS and MAAP. The temporal evolution of the predicted growth factor coincided reasonably well with the measured descriptive growth factor. However, it should be noted that the predicted GF almost always was higher than the measured DGF.

Possible reasons for the limited agreement between measured and calculated growth factors are

- ambiguous ion balance calculations, and therefore, ZSR mixing rule computations with inaccurately assigned compounds,

- limitations of the summation method,

- evaluation of the descriptive growth factor measurement for a particle diameter of 285 nm in comparison with growth factor calculations using the chemical composition of PM$_1$,

- evaporation losses of ammonium nitrate.

Despite the fact that the predicted GF was generally slightly higher than the measured DGF, good agreement was observed when sulfate was the predominant inorganic factor. Worse results with a clear overestimation of the predicted growth factors were found when the influence of nitrate became more dominant, e.g. at the beginning of the summer campaign or at the end of the winter campaign.
Overall, the measured and calculated hygroscopic growth factors show little variation during the observation periods in summer and winter. Even under variable meteorological conditions and corresponding changes in the aerosol chemical composition, the growth factors are quite robust. Hygroscopic growth seems to be more dependent from size than from various chemical compositions, as concurrent with Dusek et al. (2006).

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References


Hygroscopicity of the Paris urban aerosol

K. A. Kamilli et al.


Table 1. Overview of the density values and growth factors of the pure substances.

<table>
<thead>
<tr>
<th>Composition</th>
<th>Density [g cm(^{-3})]</th>
<th>GF for 90 % RH and 285 nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>NH(_4)NO(_3)</td>
<td>1.73 (Lide, 1993–1994)</td>
<td>1.71</td>
</tr>
<tr>
<td>(NH(_4))(_2)SO(_4)</td>
<td>1.77 (Meyer et al., 2009)</td>
<td>1.66</td>
</tr>
<tr>
<td>NH(_4)HSO(_4)</td>
<td>1.78 (Lide, 1993–1994)</td>
<td>1.67</td>
</tr>
<tr>
<td>EC</td>
<td>2 (Kostenidou et al., 2007; Gysel et al., 2004)</td>
<td>1</td>
</tr>
<tr>
<td>OC</td>
<td>1.5 (Gysel et al., 2004)</td>
<td>1.19</td>
</tr>
</tbody>
</table>
Table 2.Measured growth factors on average during summer and winter for 55%, 75%, and 90% relative humidity.

<table>
<thead>
<tr>
<th>RH in %</th>
<th>Average DGF for 285 nm, summer</th>
<th>Average DGF for 285 nm, winter</th>
</tr>
</thead>
<tbody>
<tr>
<td>55</td>
<td>1.05</td>
<td>1.04</td>
</tr>
<tr>
<td>75</td>
<td>1.13</td>
<td>1.13</td>
</tr>
<tr>
<td>90</td>
<td>1.27</td>
<td>1.35</td>
</tr>
</tbody>
</table>
Fig. 1. Time series of the DGF of 285 nm particles at a RH of 90% and the particle compounds measured with AMS and MAAP for summer (a) and winter (b).
Fig. 2. Correlation between DGF and the mass fraction (MF) of organic matter in summer (a) and the inorganic MF in summer (c) with the corresponding scatter plots (b, d).
Fig. 3. Comparison for NH$_4$ measured and predicted for ammonium bisulfate (a, c) and ammonium sulfate (b, d) in summer (top) and winter (bottom), where n denotes the slope of the regression line.
Fig. 4. HYSPLIT trajectories for the 2 periods in summer (top) and winter (bottom).
**Fig. 5.** Growth factor distributions with the mean DGF and 0.1 and 0.9 quantiles in four periods in summer (a, b) and winter (c, d). The pie charts show the corresponding volume fractions of the main aerosol compounds.
Fig. 6. Time series (a, c) and scatter plots (b, d) of the measured DGF for 285 nm particles and the calculated GF for 1 µm particles.

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ACPD
13, 14297–14330, 2013

Hygroscopicity of the Paris urban aerosol

K. A. Kamilli et al.

Abstract
Introduction
Conclusions
References

Tables
Figures

Back
Close

Full Screen / Esc

Printer-friendly Version
Interactive Discussion