| 2 | Significant contribution of organics to aerosol liquid water content in |
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| 3 | winter in Beijing, China |
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25 Abstract

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The aerosol liquid water content (ALWC), an important component of atmospheric particles, has a 27 significant effect on atmospheric optical properties, visibility and multiphase chemical reactions. In 28 this study, ALWC is determined from aerosol hygroscopic growth factor and particle number size 29 distribution (PNSD) measurements and also simulated by the ISORROPIA II thermodynamic model 30 31 with measured aerosol chemical composition data at an urban site in Beijing from 8 November to 15 December 2017. Rich measurements made during the experiment concerning virtually all aerosol 32 33 properties allow us not only to derive the ALWC but also to study the contributions by various species for which little has been done in this region. The simulated ALWC including the contribution of 34 organics and the calculated ALWC are highly correlated (coefficient of determination $R^2 = 0.92$). The 35 ALWC contributed by organics (ALWC_{org}) accounts for 30 $\% \pm 22$ % of the total ALWC during the 36 sampling period. These results suggest a significant contribution of organics to ALWC, which is rather 37 different from previous studies that showed negligible contributions by organics. Our results also show 38 that ALWC correlates well with the mass concentrations of sulfate, nitrate, and secondary organic 39 aerosols (SOA) ($R^2 = 0.66$, 0.56, and 0.60, respectively). We further noted that accumulation mode 40 particles play a key role in determining ALWC, dominating among all the aerosol modes. ALWC is an 41 exponential function of ambient relative humidity (RH) whose strong diurnal variation influence the 42 diurnal variation of ALWC. However, there is a three-hour lag between the extremes of ALWC and 43 RH values, due to the diurnal variations in PNSD and aerosol chemical composition. Finally, a case 44 study reveals that ALWC_{Org} plays an important role in the formation of secondary aerosols through 45 multiphase reactions at the initial stage of a heavy haze episode. 46

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50 1. Introduction

51 China has experienced rapid economic developments during the past few decades, resulting in 52 frequent heavy haze events. Severe air pollution may harm human health and affect the regional 53 climate through aerosol direct and indirect radiation effects (Li et al., 2016; G. X. Wu et al., 2016; Wei 54 et al., 2019a; Wei et al., 2019b). However, air pollution formation mechanisms and aerosol climate 55 effects remain highly uncertain due to the complex physical and chemical processes involved (Tao et 56 al., 2012; Y. Wang et al., 2014).

Aerosol liquid water (ALW), a component of atmospheric particles in the atmosphere, exists 57 universally and plays an important role in many atmospheric physical and chemical processes (Nguyen 58 et al., 2016). For example, ALW can influences aerosol optical properties, resulting in increased 59 extinction coefficients, lowered atmospheric visibilities, enhanced aerosol optical depths (AODs), and 60 changes in the direct climatic effect of aerosols (Dougle et al., 1996; Adams et al., 2001; Liao et al., 61 2005; Seinfeld and Pandis, 2006). Secondary aerosols (SA) are considered to be the main source of 62 particulate pollution during heavy haze events in China (Huang et al., 2014). Many studies now 63 highlight the significance of aerosol liquid water content (ALWC) in the formation of SA through 64 chemical reactions (e.g., Arellanes et al., 2006; G. Wang et al., 2016; Cheng et al., 2016). This is 65 because ALW can dilute the absolute concentration of solutes, adjust aerosol acidity, and serve as a 66 reactant, resulting in increases in trace gas (e.g., N₂O₅ and HO₂) uptake coefficients (Wahner et al., 67 1998; Bertram et al., 2009; Abbatt et al., 2012). H. Wang et al. (2017) found that the uptake coefficient 68 of N₂O₅ can be high, which is related to high ALWC in Beijing, thereby increasing the formation of 69 nitrates. ALW can also speed up the aqueous phase chemical reaction by serving as a reactor for the 70 transformation of SO₂ to sulfate (Zheng et al., 2015; G. Wang et al., 2016; Cheng et al., 2016). Some 71 72 studies have found that ALWC can facilitate the formation of secondary organic aerosols (SOA) through aqueous-phase chemistry and photochemistry (Blando et al., 2001; Surratt et al., 2007; 73 Hennigan et al., 2008; Song et al., 2019). Furthermore, observations in Beijing have shown that 74

aqueous-phase processes play a dominant role in the additional formation of oxidized SOA (Xu et al.,
2017). Overall, investigating the formation of SA and haze in North China requires an examination of
ALWC and its factors including aerosol particle number size distribution (PNSD), aerosol chemical
composition and ambient related humidity (RH) in this region.

However, directly measuring real-time ALWC is not feasible yet because of technical limitations, 79 (Kuang et al., 2018). Four indirect methods have been proposed to calculate real-time ALWC: (1) the 80 81 aerosol PNSD under dry conditions and ambient RH conditions are first measured, then ALWC is calculated as the difference between dry and ambient aerosol volumes (Stanier et al., 2004); (2) the 82 83 increased aerosol volume due to water uptake (i.e., ALWC) is calculated according to the measured dry PNSD, size-dependent aerosol hygroscopicity, and ambient RH (Kitamori et al., 2009; Bian et al., 84 2014; Tan et al., 2017); (3) the dry and ambient aerosol volumes are first estimated using the measured 85 aerosol optical enhancement and Ångström exponent, then ALWC is calculated as the difference 86 between dry and ambient aerosol volumes (Kuang et al., 2018); and (4) ALWC is simulated using 87 thermal equilibrium models such as the ISORROPIA thermodynamic model (Nenes et al., 1998), 88 Aerosol Inorganics Model models (Wexler and Clegg, 2002), the Simulating Composition of 89 Atmospheric Particles in Equilibrium model (Kim et al., 1993), and the Gibb's Free Energy 90 Minimization model (Ansari et al., 1999) with aerosol chemical composition information as input. 91

ALWC mostly depends on aerosol PNSD, chemical composition, and ambient RH. Hodas et al. 92 (2014) reported that ALWC in the Po Valley in Italy was driven by locally formed anthropogenic 93 nitrates. The implications for the lifetimes of water-soluble organic compounds and its potential 94 influence on SOA formation were also discussed. Another study also revealed that ALWC in Beijing 95 was driven by secondary inorganic aerosols (SIA; Z. Wu et al., 2018). Most previous studies have 96 focused on the interaction between inorganic salts and ALWC, but the impact of organic species on 97 ALWC been ignored to our knowledge (Blando et al., 2001; Surratt et al., 2007; Hennigan et al., 2008; 98 Carlton et al., 2014). A thorough understanding of the association of ALWC with organic aerosols in 99

100 the atmosphere is lacking.

In this study, ALWC is calculated using the indirect method (2) and simulated using the ISORROPIA II model, i.e., indirect method (4), discussed previously. The effects of inorganic aerosols, organic aerosols, PNSD, and ambient RH on ALWC are then investigated separately. We demonstrate the significant contribution of organics to ALWC in Beijing and provide evidence that the ALW contributed by organics serves as a reactor for sulfate and SOA formation.

106 2. Data and measurements

107 2.1. Sampling site

The Air Pollution and Human Health (APHH) winter field campaign took place from 8 November 108 to 15 December 2016 at the Chinese Academy of Sciences' Institute of Atmospheric Physics Tower 109 Branch in Beijing. Beijing is located in the northwest part of the North China Plain, which has 110 experienced rapid economic developments during the last few decades. A large amount of gaseous 111 112 precursors and other air pollutants are emitted in this region every year, causing serious air pollution problems. The sampling site is located in the northwestern urban area of Beijing (39.97°N, 116.37°E), 113 between the north third and fourth ring roads and surrounded by restaurants. Traffic and cooking 114 emissions are thus the main pollutants at the site. Aerosols at this site can, therefore, well represent 115 anthropogenic aerosols in highly polluted areas. Sun et al. (2013) and Y. Wang et al. (2017) provide 116 more detailed descriptions of the sampling site. 117

118 2.2. Instrumentation

Sampling instruments used during the field campaign included a scanning mobility particle sizer (SMPS) equipped with a long differential mobility analyzer (DMA; model 3081A, TSI) and a condensation particle counter (CPC; model 3772, TSI). A custom-built hygroscopicity tandem differential mobility analyzer (H-TDMA) was installed in an air-conditioned mobile container at

ground level. The temperature inside the container was maintained at 20-25°C. A high-resolution 123 aerosol mass spectrometer (HR-AMS) was set up in a sampling room located on a two-story roof, 124 about 25 m north from the container. Sampled air went through a PM_{2.5} cyclone inlet fixed on the top 125 of the container before entering the instruments. The RH of the sampled air was dried to below 20 % 126 by a dryer system consisting of a tube filled with silica gel and a Nafion dryer (model PD-70T-24ss, 127 Perma Pure Inc., USA). Various meteorological parameters, including wind speed (WS), wind 128 129 direction (WD), temperature (T), and RH, were measured from a 325-m meteorological tower located ~ 20 m west of the container. In this study, all times are reported in Beijing local time (UTC+8 h). 130

PNSDs with electrical-mobility diameters ranging from 10 to 600 nm were measured by a scanning mobility particle sizer (SMPS) at a 5-min time resolution. PNSDs were extended to diameters ranging from 0.6 to 1 μ m by fitting the measured PNSDs with functions consisting of three-mode lognormal distributions (Hussein et al., 2005). Thus generated are PNSDs with a diameter range of 10 nm to 1 μ m.

The H-TDMA system developed by the Guangzhou Institute of Tropical and Marine Meteorology 136 measured the size-dependent aerosol hygroscopic growth factor (GF). The H-TDMA system mainly 137 consists of four parts. The first part is a Nafion dryer to keep the RH of sampled air below 20 % and a 138 bipolar neutralizer (soft X-ray, model 3088, TSI Inc.) to equilibrate the particle charge (Wiedensohler 139 et al., 1988). Next, the sampled air passes through the first differential mobility analyzer (DMA1; 140 model 3081L, TSI Inc.) to produce mono-dispersed particles. In this study, the diameters were set to 141 40, 80, 110, 150, and 200 nm. The sampled air then went through a Nafion humidifier (model PD-70T-142 24ss, Perma Pure Inc., USA) used to humidify the RH of sampled air to 90 %. The last part of the H-143 TDMA is the second DMA (same model as the DMA1) and a water-based condensation particle 144 counter (model 3787, TSI Inc.), used to measure the number size distribution of humidified particles 145 in the five selected diameters. Y. Wang et al. (2017) provide a detailed introduction to the H-TDMA 146 system. 147

Size-resolved non-refractory sub-micron aerosol chemical species, including organics (Org), sulfate (SO_4^{2-}), nitrate (NO_3^{-}), ammonium (NH_4^{+}), and chloride (Cl⁻), were measured by the HR-AMS. The sampled air dried by diffusion silica gel dryers was drawn into the HR-AMS through a PM_{2.5} cyclone inlet to remove coarse particles larger than 2.5 µm. The HR-AMS was calibrated with pure ammonium nitrate following the procedures detailed in Jimenez et al. (2003). Sun et al. (2016b) provide operational details about the HR-AMS.

154 **3.** Method

155 **3.1.** ALWC calculation based on H-TDMA measurements

The ALWC is calculated based on measurements of the aerosol GF and particle number size distribution. Briefly, H-TDMA data are first used to derive the size-resolved particle GFs at various RHs. Then ALWC is calculated as the increased aerosol volume due to hygroscopic growth attributed to water uptake.

160 Chen et al. (2012) showed how to calculate size-resolved particle GFs at different RHs. First, a 161 three-mode log-normal distribution is applied to fit the measured PNSD to produce fitting parameters 162 for each mode. The hygroscopicity parameter (κ) in any mode is assumed to be constant. The H-163 TDMA-derived size-dependent κ can then be used to deduce the corresponding κ for the nucleation 164 mode, the Aitken mode, and the accumulation mode of PNSDs according to the following equation:

165
$$\kappa(D_P) = \frac{\sum_{i=1}^{3} \kappa_i \cdot N_i(D_P)}{\sum_{i=1}^{3} N_i(D_P)},$$
 (1)

where κ_i refers to the κ of the *i*th mode, and $N_i(D_p)$ refers to the number concentration of particles in the *i*th mode. According to κ -Köhler theory (Petters and Kreidenweis, 2007), κ at a certain diameter (D_d) can be calculated as

169
$$\kappa(D_{\rm d}) = ({\rm GF}^3 - 1) \cdot \left[\frac{1}{{\rm RH}} \exp\left(\frac{4\sigma_{\rm s/a}M_{\rm w}}{RT\rho_{\rm w}D_{\rm d}{\rm GF}}\right) - 1\right], \tag{2}$$

170 where RH is the control value by the humidifier in the H-TDMA system, T is the mean room

temperature of the container set to 293 K, $\sigma_{s/a}$ is the surface tension of the solution/air interface assumed to be the same as the surface tension coefficient between water and air (about 0.0728 N m⁻¹ at 293 K), M_w is the molecular weight of water, R is the universal gas constant, ρ_w is the density of water, and D_d is the diameter of the dry particles. The GF at a given RH is defined as the ratio of the humidified diameter [D_p (RH)] to D_d :

176

$$GF = D_{p}(RH)/D_{d}.$$
 (3)

177 The known κ of each mode derives the size-resolved κ at 90 % RH using Eq. (1). Substituting the size-178 resolved κ into Eq. (2) results in size-resolved GFs at various RHs. Finally, the volume of ALWC at 179 ambient RH is equal to the increased aerosol volume due to water uptake, i.e., ALWC can be calculated 180 as

181
$$ALWC_{HTDMA} = \left[\frac{1}{6}\sum_{i} n_{i} D_{p,i}^{3} \left(GF(D_{p,i}, RH)^{3} - 1\right)\right] \cdot \rho_{w} , \qquad (4)$$

where n_i refers to the particle number concentration of dry particles for the corresponding particle size range in the *i*th mode, and $D_{p,i}$ refers to the particle diameter for the corresponding particle size range.

184 **3.2.** ALWC simulations based on the ISORROPIA II model

The thermodynamic equilibrium model ISORROPIA II developed by Fountoukis and Nenes 185 (2007) using aerosol chemical composition information from the HR-AMS can simulate ALWC 186 (ALWC_{ISO}). The bulk chemical composition was used in the model. However, the ISORROPIA II 187 model only considers the contribution of inorganic species (Ca²⁺, K⁺, Mg²⁺, NH⁺, Na⁺, SO₄²⁻, NO₃⁻, 188 Cl⁻ and H₂O) on ALWC and neglects the contribution of organics. In this study, the model was set up 189 to reverse mode due to the lack of measurements of gaseous ammonia, and the phase state was assumed 190 to be stable in the model calculation. As shown in Figure S1, the simulated ALWC in stable mode is 191 similar to that in metastable mode ($R^2 = 0.99$). 192

According to the model assumptions that the aerosol curvature effect in Köhler theory is ignored, and the aerosol water uptake has no effect on ambient vapor pressure, the water activity (a_w) defined as the effective mole fraction of water is equal to the ambient RH in this model (Seinfeld and Pandis,2006):

197
$$a_w = RH \tag{5}$$

The ALWC can be calculated using the Zdanovskii–Stokes–Robinson (ZSR) mixing rule (Stokes andRobinson, 1966),

200
$$ALWC_{ISO} = \sum_{i} \frac{M_i}{m_{0i}(a_w)}, \qquad (6)$$

where M_i is the mole concentration of the *i*th species (mol m⁻³ in air), and m_{0i} (a_w) is the corresponding molality of the binary solution of the *i*th species under the same a_w with complex solution. Finally, with measured ambient RH and *T* values as input ALWC_{ISO} values under different RH and *T* conditions can be derived.

205 **3.3.** Inferring the contribution of organics to ALWC

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According to the κ -Köhler theory and the ZSR mixing rule, κ can also be expressed as the sum of the contributions of each aerosol component:

 $\kappa = \sum_{i} \varepsilon_{i} \kappa_{i} , \qquad (7)$

where ε_i and κ_i are the volume fraction and hygroscopicity of the *i*th species, respectively. Submicron aerosols mainly consist of organic and inorganic species (Carbone et al., 2013; Zieger et al., 2017). As mentioned in section 2.2, the HR-AMS measures the mass concentrations of organics and inorganics, including SO₄²⁻, NO₃⁻, NH₄⁺, and Cl⁻. The volume fraction of inorganic species can be calculated based on the ion-pairing scheme given by the following equations (Gysel et al., 2007):

214
$$n_{\rm NH_4NO_3} = n_{\rm NO_3^-}$$

215
$$n_{\rm NH_4HSO_4} = \min(2n_{\rm SO_4^{2-}} - n_{\rm NH_4^+} + n_{\rm NO_3^-}, n_{\rm NH_4^+} - n_{\rm NO_3^-}),$$

216
$$n_{(\rm NH_4)_2SO_4} = \max(n_{\rm NH_4^+} - n_{\rm NO_3^-} - n_{\rm SO_4^{2-}}, 0),$$

217
$$n_{H_2SO_4} = \max(0, n_{SO_4^{2-}} - n_{NH_4^+} + n_{NO_3^-}),$$

218
$$n_{\rm HNO_3} = 0,$$
 (8)

where *n* represents the mole numbers, and "min" and "max" are minimum and maximum values, 219 220 respectively. The κ values of the inorganic species sulfuric acid, ammonium sulfate, ammonium hydrogen sulfate, and ammonium nitrate are 1.19, 0.48, 0.56, and 0.58, respectively (Topping et al., 221 2005; Petters and Kreidenweis, 2007). So the ZSR model can be used to estimate the contribution of 222 inorganic species to the κ value. In this paper, the chloride was not taken into account in ion-pairing 223 because its source is hard to determine. This may result in a minor uncertainty in κ calculation. The 224 hygroscopicity parameter of organics (κ_{Org}) can be calculated using the volume fraction of organics 225 and the total κ value derived from the H-TDMA, according to Eq. (7). Finally, the ALWC contributed 226 by organic species (ALWC_{Org}) can be calculated as (Petters and Kreidenweis, 2007) 227

228
$$ALWC_{Org} = \frac{m_{Org}\rho_W}{\rho_{Org}} \frac{\kappa_{Org}}{\left(\frac{1}{BH} - 1\right)},$$
(9)

where m_{Org} is the organic mass concentration from the AMS (Xu et al., 2015), and ρ_{Org} is the density of organics, taken as 1.4 g cm⁻³ (Moore et al., 2011; Lathem et al., 2013; Cerully et al., 2014).

231 4. Results and discussion

4.1. Comparison of calculated and simulated ALWC

The trends in ALWC calculated based on the hygroscopic growth factor and PNSD 233 (ALWC_{HTDMA}) and simulated from ISOPPOPIA II model (ALWC_{ISO}) are generally consistent. Figure 234 1a shows that ALWC_{HTDMA} and ALWC_{ISO} agree well and that their coefficient of determination (\mathbb{R}^2) 235 is 0.89. The correlation is especially strong for RH over 90 %. However, for RH below 60 %, ALWC_{ISO} 236 is less than ALWC_{HTDMA} and even close to 0 in some cases. Bian et al. (2014) and Tan et al. (2017) 237 238 observed a similar phenomenon in northern and southern China. There are three possible explanations for these results. H-TDMA samples were humidified to 90 % RH during the field campaign, thereby 239 leading to the neglect of the deliquescence process in the ALWC_{HTDMA} calculation. This may lead to 240

overestimation of ALWC_{HTDMA} for RH below the deliquescence relative humidity (DRH). Second, the 241 assumption of constant κ in each mode may lead to small uncertainty in size-resolved GFs at different 242 RHs, resulting in small deviation in ALWC calculation. This may be another reason for the difference 243 between ALWCHTDMA and ALWCISO. In addition, the ISORROPIA II model ignores the effect of 244 aerosol shape and complex organic species on the DRH. Previous studies have suggested that the 245 particle spherical assumption and simplified aerosol chemical species in this model can overestimate 246 247 the DRH (Seinfeld and Pandis, 2006; Sjogren et al., 2007). So for RH below the simulated DRH (~60 %), particles may still be dry in the ISORROPIA II model, but may have been hydrated in the 248 249 real atmosphere. Therefore, this model underestimates ALWC. The ambient aerosol deliquescent phenomenon is rare in the North China Plain (Kuang et al., 2016). In addition, the ISORROPIA II 250 model cannot simulate water uptake by organics, which can lead to some bias between simulated and 251 calculated ALWCs. As described in section 3.3, ALWCorg can be inferred and used to discuss 252 differences between ALWC_{ISO} and ALWC_{HTDMA}. Figure 1b shows that adding ALWC_{Org} to ALWC_{ISO} 253 leads to a stronger correlation with ALWC_{HTDMA} ($R^2 = 0.92$). The correlation improves significantly 254 for RH below 60 %. This demonstrates that (1) organic species contribute significantly to ALWC, and 255 (2) the underestimation of ALWC by the ISORROPIA II model is also related to the neglect of organic 256 species in the model. 257

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259 4.2. Impact of different factors on ALWC

260 4.2.1. Impact of aerosol chemical species on ALWC

Figure 2 shows the characteristics of seven heavy pollution events selected for examination. Figures 2a and 2c display the time series of WS, WD, and ambient RH. The prevailing wind during the haze episodes was a weak southerly wind that was favorable for bringing in pollutants from the highly populated and industrialized neighboring regions to the sampling site. This is beneficial to the formation and accumulation of SA (T. Wang et al., 2010; Y. Wang et al., 2017). However, the prevailing winds during the clean events were strong northerly winds that always carried in a clean air mass, resulting in pollutants being quickly removed (Figure 2c). Note that the PM₁ mass concentration decreases somewhat in the evening during haze episodes, following the short-term change of WD from southerly to northerly. This is related to mountain-valley breezes in Beijing (Wehner et al., 2008; Gao et al., 2011; Y. Wang et al., 2017). These results demonstrate that heavy haze episodes have a strong correlation with local wind direction in Beijing.

Figures 2a and 2d show the time series of ambient RH and mass concentrations of aerosol chemical 272 273 species in PM₁. These figures suggest that the increase in inorganic and organic aerosols is synchronous with the increase in ambient RH during the heavy pollution periods (P1-P7). This is likely 274 because of a positive feedback mechanism driven by Henry's law and thermodynamic equilibrium (Z. 275 Wu et al., 2018). Figure 2b also shows that ALWC continuously increases during the pollution 276 accumulation period. On average, ALWC increases from 8 to 89 µg m⁻³ as ambient RH increases from 277 15 to 80 %, and the inorganic and organic aerosol mass concentrations increase from 15 to 120 μ g m⁻ 278 ³ and from 12 to 78 µg m⁻³, respectively. These results imply that the increase in ambient RH and 279 aerosol mass concentration are all important for the increase in ALWC. 280

Equation (4) also suggests that the absolute value of ALWC is dependent on the value of ambient 281 RH and aerosol chemical composition (i.e., the GF value). To further investigate the impact of 282 chemical composition on ALWC, the impact of RH on ALWC should be accounted for. Previous 283 studies suggest there is an exponential relationship between ALWC and RH (e.g., Z. Wu et al., 2018). 284 Here, we define the relative ALWC as the ratio of ALWC_{HTDMA} and the function of ambient RH (e^{bRH}). 285 The b is derived according to the relationship between ALWC_{HTDMA} and RH that is fitted by the 286 function $y = ae^{bx}$. Figure 3a shows the relationship between relative ALWC and primary aerosols 287 (PA) or SA mass concentrations. PA consists of primary organic aerosols (POA) and black carbon 288 (BC), and SA consists of SOA, sulfate, and nitrate. The positive matrix factorization (Paatero and 289

Tapper, 1994) was applied on the organic aerosols (OA) spectral matrices to identify POA and SOA. 290 The relative ALWC is highly correlated with SA mass concentrations ($R^2 = 0.94$) but poorly correlated 291 with PA mass concentrations ($R^2 = 0.69$). High relative ALWCs coincident with high SA mass 292 concentrations suggest that SA plays a key role in the increase in ALWC. This is likely because SA is 293 mainly generated from photochemical reactions in the daytime or reactions at night, making SA highly 294 aged with a hygroscopicity stronger than that of PA (Ervens et al., 2011; Sareen et al., 2017). SA can, 295 296 therefore, absorb more water vapor than PA in the atmosphere. The enhanced aerosol liquid water induced by SA is further favorable for the formation of SA by speeding up the atmospheric chemical 297 298 reaction rate and serving as the medium for gas-particle heterogeneous reactions (G. Wang et al., 2016; Cheng et al., 2016). This further increases the bulk aerosol hygroscopicity. This is also the reason why 299 inferred κ based on the ZSR model continuously increases during haze episodes (Figure 2c). 300

Secondary aerosols are mainly composed of nitrate, sulfate, ammonium and SOA. To determine 301 which species is the driver for ALWC in Beijing, Figure 3b shows the correlation analysis between 302 relative ALWC and the mass concentrations of different aerosol chemical species. Relative ALWC and 303 all SIA agree well [R^2 equal to 0.66 (sulfate) and 0.56 (nitrate)]. It has been reported that ALWC is 304 driven by inorganic salts with both nitrate and sulfate playing key roles in determining ALWC (Z. Wu 305 et al., 2018). ALWC also agrees well with SOA ($R^2 = 0.60$) in our study. This is unexpected because 306 the hygroscopicity of SOA is relatively lower than that of nitrate and sulfate. Some studies have also 307 suggested that the water uptake of aged organics accounts for only a few percent of the total aerosol 308 water uptake (e.g., Gysel et al., 2007; Engelhart et al., 2011). In our study, the contribution of ALWC_{Org} 309 to total ALWC is significant, accounting for 30 $\% \pm 22$ %. As shown in Figure 4, the contribution of 310 organics to total ALWC varies strongly. This is likely related with the variation in mass fraction and 311 hygroscopicity parameter of organics (κ_{org}). The mass concentration of inorganics increases more than 312 that of organics as RH increases, leading to a lower mass fraction of organics in the case of high 313 ambient RH. Figure 4 also shows ALWC_{Org} fraction increases significantly with the increase of κ_{org} . 314

All these help explain a large variation in the ALWC_{Org} contribution to total ALWC. Considering the 315 distinct ambient RH and κ_{org} between clean and polluted periods, we calculated respectively the 316 317 fraction of ALWC_{Org} during two periods. There is a higher ALWC_{Org} fraction $(33\% \pm 23\%)$ during clean periods than that during polluted periods ($26\% \pm 11\%$). Yet, there is little variability of ALWC_{Org} 318 fraction during polluted periods. The larger variability in ALWC_{Org} fraction during clean periods is 319 320 likely caused by the highly variable $\kappa_{\rm org}$ when the ambient RH is low. In summary, the contribution of organics in total ALWC varies with the variations of the mass fraction of organics and κ_{org} , and this 321 contribution is significant during both clean and polluted periods. Studies of ALWC in Beijing, 322 323 therefore, cannot neglect ALWCorg. This is different from the studies in other regions such as in the Po Valley in Italy (Hodas et al., 2014) and the eastern U.S. (Carlton et al., 2013) where the ALWC was 324 found to be only driven by nitrate and sulfate respectively. 325

An interesting phenomenon is frequently observed at the initial stage of heavy haze episodes (e.g., 326 P4, P5, P6, and P7). ALWC_{ISO} is almost close to 0, but both ALWC_{HTDMA} and ALWC_{Org} are always 327 larger than 0, and the organic aerosol mass fraction is high at this stage. These observations reveal that 328 at the initial stage of heavy haze episodes, the ALWC is mostly contributed by organic species. 329 Meanwhile, κ is not very low and increases markedly as the PM₁ mass concentration increases, which 330 is unexpected because of the lower hygroscopicity of organic aerosols compared to SIA. Therefore, 331 some highly hygroscopic substance (i.e., SA) must be generated through multiphase chemical reaction 332 at this stage. We propose that the liquid water contributed by organic species provides a reactor for the 333 transformation of gaseous precursors to SA at the initial stages of heavy haze episodes, increasing the 334 uptake of more liquid water by more SA and further accelerating the formation of heavy haze. Section 335 4.3 provides a case study to demonstrate this. 336

337 4.2.2. Impact of PNSD on ALWC

In addition to aerosol chemical composition, ALWC also depends on PNSD (Bian et al., 2014).
PNSD does not influence ALWC directly, but it's an important factor to determine the mass

concentration and the distribution of hygroscopic materials. As described in section 3.1, the nucleation mode (< 30 nm), the Aitken mode (30–110 nm), and the accumulation mode (110 nm to1 μ m) (Whitby, 1978; Birmili et al., 2001) are considered in this study. Particles with diameters greater than 1 μ m are not considered because some particles in the coarse mode are water soluble but their contribution on the ALWC is low (e.g., Hussein et al., 2004; S. Liu et al., 2008; Bian et al., 2014; Tan et al., 2017).

The contributions of nucleation mode, Aitken mode, and accumulation mode particles to 345 ALWC_{HTDMA} are about 1.0 %, 18.0 %, and 82.0 %, respectively. Figure 5 shows the correlations 346 between ALWC_{HTDMA} and the volume concentrations of different mode particles, and the average 347 348 contribution of different mode particles to ALWC_{HTDMA} (f_{ALWC}) under five different RH conditions. The R^2 and f_{ALWC} of the nucleation mode particles (left column in Figure 5) are all less than 0.1 and 349 1 %, respectively, under all RH conditions. This is likely because the volume concentration of 350 nucleation mode particles is very low, and most of these small particles are composed of hydrophobic 351 chemical species such as BC and POA. Similarly, the number concentration of Aitken mode particles 352 also shows weak correlations with ALWC_{HTDMA} ($R^2 < 0.2$) under RH < 90 % conditions, but their 353 correlation ($R^2 = 0.25$) is enhanced significantly under RH > 90 % conditions (middle column of Figure 354 5). This is because there are more aged particles in the Aitken mode which can absorb much more 355 water when the ambient RH is higher than 90 %. However, the contribution of the Aitken mode to 356 ALWC ranges from 14 % to 21 % and decreases as RH increases. ALWC_{HTDMA} is strongly correlated 357 to the volume concentration of accumulation mode particles, with R^2 and f_{ALWC} greater than 0.6 and 358 75 %, respectively, under all RH conditions (right column of Figure 5). Figure 5 also shows that ALWC 359 increases slightly as the volume concentration of accumulation mode particles increases under RH < 360 70 % conditions (slope < 0.001), but increases strongly under higher RH conditions, especially under 361 RH > 90 % conditions (slope = 0.0041). This is likely because there are more accumulation mode SA 362 formed due to multiphase chemical reactions under high ambient RH conditions. Swietlicki et al. 363 (1999) have suggested that the contribution of accumulation mode particles to ALWC is largest for all-364

365 mode particles.

In summary, the contribution of nucleation mode particles to ALWC is very low. The contribution of Aitken mode particles is much higher than nucleation mode particles and decreases with increasing ambient RH. The contribution of accumulation mode particles to ALWC is largest under all RH conditions and increases with increasing ambient RH, thus playing a key role in determining ALWC.

370 4.2.3. Impact of RH on ALWC

As discussed in 4.2.1, the absolute value of ALWC has an exponential relationship with ambient 371 RH. Figure 6 shows the relationship between ALWC and RH for different PM₁ mass concentration 372 ranges. ALWC increases slowly as RH increases under lower ambient RH conditions then sharply 373 increases when RH exceeds a critical RH value. This critical RH value is different for different PM₁ 374 mass concentrations. This is because the low RH conditions cannot provide enough water for aerosol 375 particles, even though the PNSD is dominated by accumulation mode particles with higher 376 hygroscopicity (Tan et al., 2016). This demonstrates the important influence of RH on ALWC. The 377 lower critical RH value for higher PM₁ mass concentrations (~80 %) suggests that ALWC is easily 378 formed under heavily polluted conditions. This is likely because there are more SA and accumulation 379 mode particles during pollution periods (Sun et al., 2016a; Y. Wang et al., 2017). 380

Figure 7a shows the diurnal variations of ALWC_{HTDMA} and ambient RH during the sampling 381 period. The extreme ALWC_{HTDMA} values appear at night and during the day respectively, likely related 382 to the diurnal variations of ambient RH. The elevated ambient RH at night not only increases ALWC 383 through water uptake of particles directly, but also facilitates the formation of hydrophilic particulate 384 nitrate through the speeding up of the uptake coefficient of N₂O₅ (Thornton et al., 2003; Bertram et 385 al., 2009). This can further enhance ALWC. However, although the diurnal variations of ALWC_{HTDMA} 386 387 and ambient RH are similar, the peak and nadir of ALWC_{HTDMA} (0300 LT and 1100 LT, respectively) appear three hours earlier than the peak and nadir of ambient RH (0600 LT and 1400 LT, respectively). 388 This time difference is likely related to changes in PNSD. The diurnal variation of PNSD (Figure 7b) 389

shows that the number concentrations of Aitken and accumulation mode particles begin to decrease 390 quickly at 0300 LT. ALWC also begins to decrease, although the ambient RH increases slightly at that 391 392 time. In the morning, ALWC decreases sharply following the ambient RH and PNSD changes due to the lifting planetary boundary layer height. ALWC decreases to its minimum value at ~1100 LT then 393 begins to increase quickly. However, ambient RH still decreases at that time and reaches its minimum 394 395 value at ~1400 LT. The increase in ALWC is likely associated with changes in aerosol chemical species and PNSD. Figure 7b and 7c show that there are many newly formed Aitken and accumulation mode 396 particles and that the fraction of SA increases at noon, likely because of strong photochemical 397 398 reactions. Y. Wang et al. (2017, 2018) have suggested that daytime photochemical reactions are efficient enough to enhance aerosol hygroscopicity and change the aerosol mixing state from external 399 to internal in North China through the formation of hydrophilic chemical species. All this suggests that 400 ambient RH is not the only determining factor for ALWC. PNSD and aerosol chemical composition 401 are also important for ALWC. 402

403 4.3. A case study of the impact of ALWCorg on SA formation

As discussed in section 4.2.1, a hypothesis is proposed that ALWC_{Org} maybe provide a reactor for 404 the formation of secondary species. To verify this hypothesis, the P4 case shown in Figure2 is selected 405 to further analyze the influence of ALWC_{Org} on the formation of secondary aerosols (Figure 8). Figure 406 8a shows the time series of ALWC_{HTDMA}, ALWC_{ISO}, and ALWC_{Org} during this case. On 27 November 407 2017, ALWC_{ISO} was close to 0 all day long because of the low ambient RH, but both ALWC_{HTDMA} and 408 ALWC_{Org} were always larger than 0, increasing with increasing PM₁ mass concentration (Figure 8a). 409 Figure 8a also shows that the fraction of ALWC_{Org} in ALWC_{HTDMA} was high at the initial stage of this 410 pollution case, but this fraction decreased as haze increased. This case was further divided into three 411 412 periods (Figure 8b). Organics were the most abundant chemical species during the first period (64 %), which explains the high fraction of ALWC_{Org} in ALWC_{HTDMA} at the initial stage of this haze case. The 413 pie charts in Figure 8b also show that both SOA and SIA (sulfate, nitrate, and ammonium) increases 414

from the first to third periods but POA decreases, likely related to multiphase reactions (i.e., aqueousphase reactions) due to the enhanced ALWC. Time series of f_{44} and the fraction of sulfate in total sulfur (F_{SO4}^{2-}) are also shown to further illustrate the influence of aqueous-phase reactions on aerosol chemical species. The m/z 44 signal intensity f_{44} (mostly contributed by the CO₂⁺ ion) measured by the AMS can be used as an indicator of the oxidation level in organic species (Mei et al., 2013). The sulfur oxidation ratio F_{SO4}^{2-} (Sun et al., 2006) is defined as

$$F_{\mathrm{SO}_4^{2-}} = \frac{n[\mathrm{SO}_4^{2-}]}{n[\mathrm{SO}_4^{2-}] + n[\mathrm{SO}_2]},\tag{10}$$

where $n[SO_4^{2-}]$ and $n[SO_2]$ refer to the molar concentrations of SO₄²⁻ and SO₂, respectively. Figure 8b 422 suggests that f_{44} and F_{SO4}^{2-} both increase gradually with increasing ALWC_{HTDMA} and PM₁ mass 423 concentration from 27 November to 30 November 2017. This is likely because the increase in ALWC 424 is beneficial to the oxidation of organics and the transformation of SO_2 to SO_4^{2-} , implying the 425 importance of aqueous-phase chemical reaction on haze formation in Beijing. The production of 426 secondary organic and inorganic species can further enhance aerosol hygroscopicity, increasing ALWC 427 in the atmosphere. This positive feedback is the reason behind the rapid formation of heavy haze events 428 in Beijing (G. Wang et al., 2016). A rapid increase in f_{44} and F_{SO4}^{2-} was seen during the first period at 429 night on 27 November (shown by green and red arrows in Fig. 9) when organics contributed the most 430 to ALWC. This suggests that ALWC contributed by organics may have played an important role in the 431 formation of secondary species at the initial stage of the pollution event. 432

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434 5. Conclusions

In this study, the aerosol liquid water content (ALWC) was calculated using the size-resolved aerosol hygroscopic growth factor and the particle number size distribution (PNSD) measured at a Beijing urban site during the APHH winter campaign (8 November to 15 December 2017). Also done were simulations using the ISORROPIA II model with measured aerosol chemical composition data
as input data. During the sampling period, seven heavy haze episodes were selected to investigate the
influence of different factors (PNSD, ambient RH, and aerosol chemical composition) on ALWC.

The calculated and simulated ALWC (ALWC_{HTDMA} and ALWC_{ISO}) agree well (correlation of 441 determination R^2 equal to 0.89). However, ALWC_{ISO} is much lower than ALWC_{HTDMA} for RH below 442 60 %, even approaching zero many times. This deviation is in part attributed to the neglect of the 443 444 contribution of organics to ALWC (ALWC_{Org}) in the ISORROPIA II model, contradicting with previous studies ignoring this contribution. The aerosol hygroscopicity of organics was also derived 445 446 in this study for use in calculating ALWCorg. The sum of ALWCISO and ALWCorg has a higher correlation ($R^2 = 0.92$) with the calculated ALWC (i.e., ALWC_{HTDMA}), especially for RH below 60 %. 447 This implies that organic aerosols are also an important contributor to ALWC. 448

PNSD, ambient RH, and aerosol chemical composition are all found to affect ALWC significantly. 449 Nucleation mode and Aitken mode particles have little influence on ALWC. Accumulation mode 450 particles play a key role in determining ALWC and dominate among all aerosol modes. ALWC is 451 highly related to the relative humidity (RH) when RH exceeds a critical RH value that is different for 452 different PM₁ mass concentrations. ALWC varies diurnally with its extreme values appearing at night 453 and during the day respectively. The diurnal variation of ambient RH explains this. However, there is 454 a three-hour difference between when the extreme ALWC and RH values occur, caused by the diurnal 455 variations in PNSD and aerosol chemical composition. 456

On average, ALWC_{Org} accounts for $\sim 30 \% \pm 22 \%$ of the total aerosol liquid water during the sampling period. This shows the significant contribution of organic species to ALWC. Our results suggest that ALWC is not only driven by inorganic salts but also driven by organics in Beijing. This is different from the results obtained in the Po Valley in Italy (Hodas et al., 2014) and the eastern U.S. (Carlton et al., 2013) where the ALWC is driven by nitrate and sulfate respectively. Finally, one case study was used to study the importance of ALWC_{Org} on multiphase chemical reactions. ALWC_{Org} was

| 463 | found to play an important role in the formation of secondary aerosols by speeding up aqueous-phase |
|------------|---|
| 464 | reactions at the initial stage of heavy haze. Our study is important for investigating the contribution of |
| 465 | organics to ALWC and its importance on haze formation in Beijing. |
| 466 | |
| 467 | Data availability. Data used in the study are available from the first author upon request |
| 468 | (<u>201631490012@mali.bnu.edu.cn</u>). |
| 469 | |
| 470 | Author contributions. ZL and YW designed the experiment; YW, XJ, and WX carried it out and |
| 471 | analyzed the data; other co-authors participated in science discussions and suggested additional |
| 472 | analyses. XJ and YW prepared the paper with contributions from all co-authors. |
| 473 | |
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727





Figure 1. Comparison between ALWC_{HTDMA} and (a) ALWC_{ISO} and (b) the sum of ALWC_{ISO} and ALWC_{Org}.
ALWC_{HTDMA} refers to calculated ALWC based on the measured growth factor and PNSDs, ALWC_{ISO} refers to
simulated ALWC from the ISORROPIA II model, and ALWC_{Org} refers to the inferred ALWC contributed by
organic species. The coefficient of determination R² is given in each panel. The color of the dots denotes the ambient
RH; the solid line denotes the 1:1 line.

735





Figure 2. Time series of (a) wind speed (WS, left y-axis), ambient relative humidity (RH, right y-axis), and wind

direction (WD, colored dots), (b) ALWC_{HTDMA} (in blue), ALWC_{ISO} (in red), and ALWC_{org} (in green), (c) PM₁ mass concentration (left y-axis) and hygroscopicity parameter (*k*, right y-axis) calculated using the ZSR model described by Eq. (7), and (d) mass concentrations of aerosol species in PM₁ (left y-axis) and organic aerosol mass fraction (right y-axis). Seven polluted episodes (segments of the time series with a purple background) are selected for examination.



744

Figure 3. The correlation analysis between relative ALWC and (a) primary (in black) and secondary (in red) aerosol mass concentrations, and (b) nitrate, sulfate, and secondary organic aerosol (SOA) mass concentrations. Panel (a) shows mean relative ALWCs (solid dots) with boxes showing the 25th, 50th, and 75th percentiles. The extremities show the 5th and 95th percentiles. The solid lines in (a) and the dashed lines in (b) both represent the corresponding best-fit lines from linear regression.



Figure 4. The variation of the fraction of ALWC_{0rg} in total ALWC (ALWC_{HTDMA}) with the ambient relative humidity (RH). The color of the dots denotes the hygroscopicity parameter of organics (κ_{0rg}). The boxes show the fraction of ALWC_{0rg} with the 25th, 50th, and 75th percentiles. The extremities show the 5th and 95th percentiles. The red line shows the fitting curve with the function $y = ae^{bx}$.



Figure 5. The correlation analysis between ALWC_{HTDMA} and the volume concentration of nucleation mode (left column), Aitken mode (middle column), and accumulation mode (right column) particles under different ambient relative humidity (RH) conditions. The average contribution of each mode particles to ALWC under different ambient RH conditions is denoted by *f*_{ALWC}. The red lines represent the best-fit lines from linear regression.



Figure 6. The dots show how ALWC_{HTDMA} varies with the ambient relative humidity (RH) for different PM₁ mass concentration ranges (colored symbols). The colored curves represent the best-fit lines through the data using the fitting function $y = ae^{bx}$.



Figure 7. Diurnal variations of (a) ALWC_{HTDMA} (in blue) and ambient RH (in red), (b) particle number size
distribution, and (c) the mass fraction of different chemical species. The time is in Beijing time.



770

Figure 8. Time series of (a) ALWC_{HTDMA} (in blue), ALWC_{ISO} (in red), ALWC_{Org} (in green), and RH (right y-axis), and (b) the sulfur oxidation ratio (F_{SO4}^{2-}), f_{44} , and PM₁ mass concentration (right y-axis) during the P4 case in Figure 2. The pie charts in (b) represent the average chemical compositions of PM₁ during three stages of the pollution event (denoted by brown horizontal lines). The red and green arrows in (b) indicate the rapid increase in F_{SO4}^{2-} and f_{44} at the initial stage.