1	Distinct diurnal variation of organic aerosol hygroscopicity and its relationship with
2	oxygenated organic aerosol
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24 Abstract

25 The hygroscopicity of organic aerosols (OA) is important for investigation of its climatic and environmental impacts. However, the hygroscopicity parameter κ_{OA} remains poorly characterized, 26 especially in the relatively polluted environment on the North China Plain (NCP). Here we conducted 27 28 simultaneous wintertime measurements of bulk aerosol chemical compositions of PM2.5 and PM1, and bulk aerosol hygroscopicity of PM₁₀ and PM₁ on the NCP using a capture vaporizer time-of-flight 29 aerosol chemical speciation monitor (CV-ToF-ACSM) and a humidified nephelometer system which 30 measures aerosol light scattering enhancement factor f(RH). A method for calculating κ_{OA} based on 31 f(RH) and bulk aerosol chemical composition measurements was developed. We found that κ_{OA} 32 varied in a wide range with significant diurnal variations. The derived κ_{OA} ranged from almost 0.0 to 33 0.25 with an average ($\pm 1\sigma$) of 0.08 (± 0.06) for the entire study. The derived κ_{0A} was highly correlated 34 with f_{44} (fraction of m/z 44 in OA measured by CV-ToF-ACSM), an indicator of oxidation degree of 35 OA (*R*=0.79), and the relationship can be parameterized as $\kappa_{OA} = 1.04 \times f_{44} - 0.02 (\kappa_{OA} = 0.3 \times \text{O:C} - 0.02)$ 36 0.02, based on the relationship between f₄₄ and O/C ratio for CV-ToF-ACSM). On average, κ_{0A} 37 reached the minimum (0.02) in the morning near 07:30 and then increased rapidly reaching the peak 38 value of 0.16 near 14:30. The diurnal variations of κ_{OA} were highly and positively correlated with 39 those of mass fractions of oxygenated OA (R = 0.95), indicating that photochemical processing played 40 a dominant role for the increase of κ_{OA} in winter on NCP. Results in this study demonstrate the 41 potential wide applications of humidified nephelometer system together with aerosol composition 42 measurements for investigating the hygroscopicity of OA in various environments, and highlight that 43 the parameterization of κ_{OA} as a function of OA aging processes needs to be considered in chemical 44 transport models for better evaluating the impacts of OA on cloud formation, atmospheric chemistry 45 and radiative forcing. 46

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49 **1 Introduction**

50 Aerosol hygroscopic growth plays significant roles in different atmospheric processes including 51 atmospheric radiation transfer, cloud formation, visibility degradation, atmospheric multiphase 52 chemistry and even air-pollution-related health effects, and therefore is crucial for studies on aerosol

climatic and environmental impacts. Organic materials in ambient aerosol particles, usually referred 53 to organic aerosols (OA), contribute substantially to ambient aerosol mass and often contribute more 54 than half to submicron aerosol particle mass under dry state (Jimenez et al., 2009). The hygroscopicity 55 parameter κ (Petters and Kreidenweis, 2007) of organic aerosols (κ_{OA}) is a key parameter for 56 investigating the roles of organic aerosol in radiative forcing, cloud formation and atmospheric 57 chemistry. Liu and Wang (2010) demonstrated that 50% increases in κ of secondary organic aerosol 58 59 (0.14 ± 0.07) can result in up to 40% increases in predicted cloud condensation nuclei (CCN) concentration. Rastak et al. (2017) reported that global average aerosol radiative forcing would 60 decrease about 1 W/m^2 should κ_{OA} increase from 0.05 to 0.15, which is of the same order with the 61 overall climate forcing of anthropogenic aerosol particles during the industrialization period. Li et al. 62 (2019) reported that organic aerosol liquid water contributed 18-32% to total particle liquid water 63 content in Beijing. Despite its importance, κ_{OA} has not yet been well characterized due to the 64 65 extremely complex chemical compositions of organic aerosol. Therefore, it is important to conduct more researches on the spatiotemporal variation of κ_{OA} and its relationship with aerosol chemical 66 67 compositions to reach a better characterization and come up with more appropriate parameterization schemes in chemical, meteorological and climate models. 68

The large variety in OA chemical constituents makes it difficult to directly link κ_{OA} to specific 69 organic aerosol compositions. The OA chemical composition is tightly connected to their volatile 70 71 organic precursors, which are also rich in variety and come from different natural and anthropogenic sources. OA with different oxidation levels also behave differently in respect to hygroscopic growth. 72 Thus, studies on κ_{OA} at different locations and seasons have reported distinct characteristics. Many 73 studies have investigated the influence of OA oxidation level (represented by O:C ratio or fraction of 74 m/z 44 in OA, f₄₄, which are determined from aerosol mass spectrometer measurements) on its 75 hygroscopicity (Chang et al., 2010;Lambe et al., 2011;Duplissy et al., 2011;Mei et al., 2013b;Wu et 76 al., 2013;Hong et al., 2015;Chen et al., 2017;Massoli et al., 2010) and found that the average κ_{OA} 77 generally increases as a function of organic aerosol oxidation level. However, the statistical empirical 78 79 relationship between κ_{OA} and O:C ratio or f₄₄ differs much among different studies. Several studies 80 have also analyzed the diurnal variation characteristics of κ_{OA} at different locations and seasons (Cerully et al., 2015;Bougiatioti et al., 2016;Deng et al., 2018;Deng et al., 2019;Thalman et al., 2017), 81 with some exhibiting distinct diurnal variations (Deng et al., 2018; Deng et al., 2019; Bougiatioti et al., 82

⁸³ 2016) and others not (Cerully et al., 2015). Despite this, the studies on κ_{OA} in the relatively polluted ⁸⁴ North China Plain (NCP) region are very limited (Wu et al., 2016). The diurnal characteristics of κ_{OA} ⁸⁵ on the NCP have not been reported so far. Therefore, more investigation into the diurnal variation of ⁸⁶ κ_{OA} and its relationship with OA oxidation level is required to better understand its characteristics in ⁸⁷ the NCP.

The Humidity Tandem Differential Mobility Analyzer (HTDMA) or CCN counter has been 88 widely used for aerosol hygroscopicity measurements. Both the HTMDA and size-resolved CCN 89 measurements can only be used to derive a κ within a certain size range (HDTMA: usually diameter 90 below 300 nm, with a reported highest diameter of 360 nm (Deng et al., 2019), CCN: with diameter 91 up to ~200 nm (Zhang et al., 2014; Rose et al., 2010)). The aerosol particles contributing most to aerosol 92 optical properties (Bergin et al., 2001; Quinn et al., 2002; Cheng et al., 2008; Ma et al., 2011; Kuang et 93 al., 2018) and aerosol liquid water content (Bian et al., 2014) in continental regions are usually in the 94 95 diameter range of 200 nm to 1µm, which the HTDMA and CCN hygroscopicity measurements cannot represent. Results from several studies have reported that κ_{OA} usually differentiates among particle 96 97 size (Frosch et al., 2011;Kawana et al., 2016;Deng et al., 2019). For example, Deng et al. (2019) found that κ_{OA} increases with the increases in particle dry diameter. These results further highlight a need 98 for characterization of κ_{OA} of larger particles. 99

The humidified nephelometer system which measures aerosol light scattering enhancement 100 factors is also widely used in aerosol hygroscopicity research (Titos et al., 2016). The hygroscopicity 101 parameter k retrieved from measured light scattering enhancement factor is usually referred to as 102 $\kappa_{f(RH)}$ (Chen et al., 2014; Kuang et al., 2017), which represents the overall hygroscopicity of aerosol 103 particles with their diameters ranging from 200 nm to 800 nm for continental aerosols (see discussions 104 in Sect.3.3 for physical understanding of $\kappa_{f(RH)}$). Using the retrieved $\kappa_{f(RH)}$ together with bulk 105 aerosol chemical compositions of PM_1 (particulate matter with aerodynamic diameter less than 1 μm_1 , 106 corresponding to mobility diameter of approximately760 nm assuming spherical particles and a 107 particle density of 1.7 g/cm³), κ_{OA} can be derived, representing the hygroscopicity of organic aerosol 108 particles in the diameter range of 200 to about 800 nm. In this study, the light scattering enhancement 109 110 factors of both PM₁₀ (particulate matter with aerodynamic diameter less than 10 µm) and PM₁ particles were measured at a rural site on the NCP in winter 2018, together with aerosol chemical 111 compositions by a capture vaporizer time-of-flight aerosol chemical speciation monitor (CV-ToF-112

113 ACSM). By integrating these two different measurements, κ_{OA} is derived, and the relationship 114 between κ_{OA} and OA oxidation degree, as well as the diurnal variation of κ_{OA} is elucidated.

115 **2 Site and instruments**

From 11th November to 24th December 2018, continuous measurements of physical, optical and chemical properties of ambient aerosol particles as well as meteorological parameters such as temperature, wind speed and direction and relative humidity were made at the Gucheng site in Dingxing county, Hebei province, China. The sampling site, an Ecological and Agricultural Meteorology Station (39°09'N, 115°44'E) of the Chinese Academy of Meteorological Sciences, is located between Beijing (~ 100 km) and Baoding (~40 km), two large cities on the North China Plain, and is surrounded by farmland and small residential towns.

123 2.1 Inlet system and instruments

During this field campaign, all instruments were housed in an air-conditioned container, with the 124 temperature held almost constant near 24 °C. The schematic diagram of the inlet systems for the 125 aerosol sampling instruments is displayed in Fig.1. Three inlet impactors are used for aerosol sampling, 126 two PM₁₀ inlets and one PM₁ inlet, respectively sampling ambient aerosol particles with aerodynamic 127 diameter less than 10 µm and 1 µm. Nafion driers with lengths of 1.2 m were placed downstream of 128 each PM impactor inlet, which can drop the RH of sampled air below 15%, thus, sampled aerosol 129 particles can be treated as in dry state. Additionally, downstream every PM impactor inlet an MFC 130 (mass flow controller) and a pump were added for automatic flow compensation, to ensure that each 131 impactor reaches their required flow rate of 16.7 L/min and guaranteeing for the right cut diameters. 132

Aerosol sampling instruments can be categorized into four groups according to their inlet routes. The first group (group1) downstream of the first PM_{10} inlet is comprised of only one instrument, the Aerodynamic Particle Sizer (APS, TSI Inc., Model 3321), measuring the size distribution of ambient aerosol particles with aerodynamic diameter ranging from 700 nm to 20 µm at a temporal resolution of 20 seconds. The second group (group 2) includes a humidified nephelometer system (consisting of two nephelometers and a humidifier) that measures aerosol optical properties (scattering and back



Figure 1. Schematic diagram of the inlet systems for aerosol sampling instruments

scattering coefficients at three wavelengths: 450 nm, 525 nm, 635 nm) of ambient aerosol particles in dry state (DryNeph) and under 85% RH condition (WetNeph). The third group (group3) includes two instruments, an ACSM and a scanning mobility particle sizer (SMPS; TSI model 3080). The CV-ToF-ACSM measures non-refractory particulate matter (NR-PM) species including organics, SO_4^{2-} , NO_3^- , NH_4^+ , and Cl^- with an air flow of 0.1 L/min and a time resolution of 2 minutes. Since the CV-ToF-ACSM instrument comes with a PM_{2.5} impactor, when the impactor of upstream is PM₁₀, chemical compositions of PM_{2.5} rather than of PM₁₀ were measured.

The SMPS measures particle mobility diameter size distributions with a diameter range of 12 nm to 760 nm. The inlets of group2 and group3 switch every 15 minutes, as denoted by the dashed and solid red lines in Fig.1, enabling the instruments of these two groups to alternately measure the chemical and optical properties of PM_{10} and PM_1 . The fourth group (group4) includes an AE33 aethalometer (Drinovec et al., 2015) and other aerosol instruments. Due to technical issues with the humidifier, the humified nephelometer system started to operate continuously since the 30th Nov.

152 **2.2** The humidified nephelometer system

The humidified nephelometer system we built was set up to measure dry state aerosol optical 153 properties at a fixed RH of 85%. The RH of the air sample is increased by a humidifier that consists 154 of two layers. The inner layer is a Gore-Tex tube layer passing through sampled air, while the outer 155 156 layer is a stainless-steel tube with circulating liquid water. The water vapor penetrates through the Gore-Tex tube and humidifies the sample air, while liquid water is kept from the inner layer by the 157 Gore-Tex material. Upon the switch of inlets between group 2 and group 3, delays in valve switching 158 caused instantaneous low pressure in the sample air, which broke the humidifier with the Gore-Tex 159 tube after four days of continuous operation (3rd, Dec) and flooded the WetNeph. The WetNeph was 160 fixed and recalibrated and a commercial Nafion drier (60 cm long, Perma Pure company) replaced the 161 Gore-Tex tube, which works the same way but is more resistant to low pressure. The temperature of 162 the circulating water layer is controlled by a water bath and specified by an algorithm that adjusts the 163 water temperature to maintain a relatively constant RH in the sensing volume of the WetNeph. To 164 monitor the RH in the sensing volume of the WetNeph, two temperature and RH sensors (Vaisala 165 HMP110, with accuracies of ± 0.2 °C and ± 1.7 % for RH between 0 to 90%, respectively, and 166 ± 2.5 % for RH between 90 to 100%) were placed at the inlet and outlet of the WetNeph. Defining 167 measured RHs/temperatures at the inlet and outlet of the WetNeph as RH_1/T_1 and RH_2/T_2 , the 168 according dew point temperatures T_{d1} and T_{d2} can be calculated and the average value $\overline{T_d}$ was treated 169 as the dew point of the sample air in the sensing volume of WetNeph. The sample RH is calculated 170 using $\overline{T_d}$ and the sample temperature measured by a sensor inside the sample cavity of the 171 nephelometer. 172

173 2.3 ACSM measurements and data analysis

The mass concentration and chemical composition of NR-PM species were measured with the Aerodyne ToF-ACSM which is equipped with a PM_{2.5} aerodynamic lens (Williams et al., 2010) and a

capture vaporizer (CV) (Xu et al., 2017;Hu et al., 2017) to extend the measured particle size to 2.5 µm. 176 Detailed instrument descriptions were given in Fröhlich et al. (2013) and Xu et al. (2017). The CV-177 ToF-ACSM data were analyzed with the standard data analysis software (Tofware v2.5.13, 178 https://sites.google.com/site/ariacsm/) within Igor Pro (v6.37, WaveMetrics, Inc., Oregon, USA). The 179 CV was designed with an enclosed cavity to increase particle collection efficiency (CE) at the detector 180 (Xu et al., 2017). Both laboratory and field measurements indicate that the CE of CV was fairly robust 181 and was roughly equivalent to 1. Therefore, a CE of 1 was applied to all measured species in this study 182 (Hu et al., 2017; Hu et al., 2018b) (Hu et al., 2017; Hu et al., 2018b). Relative ionization efficiencies 183 (RIEs) of 3.06 and 1.09 were used for ammonium and sulfate quantification respectively, and the 184 default values of 1.1 and 1.4 were used for nitrate and organic aerosol (OA) respectively. Compared 185 with the AMS with standard vaporizer, the CV-ToF-ACSM reports higher fragments at smaller m/z's 186 due to additional thermal decomposition associated with increased residence time and hot surface 187 collisions (Hu et al., 2018a). As a result, f₄₄ from CV-ToF-ACSM measurements is often much higher 188 than those previously reported from AMS, yet they are well correlated (Hu et al., 2018a). 189

The organic mass spectra from m/z 12 to 214 were analyzed by positive matrix factorization (PMF) 190 191 (Paatero and Tapper, 1994) with an Igor Pro based PMF evaluation tool (v3.04) (Ulbrich et al., 2009). The ion fragments m/z of 38, 49, 63 and 66 were removed from both PM₁ and PM_{2.5} PMF inputs 192 considering their small contributions to the total organic signals yet with high signal-to-noise ratios. 193 The PMF results were then evaluated following the procedures detailed in Zhang et al. (2011). After 194 carefully evaluating the mass spectral profiles, diurnal patterns and temporal variations of the OA 195 factors and comparing them with other collocated measurements, a five-factor solution was selected 196 for both PM₁ and PM_{2.5}. The five factors include four primary factors, i.e., hydrocarbon-like OA 197 (HOA), cooking OA (COA), biomass burning OA (BBOA), and coal combustion OA (CCOA), and a 198 199 secondary factor, oxygenated OA (OOA). More detailed descriptions on the PMF results will be given 200 in He et al. (in preparation).

201 2.4 Data reprocessing

The size distributions measured by APS were converted to mobility-equivalent size distributions using spherical shape assumptions and an effective particle density of 1.7 g/cm³. Note that the

designations of PM₁₀ and PM₁ are in respect to aerosol aerodynamic diameters, while the 204 corresponding mobility-equivalent cut diameters of the two impactors are approximately 7669 nm and 205 767 nm, respectively. For simplicity and consistency, we will continue to refer to them as the PM₁₀ and 206 PM₁ based on their aerodynamic diameter. For the case of PM₁ measurements, the mobility-equivalent 207 cut diameter is quite near the upper range of the SMPS size range. Considering that the cut diameter 208 of the impactor corresponds to the diameter of aerosol particles in ambient state (aerosol hygroscopic 209 growth effect needs to be taken into account) and the SMPS measures the size distributions of aerosol 210 particles in dry state, the SMPS measurements should be able to cover the full-size range of PM₁. 211 When the SMPS was sampling aerosol particles of PM₁₀, the size distributions measured by SMPS 212 and APS was merged together and truncated to an upper limit of 7669 nm to provide full range particle 213 number size distributions (PNSD). In addition, the AE33 measures aerosol absorption coefficient at 214 several wavelengths, the mass concentrations of black carbon (BC) were converted from measured 215 aerosol absorption coefficients at 880 nm with a mass absorption coefficient of 7.77 m²/g (Drinovec 216 et al., 2015). 217

Since group 2 and 3 switched between PM_1 and PM_{10} inlets every 15 minutes, all measurements were averaged over each 15-minute observation episode, resulting in valid time resolutions of 15 minutes for APS and BC PM_{10} measurements and of 30 minutes for SMPS, CV-ToF-ACSM and the humidified nephelometer system PM_1 and PM_{10} measurements, respectively. This resulted in a 15minute time lag between the averaged datasets of group 2 and group 3. To match the time of all the measurement data, the measurements of SMPS, ACSM and the humidified nephelometer system were linearly interpolated to the 15-minute time resolution of the APS data.

225 **3 Methodology**

226 **3.1** Calculations of hygroscopicity parameters κ_{sca} and κ from measurements of the 227 humidified nephelometer system

The humidified nephelometer system measures aerosol light scattering coefficients and backscattering coefficients at three wavelengths under dry state and 85% RH condition, providing measurements of the light scattering enhancement factor $f(RH, \lambda)$, which is defined as

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231 $f(\text{RH} = 85\%, \lambda) = \frac{\sigma_{sp}(\text{RH}, \lambda)}{\sigma_{sp}(\text{dry }\lambda)}$, with λ being the light wavelength. In this study, we only calculate 232 f(RH, 525 nm) and refer to it hereinafter as f(RH) for simplicity. Brock et al. (2016) proposed a 233 single parameter formula to describe $f(\text{RH}, \lambda)$ as a function of RH. Kuang et al. (2017) further 234 developed this parameterization scheme to better describe measured f(RH) by including the 235 reference RH (RH₀) in the dry nephelometer as shown in Eq.1, using which the optical hygroscopicity 236 parameter κ_{sca} can be derived from $f(\text{RH})_{measured}$.

237
$$f(\text{RH})_{measured} = (1 + \kappa_{sca} \frac{RH}{100 - RH}) / (1 + \kappa_{sca} \frac{RH_0}{100 - RH_0})$$
 (1)

An overall hygroscopicity parameter κ referred to as $\kappa_{f(RH)}$ can be retrieved from measured f(RH) with the addition of simultaneously measured particle number size distribution (PNSD) and BC mass concentration (Chen et al., 2014;Kuang et al., 2017). The idea is to conduct an iterative calculation using the Mie theory and the κ -Köhler theory together to find a $\kappa_{f(RH)}$ that closes the gap between the simulated and the measured f(RH). Details on the calculations of $\kappa_{f(RH)}$ can be found in Kuang et al. (2017).

244 **3.2** Calculations of κ_{chem} from aerosol chemical composition measurements

For the calculation of aerosol hygroscopicity parameter κ based on measured chemical composition data (κ_{chem}), detailed information on the chemical species are needed. The CV-ToF-ACSM can only provide bulk mass concentrations of SO_4^{2-} , NO_3^- , NH_4^+ , Cl^- ions and organic components. For the inorganic ions, a simplified ion pairing scheme (as listed in Table 1) was used to convert ion mass concentrations to mass concentrations of corresponding inorganic salts (Gysel et al., 2007;Wu et al., 2016).

Table 1. Densities (ρ) and hygroscopicity parameters (κ) of inorganic salts used in this study

Species	NH ₄ NO ₃	NH ₄ HSO ₄	$(NH_4)_2 SO_4$	NH ₄ Cl
$\rho (g cm^{-3})$	1.72	1.78	1.769	1.527
κ	0.58	0.56	0.48	0.93

Mass concentrations of SO_4^{2-} , NO_3^- , NH_4^+ , Cl^- are thus specified into ammonium sulfate (AS), ammonium nitrate (AN), ammonium chloride (AC) and ammonium bisulfate (ABS), with the κ values of these salts specified according to (Wu et al., 2016) and Liu et al. (2014) (Table1). For a given internal mixture of different aerosol chemical species, a simple mixing rule called Zdanovskii–Stokes– Robinson (ZSR) can be used for predicting the overall κ_{chem} on the basis of volume fractions of different chemical species (ε_i) (Petters and Kreidenweis, 2007):

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$$\kappa_{chem} = \sum_i \kappa_i \cdot \varepsilon_i$$
 (2)

where κ_i and ε_i represent the hygroscopicity parameter κ and volume fraction of chemical component *i* in the mixture. Based on Eq.2, κ_{chem} can be calculated as follows:

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$$\kappa_{chem} = \kappa_{AS}\varepsilon_{AS} + \kappa_{AN}\varepsilon_{AN} + \kappa_{ABS}\varepsilon_{ABS} + \kappa_{AC}\varepsilon_{AC} + \kappa_{BC}\varepsilon_{BC} + \kappa_{OA}\varepsilon_{OA}$$
 (3)

where κ_{OA} and ε_{OA} represent κ and volume fraction of total organics. Since black carbon is hydrophobic, κ_{BC} is assumed to be zero. With known κ_{chem} , κ_{OA} can be calculated using the following formula:

265
$$\kappa_{OA} = \frac{\kappa_{chem} - (\kappa_{AS}\varepsilon_{AS} + \kappa_{AN}\varepsilon_{AN} + \kappa_{ABS}\varepsilon_{ABS} + \kappa_{AC}\varepsilon_{AC})}{\varepsilon_{OA}}$$
 (4)

The volume concentration of organics was calculated by assuming density of POA as 1 g/cm³ and 266 density of OOA as 1.4 g/cm³ (Wu et al., 2016). For the calculation of the total volume concentration 267 (V_{tot}) , we have three approaches. The first approach is to sum up the volume concentrations of all 268 chemical species (AS, AN, ABS, AC, BC and organics), where the volume concentration of BC was 269 calculated by assuming a density of 1.7 g/cm³ (Wu et al., 2016). We refer the calculated total volume 270 concentration of aerosol particles to as $V_{tot,Chem}$. The second approach is to integrate V_{tot} from 271 measured PNSD using the equation $V_{tot,PNSD} = \int \frac{4}{3} \pi r^3 n(r) dr$, where r is the particle radius and n(r) 272 is the measured particle number concentrations. The third approach is to use the trained machine 273 learning estimator to estimate the V_{tot} based on measurements of dry nephelometer ($V_{tot,Neph}$) as was 274 introduced in Kuang et al. (2018). V_{tot} of PM₁ calculated using these three methods were compared 275 to each other and shown in Fig.S2. $V_{tot,Chem}$ correlates well with $V_{tot,PNSD}$, but it is on average 30% 276 lower than that of $V_{tot,PNSD}$. Chemical components within aerosol particles such as dust, sea salt as 277 well as metal ions cannot be detected by CV-ToF-ACSM. Since the Gucheng site is far from the ocean, 278 279 sea salt should have negligible impacts on the total mass of PM1. However, mineral dust can extend into the submicron range (Shao et al., 2007), which might be the cause for the low $V_{tot,Chem}$. 280 calculated using CV-ToF-ACSM and BC data. $V_{tot,Neph}$ also correlates well with $V_{tot,PNSD}$, but is on 281

average 16% lower than that of $V_{tot,PNSD}$. Closure studies between modelled and measured σ_{sp} and 282 σ_{bsp} at 525 nm for PM₁ and PM₁₀ aerosol particles all showed good agreement between theoretical 283 modelling results and measurements (Fig.S1), with most points falling within the 20% relative 284 deviation lines. However, modelled σ_{sp} for both PM₁ and PM₁₀ were obviously higher than measured 285 σ_{sp} , with an average relative difference of 22% and 13% between them for PM₁₀ and PM₁, respectively. 286 The result for PM₁ explains why $V_{tot,Neph}$ was lower than $V_{tot,PNSD}$. Two reasons might have 287 contributed to this discrepancy: (1) both PNSD and aerosol optical property measurements carry non-288 negligible uncertainties, with the SMPS bearing measurement uncertainty of 30% for particles larger 289 than 200 nm, which contribute most to V_{tot} (Wiedensohler et al., 2012), and the nephelometer 290 measured σ_{sp} having an uncertainty of 9% (Sherman et al., 2015; Titos et al., 2016); (2) The sampling 291 292 tube length, valves, tube angles and flow rates are different for the dry nephelometer and SMPS (e.g. much shorter tube and smaller flow rate for SMPS than those for the dry nephelometer), leading to 293 294 different wall loss and loss in semi-volatile aerosol components. ACSM and the dry nephelometer had a similar tube length and nephelometer measurements bear less uncertainty than SMPS. Thus, 295 $V_{tot,Neph}$ was chosen as V_{tot} in the calculations of Eq.4. Based on the calculated V_{tot} , the material 296 unidentified by CV-ToF-ACSM accounts for 19% of V_{tot} on average, could not be neglected in the 297 κ_{OA} calculation. Thus, Eq.4 was modified as follows: 298

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$$\kappa_{Org} = \frac{\kappa_{chem} - (\kappa_{AS} \cdot \varepsilon_{AS} + \kappa_{AN} \cdot \varepsilon_{AN} + \kappa_{ABS} \cdot \varepsilon_{ABS} + \kappa_{X} \cdot \varepsilon_{X})}{\varepsilon_{Org}}$$
(5)

where κ_X and ε_X are hygroscopicity parameter κ and volume fraction of the unidentified material. Previous studies using $V_{tot,Chem}$ as the total volume concentration of aerosol particles have avoided the discussion about influences of unidentified material by the CV-ToF-ACSM or other aerosol mass spectrometer instruments. The hygroscopicity of these unidentified materials, which might be dust or other components in continental regions, was not discussed before. Dust is nearly hydrophobic, with mineral dust showing κ in range of 0.01 to 0.08 (Koehler et al., 2009). In this paper, we arbitrarily specified κ_X to be 0.05.

307 **3.3** Can $\kappa_{f(RH)}$ represent κ_{chem} ?

According to Eq.5, the measured bulk κ_{chem} values are needed to derive κ_{0A} . Bulk aerosol chemical compositions and aerosol hygroscopicity $\kappa_{f(RH)}$ measurements are available, one might naturally jump to the conclusion of treating $\kappa_{f(RH)}$ as κ_{chem} to derive κ_{OA} because both $\kappa_{f(RH)}$ and κ_{chem} are from bulk aerosol measurements. However, the relationship between κ_{chem} , $\kappa_{f(RH)}$ and the size-resolved κ distribution needs to be clarified in order to answer the question whether $\kappa_{f(RH)}$ can accurately represent κ_{chem} . The physical meanings of used different κ representations in the following discussions are listed in Table 2.

Table 2. Different κ and their physical meanings

$\kappa_{f(\mathrm{RH})}$	A uniform κ for all particle sizes which describes $f(RH)$ accurately				
κ _{chem}	A bulk κ assuming different chemical compositions of aerosol populations				
	are internally mixed and calculated with the ZSR mixing rule				
κ_i	hygroscopicity parameter κ of chemical species <i>i</i>				
κ_{D_p}	The κ assuming different chemical compositions of particles with diameter				
•	of D_n are internally mixed and calculated with the ZSR mixing rule				

 D_p are internally mixed and calculated with the ZSR mixing rule Using V_i to represent volume concentrations of chemical species *i* and $V_i(D_p)$ to represent volume concentrations of species *i* with diameter of D_p , κ_{chem} can be derived as follows based on Eq.2,:

319
$$\kappa_{chem} = \sum_{i} \kappa_{i} \cdot \varepsilon_{i} = \sum_{i} \frac{V_{i}}{V_{tot}} \cdot \kappa_{i} = \sum_{i} \frac{1}{V_{tot}} \cdot \int \frac{d V_{i}(D_{p})}{d log D_{p}} \cdot d log D_{p} \cdot \kappa_{i}.$$
(6)

By swapping the order of summation and integration, Eq.6 can be written as:

321
$$\kappa_{chem} = \int \frac{1}{V_{tot}} \cdot \sum_{i} \frac{d V_i (D_p)}{d \log D_p} \cdot d \log D_p \cdot \kappa_i.$$
(7)

322 Considering that $\kappa_{D_p} = \sum_i \frac{dV_i(D_p)}{dV(D_p)} \cdot \kappa_i$, Eq.7 can be rewritten as:

323
$$\kappa_{chem} = \frac{1}{V_{tot}} \int \kappa_{D_p} \cdot dV(D_p)$$
 (8)

Result of Eq.8 indicates that κ_{chem} calculated using Eq.3 represents the overall hygroscopicity of aerosol particles with volume contribution as the weighting function of κ_{D_p} .

As for $\kappa_{f(RH)}$, a detailed analysis is performed here to facilitate its physical understanding. The differential form of σ_{sp} of aerosol particles in dry state can be expressed as follows:

328
$$\sigma_{sp} = \int \frac{d\sigma_{sp}}{dlog D_p} dlog D_p$$
 (9)

Based on the definition of f(RH), σ_{sp} of aerosol particles under different RH conditions can be

330 written as:

331
$$\sigma_{sp}(RH) = \int \frac{d\sigma_{sp}}{dlogD_p} \cdot f_{D_p}(RH) \cdot dlogD_p$$
 (10)

Therefore, the differential form of observed overall f(RH) can be formulated as:

333
$$f(RH) = \int \frac{1}{\sigma_{sp}} \cdot \frac{d\sigma_{sp}}{dlogD_p} \cdot f_{D_p}(RH) \cdot dlogD_p$$
 (11)

Based on this formula, the sensitivity of f(RH) on the hygroscopicity of aerosol particles with diameter D_p (κ_{D_p}) can be derived as:



Figure 2. (a) simulated $\frac{\partial f_{D_p}(RH)}{\partial \kappa_{D_p}}$; **(b)** simulated $\frac{1}{dlog D_p} \cdot \frac{\partial f(RH)}{\partial \kappa_{D_p}}$

The sensitivity of f(RH) to κ_{D_p} are determined together by the two terms in Eq. 12: (1) $\frac{1}{\sigma_{sp}} \cdot \frac{d\sigma_{sp}}{dlogD_p}$, which represents the contribution of σ_{sp} of aerosol particles in dry state with diameter D_p to total

 σ_{sp} , and (2) $\frac{\partial f_{D_p}(RH)}{\partial \kappa_{D_p}}$, which represents the sensitivity of $f_{D_p}(RH)$ to κ_{D_p} . Based on the Mie theory 339 and the κ -Köhler theory, we simulated the second term under 85% RH condition for varying D_p and 340 κ_{D_p} values (Fig.2a). In the diameter range below 200 nm, $\frac{\partial f_{D_p}(RH)}{\partial \kappa_{D_p}}$ is very high, displaying a 341 maximum near 60 nm. In this diameter range, larger κ_{D_p} generally corresponds to higher $\frac{\partial f_{D_p}(RH)}{\partial \kappa_{D_n}}$. 342 For 200 nm < D_p < 800 nm, higher and lower $\frac{\partial f_{D_p}(RH)}{\partial \kappa_{D_n}}$ appear alternatively, with all values 343 staying positive. For $D_p > 800 \text{ nm}$, maxima and minima regions appear alternatively, and $f_{D_p}(RH)$ 344 might decrease with increasing κ_{D_n} . This is because, at this diameter range, the aerosol scattering 345 efficiency has a non-monotonic response to the particle diameter increase (see Fig.2a of (Kuang et al., 346 2018)). 347

The first term of Eq.9, representing size-resolved σ_{sp} contributions of particles with diameter in 348 dry state, mainly depends on the PNSD. The average PNSD of PM₁₀ was applied in the simulation of 349 the first term using Mie theory (Fig.S3). Combining results of the first term and second term, the 350 sensitivity of f(RH) to κ_{D_n} was obtained and depicted in Fig.2b. Results reveal that f(RH) is quite 351 sensitive to the κ_{D_p} of particles within 200 to 800 nm diameter range, but almost insensitive to κ_{D_p} 352 of particles with diameters below 200 nm and above 800 nm (corresponding aerodynamic diameter of 353 about 1 μ m). For particles smaller than 200 nm, the first term was quite small especially for particles 354 smaller than 100 nm (Fig.S3), while for particles larger than 800 nm, in addition to a small first term, 355 the second term fluctuated between negative and positive values, which is why f(RH) was not 356 sensitive to the overall hygroscopicity of these larger aerosol particles. These results suggest that 357 although $\kappa_{f(RH)}$ was derived from f(RH) measurements of PM₁₀, it mainly represents the overall 358 hygroscopicity of aerosol particles with dry diameters between 200 and 800 nm for continental 359 aerosols. This result indicates that $\kappa_{f(RH)}$ derived from f(RH) measurements of PM₁₀ and PM₁ 360 should differ little from each other for measurements conducted in continental regions. 361

However, the quantitative relationship between $\kappa_{f(RH)}$ and size-resolved κ_{D_p} is still not clear. Based on Eq.11, $f_{D_p}(RH)$ can be expressed as:

364
$$f_{D_p}(RH) = \frac{d\sigma_{sp}(RH)}{d\sigma_{sp}} = \frac{\frac{1}{4}\pi \cdot (D_p \cdot g)^2 \cdot Q_{sca}(D_p \cdot g) \cdot dN}{d\sigma_{sp}}, (13)$$

where *g* is the growth factor of aerosol particles which is a function of κ_{D_p} and RH (Brock et al., 2016), i.e. $g = (1 + \kappa_{D_p} \cdot \frac{RH}{100 - RH})^{1/3}$, dN is differential form of aerosol number concentration, and Q_{sca} is the scattering efficiency as a function of D_p and *g*. The results of Kuang et al. (2018) indicate that, under dry state, Q_{sca} can be expressed as $Q_{sca} = k \cdot D_p$ with *k* varying as a function of D_p . Here, we follow this idea and express the Q_{sca} under humidified condition as $Q_{sca}(D_p, g) =$ $C \cdot D_p \cdot g$, where C is a function of D_p , κ_{D_p} and RH. Replacing *g* and Q_{sca} in Eq.13, we yield:

371
$$f_{D_p}(RH) = \frac{\frac{1}{4} \pi \cdot D_p^{3} \cdot C(\mathrm{Dp}, \kappa_{D_p}, \mathrm{RH}) \cdot (1 + \kappa_{D_p} \cdot \frac{RH}{100 - RH}) \cdot dN}{d\sigma_{sp}}, \quad (14)$$

which we can substitute into Eq.8, to obtain a new expression for f(RH):

373
$$f(\mathrm{RH}) = \int \frac{\frac{1}{4} \pi \cdot D_p^{3} \cdot C(\mathrm{Dp}, \kappa_{D_p}, \mathrm{RH}) \cdot (1 + \kappa_{D_p} \cdot \frac{RH}{100 - RH}) \cdot dN}{\sigma_{sp}}$$
(15)

374 If we define $X_c(\text{Dp}, \kappa_{D_p}, \text{RH}) = C(\text{Dp}, \kappa_{D_p}, \text{RH})/k$, and considering that $d\sigma_{sp} = \frac{1}{4} \cdot \pi \cdot D_p^2 \cdot Q_{sca}$.

375
$$dN = \frac{1}{4} \cdot \pi \cdot D_p^3 \cdot \mathbf{k} \cdot dN$$
, Eq.14 can be written as:

376
$$f(\mathrm{RH}) = \int \frac{X_c(\mathrm{Dp},\kappa_{Dp},\mathrm{RH}) \cdot (1+\kappa_{Dp},\frac{RH}{100-RH}) \cdot d\sigma_{sp}}{\sigma_{sp}}$$
(16)

The $\kappa_{f(RH)}$ is a uniform κ for aerosol particle sizes that can yield simulated f(RH) equal to the measured one. Thus, f(RH) can also be expressed as:

379
$$f(\mathrm{RH}) = \int \frac{X_c(\mathrm{Dp},\kappa_{f(\mathrm{RH})},\mathrm{RH}) \cdot (1 + \kappa_{f(\mathrm{RH})} \cdot \frac{RH}{100 - RH}) \cdot d\sigma_{sp}}{\sigma_{sp}} \quad (17)$$

Combining Eq.16 and Eq.17, the relationship between $\kappa_{f(RH)}$ and size-resolved κ_{D_p} can be derived

382
$$\kappa_{f(\mathrm{RH})} = \frac{\int X_c(\mathrm{Dp},\kappa_{D_p},\mathrm{RH})\cdot\kappa_{D_p}\cdot d\sigma_{sp}}{\int X_c(\mathrm{Dp},\kappa_{f(\mathrm{RH})},\mathrm{RH})\cdot d\sigma_{sp}} + \frac{\int (X_c(\mathrm{Dp},\kappa_{D_p},RH) - X_c(\mathrm{Dp},\kappa_{f(\mathrm{RH})},\mathrm{RH})\cdot d\sigma_{sp}}{\int X_c(\mathrm{Dp},\kappa_{f(\mathrm{RH})},\mathrm{RH})\cdot d\sigma_{sp}} \cdot \frac{100 - RH}{RH}.$$
 (18)

383 X_c values under 85% RH for different D_p and κ_{D_p} values are simulated and shown in Fig.3, based



on this result of X_c the second term of Eq.18 (which depends on the PNSD and size-resolved κ_{D_p})

Figure 3. Simulated values of X_c under 85% RH for different D_p and κ_{D_p} values. Black solid and dashed lines are two assumed size-resolved κ_{D_p} distributions.

could be calculated using the average PNSD during this field campaign and two assumed extreme 385 cases of size-resolved κ_{D_p} (solid and dashed black lines in Fig.3). For PM₁, the second term 386 corresponding to the two size-resolved κ_{D_p} cases were -0.007 and 0.008, respectively. Corresponding 387 values simulated for PM₁₀ were -0.005 and 0.004, respectively. To further investigate the possible 388 contribution range of the second term to $\kappa_{f(RH)}$, size-resolved κ_{D_p} derived by Liu et al. (2014) based 389 on size resolved chemical composition measurements in ambient atmosphere on the NCP region 390 (Fig.S4) were used with the average PNSD during this campaign to calculate values of the second term. 391 Calculated values of second term ranged from -0.005 to 0.009, with its contribution to $\kappa_{f(RH)}$ ranging 392 393 from -1.5% to 2% (0.3% on average). These results indicate that the second term was negligible in most cases, and Eq.18 could be approximated as: 394

395
$$\kappa_{f(\mathrm{RH})} \approx \frac{\int X_c(\mathrm{Dp},\kappa_{Dp},\mathrm{RH})\cdot\kappa_{Dp}\cdot d\sigma_{sp}}{\int X_c(\mathrm{Dp},\kappa_{f(\mathrm{RH})},\mathrm{RH})\cdot d\sigma_{sp}}$$
 (19)

396 X_c values shown in Fig.3 indicate that for aerosol particles in the diameter range of 200 to 800 nm 397 (which contribute most to σ_{sp} and are the part of the aerosol population that $\kappa_{f(RH)}$ is most sensitive 398 to) and for the observed κ_{Dp} range of continental aerosols (κ_{Dp} usually less than 0.5), X_c mainly 399 ranged from 0.7 to 1. Considering this, we might approximately assume X_c in Eq.18 as a constant 400 value. Then, Eq.19 can be further simplified to:

401
$$\kappa_{f(\mathrm{RH})} \approx \frac{1}{\sigma_{sp}} \int \kappa_{Dp} \cdot d\sigma_{sp}$$
 (20)

402 This result suggests that $\kappa_{f(RH)}$ can be approximately understood as the overall hygroscopicity of 403 aerosol particles with the σ_{sp} contribution as the weighting function of κ_{D_n} .

Based on results of Eqs.8 and 20, both $\kappa_{f(RH)}$ and κ_{chem} represent the overall hygroscopicity of bulk aerosol particles, however, their weighting functions of κ_{D_p} are different. Within a certain D_p range, aerosol σ_{sp} is approximately proportional to aerosol volume (Kuang et al., 2018), resulting in little difference between $\kappa_{f(RH)}$ and κ_{chem} . In this study, bulk $\kappa_{f(RH)}$ was measured for both PM₁ and PM₁₀. How much does κ_{chem} differ from $\kappa_{f(RH)}$ for PM₁ and PM₁₀ samples? Both PNSD and size-resolved κ_{D_p} distributions contribute to the difference between κ_{chem} and $\kappa_{f(RH)}$. To study



Figure 4. κ_{chem} versus $\kappa_{f(RH)}$, colors represent average Ångström exponent (Å) values of corresponding PNSD (a) corresponding to size-resolved κ_{D_p} distributions shown in Fig.3 (squares correspond to the solid line in Fig.3 and circles correspond to the dashed line in Fig.3); (b) and (c) corresponding to size-resolved κ_{D_p} distributions shown in Fig.S4 for PM1 and PM10, respectively. Gray areas represent the absolute relative differences between κ_{chem} and $\kappa_{f(RH)}$ less than 10%.

their influences in a simple and apparent way, κ_{chem} and $\kappa_{f(RH)}$ were simulated based on the two extreme cases of size-resolved κ_{Dp} distributions in Fig. 3 and five average PNSDs corresponding to five ranges of aerosol Ångström exponent (0.9-1.1,1.1-1.3,1.3-1.5,1.5-1.7,1.7-1.9) during this field campaign. In the instance of PM₁, as can be seen in Fig.4a, assuming a κ_{Dp} increasing as a function of D_p resulted in $\kappa_{chem} < \kappa_{fRH}$ (square points in Fig.4a), especially for PNSDs corresponding to larger Ångström exponents. This is because the volume contributions of small particles (e.g. particles with D_p between 100 to 300 nm) to V_{tot} are larger than their light scattering coefficient contributions

to σ_{sp} (as shown in Fig.S6), thus the hygroscopicity of small particles had larger impacts on κ_{chem} 417 than κ_{fRH} . Higher Ångström exponents generally correspond to shift in PNSD towards smaller D_p , 418 which exacerbates the contribution of small particles, further increasing the difference between κ_{chem} 419 and κ_{fRH} . For the case with κ_{D_p} decreasing as a function of D_p (circle markers in Fig.4a) it is vice 420 versa, resulting in $\kappa_{chem} > \kappa_{fRH}$. In general, for these two extreme cases of size-resolved κ_{D_p} 421 distributions, the absolute value of the relative difference between κ_{chem} and $\kappa_{f(RH)}$ ranged from 422 2.8% to 7.5% with an average of 4.8%. This result indicates that for PM₁, κ_{chem} might differ little 423 from $\kappa_{f(RH)}$ since κ_{D_p} usually varies less with D_p in ambient atmosphere than in the two assumed 424 cases (Liu et al., 2014). The average size-resolved κ_{D_p} distribution from Haze in China campaign 425 (Liu et al., 2014) indicate that κ_{D_p} varies significantly for $D_p < 250$ nm, while it varies less within the 426 427 diameter range of 250 nm to 1 µm. To further study the variation range of the relative difference between κ_{chem} and $\kappa_{f(RH)}$ under ambient conditions, the size-resolved κ_{D_p} distributions derived 428 429 from measured size-resolved chemical compositions in the NCP region (Liu et al., 2014) (shown in Fig.S5) were used in simulations and results are shown in Fig.4b. The absolute value of the relative 430 difference between κ_{chem} and $\kappa_{f(RH)}$ ranged from 0.04% to 8% with an average and standard 431 deviation of 2.8±2%, which further confirms that for PM₁ $\kappa_{f(RH)}$ can accurately represent κ_{chem} in 432 433 most cases.

For PM₁₀, values of κ_{chem} and $\kappa_{f(RH)}$ using κ_{D_p} size distributions derived from ambient measurements (Fig.S5, similar to Fig.4b) were simulated and displayed in Fig.4c. The simulated absolute values of the relative difference between κ_{chem} and $\kappa_{f(RH)}$ ranged from 0.2% to 41% with an average and standard deviation of 16±8%, with all κ_{chem} lower than $\kappa_{f(RH)}$. This is because, for PM₁₀, super-micron particles typically with low hygroscopicity (Fig.S5) contribute much more to V_{tot} than to σ_{sp} (as shown in Fig.S7). These results indicate that, for PM₁₀, $\kappa_{f(RH)}$ cannot accurately represent κ_{chem} .

441 Above analysis results indicate that $\kappa_{f(RH)}$ retrieved from light scattering measurements of PM₁ 442 represent accurately the κ_{chem} of PM₁ and can be used in Eq.5 as measured κ_{chem} for deriving κ_{OA} .

443 **4 Results and discussions**

444 **4.1 Overview of the campaign data**

The time series of ambient RH, chemical compositions of PM_{2.5} and PM₁, σ_{sp} at 525 nm of PM₁₀ 445 and PM₁ in dry state, calculated κ_{sca} and $\kappa_{f(RH)}$ values of PM₁₀ and PM₁ are shown in Fig.5. Overall, 446 the mass concentrations of NR-PM₁ and NR-PM_{2.5} ranged from 1 to 221 $\mu g/m^3$ and from 1.8 to 326 447 $\mu g/m^3$, with average concentrations of 63 and 93 $\mu g/m^3$, respectively. Measured σ_{sp} at 525 nm of 448 PM₁ and PM₁₀ ranged from 11 to 1875 Mm^{-1} and from 18 to 2732 Mm^{-1} , with average values of 449 550 and 814 Mm^{-1} , respectively. These results demonstrate that this campaign was carried out at a 450 site that is overall highly polluted, where quite clean conditions as well as extremely polluted 451 conditions were experienced during the measurement period. The mass contributions of ammonium, 452 nitrate, sulfate and organics to NR-PM2.5 and NR-PM1 are listed in Table 3 with organics being the 453 454 major fraction of NR-PM1 and NR-PM2.5.



Figure 5. Time series of ambient RH (a), chemical compositions of PM_{2.5} (b) and PM₁ (c), σ_{sp} at 525 nm of PM₁₀ and PM₁ (d), calculated κ_{sca} (e) and $\kappa_{f(RH)}$ (f) values of PM₁₀ and PM₁.

During period 1 shown in Fig.5, nitrate contributed most to inorganics, while inorganics contribute most to mass concentrations of NR-PM_{2.5} and NR-PM₁. During the period 2 shown in Fig.5, the ambient RH is relatively lower than that of the first period, ranging from 16% to 86% with an average of 49%. During this period, organics contributed most to mass concentrations of NR-PM_{2.5} and NR-PM₁, with the NR mass concentrations of PM_{2.5} and σ_{sp} at 525 nm of PM₁₀ being only 33% and 40% higher than those of PM₁.

The time series of calculated κ_{sca} and $\kappa_{f(RH)}$ are shown in Fig.5e-f. κ_{sca} of PM₁ and PM₁₀ ranged from 0.01 to 0.2, and from 0.02 to 0.17, with corresponding averages of 0.09 and 0.08, respectively. The $\kappa_{f(RH)}$ was not available from 12:00 10th Dec to 12:00 11th Dec due to the absence

of PNSD measurements. $\kappa_{f(RH)}$ of PM₁ and PM₁₀ ranged from 0.02 to 0.27, and from 0.03 to 0.26, 464 with corresponding averages of 0.12 and 0.12, respectively. These results indicate that the 465 466 hygroscopicity during this campaign was generally low, which could be associated with the high mass contributions of organics. The range as well as the average level of $\kappa_{f(RH)}$ is quite consistent with the 467 results obtained at the same site in winter 2016, suggesting the prevalent low aerosol hygroscopicity 468 conditions in winter at this site. Additionally, it can be noted that except for fog events, κ_{sca} and 469 $\kappa_{f(RH)}$ values of PM₁ are generally higher than those of PM₁₀, yet the differences are small (10% and 470 471 3.5% for κ_{sca} and $\kappa_{f(RH)}$, respectively). Although particles with diameters above 800 nm impact almost negligibly on retrieved $\kappa_{f(RH)}$ (refer to discussions in Sect3.3), it can still cause a small 472 difference between $\kappa_{f(RH)}$ of PM₁₀ and PM₁. Results of previous studies indicate that the overall 473 hygroscopicity of aerosol particles larger than 800 nm are usually low and are typically lower than the 474 overall hygroscopicity of accumulation mode particles (Liu et al., 2014), which may explain why 475 $\kappa_{f(RH)}$ values of PM₁ are generally higher than those of PM₁₀ during non-fog periods (periods with RH 476 <100%). 477

Smaaiaa	Ammonium		nitrate		sulfate		Organics	
Species	PM_1	PM _{2.5}	PM_1	PM _{2.5}	PM_1	PM _{2.5}	PM_1	PM _{2.5}
Entire	12%	12%	13%	14%	10%	11%	59%	59%
period	0.2-24%	0.1-24%	2-31%	1-32%	0.3-49%	0.2-50%%	12-99%	4-91%
Period 1	15%	16%	22%	24%	13%	14%	47%	42%
Fog	10-17%	12-18%	11-28%	16-30%	9-15%	12-16%	30-65%	37-55%
Period 1	17%	16%	23%	23%	12%	12%	43%	44%
non-fog	10-22%	7-21%	6-31%	5-32%	8-23%	7-17%	32-75%	31-69%
Daviad 2	12%	10%	11%	10%	8%	7%	64%	67%
reriou 2	0.2-20%	0.1-19%	5-30%	4-29%	0.3-16%	0.2-16%	40-82%	40-85%

478 Table 3. Average (range) mass contribution of ammonium, nitrate, sulfate and organics to NR-PM_{2.5} and NR-PM₁
479 during different periods.

480 During fog periods, a large part of submicron particles in dry state will be activated into fog 481 droplets, which are super micron particles in ambient state (see PNSD examples in Fig.S4a), exerting 482 substantial impacts on f(RH) measurements of PM₁₀ which are not detectable in the PM₁ measurements. Since for a certain particle diameter and fog supersaturation, particles with higher hygroscopicity are more readily activated, the observed PM₁₀ $\kappa_{f(RH)}$ increased during fog events and often exceeded those of PM₁ in contrast to non-fog periods (Fig.5f).

486 4.2 κ_{OA} derivations and its relationship with organic aerosol oxidation state

The results in Sect.3.3 demonstrate that $\kappa_{f(RH)}$ of PM₁ accurately represents κ_{chem} in most cases, thus a closure study between calculated κ_{chem} of PM₁ based on measured chemical compositions and measured κ_{chem} (represented by PM₁ $\kappa_{f(RH)}$) can be conducted using Eq.3 if κ_{OA} was known. A κ_{OA} of 0.06 was used in this closure test, which was calculated by Wu et al. (2016) based on aerosol chemical composition and aerosol hygroscopicity measurements. As shown in Fig.6a, the comparison between measured and calculated κ_{chem} has not achieved very good agreements. We



Figure 6. Comparison between measured and calculated κ_{chem} by assuming a κ_{0rg} of 0.06. (a) The whole period; (b) Only Period 1; (c) Only Period 2. Colors represents the mass fractions of organic aerosol in NR-PM₁ (f_{0A}), and the color bar is shown on the top.

notice that the calculated κ_{chem} was overestimated when mass fraction of organic aerosol (f_{OA}) was 493 lower than 45%, while it was underestimated when f_{OA} was higher than 45%. As described in 494 Sect.4.1, these two situations roughly correspond to Period 1 and 2, respectively. Separating the data 495 points shown in Fig.6a into Periods 1 (Fig.7b) and 2 (Fig.7c), it can be seen that all low f_{OA} data 496 497 points are found in Period 1, with most of the data points showing f_{OA} less than 50%. Although the calculated κ_{chem} during this period was on average 25% higher than the measured κ_{chem} , they were 498 highly correlated (R=0.84). A similar case was also found in Wu et al. (2013), and they concluded that 499 500 the loss of semi-volatile ammonium nitrate in the HTDMA might be the reason. The relationship

between nitrate concentration and the difference between calculated and measured κ_{chem} was investigated, which confirmed the influence of nitrate on this discrepancy (Fig.S7), and the overestimation of calculated κ_{chem} due to the volatile loss of ammonium nitrate. Since the tube length (from the splitter to inlet of instrument) of wet nephelometer was about 1 m longer than that of the CV-ToF-ACSM, there probably was more loss in ammonium nitrate in the wet nephelometer.

During Period 2, the average mass fraction of nitrate was low (11%), and the loss of ammonium nitrate had minor influence on κ_{chem} estimations (Fig.S7). However, when organic aerosol was dominant during Period 2, the calculated κ_{chem} was underestimated in most cases (Fig.6c). Previous studies have shown larger κ_{OA} for OA with higher oxidation level (Chang et al., 2010;Duplissy et al., 2011;Wu et al., 2013), which might have contributed to the underestimation in κ_{chem} . This gave us the hint that Period 2 might provide us a good opportunity to study κ_{OA} . Following the method in Sect. 3.2, κ_{OA} was derived using Eq.5, resulting in a κ_{OA} ranging from 0.0 to 0.25, with an average of



Figure 7. (a) the relationship between derived κ_{Org} and f_{44} ; (b) Comparison with previous studies.

513 0.08 ± 0.06 . This indicates that using a constant κ_{OA} value in the calculation of κ_{chem} would result 514 in a large bias. To further investigate the impact of OA oxidation level on κ_{OA} , we compared the 515 derived κ_{OA} against f_{44} , which is often used to represent the oxidation level of OA. Results show a 516 clear positive correlation (R=0.79) and a statistical relationship of $\kappa_{OA} = 1.04 \cdot f_{44} - 0.02$ (Fig.7a), 517 indicating that the degree of oxidation level is a crucial parameter determining the OA hygroscopicity. 518 Based on the relationship between f_{44} and O/C ratio for CV-ACSM (Hu et al., 2018b), 519 O:C=3.47× f_{44} +0.01, the relationship between κ_{OA} and O:C can be expressed as κ_{OA} =0.3×O:C - 520 0.02. The derived empirical relationship between κ_{OA} and f_{44} was compared to results in previous studies (Fig.7b). As mentioned in Sect.2.3, f₄₄ from CV-ToF-ACSM measurements is much higher than 521 those previously reported from AMS, but they are well correlated and the ratio between f44 of CV-ToF-522 ACSM and previous AMS instruments for ambient aerosol ranges from 1.5 to 2 with an average of 523 1.75. Therefore, to be consistent with the f_{44} in previous studies, the empirical relationship in Fig.7b is 524 changed to $\kappa_{OA} = 1.79 \cdot f_{44} - 0.03$. The κ_{OA} values are lower than that from the scheme of Chen et 525 al. (2017), but higher than those in Duplissy et al. (2011) and Mei et al. (2013a). In general, results of 526 most published studies about κ_{OA} demonstrate that hygroscopicity of organic aerosol generally 527 increases as the oxidation level of organic aerosol increases, however, the empirical mathematical 528 relationship differs much among different studies (Hong et al., 2018). These results highlight that more 529 530 studies are required to study the influence of OA oxidation level on κ_{OA} and to derive a more universal parameterization scheme that can be used in chemical transport models. 531

532 4.3 Distinct diurnal variations of κ_{0A} and its relationship with OOA

The time series of derived κ_{OA} are depicted in Fig.8a, which showed large fluctuations in a day. 533 534 The average κ_{OA} (Fig.8b) displays a distinct diurnal variation, with κ_{OA} reaching its minimum (0.02) 535 in the morning (near 07:30 LT) and increasing quickly to a maximum (0.19) near 14:30 LT. As a consequence, the water uptake abilities of organic aerosol particles changed from nearly hydrophobic 536 to moderately hygroscopic within 7 hours during daytime. Although previous results from observations 537 in Japan also revealed significant κ_{OA} diurnal variations, however, with daily minima in the afternoon 538 539 hours due to the increase of less oxygenated OA mass fractions (Deng et al. (2018) and Deng et al. (2019)), such large variability and significant diurnal variations of κ_{OA} were observed for the first 540 time on the NCP. We found that the diurnal profile of the mass fraction OOA in OA (f_{OOA}) was 541 remarkably similar to that of κ_{OA} (R=0.8, Figs. 8a and 8c), suggesting that OOA is very likely the 542 543 determining factor of κ_{OA} in winter on the NCP.



Figure 8. (a) Time series of derived κ_{0rg} and OOA mass fraction in NR-PM₁ (f_{00A}) in the right yaxis; (b) Average diurnal profile of κ_{0rg} ; (c) Scatter points of κ_{0rg} versus f_{00A} (%), and red line is the fitting line with linear regression.

544

The correlation coefficient between the average diurnal profiles of κ_{0A} and f_{00A} was 0.95, 545 which suggests that the variations in f_{OOA} was driving the significant diurnal variations of κ_{OA} . The 546 average diurnal variations of mass concentrations of identified OOA, HOA, COA, CCOA, BBOA, and 547 their mass fractions in total organic mass are shown in Fig.9a and Fig.9b, respectively. The mass 548 concentrations of HOA, CCOA and BBOA decreased rapidly from the morning time to 15:00 LT due 549 to the rising boundary layer height and also the decreased primary source emissions. The mass 550 concentrations of COA increased a little in the morning and then decreased quickly after 09:30 LT. 551 This transitory increase of COA in the morning might be associated with the cooking for breakfast. 552 However, the OOA mass increased rapidly from about 07:30 to 10:30 LT despite the boundary layer 553

development during period of time, and then remained almost constant thereafter. The rapid decreases in primary organic aerosol components and the increases in OOA concentration together resulted in a dramatic increase of f_{OOA} from ~10% at 9:00 to ~45% at 13:30 LT in the afternoon, which also



Figure 9. (a) Average diurnal profiles of mass concentrations of OOA, HOA, COA, CCOA, BBOA; (b) Average diurnal variations of mass fractions of OOA, HOA, COA, CCOA, BBOA.
corresponds to the similar increase in κ_{OA}. After 14:30 LT, the OOA mass concentration remained relatively unchanged, however, of the large increases in primary organic aerosol components also led to considerable decreases in f_{OOA} and κ_{OA}.

560 5 Conclusions

A field campaign was conducted at a rural site on the North China Plain in winter 2018. The major 561 instruments deployed were a humidified nephelometer system and a CV-ToF-ACSM for the 562 measurements of the bulk aerosol hygroscopicity of PM₁₀ and PM₁ and bulk aerosol chemical 563 compositions of PM_{2.5} and PM₁. The measured σ_{sp} at 525 nm of PM₁ and PM₁₀ in dry state ranged 564 from 11 to 1875 Mm^{-1} and from 18 to 2732 Mm^{-1} with average values of 550 and 814 Mm^{-1} , 565 respectively, suggesting a relatively polluted environment during this stusy. Retrieved $\kappa_{f(RH)}$ of PM₁₀ 566 and PM₁ ranged from 0.02 to 0.27, and from 0.03 to 0.26, with averages of 0.12 and 0.12, respectively. 567 568 The difference of $\kappa_{f(RH)}$ between PM₁₀ and PM₁ was found to be relatively small (3.5% on average), which was consistent with the physical and mathematical interpretation of $\kappa_{f(RH)}$. 569

A method for estimating κ_{OA} (organic aerosol hygroscopicity) base on f(RH) and bulk aerosol chemical composition measurements is developed. The key part of this method is that the size cut of

bulk aerosol chemical composition measurements should be PM₁ no matter the bulk $\kappa_{f(RH)}$ is 572 retrieved from light scattering enhancement measurements of PM₁ or PM₁₀. The derived κ_{OA} ranged 573 from 0.0 to 0.25 with an average of 0.08, highlighting that κ_{OA} displayed a large variability on the 574 575 NCP. Therefore, using a constant κ_{OA} could introduce a considerable uncertainty. in evaluating the climatic and environmental effects of organic aerosols The variation of κ_{OA} was highly and positively 576 correlated with the oxidation degree of OA, and κ_{OA} showed a distinct diurnal variation with the 577 minimum in the morning (0.02) and maximum in the afternoon (0.16). These results indicated the rapid 578 changes in hygroscopic properties of OA in a day by evolving from nearly hydrophobic to moderately 579 hygroscopic within 7 hours. The distinct diurnal variations of κ_{OA} were strongly associated with the 580 changes in f_{OOA}, suggesting that the rapid formation of OOA together with of the decreases in primary 581 582 organic aerosol during daytime together resulted in and the changes in κ_{0A} .

The large variability and distinct diurnal variations in κ_{OA} found in this study highlight an urgent need for more studies on the spatial and temporal variations of κ_{OA} on the NCP region, and also a better parameterization of κ_{OA} in chemical transport models to evaluate the impacts of OA on radiative forcing and CCN.

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Data availability. The data used in this study are listed in the references and a repository at 588 https://pan.baidu.com/s/16dOPuTQ568z5JRGF0jGLHQ (both python and matlab format), and also 589 590 available from the corresponding author upon request (kuangye@jnu.edu.cn) and (sunyele@mail.iap.ac.cn). 591

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

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595 Author Contributions. YK conceived and organized this paper. YC, HS, NM, YK and JT planned 596 this campaign. YK, YS and NM designed the experiments. YK and YH conducted the ACSM and 597 aerosol light scattering enhancement factor measurements. YZ and SZ conducted the particle number 598 size distribution measurements. JS and WY conducted the black carbon measurements. YH performed 599 the ACSM PMF analysis. WX, YH, YS, CZ, PZ and YC helped the data analysis, and WX helped 500 much in the language editing. YK, YH and YS prepared the manuscript with contributions from all co-501 authors. 602

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