1 A New Method for Calculating Number Concentrations of Cloud

2 Condensation Nuclei Based on Measurements of A Three-wavelength

3 Humidified Nephelometer System

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10 Abstract

The number concentration of cloud condensation nuclei (CCN) plays a fundamental role in 11 12 cloud physics. Instrumentations of direct measurements of CCN number concentration (N_{CCN}) based on chamber technology are complex and costly, thus a simple way for measuring $N_{\rm CCN}$ is needed. In 13 this study, a new method for $N_{\rm CCN}$ calculation based on measurements of a three-wavelength 14 humidified nephelometer system is proposed. A three-wavelength humidified nephelometer system 15 can measure aerosol light scattering coefficient (σ_{sp}) at three wavelengths and the light scattering 16 enhancement factor (fRH). The Angstrom exponent (Å) inferred from σ_{sp} at three wavelengths 17 provides information on mean predominate aerosol size and hygroscopicity parameter (κ) can be 18 calculated from the combination of fRH and Å. Given this, a look-up table that includes σ_{sp} , κ and 19 Å is established to predict N_{CCN} . Due to the precondition for the application, this new method is not 20 suitable for externally mixed particles, large particles (e.g. dust and sea salt) or fresh aerosol particles. 21 This method is validated with direct measurements of $N_{\rm CCN}$ using a CCN counter on the North China 22 23 Plain. Results show that relative deviations between calculated $N_{\rm CCN}$ and measured $N_{\rm CCN}$ are within 30% and confirm the robustness of this method. This method enables simpler $N_{\rm CCN}$ measurements 24 because the humidified nephelometer system is easily operated and stable. Compared with the 25

method of CCN counter, another advantage of this newly proposed method is that it can obtain $N_{\rm CCN}$ at lower supersaturations in the ambient atmosphere.

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29 1. Introduction

Cloud condensation nuclei (CCN) are the aerosol particles forming cloud droplet by hygroscopic 30 growth. CCN number concentration ($N_{\rm CCN}$) plays a fundamental role in cloud microphysics and 31 aerosol indirect radiative effect. In general, the direct measurement of $N_{\rm CCN}$ is achieved in a chamber 32 33 under super-saturated conditions (Hudson, 1989; Nenes et al., 2001; Rose et al., 2008). Due to the requirement of high accuracies of working conditions like temperatures, vapors and flow rates in 34 chambers, the direct measurement of $N_{\rm CCN}$ is complex and costly (Rose et al., 2008;Lathem and 35 36 Nenes, 2011). Thus, developments of simplified measurements of $N_{\rm CCN}$ are required. In recent years, attention has been focused on measurements of aerosol optical properties (Jefferson, 2010;Ervens et 37 al., 2007;Gasso and Hegg, 2003), which are simple and well-developed (Covert et al., 1972;Titos et 38 al., 2016). For aerosol population free of sea salt or dust, the accumulation mode aerosol not only 39 40 dominates aerosol scattering ability but also contribute most to $N_{\rm CCN}$. Thus, the calculation of $N_{\rm CCN}$ based on measurements of aerosol optical properties is feasible, and can facilitate $N_{\rm CCN}$ 41 measurement. 42

There are two kinds of methods to calculating $N_{\rm CCN}$ based on measurements of aerosol optical 43 44 properties. For the first kind, $N_{\rm CCN}$ as well as the hygroscopicity parameter (κ) can be calculated based on measurements of a humidified nephelometer system in combination with aerosol particle 45 number size distribution (PNSD) (Ervens et al., 2007; Chen et al., 2014). Thus additional 46 47 measurements of PNSD are needed. For the second kind, $N_{\rm CCN}$ is calculated based on statistical relationships between N_{CCN} and aerosol optical properties, such as scattering coefficient (σ_{sp}), 48 Angstrom Exponent (Å and single scattering albedo (SSA) (Jefferson, 2010;Shinozuka et al., 2015). 49 Å is the exponent commonly used to describe the dependence of σ_{sp} on wavelength as the formula 50 51 shows:

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$$\sigma_{\rm sp}(\lambda) = \beta \cdot \lambda^{-{\rm \AA}},$$
 (1)

53 where β is the aerosol number concentration. Coefficient of determination (R²) between measured

and calculated N_{CCN} using the first kind of method is about 0.9. For the second kind of method, R² is generally lower than 0.9, although the used instruments are cheaper and easier in operation. Applications similar to the second kind are widely used in remote sensing. As shown in Table 1, earlier studies found that the aerosol volume or aerosol PNSD retrieved from remote sensing measurements can be used to calculate N_{CCN} (Gasso and Hegg, 2003;Kapustin et al., 2006). Recently, either aerosol optical depth (AOD) or aerosol vertical profile is used to predict N_{CCN} directly(Ghan and Collins, 2004;Ghan et al., 2006;Andreae, 2009;Liu and Li, 2014).

In the statistical relationship between $N_{\rm CCN}$ and aerosol optical properties, $\sigma_{\rm sp}$ or AOD is 61 mainly the proxy of aerosol absolute concentration, while Å or SSA can be used to reveal the 62 variations of aerosol CCN activity, as shown in Table 1. Based on Kohler theory (Köhler, 63 64 1936; Petters and Kreidenweis, 2007), aerosol CCN activity is determined by aerosol size and aerosol chemical composition, and aerosol chemical composition can be defined as aerosol hygroscopicity. 65 Information about aerosol size and aerosol hygroscopicity are critical to N_{CCN} prediction and their 66 absence can lead to a deviation with factor of four (Andreae, 2009). Compared with aerosol 67 68 hygroscopicity, aerosol size is more important in determining CCN activity (Dusek et al., 2006). The value of Å can provide information on mean predominate aerosol size (Brock et al., 2016;Kuang et 69 al., 2017a). As a result, $N_{\rm CCN}$ calculation from Å and extinction coefficient is found to be accurate to 70 some extent (Shinozuka et al., 2015). As proxies for aerosol hygroscopicity, SSA or aerosol light 71 72 scattering enhancement factor (fRH) is commonly used while not so effective (Jefferson, 2010; Liu and Li, 2014). fRH is defined as: 73

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$$fRH=\sigma_{sp}(RH)/\sigma_{sp}$$
 (2)

where $\sigma_{sp}(RH)$ is the humidified σ_{sp} at a given RH. SSA is determined by the ratio between the light absorbing carbonaceous and less-absorbing components. Black carbon dominates the absorption of solar radiation and is a main hydrophobic components as well. Less-absorbing components consist of inorganic salts and acids, as well as most organic compounds, which are generally hygroscopic components. SSA correlates positively with aerosol hygroscopicity (Rose et al., 2010) but deviates significantly due to the diversity of hygroscopicity of less-absorbing components. Thus N_{CCN} calculation combining SSA, backscatter fraction and σ_{sp} still leads to

significant deviations, with $R^2 = 0.6$ (Jefferson, 2010). As for fRH, there was a study that applied 82 aerosol optical quantities (σ_{sp} or aerosol optical thickness) with fRH or SSA to calculate N_{CCN} (Liu 83 and Li, 2014). In their study, compared with the combination of SSA and aerosol optical quantities, 84 the combination of fRH and aerosol optical quantities is found to be less accurate in estimating $N_{\rm CCN}$, 85 even though fRH is directly connected with aerosol hygroscopicity (Liu and Li, 2014). This may 86 result from the significant dependence of fRH on aerosol size(Chen et al., 2014;Kreidenweis and 87 Asa-Awuku, 2014;Kuang et al., 2017a). As mentioned before, PNSD is used for better calculation of 88 κ and $N_{\rm CCN}$ from fRH in previous studies (Ervens et al., 2007; Chen et al., 2014). A new method to 89 estimate κ from fRH and Å was proposed recently (Kuang et al., 2017a;Brock et al., 2016). Based 90 on this method, fRH can be used to calculate N_{CCN} without measurements of PNSD and can be 91 92 expected to improve the $N_{\rm CCN}$ prediction just based on measurements of aerosol optical properties.

In this study, the relationship between N_{CCN} and aerosol optical properties measured by a humidified nephelometer system is studied and a new method for N_{CCN} prediction is proposed. This new method is validated based on data observed in Gucheng campaign on the North China Plain and can be expected to improve measurements of N_{CCN} due to advantages of applying nephelometers.

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98 2. Methodology

2.1. Data 99

Data in this study are mainly measured at Gucheng (39.15N, 115.74E) during autumn in 2016 on the North China Plain (NCP). Gucheng is 100km southwest from Beijing and 40km northeast from Baoding under background pollution condition in the NCP. The observation site was surrounded by farmland and about 3km away from the Gucheng town. This campaign started on 20 October and lasted for nearly one month.

Instruments used in Gucheng campaign were located in a measurement container under temperature maintained at 25 °C. Ambient aerosol was sampled and dried to relative humidity (RH) lower than 30% by an inlet system consisting of a PM10 inlet, an inline Nafion dryers and a RH and temperature sensor (Vaisala HMP110). Then the sample aerosol was separated by a splitter and directed into various instruments. During this campaign, σ_{sp} , fRH, particle size-resolved activation 110 ratio (AR) and particle number size distribution (PNSD) were obtained.

fRH as well as σ_{sp} at three wavelengths were measured by a humidified nephelometer system 111 consisting of two nephelometers (Aurora 3000, Ecotech Inc.) and a humidifier. In addition, Å can be 112 calculated directly from σ_{sp} measured by a nephelometer. The humidifier with a Gore-Tex tube 113 humidified the sample air up to 90% RH. A whole cycle of humidification lasted about 45minutes 114 from 50% RH to 90% RH. Dried σ_{sp} was obtained directly from dried sample aerosol measured by 115 116 one nephelometer and humidified σ_{sp} was obtained from humidified aerosol measured by another nephelometer. fRH can be calculated by dividing humidified σ_{sp} by dried σ_{sp} . Detailed description 117 of the humidified nephelometer system was illustrated in Kuang et al (2017a). 118

119 The particle size-resolved activation ratio (AR), defined as the ratio of $N_{\rm CCN}$ to total particles, was measured by a system mainly consisting of a differential mobility analyzer (DMA, Model 3081) 120 and a continuous-flow CCN counter (model CCN200, Droplet Measurement Technologies, USA; 121 Roberts and Nenes (2005); Lance et al., (2006)). The system selected mono-disperse particles with 122 the DMA coupled with an electrostatic classifier (model 3080; TSI, Inc., Shoreview, MN USA) and 123 measured AR of the mono-disperse particles by a condensation particle counter (CPC model 3776; 124 TSI, Inc.) and CCN counter. Ranges of particle size and supersaturation were 10-300nm and 125 0.07%-0.80%, respectively. Measurements at five supersaturations (0.07%, 0.10%, 0.20%, 0.40%) 126 127 and 0.80%) were conducted sequentially with each cycle lasted for 1 hour, and $N_{\rm CCN}$ at 0.07% supersaturation was used in this study. Due to non-idealities of CCN counter at supersaturations 128 lower than 0.10%, CCN measurement at 0.07% supersaturation was found to be the most uncertain 129 (Rose et al., 2008) and can lead to deviations of measured $N_{\rm CCN}$ in this study. Before and after the 130 131 campaign, supersaturations set in this system were calibrated using ammonium sulfate (Rose et al., 2008). More information about the system is available in Deng et al. (2011) and Ma et al.(2016). 132

PNSD with particle diameter from 9nm to 10um was measured by a mobility particle size
spectrometer (SMPS, TSI Inc., Model 3996) and an Aerodynamic Particle Sizer (APS, TSI Inc.,
Model 3321). SMPS consisted of a DMA, an electrostatic classifier and a CPC (model 3776; TSI,
Inc., Shoreview, MN USA) and measured PNSD with diameter lower than 700nm.

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7 In addition, PNSD and σ_{sp} from 2011 to 2014 at four campaigns (Wuqing in 2011, Xianghe in

138 2012 and 2013, and Wangdu in 2014) in NCP were used in this study. PNSD in these campaigns was 139 measured by a Twin Differential Mobility Particle Sizer (TDMPS, Leibniz-Institute for Tropospheric 140 Research (IfT), Germany) and an Aerodynamic Particle Sizer (APS, TSI Inc., Model 3321). A TSI 141 3563 nephelometer was used to obtain σ_{sp} at three wavelengths. Details about the four campaigns 142 can be found in Ma et al. (2011), Ma et al.(2016), Kuang et al. (2016) and Kuang et al.(2017a).

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144 2.2. Theories

Hygroscopic growth of particles at certain relative humidity can be described by κ -Köhler theory (Petters and Kreidenweis, 2007):

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$$\frac{\mathrm{RH}}{\mathrm{100}} = \frac{\mathrm{g(\mathrm{RH})^3 - 1}}{\mathrm{g(\mathrm{RH})^3 - (1 - \kappa)}} \cdot \exp(\frac{4\sigma_{\mathrm{s/a}} \cdot \mathrm{M_w}}{\mathrm{R} \cdot \mathrm{T} \cdot \mathrm{D}_{\mathrm{d}} \cdot \mathrm{g(\mathrm{RH})} \cdot \rho_{\mathrm{w}}})$$
(3)

where g(RH) is geometric diameter growth factor, κ is the hygroscopicity parameter, RH is the relative humidity; ρ_W is the density of water; M_W is the molecular weight of water; $\sigma_{s/a}$ is the surface tension of the solution–air interface, which is assumed to be equal to the surface tension of the pure water–air interface; R is the universal gas constant; and T is the temperature.

Accounting for the impact of Å, $\kappa_{\rm f}$ can be derived directly from fRH (Brock et al., 2016;Kuang et al., 2017a). A single-parameter parameterization scheme proposed by Brock et al. (2016) connects fRH and κ by the approximately proportional relationship between total aerosol volume and $\sigma_{\rm sp}$:

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$$f(RH)=1+\kappa_{sca} *RH/(100-RH)$$
 (4)

where κ_{sca} is a parameter for fitting fRH curves and is found can be used to predict κ_{f} in combination with Å in recent studies (Brock et al., 2016;Kuang et al., 2017a). This method of calculating κ_{f} based on κ_{sca} and Å was confirmed by good agreement with κ_{f} calculated from fRH and PNSD.

160 N_{CCN} can be calculated from size-resolved AR at a certain supersaturation (SS) and PNSD 161 (referred to as n(log D_{p})) as follows:

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$$N_{\text{CCN}} = \int_{\log D_{\text{P}}} \text{AR}(\log D_{\text{P}}, SS) \cdot n(\log D_{\text{P}}) d\log D_{\text{P}} \quad (5)$$

In general, size-resolved AR curves are complicated and always replaced by a critical diameter (D_c)
 to simplify calculation (Deng et al., 2013). The critical diameter is defined as:

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$$N_{\text{CCN}} = \int_{\log D_{\text{c}}}^{\log D_{\text{P,max}}} n(\log D_{\text{P}}) d\log D_{\text{P}}$$
(6)

where $D_{P,max}$ is the maximum diameter of the measured particle number size distribution. In other words, the integral of PNSD larger than D_c equals to the measured N_{CCN} . And a critical κ (κ_c) can be calculated by equation (3) and indicates CCN activity and hygroscopicity of particles.

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170 3. Results

171 3.1. Calculation of N_{CCN} based on measurements of a Humidified Nephelometer system

Free of sea salt aerosol and dust aerosol, accumulation mode aerosol dominates both the optical 172 scattering ability at short wavelengths and the CCN activity at low supersaturations, and thus a 173 reasonable relationship between σ_{sp} and N_{CCN} can be achieved. Figure 1 shows the size distribution 174 of cumulative contributions of σ_{sp} at 450nm and N_{CCN} at 0.07% with various Å and κ_c , and 175 corresponding normalized PNSDs based on data measured at the four campaigns on the North China 176 Plain. During the four campaigns, no sea salt aerosol or dust aerosol was observed(Ma et al., 177 178 2011; Ma et al., 2016; Kuang et al., 2016; Kuang et al., 2017a). For continental aerosol without sea salt or dust, Å varies from 0.5 to 1.8 and κ_c varies from 0.1 to 0.5 (Cheng et al., 2008;Ma et al., 179 2011; Liu et al., 2014; Kuang et al., 2017b). And as mentioned before, Å can be used as a proxy of 180 the overall size distribution of aerosol populations, with smaller Å indicating more larger particles. 181 In figure 1, comparisons for Å are made between 0.5 and 1.9 and for κ_c are made between 0.1 and 182 0.5. As larger particles contribute more to light scattering and CCN activation, cumulative 183 contributions of both σ_{sp} and N_{CCN} increase significantly at the diameter range of accumulation 184 mode particles. Because more hygroscopic particles are able to activate at smaller diameters, the 185 cumulative contribution of $N_{\rm CCN}$ with higher κ_c increases at smaller diameters. In general, major 186 contributions of both σ_{sp} and N_{CCN} are made by particles from 200nm to 500nm for various Å and 187 κ_c . This implies the feasibility of inferring $N_{\rm CCN}$ from aerosol optical properties. 188

Because particles smaller than 200nm can activate at supersaturations higher than 0.07% while scatter less light at wavelengths longer than 450nm, which are shown as the light color lines in Figure 1, it's obvious that significant differences will exist between cumulative contributions of σ_{sp} and N_{CCN} . This means σ_{sp} and N_{CCN} are dominated by different particles and poor correlation between σ_{sp} and N_{CCN} can be expected. Thus the method of inferring N_{CCN} from aerosol optical properties is applicable for shorter wavelength and lower supersaturations.

Furthermore, PNSD with higher Å indicates more Aitken mode particles and fewer 195 accumulation mode particles. Thus large particles contribute less for both σ_{sp} and N_{CCN} when Å are 196 higher, characterizing an increase of cumulative contribution curves at smaller diameters. In detail, 197 198 cumulative contribution curves of σ_{sp} at 1.9 Å is about 0.3 higher than these curves at 0.5 Å at the size range of 200nm to 700nm. While cumulative contribution curves of $N_{\rm CCN}$ at 1.9 Å is no higher 199 than 0.2 higher than these curves at 0.5 Å. Changes of cumulative contributions of $N_{\rm CCN}$ and $\sigma_{\rm sp}$ 200 with various Å reveal that the shape of PNSD can influence the correlation between N_{CCN} and σ_{sp} . 201 This is confirmed by previous studies in which the Å is found to play an important role in 202 calculating N_{CCN} from σ_{sp} (Shinozuka et al., 2015;Liu and Li, 2014). 203

The relationship between σ_{sp} and N_{CCN} dependent on Å and κ_c is evaluated by calculating 204 σ_{sp} and N_{CCN} with different PNSDs (classified by Å) and different κ_c . In detail, ratios of N_{CCN} to 205 σ_{sp} , referred to as AR_{sp}, are calculated to eliminate the effect of variations of particle concentrations 206 consistent at all diameters. Results at the supersaturation of 0.07% are shown in figure 2 and AR_{sp} is 207 higher than 0 and lower than 10. In general, AR_{sp} are higher for more hygroscopic particles or 208 smaller particles. As particles become more hygroscopic, more CCN can be expected when σ_{sp} is 209 fixed. As aerosol populations consist of more smaller CCN-active particles, the increase of σ_{sp} is 210 211 weaker than that of $N_{\rm CCN}$. For example, particles with diameters slightly larger than D_c contribute less to σ_{sp} than particles with diameters much larger than D_c . 212

In detail, the sensitivity of AR_{sp} to Å also changes with Å and κ_c . When Å are higher than 1.4 and κ_c is lower than 0.2, AR_{sp} is insensitive to Å. While when Å are lower than 1 and κ_c are higher than about 0.3, AR_{sp} is more sensitive to Å than κ_c . This higher sensitivity of AR_{sp} to Åreveals that, if the mean predominate size of particles is smaller, the increase of N_{CCN} due to the increase of Å mentioned in the former paragraph can be larger as a result. This is the consequence of the sensitivity of AR_{sp} to Å resulting from the variation of small CCN-active particles, as mentioned before.

Based on the lookup-table illustrated in Figure 2, N_{CCN} at the supersaturation of 0.07% can be calculated simply from Å, κ_f and σ_{sp} which can be obtained from measurements of a humidified nephelometer system. The description of this simple method is shown in figure 3. A new look-up table needs to be made for N_{CCN} estimation at other supersaturations, which should better be less than 0.07% as mentioned in the discussion of figure 1.

One critical issue about the method is the conversion of the κ_f obtained from the humidified 225 nephelometer system to the κ_c under super-saturated conditions. There are mainly two factors 226 making this conversion necessary. First, closure studies of aerosol hygroscopicity found significant 227 deviations between hygroscopicity at sub-saturated conditions and super-saturated conditions (Wex 228 et al., 2009; Irwin et al., 2010; Good et al., 2010; Renbaum-Wolff et al. 2016). Their difference can 229 230 be expected to be about 0.1 for accumulation mode aerosol(Wu et al., 2013;Whitehead et al., 2014; Ma et al., 2016). Second, κ_f indicates the hygroscopicity of total particles and can be quite 231 different from aerosol hygroscopicity at a specific diameter due to variations of size-dependent 232 particle hygroscopicity. Kuang et al. (2017a) found a difference around 0.1 between κ_f and κ 233 inferred from g(RH) measurements for accumulation mode particles whose κ_f is no larger than 0.2. 234 In this study, a simple conversion that κ_c is 0.2 higher than κ_f is used to calculate N_{CCN} , while for 235 κ_f larger than 0.2, a smaller difference of 0.1 between κ_c and κ_f should be used (Kuang et al., 236 2017a). This simplified relationship between κ_c and κ_f is a rough estimate regardless of the 237 complexity of differences of aerosol hygroscopicity measured by different instruments, but still used 238 239 in this study for two reasons. First, the accurate conversion cannot be achieved without detailed information of the particle hygroscopicity, which is difficult and complicated to measure. Second, a 240 deviation of κ_c less than 0.1 generally leads to a deviation of $N_{\rm CCN}$ less than 20% (Ma et al., 2016), 241

which is comparable with the deviation of CCN measurements. As a result, for a simple method of 242 $N_{\rm CCN}$ calculation, this simple conversion is applicable. In addition, it is important to note that the 243 244 value of the difference between κ_c and κ_f is also a rough estimate regardless of the complexity of aerosol hygroscopicity under different conditions, and the influence of $\Delta \kappa$ deviation on N_{CCN} 245 calculation needs to be further examined based on field observation. For fresh aerosol, the actual $\Delta \kappa$ 246 247 can be too large (about 4 times of kappa values for some organic components, Wex et al., 2009; Renbaum-Wolff et al., 2016) or too small (nearly zero for inorganic components and black carbon) 248 249 and thus is not suitable for the application of this method.

Besides aerosol size and hygroscopicity, aerosol mixing state can also affect aerosol CCN 250 activity. When primary aerosol emissions are strong, aerosol populations are likely to be externally 251 mixed and a realistic treatment of aerosol mixing state is critical for N_{CCN} calculation (Cubison et al., 252 2008; Wex et al., 2010). But for regions away from strong aerosol primary emissions, the influence of 253 mixing state on aerosol CCN activity is small and the assumption of internal mixing state is effective 254 for the estimation of N_{CCN} (Dusek et al., 2006;Deng et al., 2013;Ervens et al., 2010). For regions 255 256 above the boundary layer where clouds form and measurements of $N_{\rm CCN}$ are important, aerosol generally tends to be internally mixed when there is no strong vertical transport (McMeeking et al., 257 2011; Ferrero et al., 2014) and no plumes(Moteki and Kondo, 2007;McMeeking et al., 2011). In 258 addition, it should be noted that influences of aerosol hygroscopicity and aerosol size on aerosol 259 260 CCN activity are more significant than aerosol mixing state and the deviation of N_{CCN} calculation due to the assumption of aerosol mixing state is smaller than the deviation due to aerosol size and 261 aerosol hygroscopicity. In the new method of this paper, using Å and κ_c to indicate the influence of 262 aerosol size and aerosol hygroscopicity on aerosol CCN activity will increase the deviation of $N_{\rm CCN}$ 263 calculation, which is much larger than the deviation due to the assumption of aerosol mixing state. 264 As a result, the improvement of $N_{\rm CCN}$ calculation by introducing a more detailed mixing state than 265 internal mixing is limited and aerosol populations are assumed to be internally mixed for 266 simplification. Thus this method might not be applicable for regions or air masses greatly affected by 267 strong primary aerosol emissions. Furthermore, this new method cannot be applied for regions where 268 sea salt or dust prevails, as mentioned before. In summary, this method can be used to calculate $N_{\rm CCN}$ 269 for air mass tending to be dominated by aged aerosol particles like continental regions and clouds 270

271 forming heights.

272 3.2. Validation based on $N_{\rm CCN}$ measurement

273 The method for calculating N_{CCN} based on measurement of the humidified nephelometer system, including the conversion of κ_c and the lookup-table, is examined using data measured in Gucheng. 274 Overview of data in Gucheng is shown in Figure 4. From polluted periods to clean periods, 275 significant variations of N_{CCN} and σ_{sp} can be found but AR_{sp} of N_{CCN} to σ_{sp} stays around 5. On 276 October 23rd and 29th, N_{CCN} and σ_{sp} are lower than 2000#/cm³ and 500Mm⁻¹, respectively. While on 277 October 20th, 26th and November 3rd, $N_{\rm CCN}$ and $\sigma_{\rm sp}$ are higher than 2000#/cm³ and 500Mm⁻¹, 278 respectively. These variations of $N_{\rm CCN}$ and $\sigma_{\rm sp}$ are mainly due to the variation of the particle number 279 concentration rather than the shape of particle size distribution and aerosol hygroscopicity. Variations 280 of AR_{sp} result from the variations of Å and κ_c , which indicate the variations of aerosol 281 microphysical properties and chemical compositions. 282

In general, AR_{sp} is more sensitive to variations of Å than κ_c . As mentioned before, the 283 sensitivity of AR_{sp} to Å is determined by both Å and κ_f . In detail, Å during the campaign mainly 284 ranges from 0.5 to 1.5 and κ_f ranges mainly from 0.05 to 0.2, which means that κ_c ranges from 285 0.25 to 0.4. These values of Å and κ_f correspond to a significant sensitivity of AR_{sp} to Å, as the 286 lookup table shows in figure 2. The sensitivity of AR_{sp} to κ_c is much small and only notable during 287 some short periods (grey bars in Figure 4). For example, from November 5th to 7th, variations of κ_f 288 and Å are opposite and result in nearly constant AR_{sp} . And from October 30th to November 2nd, 289 consistent variations of Å and κ_f lead to greater variations of AR_{sp} than other periods. This weak 290 sensitivity of AR_{sp} to κ_f may be due to the uncertainty of κ_c calculated from κ_f based on the 291 simplified conversion. 292

Based on the lookup table of κ_c and Å, AR_{sp} is calculated and applied to calculate N_{CCN} with σ_{sp} . The calculated AR_{sp} and N_{CCN} are compared with the measured AR_{sp} and N_{CCN} shown as the green dots in Figure 5. In general, good agreements between calculations and measurements are achieved and relative deviations are within 30%. For the comparison of AR_{sp} , the system relative deviation is less than 10%. For the comparison of N_{CCN} , the slope and the correlation coefficient of the regression are 1.03 and 0.966, respectively.

299 In addition, the variation of $\Delta \kappa$ and its influence on AR_{sp} and N_{CCN} calculation are studied. As shown in Figure 6, $\Delta \kappa$ is around 0.2 and independent from Å and κ_c and over 80% of $\Delta \kappa$ ranges 300 from 0.1 to 0.3. A notable deviation of $\Delta \kappa$ can only be found when Å is higher than 1.5. High 301 values of Å represent existence of small particles, which tend to be fresh emitted and experience 302 inefficient aging processes. In this case, this simplified conversion of κ_c may not be applicable. 303 304 Furthermore, $\Delta \kappa$ with different values are applied in the new method to calculate N_{CCN} . In the first way, $\Delta \kappa$ of the κ_c conversion is set to be 0.05 higher or lower, which means $\Delta \kappa$ of 0.25 or 0.15. 305 The corresponding results are presented as the red dots and blue dots in Figure 5. In the second way, 306 307 a constant κ_c of 0.34, which is the average of κ_c values in Gucheng campaign, is used to calculate AR_{sp} and N_{CCN} , and shown as the grey dots in Figure 5. In general, differences among calculations 308 using various κ_c conversions are quite small. The $\Delta \kappa$ difference of 0.05 in κ_c conversion only 309 leads to a difference of 10% for the system relative deviation of calculated $N_{\rm CCN}$. The correlation 310 311 coefficient of the calculation using a constant κ_c is just a little lower than correlation coefficients of 312 calculations using a κ_c conversion. As a result, for data measured in Gucheng campaign, the method of calculating $N_{\rm CCN}$ is insensitive to the uncertainty of the $\kappa_{\rm c}$ conversion and a $\Delta \kappa$ of 0.2 is 313 applicable in this new method. 314

315 In this study, the insensitivity of calculated $N_{\rm CCN}$ to $\kappa_{\rm c}$ conversion is partly due to the small variation of κ_f during the campaign. However, the variation of κ_c can be quite large and cause 316 non-ignorable deviations of calculated $N_{\rm CCN}$. As previous studies of $N_{\rm CCN}$ measurement showed, the 317 variation of κ_c is often small and a constant κ_c can be used to calculate N_{CCN} accurately (Andreae 318 and Rosenfeld, 2008; Gunthe et al., 2009; Rose et al., 2010; Deng et al., 2013). Results in this study 319 320 are similar to these previous studies. But large variations of κ_c are also found in some other studies. In NCP, fluctuations of aerosol hygroscopicity during New Particle Formation events and soot 321 emissions lead to significant deviations of calculated $N_{\rm CCN}$ from average aerosol hygroscopicity (Ma 322 323 et al., 2016). Furthermore, the influence of κ_c cannot be ignored because the value of the average hygroscopicity is different in various regions during various periods. In summer of NCP, measured 324 $\kappa_{\rm f}$ at sub-saturated conditions can reach up to 0.45 when inorganic components dominate in particles 325

(Kuang et al., 2016). In this case, calculated $N_{\rm CCN}$ ignoring $\kappa_{\rm c}$ may be 10 times larger than measured N_{CCN}. To sum up, although the exact value of $\kappa_{\rm c}$ cannot be obtained from the measurement of the humidified nephelometer system, the influence of $\kappa_{\rm c}$ on $N_{\rm CCN}$ can be inferred and is found to be correct enough considering the convenience of this method. More data, especially in observations of more hygroscopic aerosol, is still needed to confirm this method.

331 4. Conclusions

 $N_{\rm CCN}$ is a key parameter of cloud microphysics and aerosol indirect radiative effect. Direct 332 measurements of N_{CCN} are generally conducted under super-saturated conditions in CCN chambers, 333 and are complex and costly. The aerosols of accumulation mode contribute most to both the aerosol 334 scattering coefficient and the aerosol CCN activity. In view of this, it is possible to predict $N_{\rm CCN}$ 335 336 based on relationships between aerosol optical properties and the aerosol CCN activity. In this study, a new method is proposed to calculate $N_{\rm CCN}$ based on measurements of a humidified nephelometer 337 system. In this method, N_{CCN} is derived from a look-up table which involves σ_{sp} , Å and κ_{f} , and 338 the required three parameters can be obtained from a three-wavelength humidified nephelometer 339 340 system.

341 Relationships between aerosol optical properties and aerosol CCN activity are investigated using datasets about aerosol PNSD measured during several campaigns in the North China Plain. The 342 relationship between σ_{sp} , Å, κ_c and N_{CCN} is analyzed. It is found that the ratio between N_{CCN} and 343 σ_{sp} , referred to as AR_{sp}, is determined by κ_c and Å. In light of this, it is possible to calculate N_{CCN} 344 based only on measurements of a three-wavelength humidified nephelometer system which provides 345 information about $\sigma_{sp}\,,$ the hygroscopicity parameter κ and Å. However, κ derived from 346 347 measurements of a humidified nephelometer system under sub-saturated conditions (termed as κ_f) differs from κ under super-saturated conditions which indicate CCN activity (termed as κ_c). As a 348 result, the conversion from κ_f to κ_c is needed. Based on previous studies of aerosol hygroscopicity 349 350 and CCN activity, a simple conversion from κ_f to κ_c with a fixed difference (referred to as $\Delta\kappa$) of 0.2 is proposed. On the basis of this simple conversion, the method of $N_{\rm CCN}$ prediction based only on 351 measurements of a humidified nephelometer system is achieved under conditions without sea salt 352

aerosol, dust aerosol, externally mixed aerosol or fresh aerosol.

This method is validated with measurements of a humidified nephelometer system and a CCN 354 counter in Gucheng in 2016. During the campaign, both $N_{\rm CCN}$ and $\sigma_{\rm sp}$ vary with the pollution 355 conditions. AR_{sp} is around 5 and changes with Å and κ_f . Based on this new method, N_{CCN} are 356 calculated to compare with its measured values. The agreement between the calculated $N_{\rm CCN}$ and the 357 measured $N_{\rm CCN}$ is achieved with relative deviations less than 30%. Furthermore, the variation of $\Delta \kappa$ 358 and its influence on N_{CCN} calculation are studied. The difference between κ_f and κ_c , was 0.2 ± 0.1 . 359 Sensitivity of calculated N_{CCN} to conversions from κ_{f} to κ_{c} is studied by applying different kinds of 360 conversions. Results show that calculated $N_{\rm CCN}$ varies little and is insensitive to the conversions, 361 362 which confirms the robustness and applicability of this newly proposed method.

This study has connected aerosol optical properties with $N_{\rm CCN}$, and also proposed a novel 363 364 method to calculate $N_{\rm CCN}$ based only on measurements of a three-wavelength humidified nephelometer system. Due to the simple operation and stability of the humidified nephelometer 365 system, this method will facilitate the real time monitoring of $N_{\rm CCN}$, especially on aircrafts. In 366 addition, measurements of the widely used CCN counter are limited to supersaturations higher than 367 0.07. In fogs and shallow layer clouds, supersaturations are generally smaller than 0.1% (Ditas et al., 368 2012; Hammer et al., 2014a, b; Krüger et al., 2014). For studying aerosol-cloud interaction, this 369 370 method is more applicable due to its applicability for calculating $N_{\rm CCN}$ at lower supersaturations than 1.0%. 371

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Aerosol PNSD (black lines), the cumulative contribution of σ_{sp} at wavelength of 450nm and 700nm (dark green lines and light green lines, respectively), the cumulative contribution of N_{CCN} at supersaturation of 0.07% (dark red and dark blue lines) and the cumulative contribution of N_{CCN} at supersaturation of 0.20% (light red and light blue lines) based on measurement in several campaigns in the North China Plain. Solid lines and dashed lines indicate Å of 1.9 and 0.5, respectively. Blue lines and red lines indicate κ_c of 0.1 and 0.5, respectively.



524

525 Figure 2.

526 Colors represent AR_{sp} (calculated as $AR_{sp} = \frac{N_{CCN}}{\sigma_{sp}}$ at 450nm wavelength and 0.07% supersaturation) 527 with different PNSDs (classified by Å values) and different κ_c .



528

529 Figure 3.







534 Overview of measurements in Gucheng in 2016. Upper plot: time series of $N_{\rm CCN}$ at the 535 supersaturation of 0.07% (red dots), $\sigma_{\rm sp}$ at the wavelength of 50nm (green dots) and their ratios 536 (black dots), referred to as AR_{sp}. Lower plot: time series of $\kappa_{\rm f}$ (red dots) and Å (green dots). The 537 grey bars are periods when the sensitivity of AR_{sp} to $\kappa_{\rm c}$ is notable.

538







541 Left plot: comparisons of calculated AR_{sp} and measured AR_{sp} with different conversions of κ_c from 542 κ_f . Right plot: regressions of calculated N_{CCN} and measured N_{CCN} with different conversions of κ_c 543 from κ_f .

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545

547 Differences between κ_c and κ_f , referred to as $\Delta \kappa$, with Å (positions of dots) and κ_f (colors of 548 dots). Bars represent percentages of $\Delta \kappa$ within different ranges.

549

Campaign	Air mass	Parameter	Caveats	Results	Reference
ICARTT ¹ in the north eastern USA and Canada	Polluted air mass	fRH and PNSD	Calculate N _{CCN} with aerosol hygroscopicity contrained by f(RH) and PNSD.	Predict $N_{\rm CCN}$ at SS > 0.3% with a 0.9 R ² .	Ervens et al., 2007

⁵⁴⁶ Figure 6.

HaChi ² on the North China Plain	Aged continental air mass	PNSD and fRH	Similar to Ervens et al., 2007. Calculate N_{CCN} with the hygroscopicity parameter constrained by f(RH) and PNSD.	Slopes around 1 and R ² around 0.9.	Chen et al., 2014
TARFOX ³ Atlantic seaboard and $ACE-2^4$	Polluted air mass	Retrieved aerosol volume from remote sensing	Predict $N_{\rm CCN}$ from aerosol volumes with empirical number-to-volume concentration ratio	Overestimate up to 5 times	Gasso and Hegg, 2003
ACE-2 in northeastern Atlantic	Diverse air mass	Backscatter or extinction profile. CCN at the surface.	Retrieve $N_{\rm CCN}$ profile from backscatter (or extinction) vertical profile assuming their ratios are the same to the ratio at surface, which can be calculated by backscatter (or extinction) and $N_{\rm CCN}$ measured at the surface	Predict $N_{\rm CCN}$ on most days for 0.1% SS and on 20%–40% of the days at 1% SS.	Ghan and Collins, 2004
ARM ⁵ Climate Research Facility central site at the Southern Great Plains	Continental air mass	Backscatter (or extinction) and RH profile. fRH and CCN at surface	Same as Ghan and Collins, 2004.	Explains CCN variance for 25%-63% of all measurements at high supersaturations	Ghan et al., 2006
TRACE-P ⁶ and ACE-Asia ⁷	Asian outflow over the western Pacific	Aerosol Index (AI, the product of ambient light extinction and Å)	Predict N_{CCN} based on empirical relationship between AI and N_{CCN}	AI relate well to CCN only with suitably stratified data	Kapustin et al., 2006
Multiple measurements	Diverse air mass	AERONET aerosol optical thickness (AOT)	Predict N _{CCN} based on empirical relationship between AOT and N _{CCN} as a power law	Predict N_{CCN} at SS > 0.3% with a 0.88 R ² , but have a factor-of-four range of N_{CCN} at a given AOT	Andreae, 2009

Four ARM sites	Polluted air mass	SSA, backscatter fraction and σ_{sp}	Estimate $N_{\rm CCN}$ from fitting parameters for the $N_{\rm CCN}$ activity spectra, which can be calculate based on their empirical relationships with aerosol optical properties.	Predict N_{CCN} with slopes around 0.9 and R^2 around 0.6.	Jefferson, 2010
Multiple ARM sites around the world	Diverse air mass	RH, fRH, SSA, AOT and σ_{sp}	Calculate N_{CCN} with σ_{sp} (or AOT) based on their empirical relationship, whose impact RH, fRH and SSA.	Achieve the best results by using σ_{sp} and SSA. Weakly affect on the σ_{sp} – N_{CCN} relationship by fRH. Deteriorate N_{CCN} –AOT relationship with increasing RH	Liu and Li, 2014
Multiple ARM sites around the world	Diverse air mass not dominated by dust	Å and extinction coefficient	Calculate N_{CCN} with light extinction based on their emperical relationship.	Deviate typically within a factor of 2.0.	Shinozuka et al., 2015

551 Table 1.

- 552 Review of studies that have used aerosol optical parameters to infer $N_{\rm CCN}$.
- ¹ International Consortium for Atmospheric Research on Transport and Transformation.
- 554 2 Haze in China.
- ³ Troposphere Aerosol Radiative Forcing Experiment.
- ⁴ Second Aerosol Characterization Experiment.
- ⁵ Atmospheric Radiation Measurement.
- ⁶ Transport and Chemical Evolution over the Pacific.
- ⁷ Aerosol Characterization Experiment–Asia.