

1 **Secondary aerosol formation alters CCN activity in the North China**
2 **Plain**

3 Jiangchuan Tao^{1, 2}, Ye Kuang^{1, 2}, Nan Ma^{1, 2}, Juan Hong^{1, 2}, Yele Sun^{3, 4, 5}, Wanyun Xu⁶, Yanyan
4 Zhang¹, Yao He³, Qingwei Luo¹, Linhong Xie^{1, 2}, Hang Su⁷, Yafang Cheng⁷

5 ¹Institute for Environmental and Climate Research, Jinan University, Guangzhou, Guangdong 511443,
6 China

7 ²Guangdong-Hongkong-Macau Joint Laboratory of Collaborative Innovation for Environmental
8 Quality, Guangzhou, China

9 ³State Key Laboratory of Atmospheric Boundary Layer Physics and Atmospheric Chemistry, Institute
10 of Atmospheric Physics, Chinese Academy of Sciences, Beijing 100029, China

11 ⁴College of Earth and Planetary Sciences, University of Chinese Academy of Sciences, Beijing 100049,
12 China

13 ⁵Center for Excellence in Regional Atmospheric Environment, Institute of Urban Environment,
14 Chinese Academy of Sciences, Xiamen 361021, China

15 ⁶State Key Laboratory of Severe Weather, Key Laboratory for Atmospheric Chemistry, Institute of
16 Atmospheric Composition, Chinese Academy of Meteorological Sciences, Beijing, 100081, china

17 ⁷Multiphase Chemistry Department, Max Planck Institute for Chemistry, Mainz 55128, Germany

18 Correspondence to: Jiangchuan Tao (taojch@jnu.edu.cn) and Nan Ma (nan.ma@jnu.edu.cn)

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20 **Abstract:**

21 Secondary aerosols (SA, including secondary organic and inorganic aerosols, SOA and SIA) are
22 predominant components of aerosol particles in the North China Plain (NCP) and its formation has
23 significant impacts on the evolution of particle size distribution (PNSD) and hygroscopicity. Previous
24 studies have shown that distinct SA formation mechanisms can dominate under different relative
25 humidity (RH). This would lead to different influences of SA formation on the aerosol hygroscopicity
26 and PNSD under different RH conditions. Based on the measurements of size-resolved particle
27 activation ratio (SPAR), hygroscopicity distribution (GF-PDF), PM_{2.5} chemical composition, PNSD,
28 meteorology and gaseous pollutants in a recent field campaign, McFAN (Multiphase chemistry
29 experiment in Fogs and Aerosols in the North China Plain), conducted during the autumn-winter
30 transition period in 2018 at a polluted rural site in the NCP, the influences of SA formation on cloud
31 condensation nuclei (CCN) activity and CCN number concentration (N_{CCN}) calculation under different
32 RH conditions were studied. Results suggest that during daytime, SA formation could lead to a
33 significant increase in N_{CCN} and a strong diurnal variation in SPAR at Super-saturations lower than
34 0.07%. During periods with daytime minimum RH exceeding 50% (high RH conditions), SA
35 formation significantly contributed to the particle mass/size changes in a broad size range of 150 nm
36 to 1000 nm, led to N_{CCN} (0.05%) increases within the size range of 200 nm to 300 nm, and mass
37 concentration growth mainly for particles larger than 300 nm. During periods with daytime minimum
38 RH below 30% (low RH conditions), SA formation mainly contributed to the particle mass/size and
39 N_{CCN} changes for particles smaller than 300 nm. As a result, under the same amount of mass increase
40 induced by SA formation, the increase of N_{CCN} (0.05%) was stronger under low RH conditions and
41 weaker under high RH conditions. Moreover, the diurnal variations of SPAR parameter (inferred from
42 CCN measurements) due to SA formation varied with RH conditions, which was one of the largest
43 uncertainties within N_{CCN} predictions. After considering the SPAR parameter (estimated through the
44 number fraction of hygroscopic particles or mass fraction of SA), the relative deviation of N_{CCN} (0.05%)
45 predictions were reduced to within 30%. This study highlights the impact of SA formation on CCN
46 activity and N_{CCN} calculation, which provides guidance for future improvements of CCN predictions
47 in chemical-transport models and climate models.

48 **1. Introduction**

49 Cloud condensation nuclei (CCN) activity of aerosol particles describes its ability to activate
50 and grow into cloud droplets at given supersaturations and thus has important impacts on cloud
51 microphysics and the aerosol indirect effect on climate. CCN activity is dependent on the
52 physicochemical properties of aerosol particles, including particle size distributions, hygroscopicity
53 (determined by chemical composition) and mixing state. Thus, atmospheric processes influencing
54 these aerosol properties may exert influences on CCN activity.

55 Secondary aerosols (SA) formation contributes greatly to aerosol populations and impacts
56 CCN properties in many ways, generally increasing CCN number concentrations (N_{CCN}) and leading
57 to changes in the CCN activity (Wiedensohler et al., 2009; Kerminen et al., 2012; Wu et al., 2015;
58 Farmer et al., 2015; Ma et al., 2016; Zhang et al., 2019 and reference therein). Differences in precursor
59 and oxidant concentrations as well as SA formation mechanisms lead to particle size growth in
60 different size ranges (Dal Maso et al., 2005; Kulmala et al., 2007; Zhang et al., 2012; Farmer et al.,
61 2015; Cheng et al., 2016; Kuang et al., 2020c), thus would impact CCN activities in different ways.
62 SA formation includes both the formation and subsequent growth of new particles (New Particle
63 Formation, NPF), and the growth of existing particles. NPF can directly provide particles large enough
64 to act as CCNs (Wiedensohler et al., 2009; Kerminen et al., 2012; Farmer et al., 2015), generally
65 affecting aerosol particles smaller than 100 nm, thereby elevating N_{CCN} at higher supersaturations
66 ($SSs > 0.2\%$) (Wiedensohler et al., 2009; Kerminen et al., 2012; Ma et al., 2016; Zhang et al., 2019 and
67 reference therein). SA formation on existing particles, especially under polluted conditions,
68 significantly adds mass to and changes the chemical composition of accumulation mode particles
69 (Farmer et al., 2015), thus affecting CCN at lower $SSs (< 0.2\%)$ (Wiedensohler et al., 2009; Mei et al.,
70 2013; Yue et al., 2016; Thalman et al., 2017; Duan et al., 2018). SSs varies greatly among different
71 clouds categories. Cumulus clouds are formed under higher SSs and are thus mostly influenced by
72 Aitken mode particles formed in NPF events (Reuter et al., 2009; Gryspeerd and Stier, 2012; Fan et
73 al., 2016; Jia et al., 2019 and reference therein). Stratus clouds and fogs that exert stronger effects on
74 climate and environment, however, are generally formed at SSs lower than 0.2%, indicating that only
75 accumulation mode particles can serve as CCN (Ditas et al., 2012; Hammer et al., 2014a, b; Krüger et
76 al., 2014; Shen et al., 2018). Numerous studies have investigated the impact of NPF on CCN (Gorden
77 et al., 2016; Ma et al., 2016; Yu et al., 2020 and reference therein), however, only few studies have
78 focused on the influence of SA formation on CCN activity of accumulation mode particles, which
79 might exhibit strong climate and environment impacts and urgently requires attention.

80 SA formation affects CCN activity of accumulation mode particles not only by enlarging their
81 size, but also by changing their chemical compositions. At a specific particle size, the CCN activity is
82 determined both by the chemical composition of particles, which originally were and stayed this size,
83 and that of particles, which grew into this size via added SA mass. These two groups of particles can
84 exert different variations to CCN activity at the same particle size (Wiedensohler et al., 2009 and
85 reference therein). In general, the SA formation can increase the hygroscopicity of particles by adding
86 chemical compounds with lower volatility and higher oxidation state, which are usually more
87 hydrophilic, thereby enhancing CCN activity of accumulation mode particles (Mei et al., 2013; Yue et
88 al., 2016). However, CCN activity may also remain unchanged (Wiedensohler et al., 2009) or be
89 weakened in some cases (Thalman et al., 2017; Duan et al., 2018). In SA formation observed in central
90 Amazon forests, Thalman et al. (2017) reported enhanced CCN activity in dry season while constant
91 CCN activity in wet season. In SA formation events under polluted conditions in Guangzhou (Pearl
92 River Delta, China), Duan et al. (2018) found that bulk CCN activity can be enhanced in summer due
93 to the formation of large and inorganic-rich particles, but weakened in winter due to the formation of
94 small and organic-rich particles, where RH seemed to have been an important factor in the variations
95 of bulk CCN activity due to different particle formation pathways. Aside from variations of particle
96 chemical composition, changes in aerosol mixing states caused by SA formation can also change CCN
97 activity (Su et al., 2010; Rose et al., 2011; Cheng et al., 2012). The fast condensation of SA components
98 on accumulation mode particles led to the turnover of soot particle mixing state from externally to
99 internally mixed, which contributed mostly to enhancements of CCN activity (Cheng et al., 2012).
100 Thus, SA formation influences the CCN activity of accumulation mode particles through its integrated
101 impacts on their size, hygroscopicity and mixing state, which requires more detailed and
102 comprehensive investigations.

103 The North China Plain (NCP) frequently experiences severe aerosol pollution due to both
104 strong emissions of primary aerosol and strong SA formation caused by the abundance of gaseous
105 precursors and oxidants (Zheng et al. ACP, 2015; Liu et al., 2010; Huang et al., 2014; Xu et al., 2019).
106 In the SA formation events on the NCP, both aqueous-phase processes and gas-phase photochemical
107 processes can play important roles, depending on atmospheric conditions such as RH (Hu et al., 2016;
108 Xu et al., 2017a; Wang et al., 2019). A recent observational study on the NCP found that SA formation
109 dominantly contributed to different particle size since SA formation mechanisms varied with RH
110 conditions (Kuang et al., 2020c). Under dry conditions, SA were mainly formed through gas-phase
111 photochemical processing and mostly added mass to accumulation mode particles. While under high
112 RH conditions or super-saturated conditions, SA was also formed in aqueous phase, contributing to

113 the formation of both accumulation mode and coarse mode particles. The difference in particle size
114 where SA formation took place and the difference in SA chemical compositions could result in distinct
115 variations of CCN activity, which has not been evaluated yet. In this study, we will study the influence
116 of SA formation on Size-resolved Particle Activation Ratio (SPAR) of accumulation mode particles
117 in the NCP under different RH conditions, which fills a gap of knowledge within CCN studies in the
118 NCP and may provide guidance for the improvement of current CCN parameterization schemes in
119 chemical-transport and climate models.

120 **2. Method:**

121 2.1. Measurements

122 2.1.1. Site

123 Under the framework of McFAN (Multiphase chemistry experiment in Fogs and Aerosols in
124 the North China Plain) (Li et al., 2021), from 16th November to 16th December 2018, physical and
125 chemical properties of ambient aerosol particles as well as meteorological parameters were
126 continuously measured at the Gucheng site in Dingxing county, Hebei province, China. This site is an
127 Ecological and Agricultural Meteorology Station (39°09'N, 115°44'E) of the Chinese Academy of
128 Meteorological Sciences, which is located between Beijing (~100km) and Baoding (~40km), two
129 mega cities in the North China Plain, and surrounded by farmlands and small towns. Measurements at
130 this site can well represent the polluted background conditions of the NCP. All aerosol measurement
131 instruments were placed in a container with temperature maintained at 24 °C, while conventional trace
132 gas instruments including CO were housed in an air-conditioned room on a two-story building located
133 ~80 meters to the south of the container, with no taller buildings between them blocking the air flow.

134 2.1.2 Instrumentation

135 In this study, ambient aerosol was sampled by an inlet system consisting of a PM10 inlet
136 (Rupprecht & Patashnick Co., Inc., Thermo, 16.67 L/min), a Nafion dryer that dried relative humidity
137 to below 30% and an isokinetic flow splitter directing the air sample to each instrument.

138 A DMA-CCNC system measured SPAR at five supersaturations (SSs), 0.05%, 0.07%, 0.2%,
139 0.44% and 0.81%, with a running time of 20 min for 0.05% and 10 min for the other SSs. This system
140 consisted of a differential mobility analyzer (DMA model 3081; TSI, Inc, MN USA), a condensation
141 particle counter (CPC model 3772; TSI, Inc., MN USA) and a continuous-flow CCN counter (model
142 CCN200, Droplet Measurement Technologies, USA; Roberts and Nenes, 2005). The system was
143 operated in a size-scanning mode over the particle size range from 9 to 400 nm. SPAR can be obtained
144 by combining the measurements of CPC and CCNC at different particle size. The sample and sheath

145 flow rate of the DMA were set to 1 lpm and 5 lpm, respectively, hence the resultant measured particle
146 diameter ranged from 9 nm to 500 nm. Since the low number concentration of particles above 300 nm
147 could lead to large uncertainty in CCNC counting, the measurements for particles larger than 300 nm
148 were excluded, except for 0.05% SS. In order to characterize the variations of particles with low
149 hygroscopicity of about 0.1, SPAR measurement up to about 400 nm is used at 0.05% SS. There are
150 12 size distribution scans during a complete 1-hour cycle, with four scans for first SS and two scans
151 for each of the rest four SSs. Only the last scan for each SS is used as the CCNC needs time for SS
152 stabilization. The SSs of CCNC were calibrated with monodispersed ammonium sulphate particles
153 (Rose et al., 2008) both before and after the campaign. The flowrates were checked regularly (every
154 few days) during the campaign, as the flows (sample flow and sheath flow) of the instrument can affect
155 both the counting of droplets and the SS in the column. A modified algorithm based on Hagen and
156 Alofs (1983) and Deng et al. (2011, 2013) was used to correct the influence of multiple-charge particles
157 and DMA transfer function on SPAR. Details about the system are described in Ma et al. (2016) and
158 the description about the inversion method can be found in the supplements.

159 Non-refractory particulate matter (NR-PM) including SO_4^{2-} , NO_3^- , NH_4^+ , Cl^- and organics with
160 dry aerodynamic diameters below $2.5 \mu\text{m}$ was measured by an Aerodyne Time-of-Flight Aerosol
161 Chemical Speciation Monitor (ToF-ACSM hereafter) equipped with a $\text{PM}_{2.5}$ aerodynamic lens
162 (Williams et al., 2010) and a capture vaporizer (Xu et al., 2017b; Hu et al., 2017a) at 2-minute time
163 resolution. The ToF-ACSM data were analyzed with the standard data analysis software (Tofware
164 v2.5.13; <https://sites.google.com/site/ariacsm/>, last access: 21 January 2020). The organic mass spectra
165 from m/z 12 to 214 were analyzed with an Igor Pro based positive matrix factorization (PMF)
166 evaluation tool (v3.04) and then evaluated following the procedures described in Zhang et al. (2011).
167 The chosen five-factor solution includes four primary factors i.e. hydrocarbon-like OA (HOA),
168 cooking OA (COA), biomass burning OA (BBOA), and coal combustion OA (CCOA), and a
169 secondary factor, i.e. oxygenated OA (OOA). More detailed descriptions on the ACSM measurements
170 and data analysis can be found in Kuang et al. (2020b) and Sun et al. (2020).

171 A Humidified Tandem differential mobility analyzer (HTDMA, Tan et al., 2013) measured the
172 size-resolved aerosol growth factor (GF) at 90% RH. The sampled particles were subsequently charged
173 by a neutralizer (Kr85, TSI Inc.) and size selected by a DMA (DMA1, model 3081L, TSI Inc.). A
174 Nafion humidifier (model PD-70T-24ss, Perma Pure Inc., USA) was used to humidify the
175 monodisperse particles with a specific diameter (D_d) at a fixed RH of $(90 \pm 0.44) \%$ and then the
176 number size distribution of the humidified particles (D_{wet}) was measured by another DMA (DMA2,

177 model 3081L, TSI Inc.) and a condensation particle counter (CPC, model 3772, TSI Inc.). Thus, GF
178 of the particles can be calculated as:

$$179 \quad GF = \frac{D_{wet}}{D_d} \quad (1)$$

180 During the campaign, four dry mobility diameters (60, 100, 150, and 200 nm) were selected for the
181 HTDMA measurements. A full scan takes about 1 hour in order to cover the four sizes. Regular
182 calibration by using standard polystyrene latex spheres and ammonium sulfate were performed to
183 ensure the instrument functioned normally. The tandem differential mobility analyzer (TDMA)
184 inversion algorithm (Gysel et al., 2009) was applied to calculate the Probability Density Function of
185 GF (GF-PDF). More details about this system can be found in Cai et al. (2018) and Hong et al. (2018).

186 Particle number size distributions (PNSDs) were measured by combining the measurements of
187 a scanning mobility particle sizer (SMPS, TSI model 3080) and an aerodynamic particle sizer (APS,
188 TSI Inc., Model 3321), that measured particle mobility diameter size distributions in the range of 12
189 nm to 760 nm and particle aerodynamic diameter size distribution in the range of 700 nm to 10 μ m,
190 respectively. A commercial instrument from Thermo Electronics (Model 48C) was used to measure
191 CO concentration. Besides monthly multipoint calibrations and weekly zero-span check, additional 6-
192 hourly zero checks were also performed for the CO instrument.

193 2.2. Data processing

194 2.2.1. Aerosol hygroscopicity and cloud activation: κ -Köhler theory

195 The ability of particles to act as CCN and its dependence on particle size and particle chemical
196 composition on CCN activity can be described by the Köhler theory (Köhler, 1936). A hygroscopic
197 parameter κ is calculated based on the κ -Köhler theory (Petters and Kreidenweis, 2007) to evaluate
198 the influence of particle chemical compositions:

$$199 \quad \kappa = \left(\frac{D_{wet}^3 - D_d^3}{D_d^3} \right) \left[\frac{1}{S} \exp \left(\frac{4\sigma_{s/a} M_w}{RT\rho_w D_{wet}} \right) - 1 \right], \quad (1)$$

200 where S represents the saturation ratio, ρ_w is the density of water, M_w is the molecular weight of water,
201 $\sigma_{s/a}$ is the surface tension of the solution/air interface, R is the universal gas constant, T is the
202 temperature, D_d is the diameter of dry particle and D_{wet} is the diameter of the humidified particle. In
203 this study, $\sigma_{s/a}$ is assumed to be the surface tension of pure water/air interface. Based on the κ -Köhler
204 theory, the surface equilibrium water vapor saturation ratio of particles with a specific κ at different
205 wet particle size can be calculated, and the maximum value of surface equilibrium saturation ratio
206 (which is generally supersaturated) is defined as the critical SS for CCN activation. As a result, the

207 variation of the critical diameter (D_a) for particles with different hygroscopicity (or GF at a specific
208 RH) at different SSs can be determined.

209 2.2.2. Aerosol growth factor and its probability density function

210 In practice, the growth factor probability density function (GF-PDF) was inverted from the
211 measured GF distribution using a TDMA_{inv} algorithm (Gysel et al., 2009). After obtaining the GF-
212 PDF, the ensemble average GF and corresponding critical diameter under a certain SS ($D_{a,GF}$) can be
213 calculated. Furthermore, the number fraction and the weighted-average GF of hygroscopic particles
214 ($\kappa > 0.1$ and $GF(90\%, 200 \text{ nm}) > 1.22$) were calculated as:

$$215 \quad NF_{\text{hygro}} = \int_{1.2}^{\infty} \text{PDF}(\text{GF}) \times d\text{GF} \quad (2)$$

$$216 \quad GF_{\text{hygro}} = \int_{1.2}^{\infty} \text{GF} \times \text{PDF}(\text{GF}) \times d\text{GF} \quad (3)$$

217 Based on the κ -Köhler theory, the hygroscopicity parameter κ and corresponding critical diameter
218 ($D_{a,\text{hygro}}$) under a certain SS for particles with GF_{hygro} can be calculated. As GF_{hygro} is higher than the
219 average GF, $D_{a,\text{hygro}}$ is smaller than $D_{a,GF}$.

220 2.2.3 Calculations of aerosol hygroscopicity from aerosol chemical-composition measurements

221 For the calculation of aerosol hygroscopicity parameter κ based on measured chemical
222 composition data (κ_{chem}), detailed information on the chemical species are needed. The ACSM can
223 only provide bulk mass concentrations of SO_4^{2-} , NO_3^- , NH_4^+ , Cl^- ions and organic components, which
224 cannot be used to calculate size resolved hygroscopicity. However, in the North China Plain,
225 accumulation mode particles are the dominant contributors to the bulk particle mass concentration (Liu
226 et al., 2014; Xu et al., 2015; Hu et al., 2017b) and thus the bulk chemical compositions can be used as
227 a proxy for that of accumulation mode particles. For the inorganic ions, a simplified ion pairing scheme
228 was used to convert ion mass concentrations to mass concentrations of corresponding inorganic salts
229 (Gysel et al., 2007; Wu et al., 2016). Thus, mass concentrations of SO_4^{2-} , NO_3^- , NH_4^+ and Cl^- are
230 specified into ammonium sulfate (AS), ammonium nitrate (AN), ammonium chloride (AC) and
231 ammonium bisulfate (ABS), for which the κ values under super-saturated conditions were specified
232 according to Petters and Kreidenweis (2007). For a given internal mixture of different aerosol chemical
233 species, the Zdanovskii–Stokes–Robinson (ZSR) mixing rule can be applied to predict the overall κ_{chem}
234 using volume fractions of each chemical species (ε_i) (Petters and Kreidenweis, 2007):

$$235 \quad \kappa_{\text{chem}} = \sum_i \kappa_i \cdot \varepsilon_i \quad (4)$$

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

$$238 \quad \kappa_{\text{chem}} = \kappa_{\text{AS}}\varepsilon_{\text{AS}} + \kappa_{\text{AN}}\varepsilon_{\text{AN}} + \kappa_{\text{ABS}}\varepsilon_{\text{ABS}} + \kappa_{\text{AC}}\varepsilon_{\text{AC}} + \kappa_{\text{BC}}\varepsilon_{\text{BC}} + \kappa_{\text{Org}}\varepsilon_{\text{Org}} \quad (5)$$

239 where κ_{BC} is assumed to be zero as black carbon is non-hygroscopic hydrophilic. κ_{org} and ε_{org} represent
240 κ and volume fraction of total organics. The values of hygroscopicity parameter for inorganic
241 compounds can be found in Table 1 of Kuang et al. (2020b). Large variations in κ_{org} has been reported
242 in former studies and a linear relationship between κ_{org} and organic aerosol oxidation state (f44) was
243 detected in our campaign (Kuang et al., 2020b), which was adopted to calculate κ_{org} in this study:

$$244 \quad \kappa_{\text{Org}} = 1.04 \times f44 - 0.02 \quad (6)$$

245 It should be noted that the κ -Köhler theory is not perfect, even for inorganic compounds.
246 Numerous studies have been focusing on the performance of its application on measurements under
247 different RH conditions (Liu et al., 2011; Wang et al., 2017). And κ_{org} used in this study was
248 determined by the measurement of humidified nephelometer at RH of 85% in Petters and Kreidenweis,
249 (2007), due to the lack of κ_{org} measured under super-saturated conditions. In this study, we focus on
250 the variations of κ values derived from HTDMA and CCN measurement during the SA formation
251 events, rather than the closure between κ values derived using different techniques, which will be
252 addressed in an upcoming study.

253 2.2.4. Fitting parameterization scheme of SPAR

254 In general, the variation in CCN activity of a particle population can be attributed to the
255 variation in the number fraction of hygroscopic particles or its hygroscopicity, which can be indicated
256 by fitting parameters of SPAR curves parameterization. SPAR curves are often parameterized using a
257 sigmoidal function with three parameters. This parameterization assumes aerosols to be an external
258 mixture of apparently hygroscopic particles that can act as CCN and non-hygroscopic particles that
259 cannot be measured by CCNC within the measured particle size range below 400 nm (Rose et al.,
260 2010). SPAR ($Ra(D_p)$) at a specific SS can be described as follows (Rose et al., 2008):

$$261 \quad Ra(D_p) = \frac{\text{MAF}}{2} \left(1 + \text{erf} \left(\frac{D_p - D_a}{\sqrt{2}\pi\sigma} \right) \right) \quad (7)$$

262 where erf is the error function. MAF is the asymptote of the measured SPAR curve at large particle
263 sizes. D_a is the midpoint activation diameter and is associated with the hygroscopicity of CCNs. σ is
264 the standard deviation of the cumulative Gaussian distribution function and indicates the heterogeneity
265 of CCN hygroscopicity. As reported by Jiang et al. (2021), based on the investigation of the

266 covariations between SPAR curves and parameterized hygroscopicity distribution, it was found that
267 the MAF can be used to estimate the number fraction of hygroscopic (thus CCN-active) particles, for
268 aerosol hygroscopicity distributions generally observed in ambient atmosphere, and thus half MAF
269 can be used to represent the number fraction of CCNs to total particles at particle size around D_a .
270 Although the influence of particles whose κ is less than 0.1 on SPAR cannot be considered in this
271 parameterization scheme, significant deviation were only found under higher SSs (Tao et al., 2020)
272 and need not to be considered under the low SSs discussed in this study.

273 To be noted, the meaning of MAF can be different regard to the SS, and SPAR measurement
274 up to about 400 nm is needed for the MAF fitting for SPAR at SS of 0.05% to represent the particles
275 with κ value higher than 0.1. For SPAR at SS of 0.8%, MAF should be 1 at 400 nm diameter. However,
276 a MAF of 1 in this case can lead to overestimations of hygroscopic particle number fraction due to the
277 significant difference between SPAR curves and sigmodal fitting curves. In the former study on SPAR
278 fitting curves in the NCP, it was found that a fitting parameterization with the combination of two
279 sigmodal fitting curves was needed for SPAR fitting at SSs higher than 0.4% (Tao et al., 2020).
280 However, in this study, we investigate SA formation on accumulation mode particles and particle CCN
281 activity at SSs below 0.1%, under which condition non-hygroscopic particles smaller than 400 nm are
282 typically CCN-inactive. The MAF fitted in the particle size range below 400 nm was used to indicate
283 the variations of SPAR that was of the main focus here in this work. In addition, due to the very low
284 N_{CCN} in particle size ranges larger than 400 nm, the deviations of N_{CCN} due to the limited range of
285 measured particle size are also very small.

286

287 3. Results

288 3.1. Overview of the measurements

289 The timeseries of meteorological parameters, SPAR, N_{CCN} at SS of 0.05% and mass
290 concentration of Non-refractory particulate matter of $PM_{2.5}$ (NR- $PM_{2.5}$), $PM_{2.5}$ SA ([inorganic](#)
291 [compounds and OOA](#)) and $PM_{2.5}$ PA (primary aerosol, defined as the sum of POA) are shown in Fig.
292 1. [The mass concentration of OOA and four POA were quantified by the ACSM PMF analysis \(Zhang](#)
293 [et al. 2011\)](#). During the campaign, $PM_{2.5}$ PA were generally lower than $100 \mu g m^{-3}$ under both high
294 and low RH periods. Meanwhile, $PM_{2.5}$ SA can approach about $400 \mu g m^{-3}$, especially during the
295 strong SA formation events under high RH conditions, but can be lower than $100 \mu g m^{-3}$ under low
296 RH conditions. Strong diurnal variations were found in SPAR with varying meteorological parameters.
297 During the whole period, the wind speed was generally lower than $4 m s^{-1}$, which is in favor of aerosol
298 particle accumulation and SA formation on existing particles. However, RH, N_{CCN} ([0.05%](#)), $PM_{2.5}$ SA
299 and NR- $PM_{2.5}$ mass concentrations revealed very distinct levels before and after 4th Dec, and thus the
300 whole campaign was divided into two stages with different RH and SA pollution conditions: higher
301 RH and stronger SA pollution before 4th Dec, and lower RH and lighter SA pollution after 4th Dec. In
302 the following discussions, the high RH stage corresponds to days before 4th Dec with daily maximum
303 and minimum RH higher than 75% and 50%, respectively. Two events that occurred during 25th Nov
304 to 27th Nov (Event 1) and 30th Nov to 2nd Dec (Event 2), respectively, displayed especially high RH
305 conditions with successive nighttime fogs (blue shaded areas). The low RH stage corresponds to the
306 period after 4th Dec with daily maximum and minimum RH below 70% and 30%, which was
307 represented by two events that occurred during 9th Dec to 11th Dec (Event 3) and 13th Dec to 15th Dec
308 (Event 4), respectively. These events were selected based on the similarity of $PM_{2.5}$ concentration and
309 evolution, while the time window was fixed to two days for the convenience of intercomparing. In
310 addition, during these events, the wind speed was generally low, the RH followed a general diurnal
311 variations and SA mass grew steadily and continuously. Thus the interference of the variations of air
312 mass and short-term local emissions can be eliminated and the influence of SA formation can be
313 highlighted. It should be noted that variations of N_{CCN} at 0.07% were similar to those at 0.05%, which
314 followed the variations of SA mass concentration. While at higher SSs, the variations of N_{CCN} differed
315 from those of SA mass concentration, especially under high RH conditions, suggesting different
316 responses of CCN activity towards distinct SA formation processes. As reported in Kuang et al.
317 (2020c), during the high RH stage aqueous phase SA formation was promoted, leading to persistent
318 increases in N_{CCN} ([0.05% and 0.07%](#)), mass concentration of NR- $PM_{2.5}$ and especially mass
319 concentration of $PM_{2.5}$ SA during Event 1 and 2. During the low RH stage, the SA formation

320 dominantly occurred in the gas-phase, that generated much less SA than aqueous-phase formation
321 (Kuang et al., 2020c). Thus, the persistent increases of N_{CCN} (0.05% and 0.07%) and $PM_{2.5}$ during
322 Event 3 and 4 was much weaker than those in Events 1 and 2. Due to the different SA mass fractions,
323 SPAR during the high RH stage was generally higher than that during the low RH stage. However, the
324 ratios between N_{CCN} (0.05%) and mass concentration of $PM_{2.5}$ SA or NR- $PM_{2.5}$, were lower during the
325 high RH period and demonstrated strong decreases, especially in Event 1 and 2. The response of CCN
326 activity and N_{CCN} (0.05%) to the different SA formation mechanisms will be discussed
327 comprehensively in the following parts.

328 3.2. The influence of different secondary aerosol formation on the diurnal variation of CCN 329 activity

330 The diurnal averages of PNSD, SPAR at SS of 0.05%, GF-PDF for 200 nm particle and mass
331 fraction of particle chemical compositions during high RH periods before 4th Dec, low RH periods
332 after 4th Dec and the four events are shown in Fig. 2, respectively. To be noted, the “high (or low) RH
333 events” is used to refer to the SA formation events under high (or low) RH conditions for convenience,
334 and it doesn't mean that RH caused variations of CCN behavior. As can be seen in Figs. 2 (1b) and
335 (2b), different variations of SPAR due to SA formations can be found during the periods with different
336 RH conditions. The average diurnal variations of these parameters for the entire high RH stage and
337 low RH stage as shown in Figs. 2 (1a-1d) and (2a-2d) revealed similar but more smoothed variations
338 as in the four selected events. The four events are discussed and intercompared in the following to
339 magnify the differences under distinct RH conditions. For accumulation mode particles, particle
340 number concentrations were higher during daytime in high RH events, while stronger diurnal
341 variations occurred in low RH events. Simultaneous daytime increases in particle SPAR in size range
342 from 200 nm to 400 nm, GF-PDF in GF range from 1.2 to 1.8 hygroscopicity and SA mass fraction
343 were found in all four events, suggesting that SA formation led to increasing hygroscopic particles
344 number concentration, which in turn enhanced particle CCN activity. This effect was more pronounced
345 in Events 1 and 2 than in Events 3 and 4. In Events 1 and 2, SPAR values were generally higher than
346 0.4 at 200 nm and reached the maximum of 1 during noontime at 300 nm. A hygroscopic mode with
347 $GF > 1.4$ was ~~be~~ found throughout the day, which dominated aerosol hygroscopicity during daytime.
348 Mass fraction of SA were generally higher than 70% and reach a maximum of 80% at noon. While in
349 Events 3 and 4, SPAR at 200 nm was lower than 0.4 at night and the maximum SPAR at 300 nm was
350 lower than 0.9. A particle mode with $GF < 1.1$ dominates particle hygroscopicity, and the mass fraction
351 of SA was lower than 60% and 30% at noon and at night, respectively. However, stronger daytime

352 increase of SA mass fraction and accordingly larger variation in SPAR was observed in Events 3 and
353 4 than in Events 1 and 2.

354 Besides SS of 0.05%, variations of SPAR at SSs of 0.07% and 0.2% are also shown in Figs.
355 S1 and S2 in the supplement. And as shown in Figs. S1 and S2, the variations of SPAR and N_{CCN}/PM
356 at SS of 0.07% are similar but lighter, compared with those at SS of 0.05%. While for SS of 0.2%, the
357 difference of SPAR between different periods or events are smaller (Fig. S1), and so did the diurnal
358 variations of SPAR and GF-PDF at particle size of 100 nm (Fig. S2). Because CCN activity at SS of
359 0.2% was strong enough (indicated by SPAR value close to 1) in particle size range where the SA
360 formation dominates, and thus the different SA formations under high or low RH conditions cannot
361 lead to significant variations of CCN activity at SS of 0.2%. In summary, based on CCN measurements
362 in this study, the RH-dependent influence of SA formation on CCN activity can be found obviously at
363 SSs of 0.05% and 0.07%. As the variations of SPAR at SS of 0.07% were quite similar to those at SS
364 of 0.05%, further analysis was only based on CCN activity at SS of 0.05%.

365 In Fig. 3a, detailed comparison of particle CCN activity during SA formation events of N_{CCN}
366 enhancements at SS of 0.05% under different RH conditions are shown as the variations of SPAR
367 curves. Particle CCN activity in Events 1 and 2 were combined due to their similar diurnal variations
368 (as shown in Fig. 2). Besides SPAR curves (Fig. 3a), corresponding fitting parameters of the SPAR
369 curve including D_a and MAF were also shown in Figs. 3b and c, respectively, as enhanced SPAR for
370 particle population can be attributed to hygroscopic particle number fraction increase (MAF increase)
371 or enhancement of hygroscopic particle hygroscopicity (D_a decrease). Same as demonstrated in Fig. 2,
372 SPAR was generally higher and thus particle CCN activity(0.05%) were generally stronger in high RH
373 events than those in low RH events. However, as shown in Fig. 3a, the difference between SPAR in
374 high and low RH events at 300 nm decreased from 0.2 to 0.1 during the SA formations, indicating for
375 a stronger enhancement in low RH events, probably due to both the stronger increase of SA mass
376 fraction and the higher nighttime PA mass fraction (Fig. 2(e)). Furthermore, in high RH events, there
377 were daytime enhancements of SPAR within the 150 to 300 nm size range, as was indicated by the
378 daytime increase of MAF and decrease of D_a , which mainly resulted from number fraction and
379 hygroscopicity increases of CCN-active particles. While in low RH events, the daytime enhancement
380 of SPAR was only observed for particles larger than 200 nm. This can be attributed to the strong
381 increase of MAF and the slight decrease of D_a , which indicates significant increasing number fraction
382 yet slightly enhanced hygroscopicity of hygroscopic particles, respectively. Overall, the enhancement
383 of SPAR was weaker but occurred at a broader particle size range in high RH events than in low RH
384 events, as shown in Fig. 3a. This is in accordance with previous the results from Kuang et al. (2020c),

385 suggesting that SA formation occurred mainly in aqueous phase within a broad particle size range (up
386 to 1 μm) in high RH events, while SA formation dominantly proceeded via gas phase reactions and
387 contributed to aerosol sizes smaller than 300 nm in low RH events. At SS of 0.05% (Fig. 3(a)), the
388 variation of SPAR from 8:00-12:00 to 12:00-16:00 in particle size smaller than 200 nm was very small
389 during low RH events, suggesting a smaller CCN activity enhancement due to SA formation compared
390 with those at high RH events. In detail, the different variations of SPAR in high and low RH events
391 indicated by MAF and D_a shown in Figs. 3(b & c) suggested different variations of hygroscopicity,
392 number fraction and size of SA particles. Before SA formation, there was a significant difference
393 between the MAF in high and low RH events, which disappeared after the SA formation. The stronger
394 variations in MAF in low RH events suggested stronger enhancement of number concentration of
395 formed SA particles. As for D_a during SA formation, there were similar, little decrease in both high
396 and low RH events, suggesting similar hygroscopicity of the SA formed under low and high RH
397 conditions. Thus differences of SPAR and the resultant N_{CCN} during low and high RH events were
398 mainly due to the different variations of number fraction of formed SA particles.

399 As there were different influences of SA formation on both CCN activity at SS of 0.05% and
400 PNSD under different RH conditions, different variation of $N_{CCN}(0.05\%)$ due to SA formation can also
401 be expected. Fig. 4 displays the diurnal variation of $\text{PM}_{2.5}$ mass concentration, volume concentration
402 (V_{conc}), number concentration (N_{conc}) and $N_{CCN}(0.05\%)$ (all divided by CO to partially compensate
403 for changes in planetary boundary layer height), as well as the $N_{CCN}/\text{PM}_{2.5}$ mass concentration ratio
404 and SPAR during high and low RH events, respectively. Variables in Fig. 4 were also presented in Fig.
405 S3 averaged for the entire high RH and low RH stages, respectively. Compared with the selected case
406 events featuring significant N_{CCN} enhancement (Fig. 4(1c-2c)), the diurnal variations averaged for the
407 entire high and low RH stages were similar, with higher levels of particle mass concentration but
408 weaker enhancement of SA and N_{CCN} , indicating similar but weakened impact of SA formation on
409 CCN activity due to the interference of other aerosol processes. Hereinafter, we discuss the variations
410 in the four events to magnify the discrepancies of SA formation under high RH and low RH conditions
411 and its distinct impact on N_{CCN} . The V_{conc} size distribution variations can be used as a proxy for the
412 evolution of NR- $\text{PM}_{2.5}$ size distributions, considering the relatively small variations in particle density
413 (ranging from 1.2 to 1.8 and with relative variations within 20% (Hu et al., 2012; Zhao et al., 2019)).
414 The variations of the ratio between N_{CCN} (in different particle size range) and the mass concentration
415 of $\text{PM}_{2.5}$ SA (referred as to N_{CCN}/SA) or NR- $\text{PM}_{2.5}$ (referred as to N_{CCN}/NR) can be used to evaluate
416 the response of N_{CCN} to SA formation.

417 During high RH events, normalized $N_{CCN}(0.05\%)$ increased by $\sim 50\%$ from 8:00 to 14:00, with a
418 similar increase in normalized $PM_{2.5}$ SA mass concentration (Fig. 4(1a)). As the $PM_{2.5}$ PA mass
419 concentration decrease was much smaller than the SA increase, the NR- $PM_{2.5}$ mass concentration
420 increase can be expected to be similar to the SA increase. As reported by Kuang et al. (2020c), SA
421 during daytime were mainly formed at larger particle sizes, featuring V_{conc} increase in the particle
422 size range of 400 to 1000 nm. In Fig. 4(1d), significant increases of particle number concentration
423 (N_{conc}) in particle size range of 150 nm to 1000 nm can be observed. At larger particle size the
424 increase of N_{conc} led to stronger increase of V_{conc} , which is why there was simultaneous but much
425 weaker increases of V_{conc} in particle size range of 150 to 300 nm compared with increases of those
426 in particle size of larger than 300 nm (Fig. 4(1b)). This suggests that $PM_{2.5}$ SA mainly contributed to
427 particle sizes of larger than 300 nm. In addition, because the SA formation enhanced hygroscopicity
428 and number fraction of CCN-active particles in particle size range of 150 to 300 nm, simultaneous
429 enhancements of SPAR can be found throughout the measured particle size range of 180 to 300 nm
430 (Fig. 4(1e)). By combining the enhancements of N_{conc} and SPAR in measured particle size ranges,
431 there were increases of N_{CCN} from 200 to 500 nm (Fig. 4(1c)). ~~While the strongest V_{conc} increases~~
432 ~~occurred at particle sizes larger than 300 nm, the N_{CCN} increased most in the particle size range of 200~~
433 ~~to 300 nm, suggesting that particles larger than 300 nm were already CCN-active before SA formation.~~
434 Thus while SA formation processes contributed to their volume (mass) and hygroscopicity increase, it
435 had no further impact on N_{CCN} . As a result, $N_{CCN} (>300 \text{ nm})/SA$, $N_{CCN} (<300 \text{ nm})/SA$, N_{CCN}
436 $(>300 \text{ nm})/NR$ and $N_{CCN} (<300 \text{ nm})/NR$ all decreased during the SA formation (Fig. 4(1f)), suggesting
437 that weakening enhancement of $N_{CCN}(0.05\%)$ in SA formation under high RH condition as SA
438 formation mainly added mass to already CCN-active particles .

439 During low RH events, weaker increases of both $N_{CCN}(0.05\%)$ and $PM_{2.5}$ SA mass
440 concentration from 8:00 to 14:00 was found (Fig. 4(2a)). At the same time, PA mass decreased by 50%
441 and the variation of total NR mass was small. Under low RH conditions, SA formation mainly
442 contributed to mass enhancements of smaller particle sizes (Kuang et al., 2020c). V_{conc} increased
443 mostly in the range of 150 to 300 nm (Fig. 4(2b)), while N_{conc} only increased within 300 nm (Fig.
444 4(2d)), suggesting that $PM_{2.5}$ SA mainly formed in particle size range ~~within~~-below 300 nm. SA
445 formation mainly enhanced number fraction of CCN-active particles in particle size of 200 to 300 nm,
446 as SPAR only revealed evident enhancement (Fig. 4(2e)) and N_{CCN} only significantly increased (Fig.
447 4(2c)) in that size range. As a result, although $N_{CCN} (>300 \text{ nm})/SA$ decreased similar as that under high
448 RH conditions, $N_{CCN} (<300 \text{ nm})/SA$ and $N_{CCN} (>300 \text{ nm})/NR$ generally stayed constant and N_{CCN}
449 $(<300 \text{ nm})/NR$ even increased during SA formation in daytime (Fig. 4(2f)). The ratio between bulk

450 N_{CCN} and mass concentration of NR-PM_{2.5} became larger due to the SA formation, suggesting that
451 stronger enhancement of N_{CCN} (0.05%) in SA formation under low RH condition, because SA
452 formation mainly added mass to CCN-inactive particles and turned them into CCN-active particles.

453 In summary, during the campaign in this study, two kinds of SA formation events were
454 observed under different RH conditions with different variations of PM and N_{CCN} at SSs lower than
455 0.07%. Under high RH conditions, there were strong SIA dominated SA formation leading to stronger
456 enhancements of CCN-active particle number fraction higher particle hygroscopicity and N_{CCN}
457 enhancements. Meanwhile, under low RH conditions, there were moderate SOA dominated SA
458 formation with moderate enhancements of CCN-active particle number fraction particle
459 hygroscopicity and N_{CCN} -enhancements. However, for a unit amount of SA formation, the increase of
460 N_{CCN} was stronger under low RH conditions and weaker under high RH conditions. This was because
461 SA formation under low RH conditions was more concentrated on particle sizes smaller than 300 nm
462 and added more mass to CCN-inactive particles, turning them into CCN-active particles. In addition,
463 strong and distinct diurnal variations of CCN activity of particles were observed during different SA
464 formation processes, whose effects on N_{CCN} calculation need to be further evaluated.

465

466 3.3. The influence of diurnal variation of CCN activity on N_{CCN} prediction

467 Since PNSD measurements are generally more simpler and more common than N_{CCN}
468 measurements, N_{CCN} is usually estimated from real-time PNSD combined with parameterized CCN
469 activity. In former sections, it was already manifested that SA formations under different RH
470 conditions led to distinct variations in PNSD and SPAR at SS of 0.05%, thus hence different variations
471 in N_{CCN} . Thus, it is important for the prediction of N_{CCN} to quantify its sensitivity towards changes in
472 PNSD and SPAR during SA formation processes under different RH conditions.

473 In this study, PNSD dominated the N_{CCN} calculation was mostly determined by PNSD, as was
474 generally found the case in former studies (Dusek et al., 2006), uring SA formation events, but
475 however, the variation of CCN activity also contributed significantly to the deviation of N_{CCN}
476 calculation during SA formation. In former discussions, CCN activity (indicated by SPAR) at
477 0.05% SS revealed significant diurnal variations during this campaign, which were different during
478 SA formations under distinct RH conditions. The ratio between of N_{CCN} calculated based on campaign
479 averaged SPAR (N_{CCN_cal}) to those measured N_{CCN} (N_{CCN_meas}) at 0.05% SS (N_{CCN_meas}) before and
480 after 4th Dec are shown in Fig. 5. SPAR is determined by the variation of D_a and MAF, which reflect
481 changes in hygroscopicity and number fraction of hygroscopic particles. Thus, to investigate the

482 ~~separate-respective~~ influences of MAF and D_a variations on N_{CCN} predictions, N_{CCN_AvgMAF} (or
483 N_{CCN_avgDa}) was calculated based on the real-time PNSD and SPAR estimated by replacing MAF (or
484 D_a) in Eq. 7 with the campaign averaged value. During the high RH stage, underestimation of daytime
485 N_{CCN_cal} can reach up to 20%, since SPAR variations due to CCN activity enhancement were not
486 considered. Similar deviations of both N_{CCN_AvgMAF} and N_{CCN_avgDa} from N_{CCN_meas} were detected,
487 suggesting that both MAF and D_a variations contributed to N_{CCN_cal} underestimation under high RH
488 conditions. During the low RH stage, up to 50% overestimation existed in $N_{CCN_AvgSPAR}$ ~~beyond-outside~~
489 SA formation ~~episodestime periods~~. Only N_{CCN_AvgMAF} displayed similar deviations from N_{CCN_meas} ~~as~~
490 $N_{CCN_AvgSPAR}$, indicating that differences between N_{CCN_cal} and N_{CCN_meas} were mainly contributed by
491 variations in MAF brought on by significant CCN-active particles number fraction growth due to SA
492 formations. To be noted, $N_{CCN_AvgSPAR}$ before and after 4th Dec ~~are-were~~ both calculated on the basis
493 based of on the averaged-SPAR averaged of over this the entire campaign (green dots in Fig. 5a) ~~rather~~
494 ~~than the averaged SPAR before or after 4th, because-since~~ the applicability of ~~the~~ campaign averaged
495 SPAR ~~on-in the~~ N_{CCN} calculations ~~in the NCP~~ was confirmed ~~in-by~~ many former studies in the NCP
496 (Deng et al., 2012; Wang et al., 2013; Ma et al., 2016). During ~~the~~ low RH periods, SPAR ~~are-was~~
497 generally lower than the campaign averaged SPAR ~~of the campaign~~ and the ratio between the
498 calculated and measured N_{CCN} ~~are-were~~ systematically higher (lasting for the whole night). In summary,
499 SA formation processes can induce significant deviation to N_{CCN} prediction that varied with RH
500 conditions, which and mainly resulted from the variation in of MAF ~~and also varied under different~~
501 ~~RH conditions~~. Thus, for accurate N_{CCN} estimations, considering the variation of MAF (changes in the
502 fraction of the hygroscopic particles aerosol hygroscopicity) is highly essential.

503 As SOA is generally considered to be more hygroscopic than POA (Frosch et al., 2011; Lambe
504 et al., 2011; Kuang et al., 2020a), the increase of hygroscopic particles or SA particles (both SIA and
505 SOA) were considered to be the cause for the increase of SPAR within 200 to 300 nm size range (Fig.
506 2). In order to account for the variations of hygroscopic particles or SA particles in N_{CCN} calculation,
507 in the following part, Number Fraction of hygroscopic particles ($GF(90\%, 200\text{ nm}) > 1.22$, NF_{hygro})
508 measured by HTDMA and Mass Fraction of SA particles (MF_{SA}) measured by ACSM in this campaign
509 were used to represent MAF variations and to provide calculation of N_{CCN} at SS of 0.05% with smaller
510 deviations combining-combined with PNSD measurement ~~with smaller deviations~~. To be noted, in
511 order to highlight the application of using MF_{SA} as estimation of MAF variations on N_{CCN} calculation,
512 the campaign averaged D_a from SPAR curves was used.

513 Based on the bulk hygroscopicity derived from particle chemical compositions measurements
514 (κ_{chem}), a critical diameter for CCN activation can be calculated based on κ -Köhler theory. ~~and based~~

515 ~~on~~ ~~With this critical diameter, which~~ $N_{CCN}(0.05\%)$ can be predicted ~~by incorporating~~ ~~combining~~
516 measured PNSD ~~with this critical diameter~~ (N_{CCN_Chem}). ~~Although there can be significant deviations~~
517 ~~for~~ κ of accumulation mode particles derived from chemical composition of the bulk aerosol ~~might~~
518 ~~bear significant uncertainties~~, which leads to significant deviations of N_{CCN} prediction. However, in
519 practice, ~~the measurements of~~ chemical compositions ~~measurements specifically for~~ accumulation
520 mode particles are not common, ~~and thus bulk aerosol~~ chemical compositions ~~of the bulk aerosol is~~ ~~are~~
521 ~~still~~ commonly applied in CCN studies ~~as substitute~~ (Zhang et al., 2014; Zhang et al., 2016; Che et al.,
522 2017; Cai et al., 2018), especially when particle hygroscopicity measurements were in lack. ~~As can be~~
523 ~~seen in~~ Fig. 6(a), ~~it can be found that~~ N_{CCN_meas} at 0.05% SS ~~wase~~ ~~can be~~ strongly underestimated
524 by N_{CCN_Chem} , especially at lower N_{CCN_meas} ($\sim 10^2 \# \text{cm}^{-3}$), which is similar to ~~the~~ results of studies
525 that encountered high fractions of organics (Chang et al., 2010; Kawana et al., 2015). This deviation
526 between N_{CCN_meas} and N_{CCN_Chem} may have resulted from the hypothesis of internal mixing state and
527 the difference of particle hygroscopicity derived by particle chemical composition measurements and
528 CCN activity. Fig. 6(b) depicts the correlation between mass fraction of SA (MF_{SA}) and MAF at 0.05%
529 SS. MF_{SA} was generally positively correlated to MAF ($r=0.8$) with slight underestimations, suggesting
530 that externally mixed SA dominated CCN-active particles. Thus, in the prediction of N_{CCN} , real-time
531 SPAR can be calculated from ~~campaign~~ average D_a and MAF assumed to ~~be~~ equal to real-time MF_{SA}
532 (N_{CCN_MF}). As displayed in Fig. 6(c), the underestimation and correlation between N_{CCN_cal} and
533 N_{CCN_meas} was improved after introducing MF_{SA} into N_{CCN} calculation. Additionally, the diurnal
534 variations of N_{CCN_cal}/N_{CCN_meas} ratio based on different methods of N_{CCN} calculation during the whole
535 campaign were shown in Fig. 6(d). ~~It can be found that~~ ~~By~~ considering ~~the~~ real-time MF_{SA} ~~variations~~,
536 the deviation of calculated N_{CCN} (Real-time MF) can be reduced throughout the day, compared ~~with~~
537 ~~to~~ N_{CCN_Chem} (Real-time Chem). Meanwhile, ~~if using~~ an averaged MF_{SA} ~~is used~~ to estimate SPAR and
538 N_{CCN} ~~could also reduce~~, ~~the~~ deviations of calculated N_{CCN} (Averaged MF) ~~can be reduced as well~~, ~~but~~
539 ~~however~~, demonstrated a much stronger diurnal variations than the deviation of N_{CCN_MF} .

540 Based on the bulk hygroscopicity derived from GF measurements (κ_{GF}) at 200 nm, ~~a critical~~
541 ~~diameter~~ D_a can be calculated based on ~~the~~ κ -Köhler theory, ~~which can be applied to predict~~ ~~and~~ N_{CCN}
542 at 0.05% SS (~~N_{CCN_GF}~~) ~~can be predicted by~~ ~~in~~ ~~combining~~ ~~combination~~ ~~with~~ measured PNSD ~~and the~~
543 ~~critical diameter~~ (~~N_{CCN_GF}~~). Fig. 7(a) ~~shows~~ ~~reveals~~ that N_{CCN_meas} were strongly underestimated by
544 N_{CCN_GF} (by more than 30%). ~~which~~ ~~This deviation~~ ~~may~~ ~~might~~ ~~also~~ have resulted ~~ed~~ from the
545 hypothesis of internal mixing state and the difference of particle hygroscopicity derived by GF and
546 particle CCN activity measured under different water vapor saturated conditions. Fig. 7(b) depicts the
547 positive correlation between NF_{hygro} and MAF at 0.05% SS, which was weaker than that between MF_{SA}

548 and MAF. Similar as before, $N_{F_{hygro}}$ was applied as a proxy for MAF in the N_{CCN} calculation, which
549 also improved the underestimation and correlation between N_{CCN_cal} and N_{CCN_meas} (Fig. 7(c)). Also,
550 the campaign averaged D_a in Fig. 5a. was used to calculate SPAR curves and N_{CCN} . The diurnal
551 variations of the N_{CCN_cal}/N_{CCN_meas} ratio based on different methods of N_{CCN} calculation during the
552 whole campaign ~~were~~ are shown in Fig. 7(d). ~~It can be found that b~~By considering the real-time
553 variation of $N_{F_{hygro}}$, the deviation of N_{CCN_NF} (Real-time NF) ~~can~~ was ~~be reduced~~ mainly reduced
554 during nighttime, compared ~~with~~ to N_{CCN_GF} (Real-time GF). Meanwhile, if applying an averaged
555 $N_{F_{hygro}}$ is used to estimate SPAR and N_{CCN} , reduced the deviations of calculated N_{CCN} (Averaged NF)
556 ~~can be reduced~~ during nighttime as well, but its deviations demonstrated ~~a~~ stronger diurnal variations
557 than ~~the deviation of those of~~ N_{CCN_NF} . If GF-PDF were directly used to calculate N_{CCN} , N_{CCN_cal} would
558 agree well with measured N_{CCN} (Fig. S4), because in this way the mixing state of aerosol would have
559 been accounted for. However, compared to the approach using GF-PDF, $N_{F_{hygro}}$ is easier to apply in
560 N_{CCN} calculation and can yield similar accuracies.

561 In summary, MAF exhibited strong diurnal variation that varied under different RH conditions
562 due to different SA formation mechanisms, which contributed most to N_{CCN} estimation deviations if
563 unaccounted for. The diurnal variations of MAF at the five measured SSs (Fig. S5) revealed significant
564 diurnal variations ~~in MAF~~ at low SSs (0.05% and 0.07%) that were dependent on RH conditions,
565 while only small diurnal variations that were insensitive to the RH conditions were detected at SSs
566 over above 0.2%. In general, MAF ~~become~~ became lower at lower SSs, especially during nighttime.
567 As the fraction of CCN-active particles were generally hygroscopic and composed of secondary
568 compounds, positive correlation was found between MAF, MF_{SA} and $N_{F_{hygro}}$. Although a good
569 prediction of N_{CCN} (0.05%) was achieved by applying an averaged MAF (Figs. 5, 6d and 7d), in practice,
570 this would still require CCN measurements or HTDMA/chemical composition- measurements as
571 proxies. Additionally, deviations of N_{CCN_cal} based on the averaged MAF can be large under low RH
572 conditions (Fig. 5c), while time-dependent MAF can eliminate a great part of these deviations. Thus,
573 by replacing MAF with real-time MF_{SA} or $N_{F_{hygro}}$ when deriving SPAR curves, the relative deviation
574 of N_{CCN} (0.05%) calculation can be reduced. The proposed N_{CCN} parameterization using MF_{SA} can also
575 be easily adopted by chemical-transport and climate models, improving their representation of N_{CCN}
576 changes due to distinct SA formation processes.

577

578 4. Conclusions

579 SA formation drives the development of haze pollution in the NCP and can result in significant
580 variations of PNSD and aerosol hygroscopicity. Studies in the NCP have shown that the mechanism
581 of SA formation can be affected by relative humidity (RH), and thus has different influences on the
582 aerosol hygroscopicity and PNSD under distinct RH conditions. The difference in particle size where
583 SA formation is taking place and the different chemical compositions of formed SA can result in
584 different variations of CCN activity. Thus, it is essential to study the influence of SA formation on
585 CCN activity of existing accumulation mode particles under different RH conditions in the NCP. As
586 N_{CCN} is often predicted based on real-time PNSD and parameterized SPAR, the influence of varying
587 SPAR in distinct SA formation processes on N_{CCN} calculation needs to be evaluated in detail.

588 Based on the measurements of CCN-activity, particle hygroscopicity, particle chemical
589 composition, PNSD during the McFAN campaign in Gucheng winter 2018, the influences of SA
590 formation on CCN activity and N_{CCN} calculation under different RH conditions were investigated
591 especially at SSs lower than 0.07%. Two kinds of SA formation events were identified under different
592 RH conditions with distinct variations in PM and N_{CCN} at 0.05% SS. Under high RH conditions, which
593 corresponds to the periods with minimum RH higher than 50% in daytime, strong SA formation and
594 $N_{CCN}(0.05\%)$ enhancements with strong hygroscopic particles and SIA dominated contribution to SA
595 (>70%) was found. While under low RH conditions, which corresponds to the periods with daytime
596 minimum RH below 30%, moderate SA formation and $N_{CCN}(0.05\%)$ enhancements with moderately
597 hygroscopic particles and SOA dominated contribution to SA was found. However, the increase of
598 N_{CCN} under the a same amount of SA formation was stronger under low RH conditions and weaker
599 under high RH conditions. This was because the formation of SA under low RH conditions was more
600 concentrated in particle size range smaller than 300 nm and added more mass to CCN-inactive particles
601 turning them into CCN-active ones after SA formation.

602 In addition, strong diurnal variations of CCN activity of particles at 0.05% SS due to the strong
603 SA formations were also observed, both varying with RH conditions. $N_{CCN}(0.05\%)$ was significantly
604 underestimated when MAF (SPAR parameter) variations were not considered. As the fraction of CCN-
605 active particles were generally hygroscopic and composed of secondary compounds, there were good
606 correlation among MAF inferred from measurements of CCN activity, particle hygroscopicity and
607 particle chemical compositions. Thus, the relative deviation of $N_{CCN}(0.05\%)$ estimation can be
608 reduced by applying measurements of particle hygroscopicity or particle chemical compositions as a
609 proxy for aerosol mixing state.

610 This study can further the understanding of the impact of SA formation on CCN activity and
611 N_{CCN} calculation, specifically for SA formations on existing particles, which can strongly affect cloud
612 microphysics properties in stratus clouds and fogs. The investigation of the influence of SA formation
613 on CCN activity of existing particles in this study is important for improving N_{CCN} parameterizations
614 in chemical-transport and climate models, so that they can account for the large variations induced by
615 SA formation processes.

616

617 **Supporting Information**

618 The supporting information is available in a separate file.

619 **Data availability.**

620 The data used in this study are available from the corresponding author upon request
621 (taojch@jnu.edu.cn and nan.ma@jnu.edu.cn)

622 **Author contributions.**

623 JT, YK and NM designed this research. JT performed the data analysis and wrote the manuscript. YC,
624 HS, NM, YK, JT, and JH planned this campaign. JT and YZ conducted the CCN measurements. YS
625 and YH conducted the ACSM measurements and the ACSM PMF analysis. JH and QL conducted the
626 HTDMA measurements. LX and YZ conducted the particle number size distribution measurements.
627 WX conducted the measurements of CO and meteorological parameters. YC, HS, YS, YK and NM
628 contributed to the revisions of this manuscript and all other coauthors have contributed to this paper in
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637 **Conflicts of interest**

638 There are no conflicts to declare.

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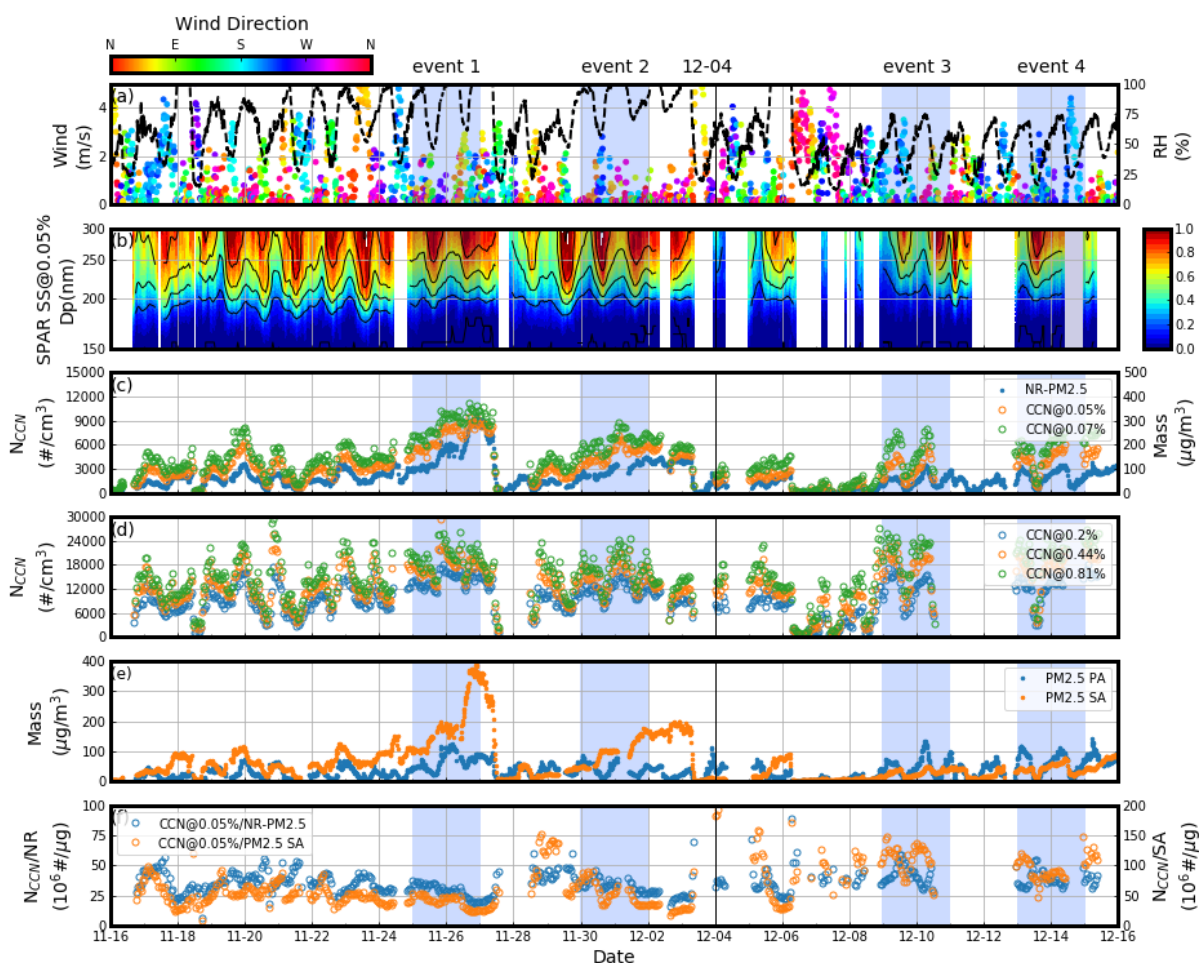
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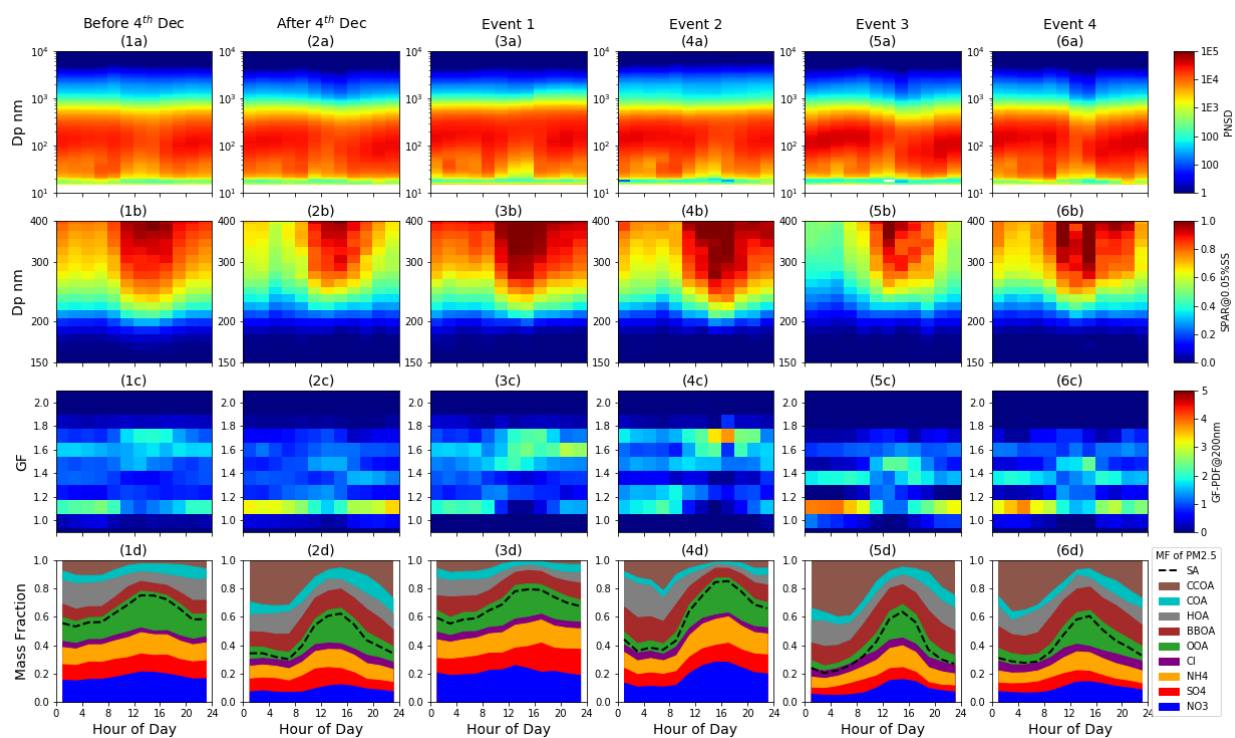
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924 **Figures:**



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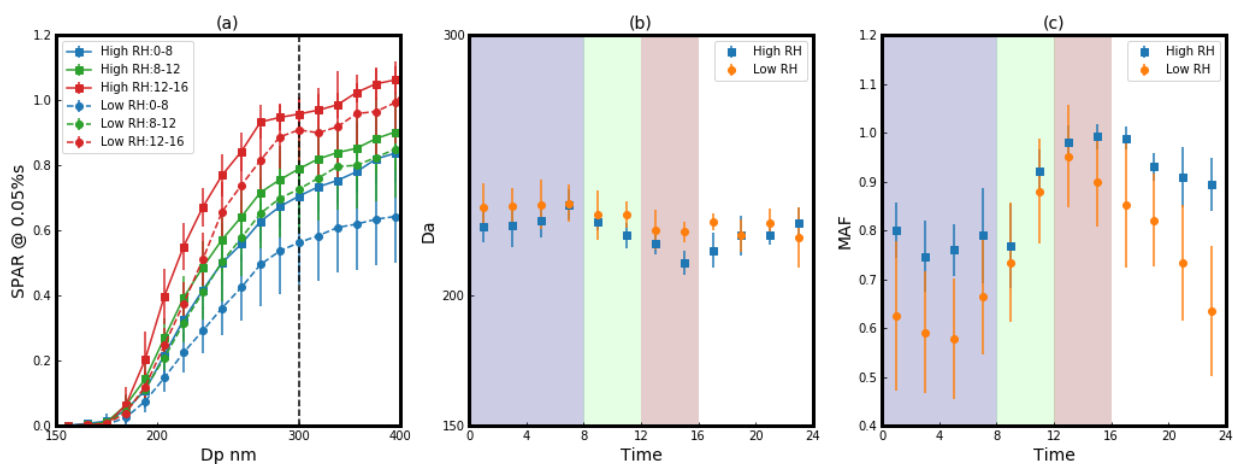
926 Fig 1. Overview of the measurements during the campaign: (a) dots represent wind speed with color
 927 indicating wind direction, and black lines represent RH; (b) SPAR under SS of 0.05%; (c) blue, green
 928 and yellow dots represent N_{CCN} under SS of 0.05% and 0.07%, and mass concentration of NR-PM_{2.5},
 929 respectively; (d) blue, green and yellow dots represent N_{CCN} under SS of 0.2%, 0.44% and 0.81%,
 930 respectively; (e) blue and yellow dots represent mass concentration of PM_{2.5} PA and PM_{2.5} SA
 931 respectively; (f) blue and yellow dots represent ratio between N_{CCN} and mass concentration of NR-
 932 PM_{2.5} and PM_{2.5} SA, respectively. There were four events with significant enhancements of N_{CCN}
 933 during the blue shaded periods.



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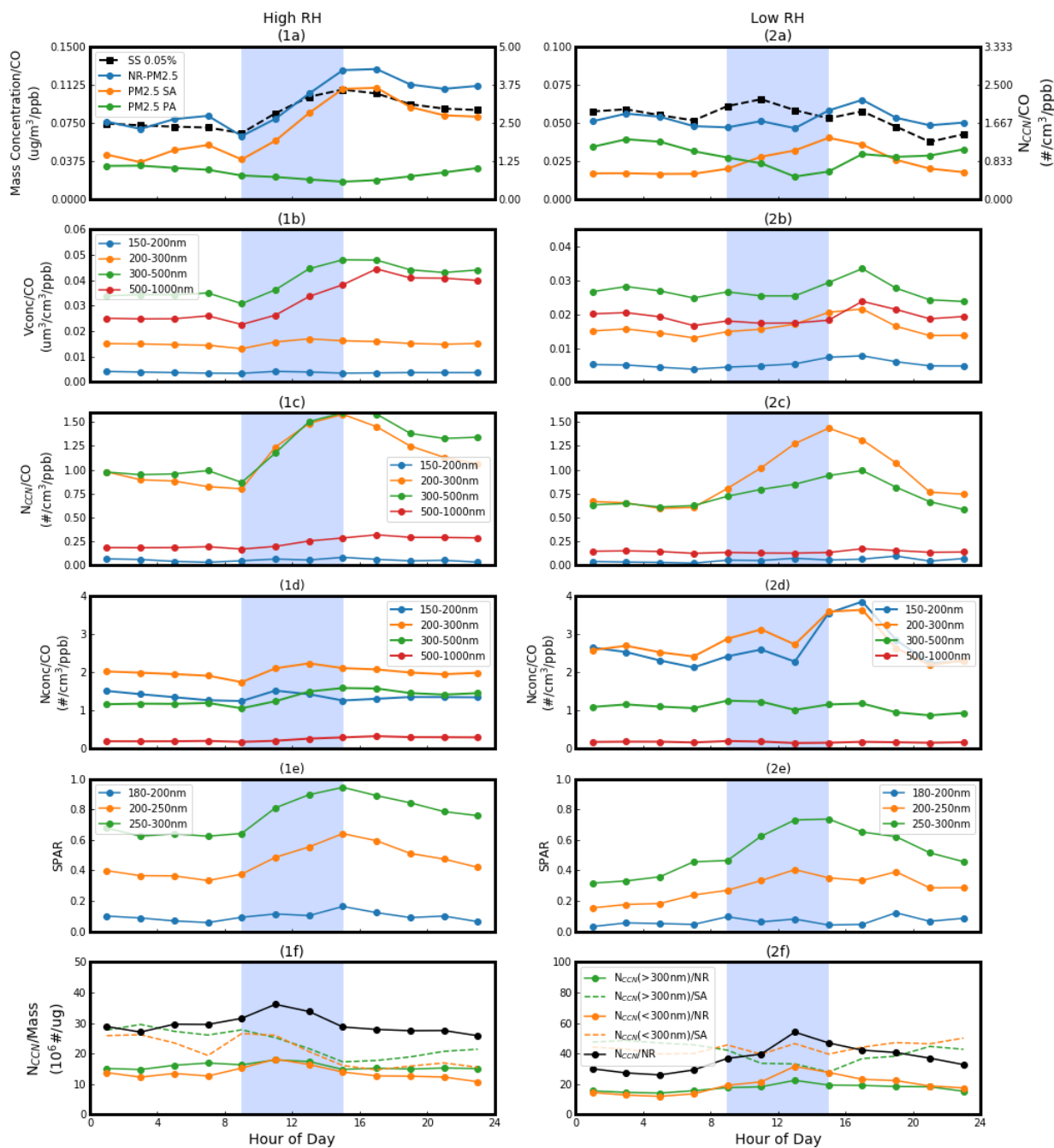
936 Fig 2. Diurnal variation of (a) PNSD, (b) SPAR at SS of 0.05%, (c) GF-PDF at 200 nm and (d) mass
 937 fraction of different PM_{2.5} chemical species during high RH periods before 4th Dec (1), low RH
 938 periods after 4th Dec (2) and the four events (3-6), including OA factors: hydrocarbon-like OA
 939 (HOA), cooking OA (COA), biomass burning OA (BBOA), coal combustion OA (CCOA), and
 940 oxygenated OA (OOA).

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943 Fig 3. (a) The averages of SPAR curves at SS of 0.05% in three different time periods (blue: 0:00-
944 8:00; green: 8:00-12:00; red: 12:00-16:00) during high (squares with solid line, event 1 and 2) and low
945 (dots with dashed line, event 3 and 4) RH events. Diurnal variation of (b) D_a and (c) MAF under high
946 (blue) and low (yellow) RH conditions. The blue, green and red shades correspond to with the three
947 periods in (a & d). Error bars indicate the standard deviations of data.



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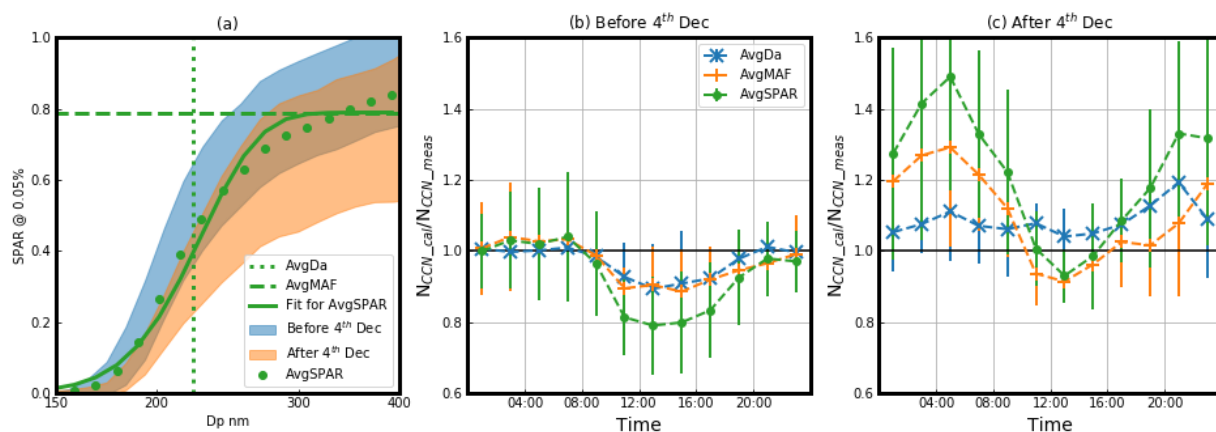
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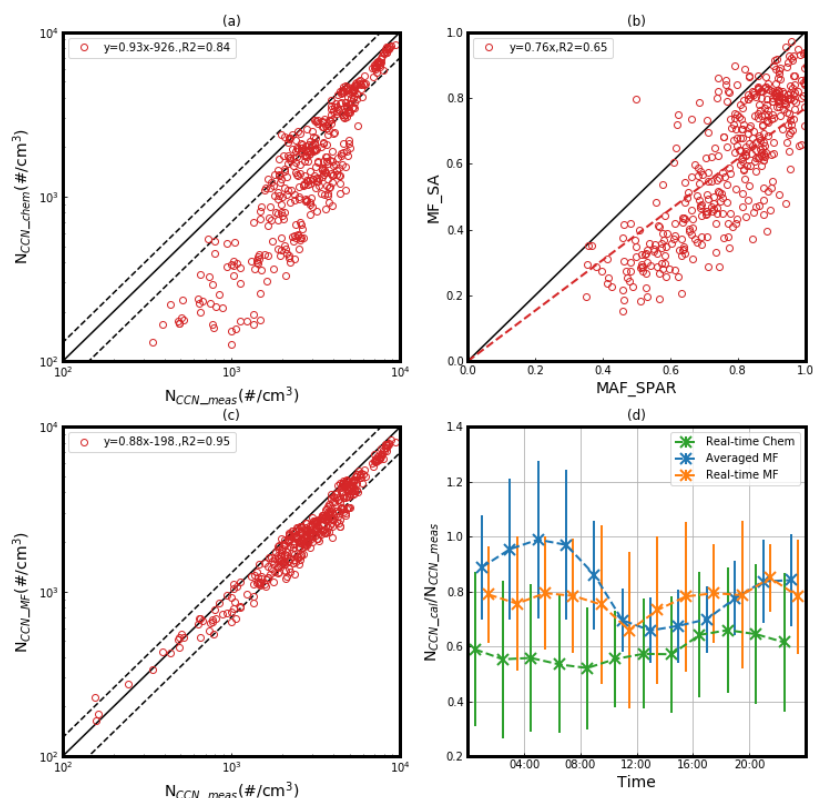
Fig 4. During different RH events, the average diurnal variation of (a) the ratios between particle mass concentration (dots with solid lines; blue: NR-PM_{2.5}; yellow: PM_{2.5} SA; green: PM_{2.5} PA) and CO concentration, and the ratio between N_{CCN} at SS of 0.05% and CO concentration (squares with solid line); (b) the ratios between particle volume concentration (Vconc) of different particle size range (indicated by colors) and CO concentration; (c) the ratios between N_{CCN} of different particle size range at SS of 0.05% (indicated by colors) and CO concentration; (d) the ratios between particle number concentration (Nconc) of different particle size range (indicated by colors) and CO concentration; (e) SPAR of different particle size range (indicated by colors); (f) the ratios between N_{CCN} at SS of 0.05% (black: bulk N_{CCN}; yellow: N_{CCN} with particle size larger than 300 nm; blue:

958 N_{CCN} with particle size smaller than 300 nm) and mass concentration of NR-PM_{2.5} SA and the ratios
959 between N_{CCN} and mass concentration of NR-PM_{2.5} (dashed lines).



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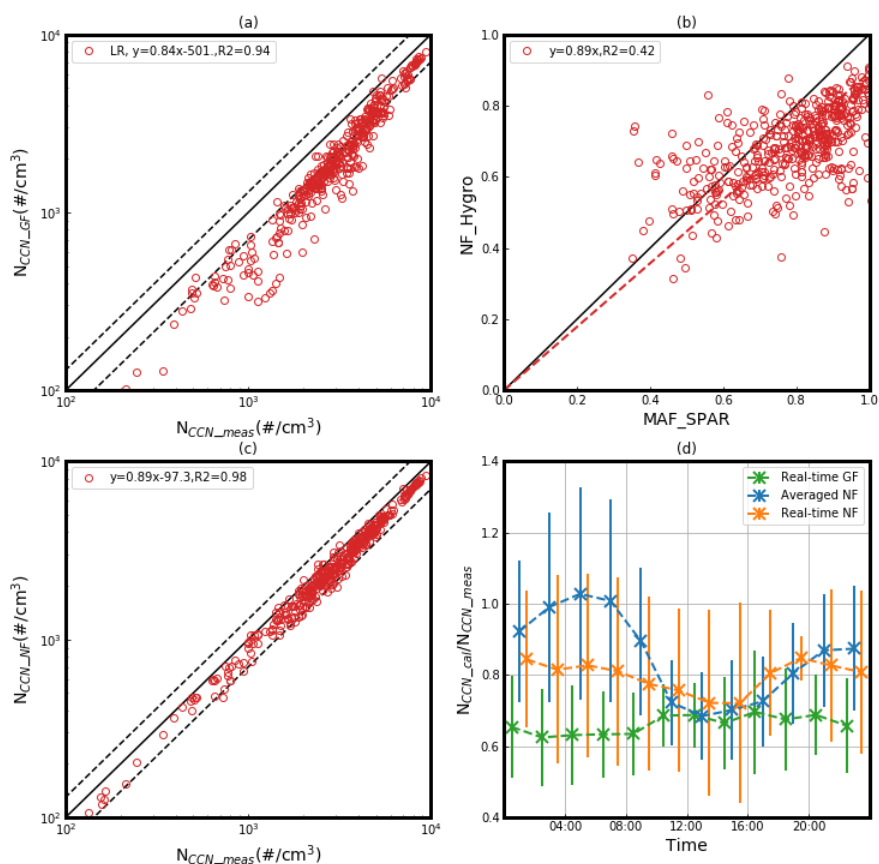
962 Fig 5. (a) The averaged SPAR at SS of 0.05% during the campaign (green scatters), the
 963 corresponding fitting curve (green line) and the averaged fitting parameters (dotted line for D_a and
 964 dashed line for MAF). The blue and yellow shaded areas represent the variations of SPAR before 4th
 965 Dec and after 4th Dec, respectively. The ratio between calculated N_{CCN} and measured N_{CCN} under (b)
 966 before and (c) after 4th Dec. Bars represent one standard deviation and colors represent different
 967 calculation of SPAR curves: green represent average SPAR during the campaign (AvgSPAR),
 968 yellow represent SPAR calculated with average D_a and real-time MAF (AvgDa) and blue represent
 969 SPAR calculated with average MAF and real-time D_a (AvgMAF).



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971 Fig 6. (a) The comparison between calculated N_{CCN} based on κ derived from bulk particle chemical
 972 compositions (N_{CCN_chem}) and measured N_{CCN} at SS of 0.05%. (b) The correlation between MAF and
 973 mass fraction of secondary aerosol (MF_{SA}). (c) the comparison between calculated N_{CCN} based on
 974 SPAR derived from real-time MF_{SA} and average D_a (N_{CCN_MF}) and measured N_{CCN} . The black dashed
 975 lines represent the relative deviation of 30%. (d) the diurnal variations of the ratio between the
 976 calculated and measured N_{CCN} during the whole campaign based on different methods (green:
 977 N_{CCN_chem} ; blue: N_{CCN} calculated based on SPAR derived from averaged MF_{SA} and average D_a ; yellow:
 978 N_{CCN_MF}).

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981 Fig 7. (a) The comparison between calculated N_{CCN} based on κ derived from bulk GF at 200 nm
 982 (N_{CCN_GF}) and measured N_{CCN} at SS of 0.05%. (b) The correlation between MAF and number fraction
 983 of hygroscopic particles (NF_{hygro} , $GF > 1.2$). (c) The comparison between calculated N_{CCN} based on
 984 SPAR derived from real-time NF_{hygro} and average D_a (N_{CCN_NF}) and measured N_{CCN} . The black dashed
 985 lines represent the relative deviation of 30%. (d) the diurnal variations of the ratio between the
 986 calculated and measured N_{CCN} during the whole campaign based on different methods (green: N_{CCN_GF} ;
 987 blue: N_{CCN} based on SPAR derived from averaged NF_{hygro} and average D_a ; yellow: N_{CCN_NF}).