

1 **Measurement Report: Cloud condensation nuclei (CCN)**
2 **activity in the South China Sea from shipborne**
3 **observations during summer and winter of 2021: seasonal**
4 **variation and anthropogenic influence.**

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22

23 **Abstract**

24 Understanding seasonal variations in cloud condensation nuclei (CCN) activity and the impact of
25 anthropogenic emissions in marine environments is crucial for assessing climate change. This study
26 presents findings from two shipborne observations conducted in the South China Sea (SCS) during the
27 summer and winter of 2021. In summer, higher particle number concentrations but lower mass
28 concentrations of non-refractory submicron particles (NR-PM₁) were observed, driven by Aitken mode
29 particle dominance. In contrast, winter showed a more balanced distribution between Aitken and
30 Accumulation mode particles. Summer particles were more hygroscopic, exhibiting higher activation
31 ratios (ARs) at all supersaturation (SS) levels. Distinct air mass periods were identified: in summer,
32 terrestrial air masses from Luzon ("Luzon" period), the Indochinese Peninsula ("Indochinese Peninsula"
33 period), and marine air masses; in winter, periods were influenced by Mainland China ("Mainland China"
34 period), a mix of Mainland China and marine air masses ("Mixed" period), and purely marine air masses.
35 The "Luzon" period in summer exhibited the highest particle number concentration, especially in the
36 Aitken mode, resulting in the highest CCN number concentration (N_{CCN}). Aerosol hygroscopicity was
37 higher during the "Indochinese Peninsula" period compared to the "Luzon" period, leading to a higher
38 bulk AR due to the combination of higher hygroscopicity and a greater fraction of accumulation mode
39 particles. The "Mainland China" period in winter showed a high nitrate fraction in the NR-PM₁, but the
40 inorganic fraction was similar to it in "Luzon" period, resulting in comparable hygroscopicity at low SS
41 to the "Luzon" period. However, smaller particle hygroscopicity was significantly lower in the
42 "Mainland China" period compared to summer. The "Mixed" period in winter exhibited a higher fraction
43 of accumulation mode particles, causing a higher bulk AR compared to the "Mainland China" period.
44 Overall, summer terrestrial air masses increased the Aitken mode particle and CCN concentration, while
45 winter terrestrial air masses led to higher concentration of large particles and lower hygroscopicity of
46 fine particles. CCN closure analysis, considering aerosol composition and mixing state, revealed that
47 summer aerosols were primarily internally mixed, whereas smaller aerosols in winter were primarily
48 externally mixed. The potential effect of undetected sea salt may lead to an underestimation of aerosol
49 hygroscopicity in summer. This study highlights significant seasonal differences in aerosol properties
50 and the impact of different types of terrestrial air masses on CCN activity in the SCS, contributing to our
51 understanding of regional climate influences.

52 **1.Introduction**

53 Aerosols can act as cloud condensation nuclei (CCN), influencing cloud formation, lifespan, and
54 albedo, thus indirectly impacting global radiative balance (Fletcher et al., 2011; Albrecht, 1989). The
55 aerosol-cloud interaction currently represents the largest uncertainty in radiative forcing within climate
56 models, ranging from -1.7 to -0.3 W m⁻² (IPCC, 2021). This uncertainty can be partially attributed to the
57 significant spatiotemporal variability in the aerosol size distribution and the ability of atmospheric
58 aerosol particles acting as CCN (CCN activity) (Fitzgerald, 1973; Jimenez et al., 2009; Sihto et al., 2011).
59 Thus, field measurements of aerosol size distribution and physicochemical properties are needed to better
60 understand the radiative forcing exerted by atmospheric aerosol particles.

61 Previous studies suggest that particle number size distribution (PNSD) is a primary factor
62 influencing CCN concentrations (Dusek et al., 2006; Rose et al., 2010; Pöhlker et al., 2016; Burkart et
63 al., 2011). The PNSD can account for 84–96% of the variability in the CCN concentrations (N_{CCN}) (Dusek
64 et al., 2006), while CCN activities may also play a significant role in the N_{CCN} (Quinn et al., 2008; Cai
65 et al., 2018; Ovadnevaite et al., 2017; Liu et al., 2018; Crosbie et al., 2015), which are primarily governed
66 by the particle size, chemical composition, mixing state, surface tension, and hygroscopicity (Köhler,
67 1936; Seinfeld and Pandis, 2016). Among these factors, the impact of hygroscopicity on CCN activities
68 has received great attention in recent years (Petters and Kreidenweis, 2007; Ajith et al., 2022; Rose et al.,
69 2010). Petters and Kreidenweis (2007) proposed the κ - Köhler theory based on the Köhler theory to
70 quantify the ability of aerosol particles to absorb moisture and become CCN based on the aerosol
71 hygroscopicity parameters (κ). Ajith et al. (2022) showed that 64% of particles can be activated as CCN
72 when κ is equal to 0.37, whereas when κ decreases to 0.23, only 48% of particles can be activated in the
73 tropical coastal area.

74 Significant seasonal variations in PNSD and hygroscopicity under both terrestrial and marine
75 environments were observed in previous field observations, leading to the seasonal variations in N_{CCN}
76 (Crosbie et al., 2015; Schmale et al., 2018; Burkart et al., 2011; Bougiatioti et al., 2009; Sihto et al., 2011;
77 Leena et al., 2016; Ross et al., 2003; Gras and Keywood, 2017; Quinn et al., 2019). Crosbie et al. (2015)
78 revealed that in the urban area of Arizona particles had larger sizes, higher hygroscopicity, and N_{CCN} was
79 also higher during winter, while a higher abundance of smaller particles was observed during summer
80 owing to stronger photochemical reactions. In pristine environments like mountain, coastal, and forested

81 regions, seasonal variations in N_{CCN} and PNSD were more pronounced than urban and rural areas
82 (Schmale et al., 2018). Pöhlker et al. (2016) observed significant differences in N_{CCN} between the wet
83 and dry seasons in the Amazon rainforest, while the κ values remained relatively stable. They also noted
84 increased particle concentrations and aerosol hygroscopicity, both subject to the impact of long-range
85 transport originating from anthropogenic emissions. Observations in marine areas during different
86 seasons are relatively scarce compared with those in inland areas. Gras (1995) found that both particle
87 concentration and N_{CCN} in the Southern Ocean reached their peaks during summer and gradually decrease
88 to their valleys in winter. Quinn et al. (2019) showed that sea spray aerosols make a relatively significant
89 contribution to N_{CCN} only during winter in the Western North Atlantic, while in other seasons, the primary
90 contribution comes from biogenic aerosols oxidized from dimethyl sulfide (DMS). Zheng et al. (2020)
91 revealed that sulfate dominates the particle condensational growth to CCN sizes during summer in the
92 Eastern North Atlantic, while secondary organic aerosols played a significant role in particle growth
93 throughout all seasons. These results indicate that CCN activity and concentration could vary in a large
94 range during different seasons. Thus, further observations across different seasons in marine
95 environments are needed to enhance our understanding of marine CCN activities and their seasonal
96 variations.

97 The South China Sea (SCS), located in Southeast Asia and bordered by China, the Indochinese
98 Peninsula, and Maritime Southeast Asia, is significantly influenced by air pollutants transported through
99 terrestrial air masses. Studies have shown that these pollutants play a crucial role in determining aerosol
100 concentration and properties in the region (Atwood et al., 2017; Xiao et al., 2017; Geng et al., 2019;
101 Liang et al., 2021; Sun et al., 2023; Qin et al., 2024). For instance, Xiao et al. (2017) reported that 69.7%
102 of nitrate and 57.5% of sulfate in the SCS originated from fossil fuel combustion, particularly coal
103 burning in Chinese coastal regions. Additionally, Liang et al. (2021) and Sun et al. (2023) observed an
104 increase in the organic fraction and concentration of submicron aerosols when the region was influenced
105 by terrestrial air masses from Mainland China and the Indochinese Peninsula in the northern SCS. Further
106 studies highlighted the variation in aerosol properties under different air mass influences. Atwood et al.
107 (2017) found a significant bimodal particle distribution with a κ value of 0.65 in the southern SCS under
108 marine air mass influence, whereas a unimodal distribution with a κ of 0.4 was observed under
109 continental air mass influence.

110 The SCS experiences a typical monsoon climate with distinct seasonal wind direction changes
111 (Wang et al., 2009). The northeast monsoon, occurring from November to March, is characterized by
112 stronger average wind speeds and longer period compared to the southwest monsoon, which dominates
113 from June to August. The transitional periods occur from April to May and September to October. During
114 the northeast monsoon, air pollutants are primarily transported to the SCS by terrestrial air masses from
115 China (Xiao et al., 2017; Liu et al., 2014; Geng et al., 2019). In contrast, during the summer, pollutants
116 mainly originate from terrestrial air masses from the Indochinese Peninsula and Maritime Southeast Asia
117 (Geng et al., 2019; Liang et al., 2021; Sun et al., 2023). These varying sources of anthropogenic emissions
118 exert different impacts on CCN activity differently across seasons. Additionally, when the marine
119 boundary layer over the SCS is influenced by various natural and anthropogenic sources, resulting in
120 altered aerosol properties, the characteristics of cumulus clouds are correspondingly affected (Miller et
121 al., 2023). This indicates that aerosol-cloud interactions vary between winter and summer seasons.
122 However, due to limited observational data, our understanding of seasonal variations in CCN activity in
123 the SCS remains incomplete. Conducting comprehensive observational studies on CCN activity across
124 different seasons is essential for improving our understanding of aerosol-cloud interactions on the SCS.

125 In this study, we conducted two shipborne observations in the SCS during summer (May 5–June 9,
126 2021) and winter (December 19–29, 2021). Our observations with online instruments focused on
127 measuring aerosol chemical composition, PNSD, and CCN activation in the region. Our results provide
128 valuable insights into the differences in CCN activity between winter and summer, as well as the
129 influence of different types of terrestrial air masses on CCN activity in the SCS across different seasons.

130 **2. Methodology**

131 **2.1 Cruise information and onboard measurements**

132 **2.1.1 Cruise information**

133 This study consists of two research cruises conducted during the summer and winter of 2021,
134 respectively. These two cruises were interdisciplinary scientific expeditions, integrating fields such as
135 marine geology, oceanography, and atmospheric environment. The primary objective in atmospheric
136 environment was to investigate the impact of summer and winter monsoons on the atmospheric
137 environment of the South China Sea (SCS). The summer and winter cruises were carried out respectively
138 by the vessels "Tan Kah Kee" and "Sun Yat-sen University". The "Tan Kah Kee" is an oceanographic

139 research vessel with a length of 77.7 meters, a beam of 16.24 meters, and a displacement of 3611 tons.
140 The "Sun Yat-sen University" is a comprehensive oceanographic training vessel with a total length of
141 114.3 meters, a beam of 19.4 meters, and a displacement of 6880 tons.

142 The first cruise was from May 5th to June 9th, 2021. The cruise started from Xiamen Port and
143 traversed from the northern to the central-southern South China Sea, and then circled back near Hainan
144 Island, and finally returned to Xiamen Port. The second cruise was from December 19th to December
145 29th, 2021. It began from Gaolan Port in Zhuhai and reached the vicinity of Yongxing Island, and
146 ultimately returned to Gaolan Port (Fig. 1a). Unfortunately, due to adverse weather conditions, such as
147 strong winter monsoon winds causing poor sea conditions, and the fact that it was the first scientific
148 deployment of the research vessel Sun Yat-sen University, the winter cruise had a shorter duration and
149 covered a narrower spatial range, remaining only in the northern SCS (Fig. S1), compared to the summer
150 cruise. On both cruises, most of the instruments were housed in a single compartment and the sampling
151 lines were extended from the window of the compartment to the height of the ship's bridge (~17 m above
152 sea level) (Fig. 1a).

153 **2.1.2 Size-resolved cloud condensation nuclei activity measurement**

154 The size-resolved CCN activity was measured using the scanning mobility CCN analysis (SMCA)
155 method proposed by Moore et al. (2010), employing a combination of a scanning mobility particle sizer
156 (SMPS) system and a cloud condensation nuclei counter (CCNc-200, DMT Inc., USA) (Fig. S2). The
157 SMPS system consisted of a differential mobility analyzer (DMA; model 3082, TSI, Inc.) and a
158 condensation particle counter (CPC; model 3756, TSI Inc.). The SMPS and the CCNc system were used
159 to measure PNSD and size-resolved CCN number concentration at a mobility size range of 10–500 nm
160 and 10–593 nm in summer and winter campaign, respectively. Unfortunately, due to the malfunction of
161 flow sensor in the column B on both cruises, only the data from column A is presented in this study.
162 During the SMCA measurement, the particles were first passed through a Nafion dryer to remove
163 moisture, then neutralized using a neutralizer. After that, they were subjected to size selection with a
164 DMA. The particles were then split between a CPC (1 L min⁻¹) for particle concentration measurement
165 and a CCNc (0.5 L min⁻¹) for CCN measurement at a specific supersaturation (SS). To maintain sample
166 flow through the DMA, dilution air (0.5 L min⁻¹) was added to the CPC inlet stream. The effect of the
167 dilution air was accounted for in the PNSD data processing (Fig. S2). The supersaturation of the CCNc

168 was set at 0.2 %, 0.4 %, and 0.7 % in summer campaign and 0.1%, 0.2 %, 0.4 %, and 0.7 % in winter
169 campaign, respectively. During the measurement process, each supersaturation level was held constant
170 for 20 minutes, with the DMA completing a full scanning cycle every 5 minutes. During the
171 measurements, supersaturation levels varied incrementally between 0.1% and 0.2%, 0.2% and 0.4%, and
172 0.4% and 0.7%, with temperature stabilization times ranging from a few seconds to several tens of
173 seconds. However, reducing the supersaturation from 0.7% to 0.1% or 0.2% required approximately 5
174 minutes for stabilization. For data processing, only instances where the temperature remained stable
175 throughout the DMA scanning phase were included in the analysis. Before the measurements, the CCNc
176 was calibrated with ammonium sulfate ((NH₄)₂SO₄) particles at each set SS. Detailed description of the
177 instrument configuration and calibration can be found in Cai et al. (2018). The uncertainty in the
178 instrument's measurement of size-resolved particle number concentration is approximately 5%-6%
179 (Morre et al. 2010).

180 **2.1.3 Aerosol chemical composition measurement**

181 The chemical composition of atmospheric non-refractory submicron particulate matter (NR-PM₁),
182 including sulfate, nitrate, organics, ammonium, and chloride, was measured using an online time-of-
183 flight ACSM (ToF-ACSM; Aerodyne Inc., USA). The sampling time of the ToF-ACSM was
184 approximately 10 min. The ionization efficiency (IE) and relative ionization efficiency (RIE) values of
185 the instrument were calibrated using ammonium nitrate (NH₄NO₃) and ammonium sulfate ((NH₄)₂SO₄)
186 both before the start and after the completion of the campaigns. The calibration gives an IE value of
187 103.4 ions pg⁻¹ and 98.9 ions pg⁻¹ for nitrate in summer and winter cruises, respectively. The RIE values
188 for ammonium were 3.31 and 3.33 during the summer and winter, respectively, while the ones for sulfate
189 were 1.02 and 0.81 during the summer and winter, respectively. The collection efficiency (CE) was
190 determined as shown in Sun et al. (2023) and time-independent CE values were used in this study.
191 Detailed CE calculation and discussion can be found in the supplementary (Text S1, and Fig. S3). The
192 values obtained using the time-independent CE method show a deviation of approximately 3% compared
193 to those obtained with a constant CE of 0.5. Assuming an average aerosol density of 1.5 g cm⁻³ (Geller
194 et al., 2006), the mass concentrations measured by the SMPS and ToF-ACSM exhibit a strong overall
195 correlation, with correlation coefficients of 0.84 in summer and 0.93 in winter. During the pre-onset
196 phase of the summer monsoon (prior to May 24), periodic discrepancies were observed between the

197 ACSM and SMPS data, likely due to the influence of refractory aerosol (ie. Black carbon). This issue is
198 discussed in detail in Text S1. The black carbon (BC) mass concentrations were measured using an
199 aethalometer (Model AE33, Magee Scientific, USA) with a 1-minute time resolution (Drinovec et al.,
200 2015). Notably, the BC mass concentrations obtained from AE33 are referred to as equivalent BC mass
201 concentrations, as they represent the combined light absorption of BC at 880 nm. Prior to entering the
202 AE33, the sampled air was passed through a PM_{2.5} cyclone (BGI Inc., Waltham, MA, USA) to exclude
203 particles larger than 2.5 μm.

204 **2.1.4 Meteorological parameter measurements**

205 The meteorological elements, including temperature, relative humidity (RH), wind speed, and wind
206 direction, were measured by the combined automatic weather station (AWS430, Vaisala Inc., Finland)
207 onboard the vessels (Sun et al., 2024). During the winter cruises, meteorology data before 12.22 was
208 missed due to the calibration for the automatic weather station (WXT536, Vaisala Inc., Finland) before
209 12.22. The timeseries of meteorological data were presented in Fig. S5. The AWS430 provides
210 measurement accuracies of ±2% for wind speed, ±2% for wind direction, ±0.3°C for temperature, and
211 ±1% for relative humidity (within the range of 0–90%). Similarly, the WXT536 offers accuracies of ±3%
212 for wind speed, ±3% for wind direction, ±0.3°C for temperature, and ±3% for relative humidity (within
213 the range of 0–90%) (www.vaisala.com).

214 **2.2 Data analysis**

215 **2.2.1 CCN activation**

216 The size-resolved number concentration of total particle and cloud condensation nuclei were
217 obtained from the SMPS and CCNc through the SMCA method. The activation diameter was determined
218 by fitting the activation ratio (AR, N_{CCN}/N_{CN}) and dry diameter at each supersaturation through the
219 following equation:

$$220 \quad AR = \frac{B}{1 + \left(\frac{D_P}{D_{50}}\right)^C}, \quad (1)$$

221 where AR indicates the size-resolved AR, D_P represents dry particle diameter (nm); B, C, and D_{50} are the
222 three fitting parameters, representing the asymptote, the slope, and the inflection point of the sigmoid,
223 respectively (Moore et al., 2010). The D_{50} parameter, also known as the critical diameter, corresponds to

224 the particle size at which 50% of the particles are activated at a specific SS. The fitting results from
225 SMCA method measured in this study are presented in Fig. S6.

226 The hygroscopicity parameter (κ) which represents CCN activity according to κ -Köhler equation is
227 calculated as follows (Petters and Kreidenweis, 2007):

$$228 \quad \kappa = \frac{4A^3}{27D_{50}^3(\ln S_c)^2}, \quad A = \frac{4\sigma_{s/a}M_w}{RT\rho_w} \quad (2)$$

229 where ρ_w is the density of pure water (about 997.04 kg m^{-3} at 298.15 K), M_w is the molecular weight of
230 water ($0.018 \text{ kg mol}^{-1}$), $\sigma_{s/a}$ corresponds to the surface tension of the solution-air interface and is assumed
231 to be equal to the surface tension of pure water ($\sigma_{s/a}=0.0728 \text{ N m}^{-1}$ at 298.15 K), R is the universal gas
232 constant ($8.314 \text{ J mol}^{-1} \text{ K}^{-1}$), T denotes thermodynamic temperature in kelvin (298.15 K), and D_{50} is the
233 critical diameter (in m). Additionally, it is noting that the estimated κ values refer to particles with the
234 D_{50} .

235 According to κ -Köhler theory, in the following discussion, the hygroscopicity of small particles is
236 associated with hygroscopicity at high SS, whereas the hygroscopicity of large particles is linked to
237 hygroscopicity at low SS.

238 During part of the summer measurement period, the D_{50} at 0.7% supersaturation ranged between 30
239 and 40. However, due to lower concentrations during these times, instrument noise introduced greater
240 measurement uncertainty, as demonstrated in Fig. S7. Consequently, the average D_{50} and κ at 0.7% SS
241 are not included in Table 1.

242 2.2.2 Closure Method

243 According to Petters and Kreidenweis. (2007), κ can be predicted by a simple mixing rule based on
244 chemical volume fractions:

$$245 \quad \kappa_{sim} = \sum_i \varepsilon_i \kappa_i \quad (3)$$

246 where ε_i and κ_i are the volume fraction and hygroscopicity parameter for the specific dry component
247 in the mixture. We obtained ε from aerosol chemical composition measured by the ToF-ACSM. In this
248 study, κ for $(\text{NH}_4)_2\text{SO}_4$ (0.48), NH_4NO_3 (0.58), and NaCl (1.1) represent the κ of SO_4^{2-} , NO_3^- , and Cl^-
249 provided by the ToF-ACSM (Huang et al., 2022). Besides, the κ of organic was 0.1 at this study according
250 to Huang et al. (2022). The density of $(\text{NH}_4)_2\text{SO}_4$, NH_4NO_3 , NaCl and organic are 1769 kg m^{-3} , 1720 kg
251 m^{-3} , 2165 kg m^{-3} , and 1400 kg m^{-3} (Huang et al., 2022; Gysel et al., 2007).

252 2.2.3 CCN concentration and activation ratio calculation

253 Due to the malfunction of the column B, the CCN concentration (N_{CCN}) was calculated based on the
254 size-resolved AR at a specific SS from SMCA method and observed particle number concentration. It
255 can be calculated by the following equation (Cai et al., 2018):

$$256 N_{CCN}(SS) = \int_0^{\infty} AR(SS, D_p) N_{CN}(D_p) dD_p \quad (4)$$

257 where $N_{CCN}(SS)$ is the CCN concentration at a specific SS, $AR(SS, D_p)$ is the ratio of N_{CCN} at a specific
258 SS to N_{CN} on a specific diameter from the SMCA method and $N_{CN}(D_p)$ is the particle number
259 concentration at a specific diameter (D_p). Due to the absence of direct measurements for total N_{CCN} , we
260 refer to the N_{CCN} derived from Eq. (4) as observed values ($N_{CCN,obs}$) in this study. Previous research has
261 shown that this method (size-resolved CCN from one column in CCNc-200) provides results closely
262 matching those obtained from direct measurement (from another column in CCNc-200), supporting its
263 reliability (Meng et al., 2014; Lathem and Nenes, 2011).

264 The N_{CCN} (referred as $N_{CCN,sim}(SS)$) can be predicted by D_{50} from closure method ($D_{50,sim}(SS)$)
265 and N_{CN} according to following equation (Jurányi et al., 2011):

$$266 N_{CCN,sim}(SS) = \int_{D_{50,sim}(SS)}^{\infty} N_{CN}(D_p) dD_p \quad (5)$$

267 where the $D_{50,sim}(SS)$ is calculated based on the eq. (2) and (3).

268 The bulk AR at a specific SS can be calculated by:

$$269 AR(SS) = \frac{N_{CCN,obs}(SS)}{N_{CN,tot}} \quad (6)$$

270 where the $N_{CN,tot}$ represents the total particle number concentration.

271 To investigate the impact of the fraction and mixing state of aerosol on N_{CCN} , two CCN simulation
272 schemes are applied in this study (Patel et al., 2021).

273 (1) Internal-mixed scheme: the aerosol composition from the ToF-ACSM was assumed to be size-
274 independent and internally mixed. All aerosols have an identical chemical composition in the
275 whole size range. N_{CCN} is calculated by κ_{sim} and measured PNSD according to Eq. (2), Eq. (3),
276 and Eq. (5) (Fig. S8a).

277 (2) External-mixed scheme: the aerosol composition from the ToF-ACSM was assumed to be size-
278 independent and externally mixed. Four types of aerosols ($(NH_4)_2SO_4$, NH_4NO_3 , NaCl and
279 organic) are assumed to have the same proportion for all sizes. The D_{50} from each species was

280 calculated by Eq. (2) according to their κ values mentioned in 2.2.2. N_{CCN} is calculated
281 according to the Eq. (5) (Fig. S8b and Table S1).

282 To access the simulation result from these two schemes, normalized mean bias (NMB) was used in
283 this study:

$$284 \quad NMB = \frac{\sum(N_{CCN,sim} - N_{CCN,obs})}{\sum N_{CCN,obs}} \quad (7)$$

285 where $N_{CCN,sim}$ is the simulated N_{CCN} from two schemes, and $N_{CCN,obs}$ is the observed N_{CCN} .

286 2.2.4 Fitting of log-normal modes to particle number size distributions

287 The multi-lognormal distribution function (Eq. (8)) is used to parameterize and optimize the
288 descriptions of the measured PNSD (Heintzenberg, 1994) and is widely applied in aerosol research (Cai
289 et al, 2020; Boyer et al., 2023; Zhu and Wang, 2024). An automatic mode-fitting algorithm (Hussein et
290 al., 2005) is used to generate the model-fitted results.

$$291 \quad f(D_p, \bar{D}_{pg,i}, N_i, \sigma_{g,i}) = \sum_{i=1}^n \frac{N_i}{\sqrt{2\pi} \log(\sigma_{g,i})} \times \exp \left[-\frac{[\log D_p - \log \bar{D}_{pg,i}]^2}{2(\log \sigma_{g,i})^2} \right] \quad (8)$$

292 where D_p is the diameter of a particle. Each lognormal mode is characterized by three parameters:
293 the mode number concentration (N_i), geometric variance ($\sigma_{g,i}$), and geometric mean diameter (GMD,
294 $\bar{D}_{pg,i}$). The total number of lognormal modes used to describe the PNSD is denoted by n . These modes
295 are fitted using an algorithm applied to each particle size distribution, with one to three log-normal
296 distributions used per time step. The algorithm classifies the PNSD into nucleation, Aitken, and
297 accumulation modes based on their geometric mean diameters (GMDs). The GMD for nucleation modes
298 (GMD1) typically ranges from 3 to 30 nm, for Aitken modes (GMD2) from 30 to 100 nm, and for
299 accumulation modes (GMD3) above 100 nm (Heintzenberg, 1994; Hussein et al., 2005; Zhu and Wang,
300 2024).

301 2.2.5 Backward trajectory simulation and cluster analysis

302 Backward trajectory calculations were performed using the MeteoInfo, an open-source software
303 (Wang, 2014) to determine potential source origins. Weekly GDAS1 (Global Data Assimilation System
304 at a resolution of 1°) files were downloaded from the NOAA Air Resource Laboratory (ARL) website
305 (<https://www.ready.noaa.gov/gdas1.php>). The calculation of backward trajectories is performed every
306 1hour based on the location mentioned below, generating 72-hour backward trajectories at 500m.

307 To clarify the sources of air masses, the cluster analysis was applied in this study, which was
308 performed by TrajStat, a plug-in module of MeteoInfo, based on k-means method
309 (http://meteothink.org/docs/trajstat/cluster_cal.html). According to the report by the China
310 Meteorological Administration (Chao et al., 2022), the summer monsoon in 2021 broke out during the
311 sixth pentad of May. Therefore, based on the timing of the monsoon onset and the actual trajectory of the
312 ship, we selected two representative midpoints of the ship track for backward trajectory calculations and
313 cluster analysis in summer: the midpoint of the ship's track before the onset of the summer monsoon
314 (May 5-23) and the midpoint of the track after the summer monsoon began (May 24-June 9). In the
315 winter cruise, backward trajectories calculation and cluster analysis were performed at two specific
316 locations: the ship's anchorage near Big Ten-thousand Mountain Island (December 19-22 and December
317 27-29) and the midpoint between Dawan Mountain Island and Yongxing Island (December 23-26). To
318 ensure the accuracy of the backward trajectory calculations and cluster analysis, we compared the
319 trajectories at the midpoints with those from the ship's actual locations to verify consistency in air mass
320 sources (Fig. S9). Minor discrepancies may exist between the air mass origins at certain midpoints and
321 the actual ship locations. However, overall, the air mass origins at the midpoints are representative of
322 those at the actual locations. We further examined the trajectories for each cluster to verify their alignment
323 with the air mass origins they represent (Fig. S10). The results demonstrate that cluster analysis was well
324 conducted. Additionally, figure S10 illustrates the average altitude variation as the age in hours increases
325 across different periods. During summer, the altitude of the clusters remained below 880 hPa, indicating
326 that they resided within the boundary layer (about 800 hPa). While in winter, the altitude of the clusters
327 was higher than in summer, especially for the cluster during the mixed period (peaked at about 755 hPa).
328 However, these clusters were generally within or close to the boundary layer. These results suggest that
329 the back trajectories could represent the characteristics of the air masses originating from these specified
330 regions.

331 **2.2.5 Data quality control**

332 To ensure reliable atmospheric samples in the SCS and mitigate the influence of research vessel
333 emissions, we applied the following data processing procedures (Huang et al., 2018; Cai et al., 2020;
334 Liang et al., 2021).

335 Firstly, we identified organic compounds, black carbon (BC), and small particulate matter (41.4 nm
336 particles) as indicators of ship emissions, recognizing their sudden peak values as indicative of the ship's
337 own emissions.

338 Secondly, we accounted for the relative positions of the ship's chimney and the sampling tube.
339 During the summer cruise, we excluded data corresponding to a relative wind direction (with respect to
340 the ship's bow) between 150° and 270° and a relative wind speed (with respect to the ship's speed) of less
341 than 2.5 m s⁻¹ (Fig. S12a, Fig. S13a1, and Fig. S14a-c). During the winter cruise, we excluded data for a
342 relative wind direction between 150° and 220° and a relative wind speed of less than 2.5 m s⁻¹ (Fig. S12b,
343 Fig. S13b1, and Figs. S14d-f).

344 Applying these criteria, 74.8% of the data in summer and 92.2% in winter (both at 10-minute
345 resolution) were classified as “clean” and retained for analysis. The timeseries of data before and after
346 quality control is shown in Fig. S15.

347 **3. Results and discussion**

348 **3.1 CCN concentration and aerosol characteristics over SCS in summer and winter**

349 Figure 2 presented the timeseries of PNSD (a1 and a2), NR-PM₁ mass concentrations and fractions
350 (b1 and b2, c1 and c2), number concentrations of CCN (d1 and d2), and hygroscopicity κ -values (e1 and
351 e2) during two campaigns in summer and winter. During the summer cruise, we observed two distinct
352 periods around the onset of the summer monsoon. The South China Sea (SCS) summer monsoon began
353 in the sixth pentad of May (Chao et al., 2022). In winter, the influence of the winter monsoon persisted
354 throughout the entire observation period (Fig. 1c). Despite our measurements being limited to the
355 northern SCS in winter, the impact of the Northeast Monsoon on the SCS was evident.

356 The average particle number concentration in summer (6966 cm⁻³) was higher than in winter (4988
357 cm⁻³), primarily due to the higher number concentration of Aitken-mode particles in summer (Fig. 3a-b).
358 In summer, particles were concentrated in smaller sizes, whereas in winter, particle size distribution was
359 relatively balanced between the Aitken mode (2185 cm⁻³) and the accumulation mode (2176 cm⁻³) (Fig.
360 3a-b).

361 The average mass concentration of NR-PM₁ was 3.76 $\mu\text{g m}^{-3}$ in summer and increased to 9.39 $\mu\text{g m}^{-3}$
362 in winter (Fig. 3c-d). In summer, the dominant aerosol component was sulfate (45.5%), followed by
363 organics (35.8%), ammonium (12.9%), nitrate (4.0%), and chloride (1.9%) (Fig. 3c), similar to the

364 pattern observed in the northern SCS during the summer of 2018 (Fig. 3e) (Liang et al., 2021). However,
365 in winter, organics became the predominant aerosol component (37%), with nitrate (22.2%) replacing
366 sulfate (18.9%) as the highest proportion of inorganic components (Fig. 3d). Although N_{CN} was higher
367 in summer than in winter, the particle volume size distribution indicates that a higher fraction of particles
368 was concentrated in larger size in winter, which significantly influenced mass concentration, resulting in
369 a higher NR-PM₁ concentration (Fig. S16).

370 The average number concentration of cloud condensation nuclei (N_{CCN}) in summer was higher than
371 in winter at all supersaturation (SS) levels (Table 1). The ratio of N_{CCN} between summer and winter was
372 smaller at high SS ($N_{CCN, winter}/N_{CCN, summer} = 0.51$ and 0.54 at 0.4% SS and 0.7% SS, respectively)
373 compared to low SS ($N_{CCN, winter}/N_{CCN, summer} = 0.62$ at 0.2% SS), likely due to the significant difference in
374 number concentration of Aitken-mode particles between the two seasons (Fig. 3a-b). Compared to the
375 observation in the Yellow Sea, a region similarly influenced by terrestrial air masses from mainland
376 China, the N_{CCN} were lower in winter, while in summer, the N_{CCN} were more comparable to those
377 observed in the Yellow Sea (4821 cm^{-3} at 0.63% SS) (Park et al., 2018).

378 The aerosol hygroscopicity (κ) was higher in summer than that in winter (Table 1). Besides, the
379 hygroscopicity pattern varied between seasons: in summer, κ increased with SS (from 0.47 to 0.54
380 between 0.2% SS and 0.4% SS), while in winter, κ decreased with SS (from 0.50 to 0.15 between 0.1%
381 SS and 0.7% SS) (Fig. 3a-b). This contrasting trend may be related to the reduced sulfate fraction in
382 smaller sizes during winter, as sulfate production via DMS oxidation is diminished due to lower sea
383 surface temperatures in winter (18.0°C) compared to summer (29.3°C), which in turn inhibits DMS
384 production by phytoplankton (Bates et al., 1987; Kouvarakis and Mihalopoulos, 2002). Additionally, it
385 could be linked to the mixing state of the particles, with further discussion provided in the following
386 sections. The winter κ pattern was similar to observations in the Western North Pacific (Table 1) (Kawana
387 et al., 2020). Additionally, the winter κ values were comparable to those in Guangzhou (Cai et al., 2020),
388 adjacent to the SCS, indicating that the northern SCS is influenced by air masses from Mainland China
389 under the significant influence of the Northeast Monsoon during winter.

390 **3.2 Anthropogenic influence on CCN concentration in different seasons**

391 Cluster analysis revealed distinct periods influenced by various air masses. In summer, three
392 terrestrial air mass sources were identified: Luzon Island (referred to as “Luzon”), Palawan Island, and

393 the Indochinese Peninsula, along with a marine air mass source (Fig. 4a). Given the limited influence of
394 air masses from Palawan Island, this period was excluded from the study. Consequently, the study
395 focused on periods dominated by air masses from Luzon (“Luzon” period), the Indochinese Peninsula
396 (“Indochinese Peninsula” period), and marine sources (“Marine-s” period). In winter, the air mass
397 sources included Mainland China, a mixture of Mainland China and the South China Sea (referred to as
398 “Mixed”), and a marine source (Fig. 4b). These were classified as the “Mainland China” period, “Mixed”
399 period, and “Marine-w” period, respectively.

400 As shown in figure 5, terrestrial air masses could significantly affect the aerosol chemical
401 composition in the SCS, resulting in higher NR-PM₁ mass concentration and a higher fraction of organic
402 compounds compared to those influenced by marine air masses. Additionally, the particles number
403 concentration in the accumulation mode and the N_{CCN} at low supersaturation (SS) were higher during
404 periods influenced by terrestrial air masses (“Luzon” period) than those during marine air mass periods
405 (Table 2). Notably, we were able to obtain an accurate D₅₀ at 0.7% supersaturation only during the “Luzon”
406 period in summer. Due to the relatively lower hygroscopicity compared to other summer periods, the
407 corresponding D₅₀ at 0.7% SS ranged between 40 and 60 nm, with relatively high concentration of CN
408 and CCN (Fig. S7), allowing for a more precise measurement of D₅₀. As a result, the κ at 0.7% SS shown
409 in Fig. 7 was specific to the Luzon period in summer.

410 In summer, the “Luzon” period exhibited the highest N_{CN}, attributed to the elevated particle
411 concentration in the Aitken mode, compared to all other periods in both summer and winter (Fig. 6a and
412 Table 1). This high fraction of Aitken mode particles led to the lowest bulk AR among the summer
413 periods (Fig. 7a), as a larger fraction of particles centered on a size range lower than the D₅₀ (Fig. 7b).
414 Furthermore, the prevalence of a higher fraction of Aitken mode particles during terrestrial air mass
415 periods is commonly correlated with the influence of fresh anthropogenic emissions (Beddows et al.,
416 2015), which could lower the hygroscopicity and consequently suppress the bulk AR.

417 In the “Indochinese Peninsula” period, the N_{CN} was lower than it in the “Marine-s” period (Table
418 2). This difference was mainly due to the variation of Aitken mode particles, while accumulation mode
419 particles were higher during the “Indochinese Peninsula” period than in “Marine-s” period (Table 2). The
420 “Marine-s” period primarily occurred during the transition phase before the onset of summer monsoon,
421 when wind direction shifted from east (Luzon Island direction) to southwest (Indochinese Peninsula
422 direction). Anthropogenic emissions from Luzon Island still affected the marine atmosphere, leading to

423 higher concentrations of Aitken mode particles compared to the “Indochinese Peninsula” period (Table
424 2). The higher fraction of accumulation mode particles and higher hygroscopicity during the
425 “Indochinese Peninsula” period resulted in a higher bulk AR compared to the “Luzon” period. Despite a
426 higher organic fraction in NR-PM₁ during the “Indochinese Peninsula” period (Fig. 5), hygroscopicity
427 was still higher due to a higher oxidation degree of organics, indicated by a higher m/z 44 to 43 ratio
428 (5.87 compared to 5.60 in the “Luzon” period) (Lambe et al., 2011; Jimenez et al., 2009). Additionally,
429 higher wind speeds during this period (7.26 m s⁻¹ compared to 3.18 m s⁻¹ in the “Luzon” period)
430 potentially led a higher fraction of sea salt (Huang et al., 2022), resulting a higher aerosol hygroscopicity.
431 Unfortunately, owing to instrument limit, sea salt cannot be detected by the ToF-ACSM.

432 In winter, nitrate accounted for the highest fraction of NR-PM₁ (25.4%) during the “Mainland
433 China” period compared to other periods (Fig. 5d). Due to similar hygroscopicity between nitrate and
434 sulfate, as well as comparable inorganic fractions between the “Mainland China” and “Luzon” periods,
435 κ at 0.2% SS was also similar between these two periods (0.30 and 0.33, respectively) (Fig. 7b). However,
436 aerosol hygroscopicity at small sizes (high SS) was much lower in the “Mainland China” period than in
437 the “Luzon” period (Fig. 7b), contributing to the low bulk AR in the “Mainland China” period (Fig. 7a).
438 The BC mass concentration was higher during the “Mainland China” period (2.25 $\mu\text{g m}^{-3}$) compared to
439 the “Luzon” period (0.72 $\mu\text{g m}^{-3}$). This suggests that the lower hygroscopicity in smaller particles during
440 the “Mainland China” period may be attributed to a larger fraction of hydrophobic BC. Additionally, as
441 discussed in Section 3.1, the reduced biological activity during winter, which results in a decline in the
442 fraction of small-particle sulfate and an increase in the fraction of organics, may also contribute to this
443 low hygroscopicity in small particles (at high SS, fig 7b). The similar fractions of Aitken mode and
444 accumulation particles indicated that PNSD could not fully explain the low bulk AR in the “Mainland
445 China” period. Overall, lower N_{CN} and bulk AR in the “Mainland China” period compared to the “Luzon”
446 period resulted in a lower N_{CCN} .

447 During the “Mixed” period, N_{CCN} was lower than in the “Mainland China” period, which can be
448 attributed to the decreased N_{CN} (Table 2). However, accumulation mode particles dominated, unlike in
449 other terrestrial air mass periods (Fig. 6), resulting in a significantly higher bulk AR compared to the
450 “Mainland China” period. Organic aerosol hygroscopicity was also higher during the “Mixed” period,
451 supported by a higher m/z 44 to 43 ratio (3.88 vs. 3.10 in the “Mainland China” period), which explains
452 the greater hygroscopicity despite a higher organic fraction in NR-PM₁. Additionally, the lower BC

453 concentration in the “Mixed” period ($1.20 \mu\text{g m}^{-3}$ vs. $2.25 \mu\text{g m}^{-3}$ in the “Mainland China” period)
454 suggests a smaller BC fraction. Moreover, the higher wind speeds during the “Mixed” period (10.77 m s^{-1}
455 vs. 7.14 m s^{-1} in the “Mainland China” period) could have increased the sea salt fraction, further
456 enhancing aerosol hygroscopicity.

457 3.3 CCN closure analysis

458 The CCN closure method is a widely used approach that connects CCN activity with aerosol
459 chemical composition (Cai et al., 2018; Meng et al., 2014; Deng et al., 2013). Studies have demonstrated
460 that the aerosol mixing state is crucial for accurately parameterizing CCN activity (Su et al., 2010; Wang
461 et al., 2010; Ervens et al., 2010). Moreover, the CCN closure method provides a framework for
462 investigating the influence of aerosol mixing states on CCN activity (Padró et al., 2012; Wang et al.,
463 2018; Patel et al., 2021). In this study, we applied two schemes based on the CCN closure method, as
464 described in Section 2.2.3, which consider aerosol composition and mixing state. The fitting parameters
465 and coefficient of determination (R^2) are presented in Table 3, while the fitting plots for both schemes
466 are shown in Figures S17 and S18. Besides, the NMB from these schemes was presented in Fig. 8.

467 In summer, the NMB was always lower than 0, which indicated that simulated aerosol
468 hygroscopicity was lower than observed value (Fig. 8). Sea salt which cannot be detected by the ToF-
469 ACSM may account for higher fraction in summer due to low aerosol concentration in summer (Fig. 3c),
470 resulting in the underestimation of aerosol hygroscopicity. The NMB exhibits similar trends with changes
471 in SS in all three periods in summer. Better fitting result appeared at high SS, which indicated a greater
472 underestimation of the hygroscopicity of larger particles. Besides, “Internal-mixed” scheme had more
473 precious result than it in “External-mixed” scheme in summer (Fig. 8), suggesting the aerosol was
474 primary internally mixed in summer.

475 In winter, the “External-mixed” scheme always showed a better result than “Internal-mixed” scheme
476 at high SS (0.4% SS and 0.7% SS), indicating that particles in small size were mainly externally mixed.
477 Considering the low hygroscopicity of small-sized particles in winter, it is likely that a significant fraction
478 of these particles consists of externally mixed BC, which probably originated from fresh anthropogenic
479 emissions and remains unmixed with other inorganic salts and organics. As BC ages, inorganic and
480 organic components adhere to it, which would lead to the increase of diameter and particles tended to be
481 internally mixed (Sarangi et al., 2019). This transition resulted in higher hygroscopicity in large-sized

482 particles compared to the smaller-sized particles. Besides, overestimation of aerosol hygroscopicity at
483 high SS could be owing to a higher fraction of non- or less- hygroscopic component (such as organic and
484 BC) at small particle sizes. The predicted N_{CCN} at 0.1% SS are 20%-40% lower than the observed
485 concentrations, whereas the predictions at 0.2% SS more closely match the observed values (Fig. 8). This
486 discrepancy may be due to the higher fraction of sea salt in larger particles. However, due to instrumental
487 limitations, the ToF-ACSM cannot detect BC and sea salt. Future observations including BC and sea salt
488 are needed to better assess their effects on aerosol hygroscopicity in the South China Sea (SCS). In
489 addition, further study of size-resolved aerosol composition can also enhance the understanding on CCN
490 activity in the SCS.

491 **4. Conclusion**

492 In this study, we investigated the seasonal variations of cloud condensation nuclei (CCN) activity
493 in the South China Sea (SCS) and explored the impact of anthropogenic emissions. Shipborne
494 observations were conducted during the summer (May 5–June 9) and winter (December 19–29) of 2021.
495 We measured CCN activity, chemical composition, and particle number size distribution (PNSD) using
496 several onboard instruments, including a ToF-ACSM, a CCNc, an SMPS, and an AE33. Observations
497 included periods before and after the summer monsoon onset and periods influenced by the winter
498 monsoon.

499 Our results show that particle number concentration (N_{CN}) and CCN number concentration (N_{CCN})
500 were higher in summer than in winter, while the mass concentration of non-refractory submicron
501 particulate matter (NR- PM_{10}) was lower in summer. This difference is primarily attributed to the
502 predominance of Aitken mode particles in summer, contrasted with a higher concentration of
503 accumulation mode particles in winter. Additionally, aerosol hygroscopicity and bulk AR were found to
504 be higher in summer than in winter.

505 Backward trajectory and cluster analyses identified distinct influences from various air masses. In
506 summer, we identified periods affected by terrestrial air masses from Luzon Island (the “Luzon” period)
507 and the Indochinese Peninsula (the “Indochinese Peninsula” period), alongside a period influenced by
508 marine air masses (the “Marine-s” period). In winter, the periods were influenced by terrestrial air masses
509 from Mainland China (the “Mainland China” period), a mix of Mainland China and marine sources (the

510 “Mixed” period), and marine air masses (the “Marine-w” period). Terrestrial air mass periods exhibited
511 higher NR-PM₁ mass concentrations, organic fractions, and N_{CCN}, particularly at low supersaturation,
512 compared to those influenced by marine air masses.

513 During the “Luzon” period, high N_{CCN} was observed, attributed to high N_{CN}, especially in the Aitken
514 mode. This high concentration in the Aitken mode resulted in a low bulk AR at 0.2% SS, indicating a
515 higher fraction of primary organic aerosol with low hygroscopicity. This caused lower overall
516 hygroscopicity compared to other summer periods. The lower ratio of m/z 44 to 43 also suggested a
517 lower oxidation degree of organics in this period. In the “Indochinese Peninsula” period, a higher fraction
518 of the accumulation mode particles compared to the “Luzon” period led to a higher bulk AR, combined
519 with increased hygroscopicity.

520 In winter, the “Mainland China” period was characterized by a high nitrate fraction in the NR-PM₁.
521 The similar inorganic fractions in the NR-PM₁ between the “Mainland China” and “Luzon” periods
522 resulted in comparable aerosol hygroscopicity at low supersaturation (0.2% SS). However, at higher
523 supersaturation levels (0.4% and 0.7% SS), the “Mainland China” period demonstrated significantly
524 lower hygroscopicity, which led to a reduced bulk AR at elevated supersaturation. During the “Mixed”
525 period, accumulation mode particles predominated, leading to a high bulk AR. This indicated an aging
526 process during transport, with more oxidized organics and higher aerosol hygroscopicity. The lower black
527 carbon (BC) fraction and the higher sea salt fraction from high wind speed contributed to higher
528 hygroscopicity in the “Mixed” period compared to the “Mainland China” period, despite the high organic
529 fraction.

530 The CCN closure analysis, considering aerosol composition and mixing state, revealed that aerosols
531 in summer were primarily internally mixed, while in winter, small-sized aerosols were primarily
532 externally mixed. This distinction is crucial for climate models predicting N_{CCN} in the SCS. The
533 underestimation of aerosol hygroscopicity in summer suggests that the effect of sea salt should be
534 considered.

535 Our study highlights significant seasonal differences in CCN activity in the SCS and the influence of
536 different types of terrestrial air masses. Future measurements including size-resolved aerosol
537 composition and obtain more precise measurements of BC and sea salt are needed to better understanding
538 CCN activity in this region. Additionally, our observation in winter focused on the CCN activity over the
539 northern SCS, while the influence of air masses from Mainland China in remote SCS was still unclear.

540 Further observations in remote SCS areas could help clarify the anthropogenic influence during winter
541 under the effect of the winter monsoon.

542

543 *Data availability.* Data from the measurements are available at [https://doi.org/10.6084/m9.figshare.](https://doi.org/10.6084/m9.figshare.25472545)
544 [25472545](https://doi.org/10.6084/m9.figshare.25472545) (Ou et al., 2024).

545

546 *Supplement.* The supplement related to this article is available online.

547

548 *Author contributions.* **HO, MC, and JZ** designed the research. **YZ, XN, BL, and CS** performed the
549 measurements. **HO, MC, QS, and SM** analyzed the data. **SZ and HW** provided useful comments on the
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551

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

553

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563

564

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Location	period	N_{CN} (cm ⁻³)	N_{CCN} (cm ⁻³)	Hygroscopicity (κ)	Bulk AR	D_{50} (nm)	Reference
South China Sea	2021.05.05- 2021.06.09	6966±9249	2640±3639 (0.20% SS)	0.47±0.21 (0.20% SS)	0.37±0.16 (0.20% SS)	96±19 (0.20% SS)	This study
			4392±6415 (0.40% SS)	0.54±0.21 (0.40% SS)	0.63±0.17 (0.40% SS)	57±9 (0.40% SS)	
			5215±6862 (0.70% SS)	0.87±0.17 (0.70% SS)			
Northern South China Sea	2021.12.19- 2021.12.29	4988±3474	1086±691 (0.10% SS)	0.50±0.21 (0.10% SS)	0.23±0.09 (0.10% SS)	145±18 (0.10% SS)	This study
			1625±1110 (0.20% SS)	0.31±0.10 (0.20% SS)	0.33±0.12 (0.20% SS)	107±13 (0.20% SS)	
			2218±1503 (0.40% SS)	0.19±0.05 (0.40% SS)	0.44±0.13 (0.40% SS)	79±7 (0.40% SS)	
Northern South China Sea	2018.08.06- 2018.08.27	3463		0.38±0.09 (0.18% SS)			Cai et al., 2020
			1544 (0.34% SS)	0.40±0.08 (0.34% SS)	/	/	
				0.38±0.08 (0.59% SS)			
Remote South China Sea	2012.09.14- 2012.09.26	503±455	450±388 (0.14% SS)		0.47±0.16 (0.14% SS)		Atwood et al., 2017
			675±516 (0.38% SS)	0.54±0.14 (0.14% SS)	0.72±0.17 (0.38% SS)	/	
			698±555 (0.53% SS)	0.50±0.21 (0.38% SS)	0.79±0.15 (0.53% SS)		
Western North Pacific	2015.03.04- 2015.03.26	/	/	0.75±0.21 (0.11% SS)	0.40±0.22 (0.11% SS)		Kawana et al., 2020
				0.51±0.16 (0.24% SS)	0.50±0.22 (0.24% SS)	/	
				0.45±0.16 (0.60% SS)	0.70±0.23 (0.60% SS)		
Guangzhou	2014.11- 2014.12	/	3103±1913 (0.10% SS)	0.37±0.11 (0.10% SS)	0.26±0.10 (0.10% SS)	156 ± 19 (0.1% SS)	Cai et al., 2018
			5095±2972 (0.20% SS)	0.29±0.09 (0.20% SS)	0.41±0.14 (0.20% SS)	107 ± 17 (0.2% SS)	
			6524±3783 (0.40% SS)	0.18±0.07 (0.40% SS)	0.53±0.15 (0.40% SS)	78 ± 15 (0.4% SS)	
			7913±4234 (0.70% SS)	0.15±0.06 (0.70% SS)	0.64±0.13 (0.70% SS)	58 ± 11 (0.7% SS)	

Yellow Sea	2017.04- 2017.05	7622± 4038	4821±1763 (0.63% SS)	/	/	/	Park et al., 2018
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835 Table 1. The number concentration of particle and cloud condensation nuclei at different supersaturation (SS), the hygroscopicity and bulk activation ratio (AR), and activation
836 diameter (D_{50}) at different SS in different studies.

Cluster	Summer			Winter		
	Indochinese Peninsula	Luzon	Marine	Mainland China	Marine	Mixed
N_{CCN} (cm^{-3})						
0.1% SS	\	\	\	1359±669	439±223	945±400
0.2% SS	1200±787	4066±4748	1135±800	2058±1095	614±318	1460±514
0.4% SS	1650±1187	7804±8608	1812±1052	2792±1478	830±424	1801±640
0.7% SS	2239±1367	10480±9741	2515±1523	3514±1841	1024±463	2101±757
N_{CN} (cm^{-3})						
Total	2699±2147	14674±13844	3033±2366	6875±3263	1728±465	2918±1204
Nucleation	111±206	1543±3341	238±426	893±925	214±281	141±191
Aikten	1156±1261	8653±8815	1668±1526	3089±2017	732±337	806±427
Accumulat ion	1434±1444	3764±4157	1121±929	2923±2440	781±313	1975±831
Bulk AR						
0.1% SS	\	\	\	0.21±0.07	0.26±0.10	0.32±0.04
0.2% SS	0.49±0.13	0.31±0.17	0.40±0.13	0.30±0.09	0.36±0.14	0.51±0.05
0.4% SS	0.73±0.09	0.55±0.18	0.68±0.14	0.40±0.10	0.49±0.16	0.63±0.06
0.7% SS	0.98±0.15	0.76±0.16	0.90±0.13	0.50±0.09	0.61±0.18	0.73±0.06

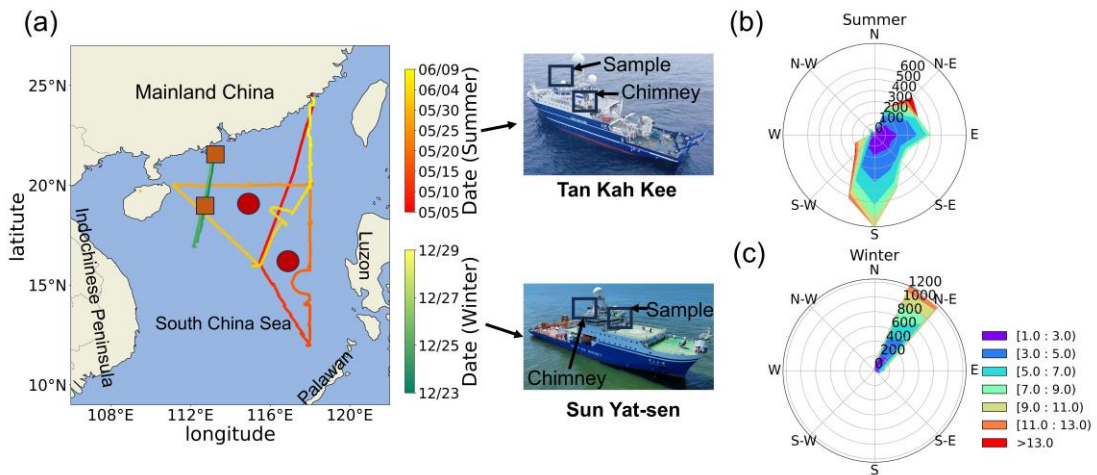
838 Table 2. The number concentration of particle, cloud condensation nuclei, and bulk activation ratio in
839 different periods.

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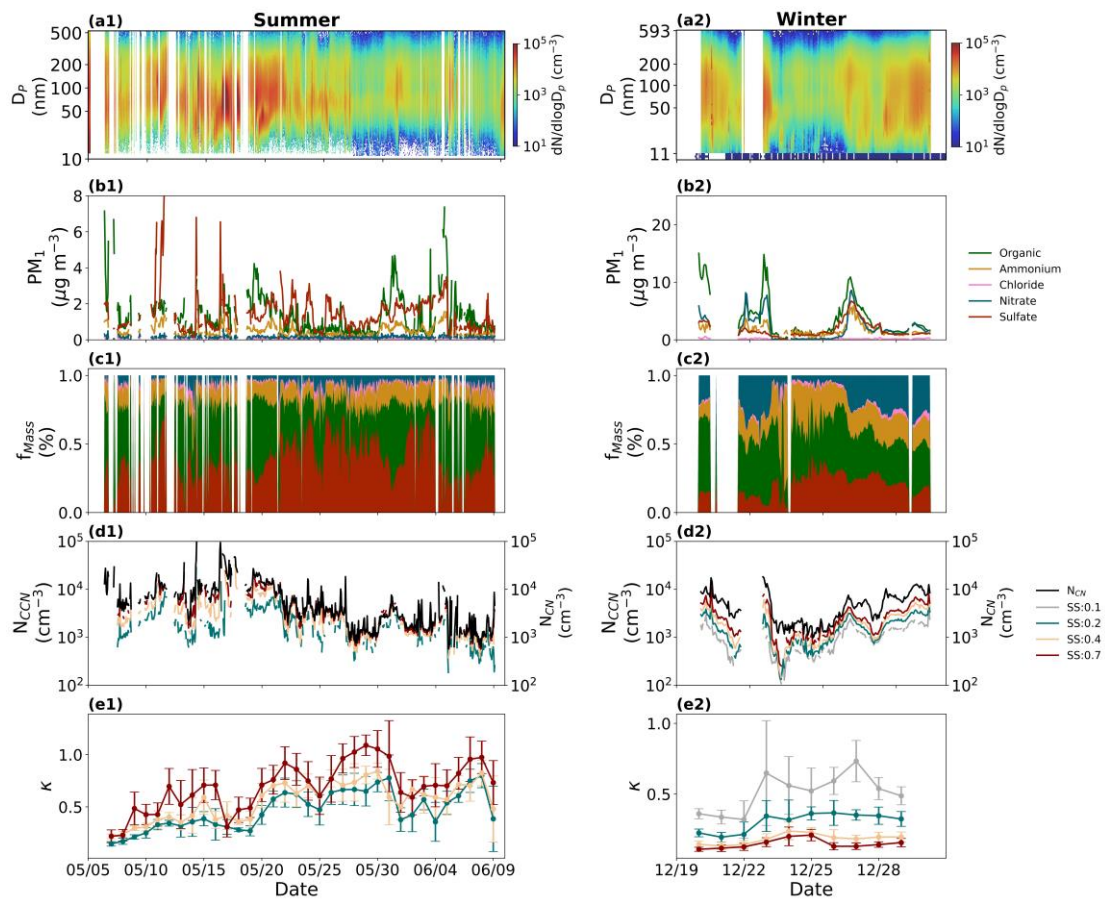
Cluster	Summer			Winter		
	Luzon	Indochinese Peninsula	Marine	Mainland China	Mixed	Marine
Internal scheme						
0.1% SS	\	\	\	0.91 (0.97)	0.72 (0.95)	0.71 (0.94)
0.2% SS	0.83 (0.89)	0.89 (0.81)	0.76 (0.96)	1.13 (0.96)	1.01 (0.99)	0.94 (0.97)
0.4% SS	0.90 (0.96)	0.90 (0.98)	0.89 (0.97)	1.34 (0.97)	1.14 (0.98)	1.04 (0.98)
0.7% SS	0.91 (0.93)	0.96 (0.92)	0.88 (0.98)	1.38 (0.97)	1.16 (0.99)	1.04 (0.96)
External scheme						
0.1% SS	\	\	\	0.80 (0.97)	0.62 (0.95)	0.59 (0.94)
0.2% SS	0.74 (0.88)	0.77 (0.79)	0.80 (0.96)	1.01 (0.97)	0.90 (0.99)	0.81 (0.97)
0.4% SS	0.78 (0.93)	0.80 (0.97)	0.82 (0.96)	1.23 (0.97)	1.05 (0.98)	0.95 (0.98)
0.7% SS	0.80 (0.92)	0.89 (0.92)	0.80 (0.98)	1.24 (0.98)	1.11 (0.99)	1.00 (0.96)

842 Table 3. The slope and coefficient of determination (in parentheses) in CCN closure analysis at
843 different supersaturations in different periods.

844

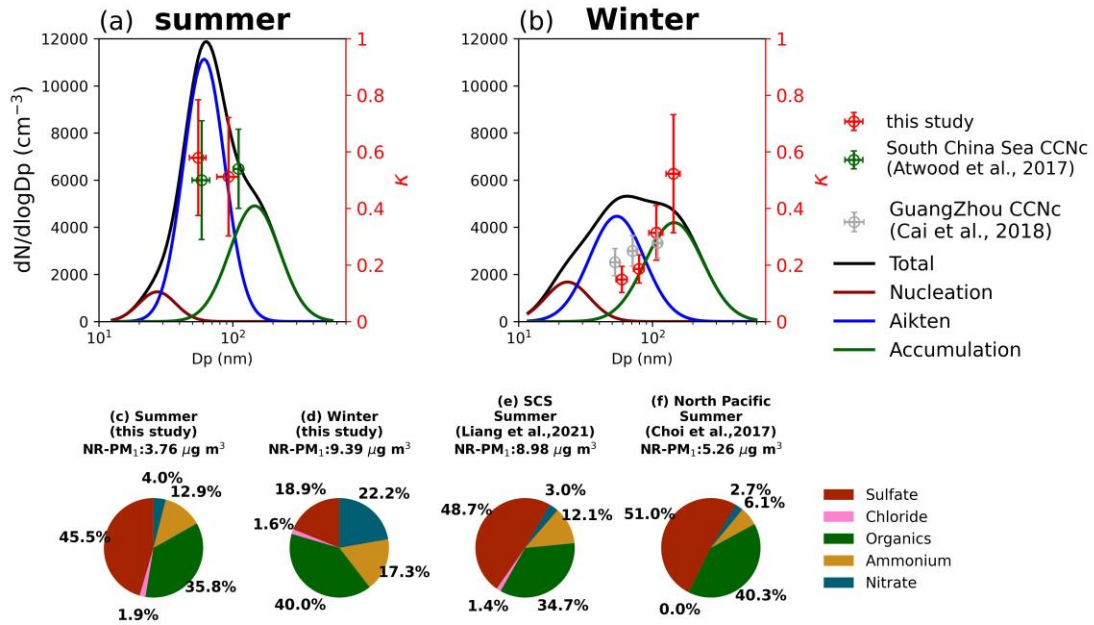


847 Figure 1. The cruises of two shipborne observations, and the location of sample line and chimney of Tan
 848 Kah Kee, and Sun Yat-sen scientific vessel (a); Wind rose of the wind direction and wind speed in
 849 summer and winter cruises; The radius represents the frequency of wind direction occurrences, and the
 850 shaded areas indicate wind speed (b) and (c). The red circles are the midpoints of the ship trajectory
 851 selected for backward trajectory and cluster analysis in summer and the orange squares are the midpoints
 852 of the ship trajectory selected for backward trajectory and cluster analysis in winter.



854

855 Figure 2. Timeseries of (a) particle number size distribution, (b) mass concentration of NR-PM1, and (c)
 856 its fraction, (d) number concentration of total particle and cloud condensation nuclei under the
 857 supersaturation of 0.1%, 0.2%, 0.4%, and 0.7%, and (e) aerosol hygroscopicity. The number 1 in figure
 858 number means timeseries in summer and number 2 means it in winter.



859

860

Figure 3. Particle number size distribution in summer (a) and winter (b); The red markers represent the activation diameters and hygroscopicity parameters corresponding to 0.1%, 0.2%, 0.4%, and 0.7% supersaturations in this study (without 0.1% in summer). The green markers represent the hygroscopicity parameters reported in Atwood et al. (2017) for the southern South China Sea during summer. The gray markers represent the hygroscopicity parameters documented in Cai et al. (2018) for the Pearl River Delta region during winter. The fraction of NR-PM₁ in summer (c) and winter (d) in this study, in northern SCS reported by Liang et al. (2021) (e), and in North Pacific reported by Choi et al. (2017) (f).

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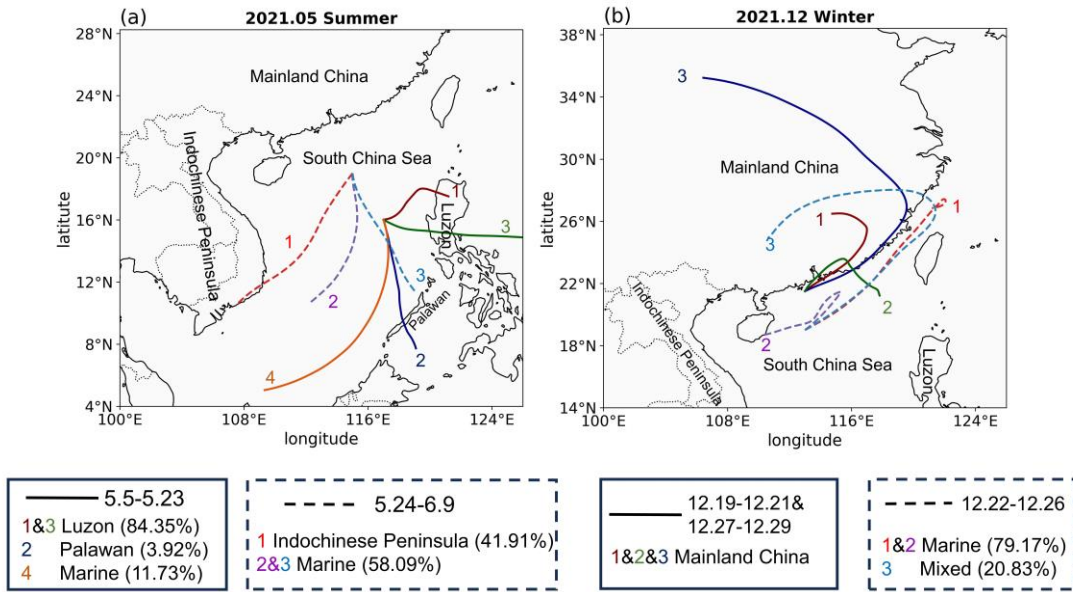
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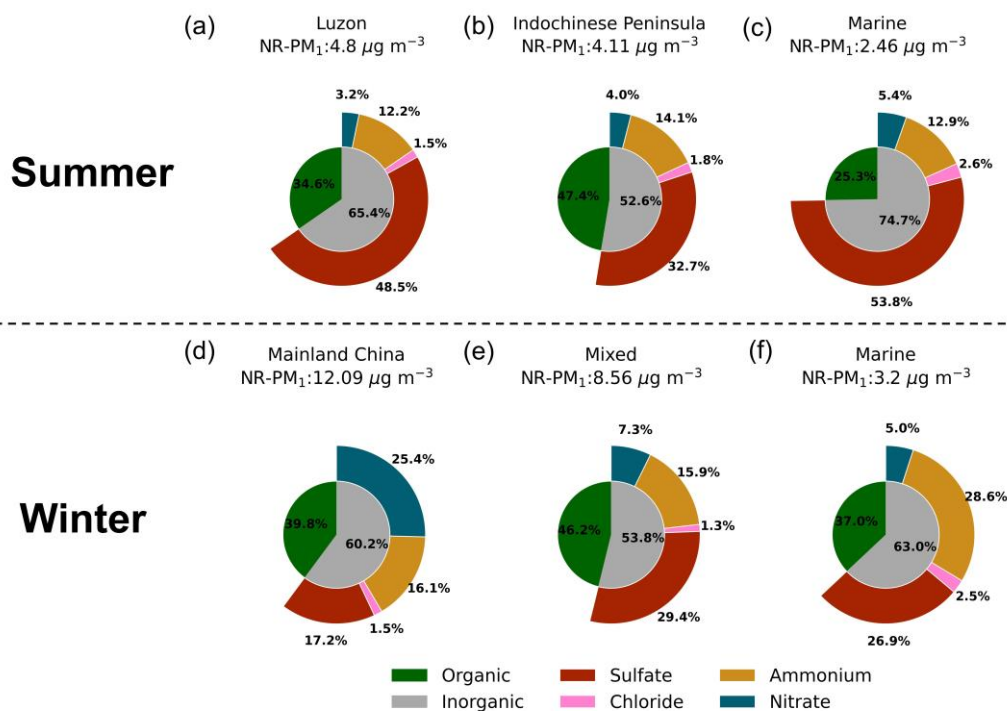
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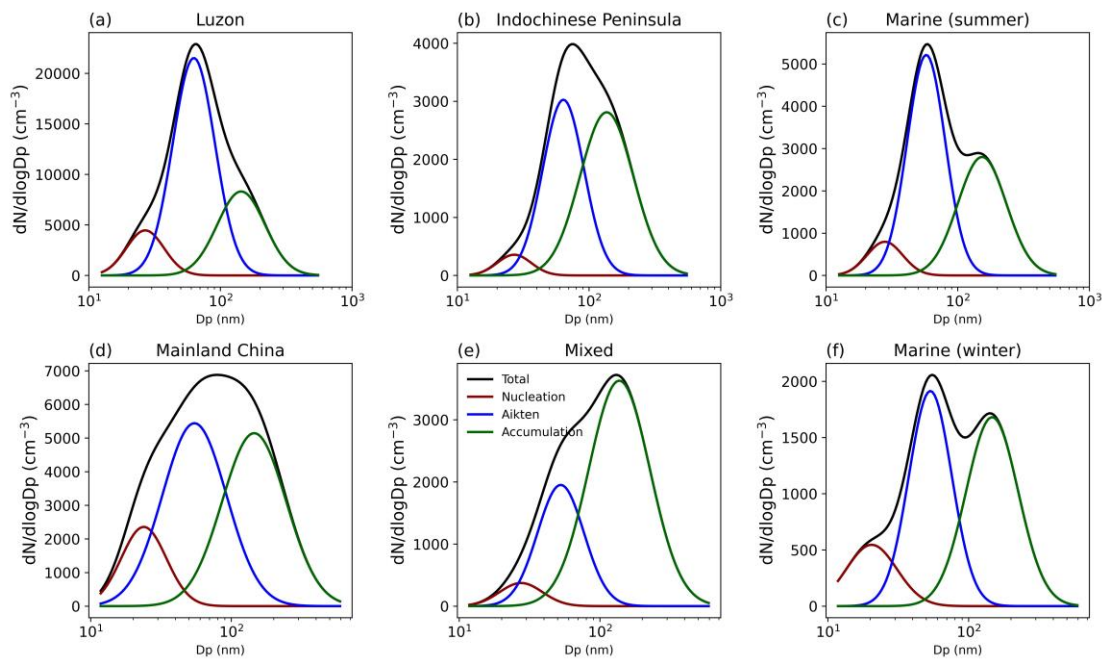
868 Figure 4. The cluster analysis results in summer (a) and winter (b). The solid line in summer means
 869 cluster analysis from May 5 to May 24 and the dash line in summer means cluster analysis from May 25
 870 to June 9; The solid line in winter means cluster analysis from Dec 19 to Dec 21 and Dec 27 to Dec 29,
 871 and the dash line in winter means cluster analysis from Dec 22 to Dec 26.



873

874 Figure 5. The fraction of NR-PM₁ in “Luzon” period (a), “Indochinese Peninsula” period (b), and875 “Marine-s” period (c) in summer. The fraction of NR-PM₁ in “Mainland China” period (d), “Mixed”

876 period (e), and “Marine-w” period (f) in winter.



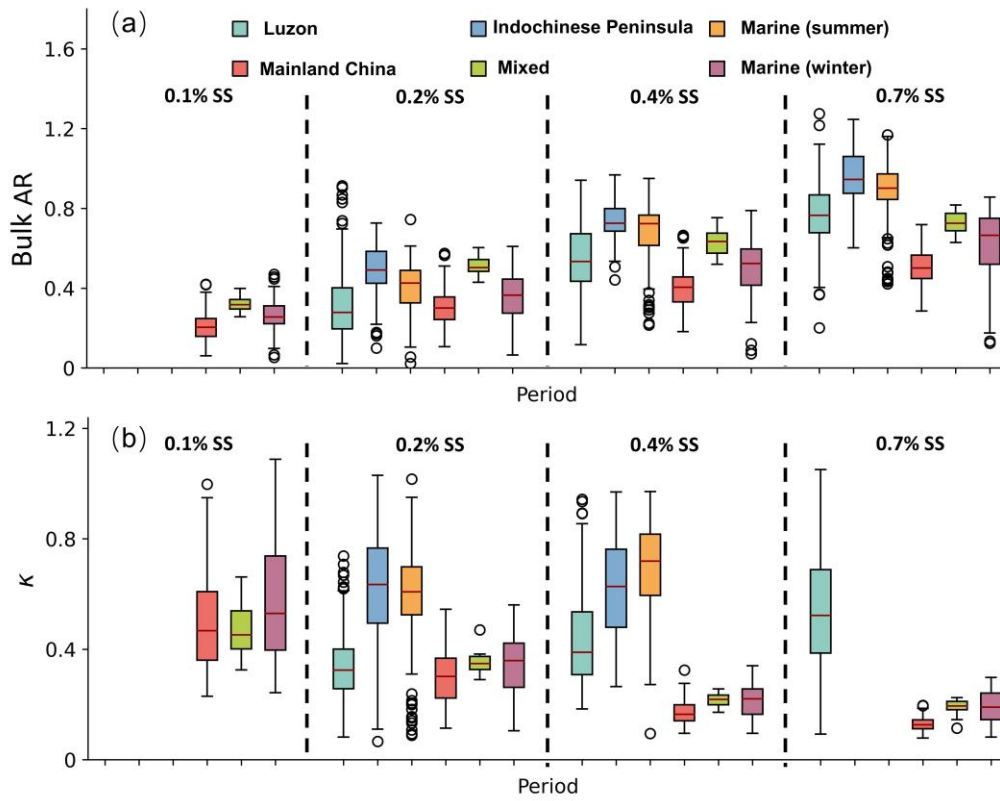
878

879 Figure 6. The particle number size distribution (PNSD) in “Luzon” period (a), “Indochinese Peninsula”

880 period (b), and “Marine-s” period (c) in summer. The PNSD in “Mainland China” period (d), “Mixed”

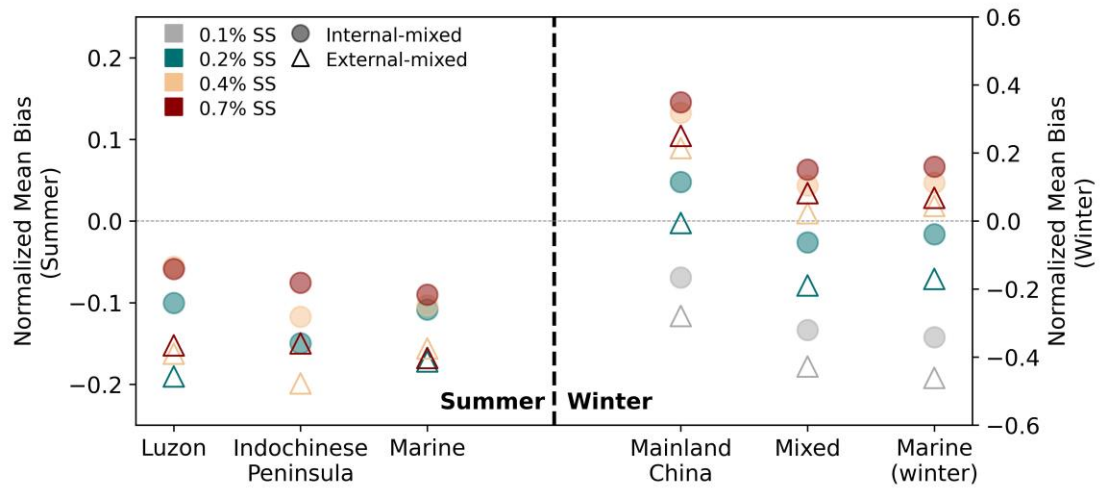
881 period (e), and “Marine-w” period (f) in winter.

882



883

884 Figure 7. The bulk activation ratio (AR) at different supersaturation (SS) in different periods (a); The
885 aerosol hygroscopicity (κ) at different supersaturation (SS) in different periods (b).



886

887 Figure 8. The normalized mean bias (NMB) calculated by “Internal-mixed” scheme and “External-mixed”
 888 scheme according to CCN closure method. The marker of circle means “Internal-mixed” scheme and the
 889 marker of triangle means “External-mixed” scheme. Different colors mean different supersaturations.

890