1	Effects of continental emissions on Cloud Condensation
2	Nuclei (CCN) activity in northern South China Sea during
3	summertime 2018
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21	Abstract. Aerosol particles in marine atmosphere have been shown to significantly affect cloud
22	formation, atmospheric optical properties, and climate change. However, high temporally and spatially
23	resolved atmospheric measurements over sea are currently sparse, limiting our understanding of aerosol
24	properties in marine atmosphere. In this study, a ship-based cruise campaign was conducted over northern
25	South China Sea (SCS) region during summertime 2018. Chemical composition of non-refractory PM ₁

26 (NR-PM₁), particle number size distribution (PNSD) and size-resolved cloud condensation nuclei (CCN)

27	activity were measured by a time-of-flight aerosol chemical speciation monitor (ToF-ACSM), and the
28	combination of a cloud condensation nuclei counter (CCNc) and a scanning mobility particle sizer
29	(SMPS), respectively. Overall, aerosol particles exhibited a unimodal distribution centering at 60~80 nm
30	and chemical composition of the NR-PM $_1$ was dominated by sulfate (~46%) which likely originated from
31	anthropogenic emissions rather than dimethyl sulfide (DMS) oxidation. Two polluted episodes were
32	respectively observed at the beginning (P1) and at the end (P2) of the campaign and both were
33	characterized by high particle number concentrations (N _{CN}) which originated respectively from local
34	emissions and from emissions in inland China via long range transport as shown by back trajectory
35	analysis. The concentrations of trace gases (i.e., O_3 , CO , NO_X) and particles (N_{CN} and N_{CCN} at ss=0.34%)
36	were elevated during P2 and decrease with the offshore distance, further suggesting important impacts
37	of anthropogenic emissions from the inland Pearl River Delta (PRD) region on the northern SCS. Two
38	relatively clean periods (C1 and C2) prior to and after tropical storm Bebinca were classified due to
39	substantial removal of pollutants by strong winds and rainfalls accompanying with the storm. During C1
40	and C2 periods, the air was affected by air masses from southwest and from Indo-China Peninsula,
41	respectively. Chemical composition measurements showed an increase of organic mass fraction during
42	P2 compared to C2; however, no obviously different κ values were obtained from the CCNc
43	measurements, implying that the air masses carried pollutants from local sources during long range
44	transport. We report an average value of about 0.4 for aerosol hygroscopicity parameter κ which falls
45	within the literature values (i.e., 0.2-1.0) for urban and remote marine atmosphere. In addition, our results
46	showed that the CCN fraction (N $_{\rm CCN}/N_{\rm CN,tot}$) and the κ values obtained from the CCNc measurements
47	(ss=0.34%) had no clear correlation either with the offshore distance or with concentrations of the
48	particles. Our study highlights dynamical variations of particle properties and the impact of long range

- 49 transport from the China continent and Indo-China Peninsula on the northern SCS region during
- 50 summertime.

1 Introduction

52	Aerosol particles directly affect global radiation balance by scattering and absorbing solar radiation.
53	Meanwhile, they can alter cloud microphysics, lifetime, and albedo, indirectly affecting heat transfer
54	through atmosphere (Stocker, 2013). However, high uncertainties still exist on their contributions to the
55	climatic impact, partly owing to our limited knowledge on spatial and temporal distribution of aerosol
56	particles and their properties in various environments. Thus, it is essential to conduct field measurements
57	under different environments to obtain chemical and physical properties of particles, including chemical
58	composition, particle number size distribution (PNSD), and cloud condensation nuclei (CCN) activity,
59	in order to better understand the radiation forcing induced by aerosol particles.
60	The CCN activity describes how particles grow into cloud droplets and further affect cloud
61	development. Whether particles can be activated as CCN is determined by their chemical composition,
62	hygroscopicity, size, and ambient supersaturation (ss). Generally, the CCN activity can be described by
63	Köhler theory based on the water activity in solution, surface tension, molecular weight of water,
64	temperature, and diameter of the particle (Köhler, 1936). Alternatively, the hygroscopicity parameter κ
65	proposed by Petters and Kreidenweis (2007) can be used to characterize the CCN activity. Aerosol
66	hygroscopicity describes the ability of particles to grow by absorbing moisture in ambient environments.
67	The κ values can be measured in subsaturation (RH<100%) condition by the hygroscopicity-tandem
68	differential mobility analyzer (HTDMA) measurements or in supersaturation (RH>100%) by the cloud
69	condensation nuclei counter (CCNc) measurements.
70	Field measurements for the CCN activity have been conducted primarily in terrestrial environments
71	(e.g., urban cities, forested areas, and remote countryside areas) (Rose et al., 2010; Wang et al., 2010;
72	Cerully et al., 2011; Pierce et al., 2012; Hong et al., 2014; Cai et al., 2018). Cerully et al. (2011) reported

73	κ values ranging from 0.1 to 0.4 in forest during the 2007 EUCAARI campaign and concluded that the
74	κ values obtained from the HTDMA measurements were generally 30% lower than those from the CCNc
75	measurements. Wang et al. (2010) showed that the mixing state of particles was important in predicting
76	the CCN number concentration (N_{CCN}). Cai et al. (2018) found that the CCN activity increased by
77	decreasing the surface tension through increase of organic fractions in particles based on the
78	measurements of the CCN activity, hygroscopicity, and chemical composition in the Pearl River Delta
79	(PRD) region. Progresses on the aforementioned field measurements conducted in the continental
80	environments have substantially improved our understanding of the influence of aerosols in global
81	radiation forcing and precipitation under the terrestrial environments.
82	Aerosol particles in the marine atmosphere, on the other hand, have been well known to significantly
83	affect cloud development, atmospheric optical properties, and climate change (Johnson et al., 2004;
84	Ackerman et al., 2004; Mulcahy et al., 2008). Fewer field measurements were conducted in the oceanic
85	atmosphere than those in land, leading to less characterization of marine aerosol particles. Remote
86	sensing and ship-based cruise methods are two typical approaches employed to measure aerosol
87	properties in marine environments (Durkee et al., 1986; Kim et al., 2009; Lehahn et al., 2010; Huang et
88	al., 2018). Compared to ship-based measurements, remote sensing covers spatially a larger area and
89	temporally a longer period which are essential in the characterization of marine aerosols. For example,
90	Reid et al. (2013) employed remote sensing to describe long range transport patterns in the Southeast
91	Asia. The aerosol size information was compared between the retrievals from Moderate Resolution
92	Imaging Spectroradiometer (MODIS) and the measurements from ground-based radiometers such as
93	Aerosol Robotic Network (AERONET) over ocean (Kleidman et al., 2005). However, extensive cloud
94	coverages over oceanic region can significantly affect the quality and availability of satellite

95	measurements. Meanwhile, dry bias or clear-sky bias also challenge satellite measurements for obtaining
96	accurate data (John et al., 2011; Reid et al., 2013; Choi and Ghim, 2017). Moreover, remote sensing using
97	satellite sensors is limited in providing high time resolution (i.e., minutes), high spatial resolution (i.e.,
98	within tens of meters in dimension) data and specific particle properties (i.e., hygroscopicity and
99	chemical composition). Although ship-based measurements are limited in spatial coverage, they can
100	provide higher spatial and temporal resolution for obtaining comprehensive physical and chemical
101	properties of gas and aerosol particles. Huang et al. (2018) measured chemical composition of particles
102	with a high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS) over the Atlantic Ocean
103	aboard a campaign ship and found that about 19% of organics originated from continental long-range
104	transport. Kim et al. (2009) found that particle size distribution varied in a dynamic range, depending on
105	the meteorological conditions over the Yellow Sea and the East China Sea. Atwood et al. (2017) showed
106	that biomass burning, anthropogenic pollution from continent and ship emissions would affect the remote
107	South China Sea during the southwestern monsoon (SWM) season. However, few ship-based campaigns
108	are available in the literature on measurements of atmospheric composition including gases and aerosol
109	particles, especially in several important China sea regions (e.g., SCS).
110	The air over northern SCS is affected by anthropogenic pollution from the adjacent Pearl River
111	Delta region, China inner continent, and Indo-China Peninsula (Zhang et al., 2018). Furthermore, as one

116 wind to this region, which helps to remove air pollutants. However, on one hand, it has been found that

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of the most important and busy trading regions in China, the PRD and the northern SCS are subjected to

severe air pollution due to emissions from heavy loadings of cargo ships and fishing vessels (Lv et al.,

2018). Special weather patterns are dominant in the SCS during summertime which are characterized by

SWM and occasionally affected by typhoons. Typically, typhoon brings heavy precipitation and strong

117 downdrafts prior to a typhoon usually affect negatively atmospheric diffusion, leading to the 118 accumulation of the air pollutants in the region (Feng et al., 2007). On the other hand, marine background 119 particles and emissions from Indo-China Peninsula are brought into this region through SWM. As a result, 120 the physical and chemical properties of marine aerosol particles vary dynamically which can be 121 distinguished from those of continental particles. Differences (i.e., physical and chemical properties, life 122 cycle) between the two types of aerosol particles reflect different transport pathways and source origins 123 which are not well known. In addition, lack of understanding on aerosol characteristics will inevitably 124 hinder our ability to evaluate the impacts of aerosol particles on global radiation forcing and atmospheric 125 processes. Thus, ship-based field measurements are urgently needed in this region in order to understand 126 the CCN activity, chemical composition, particle size distribution, and their relationships with 127 continental and marine air masses. 128 In this study, we report results from a recent ship-based cruise measurement in the northern SCS 129 during summertime 2018. During the campaign, size-resolved CCN activity, chemical composition, and 130 particle number size distribution were measured by a CCNc, a time-of-flight aerosol chemical speciation 131 monitor (ToF-ACSM) and a scanning mobility particle sizer (SMPS), respectively. Temporal and spatial 132 distributions of the aerosol chemical and physical properties and impact of different air masses on the 133 properties were investigated. Our results provide valuable knowledge on the effects of long range 134 transport and on the atmospheric processes in the SCS.

136 2 Methodology

137 2.1 Ship-based campaign

138 The cruise campaign is a routine comprehensive exercise organized by Sun Yat-sen University (SYSU) during summertime 2018 (6th to 27th August) including a variety of multidisciplinary sciences 139 140 (i.e., atmosphere, ocean, chemistry, geology, and biology). The round-trip journey started and ended at 141 Huizhou port (22°43' N, 114°36' E), which is about 140 km from Guangzhou, traveling towards northern 142 SCS with an area between 19°37' N to 22°43' N and 113°44' E to 118°12' E. The ship track includes two 143 routes during which the vessel was anchored near the port due to tropical storm Bebinca as its track was shown in Fig. 1a, along with the complete, color-coded ship track. The first route started 7th August from 144 145 the port and arrived northeast of Dongsha Islands (20°45' N 118°12' E) on 10th August 2018, and then returned to anchor near the port during the typhoon period (11th to 15th August). The second route left the 146 147 port on 15th August toward Hong Kong and arrived at its south in the afternoon (18:00 local time, LT). The vessel then headed southeast for about 42 hours on 18th August and turned toward Dongsha Islands. 148 It anchored at several sites around this sea area and then returned on 24th August following a similar 149 150 pathway as the first route to Huizhou port on 27th August.

A commercial vessel with a capacity of 8000 ton was employed for the routine summer measurement campaign whose schematic diagram was shown in Fig. 1b. An air conditioned (T=298K) sea container of about 30 m² housed all the instruments which was listed in Table 1 and was placed in the front deck of the vessel. Trace gases, including O_3 , SO_2 , CO, NO_X (NO and NO_2), were measured by gas analyzers (model T400U, T100U, T300, and T200U, Teledyne API Inc., USA, respectively). Detailed descriptions of the major instruments used in the campaign could be found in the following subsection.

157	The aerosol sampling port with a $PM_{2.5}$ cyclone inlet was made of a 5 m long 3/8" o.d. stainless-steel
158	tube which extended outside of the container with an inclination angle of 45° to the deck. The inlet is
159	about 2.5 m above the deck and 1.5 m away from the container. All aerosol sampling flows firstly passed
160	through a Nafion dryer (model MD-700, Perma Pure Inc., USA) to reach a relative humidity (RH) lower
161	than 30%. The gas sample inlet made of a 2 m long 1/4" o.d. Teflon tube with a similar inclination angle,
162	also extended outside of the container.

164 2.2 Origins of air masses by HYSPLIT

165	The HYbrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model developed by
166	National Oceanic and Atmospheric Administration (NOAA) was used to investigate trajectories of air
167	movement for identification of source origins which might affect the northern SCS region during the
168	campaign. The model calculated the 72 hours back trajectories of air masses at 6 hours intervals arriving
169	at the campaign vessel. The arrival height of the trajectories was set to be 150 m, 500 m, and 1000 m
170	above the ground level, a reasonable representative of the air masses. The Global Data Assimilation
171	System (GDAS) $1^{\circ} \times 1^{\circ}$ meteorological data was employed to drive the HYSPLIT.

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173 **2.3 Measurements**

174 2.3.1 Size-resolved cloud condensation nuclei activity

The size-resolved CCN activity was measured with combination of a homemade scanning mobility
particle sizer system and a cloud condensation nuclei counter (model CCNc-200, DMT Inc., USA). The

177	homemade SMPS system consisted of a differential mobility analyzer (DMA, model 3081L, TSI., Inc.)
178	and a condensation particle counter (CPC, model 3787, TSI Inc.). The CCNc-200 has two parallel cloud
179	columns (column A and B) which measure the CCN concentrations (N $_{\rm CCN}$) at two specific ss at the same
180	time. Only the N_{CCN} measured by column A was discussed in this study. During the measurements, the
181	SMPS system was operated in a scanning mode. The sample particles after the Nafion dryer were firstly
182	neutralized by a X-ray neutralizer (model 3088, TSI., Inc., USA) and were subsequently classified by the
183	DMA. The selected particles were split into the CPC for measurements of total particle number
184	concentration (with a flow rate of 0.6 LPM) and the CCNc for measurements of the CCN number
185	concentration at a specific supersaturation (with a flow rate of 0.5 LPM). The SMPS and the CCNc
186	system were set to measure particle number size distribution and size-resolved CCN number
187	concentration at a mobility size range of 10-400 nm. The supersaturation of the CCNc was set to be
188	0.18%, 0.34%, and 0.59%. Before the measurements, the CCNc-200 was calibrated with ammonium
189	sulfate ((NH ₄) ₂ SO ₄) particles at three ss (0.18%, 0.34%, and 0.59%), detailed description of the
190	calibration could be found in Cai et al. (2018). The SMPS system was also calibrated with standard
191	polystyrene latex spheres (PSL, with a size of 20 nm, 50 nm, and 200 nm) prior to the campaign.
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193 2.3.2 Aerosol chemical composition

An Aerodyne time-of-flight aerosol chemical speciation monitor was deployed to measure bulk nonrefectory PM₁ chemical composition during the campaign. The ToF-ACSM can provide mass concentration of sulfate, nitrate, ammonium, chloride, and organics, except non-refectory components such as sea salt, black carbon, and crustal species. Detailed description of ToF-ACSM can be found in Fröhlich et al. (2013) and only a brief introduction relevant to this work was given here. During the

199	campaign, the measurement cycle of the ToF-ACSM was set to be about 10 min and the mass resolving
200	power was about 160. The sample flow dried by the Nafion dryer entered an automatic three-way valve,
201	of which one way was directly connected to the lens system and the other way was connected to a filter
202	before entering the aerodynamic lens. By switching the automatic valve periodically, the instrument can
203	measure the total signal without a filter and the background signal with a filter, thus the net signal
204	representing the chemical composition of the aerosol particles can be obtained. The aerodynamic lens
205	system removes particles larger than 1 μm (at aerodynamic diameter, $D_{\text{VA}})$ and has a relative low
206	transmission for small particles ($D_{VA} < 50$ nm). Monodisperse pure ammonium nitrate (NH ₄ NO ₃) and
207	ammonium sulfate ((NH ₄) ₂ SO ₄) particles generated by a homemade atomizer and then selected by a
208	DMA (about 300 nm in diameter) were used to calibrate the relative ionization efficiency (RIE) value of
209	NH_4 (RIE_{NH_4}) and SO_4 (RIE_{SO_4}) at the beginning and at the end of the campaign.

211 2.4 Data processing of CCN activation

The size-resolved N_{CN} and N_{CCN} measured by the SMPS and CCNc-200 system was used to calculate the activation ratio (AR), which was defined as the ratio of N_{CCN} to N_{CN} at each size bin. The size-resolved ARs were inverted based on the method described by Moore et al. (2010). The AR spectrum was then fitted using a three-parameter fit:

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$$\frac{N_{CCN}}{N_{CN}} = \frac{B}{1 + (\frac{D_p}{D_{TO}})^C}, \qquad (1)$$

where D_p represents dry particle diameter (nm), B, C and D_{50} are the three fitting parameters which represent the asymptote, the slope, and the inflection point of the sigmoid, respectively (Moore et al., 2010). The D_{50} is called the critical diameter, where 50% of the particles are activated at a specific ss. A hygroscopicity parameter κ which represents the CCN activity was calculated from the critical saturation ratio (Sc) and D₅₀ from the following equation (Petters and Kreidenweis, 2007):

222
$$\kappa = \frac{4A^3}{27D_{50}^3(\ln Sc)^2}$$
, $A = \frac{4\sigma_{s/a}M_W}{RT\rho_W}$, (2)

where ρ_w is density of pure water (about 997.04 kg m⁻³ at 298.15K), M_w is molecular weight of water (0.018 kg mol⁻¹), $\sigma_{s/a}$ is surface tension of the solution/air interface which is assumed to be value of pure water ($\sigma_{s/a} = 0.0728$ N m⁻¹ at 298.15K), R is the universal gas constant (8.314 J mol⁻¹ K⁻¹), T is thermodynamic temperature in Kelvin (298.15K), and D₅₀ is the critical diameter (in meter).

228 3 Results and Discussion

229 3.1 Overview

230 Figure 2 shows number size distribution (a), mass concentration and fraction (b and c), number 231 concentration of CCN (d), and hygroscopicity parameter (e) measured by different instruments during 232 the campaign. The particle sizes were predominantly larger than 10 nm, implying that no new particle 233 formation events were observed during the campaign. Furthermore, the distribution exhibited mainly 234 unimodal characteristics which peaked at a size range of about 60-80 nm. The average number 235 concentration was about 3400 cm⁻³, which was in general lower than that in inland PRD region (Cai et 236 al., 2017) and slightly lower than the ship measurement (4335 cm⁻³) over the East China Sea (Kim et al., 237 2009). However, two relative polluted periods were classified with high particle number concentrations at the beginning (6th-8th August, defined as P1 with a particle size peaking at about 80 nm) and at the end 238 239 (25th-26th August, defined as P2 peaking at about 100 nm) of the campaign. In contrast, two relatively clean periods were identified in between (9th-10th August, defined as C1 and 19th-21st August, defined as 240 241 C2).

242	Temporal profile of the mass concentration (Fig. 2a) measured by ToF-ACSM was consistent with
243	that of PNSD, which showed the highest concentration on 25th August. The total measured mass
244	concentration of NR-PM ₁ varied dramatically from 0.92 to 85.08 μ g m ⁻³ , with a median of 7.97 μ g m ⁻³ .
245	Mass concentrations of $PM_{2.5}$ were reported over the same region during Cruise I (27.6 $\mu g\ m^{\text{-}3})$ and
246	Cruise II (10.10 μ g m ⁻³) in Zhang et al. (2007). The mass concentration in our measurements was higher
247	than that in clean marine atmosphere (from 0.27 to 1.05 μ g m ⁻³) reported at the coastal station, Ireland
248	(Ovadnevaite et al., 2014) and the atmosphere over the Atlantic Ocean (Huang et al., 2018). Mass
249	concentration of SO_4^{2-} varied from 0.35 to 33.20 µg m ⁻³ , with a median of 3.66 µg m ⁻³ , which falls in a
250	range of previous measurement in Dongsha Islands (1.3 to 5.5 μ g m ⁻³ , Chuang et al., 2013). The average
251	mass fraction of NR-PM1 during the campaign was dominated by sulfate (46%), followed by organics
252	(35%), ammonium (14%), nitrate (3%), and chloride (2%), which was similar to the measurement over
253	the Atlantic Ocean (Huang et al., 2018). The chemical composition over northern SCS was quite different
254	from that at the urban site which was dominated by organics largely from anthropogenic sources (Cai et
255	al., 2017). A higher mass fraction of sulfate in the marine atmosphere may probably be attributed to
256	anthropogenic emissions (such as nearby ship emissions) rather than oxidation of dimethyl sulfide (DMS)
257	emitted from the ocean. The oxidation of DMS leads to formation of sulfur dioxide and methansulfonic
258	acid (MSA) both of which can be further oxidized to produce non-sea-salt (NSS) sulfate in marine
259	atmosphere. Oxidation of SO ₂ from ship emissions or inland transport can also be a major source of NSS
260	sulfate (Savoie et al., 2002). As an intermediate between DMS and sulfate, MSA in principle can be
261	detected by ToF-ACSM, although resolution of the instrument is low. Preliminary results show that the
262	fraction of sulfate from DMS oxidation is far below that from ship emissions. An early study showed
263	that anthropogenic sulfate accounted for about 81-97% of NSS sulfate over China Sea (Gao et al., 1996).

264	A ratio of 15-655 NSS sulfate to MSA in PM2.5 was reported in the northern South China Sea (Zhang et
265	al., 2007), much higher than that (18-20) in the remote marine (Savoie et al., 2002). Here we employed
266	the Modern-Era Retrospective analysis for Research and Applications, Version 2 (MERRA-2) to analyze
267	the distribution of ratio of sulfate to MSA at 925 hPa during the measurement period (GMAO, 2015).
268	The results were shown in Fig. S2 and the ratio ranged from 100 to 10000 over the SCS, much higher
269	than that in the remote Pacific Ocean (1-50). In addition, it also increases with latitude, indicating that
270	the anthropogenic emission is likely the major source of the total sulfate in the northern SCS region.
271	The number concentrations of CCN (N _{CCN} at ss=0.18%, 0.34%, and 0.59%) and total particles (N _{CN})
272	were shown in Fig. 2d. The N_{CN} values during the two polluted periods (P1 and P2) were significantly
273	higher than the average N_{CN} (3463 cm ⁻³) over the whole campaign period and those from other marine
274	measurements (Cai et al., 2017; Kim et al., 2009). This average value falls between the smoke type (2280
275	cm ⁻³) and the port type (4890 cm ⁻³) measured over the remote South China Sea (Atwood et al., 2017).
276	Note that since the abnormally spiked signals which were probably caused by emissions of the nearby
277	ships or the ship itself were removed in the data processes, the high $N_{\rm CN}$ values during those episodes
278	were likely attributed to regional pollution or long range transport from continents. For consistency, we
279	removed spikes likely associated with smoking, emissions from the ship itself and other adjacent ships
280	and cooking from further data analysis, including either abrupt high number concentrations of particles
281	(measured by SMPS), organics (measured by ToF-ACSM), and NO_{X} (measured by the NO_{X} monitor)
282	(Detailed criteria can be referred to descriptions and Fig. S1 in supplementary). In general, the $N_{\rm CCN}$
283	values at the three supersaturations increased with increase of the N_{CN} . The average value of N_{CCN} (1544
284	cm ⁻³ , ss= 0.34%) was similar with the simulated value (1000-2000 cm ⁻³ , ss= 0.4%), suggesting the model
285	simulation could satisfactorily predict the N_{CCN} in this region (Yu and Luo, 2009). Although the N_{CCN}

286	and $N_{\mbox{CN}}$ were relatively higher in P1 and P2 than the average value, they remained overall low during
287	the campaign compared to those from the inland PRD sites. The N_{CCN} values in P1 were lower than those
288	in P2 with similar values of $N_{\mbox{\tiny CN}}$ in both P1 and P2, suggesting a lower activation fraction in P1 than in
289	P2, which cloud be attributed to relatively high fractions of smaller particles and a lower hygroscopcity
290	in P1. As discussed above, particles peaked at a smaller size in P1, leading to fewer particles larger than
291	$D_{50}.$ The time series of the κ values calculated using Eq. 2 show that the aerosol hygroscopicity was lower
292	at the beginning of the campaign, leading to a lower CCN activity in P1. The measurements could be
293	affected by local fresh emissions with lower hygroscopic particles in urban since the ship was anchored
294	near Huizhou port and Hong Kong during P1, similar to lower hygroscopicity for urban particles
295	previously measured by Cai et al. (2017). Furthermore, low particle hygroscopicity was found from 11 th
296	August to 15 th August when the ship was sheltered at the port from the tropical storm Bebinca.
297	Aerosol hygroscopicity, an important parameter affecting CCN activity, can vary largely in its
297 298	Aerosol hygroscopicity, an important parameter affecting CCN activity, can vary largely in its values under different environments due to a variety of particle sources (Adam et al., 2012; Liu et al.,
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298 299	values under different environments due to a variety of particle sources (Adam et al., 2012; Liu et al., 2014; Hong et al., 2014; Wu et al., 2013; Cai et al., 2017). Comparison of the hygroscopicity parameter
298 299 300	values under different environments due to a variety of particle sources (Adam et al., 2012; Liu et al., 2014; Hong et al., 2014; Wu et al., 2013; Cai et al., 2017). Comparison of the hygroscopicity parameter κ obtained from this study, urban Guangzhou, remote marine Okinawa, remote South China Sea, and
298 299 300 301	values under different environments due to a variety of particle sources (Adam et al., 2012; Liu et al., 2014; Hong et al., 2014; Wu et al., 2013; Cai et al., 2017). Comparison of the hygroscopicity parameter κ obtained from this study, urban Guangzhou, remote marine Okinawa, remote South China Sea, and mountain Goldlauter was shown in Fig. 3. The κ_{median} values obtained from this study (around 0.4) fall
298 299 300 301 302	values under different environments due to a variety of particle sources (Adam et al., 2012; Liu et al., 2014; Hong et al., 2014; Wu et al., 2013; Cai et al., 2017). Comparison of the hygroscopicity parameter κ obtained from this study, urban Guangzhou, remote marine Okinawa, remote South China Sea, and mountain Goldlauter was shown in Fig. 3. The κ_{median} values obtained from this study (around 0.4) fall between those at the continental sites (Guangzhou and Goldlauter) and remote marine measurement
298 299 300 301 302 303	values under different environments due to a variety of particle sources (Adam et al., 2012; Liu et al., 2014; Hong et al., 2014; Wu et al., 2013; Cai et al., 2017). Comparison of the hygroscopicity parameter κ obtained from this study, urban Guangzhou, remote marine Okinawa, remote South China Sea, and mountain Goldlauter was shown in Fig. 3. The κ_{median} values obtained from this study (around 0.4) fall between those at the continental sites (Guangzhou and Goldlauter) and remote marine measurement (remote South China Sea and Okinawa) and are barely dependent on particle sizes whose pattern is quite
298 299 300 301 302 303 304	values under different environments due to a variety of particle sources (Adam et al., 2012; Liu et al., 2014; Hong et al., 2014; Wu et al., 2013; Cai et al., 2017). Comparison of the hygroscopicity parameter κ obtained from this study, urban Guangzhou, remote marine Okinawa, remote South China Sea, and mountain Goldlauter was shown in Fig. 3. The κ_{median} values obtained from this study (around 0.4) fall between those at the continental sites (Guangzhou and Goldlauter) and remote marine measurement (remote South China Sea and Okinawa) and are barely dependent on particle sizes whose pattern is quite similar to those in Okinawa. Moreover, a κ value was respectively reported to be in a range of 0.22-0.65

308	Pacific and Southern Oceans (Berg et al., 1998). In contrast to maritime environments (i.e., SCS and
309	Okinawa), the κ_{median} values in Guangzhou (0.21-0.31) are much lower and increase obviously with
310	particle sizes. The low hygroscopicity for small particles in Guangzhou was attributed to local emissions
311	from traffic and industry (Cai et al., 2017). The cruise in this campaign is in an offshore region where
312	the air is affected by anthropogenic emissions from the adjacent inland PRD region, leading to medium
313	values of aerosol hygroscopicity between urban and marine background regions.

3.2 Temporal and spatial distributions

316	As discussed above, the air over the offshore northern SCS is affected by local emissions from
317	inland PRD regions. The shoreline along Huizhou port is roughly 45° inclined to the latitude (from South
318	to North) and it is reasonable to assume that the concentrations of the air pollutants originating from local
319	emissions are generally dependent on the distance offshore which can be roughly represented by the
320	latitude in this study. Hence in this section, the temporal and spatial concentration distributions of air
321	pollutants (particles and gases) were presented with latitude and the dates were color-coded, representing
322	from the beginning (dark blue) to the end (dark red) of the cruise (Fig. 4). The concentrations of trace
323	gases (O ₃ , CO, and NO _x), N_{CN} , and N_{CCN} (ss=0.34%) were higher during the late half than during early
324	half of the campaign, while SO ₂ concentration varied in an opposite way, suggesting that the sources of
325	the air pollutants or the air masses were different at the beginning and at the ending of the campaign. In
326	particular, the aforementioned quantities increased substantially with latitude (the higher the latitude the
327	closer to the shore) from 19th to 26th August, indicating that the air masses from inland China could affect
328	the northern SCS region during this period. However, the $N_{CCN}/N_{CN,tot}$ and κ values (ss=0.34%) showed

329	almost no pattern (Figs. 4g and 4h), except that the $N_{CCN}/N_{CN,tot}$ values were both high (about 0.8) at the
330	beginning and at the end of the cruise. The $N_{\text{CCN}}/N_{\text{CN,tot}}$ was defined as the ratio of number concentration
331	of cloud condensation nuclei and total aerosol particles at a specific ss. The κ values were observed to be
332	relatively low when the vessel located at a latitude of about 22°N corresponding to 6th and 26th August,
333	suggesting that the air was affected by local fresh emissions which increased the organic content of the
334	particles. Interestingly, a higher value on 26 th August than on 6 th August was clearly shown (Fig. 4g) due
335	probably to larger averaged particle sizes on 26th August (about 110 nm) which were more easily
336	activated than smaller particles on 6 th August (about 60-90 nm).
337	To further investigate the effects of local emissions on aerosol particles over northern SCS, the
338	correlations of SO ₂ , CO, NOx concentration, N_{CCN} , $N_{CCN}/N_{CN,tot}$, κ with N_{CN} were explored (Fig. 5). The
339	variation of SO ₂ concentration was independent of N_{CN} , suggesting that SO ₂ did not share the same source
340	with particles. The CO concentration is positively correlated with $N_{\rm CN}$ during the second half of the cruise,
341	while no obvious correlation is observed during the first half, implying that sources of particles could be
342	different during the two periods. The correlation during the second half of the cruise indicates that the
343	particles might share the same source with CO which was attributed to biomass burning or anthropogenic
344	emissions. An excellent correlation between NO_X concentration and N_{CN} was shown in all ranges of
345	particle number concentrations, implying that the aerosol particles might originate from the same source
346	as NO_{X} which was likely attributed to traffic and industry in the continental PRD region. The $\mathrm{N}_{\mathrm{CCN}}$ was
347	observed to follow two distinct trends for the first and second half of the cruise which show in general a
348	higher activation efficiency during the second half of the campaign, especially when N_{CN} is greater than
349	about 7000 cm ⁻³ , further validated by a much higher $N_{CCN}/N_{CN,tot}$ ratio against N_{CN} as shown in Fig. 5e.
350	As discussed in the previous paragraph, distinct κ values were seen at the very beginning and at the end

351 of the campaign, suggesting that the properties and sources of the particles could be different as will be 352 further discussed in the case study below.

353

354 3.3 Case Study

355 In section 3.1, we classified four periods (all in August) based upon particle number concentration, corresponding to P1 (6th to 8th), C1 (9th to 10th), C2 (19th to 21st), and P2 (25th to 26th) as shown in Fig. 6. 356 357 During the two clean periods (C1, before Bebinca; C2, after Bebinca), the vessel travelled around 358 northeast of Dongsha islands where the particle number concentrations remained relatively low which 359 were not affected by the continental emissions from the PRD region. However, high number 360 concentrations of particles were observed during P1 when the vessel was close to the shore where the air 361 was substantially affected by local emissions from either Hongkong or Huizhou. During the last two days 362 in P2, even higher particle number concentrations were observed, suggesting that the pollutants might 363 originate from inland continent via long range transport. 364

We performed HYSPLIT to investigate the source origins of the air pollutants according to 365 movement of air masses during the campaign (Fig. 7). The backward trajectories during P1 showed that 366 the air masses were mainly from east and south and when arriving at the location of the vessel, the air 367 masses were stagnant on the shore, suggesting that the pollutants might originate from local emissions. Interestingly, particle number concentrations were low during 11th to 15th August when the vessel was 368 369 sheltered from Bebinca, due probably to the arrival of the typhoon which caused high wind speeds and 370 brought rainfall in the northern SCS, resulting in removal of air pollutants in Huizhou and in Hong Kong. 371 The air masses over northern SCS originated from southwest (C1) or from Indo-China Peninsula (C2) 372 due to summer monsoon during the two clean periods (Fig. 7). The air masses moved northerly during P2 and brought high concentrations of particles from inland China to PRD region, and then further to the
northern SCS (Fig. 7).

375	Chemical speciation by ToF-ACSM showed that the mass fractions of aerosol composition were
376	substantially different during C1, C2, and P2, except for nitrate whose fraction remain almost constant
377	among the above three periods (Fig. 8). Note that the mass fraction during P1 was not available for
378	comparison due to instrumental failure. Even the mass fractions during the two clean periods were
379	distinctly different, in particular, those of organics (26% for C1 vs 40% for C2), ammonium (19% for C1
380	vs 12% for C2), and chloride (7% for C1 vs 2% for C2), although the particle composition was dominated
381	by sulfate which was almost equal in mass fraction (44% for C1 vs 42% for C2). The mass fraction
382	during C1 was dominated by sulfate, followed by organics, ammonium which was similar to that in
383	remote marine region (Cai et al., 2017). The mass fraction of sulfate in the NR-PM1 during C1 and C2
384	was also similar to the previous study (44% and 43% in $PM_{2.5}$ for Cruise I and II, respectively) over the
385	northern SCS (Zhang et al., 2007). Although the mass fraction was still dominated by sulfate, a
386	substantially increasing fraction of organic (increase of 26% for C1 to 40% for C2) was observed. This
387	increase in organic fraction was likely attributed to the air masses passing through Indo-China Peninsula
388	which brought significant local sources. In contrast to the clean periods, the mass fraction in the NR-PM1
389	during P2 was dominated by organics (47%), followed by sulfate (33%) and ammonium (13%), similar
390	to that in urban areas (Huang et al., 2014), indicating that air masses from the north could bring
391	continental particles in inland China to the northern SCS.

The particle number size distribution (PNSD) was measured by the custom-made SMPS which was described in the methodology section. The average particle number concentrations during P1 and P2 (9239 and 10088 cm⁻³ respectively) were much higher than those during the clean periods (1826 and

395	1683 cm ⁻³ for C1 and C2 respectively). In addition, the PNSD during the pollution periods was
396	characterized by an obvious accumulation mode that was attributed to secondary aerosols (Fig. 9), while
397	the one during the clean periods has a smaller and a less obvious accumulation mode and a more obvious
398	Aitken mode which was more related to marine background particles (Cai et al., 2017; Atwood et al.,
399	2017; Kim et al., 2009). The median diameters and concentration of the accumulation mode during C1
400	and C2 was similar to those previously reported in South China Sea (Reid et al., 2015). Note that the
401	fitted nucleation modes for both clean and pollution periods were barely seen due to the obviously low
402	concentrations of particles in this mode. The lognormal median diameters for the Aitken mode (70.4 nm)
403	and the accumulation mode (165.7 nm) during P2 were respectively larger than those (48.6 nm and 143.1
404	nm) during P1, implying more aging processes and particle growth in the long range transport from the
405	inland continent. Furthermore, a wider accumulation mode during C2 than during C1 was observed,
406	implying more complex sources for larger size particles which could probably be attributed to biomass
407	burning or anthropogenic activities across Indo-China Peninsula. The backward trajectories during C2
408	pass through the burning regions in Southeast Asia (e.g., Viet Nam, Laos, Cambodia etc.), also supporting
409	this conjecture. However, more solid evidences are needed since the observation of biomass burning
410	tracers (such as K and levoglucosan) is missing in this campaign.
411	The CCN activity parameters (average N_{CCN} , D_{50} , and $N_{CCN}/N_{CN,tot}$ at ss=0.18%, 0.34%, and 0.59%)
44.0	

7139 cm⁻³, much higher than the simulated annual mean values in northern SCS region (1000-2000
cm⁻³, ss=0.4%, Yu and Luo, 2009). It implies that the continental emissions could have significant impact

during each period were summarized in Table 2. The N_{CCN} (ss=0.34%) during P1 and P2 were 3969 and

- 415 on the CCN concentration over this region. Although the mass fractions of chemical composition for C1,
- 416 C2, and P2 were quite different among those periods, no significant differences of the hygroscopicity

417	parameter κ values were seen, indicating particles with a size range of 30-120 nm were less affected by
418	long range transport from Indo-China Peninsula or inland China continent. The calculated median κ
419	values based on the measured D_{50} ranged from 0.32 to 0.41 and no significant differences in diameters
420	and periods were observed (Fig. S3), suggesting that the high mass fractions of organics during C2 might
421	be distributed in larger particle sizes (Fig. 8). The D_{50} values during P2 were smaller at all supersaturation
422	ratios, suggesting higher hygroscopicity and CCN activity during this period. In addition, the $N_{CCN}/N_{CN,tot}$
423	and N_{CCN} during P2 was larger than during P1, owing to a larger number fraction of accumulation mode
424	and a higher hygroscopicity. Meanwhile, the median κ values fell in a range of 0.12-0.19 during P1,
425	significantly lower than those during three other periods but similar to the values measured in urban cities
426	(Tan et al., 2013; Jiang et al., 2016; Cai et al., 2018). Such lower values of hygroscopicity were probably
427	contributed from local emissions originating from inland urban cities or heavy duty ships. More cruise
428	campaigns are hence needed to identify the source origins of marine aerosols over the SCS region.
428 429	campaigns are hence needed to identify the source origins of marine aerosols over the SCS region. The mixing state and heterogeneity of particles can affect the steepness of the activation curves (Cai
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429 430 431 432	The mixing state and heterogeneity of particles can affect the steepness of the activation curves (Cai et al., 2018). A steeper curve indicates that particles intend to be internally mixed and have a higher similarity in hygroscopicity. The average activation curves at 0.18% ss during the P1, C1, C2 and P2 periods are shown in Fig. S4. The parameter C (in Eq. 1) can be used to present the steepness of activation
429 430 431 432 433	The mixing state and heterogeneity of particles can affect the steepness of the activation curves (Cai et al., 2018). A steeper curve indicates that particles intend to be internally mixed and have a higher similarity in hygroscopicity. The average activation curves at 0.18% ss during the P1, C1, C2 and P2 periods are shown in Fig. S4. The parameter C (in Eq. 1) can be used to present the steepness of activation curve. A small C value indicates a steep activation curve. The C values during P1, C1, C2 and P2 periods
 429 430 431 432 433 434 	The mixing state and heterogeneity of particles can affect the steepness of the activation curves (Cai et al., 2018). A steeper curve indicates that particles intend to be internally mixed and have a higher similarity in hygroscopicity. The average activation curves at 0.18% ss during the P1, C1, C2 and P2 periods are shown in Fig. S4. The parameter C (in Eq. 1) can be used to present the steepness of activation curve. A small C value indicates a steep activation curve. The C values during P1, C1, C2 and P2 periods were -8.5, -14.3, -13.7 and -10.6, respectively. The smooth curve and the largest C value during P1
 429 430 431 432 433 434 435 	The mixing state and heterogeneity of particles can affect the steepness of the activation curves (Cai et al., 2018). A steeper curve indicates that particles intend to be internally mixed and have a higher similarity in hygroscopicity. The average activation curves at 0.18% ss during the P1, C1, C2 and P2 periods are shown in Fig. S4. The parameter C (in Eq. 1) can be used to present the steepness of activation curve. A small C value indicates a steep activation curve. The C values during P1, C1, C2 and P2 periods were -8.5, -14.3, -13.7 and -10.6, respectively. The smooth curve and the largest C value during P1 suggest that particles had a higher degree of external mixing and higher heterogeneity, owing to the local

emissions. The activation curve during P2 period was smoother than C1 and C2 but steeper than P1,
indicating that the particles during this period could be a mixture of aged particles from China inland and
fresh particles from onshore emissions.

442

443 4 Conclusions

444 As an annual routine exercise for SCS expedition during summertime, the 2018 cruise campaign 445 organized by Sun Yat-sen University is a comprehensive and interdisciplinary field measurement 446 involving atmosphere, ocean, geology, biology, and chemistry etc. The measurement includes stationary 447 and navigating observations based on compromise among multiple disciplines. For atmospheric 448 measurements, several key scientific questions are emerging to be addressed over SCS region, including 449 sources of air pollutants (gases and particles) in marine atmosphere, impacts of biomass burning from 450 southeastern Asia and summer monsoon on atmospheric chemistry and physics in SCS region. In this 451 study, the CCN activity, chemical composition, and particle number size distribution over northern SCS 452 were measured using several onboard instruments including a ToF-ACSM, a CCNc, a SMPS, several 453 monitors for trace gases (i.e., SO₂, NO_X, CO, and O₃). On one hand, lower concentrations of key trace 454 gas pollutants and particle number or mass were observed in atmosphere of SCS than those in urban areas in PRD region, consistent with previously reported values for background marine atmosphere. 455 456 Overall, chemical composition of NR-PM₁ was dominated by sulfate (46%) and the PNSD showed 457 unimodal distribution centering at about 60-80 nm and the hygroscopicity κ values being higher than 458 those in urban areas. On the other hand, characteristics of air pollutants (e.g., concentrations, physical 459 and chemical properties) show substantially variations during summer monsoon season, depending on

460 source origins. Characteristics similar to continental aerosols were shown when air masses originate from 461 inland China continent or Indo-China peninsula possibly via long range transport, leading to increase of 462 organic fraction in chemical composition and decrease of hygroscopicity which might be attributed to 463 picking up locally emitted and fresh pollutants during transport. Furthermore, low hygroscopicity κ values were shown when the air was affected by local fresh emissions and in this case the number 464 465 concentration of particles increased with decrease of offshore distance. In addition, concentrations of 466 both NO_X and CCN concentrations were well correlated with the total concentration of particles. Interestingly, a tropical storm Bebinca was caught in the middle of the campaign, resulting in two 467 468 relatively clean periods (C1 and C2). These clean periods were likely attributed to strong wind and 469 rainfalls brought by the storm which could obviously blow away or wash out pollutants in northern SCS 470 region.

471 Our results suggest that aerosol properties and trace gases concentration over northern SCS is 472 complex and substantially variable. The median hygroscopicity k values of the particles in northern SCS 473 were measured to be about 0.4, in the range of between those in the remote northwestern Pacific Ocean 474 and those in urban PRD region, implying that particles in northern SCS could be a mixture of marine 475 background and anthropogenic particles from continents (e.g., Indo-China peninsula and inland China continent). Concentrations of aerosol particles and trace gases exhibit complex temporal and spatial 476 477 distribution. Concentrations of trace gases (i.e., O₃, CO, and NO_X except SO₂), particles (i.e., N_{CN} and 478 N_{CCN}) were higher at the beginning (pollution episode: P1) than at the end (pollution episode: P2) of the 479 campaign, implying different source origins for the two periods. At the beginning of the campaign, the 480 air was likely affected by local fresh emissions from Huizhou, leading to increase of concentrations of 481 both measured trace gases (except SO₂) and particles with decrease of offshore distance. Meanwhile,

482	concentration of NO_{X} had a good correlation with the N_{CN} , suggesting they might originate from the
483	same sources. Similarly, at the end of the campaign, concentrations of both measured trace gases (except
484	SO ₂) and particles also increased with decrease of offshore distance, while because of more larger
485	particles, higher fractions of particles were activated at the end than at the beginning of the campaign.
486	We attributed the source origin during this period to inland China content via long range transport with
487	additional local fresh pollutants during transport process, leading to barely clear patterns for both
488	$N_{CCN}/N_{CN,tot}$ and D_{50} at all applied ss (ss=0.18, 0.34, and 0.59%). Furthermore, our results indicate that
489	biomass burning from southeastern Asia may have important impacts on chemical composition and
490	properties of aerosol particles over northern SCS, in particular, leading to increase of organic mass
491	fractions and decrease of hygroscopicity $\boldsymbol{\kappa}$ values and hence affecting CCN activity in the region. Our
492	study highlights the necessity for performing more intensive ship-based atmospheric measurements in
493	order to better understand marine aerosols and air pollution in SCS region.
494	

Data availability. Data from the ship-based cruise measurements are available upon request (Jun Zhao
496 via <u>zhaojun23@mail.sysu.edu.cn</u>).

Supplement. The supplement related to this article is available online at xxx.

500 Author contributions. MC, JZ, and HT designed the research. MC and BL performed the ship-based

- 501 cruise measurements. XC performed sulfate/MSA analysis. MC, JZ, HT, BL, and QS analyzed the data.
- 502 MC, JZ, and HT wrote the paper with contributions from all co-authors.

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

506	Acknowledgements. We acknowledges support from National Key Project of MOST (2017YFC0209502,
507	2016YFC0201901, 2016YFC2003305), National Natural Science Foundation of China (NSFC)
508	(91644225, 21577177, 41775117), Science and Technology Innovation Committee of Guangzhou
509	(201803030010), the "111 plan" Project of China (Grant B17049), Scientific and Technological
510	Innovation Team Project of Guangzhou Joint Research Center of Atmospheric Sciences, China
511	Meteorological Administration (Grant No.201704). Additional support from the crew of the vessel and
512	from Southern Marine Science and Engineering Guangdong Laboratory (Zhuhai) is greatly
513	acknowledged.

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- 645

-6	Table 1. Summary of the instruments	s used in the campaign.
	Instruments	Parameters
	ToF-ACSM	NR-PM ₁
	SMPS+CCNc	PNSD (9-415 nm), Size-resolved CCN
		Activation Ratio (at ss=0.18%, 0.34%, and 0.59%)
	CO Monitor	CO concentration
	SO ₂ Monitor	SO ₂ concentration
	O ₃ Monitor	O ₃ concentration
	NO _X Monitor	NO _X , NO, NO ₂ concentration

Table 1. Summary of the instruments used in the campaigr

 $\label{eq:ccn} \textbf{Table 2. Summary of average } N_{CCN}, D_{50}, \text{ and } N_{CCN}/N_{CN, tot} \text{ at } 0.18\%, 0.34\%, \text{ and } 0.59\% \text{ ss during P1},$

Period	SS	0.18%	0.34%	0.59%
P1	N _{CCN} (# cm ⁻³)	1825	3969	7198
	D ₅₀ (nm)	132	96	65
	N _{CCN} /N _{CN,tot}	0.19	0.34	0.49
C1	N _{CCN} (# cm ⁻³)	566	978	1330
	D ₅₀ (nm)	105	67	49
	N _{CCN} /N _{CN,tot}	0.31	0.54	0.71
C2	N _{CCN} (# cm ⁻³)	536	844	1183
	D ₅₀ (nm)	108	68	48
	N _{CCN} /N _{CN,tot}	0.32	0.55	0.73
P2	N _{CCN} (# cm ⁻³)	4969	7140	8679
	D ₅₀ (nm)	101	65	49
	N _{CCN} /N _{CN,tot}	0.49	0.74	0.85

650 C1, C2, and P2.

653 FIGURE CAPTIONS

Figure 1. Ship track and tropical storm Bebinca track during the campaign (a), and schematic diagram of

the vessel showing the location of the sea container which housed the onboard instruments during the

656 campaign (b).

- 657 Figure 2. Temporal profiles of the meausred particle number size distribution (a), mass concentration (b)
- and mass fraction (c) of chemical composition, N_{CCN} and N_{CN} (d) and the daily averaged κ values with
- the upper and lower error bars (e). No data were shown between 6th and 8th August due to the instrumental
- failure of the TOF-ACSM.

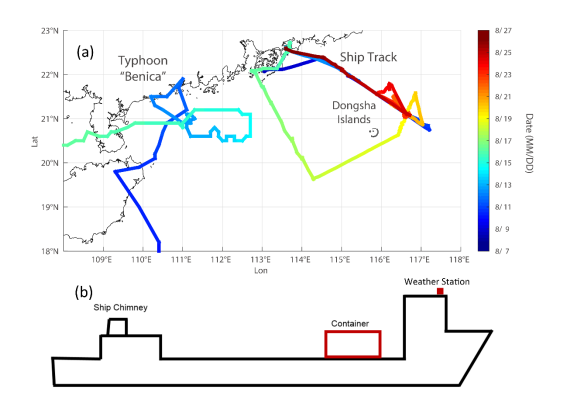
- site, at marine background Okinawa site, and the mean and standard deviation κ values measured over
- 663 remote South China Sea and at mountain Goldlauter site. The κ values over South China Sea were
- obtained from CCNc measurements (ss=0.18%, 0.34%, and 0.59%, in blue). The κ values in urban
- Guangzhou were obtained from CCNc (ss=0.1%, 0.2%, 0.4%, and 0.7%, in orange) and HTDMA
- 666 measurements (in purple). The κ values in marine region Okinawa were obtained from HTDMA
- 667 measurements (in green). The κ values in remote South China Sea were obtained from CCNc (ss=0.14%
- and 0.38%, in orange). The κ values in mountain Goldlauter site were obtained from CCNc (ss=0.07%,
- 669 0.10%, 0.19% and 0.38%, in black).
- 670 Figure 4. Concentrations of SO₂ (a), O₃ (b), CO(c), NO_X (d), N_{CN} (e), N_{CCN} (f), N_{CCN}/N_{CN,tot} at 0.34% ss
- 671 (g), and κ at 0.34% ss (h) as a function of latitude. The data points were color-coded according to date.
- Figure 5. Correlations of SO₂(a), CO(b), NO_X (c), N_{CCN} (d), AR at 0.34% ss (e), and κ at 0.34% ss (f)
- 673 with N_{CN}. The data were plotted according to color-coded dates.
- Figure 6. The ship track during P1, C1, C2 and P2 periods.

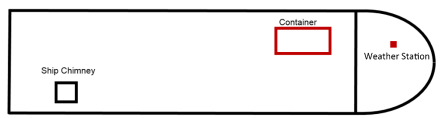
675	Figure 7. The	72 h backward	trajectories	arriving at	the l	ocation of the	vessel v	vith three l	neights (150 m.
		,								

500 m, and 1000 m) during P1, C1, C2, and P2, respectively. The dots represent the fire spots detected

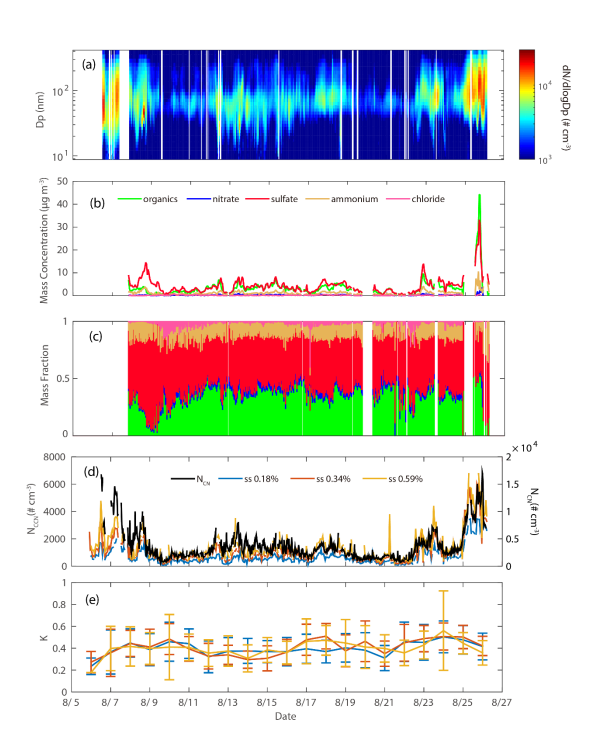
by MODIS.

- Figure 8. The average mass fraction of NR-PM₁ composition during the C1, C2 and P2 periods.
- Figure 9. The average and standard deviation (shaded area) PNSD, along with trimodal lognormal fitted
- 680 modes (dash color lines). The average N_{CN} during each period and the median size of each lognormal fit
- 681 were shown.

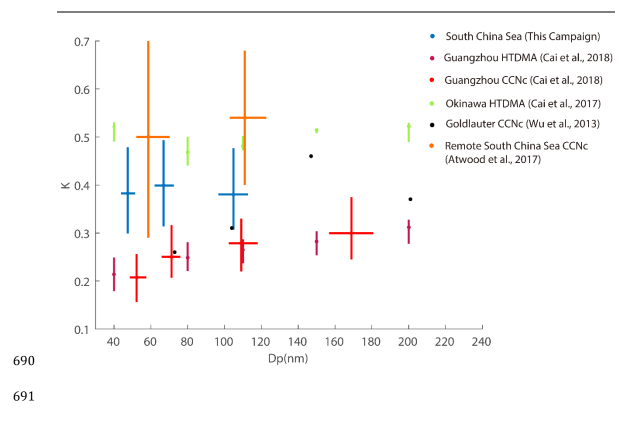




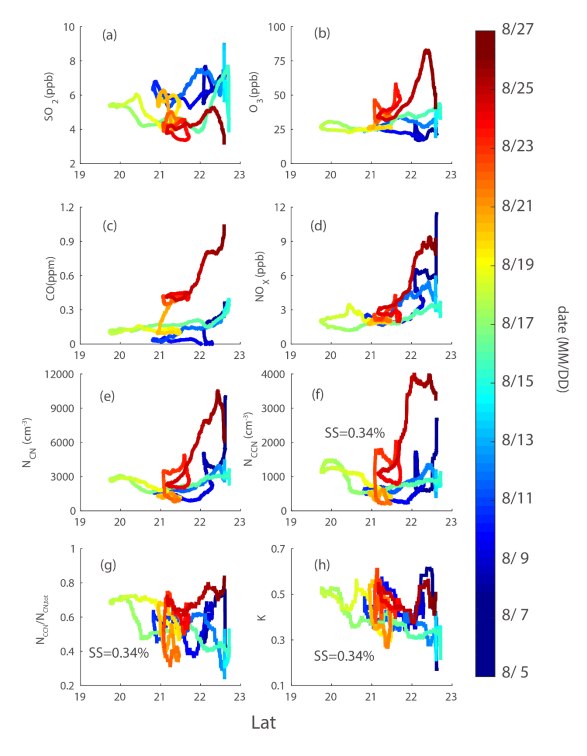
684 Fig. 1.



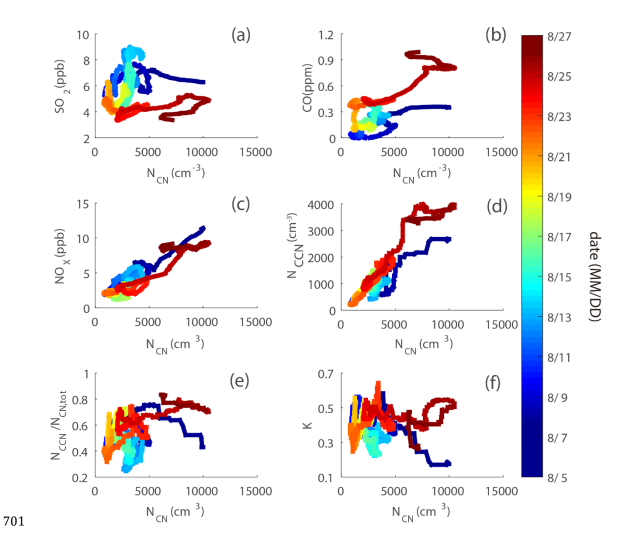
688 Fig. 2.



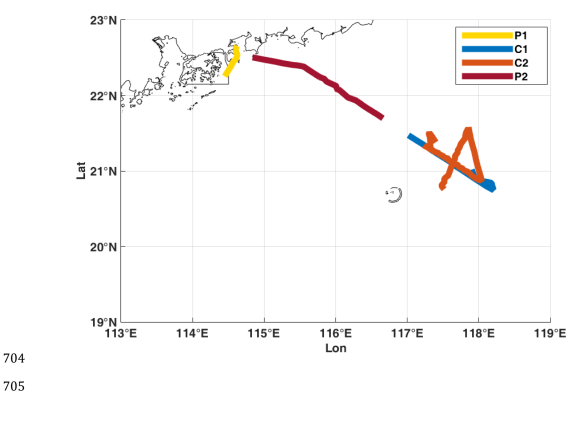
- 692 Fig. 3.



696 Fig. 4.







706 Fig. 6.

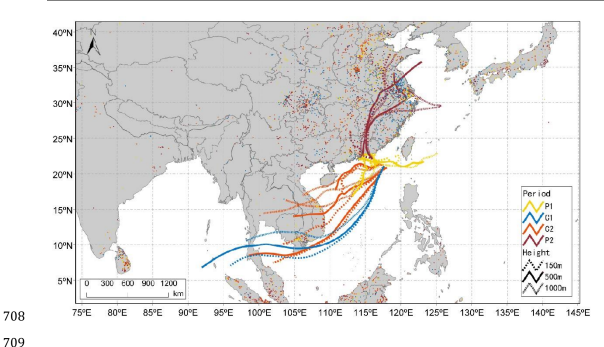
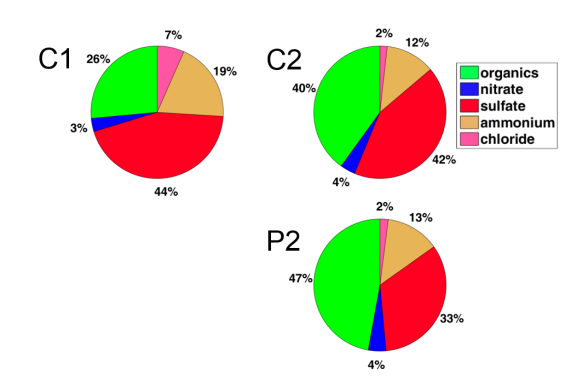
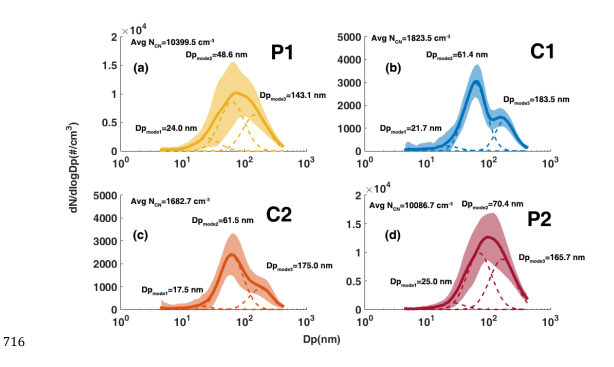




Fig. 7.



- 714 Fig. 8.





718 Fig. 9.