Comments to the Author:

The authors have responded most of the referee comments. However, when looking at the revised paper, there are still a few issues that should be considered before accepting this paper for publication. They are listed below.

We thank the editor for carefully reviewing the manuscript, and all the comments have been addressed.

In the introduction the authors discuss marine aerosols in very general terms, but then mention emission trends that are not true in general in the global atmosphere (lines 97-100). Please either mention explicitly here that you refer to Asian emissions, or revise the text according to what has taken place in different continents (In Europe and NA, for example, SO2 emissions have decreased for several decades and also NOx emissions have evolved very differently from those in Asia).

Response: The emission trend is mainly applied to China, and this has been added in the revised manuscript.

line 161: Surface tension is a composition-dependent variable, not a constant. It is usually set constant when applying the equation to calculate the hygroscopicity parameter cappa, but that is a different thing. Please correct the text.

Response: This has been revised as follows:

 $\sigma_{s/a}$ represents the surface tension over the interface of the solution and air with the value of 0.072 J m⁻² applied in this study

Lines 362-369: The minimum diameter of particles at SS of 0.2-1 % is somewhere in the range 50-100 nm, so statement about nanometer-size particles on lines 366-367 is very confusing. Why to talk about nanometer-size (sub-10 nm) particles here? Is this statement even correct? There is often more organics in sub-100 nm particle compared with accumulation mode particles, but the situation in the sub-10 nm size range may be totally different. Please be more careful here what size ranges you are really referring to here.

Response: This has been revised as follows:

In general, the fraction of organics in the nanometer particles increases with decreasing particle size from ~ 100 nm to ~ 50 nm

Section 3.6, end of Introduction and Figure 7: I would recommend using the terms "relationship" or "regression equation" rather than "correlation" here. The term "correlation equation" does not sound correct at all.

Response: This has been revised based on the reviewer's suggestion.

The text still contains many grammatical problems (lack of articles, wrong prepositions etc.). Please check out the text once more with the help of a native English speaker.

Response: We have asked a native speaker to check the grammar issues.

		Style Definition: Comment Text: Font: (Default) Tahoma, 8 pt
1	Variations in N_{cn} and N_{ccn} over $\frac{China}{China}$ marginal seas $\frac{in\ China}{China}$ related to marine	Style Definition: Comment Reference: Font: (Default) Tahoma, 8 pt
2	traffic emissions, new particle formation and aerosol aging-	Tanonia, o pt
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3	Yang Gao ^{1,2**} , Deqiang Zhang ^{1#} , Juntao Wang ¹ , Huiwang Gao ^{1,2} and Xiaohong Yao ^{1,2*}	
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10	*Correspondence to yanggao@ouc.edu.cn ; xhyao@ouc.edu.cn	
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Abstract_

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In this study, a cruise campaign was conducted over China marginal seas in China to measure the concentrations of condensation nuclei (Ncn), cloud condensation nuclei (N_{cen}) and other pollutants during from day of year (DOY) 110 to DOY 135 of 2018. With exhaustedly excluded The ship self-ship emission signals, were exhaustively excluded, and the mean values of N_{ccn} during the cruise campaign were found to slightly increased increase from $3.2 \pm 1.1 \times 10^3$ cm⁻³ (mean \pm standard) at supersaturation (SS) of 0.2% to $3.9 \pm 1.4 \times 10^3$ cm⁻³ at SS of 1.0%, and the mean value for N_{cn} was 8.1 ± 4.4 $\times 10^3$ cm⁻³. Data analysis showed that marine traffic emissions apparently vielded a large contribution argely contributed to the increase of In Ncn in the daytime, especially in the marine atmospheres over their heavily travelled traveled sea zones; however, the fresh sources hadmade no clear contribution to the increase of Nccn. This finding was supported by the quantitative relations between N_{cn} and N_{ccn} at SS=0.2-1.0% against the mixing ratios of SO₂ in the ship self-ship emission plumes, i.e., a 1 ppb increase in SO₂ corresponds corresponded to a 1.4×10⁴ cm⁻³ increase in N_{cn7} but only a 30-170 cm⁻³ ³ increase in N_{ccn} possibly because of abundant organics in the aerosols. The smooth Smooth growth of can be observed in the marine traffic-derived particles can be observed, reflecting aerosol aging. The estimated hygroscopicity parameter (κ) values were generally as high as 0.46-0.55 under the dominant onshore winds, suggesting that inorganic ammonium aerosols likely actingacted as the major contributor to N_{ccn} largely through aerosol aging processes largely decomposed of decomposing organics. Moreover, the influences of the <u>new</u> transported new particles from the continent on <u>the</u> N_{cn} and N_{ccn} in the marine atmosphere were also investigated.— Key words Keywords: Ncn; Nccn; marine traffic emissions; hygroscopicity parameter;

1. Introduction

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Oceans occupy approximately 2/3 of the Earth's surface, and water evaporation from oceans acts as theis a major source of moisture in the atmosphere. Aerosol-cloud interactions in marine atmospheres, covering ranging from tropics tropical to polar regions, thereby attracthave attracted great attentions attention in the past few decades due to their impact on-the climate change (Huebert et al., 2003; Yu and Luo, 2009; Quinn and Bates, 2011; Wang et al., 2014; Brooks and Thornton, 2018; Rosenfeld et al., 2019). However, large uncertainties still exist in various marine atmospheres, e.g., the sources of aerosols, the concentrations of bulk cloud condensation nuclei (CCN) and aerosol CCN activation under various of supersaturation, etcsupersaturations. (Clarke et al., 2006; Decesari et al., 2011; Quinn and Bates, 2011; Saliba et al., 2019; Rosenfeld et al., 2019). The These uncertainties are mainly determined by limited observations in marine atmospheres, although a few additional observations of the number concentrations of aerosola (Ncn) and CCN (Nccn) were recently reported in different marine atmospheres, e.g., over the Mediterranean (Bougiatioti et al., 2009), Sea of Japan (Yamashita et al., 2011), Bay of Bengal (Ramana and Devi, 2016), coast of California (Ruehl et al., 2009) and the Northwest Pacific Ocean (Wang et al., 2019), etc.).

Besides In addition to sea-spray aerosols and secondarily formed aerosols from seaderived gaseous precursors (O'Dowd et al., 1997; Clarke et al., 2006; Quinn and Bates, 2011; Blot et al., 2013; Fossum et al., 2018), marine traffics also emit atraffic emits large amountamounts of aerosols and reactive gases (Chen et al., 2017). These pollutants may also directly or indirectly contribute to CCN—therein, to some extent (Langley et al., 2010). In addition, the long-range transport of continental aerosols has been widely reportedly acted reported to act as an important source of CCN in marine atmospheres (Charlson et al., 1987; Huebert et al., 2003; Fu et al., 2017; Royalty et al., 2017; Sato and Suzuki, 2019; Wang et al., 2019). The continent-derived aerosol particles observed in marine atmospheres usually mix with different sources, such as biomass burning, dust and anthropogenic emissions (Feng et al., 2017; Lin et al., 2015; Guo et al., 2014; Guo et al., 2016). An appreciable fraction of organics reportedly exists in marine aerosols and continental aerosols upwind of oceans (O'Dowd et al., 2004; Feng et al., 2012; Quinn et al., 2015; Feng et al., 2016; Song et al., 2018; Ding et al., 2019). However, ammonium sulfate aerosols have been frequently reported to dominantly contribute to CCN-related aerosols in many marine atmospheres and lead to hygroscopicity parameter<u>parameters</u> (κ) larger than 0.5 (Mochida et al., 2010; Cai et al., 2017; Fu et al., 2017; Royalty et al., 2017; Phillips et al., 2018). A question is automatically naturally raised, i.e., where do particulate organics go in the marine aerosols enriched in ammonium sulfate? Anthropogenic emission emissions in China such as SO₂, and NO_x in general increase have generally increased since the 1980s, until and recently started to decrease, i.e., SO2 startstarted to decrease fromin 2006 (Li et al., 2017), whereas NOx started to decrease sincein 2011 (Li et al., 2017; Liu et al., 2016). Together with the influence of the Asian Monsoon, the marginal seas of China are, therefore, inevitably affected by the outflow of continental aerosols (Guo et al., 2016; Feng et al., 2017). Observations of N_{cn} and N_{ccn} in marine atmospheres over China marginal seas helpshelp to resolveaddress the data scarcity, understand their the sources and dynamic changes in these parameters and better service the study of their potential climate impacts.—

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In this study, cruise campaigns were conducted to measure the N_{cen}, N_{en}, particle number size distributions, gaseous pollutants and aerosol composition of water-soluble ionic species over the marginal seas from 20 April 2018 (day of year (DOY) 110) to 15 May 2018 (DOY 135), traveling from the East China seaSea to the South China sea,Sea and returning to the Yellow seaSea. Spatiotemporal variations in the N_{en}, N_{een} and CCN activities of the aerosol particles were studied. The *Kappa* values of the aerosol particles

from DOY 110 to DOY 118 over the marine <u>environments</u> were calculated and analyzed. Finally, we tried to establish <u>the correlations relationship</u> of N_{cn} and N_{ccn} with <u>the mixing</u> ratios of SO_2 in self-ship plumes and ambient marine air. The <u>correlation regression</u> equations are valuable for the estimation of N_{cn} and N_{ccn} from SO_2 when the direct observations of N_{cn} and N_{ccn} are not available.

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2. Experimental design

2.1 Instruments and data sources

A cruise campaign was conducted across China marginal seas in China from DOY 110 to DOY 135 of 2018 (Fig. 1a, b). A suite of instruments including a Fast Mobility Particle Sizer fast mobility particle sizer (FMPS, TSI Model 3091), CCN counter (CCNC, DMT Model 100), Condensation Particle Countercondensation particle counter (CPC, TSI Model 3775), gas analyzers, Ambient Ion Monitor Ionambient ion monitor-ion chromatography (AIM-IC), etc., were onboard athe commercial cargo ship Angiang 87 for measurements. The FMPS was used to measure the particle number size distributions with mobility diameters from 5.6 nm to 560 nm in 32 channels at 1second temporal resolution with an inlet flow of 10 L min⁻¹. The CPC was used to report the N_{cn} ranging from 4 nm (50% efficiency) to 3000 nm (N_{cn}) in 2-second time resolution with an inlet flow of 1.5 L min⁻¹. The N_{cn} was then used to calibrate the particle number size distributions simultaneously measured by the FMPS, on the basis of the procedure proposed by Zimmerman et al. (2015). Due to the severe oceanic conditions and humid weather conditions, the FMPS and CPC were out of service after DOY 118 and DOY 122, respectively. Prior to the campaign, the CCNC was calibrated with ammonium sulfate particles based on the standard procedure detailed at by Rose et al. (2008). The calibration curve wasis shown in Fig. S1. The total flow rate of CCNC was 0.45 L min⁻¹, with a ratio of sample to sheath at 1/10, and five super saturations (SS) conditions were selected, including 0.2-%, 0.4-%, 0.6-%, 0.8-%, and 1.0-%. More detailed information about the measurement of N_{ccn} can be found in Wang et al. (2019).

During the experiment, ambient particles were first sampled through a conductive tube (TSI, US) and a diffusion dryer filled with silica gel (TSI, US); and then splitted intesplit for analysis by means of different instruments with a splitter. All instruments were placed in an air-conditioned container on the deck of the ship, with an inlet height of approximately 6 m above the sea level. Regarding the gas analyzers, the ambient O₃ (Model 49i, Thermo Environmental Instrument Inc., USA C-series), SO₂ (Model 43i, Thermo Environmental Instrument Inc., USA C-series), and NO_x (Model 42i, Thermo Environmental Instrument Inc., USA C-series) were measured in mixing ratios with a temporal resolution of one-_minute. The CCNC and gas analyzers were operated properly throughout the entire campaign. The same was true for the AIM-IC, which was used to measure the water-soluble ionic species in the ambient particles lessized smaller than 2.5 μm.

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In this study, the Hybrid Single-Particlehybrid single-particle Lagrangian Integrated Trajectoryintegrated trajectory (HYSPLIT) model from the NOAA Air Resources Laboratory was used to track the particle sources. The input of HYSPLIT, such as wind speed and wind direction, was obtained from the National Center for Environmental Prediction (NCEP) Global Data Assimilation System (GDAS) with a spatial resolution of 0.5 degree. degrees.

The hygroscopicity parameter (κ) was calculated according to the method proposed by Petters and Kreidenweis (2007).

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$$\kappa = \frac{4A^3}{27D_d^3 \; \ln^2 S_C} \; , \; A = \frac{4\sigma_{S/a} \; M_W}{RT\rho_W}$$

where D_d is the dry diameter, S_C is the super saturation supersaturation, M_w is the molecular weight of water, $\sigma_{s/a}$, a constant of 0.072 J m⁻², represents the surface tension over the interface of the solution and air with the value of 0.072 J m⁻² applied in this study. R is the universal gas constant, T is the ambient temperature and ρ_w is

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the critical diameter for CCN activation (Dcrit). Dcrit was defined as the particle diameter 188 189 down to which by integrating from the largest diameter with the integrated number 190 concentration equalsequal to the CCN concentration (Hung et al., 2014; Cheung et al., 191 2020). The FMPS hashad a low size resolution, particularly at the size greater than 192 90 nm, which doesn'tdid not allow accurately calculating accurate calculation of the Kappa values at SS=0.2%. At SS=0.6% and 0.8%, the Kappa value was not calculated 193 194 considering the complication in the explanation of the value, possibly reflecting the 195 combined effects of particle size, mixing state and chemical composition.— 196 197 2.2 Separating ambient signals of N_{cn} and N_{ccn} from ship self-ship emissions 198 The data measured during the cruise campaign were frequently interfered by subject to 199 interference from self-ship emission signals-from the ship. The N_{cn} and N_{ccn} over the 200 marginal seas were first distinguished based on the source of the ambient environment or the ship self-ship emission.emissions. The data measured at 18:00-24:00 on DOY 201 115 were are used to illustrate the separation (in Fig. 2), with, and the size distribution 202 of the particle number concentration during DOYDOYs 110-118 is shown in Fig. S2-203 S10 in the supporting information. At 18:00-21:11 LT (Local Time), the local time), a 204 Formatted: Font: Symbol 205 low N_{cn} of 5.8±0.4×10³ cm⁻³ werewas observed. The accumulation mode dominated inthe particle number concentration with thea median mobility mode diameter atof 206 Formatted: Font: Symbol 207 105±4 nm (Fig. 2a). Afterwards, the N_{cn} rapidly increased by over one order of 208 magnitude (Fig. 2b). The dominant particle number concentration mode changed from 209 accumulation mode to Aitken mode, with the median mobility diameter of the Aitken Formatted: Font: Symbol mode stabilized at 47±4 nm infor approximately 90% of the time. The rapid increase in 210 211 N_{cn} and the change in the mode size indicated the signal of the emissions of the ship 212 emission itself. The ship emission interference after 21:11 was also supported 213 by additional evidencesevidence, e.g., a large decrease in the activation ratio (AR), defined as the quotient of N_{ccn} and N_{cn}, from >0.5 to <0.2 at SS=0.4% (Fig. 2c) due to 214 215 a large increase of N_{cn} but a much smaller magnitude enhancement of N_{ccn} (Fig. 2b), Formatted: Font: Symbol

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a rapid increase of NO_x from <10 ppb to 192±99 ppb, NO/NO₂ from <0.1 to 0.7±0.3,

as well as and SO₂ from <2 ppb to 6.2±2.4 ppb. The large Large changes were expected

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because the ship smoke stock was <u>only</u> approximately <u>only</u> 10 meters away from these detectors. Thus, based upon the <u>featurefeatures</u> described above, certain criteria were designed in this study to identify <u>ship</u> self-<u>ship</u> emission signals so as to separate <u>them</u> from ambient signals, i.e., a net increase in N_{cn} beyond 5×10^4 cm⁻³ in five minutes, thea median mobility mode diameter <u>around of approximately</u> 50 nm, NO₂>30 ppb and NO/NO₂>0.5.

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3. Results and discussion

226 3.1 Spatiotemporal variations in ambient N_{cn} during the cruise period—

Fig. 3 shows athe time series of minutely averaged distributions of N_{cn}, N_{ccn} and AR at

228 SSSs of 0.4% and 1.0% from DOY 110 to DOY 135 2018, when after the ship self-

229 ship-emission signals had been exhaustedlywere exhaustively removed.—

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When the spatiotemporal variations in N_{cn} were examined during the first half of the cruise period (Fig. 3a), their was found that N_{cn} spanned a broad range of 0.2-4.5×10⁴ cm⁻³ with thean average value of $8.1 \pm 4.4 \times 10^3$ cm⁻³. Specifically, the N_{cn} were was only $6.5 \pm 0.8 \times 10^3$ cm⁻³ at 00:00-06:00 LT on DOY110DOY 110 when the ship anchored at the Yangtze River estuary near Shanghai (Fig. 1). The low N_{cn} values were comparable to the mean value of N_{cn} (5.4×10³ cm⁻³) in the marine-air cases during January-December 2010 in Shanghai reported by Leng et al. (2013). The Ncn <u>largely greatly</u> increased to $1.9 \pm 0.7 \times 10^4$ cm⁻³ at 08:00-21:00 LT on <u>DOY110DOY</u> 110 when the ship cruised across the Yangtze River estuary. The measured particles in the number concentration were dominantly distributed atin the Aitken mode on that day. while the median Aitken mode diameter shifted from 49±5 nm at 00:00-06:00 to 40±5 nm at 08:00-21:00 (Fig. S2). The Yangtze River estuary contains several world-class ports and is heavily travelled traveled by marine traffics in the daytime (Chen et al., 2017). Since the onshore wind dominated on that day (not shown), the increase in N_{cn} and the decrease in the median Aitken mode diameter at 08:00-21:00 LT possibly reflected the increased contribution from marine traffic emissions. Marine traffics The marine traffic visibly decreased when the ship left the Yangtze River estuary toward the south. The N_{cn} were value then significantly decreased, i.e., to $9.5 \pm 4.4 \times 10^3$ cm⁻³ in Formatted: English (United States)

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249 the marine atmosphere over the sea zone in Zhejiang Province (atfor 07:00 LT on DOY111DOY 111 - 17:00 LT on DOY 114), with P<0.01. The N_{cn} further decreased to 250 the lower values of $5.8 \pm 1.7 \times 10^3$ cm⁻³ in the marine atmosphere over the sea zone in 251 Fujian Province (atfor 18:00 LT on DOY 114 - 14:00 LT on DOY 117), All 252 these values were, however, 1-2 orders of magnitude largergreater than the background 253 values in remote clear marine atmospheres, e.g., <300 particle cm⁻³ without the 254 255 influence of industrial activities in the western Pacific and the tropical Pacific (Ueda et al., 2016) and those reported by Quinn and Bates (2011) and Saliba et al. (2019), 256 indicating overwhelming contributions from non-seanonsea-spray aerosols including 257 258 marine traffic emissions, the long-range continental transport, newly formed particles in marine atmospheres, etc. As reported, atmospheric particles over 259 260 China-marginal seas in China can be further transported to the remote northwest Pacific 261 Ocean (NWPO) in spring under westerly winds, e.g., the Ncn observed over the NWPO in March-April 2014 werewas as high as $2.8\pm1.0 \times 10^3$ 262 approxiately approximately half of that over China marginal seas in China observed in 263 264 March 2014 (Wang et al., 2019).-265 The mean value of N_{cn} (8.1 \pm 4.4 \times 10³) observed in this study was close to that of 7.6 266 267 $\pm 4.0 \times 10^3$ cm⁻³ (the number concentrations of particles larger than 10 nm) observed over the eastern part of the Yellow seaSea in spring 2017 reported according by Park et 268 269 al. (2018). They attributed the high number concentrations of particles within 270 nucleation and Aitken modes to the long-range transport of air pollutants over eastern 271 China under the influence of westerly winds. Consistently, larger values of N_{cn} were 272 frequently observed in the continental atmospheres upwind of the Yellow seaSea, e.g., the mean values of $1.8 \pm 1.4 \times 10^4$ cm⁻³ in May 2013 in Qingdao, a coastal city in 273 proximity to the Yellow Sea (Li et al., 2015), 3.18×10⁴ cm⁻³ in February-August 2014 274 in Beijing (Dal Maso et al., 2016), and 1.0×10⁴ cm⁻³ in continental-air cases during 275 January-December 2010 in Shanghai (Leng et al., 2013).-276 277 278

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3.2 Spatiotemporal variations in ambient N_{ccn} during the cruise period

279 280 N_{ccn} data were generally available during the entire campaign (Fig. 3b). The mean values of N_{ccn} over China marginal seas in China during the DOY 110 to DOY 135, 281 2018—were, ranged from $3.2 \pm 1.1 \times 10^3$ cm⁻³ to $3.9 \pm 1.4 \times 10^3$ cm⁻³ under SSSSs 282 ranging from 0.2% to 1.0%(% (Table 1), which is two to four times larger than the N_{ccn} 283 at the same SS over the NWPO in March-April 2014 (Wang et al., 2019), and much higher, i.e., 1-2 orders of magnitude, than the pristine marine background values (Quinn 284 285 and Bates, 2011). As-was discussed in the previous section, the mean N_{cn} in this study $(8.1 \pm 4.4 \times 10^3 \text{ cm}^{-3})$ was comparable to that of N_{en} $(7.6 \pm 4.0 \times 10^3 \text{ cm}^{-3})$ over the 286 287 Yellow Sea in spring 2017 in Park et al. (2018); however, the comparison of the mean N_{ccn} reveals that the mean value (3.6±1.2×10³ cm⁻³) at SS of 0.6% in this study was 288 approximately 25% smaller than that (4.8×10³ cm⁻³ at a similar SS of 0.65%) in Park 289 et al. (2018), which was likely a result of long range transport, considering the 290 observations made a relatively distantlong distance (i.e., 500-600 km) observations 291 292 away from the land depicted in Fig. 1 of Park et al., 2018, and the subsequently higher 293 extent of aerosol aging. The N_{ccn} under SS of 0.2% in this study (3.2±1.1×10³) is comparable to that $(3.1\pm1.9\times10^3)$ byof Li et al. (2015) in the continental atmosphere 294 of Qingdao in May 2013; however, the increment of N_{ccn} with the increase of increasing 295 SS was much weaker in our study, resulting in onan average of 36% smaller _in Nccn 296 under <u>SSSSs</u> of 0.4% to 1.0% compared to <u>that of</u> Li et al. (2015). <u>Consistently, the The</u> 297 298 sensitivity differences of N_{ccn} to SS between the relatively clean (i.e., N_{cn} (8.1 \pm 4.4 \times 10³) in this study) and polluted (with Ncn of 1.8 \pm 1.4×10⁴ cm⁻³) 299 environments in Li et al. (2015) iswere also reported by Nair et al. (2019), 300 who found little sensitivity of N_{ccn} to changes in SS over the equatorial Indian Ocean 301 (<6°N) with relatively clean air, and much larger enhancement of N_{ccn} with the increase 302 of increasing SS in polluted marine atmospheres (> 6 °N). 303 304 305

In addition, the N_{ccn} at SSSs from 0.1% to 1.0% during the period with high NH₄⁺ (17:00 LT on DOY 114 to 10:00 LT on DOY 120) is statistically significantsignificantly higher (P<0.01) in comparison tothan that during the poor NH₄⁺ period (11:00 LT on DOY 120 to 7:00LT₀₀ LT on DOY 136; Fig. 3b). More specifically, a large increase in NH₄⁺ concentration, with a mean concentration of 6.3±2.5 μg m⁻³, can be observed during the period from 17:00 LT on DOY 114 to 10:00 LT on DOY 120 (Fig. 3b). The

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312 SSSS ranging offrom 0.2% to 1.0%. In contrast, after DOY 120, the concentration of NH₄⁺ (0.67±0.70 μg m⁻³) substantially decreased by almost 90%, during which the 313 314 mean N_{ccn} at each SS showed statistically significant decreased decreases of 8% to 15%, implicative of the vital contribution to CCN of secondary ammonium salt aerosols. to 315 CCN. 316 317 318 Another feature depicted in Fig. 3b is that the N_{ccn} during the low NH₄⁺ period may 319 even exceed the maximal value of N_{ccn} during the high NH₄⁺ period. To elucidate the underlying mechanism, the $N_{\text{cen}_{\overline{2}}}$ values under each SS, was were composited and 320 321 compared between for the days with NH₄⁺ concentration concentrations higher than the 322 upper quartile and the days in the lower quartile, yielding some interesting findings. At 323 SS=0.2%, the composited N_{ccn} under the high NH₄⁺ period was higher than that during the low NH₄⁺ period with a statistical significance level of 0.01. There was no 324 325 significant difference in Neen-between the Neen values of the two composited composite 326 periods at SS values of 0.4% and 0.6%. However, the composited Nccn (i.e., only selection of the upper quartile) during the high NH₄⁺ period was significantly lower 327 328 than the composited value during the low NH₄⁺ period with for P<0.01, e.g., 5.1 ± 0.5×10^3 cm⁻³ versus $5.3 \pm 0.7 \times 10^3$ cm⁻³ at SS= $0.8 \frac{96}{5} \frac{9}{0}$ and $5.2 \pm 0.5 \times 10^3$ cm⁻³ versus 329 $5.7 \pm 0.7 \times 10^3$ cm⁻³ at SS =1.0%. During the low NH₄⁺ period, the marine atmospheres 330 331 over the observational zones may sometimes receive strong continental inputs and/or 332 marine traffic emissions, leading to the larger N_{ccn}. Enhanced The enhanced formation of ammonium salt aerosols during the high NH₄⁺ period likely canceled out or even 333 334 overwhelmed the effects of the continental inputs and/or marine traffic emissions in 335 increasing on Nccn at SS=0.2%.-In addition, fresh marine traffic emissions likely <u>yieldedmade</u> a negligible contribution 336 to N_{ccn} in the marine atmosphere because of <u>athe</u> large <u>amountamounts</u> of aged aerosols 337 338 from various sources therein. For example, the mean values of N_{ccn} were 3.2×10³ cm⁻³ and 4.5×10³ cm⁻³ at SS=0.4% and 1.0% at 08:30-11:30 on DOY110DOY 110, 339 respectively. They These values were almost the same as the 3.2×10³ cm⁻³ at SS=0.4% 340

mean N_{ccn} during this period varied from $3.5 \pm 1.0 \times 10^3$ cm⁻³ to $4.0 \pm 1.1 \times 10^3$ cm⁻³ at

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341 and 3.8×10³ cm⁻³ at SS=1.0% before 06:00 on that day. The mean values of N_{cn}, however, largely greatly increased from $6.5 \pm 0.8 \times 10^3$ cm⁻³ before 06:00 to $1.3 \pm$ 342 Formatted: English (United States) 0.3×10⁴ cm⁻³ at 08:30-11:30 when the ship cruised across the Yangtze River estuary 343 (Fig. 3b).-344 345 346 3.3 Spatiotemporal variations in CCN activation and Kappa values 347 The AR values at SSSs of 0.4% and 1.0% wereare examined in thethis section, as 348 shown in Fig. 3c. At SS=0.4%, the AR values largely varied from 0.06 to 0.92 with thea median value of 0.51. Specifically, the AR values narrowly varied around 0.51 \pm 0.04 349 Formatted: English (United States) at 00:00-06:00 LT on DOY110DOY 110. At 08:00-21:00 LT on that day, when the ship 350 Formatted: English (United States) 351 cruised across the Yangtze River estuary, the AR values were substantially decreased to Formatted: English (United States) 0.26 ± 0.06 concurrently with an approximate 200% increase in the N_{cn} values, i.e., N_{cn} 352 Formatted: English (United States) 353 values of $6.5 \pm 0.8 \times 10^3$ cm⁻³ at 00:00-06:00 LT and $2.0 \pm 0.7 \times 10^4$ cm⁻³ at 08:00-09=0.00Formatted: English (United States) 354 21:00 LT on DOY110 (Fig. 3a). The AR values then exhibited an oscillating Formatted: English (United States) Formatted: English (United States) increase from DOY 111 to DOY113. LowDOY 113. A low AR valuesvalue of 0.12 ± 355 Formatted: English (United States) 356 0.04 werewas suddenly observed at 10:00-18:00 LT on DOY114DOY 114 in the presence of strong new particle signals transported from the upwind continental 357 atmosphere, as discussed later. The AR values, however, reached 0.34± 0.04 at 06:00-358 359 08:00 LT and 0.39 ± 0.08 at 19:00-24:00 LT on $\frac{\text{DOY}114}{\text{DOY}114}$, with the new particle 360 signals largely reduceddecreased. Even excluding the AR values on DOY 114, a significant difference was still obtained between the AR values of 0.61 ± 0.12 during 361 the high NH_4^+ period and those of 0.55 \pm 0.17 during the low NH_4^+ period. 362 363 Enhanced The enhanced formation of ammonium salts seemingly increased the CCN activity to some extent. At SS=1.0%, the AR values showed large 364 fluctuation fluctuations with thea median value of 0.57± 0.17 (Fig. 3c), and the 365

To minimize the impact from the particle sizes, the Kappa values were further

temporal trend was similar to that at SS=0.4%.—

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resolution usually exhibited exhibit a broad distribution, reflecting the complexity due to various of factors. To reveal the key factors in determining the Kappa values inon a large spatiotemporal scale, the daily Kappa values of atmospheric aerosols were estimated, on the basis of the daily mean N_{cen} and the size distributions of the particle number concentration from DOYDOYs 110-118 (Fig. 3c). Please note that for DOY 110, considering the large differences of the particle number concentration between 00:00-06:00 and 08:00-21:00 (Fig. S2), the Kappa values were calculated separately for these two periods. At SS=0.4% (green dashed line in Fig. 3c), the estimated Kappa values were as high as 0.66 at 00:00-06:00 LT, while itthey decreased to 0.37 at 08:00-21:00 LT on DOY110DOY 110. The Kappa value varied narrowly from 0.46 to 0.55 on DOYDOYs 111-113, 115 and 117, implying that inorganic aerosols such as completely and incompletely neutralized ammonium salts may vield amake large contribution to the N_{ccn}. These values were generally consistent with the reported observations in most of marine atmospheres. For example, Cai et al. (2017) reported thea Kappa value around of approximately 0.5 for particles with sizes of 40-200 nm at a marine site in Okinawa and that sulfate to bewas the dominant component of aerosol particles on 1-9 November 2015, and a similar Kappa value in spring 2008 over this site was reported by Mochida et al. (2010)- over this site. Royalty et al. (2017) reported Kappa values for 48, 96, and 144 nm dry particles to beof 0.57 ± 0.12 , 0.51 ± 0.05 0.09, and 0.52 ± 0.08 in the subtropical North Pacific Ocean and sulfate-like particles contributing at most 77-88% to the total aerosol number concentration. Kappa values overOver the Atlantic Ocean-, Kappa values of approximately 0.54 ± 0.03 were observed around 0.54 ± 0.03 for 284 nm particles (Phillips et al., 2018).—

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398 399 The estimated *Kappa* values sometimes reached 0.66-0.67 (i.e., on DOY 116), which may be related to unidentified factors. For example, O'Dowd et al. (2014) proposed that some organics derived from sea-spray aerosols may also increase the N_{ccn5} to some extent, by reducing the surface intensiontension, leading to an increase of in the *Kappa* values. A small fraction of sea-salt aerosols in submicron particles may also increase the *Kappa* values since its *Kappa* value was as high as 1.3 (O'Dowd et al., 1997;

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400 O'Dowd et al., 2004). The A Kappa value of 0.29 was obtained on DOY118, DOY 118, 401 which is close to the Kappa values widely observed for continental atmospheric 402 aerosols (~0.3) (Andreae and Rosenfeld, 2008; Poschl et al., 2009; Rose et al., 2010). 403 The estimated Kappa value largely decreased to 0.15 on DOY114DOY 114 when new 404 particle formation (NPF) occurred, with; see section 3.5 for detailed discussion in section 3.5. Moreover, at an SS of 1.0%, the estimated Kappa value was always smaller 405 406 than 0.2. The Kappa values of organics waswere commonly assumed asto be 0.1 407 (Rose et al., 2011; Cai et al., 2017; Singla et al., 2017). In general, the fraction of organics in the nanometer particles increases with decreasing particle sizes from 408 <u>~</u>100 nm to 40-50 nm (Rose et al., 2010; Rose et al., 2011; Crippa et al., 2014; 409 Cai et al., 2017). A combination of the two factors likely led tothe overall Kappa values 410 411 estimated at SS=1.0% to be much lower. However, the direct measurements of the 412 chemical composition of nanometer particles are needed to confirm thethese arguments. 413 414 3.4 Particle number size distributions and CCN activation associated with marine 415 traffic emissions and aerosol aging 416 The particle number size distributions during **DOYDOYs** 110-118, shown in Fig. 4, can 417 be in generalgenerally classified into two categories. Category 1 occurred on DOY110 DOYs 110-114, when particle number concentrations were mainly distributed 418 419 at thein Aitken mode, whereas the accumulation mode was generally undetectable. Category 2 occurred on **DOY115**DOYs 115-118, when the accumulation mode 420 cancould be clearly identified and generally dominated over the Aitken mode. Hoppel 421 422 (1986) proposed that cloud-modified aerosols to beare mainly distributed at 80-150 nm 423 in the remote tropical Atlantic and Pacific oceans. Cloud-modified aerosols are quite 424 common in remote marine atmosphereatmospheres, likely leading to the dominatedominant accumulation mode particles to bebeing observed 425 426 DOY115 DOYs 115-118. Occasionally, the Aitken mode dominated over the accumulation mode-on some day, such as on DOY 118. To further dive into investigate 427

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429 116 and DOY118DOY 118 were selected.— 430 431 On DOY 112, the Aitken mode particles accounted for approximately 60% of the total 432 particle number concentration (Fig. 5a), with median Aitken mode diameters aroundof Formatted: Font: Symbol 433 approximately 54±8 nm, LikeSimilar to the observations over the Yangtze River estuary, Formatted: Font color: Black the mean value of N_{cn} increased by approximately 50% concurrently with a decrease in 434 Formatted: English (United States) the median Aitken mode diameters by ~9 nm at 05:30 - 11:40 LT against compared to 435 Formatted: Font color: Black 436 those atin the early morning before 05:30 LT (Fig. 5b).— Concomitantly, the AR values Formatted: Font: Symbol decreased to 0.31±0.09 at SS of 0.4%, with similar AR decreased decreases at SS of 1.0%, 437 438 and the lowest AR and Kappa values occurring occurred at 06:00-07:00 LT at SSSSs of 439 both 0.4% and 1.0%. All these results pointed towards indicated that the increase in 440 Aitken mode particles at 05:30 - 11:40 LT to bewas likely derived from enhanced 441 marine traffic contributions carried by the onshore wind from the south (Fig. S11). 442 During other timetimes on DOY112DOY 112, the onshore wind may also carry the Formatted: Font color: Black marine-traffic-derived particles to the observational sea zones. However, the marine-443 Formatted: Font color: Black traffic-derived particles likely aged to some extent, e.g., the median Aitken mode 444 Formatted: Font color: Black diameters exhibited an oscillating increase from approximately 50 nm at 19:00 to 445 446 approximately 70 nm at 24:00 LT with thea particle growth rate of ~4 nm hour⁻¹. The Formatted: Font: Symbol AR values, however, narrowly varied around 0.47±0.03 at SS=0.4% and 0.52±0.05 at 447 Formatted: Font: Symbol SS=1.0% during the particle growth period. The Kappa values at SS=0.4% gradually 448 Formatted: Font color: Black 449 decreased from 0.56 at 19:00 to 0.41 at 23:00 LT, reflecting more aged marine-traffic 450 _derived particles growing into CCN size. sizes. 451 452 On DOY 116, the accumulation mode particles instead of Aitken mode particles 453 dominantly contributed to N_{cn} rather than Aitken mode particles (Fig. 5d), under the marine air influence from the northeast (Fig. S13). The median accumulation mode 454 Formatted: Font: Symbol diameters narrowly varied around 135±5 nm at 01:00-13:00 LT and 102±5 nm at 16:20-455 Formatted: Font: Symbol 456 24:00 LT with thea transition period in between (Fig. 5e). The AR and Kappa values, however, showed no statistically significant differenced during the two 457

periods at SSSSs of 0.4% and 1.0%, implying that the size change in the accumulation

mode particles showedhad a negligible influence on the CCN activation. HourlyThe 459 460 hourly variations in the AR and Kappa values may be associated with other factors, e.g., 461 chemical composition, and mixing state, etc. (Gunthe et al., 2011; Rose et al., 2011).-462 463 On DOY 118, under the influence of mixture mixtures from the marine and coastal areas 464 from the northeast (Fig. S14), the accumulation mode particles generally dominated the 465 contribution to N_{cn} while the reverse was true inon some occasions (Fig. 5g, h). The 466 median accumulation mode diameters exhibited an oscillating increase from 467 approximately 100 nm to 130 nm at 00:00-08:00 LT, narrowly varied around 133±5 nm at 08:00-13:00 LT, and then exhibited an oscillating decrease down to approximately 468 100 nm at 20:00 LT. The AR values and Kappa values at SS=0.4%, however, exhibited 469 470 an inverted bell- shape with the lowest values at 0.31 and 0.11 at 13:00. The decreases 471 in the AR values and Kappa may be related to organic condensed condensation on the accumulation mode particles since the median accumulation mode diameters were 472 473 almost the largest at 13:00. The number concentration of Aitken mode particles was 474 evidently enhanced at 14:00-15:00, but the influence on the AR values and Kappa 475 values at SS=0.4% was undetectable (Fig. 5i).-476 477 3.5 The long-range transport of <u>new</u> grown new-particles on DOY 114 478 No hour-long sharp increase-increases were observed in the number concentration of 479 the nucleation mode particles (< 20 nm) was observed during the period from DOY 110 480 to DOY 118, except on DOY114 DOY 114 (Fig. 4). According to the conventional definition of NPF events (Kulmala et al., 2004; Dal Maso et al., 2005), the occurrence 481 frequency of NPF events was low in this study. Unlike continental atmospheres where 482 a high occurrence frequency of NPF events has been observed globally in spring 483

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(Kulmala et al., 2004; Kerminen et al., 2018), a low occurrence frequency reportedly

occurred over the seas during the "Meiyu (plum-rain) season" in spring because of frequent rainy, foggy or cloudy weather conditions (Zhu et al., 2019). Lack The lack of

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bewere mainly contributed by primarilyfrom primary emitted acrosols and their aged 488 489 products.-During the period of 10:00-18:00 LT on DOY 114, athe large increase in the number 490 concentrations of Aitken mode particles (Fig. 6a) likely reflected the long-range 491 492 transport of <u>new grown new particles</u> from upwind continental atmospheres (Fig. S12). 493 The size distributions of the particle number concentration showed a dominant Aitken 494 mode at 10:00-18:00 LT, when the spatiotemporal variations in N_{cn} and median Aitken 495 mode diameters exhibited bell-shapeshaped patterns (Fig. 6b). The median Aitken 496 mode diameters increased from 26 nm at 10:00 LT to 33 nm at 12:00-13:00 LT and then 497 decreased to 20 nm prior to the signal disappearance, likely reflecting the growth and shrinkage of the Aitken mode particles (Yao et al., 2010; Zhu et al., 2019). The median 498 499 Aitken mode diameters were evidently smaller than the values, i.e., 40-50 nm for the 500 Aitken mode particles, observed over the Yangtze River estuary on DOY 112 (Fig. 5a). Moreover, the number concentrations of the 20-40 nm particles increased by 5.8 times 501 at 12:00-13:00 LT compared to the mean value at 06:00-09:00 LT, while the total 502 number concentrations of particles greater than 90 nm increased by only 67%. These 503 504 results implied that the largely increased large increases in the number concentrations 505 of Aitken mode particles with a dynamic change in the mode diameter observed at 10:00-18:00 LT unlikely to bewere not likely caused by primarilyprimary emitted and 506 aged particles from marine traffic emissions or other combustion sources. The 507 508 observations of the gaseous and particulate species, during the same period, implied that the air masses to bewere well-aged and less polluted. For instance, the measured 509 hourly average mixing ratios of SO₂ waswere no larger than 1.2 ppb (Fig. 6c), and the 510 hourly average concentrations of NH₄⁺ in PM_{2.5} were smaller than 2 µg m⁻³ (Fig. 3b). 511 512 In addition, the concentrations of K⁺ were below 0.3 µg m⁻³, suggesting negligible

Before 09:00 LT, a much weaker spike of nucleation mode particles was intermittently observed (Fig. 6a). The weak and intermittent NPF seems to occur in the marine atmospheres before 09:00 LT when no apparent growth of new particles was observed.

contributions from biomass burning (Fig. 6e).-

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Possibly due to the transport from the continent (Fig. S12) and an increase in the condensational sink aroundat approximately 10:00 am (Fig. 6a), the weak NPF signal gradually dropped to a negligible level half an hour later, concomitant with a large increase in the number concentrations of Aitken mode particles at 10:00-18:00 LT.

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 N_{ccn} at SS=0.4% increased from 1.2×10^3 cm⁻³ at 06:00-09:00 LT to the peak value of 2.3×10³ cm⁻³ at 12:00 LT, with an increase of 92%, and N_{ccn} at SS=1.0% increased from 1.6×10^3 cm⁻³ to 4.0×10^3 cm⁻³, with an increase of 150% (Fig. 6d). The net increase in N_{ccn} at SS=0.4% likely reflected the contribution from pre-existing particles since new particles with the diameter diameters less than 50 nm were unlikely to be activated as CCN at such low SS (Li et al., 2015; Wu et al., 2016; Ma et al., 2016). The larger net increase in N_{ccn} at SS=1.0% may reflect the contributions mixed from preexisting particles and new grown new particles. The high SS can activate particles as CCN with diameters down to 40 nm (Dusek et al., 2006; Li et al., 2015). The invasion of new grown new particles also led to the large decreases in the AR values largely decreased from 0.3 to 0.1 at SS=0.4%, on and from 0.4 to 0.2 at SS=1.0% (Fig. 6e). After 18:00 LT, the AR values retuned to 0.3-0.4 at SS=0.4% and 0.4-0.6 at SS=1.0%. When the The calculated Kappa values were examined (Fig. 6c), they decreased 6c) and were found to decrease from 0.4 to 0.1-0.2 at SS=0.4%. The This value returned to 0.3 at 18:00-19:00 LT (FMPS-was temporarily malfunctioned after 19:20 LT). The Kappa values were below 0.2 at SS=1.0% on that day. The decreases in the AR values and Kappa values at the two SS were likely caused by organic vapor condensed on preexisting particles and new particles (Wu et al., 2016; Zhu et al., 2019).—

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3.6 Correlations Relationship Relationship of N_{cn} and N_{ccn} with SO₂ in <u>ship</u> self-ship

plumes and ambient air-

When <u>ship</u> self-<u>ship</u>-emission signals were detected, the observational values included a combination of contributions from <u>ship</u> self-<u>ship</u> emissions and ambient concentrations. Although <u>the</u> ambient N_{cn} was negligible in comparison with <u>the</u> N_{cn}

derived from the ship self-ship-emissions, itthis was not the case for N_{cen} and SO₂. Based on the minutelyper minute data, the signal was considered asto be vessel-self emission-emissions when both N_{cn} was greater than 50,000 cm⁻³ and SO₂ was greater than 5 ppb. The composited data waswere then used to derive the hourly average N_{cn}, N_{cen} and SO₂, which was then subtracted by the ambient hourly mean value during the preceding hour with relatively clean conditions (i.e., concentration concentrations of N_{cn} lower than 10,000 cm⁻³, and SO₂ lower than 2.5 ppb). Please note that uncertainties exist in terms of the criteria and separation between the ship self-shipsignals and ambient signals; however, minimal impact is expected in the relationship examined below.

Fig. 7a showed correlationsshows the relationship of N_{cn} and N_{cen} with the mixing ratio of SO₂ in the ship self-ship plumes, prefixed by Δ for N_{cn}, N_{cen} and SO₂ to implicate the removal of ambient signals. A good correlation of 0.66 for R² (P<0.01) wasis obtained, and the slope indicates that N_{en}the increase in N_{cn} by 1.4 × 10⁴ cm⁻³ for each ppb increase of SO₂ resulted from ship emissionemissions (Fig. 7a). High emissions of N_{cn} were generally reported in engine exhausts withwhere high sulfur-content diesel to bewas used (Yao et al., 2005; Yao et al., 2007). In regard of The N_{cen} at SS of 0.2% to 1.0% (Fig. 7b), it increases) increased from 30 cm⁻³ to 170 cm⁻³ per 1 ppb increase of SO₂, showing statistical a statistically significant correlation at the 99th confidence level. The contribution ratioratios of SO₂ to N_{cen} iswere 0.002 (SS of 0.2%), 0.004 (SS of 0.4%) and 0.012 (SS of 1.0%) to that of N_{cn}, which is in general consistent with thea previous study by Ramana and Devi (2016), in which a range of 0.0012–0.57 was observed for CCN at 0.4% in Bay of Bengal during Aug 13–16, 2012.—

The <u>correlationsrelationship</u> of hourly averaged N_{cn} and N_{ccn} with SO₂ in ambient air were examined and <u>showedare shown</u> in Fig. 7c, d. The data <u>waswere</u> segmented into pieces based on SO₂ with <u>an</u> interval of 0.2 ppb. A good correlation <u>was obtained</u> between the averaged N_{cn} and SO₂ <u>were obtained</u> with <u>an</u> R² of 0.80 (P<0.01), and <u>a</u>

576	1 ppb increase in SO ₂ likely increased N_{en} by 1.6×10^3 cm ⁻³ (Fig. 7c). The increase in			
577	N _{cn} with SO ₂ may reflect the contribution from primary emissions. An The intercept was,			
578	however, as large as 3.9×10^3 cm ⁻³ , likely representing the contribution from well-aged			
579	aerosols.—			
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581	$\underline{\text{HourlyThe hourly}} \ \text{averaged} \ N_{ccn} \ \text{at different SS generally increased with } \underline{\text{increase}}$			
582	ofincreasing ambient SO ₂ (Fig. 7d). A good correlation was obtained between the			
583	averaged N_{ccn} and SO_2 -were obtained, with R^2 =0.78-0.91 (P<0.01), \underline{A} 1 ppb increase			
584	in SO ₂ likely increased N_{cen} by 0.6×10^3 to 0.8×10^3 cm ⁻³ at SSSS from 0.2% to 1.0%.			
585	The increase in N_{cen} with SO_2 may also reflect the contribution from primary emissions.			
586	The intercepts of 2.2×10^3 - 2.7×10^3 cm ⁻³ at different SS were likely contributed by			
587	well-aged aerosols. $\underline{\text{The}\underline{\text{This}}}$ relationship may be used as an estimation of $\underline{\text{the}}\underline{\text{N}}_{cen}$ in			
588	marine atmospheres over China marginal seas, in China when no measurements of CCN			
589	$\underline{\text{were}\underline{\text{are}}} \text{ available}\underline{\text{a}} \text{ whereas }\underline{\text{the}} \text{ ambient SO}_2 \text{ can be estimated from web-based satellite}$			
590	data.—			
591	4. Conclusions			
592	$\underline{\textbf{Spatiotemporal}}\underline{\textbf{The spatiotemporal}} \ \ \text{variations in ambient} \ \ N_{cn} \ \ \text{and} \ \ N_{cen} \ \ \text{were studied}$			
593	during a cruise campaign on DOYDOYs 110-135 over China-marginal seas in China			
594	The mean values of $N_{cn}~(8.1\times10^3~cm^{\text{-}3})$ and $N_{ccn}~(3.2~\text{-}3.9\times10^3~cm^{\text{-}3})$ at $\frac{\text{SSSS}}{\text{SSS}}$ of 0.2%			
595	1.0% were approximately one order of magnitude larger than those in remote clear			
596	marine atmospheres, indicating overwhelming contributions from non-seanonsea-spray			
597	aerosols such as marine traffic emissions, the long-range continental transport and			
598	others.			
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600	Observed The observed ship self-ship emission signals showed that fresh marine traffic			
601	emissions can be important sources of N_{cn_7} but a-minor sources of N_{cen} in the			
602	marine atmosphere. The signals showed that <u>a</u> l ppb increase in SO ₂			
603	corresponds corresponded to a 1.4×10 ⁴ cm ⁻³ increase in N _{cn} and a 30-170 cm ⁻³ increase			
604	in Non at SS=0.2-1.0% Data analysis showed that marine traffic emissions largely			

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605 increased N_{cn} over their heavily travelled traveled sea zones in the daytime.-606 607 In ambient marine air, the growth of marine traffic-derived particles led to a decrease in the estimated bulk kappa values at 0.4%%, possibly because some of these particles 608 609 enriched in organics grew into CCN size. However, strong formation of ammonium 610 salts led to aerosol aging, and significantly increased N_{ccn} at SS of 0.2-1.0% in 611 comparison with those observed during the period poor in ammonium salt aerosols in Formatted: English (United States) 612 PM_{2.5} with P<0.01. The estimated bulk Kappa values from the daily average values 613 varied from 0.46 to 0.55 at SS=0.4% in most of marine atmospheres, indicating that inorganic ammonium aerosols may dominantly contribute to the N_{ccn} at SS of 0.4%. 614 Formatted: English (United States) 615 The particle number size distributions showed that the high bulk Kappa values could 616 be related to cloud-modified aerosols, which likely led to a large extent of degradation Formatted: English (United States) 617 of organics and subsequently lost subsequent loss, from the particle phase.— 618 Humid marine ambient air led to NPF events rarely occurring therein occurred in the 619 620 humid ambient marine air. The dominant onshore winds occurred during most of the measurement periods, and should carrylikely carried primary aerosols and their aged 621 622 products rather than secondarily formed aerosols to the observational zone. During an 623 occasion when offshore winds blew from the northwest (Fig. S12), new particle signals 624 transported from the continent can be clearly observed. However, the NPF in the marine atmosphere was too weak to be important. The new transported new particles from the 625 continent yielded the maximal $\frac{increase increases}{in N_{ccn}}$ in N_{ccn} by of 92% at SS of 0.4% and 626 Formatted: Font color: Black 150% at SS of 1.0%. However, consistent with those reported in the literature, the 627 628 estimated kappa Values largely decreased from 0.4 to 0.1-0.2 at SS=0.4% during 629 most time of the continent-transported transporting NPF event because of the

Author contributions. YG and XY designed the research, YG, DZ and XY performed

kappaKappa value of the organic condensation vapor was as low as 0.1.—

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

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- 634 the analysis, JW and HG helped on the interpretation of the results, and all co-authors
- 635 contributed to the writing of the paper.

636 Acknowledgment

- This research is supported by the National Key Research and Development Program in
- 638 China (grant no. 2016YFC0200504) and the Natural Science Foundation of China
- 639 (grant no. 41576118).

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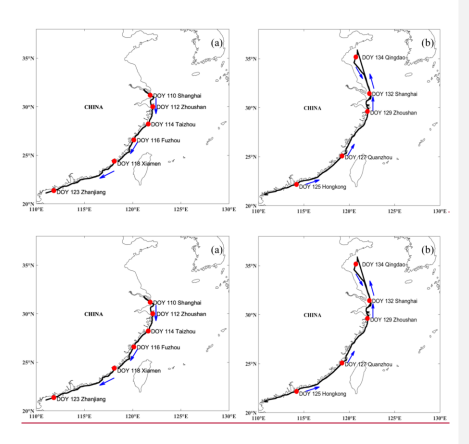
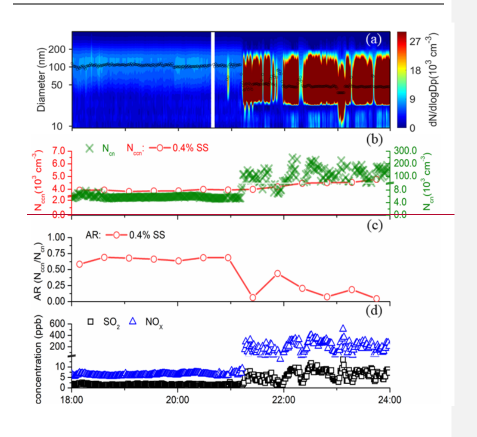


Fig 1 The ship track during the campaign of 2018, and where the blue arrows represented represent the sailing direction, with the southward track (a) and northward track (b).



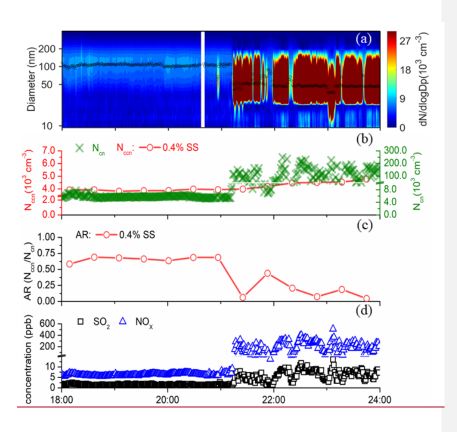


Fig 2 Contour plot of particle number size distribution with the median mobility mode diameter shown in black hollow circles (a), time series of minutelyper minute N_{cn} and half-hourly N_{ccn} at SS=0.4% (b), half-hourly AR values at SS=0.4% (c), SO₂ and NO_x at nighttime on DOY 115.

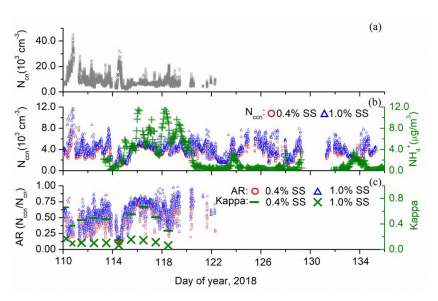


Fig 3 Time series of minutelyper minute N_{cn} from DOY 110 to 122 (a), minutelyper minute N_{cn} at SS of 0.4% and 1.0% during DOY 110-135 and hourly NH₄⁺ during DOY 113-135 (b), and minutelyper minute AR at SS of 0.4% and 1.0% during DOY 110-122 and daily *Kappa* values at SS of 0.4% and 1.0% from DOY 110 to 118 due to data availability (c). Please note that for Fig. 3c, most *Kappa* values were based on a daily scale, except on DOY 110, during which two *Kappa* values were calculated from 00:00-06:00 and 08:00-21:00, respectively.

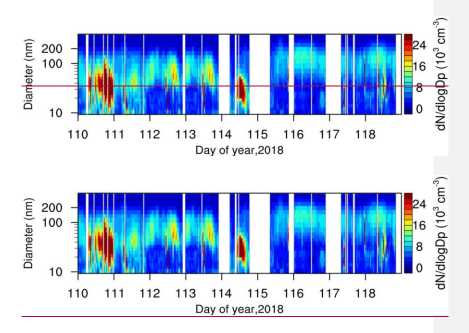


Fig 4 Contour plot of particle number size distribution on DOY 110-118 with <u>ship-emission signals removed.-</u>

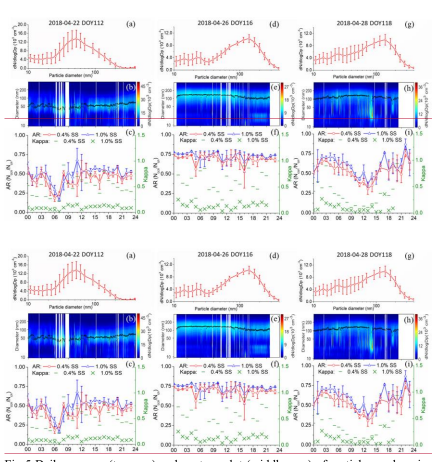


Fig 5 Daily average (top row) and contour plot (middle row) of particle number size distributions, and time series of hourly averaged AR at SS of 0.4% and 1.0% and *Kappa* value on DOY112, DOY116 DOY 112, DOY 116 and DOY118 DOY 118. The bars represent the standard deviation, with the mean indicated by the hollow circles.—

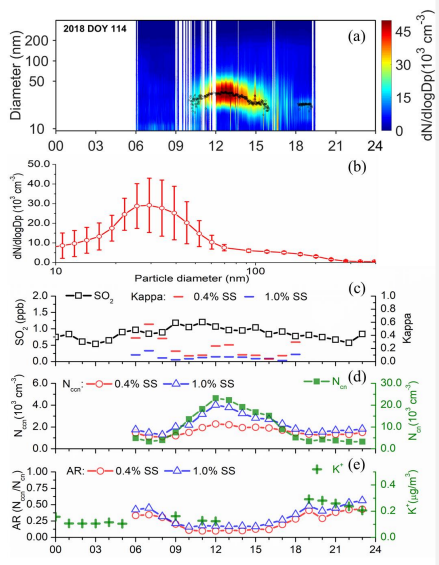


Fig 6 Contour plot of particle number size distributions for the day of DOY 114 2018 (a), the size distributions of <u>the particle number concentration during 10:00 -18:00 LT DOY 114 2018 (b)</u>, time series of hourly averaged SO_2 and *Kappa* values at SS of 0.4% and 1.0% (c), N_{ccn} at SS of 0.4% and 1.0% (d), and AR values at SS of 0.4% and 1.0% and K^+ (e) for the day of DOY 114 2018.

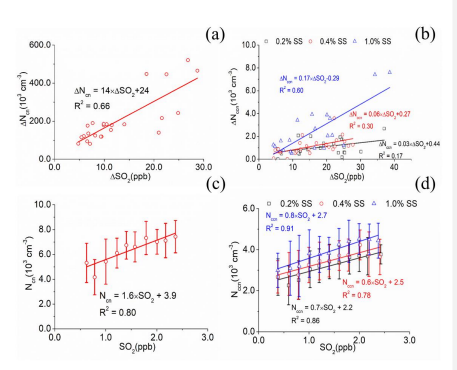


Fig 7 CorrelationsRelationship of hourly averaged N_{cn} and N_{ccn} with SO_2 at SS of 0.2%, 0.4% and 1.0%. For Fig. 7a, b, $\triangle N_{cn}$, $\triangle N_{ccn}$ and $\triangle SO_2$ reflectsreflect the impact fromof the ship self-ship emission after the removal of the ambient concentration. For Fig. 7c, d. each bar indicates the standard deviation with the mean value marked as the hollow circles (or triangles, squares), and the interval of SO_2 is 0.2 ppb for each bar.

Table 1. N_{cn} and N_{ccn}, AR and SO₂ mixing ratios on DOY 110-135, 2018, over Chinamarginal seas in China. Please note that Ncn and AR are from 110-122, 2018.

Variables Variable	Supersaturation	Ranges Range	Mean ± standard
	(SS)		deviation
N _{cn} (×10 ³ cm ⁻³)		2.0-45	8.1±4.4
	SS=0.2%	0.4-8.8	3.2±1.1
	SS=0.4%	0.5-9.4	3.4±1.1
N_{ccn} (×10 ³ cm ⁻³)	SS=0.6%	0.5-8.6	3.6±1.2
	SS=0.8%	0.5-11	3.8±1.2
	SS=1.0%	0.6-12	3.9±1.4
	SS=0.2%	0.06-0.89	0.49±0.17
	SS=0.4%	0.06-0.92	0.51±0.17
AR	SS=0.6%	0.10-0.94	0.54±0.17
	SS=0.8%	0.08-0.95	0.56±0.17
	SS=1.0%	0.11-0.98	0.57±0.17
SO ₂ (ppb)		0.25-9.7	1.7±1.1

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