

1 **In-depth study of the formation processes of single atmospheric particles in**
2 **the southeastern margin of Tibetan Plateau**

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13

14 **Abstract**

15 The unique geographical location of the Tibetan Plateau (TP) plays an important role in
16 regulating global climate change, but the impacts of the chemical components and
17 atmospheric processing on the size distribution and mixing state of individual particles are
18 rarely explored in the southeastern margin of the TP, which is a transport channel for
19 pollutants from Southeast Asia to the TP during the pre-monsoon season. Thus a
20 single-particle aerosol mass spectrometer (SPAMS) was deployed to investigate how the local
21 emissions of chemical composition interact with the transporting particles and assess the
22 mixing state of different particle types and secondary formation in this study. The TP particles
23 were classified into six distinct types, mainly including the largest fraction of the
24 rich-potassium (rich-K) type in the total particles (30.9%), followed by the biomass burning
25 (BB) type (18.7%). Most particle types were mainly transported from surroundings the
26 sampling site and along the Sino-Myanmar border; but the air mass trajectories from
27 northeastern India and Myanmar show a greater impact on the number fraction of BB (31.7%)
28 and Dust (18.2%) types, respectively. Then, the two episodes with high particle
29 concentrations showed that the differences in the meteorological conditions in the same
30 trajectory clusters could cause significant changes in chemical components, especially the
31 Dust and EC-aged types, which changed by a sum of 93.6% and 72.0%, respectively.
32 Ammonium and Dust particles distribute at a relatively larger size (~ 600 nm), but the size
33 peak of other types is present at ~ 440 nm. Compared with the abundant sulfate ($^{97}\text{HSO}_4^-$),
34 the low nitrate ($^{62}\text{NO}_3^-$) internally mixed in TP particles is mainly due to the fact that nitrate
35 is more volatilized during the transport process. The formation mechanism of secondary
36 speciation demonstrate that the formation capacity of atmospheric oxidation is presumably
37 affected by the convective transmission and the regional transport in TP. However, the
38 relative humidity (RH) could significantly promote the formation of secondary species,
39 especially $^{97}\text{HSO}_4^-$ and $^{18}\text{NH}_4^+$. This study provides new insights into the particle
40 composition and size, mixing state, and aging mechanism in high time resolution over the TP
41 region.

42 **Keywords**

43 Southeastern Tibetan Plateau, Individual particles, Chemical characteristics, mixing state,

44 Secondary formation

45 **1 Introduction**

46 Atmospheric aerosols have complex components and sources and can be coated with
47 inorganic or organic materials during transport and atmospheric processing (Crippa et al.,
48 2013), and then its sizes, chemical compositions, mixing states and optical properties would
49 change greatly, leading to its influence in the atmosphere more uncertain (Jacobson, 2002;
50 Zaveri et al., 2010; Matsui, 2016; Budisulistiorini et al., 2017; Ma et al., 2012). Currently, the
51 influences of the complex chemical components on aerosol size and mixing state show large
52 regional differences due to the variations in the pollution sources, atmospheric formation
53 mechanism and meteorological conditions, which have been widely studied in an urban area
54 at a low altitude (Pratt et al., 2011; Liu et al., 2020a; Xu et al., 2017; Wang et al., 2022).
55 However, Liu et al (2020) have found that the migration or formation of low-volatile
56 component (such as nitrate and organic matter) could effectively be reduced due to
57 evaporation during the upward transportation process, which further alter the chemical
58 compositions and the particle sizes. The transportation of the aerosols to a relatively cleaner
59 environment prevails the formation of secondary chemicals at a high altitude (Liu et al.,
60 2020b). Therefore, a comprehensive investigation of the detailed characteristic of aerosol
61 formation and mixing states is required to understand their environmental effects in low-, and
62 high-altitude.

63 As a typical high-altitude region, the Tibetan Plateau (TP) has the highest and largest
64 mountain area in the world, which is the most sensitive and obvious indicator of climate
65 change in the entire Asian continent (Liu et al., 2017; Chen and Bordoni, 2014; Immerzeel et
66 al., 2010). Numerous studies have shown that the melting and retreat of glaciers in the TP
67 regions is accelerating in recent decades, largely attributed to anthropogenic emissions, such
68 as greenhouse gases and aerosols (Luo et al., 2020; Hua et al., 2019). Atmospheric aerosols
69 also can act as cloud condensation nuclei to impact the local hydrological cycles and
70 monsoon patterns by changing the microphysical properties and life span of clouds (Qian et
71 al., 2011; Gettelman et al., 2013; Kumar et al., 2017). The southern part of the TP is always
72 affected by the transport of more polluted air from South Asia along the mountain valleys,

73 especially during the pre-monsoon (i.e., March-May) with the southwest prevailing wind
74 (Chan et al., 2017; Zhao et al., 2017; Han et al., 2020). Most studies have focused on the
75 optical characteristics within the TP; however, only a few research has been conducted on
76 aerosol components.

77 Present researches on aerosol components over the TP mostly focus on exploring the
78 influence of light-absorbing carbon aerosols and dust particles on climate change by optical
79 or offline sampling methods (e.g., Wang et al., 2019a; Liu et al., 2021). There is a lack of
80 studies on the chemical composition, mixing states, and formation mechanism of aerosols in
81 the southeast margin and even the entire TP, especially using high-time resolved
82 measurements. Although time-integrated sampling with filter collection followed by
83 laboratory analyses has been widely adopted for the chemical characterization of aerosols (Li
84 et al., 2022a; Shen et al., 2015; Zhang et al., 2013), the drawbacks of the traditional approach
85 need to get attention, including the low time resolution, high detection limit, and time- and
86 labor-intensive procedures. Therefore, more advanced aerosol measurement equipment with
87 high-time resolution are developed, for example the aerosol chemical speciation monitor
88 (ACSM) and aerosol mass spectrometer (AMS) (Ng et al., 2011; Canagaratna et al., 2007) are
89 mainly achieved the online observation datasets of non-refractory submicron aerosol
90 (including the mass concentration of sulfate, nitrate, ammonium, chloride and organic; and
91 mass spectral of organic). This is beneficial to recognize the dynamic processes of source
92 emission of organic matter in the atmosphere (Du et al., 2015; Zhang et al., 2019a).
93 Meanwhile, aerosol time-of-flight mass spectrometry (ATOFMS) (Dall'Osto et al., 2014) and
94 single particle aerosol mass spectrometer (SPAMS) (Zhang et al., 2020) are popular for
95 characterizing atmospheric individual particles. These devices can determine the chemical
96 composition and size distribution of the particles in detail, such as the dynamic processes of
97 chemical aging, mixing state and transporting (Liang et al., 2022; Li et al., 2022b; Zhang et
98 al., 2019b). To the best knowledge, the advanced measurement device has not yet been
99 applied for the studies conducted in TP, leading to a lack of in-depth research on the PM_{2.5}
100 pollution in TP, especially in the southeastern margin, which hinders our understanding of the
101 distribution characteristics and formation mechanism of aerosol components in high-altitude
102 regions.

103 The southeastern margin of the TP is an important transitional zone between the
104 high-altitude TP and the low-altitude Yungui Plateau (Wang et al., 2019a; Zhao et al., 2017),
105 is an ideal place for investigating the impacts of pollutants transport and formation in the
106 high-altitude zone. In this study, continuous field observation of individual particles (SPAMS)
107 was deployed on the southeastern margin of the TP during the pre-monsoon period, to (i)
108 investigate the changes of chemical characteristics between transport and local fine particles
109 during pre-monsoon, (ii) determine the size distributions and mixing states of different
110 particle types, and (iii) assess the contributions of photooxidation and aqueous reaction to the
111 formation of the secondary species. These results would expand our understanding of the
112 chemical components, size distribution, mixing state and aging pathways of aerosols in the
113 high-altitude areas over the TP and surrounding areas.

114 **2 Methodology**

115 **2.1 Sampling site**

116 Intensive one-month field observation was deployed at the rooftop (~ 10 m above ground
117 level) of the Lijiang Astronomical Station, Chinese Academy of Sciences (3260 m above sea
118 level; 26°41'24"N, 100°10'48"E), Gaomeigu County, Yunnan Province, China, during the
119 pre-monsoon period (from April 14th to May 13th, 2018). The nearest residential area is the
120 Gaomeigu village (3–5 km away) with a small population size of 113 residents in 27
121 households. Villagers earn a living by farming (e.g., potato and autumn rape), and biomass is
122 the major domestic fuel (Li et al., 2016). The sampling site is surrounded by rural and
123 mountainous areas and has no obvious industry or traffic emissions. During the total
124 observation period, the average temperature (T) and relative humidity (RH) are $8.4 \pm 3.1^\circ\text{C}$
125 and $69\% \pm 21\%$, respectively. The wind speed (WS) is $2.2 \pm 1.2 \text{ m}\cdot\text{s}^{-1}$ with the prevailing
126 wind in the north and northeastern (Fig. S1).

127 **2.2 On-line instrument**

128 A detailed operational principle and the calibrations of the single-particle aerosol mass
129 spectrometer (SPAMS, Hexin Analytical Instrument Co., Ltd., Guangzhou, China) has been
130 described elsewhere (Li et al., 2011). Briefly, individual particles are drawn into SPAMS

131 through a critical orifice. The particles are focused and accelerated, then aerodynamically
132 sized by two continuous diode Nd: YAG laser beams (532 nm), subsequently desorbed and
133 ionized by a pulsed laser (266 nm) triggered exactly based on the velocity of the specific
134 particle. The generated of positive and negative molecular fragments are recorded with the
135 corresponding size of individual particles. In summary, a velocity, a detection time, and an
136 ion mass spectrum are recorded for each ionized particle, while there is no mass spectrum for
137 not ionized particles. The velocity could be converted to d_{va} based on a calibration using
138 polystyrene latex spheres (PSL, Thermo Scientific Corp., Palo Alto, USA) with predefined
139 sizes. The average ambient pressure is 690 hPa (in a range of 685–694 hPa) during the
140 measurements and calibration. A hollow silicone dryer was installed in front of the inlet. This
141 reduces the uncertainty of particle collection efficiency due to the changes of humidity in
142 sampled airs. Particles measured by SPAMS mostly are within the size range of vacuum
143 aerodynamic diameter (d_{va}) 0.2–2.0 μm . This SPAMS-specific size distribution is
144 semi-quantitative evaluated the relative concentration and contribution of each particle type,
145 mainly due to it largely dependence on the particle-detection efficiency (Allen et al., 2000;
146 Yang et al., 2017). The characteristics of SPAMS-specific size distribution are statistical
147 results, while the comparison of the relative distribution and number fraction of different
148 particle types in each size bin are significant.

149 Meteorological parameters, including the temperature ($^{\circ}\text{C}$), RH (%), WS ($\text{m}\cdot\text{s}^{-1}$), and
150 wind direction (WD) were continuously achieved using an automatic weather station (Model
151 MAWS201, Vaisala HydroMet, Helsinki, Finland) in a 5-min resolution, and the planetary
152 boundary layer (PBL) was acquired from the website (<https://rda.ucar.edu/datasets/ds083.2>,
153 last access: 17 April 2020) in a 1-hour resolution. Gaseous concentrations (ppbv) were
154 obtained using a multiple gas analyzer (Thermo Scientific Corp.), including ozone (O_3 , model
155 49i) and nitrogen oxides (NO_x , model 42i) in a 5-min resolution. The SPAMS and gas
156 analyzers are co-located in the same position, and the weather station was uncovered outside
157 ~ 5 m from the sampling house. Time series of SPAMS particles, gaseous concentrations (NO ,
158 NO_x , O_3 , and CO) and meteorological parameters (PBL, temperature, RH, WD, and WS)
159 were shown in Fig. S2.

160 **2.3 Individual particle classification**

161 During the observation period, a total of 461,876 ambient particles with the size (d_{va}) of
162 0.2–2.0 μm were collected, including 55,583 in Episode 1 (E1; from April 18th 08:00 local
163 time (LT) to April 19th 08:00 LT) and 62,110 in Episode 2 (E2; from April 26th 17:00 LT to
164 April 28th 02:00 LT). The analyzed particles are classified into 1,557 groups using an adaptive
165 resonance theory neural network (ART-2a) with a vigilance factor of 0.8, a learning rate of
166 0.05, and 20 iterations (Song et al., 1999). Finally, eight major particle clusters [i.e.,
167 potassium-rich (rich-K), biomass burning (BB), organic carbon (OC), Ammonium, aged
168 element carbon (EC-aged), Dust, sodium (Na)-potassium (K)-containing (NaK-SN), and iron
169 (Fe)-lead (Pb)-containing (Metal)] with distinct chemical patterns were manually combined,
170 which represent $\sim 99.7\%$ of the population of the detected particles. The remaining particles
171 are grouped as “Other”. The characteristics of the positive and negative mass spectra (MS) of
172 each particle type are shown in Fig. S3. A detailed description of classification criteria for
173 individual particles and the characteristic ion fragments for each particle type can be found in
174 Text S1. The criteria used for searching the some secondary species in the SPAMS datasets
175 are summarized in Table S2.

176 **2.4 Trajectory-related analysis**

177 To determine the influence of regional transport on different particles at the southeastern
178 margin of the TP, the trajectory clusters analysis was carried out using the 72-h backward air
179 mass trajectories at arrival heights of 500 m above ground level. The trajectories were
180 calculated with the Hybrid Single-Particle Lagrangian Integrated Trajectory model (Draxler
181 and Hess, 1998), and the meteorological data were obtained from the Global Data
182 Assimilation System (GDAS; <ftp://arlftp.arlhq.noaa.gov/pub/archives/gdas1>, last access: 6
183 April, 2022). The cluster analysis employs a Euclidean-oriented distance definition to
184 differentiate and cluster the major spatial features of the inputting trajectories. Details of the
185 trajectory clustering method can be found in Sirois and Bottenheim (1995). To investigate the
186 effects of transport on the chemical characteristic of the individual particles, trajectories with
187 particle number concentrations high than the 75th percentile are considered as pollution (Liu
188 et al., 2021).

189 3 Results and Discussion

190 3.1 Characteristics of particle composition

191 Table 1 summarizes the number concentrations, relative percentages and characteristic
192 ions of each particle type. The most dominant particle type in Gaomeigu during pre-monsoon
193 is rich-K, accounting for an average of 30.9% of the total resolved particles, followed by BB
194 (18.7%), OC (12.8%), Ammonium (11.9%), EC-aged (10.9%) and Dust (10.7%). Their
195 characteristics of mass spectrum and possible sources are described in supplemental
196 information of text S1 in detail. Similar to the results of some studies in urban areas, rich-K
197 or carbonaceous-containing type is the dominant particle type (15-50%) (Xu et al., 2018;
198 Wang et al., 2019b; Li et al., 2022). Combined with the previous studies and the
199 characteristics of the mass spectrum (Fig. S3a) in this study, the rich-K particles are
200 contributed by biomass burning and traffic emission, because that extensive works usually
201 identify abundant $^{39}\text{K}^+$ signals for biomass burning (Pratt et al., 2011; Chen et al., 2017),
202 while the presence of phosphate (m/z $^{79}\text{PO}_3^-$) indicates the vehicle exhaust (Yang et al., 2017).
203 The results of the correlation between seven variables (Fig. S4) show that rich-K type is
204 strongly correlated with Ammonium ($r=0.84$) and EC-aged ($r=0.90$) types, follow well
205 correlated with OC ($r=0.70$) and BB ($r=0.68$) types, further demonstrate that rich-K particles
206 type is from traffic emission and biomass burning, and is affected by secondary formation
207 during the atmospheric aging in southeastern TP. It is worth noting that few research have
208 captured the high proportion of Ammonium particles as shown in this study (Shen et al., 2017;
209 Xu et al., 2018), which is ascribed to the conversion of ammonia (NH_3) precursor emitted
210 from large-scale agricultural activities and mountain forest (Engling et al., 2011; Li et al.,
211 2013). It is necessary to point out that 60% of Ammonium particles contain signals of amine
212 fragment (m/z 58, $\text{C}_2\text{H}_5\text{NH}=\text{CH}_2^+$), implying their similar formation pathway (Zhang et al.,
213 2012). Moreover, the amine-containing particle represented 12.5% of the total ambient
214 particles, which is significantly higher than that in some urban areas at low altitudes (around
215 2%) (Cahi et al., 2012; Zhang et al., 2015; Li et al., 2017) but is comparable to that at
216 observed sites with high RH, or during fog and cloud events at a high altitude ($> 9\%$) (Roth et
217 al., 2016; Lin et al., 2019). This suggests that the formation of amines under high RH and fog

218 conditions might exist in the Gaomeigu area (with an altitude of 3260 m), for example, the
219 high relative fraction of amine-containing particle corresponds to a high RH (Fig. S5), and
220 the existence of amine sources govern the ammonium formations (Bi et al., 2016; Rehbein et
221 al., 2011). The relatively larger fraction of Dust particles is related to the short-time
222 occurrences of dust events in spring (Fig. S6), leading to a wide contribution ranging between
223 10% and 70% in the period of 19:00 LT on April 16th to 10:00 LT on April 17th.

224 Fig. 1 shows the diurnal variations of each particle type. The rich-K, BB and OC
225 particles decrease after midnight until 06:00 LT, possibly explained by the curtailment of
226 local traffic and biomass-burning activities at nighttime even though both the planetary
227 boundary layer (PBL) height and WS decrease (Fig. S7). Then, their concentrations rapidly
228 increase in the morning (around 07:00 LT) due to more pollutants from biomass burning and
229 traffic emissions at the upwind region. The increases of PBL height and WS also lead to the
230 transported of air pollutants from the surrounding regions to the sampling site (Liu et al.,
231 2021). At 11:00 LT, the particle concentrations sharply decrease till 16:00–17:00 LT, caused
232 by the pollutant dispersion with the continuing increases of the PBL height and WS.
233 Increasing trends are observed after 17:00 LT due to the pollutant accumulation with the
234 reduction of PBL height and WS. In contrast, the Ammonium, EC-aged and Dust particles
235 show a unimodal pattern of the daily diurnal variation (Fig. 1d–f). From 00:00 to 06:00 LT,
236 minor fluctuation of particle concentrations of Ammonium, EC-aged and Dust is observed for
237 these particle types. After that, their levels continuously elevate until ~11:00 LT due to the
238 regional transport, traffic emission and fugitive dust (Text S2). While the PBL height and WS
239 increase continuously, the Ammonium, EC-aged and Dust types begin to decline from 12:00
240 to 17:00 LT. The subsequent increases of these three types after 17:00 LT are attributed to the
241 reduction of PBL height, as a result of the accumulation of pollutants in the near-surface
242 atmosphere.

243 Based on the transport pathways, four air mass clusters are identified to investigate the
244 effect of regional transport on the major particle types (i.e., rich-K, BB, OC, Ammonium,
245 EC-aged and Dust) (Fig. 2). Cluster 1, 3 and 4 are originated from northeastern Myanmar,
246 accounting for 59.8%, 33.2% and 4.6% of the total trajectories, respectively. Cluster 1 had an

247 average percentage of 32.7%, 18.5%, 12.0%, 12.5%, 11.1% and 8.9%, respectively, on the
248 rich-K, BB, OC, Ammonium, EC-aged and Dust particles (Table S1). Cluster 3 and 4 have
249 the comparable contributions of OC (15.5% and 12.5%, *respectively*), increased of BB
250 (19.3% and 26.8%, *respectively*) and decreased of rich-K (26.8% and 25.2%, *respectively*),
251 Ammonium (10.4% and 7.7%, *respectively*) and EC-aged (7.7% and 6.3%, *respectively*), to
252 those of Cluster 1, but with a high contribution of Dust (16.6%), which refer Cluster 3 and 4
253 to as dust and biomass burning pollution. However, Cluster 1 is more influenced by
254 compound pollution, mainly including secondary formation, biomass burning, and traffic
255 emissions. The diurnal variations of the BB and OC fractions are similar which rapidly
256 elevate at 07:00 LT (Fig. S8) due to the increased contribution of biomass burning and traffic
257 emissions from Cluster 1, Ammonium and EC-aged particles (peak at 07:00 LT) caused by
258 the effect of Cluster 1 and 3 together. A stable diurnal variation of rich-K fraction is mainly
259 due to its large proportion and diverse sources. The similar diurnal trend of Clusters 3 and 4
260 are both associated with dust contributions, which decrease at 04:00 LT and increase at noon.
261 The increased nighttime particles could be attributed to the pollutant accumulation with the
262 decreased PBL height. Cluster 2 originate from the northeastern India and passes over
263 Bangladesh. This cluster accounts for only 2.4 % of the total trajectories, in which ~30.8%
264 and ~35.9% are mainly associated with the rich-K and BB particles, respectively. Although
265 Cluster 2 and 4 are composed of a small fraction of total trajectories (2.4% and 4.6 %,
266 respectively), BB and dust particles are identified as the major pollutants, suggesting
267 significant influences from India and northeastern Myanmar during the campaign.

268 A more in-depth investigation of the characteristics of the main particle types in the
269 southeastern Tibet Plateau was conducted during two episode periods when the number
270 concentration of particles was high (i.e., E1: from 08:00 LT April 18th to 08:00 LT April 19th,
271 2018; E2: 17:00 LT April 26th to 02:00 LT April 28th, 2018) (Fig. S6). Even though the two
272 episodes are contributed by Cluster 1, the chemical components show significant differences
273 (Table 1). During E1, the average fractions of the rich-K, BB, OC, Ammonium, EC-aged, and
274 Dust particle are 29.0%, 11.5%, 8.1%, 17.5%, 10.0% and 20.3%, respectively, different from
275 39.3%, 14.2%, 10.0%, 13.5%, 17.2%, and 1.3% respectively, during E2. It can be seen that

276 the major changed factor of the Dust particle is 93.6% lower during E2 than E1, whereas the
277 EC-aged particle shows a reversible of 72.0% higher during E2. Meanwhile, rich-K, BB and
278 OC particles also increase by 35.5%, 23.5% and 23.4% respectively during E1 compared to
279 E2. For the air mass clusters (Fig. S9), E1 and E2 exhibit minor differences, mostly
280 originating from northern Myanmar and the Sino-Burmese border, but not identical regions.
281 The Dust particles that are much lower during E2 than E1 could be explained by higher WS
282 (on average of 2.7 ± 1.0 m/s versus 0.4 ± 0.5 m/s) (Fig. S9) and PBL height (771 ± 717 m
283 versus 560 ± 549 m) (Fig. S10). The Dust particles are mainly formed by re-suspension in the
284 local areas. In addition, the quick thrown-up dust belongs to more coarse size particles, which
285 are out of the detection range of the SPAMS. However, due to the larger dust particles
286 deposited more easily under the low WS and the stagnant air conditions during E1, more
287 suspended dust particles of small size fall in the detection range of SPAMS. Moreover, the
288 increased PBL height and WS could speed up the transportation of pollutants from multiple
289 sources (e.g., traffic and biomass burning emissions) to the observation site, leading to
290 elevate the fraction of EC-aged, rich-K, BB, and OC particles during E2. The decreased
291 Ammonium fraction during E2 is potentially explained by the reductions in the secondary
292 pollutant formation with declines of RH (from $73.9\% \pm 23.9\%$ to $53.1\% \pm 14.9\%$), in
293 comparison to those during E1.

294 **3.2 Characteristics of SPAMS-specific size distribution and mixing state**

295 The SPAMS-specific size distributions of all particle types are shown in Fig. 3.
296 According to the characteristics of the average MS (Text S1 and Fig. S3), rich-K, BB, OC
297 and EC-aged particles originated from the similar sources of solid-fuel combustion or vehicle
298 emission. Their SPAMS-specific size distribution thus presents within a small-scale (~ 440 nm)
299 (Fig. S11a). However, the relative percentage of each particle type is distinct with different
300 size ranges, possibly due to the unique atmospheric processing. For example, as shown in Fig.
301 3a, the proportions of rich-K and BB types increases along with the increase in particle size
302 from 200 to 420 nm, then decrease. OC and EC-aged types are mainly distributed in
303 relatively small sizes, and their proportions gradually decrease when the size ranges become
304 larger. Ammonium and Dust types are mainly distributed in large sizes of ~ 600 nm (Fig. S11a).

305 The proportion of Ammonium particles gradually increases with the increase of particle size
306 and peaks at 740 nm, the relatively large SPAMS-specific size distribution is ascribed to the
307 intense atmospheric aging during regional transport (Text S1). The proportion of Dust
308 particles gradually increases with a size > 560 nm and peaks at 1.48 μm . This is consistent
309 with the fact that dust is a coarse particle, generally formed at the roadside and fly ash.

310 Compared with the SPAMS-specific size distribution of the total particles, the peak
311 values of the six main particle types show minor differences (< 80 nm) during the two
312 different episode periods (Fig. 11b,c). However, the percentage of the six particle types
313 distribute in wider size ranges during E2 than during E1 possibly due to the more intensive
314 atmospheric aging. Similarly, during the two episodes (Fig. 3b,c), the relatively high fraction
315 of the rich-K and BB particles are more affected by the primary emissions when their peak
316 value of SPAMS-specific size distribution concentrate at < 300 nm, and > 300 nm are more
317 related to the aging process (Li et al., 2022b; Bi et al., 2011). Relatively greater fluctuation
318 for the large-size fractions (> 1.1 μm) could be explained by the low particle concentration (a
319 number less than 20). It should be pointed out that further application of this method would
320 require a co-located particle-sizing instrument to scale the size-resolved particle detection
321 efficiency. Both particle composition and size-dependent are the predominant impacting
322 factors on the particle detection efficiency of the SPAMS (Wenzel et al., 2003; Yang et al.,
323 2017; Healy et al., 2013).

324 To investigate the mixing state of the secondary species in the six main particle types,
325 the number fractions of six secondary markers ($^{97}\text{HSO}_4^-$, $^{195}\text{H}(\text{HSO}_4)_2^-$, $^{62}\text{NO}_3^-$, $^{18}\text{NH}_4^+$,
326 $^{58}\text{C}_2\text{H}_5\text{NHCH}_2^+$ and $^{89}\text{HC}_2\text{O}_4^-$) are calculated (Fig. 4). The presences of amine (m/z
327 $^{58}\text{C}_2\text{H}_5\text{NHCH}_2^+$) and sulfuric acid (m/z $^{195}\text{H}(\text{HSO}_4)_2^-$) signals are possibly indicative of the
328 water uptake (Chen et al., 2019) and acidic property of the particles (Rehbein et al., 2011),
329 respectively. The mixing states are obtained by the ratio of the number concentration of the
330 selected ions to each particle type.

331 The most abundant of $^{97}\text{HSO}_4^-$ and $^{18}\text{NH}_4^+$ fraction are seen in Ammonium (99% and
332 94%, *respectively*) and EC-aged (92% and 31%, *respectively*) particles, whereas much low
333 fraction of $^{62}\text{NO}_3^-$ is found (2% and 7%, *respectively*). These suggest that ammonium sulfate
334 is not a predominant form instead of ammonium nitrate (Zhang et al., 2013). The high

335 contribution of $^{97}\text{HSO}_4^-$ in EC-containing particles also suggests a significant influence of
336 anthropogenically emitted sulfate precursors (e.g., SO_2) on the aging of EC-containing
337 particles at the high altitude (Peng et al., 2016; Zhang et al., 2017a). Meanwhile, relatively
338 high number fractions of $^{195}\text{H}(\text{HSO}_4)_2^-$ and $^{58}\text{C}_2\text{H}_5\text{NHCH}_2^+$ are also observed in Ammonium
339 (63% and 60%) and EC-aged (4% and 19%) particles. These abundant mixtures potentially
340 represent the high hygroscopicity of Ammonium and EC-aged particles, and their ability to
341 neutralize the acidic particles of Ammonium particle (Sorooshian et al., 2007). Then, a
342 moderate fraction of $^{97}\text{HSO}_4^-$ and $^{18}\text{NH}_4^+$ are seen on the rich-K (65%, 7%) and OC (56%,
343 4%) particles. In contrast, more $^{62}\text{NO}_3^-$ fraction contribute to the rich-K (38%) and OC (68%)
344 particles, mainly affected by vehicle emissions and biomass burning (Text S1). Following BB
345 (18%) and Dust (6%) particles are found in a relatively low fraction of $^{97}\text{HSO}_4^-$, while the
346 moderate $^{62}\text{NO}_3^-$ accounts for 45% of the BB particle but only 3% of the Dust particle.
347 Combined with the results of the minor $^{18}\text{NH}_4^+$ fraction (<1%) in BB and Dust particles
348 suggests a relatively low degree of aging. In addition, oxalate ($^{89}\text{HC}_2\text{O}_4^-$), a representative
349 component of secondary organic formation is mainly mixed with BB (13%) and rich-K (12%)
350 particles. This is because the substantial precursors of oxalic acid, including acetate
351 ($^{59}\text{C}_2\text{H}_3\text{O}_2^-$), methylglyoxal ($^{71}\text{C}_3\text{H}_3\text{O}_2^-$), glyoxylate ($^{73}\text{C}_2\text{HO}_3^-$), are emitted from biomass
352 burning, and then oxalate heterogeneously formed in BB related particles (Zhang et al.,
353 2019b; Zauscher et al., 2013). A relatively low fraction (<5%) of oxalate-containing particles
354 in OC, Ammonium, EC-aged and Dust particles is potentially limited by the contributions of
355 precursor oxalic acid.

356 Compared to the mixing state of individual particle in urban or suburban areas that are
357 located close to emission sources (Chen et al., 2016; Dall'Osto and Harrison, 2012; Zhang et
358 al., 2017a; Li et al., 2022b), the high fractions of sulfate and ammonium at the high altitude
359 area demonstrate a high degree of aging of the individual particles, whereas the low fraction
360 of nitrate with high volatility indicates its loss during transportation processing.

361 The number fractions of six markers in the four trajectories were used to further
362 investigate the impacts of regional transport. As shown in Fig. 5(a,c), the dominant mixing
363 ion types in each particles (except for Dust) are similar among the four clusters. For Cluster 1,
364 the number fractions of $^{97}\text{HSO}_4^-$ and $^{89}\text{HC}_2\text{O}_4^-$ have larger values in five particle types

365 (except for Dust type) than those in other trajectories. Similar to Cluster 1, Cluster 3 and 4 are
366 impacted by regional transport from northeastern Myanmar, the fractions of the six markers
367 also similar in OC, Ammonium, and EC-aged types. However, $^{97}\text{HSO}_4^-$ in Cluster 3 and 4 is
368 reduced in rich-K, BB and Dust types, while $^{62}\text{NO}_3^-$ is increased in rich-K and decreased in
369 Dust types, compared with Cluster 1. As discussed in Section 3.1, these results demonstrate
370 that the aging degree of Cluster 3 and 4 might be lower than that of Cluster 1. For Cluster 2,
371 the fraction of $^{97}\text{HSO}_4^-$ is obviously decreased in rich-K, BB and EC-aged types but slightly
372 increased in Dust type (Fig. 5f). Such pattern inverse the observations in rich-K, OC and Dust
373 types for $^{62}\text{NO}_3^-$ ions. These variations in Cluster 2 are more likely due to the influences of
374 biomass-burning activities from surrounding the sampling site, rather than regional transport.
375 Furthermore, Cluster 2 is associated with regional transport from northeastern India along the
376 afternoon to nighttime (from 15:00 LT on 11 May to 07:00 LT on 12 May), which is
377 favorable to the nitrate formation N_2O_5 by heterogeneous hydrolysis (Wang et al., 2017; Ding
378 et al., 2021). However, these cases are infrequent, as only 2% of trajectories are associated
379 with Cluster 2.

380 During E1, more than 50% of $^{97}\text{HSO}_4^-$ fractions are mixed in the rich-K (81%), OC
381 (62%), Ammonium (100%), EC-aged (98%) particles (Fig. S12), low in BB (37%) and Dust
382 (4%) particles. Dissimilar with E1, the number fraction of $^{97}\text{HSO}_4^-$ in Dust increases to 34%
383 during E2, potentially associated with the enhancement by secondary formation during
384 regional transport. However, the mixing state of $^{195}\text{H}(\text{HSO}_4)_2^-$, $^{62}\text{NO}_3^-$, NH_4^+ and oxalate
385 fractions are similar between the two episode events. The $^{58}\text{C}_2\text{H}_5\text{NHCH}_2^+$ fractions are
386 significantly higher in E2 than E1 for Ammonium (67% versus 31%) and EC-aged particles
387 (48% versus 17%), due to the relatively higher hygroscopic behavior (i.e., RHs) (Sorooshian
388 et al., 2007).

389 **3.3 Formation process of the high number concentration particle episodes**

390 Photochemical oxidation and aqueous-phase reaction are the key formation pathways of
391 secondary species (Link et al., 2017; Xue et al., 2014; Jiang et al., 2019). The oxidant O_x (O_3
392 + NO_2) concentration and RH usually serve as indicators of the degree of photochemical
393 oxidation (Wood et al., 2010) and aqueous-phase reaction (Ervens et al., 2011), *receptively*,
394 though the current O_x and RH conditions obtained using the in-situ measurement are not

395 indicative of the past conditions experienced by the particle. Thus, the relative number
396 fractions of $^{43}\text{C}_2\text{H}_3\text{O}^+$, $^{89}\text{HC}_2\text{O}_4^-$, $^{62}\text{NO}_3^-$, $^{97}\text{HSO}_4^-$ and $^{18}\text{NH}_4^+$ -containing particles to the total
397 detected particles were selected to provide a rough speculative of the secondary formation
398 mechanism in TP ambient conditions (Liang et al., 2022). The correlations of the number
399 fraction of each secondary species with the O_x concentrations (O_x) during daytime (from
400 06:00 to 20:00 LT) and RH during nighttime (from 20:00 to 06:00 LT next day) are used to
401 reveal the formation pathways during the two episodes (Li et al., 2022).

402 As illustrated in Fig. 6, for E1, $^{43}\text{C}_2\text{H}_3\text{O}^+$, $^{89}\text{HC}_2\text{O}_4^-$, $^{97}\text{HSO}_4^-$, and $^{18}\text{NH}_4^+$ show
403 significant negative linear correlations with O_x ($p < 0.01$), and the correlation strengths range
404 from moderate to strong ($r = -0.51$ to -0.81). However, the $^{62}\text{NO}_3^-$ fraction shows an upward
405 trend with an insignificant correlation ($r = 0.33$, $p > 0.05$) with the increase in O_x
406 concentration. For E2, $^{43}\text{C}_2\text{H}_3\text{O}^+$ shows weak correlation with O_x ($r = 0.37$, $p > 0.05$), but
407 strong correlations with $^{89}\text{HC}_2\text{O}_4^-$, $^{97}\text{HSO}_4^-$, and $^{18}\text{NH}_4^+$ ($r = 0.81\sim 0.92$, $p < 0.01$). It should
408 be noted that $^{62}\text{NO}_3^-$ has a strong negative correlation ($r = -0.85$, $p < 0.01$) with O_x . In general,
409 the opposite linear relationship between secondary aerosol and O_x during E1 and E2 might be
410 influenced by reasons of i) the relatively low secondary formations because of the small
411 amount of precursors emitting from anthropogenic activities around the sampling site (Li et
412 al., 2016); ii) higher dilution rate of the particles formed in the atmosphere with the rapid rise
413 of PBL height during E1 than E2 (Fig. S13a); iii) the degrees of contributions of regional
414 transport due to the low WS ($0.5 \pm 0.6 \text{ m s}^{-1}$) during E1 and the high WS ($3.1 \pm 1.0 \text{ m s}^{-1}$)
415 during E2, respectively (Fig. S9). Therefore, for E1, the increases of NO_3^- fraction could be
416 influenced by the local nitrate formation, while the declines of other secondary components
417 should be ascribed to the reduced contribution of regional transport. For E2, the decreased of
418 NO_3^- fraction could be caused by the relatively higher volatilization loss of nitrate than other
419 components through the regional transport. Additionally, previous study proves that the
420 formations of organic nitrate species (such as $^{27}\text{CHN}^+$, $^{30}\text{NO}^+$, $^{43}\text{CHO}_1\text{N}^+$ and CHO_xN^+)
421 through the $\text{NO}+\text{RO}_2$ pathway dominate 80% of the total nitrate production in tropical
422 forested regions during summertime (Alexander et al., 2009). Aruffo et al (2022) also found
423 that low NO_x (i.e. $< 6 \text{ ppbv}$), compared to $2.3 \pm 0.8 \text{ ppbv}$ in this study, could even promote
424 the particle-phase partitioning of the lower volatility of organonitrates. These results suggest

425 that the secondary organic species have different formation capacities through
426 photo-oxidation reactions, among which the rate of HSO_4^- formation (slop=0.017) is the
427 highest. Increased with O_x concentration during E2, the concentration levels of secondary
428 organic species of $\text{C}_2\text{H}_3\text{O}^+$ (18%-28%) imperceptibly rise, while the oxalate fraction
429 significantly increase by 7%-20%.

430 Considering that the oxalate is abundant mixed in rich-K (14%), BB (15%), EC-aged
431 (5%), and Dust (6%) particles in Cluster 1 (Fig. 5), and the increased contributions of rich-K
432 (39.3%), BB (14.2%) and EC-aged (17.2%) types during E2 (Table 1), the apparent formation
433 of oxalate might be due to the enhancement of regional transport. Particularity, this presents
434 the nearby biomass burning and combustion activities produce more precursor species of
435 oxalate (Sullivan et al., 2007; Kundu et al., 2010; Zhang et al., 2017b).

436 Fig. 7 illustrates that the number fractions of $^{43}\text{C}_2\text{H}_3\text{O}^+$, $^{89}\text{HC}_2\text{O}_4^-$, $^{97}\text{HSO}_4^-$, and $^{18}\text{NH}_4^+$
437 have moderate to strong positive correlations with RH ($r = 0.70\sim 0.81$, $p < 0.01$ or 0.05) in the
438 nighttime during the two episodes, except that $^{43}\text{C}_2\text{H}_3\text{O}^+$ during E2 ($p = 0.48$) and $^{89}\text{HC}_2\text{O}_4^-$
439 during E1 ($p = 0.12$). Furthermore, $^{62}\text{NO}_3^-$ fraction has no obvious changes with insignificant
440 correlation with RH during E1 ($p = 0.43$) and presents a moderate negative correlation with
441 RH ($r = 0.69$, $p < 0.01$) during E2. As shown in Fig. 7e, the highest aqueous formation rate of
442 HSO_4^- is mainly due to the properties of low volatile and high hygroscopic of sulfate (Wang
443 et al., 2016; Zhang et al., 2019c; Sun et al., 2013). Compared with that during E2
444 (slop=0.014), the decreased formation rate of HSO_4^- during the E1 (slop=0.009) may be
445 because the decreases of aerosol acidity in higher RH $> 80\%$ (Huang et al., 2019; Meng et al.,
446 2014; Tian et al., 2021). And the increased contributions of regional transport due to the high
447 WS ($2.4 \pm 0.8 \text{ m s}^{-1}$) during E2 are compared with the low WS ($0.08 \pm 0.08 \text{ m s}^{-1}$) during E1
448 (Fig. S9). The fair production rate of NH_4^+ during the E1 (slop= 0.005) and E2 (slop=0.006)
449 demonstrate that an aqueous-phase reaction could effectively promote ammonium formation.
450 Meanwhile, a slightly larger slop of NH_4^+ during E2 could be also affected by the increased
451 contributions of regional transport. Compared with those during E1, the inverse generation
452 rates of two secondary organic species (i.e., $\text{C}_2\text{H}_3\text{O}^+$ and HC_2O_4^-) during E2 are possibly
453 caused by the different formation pathways with a variety of RH levels or distinct regional
454 transports. For example, $\text{C}_2\text{H}_3\text{O}^+$ shows a strong correlation with RH ($r = 0.70$, $p < 0.05$)

455 during E1 (slop=0.003) but has insignificant correlation during E2. This could be explained
456 by high RHs that could effectively promote secondary organic formation during E1. In
457 addition, the HC_2O_4^- fraction increases slightly (9.7-13.1%) during E1 is potentially ascribed
458 to more abundant Dust-type particles (20.3%) which compose of high calcium (Ca) (Fig. S14)
459 that favor the formation of metal oxalate complexes (i.e., Ca oxalate). At high RHs ($93.4 \pm$
460 7.6%), if oxalate ions are dissolved in the aqueous phase with the presence of Ca ions, the Ca
461 oxalate complexes can precipitate because of their low hygroscopic and insoluble natures
462 (Furukawa and Takahashi, 2011). This could offset the oxalate formation in the
463 aqueous-phase reaction. However, significant linear increases (slop=0.003) with RH ($r = 0.81,$
464 $p < 0.01$) during E2 demonstrate that the aqueous-phase reaction effectively promotes the
465 oxalate formation (Cheng et al., 2017; Meng et al., 2020). No obvious change and
466 insignificant correlation between $^{62}\text{NO}_3^-$ and RH are found during E1, potentially attributed
467 to the decreases of NO_2 concentration (3.7 ± 0.4 ppbv) in the local atmosphere. Meanwhile,
468 high RHs could promote organonitrates formation (Fang et al., 2021; Fry et al., 2014). The
469 linearity between $^{62}\text{NO}_3^-$ and RH ($r = 0.69, p < 0.01$) significantly decreases during E2,
470 mostly due to the losses of the volatile compound through the regional transport (Fig. S15).

471 **4 Conclusions**

472 This study presents the chemical composition, size distribution, mixing state and
473 secondary formation of individual particles in the southeastern margin of TP, China during
474 the pre-monsoon season using a high-resolution SPAMS. The finding shows that the rich-K
475 (30.9%) and BB types (18.7%) are the two dominant aerosol particles in the remote area;
476 followed by the OC (12.8%), Ammonium (11.9%), EC-aged (10.9%), and Dust (10.7%) types;
477 the NaK-SN, Metal and Others particle types contributed 0.3–2.8% to the total ambient
478 particles. By interpreting the mass spectra and diurnal trends, the major particle types are
479 mainly from traffic emission, biomass burning, secondary formation and fly ash, while the
480 dynamics of the PBL height could also affect their contributions. The observed change in the
481 number fraction of the particle types was mainly influenced by air mass (97.61% of the total
482 trajectories) from northeastern Myanmar, and significantly contributed to rich-K and BB
483 types. The particle types show distinct size distributions. The two critical particle types of
484 rich-K and BB appear in a unimodal pattern, the fractions of OC and EC-aged gradually

485 decrease with the increase of the particle sizes, but Ammonium and Dust types show the
486 opposite. Sulfate is the major secondary species and is highly mixed with rich-K, Ammonium
487 and EC-aged types. Nitrate has a relatively low mixing ratio due to its higher volatility than
488 sulfate during regional transportation, except for BB and OC types. During the entire study
489 campaign, two episodes with the high number concentration of particles occur but with
490 significant differences in each particle fraction due to the different meteorological conditions
491 (RH, WS, etc.). Meanwhile, the different meteorological conditions also lead to an inverse
492 linear correlation between the indicators of secondary formation, including $C_2H_3O^+$, $HC_2O_4^-$,
493 NH_4^+ , NO_3^- and HSO_4^- . and O_x (O_3+NO_2) during episode 1 and 2 periods; however, they
494 present a positive linear correlation with relative humidity (RH), except for NO_3^- shown the
495 negative linear correlation with RH due to the low precursors concentration and potential
496 organonitrates formation. These results demonstrated that the capacity of atmospheric aging
497 of photo-oxidation and aqueous reaction have complex influencing factors. Although the
498 detailed formation pathways and their percentage contributions to secondary species are not
499 quantitatively estimated in this study, our results have important implications for the various
500 possibilities affecting the characteristic of chemical components, size distribution, mixing
501 states, and formation mechanism of aerosols in the southeast TP. More depth investigations
502 concerning the evolution mechanisms of secondary aerosols are encouraged since TP is a
503 significant regulator to global climate change.

504 *Data availability.* The data presented in this study are available at the Zenodo data archive
505 <https://doi.org/10.5281/zenodo.7336857>.

506

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

508

509 *Author contributions.* QW and JC designed the campaign. WR conducted field measurements.
510 LL, QW, JT, and YZ made data analysis and interpretation. LL and QW wrote the paper. All
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512

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865 <https://doi.org/10.1016/j.scitotenv.2017.04.101>, 2017.

866

867 Table 1. The number concentrations, average percentages and characteristic ions of nine types
 868 of particles during the entire campaign, and the average percentages of the major six particle
 869 types during two episodes.

Type	Number count	Fraction in total (%)	Episode 1 (%)	Episode 2 (%)	Tracer ions
rich-K	151040	30.9	29.0	39.3	$^{39}\text{K}^+$, $^{26}\text{CN}^-$, $^{42}\text{CNO}^-$, $^{46}\text{NO}_2^-$, $^{62}\text{NO}_3^-$, $^{97}\text{HSO}_4^-$
BB	91322	18.7	11.5	14.2	$^{39}\text{K}^+$, levoglucosan ($^{45}\text{CHO}_2^-$, $^{59}\text{C}_2\text{H}_3\text{O}_2^-$, $^{71}\text{C}_3\text{H}_3\text{O}_2^-$, $^{73}\text{C}_3\text{HO}_3^-$), $^{26}\text{CN}^-$, $^{35,37}\text{Cl}^-$, $^{42}\text{CNO}^-$, $^{46}\text{NO}_2^-$, $^{62}\text{NO}_3^-$, $^{97}\text{HSO}_4^-$
OC	62446	12.8	8.1	10.0	$^{27}\text{C}_2\text{H}_3^+$, $^{37}\text{C}_3\text{H}^+$, $^{38}\text{C}_3\text{H}_2^+$, $^{39}\text{K}^+/\text{C}_3\text{H}_3^+$, $^{43}\text{C}_2\text{H}_3\text{O}^+$, $^{51}\text{C}_4\text{H}_3^+$, $^{26}\text{CN}^-$, $^{42}\text{CNO}^-$, $^{46}\text{NO}_2^-$, $^{62}\text{NO}_3^-$, $^{97}\text{HSO}_4^-$
Ammonium	58317	11.9	17.5	13.5	$^{12}\text{C}^+$, $^{18}\text{NH}_4^+$, $^{39}\text{K}^+$, $^{58}\text{C}_2\text{H}_5\text{NHCH}_2^+$, $^{97}\text{HSO}_4^-$, $^{195}\text{H}(\text{HSO}_4)_2^-$
EC-aged	53337	10.9	10.0	17.2	C_n^\pm (n=1~5), $^{39}\text{K}^+$, $^{97}\text{HSO}_4^-$
Dust	52533	10.7	20.3	1.3	$^{40}\text{Ca}^+$, $^{56}\text{CaO}^+$, $^{16}\text{O}^-$, $^{17}\text{OH}^-$, $^{76}\text{SiO}_3^-$, $^{79}\text{PO}_3^-$
NaK-SN	13726	2.8	na	na	$^{23}\text{Na}^+$, $^{39}\text{K}^+$, $^{62}\text{NO}_3^-$, $^{97}\text{HSO}_4^-$
Metal	4672	1.0	na	na	$^{51}\text{V}^+$, $^{56}\text{Fe}^+$, $^{64,66,68}\text{Zn}^+$, $^{206,207,208}\text{Pb}^+$
Others	1580	0.3	na	na	No obvious characteristic peaks

870

871 **Figure captions:**

872 **Figure 1.** Box and whisker diurnal plots of the number concentration of the main particle
873 types (a) rich-Potassium (K), (b) Biomass burning (BB), (c) Organic carbon (OC), (d)
874 Ammonium, (e) Element carbon (EC)-aged, (f) Dust in hourly resolution. The lower, middle,
875 and upper lines of the boxes denote the 25th, 50th, and 75th percentiles. The lower and upper
876 whiskers represent the 10th and 90th percentiles, respectively. Average values are shown in
877 white dots.

878 **Figure 2.** Maps of the mean HYSPLIT back trajectory clusters (72 h) at the height of 500 m
879 during the whole field observation. Embedded pie chart represents the relative fraction of
880 each particle type in the four clusters.

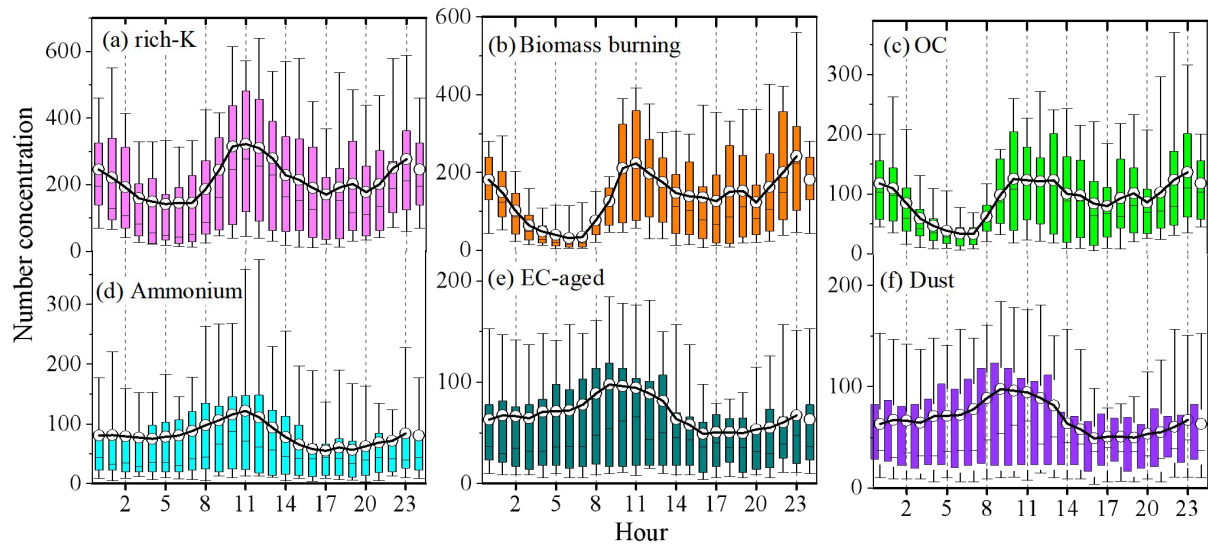
881 **Figure 3.** SPAMS-specific size distributions of the relative number fraction (%) of the total
882 particles for nine groups during (a) the total sampling campaign and two episodes of (b) E1
883 and (c) E2.

884 **Figure 4.** Number fractions of secondary markers associated with the six particle types
885 (rich-K, BB, OC, Ammonium, EC-aged, Dust) during the whole observation. Secondary
886 species include sulfate ($^{97}\text{HSO}_4^-$), sulfuric acid ($^{195}\text{H}(\text{HSO}_4)_2^-$), nitrate ($^{62}\text{NO}_3^-$), ammonium
887 ($^{18}\text{NH}_4^+$), amine ($^{58}\text{C}_2\text{H}_5\text{NHCH}_2^+$), and oxalate ($^{89}\text{HC}_2\text{O}_4^-$) ions.

888 **Figure 5.** Number fractions of secondary markers associated with the six particle types (i.e.,
889 rich-K, BB, OC, Ammonium, EC-aged, and Dust) in four clusters. Secondary species
890 abbreviations as in Figure 4.

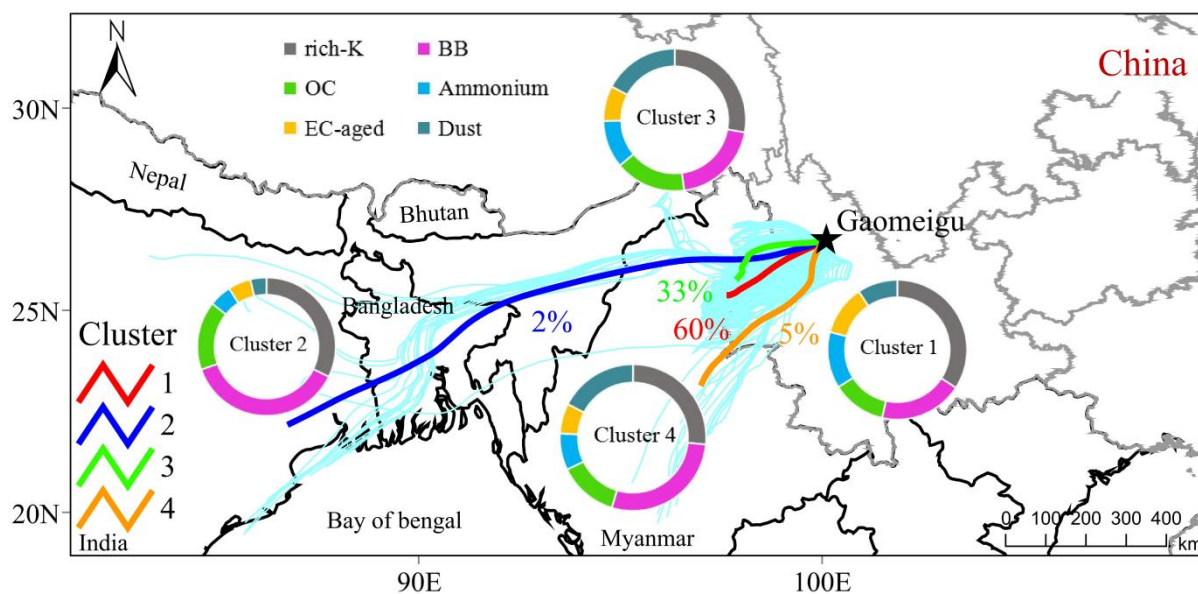
891 **Figure 6.** Correlations between the relative number fractions of the secondary species (a)
892 $^{43}\text{C}_2\text{H}_3\text{O}^+$, (b) $^{89}\text{HC}_2\text{O}_4^-$, (c) $^{18}\text{NH}_4^+$, (d) $^{62}\text{NO}_3^-$, (e) $^{97}\text{HSO}_4^-$ and O_x concentration during E1
893 (blue square) and E2 (red dot).

894 **Figure 7.** Correlations between the relative number fractions of the secondary species (a)
895 $^{43}\text{C}_2\text{H}_3\text{O}^+$, (b) $^{89}\text{HC}_2\text{O}_4^-$, (c) $^{18}\text{NH}_4^+$, (d) $^{62}\text{NO}_3^-$, (e) $^{97}\text{HSO}_4^-$ and relative humidity (RH)
896 during E1 (cyan dot) and E2 (orange square).



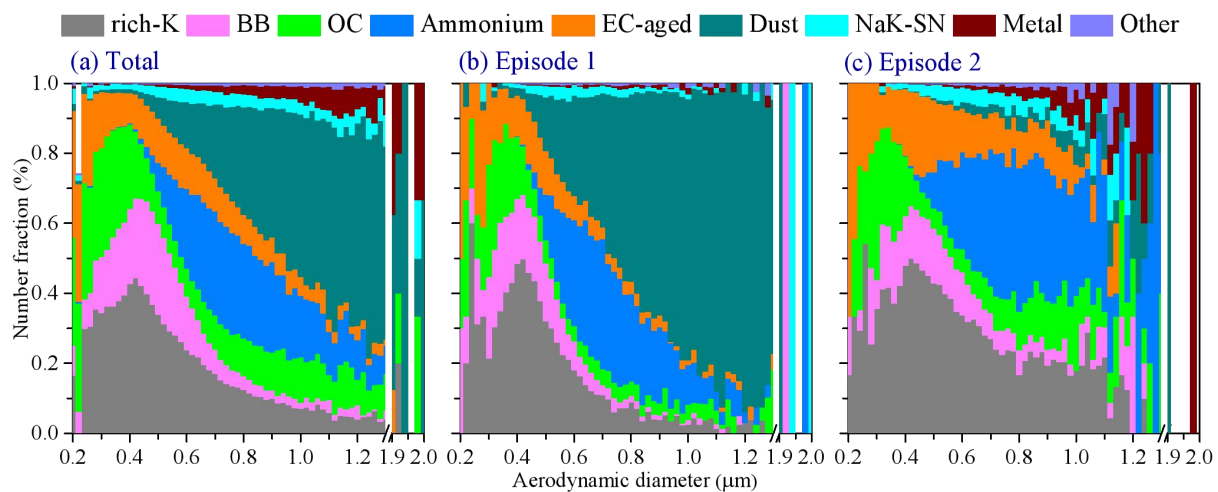
897

898 Figure 1. Box and whisker diurnal plots of the number concentration of the main particle types (a)
 899 rich-Potassium (K), (b) Biomass burning (BB), (c) Organic carbon (OC), (d) Ammonium,
 900 Element carbon (EC)-aged, (f) Dust in hourly resolution. The lower, middle, and upper lines of the boxes denote the
 901 25th, 50th, and 75th percentiles. The lower and upper whiskers represent the 10th and 90th percentiles,
 902 respectively. Average values are shown in white dots.



903

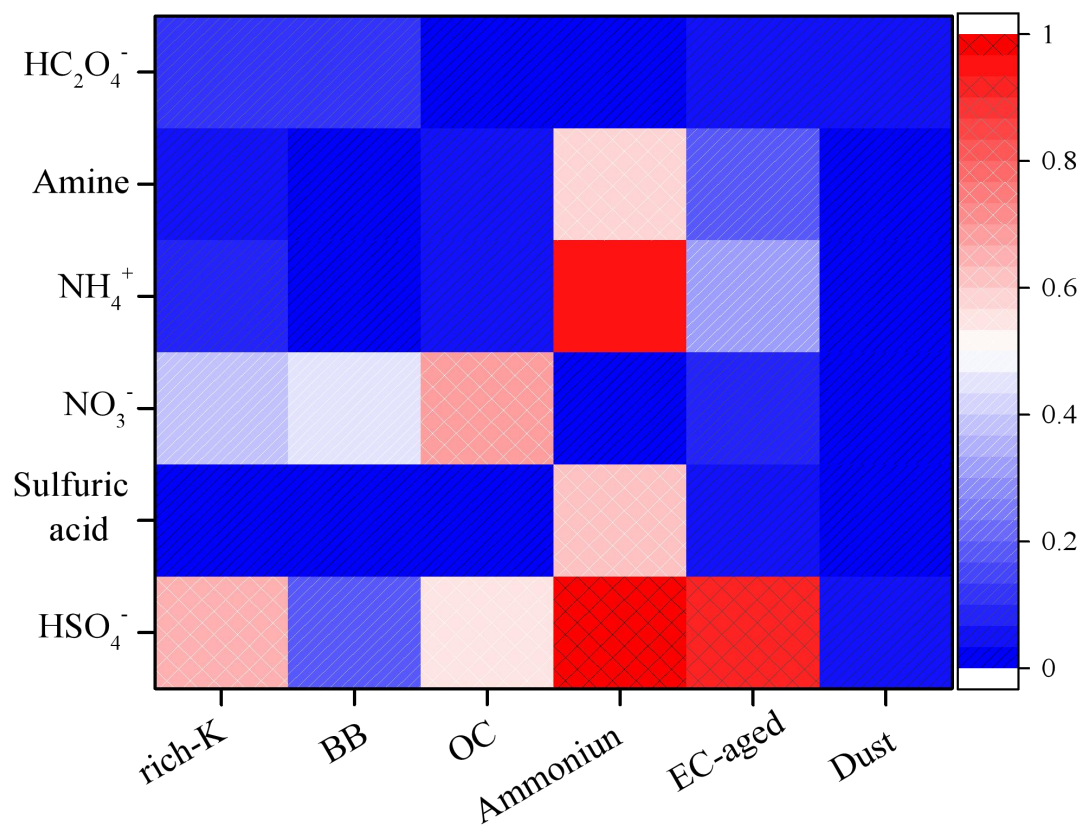
904 Figure 2. Maps of the mean HYSPLIT back trajectory clusters (72 h) at the height of 500 m during the
 905 whole field observation. Embedded pie chart represents the relative fraction of each particle type in the
 906 four clusters.



907

908 Figure 3. SPAMS-specific size distributions of the relative number fraction (%) of the total particles for

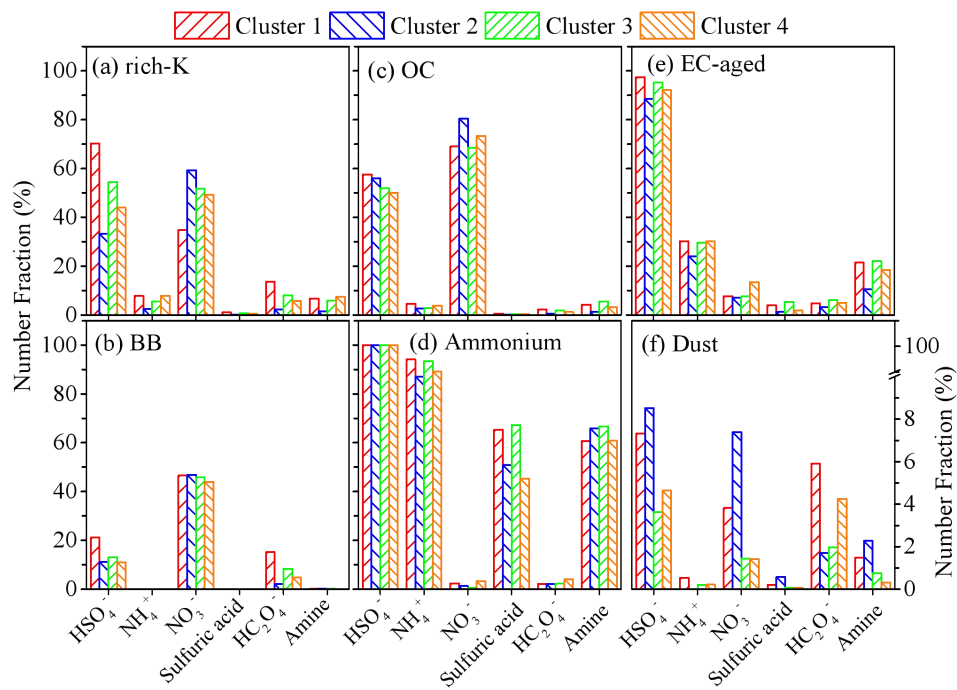
909 nine groups during (a) the total sampling campaign and two episodes of (b) E1 and (c) E2.



910

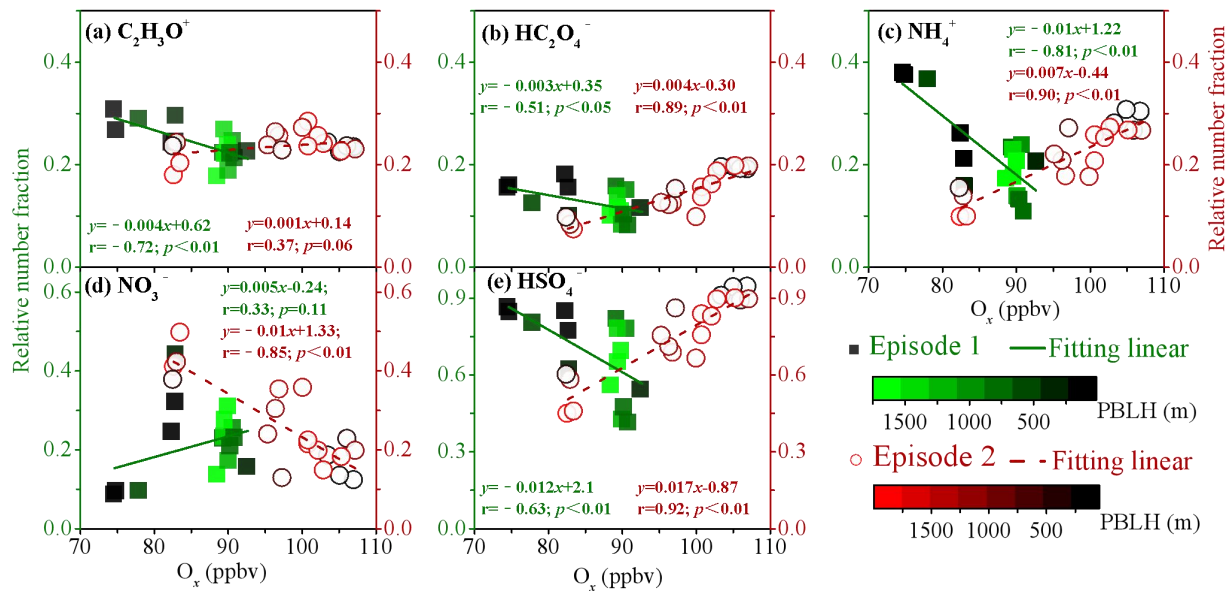
911 Figure 4. Number fractions of secondary markers associated with the six particle types (rich-K, BB, OC,
 912 Ammonium, EC-aged, Dust) during the whole observation. Secondary species include sulfate (⁹⁷HSO₄⁻),
 913 sulfuric acid (¹⁹⁵H(HSO₄)₂⁻), nitrate (⁶²NO₃⁻), ammonium (¹⁸NH₄⁺), amine (⁵⁸C₂H₅NHCH₂⁺), and oxalate
 914 (⁸⁹HC₂O₄⁻) ions.

915



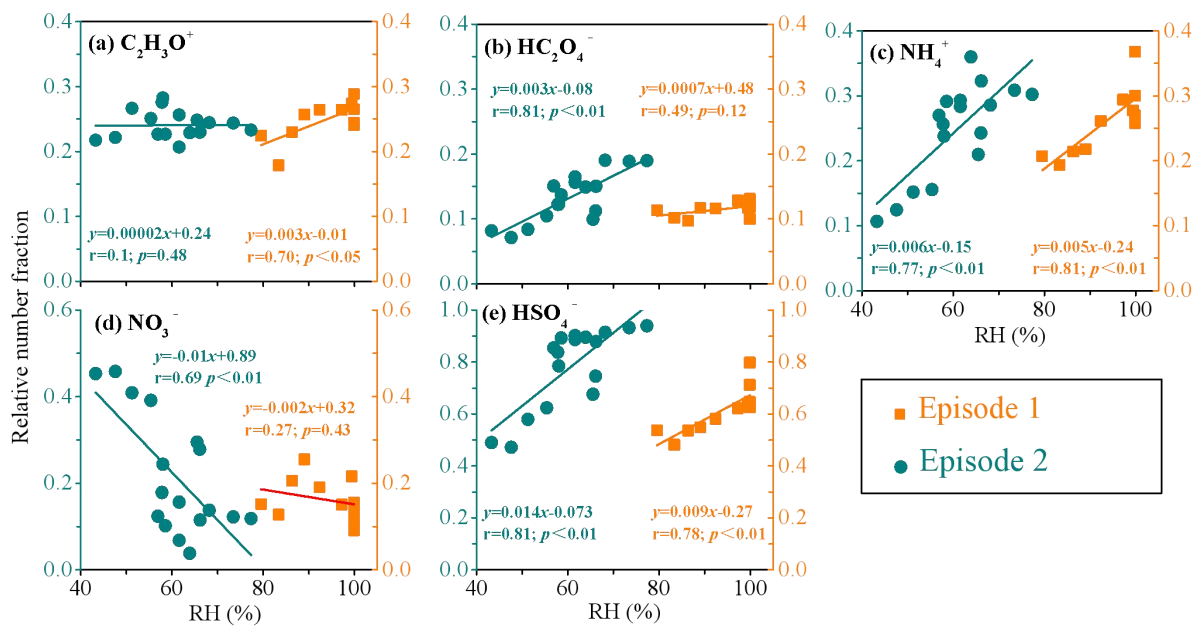
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 919

Figure 5. Number fractions of secondary markers associated with the six particle types (i.e., rich-K, BB, OC, Ammonium, EC-aged, and Dust) in four clusters. Secondary species abbreviations as in Figure 4.



920

921 Figure 6. Correlations between the relative number fractions of the secondary species (a) $^{43}C_2H_3O^+$, (b)
 922 $^{89}HC_2O_4^-$, (c) $^{18}NH_4^+$, (d) $^{62}NO_3^-$, (e) $^{97}HSO_4^-$ and O_x concentration during E1 (blue square) and E2 (red
 923 dot).



924

925 Figure 7. Correlations between the relative number fractions of the secondary species (a) $^{43}C_2H_3O^+$, (b)
 926 $^{89}HC_2O_4^-$, (c) $^{18}NH_4^+$, (d) $^{62}NO_3^-$, (e) $^{97}HSO_4^-$ and relative humidity (RH) during E1 (cyan dot) and E2
 927 (orange square).

928