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1	Chemical compositions of fog and precipitation at Sejila Mountain in the										
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27 Abstract:

Chemical compositions of fog and rain water were measured between July 2017 and September 2018 at Sejila Mountain, southeast Tibet, where fog events frequently occurred in original fir forests at altitude 3950 m. Fog water samples were collected using a Caltech Active Strand Cloud Collector (CASCC), and rain samples were collected using a precipitation gauge. Differences were observed between fog water and rain composition for most analyzed ions. Ion abundance in fog water was $Ca^{2+} >$

 $Cl^{-} > Na^{+} > SO_{4}{}^{2-} > Mg^{2+} > NH_{4}{}^{+} > K^{+} > NO_{3}{}^{-} \text{ whereas an order of } Ca^{2+} > Na^{+} > Cl^{-} > Na^{+} > Na^$ 34 $Mg^{2+} > SO_4^{2-} > NO_3^{-} > K^+ > NH_4^+$ was observed for rain water. All ion concentrations 35 were higher in fog water than in rain water. Additionally, Ca²⁺ was the dominant 36 cation in both fog and rain samples, accounting for more than half of all measured 37 cations. NH_4^+ and SO_4^{2-} concentrations were notable for being higher in fog than rain 38 water when compared with other ions. For trace elements, Al, As, Mn and Se were the 39 most abundant elements in fog water; only Al and As were detected in rain water. 40 41 Seventy-two hour back-trajectory analysis showed that air masses during fog and/or rain events mainly came from the south of Sejila Mountain. Spearman correlation 42 analysis and source contribution calculations indicated that both marine and terrestrial 43 sources contributed to the observed ion concentrations. Considering the higher 44 concentrations of NH_4^+ and higher ratio of NH_4^+/NO_3^- measured in fog than in rain, 45 we suggest that quantification of fog nitrogen deposition and its ecological effect in 46 this area should be given more attention. 47

Keywords: Fog water; Ion concentration; Trace elements; Emission source; Southeast
Tibet

50 **Capsule:** Comparison of water-soluble ions in fog and rainwater in southeast Tibet.

51

52 **1. Introduction**

53 Clouds and fogs play an important role in processing pollutants and other trace chemical species in the atmosphere. Scavenging of particles and soluble trace gases 54 by cloud and fog droplets, followed by direct droplet deposition to the ground or 55 incorporation into precipitation, represent important pathways for pollutant removal 56 from the atmosphere (Liu et al., 2004). In order to assess the impacts of cloud and fog 57 on various ecological environments, numerous studies of cloud and fog water 58 chemical composition have been conducted in Europe (e.g., Collett et al., 1993a; Blas 59 et al., 2010; Blas et al., 2008; Malcolm et al., 2003; Giulianelli et al., 2014), North 60 America (Waldman et al., 1985; Weathers et al., 1988; Collett et al., 1990, 1991a; 61 62 Saxena and Lin, 1990; Straub et al., 2012, Herckes et al., 2015, Templer et al., 2015, Straub, 2017), South America (Weathers et al., 2000; Strater et al. 2010), and Asia 63

(Aikawa et al., 2005; Kim et al., 2006; Beiderwieden et al., 2007; Sheu and Lin, 2011; 64 Wang et al., 2011; Shen et al., 2012). In many cases, clouds and fogs have high ion 65 concentrations and can be 5-20 times more concentrated than rain water (Anderson et 66 al., 1999; Beiderwieden et al., 2007). Although the direct hydrologic input from fogs 67 and intercepted clouds is typically much lower than from rain and snow, the higher 68 solute concentrations mean that cloud and fog deposition should not be ignored when 69 considering nutrient and pollutant input, especially for high elevation ecosystems 70 71 (Collett et al., 1990, 1993a; Aleksic et al., 2009).

Montane cloud forests are defined as forests that are frequently covered in cloud 72 or mist (Hamilton et al., 1995). In the forest ecosystems of higher mountains, fog 73 deposition has been recognized as an important component of hydrological and 74 nutrient cycling (Dawson et al., 1998; Elias et al., 1995; Weathers et al., 2000). For 75 example, Lovett (1982) found that the input of NH_4^+ and NO_3^- through fog deposition 76 was 4 times that by rainfall for a fir forest ecosystem in New Hampshire. In contrast, 77 the water input via fog deposition was only 18-23% of the corresponding 78 79 precipitation amount (Yamaguchi et al., 2015). Although the chemical characteristics of fog water were highly variable in different regions, the contribution of fog water 80 was found to be crucial in the water and nutrient cycle of the forest ecosystem (Fuzzi 81 et al., 1988; Basset et al., 1991; Mueller et al., 1991a; 1991b; Berresheim et al., 1993; 82 83 Chang et al., 2006; Klemm et al., 2007; Novak et al., 2015). Moreover, leaves could gather water and ions by direct contact with fog, and even become the main source of 84 plant nutrients (Hutley et al., 1997; DeFelice, 2002; Liu et al., 2004). Additionally, the 85 chemistry of fog and rain were usually different even at the same area, because 86 87 precipitation incorporates cloud condensation nuclei (aerosol particles) when cloud droplets are formed at higher altitudes but often experiences dilution by significant 88 water vapor condensation (e.g., Collett et al., 1991b) whereas fog droplets at the 89 surface condense in lower boundary layer air (Templer et al., 2015; Novak et al., 90 2015). The pH value of fog water was often lower than rain water, and acid damage to 91 92 vegetation by fog water (Waldman et al., 1982) can be considerable because plant 93 leaves are immersed in fog for long periods of time (Waldman et al., 1982; Charlson

et al., 1982; Wrzesinsky et al., 2000; Adzuhata et al., 2001). As a result, the chemical
characteristics of fog water at montane forests have been highlighted by researchers in
several regions (Aleksic et al., 2009; Polkowska et al., 2014; Michna et al., 2015;
Wang et al., 2015). Köhler et al. (2015) reported that atmospheric water and element
inputs increased with the rise of elevation in temperate mountain forests, attributable
mainly to fog and cloud deposition.

The above-ground carbon reserves of forests in Tibet can exceed 250 t hm⁻². 100 101 Moreover, the forest with the highest monthly average net productivity is the dark coniferous forest near an altitude of 4000 m (a.s.l.) in Nyingchi city (He, 2008). The 102 chemical composition of rainwater and fogwater in this region has not previously 103 been reported; consequently, potential impacts of pollutant and nutrient deposition on 104 these forests are unquantified. Considering the unique geographical location of the 105 Qinghai-Tibetan plateau and the relatively low anthropogenic disturbance in this 106 region, understanding the chemical composition of cloud/fog water and rain water in 107 this region is not only significant to the nutrient circulation of the local forest 108 109 ecosystem, but also could provide useful insight into long range transport of air pollutants. Here we select a high elevation native fir forest as a representative region 110 to investigate the fog chemistry at Sejila mountain, with the objectives of 1) 111 understanding fog and rainfall chemical characteristics in a southeast Tibet forest 112 ecosystem; 2) clarifying the possible sources of inorganic ions in fog and rainfall in 113 the area. We hypothesize that fog interception of the forest will be frequent and that 114 concentrations of key solutes will be higher in fog water than in rainfall, pointing to a 115 need to further explore potential ecosystem impacts of fog deposition. 116

117 2. Materials and methods

118 2.1. Site description

The sampling site was established at the National Field Scientific Observation Station of the Alpine Forest Ecosystem (29.65° N, 94.72° E 3950 m a.s.l.) on Sejila Mountain. Sejila Mountain is a nationally protected area because of its primeval fir forest with a variety plants and animals. The climate is dominated by warm air currents from the Indian Ocean, with air masses arriving mainly from a southerly

direction during April to October, when fog events are likely to occur. The annual 124 average temperature was -0.73 °C and the annual average precipitation was 125 approximately 1000 mm. This station was surrounded by forest and there were no 126 nearby agricultural and industrial pollution sources except a state road (#318). Fog 127 events occurred mainly in the evening, lasting sometimes till the following morning. 128 Unfortunately, fog events were often coupled with rain events in the night, however, 129 our experiment condition was limited, even continuous power supply was not 130 131 guaranteed in our research station, and we were unable to identify fog events during nighttime. In order to avoid contamination of fog samples by rainfall, only samples 132 collected in the daytime were included in this study. Location and sampling 133 environment are shown in Fig. 1. 134

135 2.2. Sample collection

Fog samples were collected from July 2017 to September 2018. No fog events 136 were observed in any part of the research area in July 2017 or between November 137 2017 and March 2018, although frost or rime could be seen. Fog was observed almost 138 139 every day at the mountain hillside forest area during the experimental period. However, due to experimental constraints, 35 fog water samples were obtained from 140 35 individual fog events during the sampling period, with most of those samples 141 collected in July, August and September. A Caltech Active Strand Cloud Collector 142 143 (CASCC) was used to collect fog samples (Demoz et al., 1996). The CASCC family of fog collectors has been widely used for fog sampling (e.g., Collett et al., 1990, 144 2002; Bator and Collett, 1997; van Pinxteren et al., 2016; Li et al., 2011; Guo et al., 145 2012). Air was drawn by a fan into the CASCC at 24.5 m³ min⁻¹. Six rows of 508 μ m 146 Teflon strands were used to capture droplets by inertial impaction; the 50% lower size 147 cut for the collector corresponded to a drop diameter of 3.5 µm. Collected droplets 148 were drawn down the strands by aerodynamic drag, accumulated in a Teflon 149 collection trough, and flowed through a Teflon tube to a pre-cleaned sample bottle. 150 Fog samples were collected with the CASCC when fog interception events were 151 observed at the site and continued until the fog disappeared. After sampling, all 152 collected fog water samples were mixed as one sample for each individual fog event, 153

then transferred to clean polyethylene bottles (50 ml) for storage. The CASCC was
cleaned in late afternoon (e.g. 6 p.m.) in order to collect fog which mainly occurred in
the evening and morning. In addition, the national road No. 318 in Sejila Mountain
section was often closed to vehicles between 12:00 p.m. (midnight) and 7:00 a.m.
(morning) next day. Fog collection was not carried out during or after precipitation
events to avoid contamination by rain water.

Rain samples were collected during the same study period (from July 2017 to 160 161 September 2018 with the exception of the period between November 2017 and March 2018) by continuously open rain gauges (Fuyuanming Inc., Tianjin, China). All parts 162 are made of stainless steel except the glass water collection bottle. Precipitation (snow 163 and rain) samples were collected by the rain gauges during the monitoring period. 164 Since the collector remained open, rainwater concentrations measured in this study 165 reflect additional inputs from dry deposition of aerosols and trace gases to the 166 collector surface. After each precipitation event, the samples were thoroughly stirred 167 and immediately transferred to clean polyethylene bottles (50 ml) for storage. The 168 169 rainwater collection bottle was rinsed with deionized water to eliminate cross-event contamination. Further details of measuring methods and collection are given in our 170 previous studies (Xu et al., 2015, 2018, 2019a) 171

172 2.3. Sample analysis and quality control

After each sample collection, fog water or rain samples were immediately taken 173 into the laboratory and filtered with a 0.45 µm pore size cellulose acetate filter 174 (Tengda Inc., Tianjin, China), then divided into three equal parts that were used to 175 measure sample pH and concentrations of water-soluble ions (Na⁺, NH₄⁺, K⁺, Mg²⁺, 176 Ca^{2+} , Cl^- , NO_3^- , and SO_4^{2-}) and trace elements (Mn, V, Cr, As, Se, Pb, Cd, Al). The 177 pH was measured by a pH meter (SG3, Mettler Toledo Company) as soon as possible 178 after filtration (pH meter calibrated against pH 4.00 and 6.86 buffers). The other two 179 sample portions were stored in a refrigerator at 4°C prior to ion and element analysis; 180 all samples were analyzed within 48 hours. 181

Water-soluble ions were analyzed by ion chromatography (IC, DionexCorporation). Anions were analyzed by an ICS-2100 ion chromatograph equipped

with a Dionex Ionpac AG11/AS11 guard/separation column pair using a potassium 184 hydroxide eluent. Cations were analyzed on a DX-600 ion chromatograph equipped 185 with a Dionex AG12A/CS12A guard/separation column pair using a methanesulfonic 186 acid eluent; the detection limit was 0.02 μ eg L⁻¹ for Na⁺ and K⁺, 0.03 μ eg L⁻¹ for Mg²⁺ 187 and Ca^{2+} , 0.06 µeq L⁻¹ for NH₄⁺, and 0.01 µeq L⁻¹ for Cl⁻, NO₃⁻, and SO₄²⁻. No rain or 188 fog samples were below detection limits. Trace elements were analyzed by 189 inductively coupled plasma atomic emission spectrometry (ICP-AES, JOBIN-YVON), 190 the detection limit was 0.001 mg L⁻¹. H⁺ concentrations were calculated from the 191 measured pH values. 192

193 2.4. Backward trajectory analysis

Air mass back-trajectory analysis was performed using the Hybrid-Single 194 Particle Integrated Trajectory Model (HYSPLIT 4) 195 (http://ready.arl.noaa.gov/HYSPLIT.php), provided by the Air Resource Laboratory of 196 the National Oceanic and Atmospheric Administration (NOAA). Meteorological data 197 were input from the Global Data Assimilation System (GDAS1 global, 2006 to 198 present-ftp://arlftp.arlhq.noaa.gov/pub/archives/gdas1/) with a model resolution of 199 $1^{\circ} \times 1^{\circ}$. Three-day (72 h) backward trajectories were calculated at 6 h intervals (06:00, 200 12:00, 18:00, 24:00 UTC) on sampling days, with an arrival height of 500 m (above 201 ground level) for rain events and of 100 m for fog events. Cluster analysis was 202 203 performed using the trajectories based on the total spatial variance (TSV) method (Draxler et al., 2012). 204

205 2.5. Statistical analysis

In this study, the Spearman correlation values between the sum of total cations 206 $(Na^{+} + NH_{4}^{+} + K^{+} + Mg^{2+} + Ca^{2+})$ and sum of total anions $(Cl^{-} + NO_{3}^{-} + SO_{4}^{2-})$ were 207 0.874 and 0.964 for fog and rainwater, respectively, suggesting that the measured ion 208 balance in this experiment was credible. Additionally, the ratios of total cations to 209 anions in fog and rain water were 3.1 and 2.8, respectively, suggesting that one or 210 more major anions were missing in our determination. It has been reported that HCO_3^{-1} 211 and CO_3^{2-} were the dominant anions in precipitations and major rivers in 212 Qinghai-Tibet plateau (Li et al., 2007; Xiang et al., 2009; Yang et al., 2012). The pH 213

value of atmospheric water is 5.6 for equilibrium with atmospheric carbon dioxide 214 (Charlson and Rodhe, 1982). The amount of HCO_3^- was estimated using the following 215 equation $HCO_3 = 10^{(pH-5.05)}$ (Xing et al., 2017), yielding estimated HCO_3^{-1} 216 concentrations on average of 24.2 and 16.9 μ eq L⁻¹ for rain and fog samples, 217 respectively. These concentrations are not sufficient to explain the missing anions. 218 Similar results were also reported for Qinghai-Tibet plateau in previous study (Li et 219 al., 2007; Liu et al., 2015; Xing et al., 2017). Therefore, the relatively low total anion 220 concentration was attributed to unmeasured organic acids (Migliavacca et al., 2005) 221 and other unknown components. At the high pH values of these rain and fog samples, 222 gas phase organic acids such as formic and acetic acid are highly soluble. 223

The Enrichment Factor (EF) has been widely used to examine the source 224 contributions of major ions in both fog and rain water (Hissler and Probst 2006; Cong 225 et al., 2010; Liu et al., 2015; Xu et al., 2017). Generally speaking, Na⁺ can be 226 considered as a typical marine source tracer while Ca^{2+} derives mainly from the 227 continental crust. In this study, Na⁺ and Ca²⁺ were therefore used as background tracer 228 229 elements for marine and continental sources, respectively. The background of sea source elements is described in Keene et al., (1986) and continental source elements 230 in Taylor (1964). 231

The EF of an ion concentration in fog (rain) relative to the concentration in sea was estimated as follow (Liu et al., 2015):

234

$$EF_{sea} = \frac{[X/Na^+]_{sample}}{[X/Na^+]_{sea}}$$

 $EF_{sea} = \frac{[X/Ca^{2+}]_{sample}}{[X/Ca^{2+}]_{soil}}$

235 Similarly the EF of an ion concentration in fog (rain) relative to the ion in the 236 continental crust was estimated as follow (Liu et al., 2015):

237

The sources of ions can be divided into ocean, land and human activities. In this study, the follow equations were used to calculate the ion sources in fog and rain water (Liu et al., 2015):

$$\%SSF = \frac{[X/Na^+]_{sea}}{[X/Na^+]_{sample}}$$

$$\% CF = \frac{[X/Ca^{2+}]_{soil}}{[X/Ca^{2+}]_{sample}}$$

AF = 100%-%SSF-%CF

SSF refers to the sea salt fraction, CF to the crustal fraction and AF to the 241 anthropogenic fraction; it should be noted that, if the SSF (CF) ratio was greater than 242 1, SSF (CF) was calculated as 1 minus CF (SSF). 243 Spearman correlation and principal component analyses have been widely used 244 to determine natural and anthropogenic source contributions to chemical composition 245 of precipitation (Cao et al., 2009; Sheu and Lin, 2011; Yue et al., 2014). In this study, 246 Spearman correlation and principal component analysis were conducted using the 247 SPSS software package, version 20.0 (SPSS Inc., Chicago, IL). 248 3. Results and discussion 249 3.1. Fog and rain chemistry characterization 250 As shown in Table 1 and Fig. 2, the relative composition of fog water and rain 251 water was similar, while all mean ion concentrations observed were higher in fog 252 water than in rainwater. The concentration abundance order of ion concentrations in 253 fog water was $Ca^{2+} > Cl^- > Na^+ > SO_4^{-2-} > Mg^{2+} > NH_4^+ > K^+ > NO_3^-$ and an order of 254 $Ca^{2+} > Na^+ > Cl^- > Mg^{2+} > SO_4^{2-} > NO_3^- > K^+ > NH_4^+$ was observed in rain water. 255 Additionally, average total cations summed to 0.23 and 0.41 meq L⁻¹ while total 256 anions summed to 0.07 and 0.14 meq L^{-1} in rainwater and fog samples, respectively. 257 The mean concentrations of NH₄⁺, SO₄²⁻, K⁺, Ca²⁺, Cl⁻, NO₃⁻, Mg²⁺, Na⁺ in fog were 258 4.6, 2.6, 2.3, 1.9, 1.9, 1.5, 1.3 and 1.3 times those in rainwater, respectively. The 259 percentages of Ca^{2+} , NH_4^+ and K^+ in total cations were higher in fog than in rainwater. 260 NH_4^+ increased from 3% in rain to 8% in fog water. The percentage of SO_4^{2-} was also 261

263 and NO_3^- (Fig. 2a and b).

262

In high-elevation environments, cloud and fog water have generally been recognized as being more acidic than rain (Anderson et al., 1999; Collett et al., 2002; Weathers et al., 1988; Herckes et al., 2002). Characterizing the acidity of fog and rain water is crucial to better understanding inputs of acidic deposition at high-elevation

higher in fog than in rainwater. By contrast, opposite behaviors were observed for Cl⁻

locales. Higher acidity was observed in these Tibetan fog samples than in collected precipitation samples; but the difference was relatively small (6.4 versus 6.2). One possible explanation is that pH values in fog and rain water were strongly influenced by high Ca^{2+} concentrations.

The effect of cloud-precipitation concentration differences has previously been 272 highlighted by other studies (Lovett et al., 1982; Collett et al., 1990, 1991a). 273 Generally speaking, ion concentrations in fog water have been found to be higher than 274 275 those in rain water; however, the differences have varied largely across regions(Chang et al., 2006; Aleksic et al., 2009). For instance, large variations for all ion 276 concentrations between fog and rain water were found in NE Taiwan (Chang et al., 277 2006; Beiderwieden et al., 2007), the concentrations of Na⁺, NH₄⁺, Mg²⁺, Ca²⁺, Cl⁻, 278 NO_3^- and SO_4^{2-} in fog water were 53, 13, 9.4, 5.7, 15, 15 and 14 times higher than 279 those in rain water, respectively. In contrast, minor differences of ion concentrations 280 between fog and rain water were reported at Mangdang Mountain, China (Huo et al., 281 2010), the concentrations of Na⁺, NH₄⁺, K⁺, Mg²⁺, Ca²⁺, Cl⁻, NO₃⁻ and SO₄²⁻ in fog 282 water were 5.0, 1.6, 1.7, 2.4, 3.7, 2.0, 3.5 and 1.6 times higher than those in rain water 283 for that study, respectively. Compared to variations of ion conentrations between fog 284 and rain water in other regions (Bridges et al., 2002; Aleksic et al., 2009; Straub et al., 285 2012), the differences of ion concentrations in Sejila were minor. One reason might be 286 that ion concentrations in fog water were low due to a lack of major nearby emission 287 sources from human activity (Table 2). A second factor is that precipitation samples 288 were collected by continuously open rain gauges, resulting in some dry deposition 289 being mixed in those samples, such that the measured ion concentrations in rain 290 samples should be higher than in the local precipitation. 291

Compared with previous studies (**Table 2**), high Ca^{2+} concentrations were found in fog water samples. High Ca^{2+} concentrations are expected to be found in the Qinghai-Tibet plateau ambient environment, as with aerosol (Cong et al., 2007), rivers (Huang et al., 2008; Xiang et al., 2009), and soil (Li et al., 2009). Most areas of the Qinghai-Tibetan plateau are above 4,000 m altitude, with sparse vegetation, bare land and relatively high winds, and the weathering of the rocks on top of the mountain is

severe. Thus, concentrations of particulate Ca²⁺ in the atmosphere are relatively high 298 (Zhang et al., 2001) and expected to contribute to high Ca²⁺ concentrations in local 299 fog water. Moreover, the high Ca^{2+} concentrations reflect abundant atmospheric dust 300 particles that can be efficiently incorporated into precipitation though in-cloud and 301 below-cloud scavenging processes (Huang et al., 2014). As a result observed Ca²⁺ 302 concentrations in precipitation in the Tibetan plateau have been extremely high (Li et 303 al., 2007; Yang et al., 2012; Li et al., 2015; Liu et al., 2015). HCO₃⁻ was reported as 304 the main anion in precipitation, accounting for as much as 62% of the total anion 305 concentration. HCO_3^{-1} was not measured in our work, one factor contributing to total 306 measured anion concentrations much lower than total cation concentrations in both 307 fog and rain samples (Table 1). 308

 NH_4^+ concentrations in fog water averaged 4.6 times higher than in rain water 309 (Table 1). Often, rain and fog have different origins, with precipitation falling from 310 clouds formed at higher altitudes and fog forming in lower boundary layer air. High 311 NH_4^+ concentrations in fog water might reflect high atmospheric NH₃ concentrations 312 from local agricultural activity at our study site. Wang et al. (2019) has reported that 313 concentrations of NO₃⁻-N and NH₄⁺-N in precipitation were similar in southeast Tibet, 314 but atmospheric NH₄⁺-N (sum of gaseous NH₃ and particulate NH₄⁺) concentrations 315 were much higher than NO₃⁻-N (sum of gaseous NO₂, HNO₃ and particulate NO₃⁻) 316 concentrations. It is worth noting that NH₄⁺ concentrations measured in this study 317 were relatively low compared with those in other areas (Table 2). One possible 318 explanation is that NH₃ volatilization from soil is limited by low temperature at the 319 altitude of 4000 m. SO_4^{2-} and NO_3^{-} mainly come from oxidation of sulfur dioxide and 320 nitrogen oxides, emitted by a wide range of anthropogenic sources, such as coal and 321 fuel combustion (Yang et al., 2012; Xu et al., 2019b). High concentrations of SO_4^{2-} 322 and NO₃⁻ have, not surprisingly been reported in areas near industrial districts 323 (Bridges et al., 2002; Strater et al., 2010). In this study, both SO_4^{2-} and NO_3^{-} 324 concentrations were relatively low (Table 2), consistent with a lack of substantial 325 anthropogenic activities in the study area. 326

327 3.2. Concentration of trace elements in fog and rain water

Although trace element concentrations are much lower than the major ion 328 concentrations, they can still play an important role in aqueous reactions in clouds 329 (e.g., Mancinelli et al., 2005; Harris et al., 2013). The composition of trace elements 330 in the atmosphere is mainly attributed to anthropogenic emissions, biogenic emissions, 331 biomass buring, and soil dust (Viana et al., 2009). Trace element concentrations 332 measured in fog water in this study are shown in Table 3, where concentrations of Al, 333 As, Mn and Se are seen to be higher than those of Pb and V. Cd and Cr were also 334 335 measured in the study, but concentrations were below our detection limit. Mn and Se, essential micronutrient for plants, averaged 13.8 and 16.4 μ g L⁻¹. Mn is also an 336 efficient catalyst for in-cloud reactions such as the oxidation of dissolved SO₂ to 337 sulfate (Rao and Collett, 1998; Harris et al., 2013). Al and As, considered as toxic 338 metals, had mean concentrations of 19.5 and 10.8 μ g L⁻¹, respectively. All trace 339 elements in rain water were below the detection limit except Al and As, with mean 340 average concentrations of 2 and 6 μ g L⁻¹, respectively. 341

Compared with more economically developed areas in China, such as Mt. Lu and 342 343 Mt. Tai (**Table 3**), all Sejila mountain trace element concentrations were lower except for Se. Al, Ca, and Mn all can be derived from terrestrial sources (Cong et al., 2010). 344 In this study, the correlation (p < 0.05) between Al and Ca was 0.469, while the 345 correlations between Al and Mn, and Ca and Mn reached 0.784 and 0.797 (both 346 p<0.01), respectively. These results suggest that those elements came mainly from 347 terrestrial sources in our research area. Pb and Se are considered primarily as 348 anthropogenic source elements. Considering the low anthropogenic activities in our 349 research area, concentrations of trace elements should be lower than in highly 350 351 populated regions. Interestingly, high concentraitons of As and Se were found in this 352 study; one explanation could be that those elements mainly derived from local crustal material. Similar results were found in Qinghai-Tibetan plateau observations in other 353 previous studies (Cong et al., 2007; Wang et al., 2012). Vehicle exhaust might also be 354 a factor contributing to local atmospheric heavy metal concentrations because our 355 monitoring station was located close to the national road 318# (Tinus et al., 2010; 356 Srimuruganandam et al., 2011). 357

358 *3.3. Source assessment of ions in fog and rain water*

Air mass back-trajectories are frequently used to trace regional transport of 359 atmospheric pollutants (Sun et al., 2015; Yue et al., 2014). In this study, 72 hour 360 back-trajectories for each fog event (rainfall event) from August 2018 to September 361 were computed. As shown in Fig. S1a, almost all air masses arrived from a southerly 362 direction when fog events occurred. Additionally, all air masses from different 363 directions were influnced by surface sources. As shown in Fig. S1b. major air mass 364 365 transport patterns for rainfall events could be divided into 4 categories, the percentage of south, southwest, west and northwest source regions were 43%, 37%, 7% and 13% 366 respectively, similar to findings in previous studies (Liu et al., 2015; Wang et al., 367 2018). These results reflect that southeast Tibet is influenced by the southwest 368 monsoon and air mass transport from the Indian Ocean (Yang et al., 2012). Not 369 surprisingly, therefore, high Na⁺ concentrations were found in both rainfall and fog 370 samples (Table 1). 371

Concentrations of water soluble inorganic ions have been extensively studied 372 373 because they often dominate fog composition in areas with a continental background and high air pollution (Collett et al., 2002; Marinoni et al., 2004). Spearman 374 correlation analyses have been widely used to examine the relationships among 375 various water-soluble inorganic ions. The correlations between different ions in fog 376 water were all significant (Table S1), perhaps reflecting simultaneous dilution 377 (concentration) of all fog solutes with increasing (decreasing) fog liquid water content. 378 In contrast, the correlation between more than half of the ions was not significant in 379 rain water (Table S2). 380

Principal component analysis (PCA) as a multivariate statistical method is commonly used to find a small number of factors from a data set of many correlated variables (Brereton, 2003). The original data matrix is decomposed into the product of a matrix of factor loadings and a matrix of factor scores plus a residual matrix. The residual matrix contains the part of variance of the data set that cannot be explained by common factors (e.g. analytical uncertainties). On the basis of the correlation matrix, orthogonal factors, sorted by descending order (Marengo et al., 1995) are

extracted solving an eigenvalue problem. In general, the number of extracted factors 388 is lower than the number of measured features. After rotation of the factor loading 389 matrix, the factors can often be interpreted, for example, as origins or common 390 sources of pollutants. Based on the varimax rotated PCA, we characterized fog 391 samples using three factors which collectively explain 90.0% of the total variance 392 (**Table 4**). In fog water, PC1 is dominated by Mg^{2+} , Ca^{2+} , NO_3^{-} and SO_4^{-2-} , and thus 393 can be regarded as crustal/soil dusts combined with transported anthropogenic 394 emissions. PC2 features high fractions of Na⁺ and Cl⁻, indicating a marine source. PC3 395 contained high levels of NH_4^+ and K^+ , which could be explained by local source 396 emissions from agriculture and residential biomass burning which is commonly used 397 for local cooking and domestic heating. For rain water, three factors were identified 398 with the combined explained variance more than 89%. PC1 had high loading for Mg^{2+} , 399 Ca^{2+} , Na^+ , NO_3^- and SO_4^{2-} , and may be a combination of dust, sea salt, and long range 400 transported anthropogenic emissions. PC2 shows high loadings for K⁺ and Cl⁻, 401 suggesting a biomass burning source. The third component was NH_4^+ , suggesting 402 403 contributions from an agricultural source, perhaps dominated by local emissions as described above. 404

Recognizing that Na⁺ is typically associated with a marine source and Ca²⁺ with 405 a terrestrial source, EF is frequently used to evaluate ion enrichments in precipitation. 406 If the EF value of an ion is higher (lower) than 1, this ion could be seen as enriched 407 (diluted) in the environment relative to the reference source composition. In this study, 408 NH_4^+ , NO_3^- and $SO_4^{2^-}$ were all enriched for marine sources and terrestrial sources 409 (Table S3), suggesting that concentrations of those ions are influenced by human 410 activities. NO_3^- and SO_4^{2-} might come from India (Fig. S1) where anthropogenic 411 emissions of NO_x and SO_2 to the atmosphere significantly increased from 2001 to 412 2015 (Paulot et al., 2018). NH_4^+ might result from high ambient particulate NH_4^+ 413 concentrations in summer in southeast Tibet (Wang et al., 2019) as well as from 414 scavenging of local agricultural NH₃ emissions. Both rain and fog showed similar 415 patterns for ion sources: Na^+ , Mg^{2+} and Cl^- were mainly from a marine source, K^+ and 416

417 Ca^{2+} were mainly from a terrestrial source, and NH_4^+ , NO_3^- and SO_4^{-2-} were dominated 418 by anthropogenic emission (**Table 5**).

419 **4. Summary and conclusions**

420 This study analyzed the chemical characteristics of fog water and precipitation in the southeast Tibet and examined potential sources of different ions. Air masses 421 associated with both fog and precipitation events arrived mainly from a southerly 422 direction, suggesting potential impacts of species transported over long distances from 423 India or the Indian Ocean. Concentrations of all measured ions were enriched in fog 424 water over rain, but both were dominated by Ca^{2+} . The major ion sources for fog and 425 precipitation showed a similar pattern. Na⁺ and Mg²⁺ were mainly from marine 426 sources, Ca^{2+} was contributed mainly from abundant crustal dust aerosols, and SO_4^{2-} 427 and NO_3^- were dominated by transported anthropogenic emissions. Enriched NH_4^+ 428 concentrations in fog water appeared to reflect local agricultural emissions of NH₃ 429 while K^+ and Cl^- appeared to come from local residential biomass burning sources. 430 The most abundant trace elements in fog water were Al, As, Mn and Se, whereas only 431 Al and As were detected in rain water. 432

Because these are the first measurements of this kind in the region and because the dataset is still relatively limited in scope, there remains a need for further research, including better characterization of fog occurrence, liquid water content, and composition over longer time periods. The enrichment of pollutant and nutrient species in fog over rain and the frequent interception of fogs by the local forest certainly points to a need for better examining potential contributions of occult (fog) deposition to nutrient and pollutant deposition budgets for the local ecosystem.

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451

452 **Author contributions**

- 453 Xuejun Liu designed the research. Wei Wang and Wen Xu conducted the research
- 454 (collected the data and performed the measurements) and wrote the manuscript. All
- 455 authors were involved in the discussion and interpretation of the data as well as the
- 456 revision of the manuscript.

457 **Competing interests**

458 The authors declare that they have no conflict of interest.

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839	Figure captions
840	Figure 1. Location and sampling environment of the study site at Sejila Mountain,
841	southeast Tibet
842	Figure 2. Comparison of the percentage of cations (a) and anions (b) based on
843	equivalents in fog and precipitation samples
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Figure 2





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Na* NH4* K Mg* Ca* Cf NO3* SO4* HT cations Rain 58.5±26.9 7.13±2.93 13.1±5.2 28.7±17.3 123±83 40.1±13.9 13.9±20.2 19.4±17.5 0.47±0.25 230 Fog 75.2±26.6 32.9±20.3 29.4±13.9 38.0±15.0 233±97 75.8±29.0 20.6±16.1 50.0±48.0 0.72±0.59 408		+			2+	~ 2+			2		Total	Т
Rain 58.5±26.9 7.13±2.93 13.1±5.2 28.7±17.3 123±83 40.1±13.9 13.9±20.2 19.4±17.5 0.47±0.25 408 Fog 75.2±26.6 32.9±20.3 29.4±13.9 38.0±15.0 233±97 75.8±29.0 20.6±16.1 50.0±48.0 0.72±0.59 408 V		Na⁺	$\mathrm{NH_4}^+$	K⁺	Mg²⁺	Ca ²⁺	Cl	NO ₃ ⁻	$SO_4^{2^2}$	H^{+}	cations	a
Fog 75.2±26.6 32.9±20.3 29.4±13.9 38.0±15.0 233±97 75.8±29.0 20.6±16.1 50.0±48.0 0.72±0.59 408	Rain	58.5±26.9	7.13±2.93	13.1±5.2	28.7±17.3	123±83	40.1±13.9	13.9±20.2	19.4±17.5	0.47±0.25	230	
ACCEPTED MARINES	Fog	75.2±26.6	32.9±20.3	29.4±13.9	38.0±15.0	233±97	75.8±29.0	20.6±16.1	50.0±48.0	0.72±0.59	408	
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Monitoring site	Na^+	$\mathrm{NH_4}^+$	\mathbf{K}^+	Mg^{2+}	Ca ²⁺	Cl	NO ₃ ⁻	SO4 ²⁻	pН	Year	Reference
Northern Poland	19.4	16.5	5.9	1.5	8.7	18.1	14.7	17.8	6.1	2010	Polkowska et al. (2014)
Northeastern Taiwan	29.5	22.2	9.00	12.9	30.0	34.3	27.9	108	4.3	2000-2001	Chang et al. (2002)
Norway Sundsbø	69.7	3.90	1.90	11.7		102	5.3	13.1	5.0	2011	Wang et al. (2015)
Norway Bakka	75.1	10.3	2.60	14.2	5.5	106	<1	14.8	5.0		
Norway Hakadal	73.3	87.1	13.5	11.9	31.4	47	61.7	42.8	4.7		
whiteface mountain	3.10	117	1.60	5.00	17.8	5.60	79.9	200	-	1994-2006	Aleksic and Dukett, (2010)
Mangdang Mountain	25.0	63.0	5.00	12.0	92.0	12.0	105	63.0	4.8-6.0	2009	Huo et al. (2010)
Germany	35.0	216	6.10	5.10	9.80	30.0	164	43.0	4.3	2010	van Pinxteren et al. (2016)
South Korea	69.0	173	10.1	22.2	59.8	74.9	93.0	176	4.4	2002-2003	Kim et al. (2006)
Southern China	21.9	276	14.6	9.90	62.2	27.9	182.8	341	3.8	2011-2012	Sun et al. (2015)
NE taiwan	58.6	235	10.3	18.8	34.1	42.8	179	401	3.6	2006	Beiderwieden et al. (2007)
Swiss 2330 m a.s.1	43.0	143	5.00	12.6	46.8	10.6	87.0	72.3	6.4	2006-2007	Michna et al. (2015)
Swiss 1650 m a.s.l	44.3	249	10.3	13.9	35.4	20.5	200	105	6.6		
Swiss 682 m a.s.l.	78.0	1030	96.1	139	117	44.0	347	334	7.3		
Swiss	20.0	045	72.0	511	22.0	22.0	204	207	7.0		
(inside forest canopy)	38.2	845	72.0	54.1	32.8	32.8	294	297	1.2		
Czech	64.4	203	19.4	404	136	155	726	1250	3.0	1995-1996	Bridges et al.(2002)
Poland	130	230	50.0	31.0	100	140	170	220	4.1-5.1	2005-2006	Blas et al. (2010)
Ailaoshan Mountain	6.09	727	25.3	7.41	27.5	24.3	69.2	185	4.05	2015	Nieberding et al., 2018
Cervenohorske sedlo	26.1	44.3	7.7	<8.2	25	<19.7	<29	127	5.2	2002	Zapletal,et al.,2007
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Table 2. Comparison of concentrations ($\mu eq L^{-1}$) of chemical composition in fog water in previous studies.

Sites	Al	As	Mn	Pb	Se	V	Cd	Cr	Year	References	
Lu Mountain	111	20.4	16.4	54.4	7.40	5.90	1.66	8.37	2011-2012	Sun et al. 2015	
Tai Mountain	157	13.7	42.8	46.2	n.a.	n.a.	3.08	0.93	2007	Liu et al. 2012	
Yangtze River	391	3.38	40.0	5.83	2.10	3.65	1.37	4.96	2015	Xu et al. 2017	
Northern Poland	9.50	n.a.	1.86	1.06	n.a.	n.a.	7.78	n.a.	2010	Polkowska et al. 2014	
Northern Chile	n.a.	9.10	<200	<10	<5	n.a.	< 0.5	<2.5	2008	Sträter et al. 2010	
Elden Mountain	16.6	0.80	34.0	0.40	2.00	3.20	n.a.	1.70	2005-2007	Hutchings et al. 2009	
Sejila Mountain	19.5	10.8	13.8	1.23	16.4	4.38	n.a.	n.a.	2017-2018	This study	
Note: n.a. denotes not available.											

Table 3. Comparison of concentrations (μ g L-1) of trace elements in fog water at Sejila Mountain with other areas

		Na ⁺	$\mathrm{NH_4}^+$	\mathbf{K}^+	Mg^{2+}	Ca ²⁺	Cl	NO ₃	SO4 ²⁻	Variance (%)	Cumulative (%)
Fog	PC1	0.329	0.515	0.129	0.731	0.685	0.272	0.879	0.941	71.1	71.1
	PC2	0.888	0.105	0.584	0.473	0.414	0.874	0.254	0.252	12.5	83.6
	PC3	0.230	0.683	0.749	0.443	0.497	0.193	0.302	0.110	6.6	90.0
	СТ	0.951	0.743	0.919	0.955	0.887	0.875	0.928	0.960		
	PC1	0.656	0.072	-0.062	0.957	0.927	0.567	0.857	0.970	59.8	59.8
Rain	PC2	0.518	0.182	0.897	-0.028	0.229	0.700	0.212	0.075	20.2	80.0
	PC3	0.334	0.938	0.290	-0.169	0.182	-0.092	0.294	0.004	9.4	89.3
	СТ	0.811	0.918	0.893	0.945	0.945	0.820	0.865	0.947		

Table 4. Varimax-rotated principal component analysis of major ions of major ions in fog and rain at Sejila Mountain.

Note: PC1, PC2 and PC3 indicate the first, second and third component, respectively. CT means communality.

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		Na ⁺	$\mathrm{NH_4}^+$	K^+	Mg^{2+}	Ca ²⁺	Cl	NO ₃	SO4 ²⁻
	SSF	100	2.0	3.3	84.2	1.7	77.1	2.2	8.7
Fog	CF		0.5	96.7	15.8	98.3	0.5	0.8	3.6
	AF		97.5			2	22.4	97.0	87.7
	SSF	100	7.3	5.8	86.7	2.5	99.5	2.6	17.4
Rain	CF		1.2	94.2	13.3	97.5	0.5	0.6	5.0
	AF		91.5					96.8	77.6

Table 5. Source contributions (%) for major ions in fog and precipitation at Sejila Mountain.

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Highlights

- 1. Characteristics of ions and trace elements in fog water in the Sejila mountain of southeast Tibet were reported.
- 2. Ion and trace element concentrations were higher in fog water than in rain water but pH values were on the contrary.
- 3. Ratios of NH_4^+ -N and NO_3^- -N were much higher in fog water than in rain water.
- 4. Local source contributed mainly to atmospheric ions in the Sejila mountain.

Competing interests

The authors declare that they have no conflict of interest.