1 Unexpectedly high concentrations of atmospheric mercury

2 species in Lhasa, the largest city on the Tibetan Plateau

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25 Abstract

26 Lhasa City is located in the central Tibetan Plateau and is the most densely populated area. As the first continuous monitoring of atmospheric mercury (Hg) species in a city on the Tibetan Plateau, 27 28 our monitoring in Lhasa showed that the concentrations of gaseous elemental Hg (GEM), gaseous 29 oxidized Hg (GOM), and particle-bound Hg (PBM) during subsequent of the Indian Summer 30 Monsoon (S-ISM) period were 2.73 ± 1.48 ng m⁻³, 38.4 ± 62.7 pg m⁻³, and 59.1 ± 181.0 pg m⁻³, respectively. During the Westerly Circulation (WEC) period, the GEM, GOM and PBM 31 32 concentrations were 2.11 ± 2.09 ng m⁻³, 35.8 ± 43.3 pg m⁻³, and 52.9 ± 90.1 pg m⁻³, respectively. The GOM and PBM concentrations were higher than those of previous monitoring on the Tibetan 33 Plateau and other provincial capitals in China. Typical high-value occurrence processes were studied 34 to investigate random events with high atmospheric Hg concentrations in Lhasa. Combustion event 35

36 nearby or further away may be the main contributor of the high-concentration events. The lowest GEM concentrations occurred in the afternoon and persistently high concentrations were observed 37 38 at night. The changes in GEM concentrations were consistent with the trends of other pollutant 39 concentrations and contradictory to those of the wind speed. The high GEM concentrations at night 40 can be attributed to the lower boundary layer height and lower wind speed. For both GOM and PBM, 41 higher GOM concentrations occurred during the day and PBM during the night. The results of the 42 principal component analysis indicated that local sources and wind speed are important factors 43 influencing atmospheric Hg concentrations in Lhasa. The trajectory simulation showed that the 44 source of the GEM in Lhasa gradually shifted from the south to the west of Lhasa from the S-ISM 45 to the WEC periods, while both the southern and western sources were important in the late WEC 46 period. The concentrations and change patterns of Hg species in Lhasa were significantly different 47 than those at other monitoring sites on the Tibetan Plateau. Monitoring Hg species in Lhasa shows 48 the possible maximum anthropogenic influences on the Tibetan Plateau and demonstrates the 49 dramatic effect of wind on changes in urban atmospheric Hg concentrations.

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51 **1. Introduction**

52 Mercury (Hg) has received worldwide attention owing to its high toxicity and bioaccumulation. 53 Atmospheric mercury (Hg) exists in three different forms: atmospheric gaseous elemental Hg 54 (GEM), gaseous oxidized Hg (GOM), and particle-bound Hg (PBM) (Selin, 2009). They exhibit 55 different behaviors in the environment owing to their various chemical properties (Selin, 2009; 56 Travnikov et al., 2017; Lindberg and Stratton, 1998; Seigneur et al., 2006). Many established 57 monitoring networks for atmospheric Hg exist in North America and Europe (Stylo et al., 2016) 58 including the Atmospheric Mercury Network (AMNet; Gay et al., 2013), the Global Mercury 59 Observation System (GMOS; Sprovieri et al., 2013; Sprovieri et al., 2016), the Canadian 60 Atmospheric Mercury Network (CAMNet; Kellerhals et al., 2003), and the Arctic Monitoring 61 Assessment Programme (AMAP; https://mercury.amap.no/) (Gay et al., 2013; Sprovieri et al., 2013; 62 Sprovieri et al., 2016; Kellerhals et al., 2003). They have been operating for decades and have 63 provided a large amount of atmospheric Hg data. Compared to Europe and the United States, independent research teams have conducted monitoring work in China based on different research 64

65 interests (Fu et al., 2012a; Fu et al., 2008; Fu et al., 2016a; Fu et al., 2019; Fu et al., 2016b; Liu et al., 2011; Feng and Fu, 2016; Feng et al., 2013; Wang et al., 2015; Hu et al., 2014; Ci et al., 2011; 66 67 Duan et al., 2017; Liu et al., 2002; Yin et al., 2018; Yin et al., 2020b; Lin et al., 2022; Lin et al., 68 2019). Most monitoring stations are set up only in developed regions, such as eastern and central 69 China, owing to operational difficulties in remote areas. Few studies on atmospheric Hg in western 70 China have been reported; thus, little is known about the overall level of atmospheric Hg in western 71 China. To better employ the Minamata Convention and verify the effect of the implementation of 72 the Convention, monitoring atmospheric Hg concentrations around the globe is significant and can 73 aid in identifying the global Hg transport pattern.

74 The Tibetan Plateau is in the mid-latitudes of the Northern Hemisphere (in central Asia) and is 75 an important area for studying the global Hg circulation. Owing to the high altitude and rough living 76 conditions, there is little Hg research on the Tibetan Plateau. This area is less developed and there 77 are few industrial activities; therefore, it is generally considered a clean region and can be treated 78 as a background condition. However, there are large tourist cities in this area, such as Lhasa, where 79 the number of tourists reached 40,121,522 in 2019 (Tibet Bureau of Statistics, 2020). Local cement 80 production in Tibet reached 10.81 million tons in 2019 (Tibet Bureau of Statistics, 2020). 81 Meanwhile, although the high altitude makes the Tibetan region a natural barrier between inland 82 China and the Indian subcontinent (Qiu, 2008; Yao et al., 2012; Pant et al., 2018), the Tibetan Plateau 83 is potentially influenced by the Indian summer monsoon (ISM) and the Westerly circulation (WEC). 84 Trans-boundary inputs of atmospheric pollutants to the Tibetan Plateau have been demonstrated in 85 pollutant studies such as with persistent organic pollutants and black carbon (Yang et al., 2018; Li 86 et al., 2016b; Zhang et al., 2015b; Pokhrel et al., 2016; Wang et al., 2018; Zhang et al., 2015a; Feng 87 et al., 2019; Zhu et al., 2019). Our previous study on atmospheric Hg in the Qomolangma region 88 (QNNP) also suggested that atmospheric Hg from India can be transported and affect atmospheric 89 Hg concentrations on the Tibetan Plateau as a result of the Indian monsoon (Lin et al., 2019). Hence, 90 it remains unclear whether the Tibetan Plateau can be treated as a background area for studying 91 atmospheric Hg, and further monitoring data are required. Monitoring in the largest cities on the 92 Tibetan Plateau will provide important information and corroboration to address this query.

93 In previous study, Yin et al. (2018) reported GEM concentration data for the Namco region on

94 the Tibetan Plateau from 2012-2014 and found that the GEM concentration at Namco was $1.33 \pm$ 95 0.24 ng m⁻³, which is lower than the mean GEM concentration in the Northern Hemisphere 96 (Lindberg et al., 2007; Slemr et al., 2015; Venter et al., 2015; Sprovieri et al., 2016; Lan et al., 2012). 97 Our previous study at QNNP (Lin et al., 2019) showed that the atmospheric Hg concentrations were 1.42 ± 0.37 ng m⁻³, 21.4 ± 13.4 pg m⁻³, and 25.6 ± 19.1 pg m⁻³ for GEM, GOM, and PBM, 98 99 respectively, close similar to the average GEM concentrations in the Northern Hemisphere (Lindberg et al., 2007; Slemr et al., 2015; Venter et al., 2015; Sprovieri et al., 2016; Lan et al., 2012). 100 101 The concentrations of atmospheric Hg species in Nyingchi, in the southeast Tibetan Plateau, were very low (1.01±0.27 ng m⁻³, 12.8±13.3 pg m⁻³, and 9.3±5.9 pg m⁻³ for GEM, GOM, and PBM, 102 103 respectively), which may be affected by heavy wet deposition and the large amounts of vegetation 104 in the Yarlung Zangbu/Brahmaputra Grand Canyon (Lin et al., 2022). However, Namco, QNNP 105 and Nyingchi are remote areas on the Plateau, with few populations and industries. During previous 106 studies in Lhasa, the largest city on the Plateau, only dry and wet depositions of atmospheric Hg 107 were analyzed. Monitoring of atmospheric Hg particulate matter (Huang et al., 2016a) indicated that Lhasa has mean particulate Hg levels as high as 224 pg m⁻³ (ranging from 61.2 to 831 pg m⁻³), 108 109 which is much higher than expected. Huang et al. (2013) measured the wet deposition of atmospheric Hg in Lhasa in 2010 and showed that the wet depositions of total Hg concentration and 110 particulate Hg concentration were higher during the non-monsoon period than that during the 111 112 monsoon period. The active Hg was higher during the monsoon than during the non-monsoon period, and they concluded that the wet deposition of Hg originated mainly from local sources. This 113 114 indicates that atmospheric Hg concentrations in Lhasa may be elevated and further detailed 115 monitoring is needed.

In this study, we conducted a high-time-precision atmospheric Hg species monitoring system in Lhasa. We performed continuous measurements of GEM, GOM, and PBM concentrations from subsequent of the Indian Summer Monsoon (S-ISM) period to the WEC period from 2016 to 2017. Based on literature research, this is the first continuous monitoring of atmospheric Hg species in a city on the Tibetan Plateau, and the influence of human activities, meteorological factors, and longrange transportation of pollutants on the diurnal variation of atmospheric Hg in Lhasa is discussed. We combined monitoring with other pollutant concentrations to explore the main factors influencing the local atmospheric Hg concentrations. To determine the detailed source profile of atmospheric Hg, we combined real-time Hg monitoring data with backward trajectory and cluster analyses. This study can help understand atmospheric Hg characteristics in the city of the Plateau and provide scientific support for managerial decision-making.

- 127 **2.** Material and methods
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Mater far and methods

2.1 Atmospheric Hg monitoring sites

129 The monitoring site for atmospheric Hg species in Lhasa was set up on the top floor of the 130 Lhasa station office building in the Institute of Tibetan Plateau Research, Chinese Academy of Sciences, in western Lhasa City (29.64°N, 91.03°E, 3650 m above sea level; Figure 1, Figure S1). 131 132 The monitoring in Lhasa included the subsequent Indian Summer Monsoon (S-ISM, August 29 to September 30, 2016) and Westerly Circulation (WEC, October 1, 2016 to February 2, 2017) periods 133 134 from August 29, 2016 to February 2, 2017. To better understand the changes of atmospheric Hg concentrations in different periods, the WEC period was further divided into WEC1 (October 1 to 135 136 December 30, 2016) and WEC2 (January 1 to February 2, 2017) periods. Lhasa is located in the central region and is the largest city on the Tibetan Plateau, covering an area of approximately 60 137 138 km². The Lhasa population in 2019 was 720,700, accounting for approximately 20.6% of the total 139 population of the Tibet Autonomous Region (Tibet Bureau of Statistics, 2020). The entire city is in a flat river valley surrounded by mountains up to 5,500 m above sea level. During the ISM period 140 141 (from May to September), the low pressure on the Tibetan Plateau attracts the summer monsoon 142 from the Indian Ocean to the Plateau, exhibiting a wetter monsoon season (Qiu, 2008). During the 143 non-monsoon season (from October to April), the large-scale atmospheric circulation on the Tibetan 144 Plateau is mainly under the control of westerly winds, which largely come from the inland areas of 145 Central Asia, presenting a drier season in Lhasa during this time (Huang et al., 2010; Guo et al., 146 2015). According to previous studies, the air quality in Lhasa may be influenced by local emissions 147 from anthropogenic activities (e.g., power plants, cement facilities, vehicular traffic, and religious 148 activities) (Li et al., 2008; Cong et al., 2011; Huang et al., 2010; Guo et al., 2015; Luo et al., 2016; 149 Li et al., 2016a) and long-range transboundary atmospheric transport (Huang et al., 2016a; Huang 150 et al., 2016b).



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Figure 1 Location of the Lhasa station in the Institute of Tibetan Plateau Research,
 Chinese Academy of Sciences (red star).

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155 **2.2 GEM, GOM, and PBM active monitoring**

156 Monitoring in Lhasa was performed using Tekran models 2537 B, 1130, and 1135 (Tekran Inc., 157 Toronto, Canada) for real-time continuous measurements of GEM, GOM, and PBM concentrations. Model 2357b is the main analytical module used to analyze Hg concentrations employing the cold 158 159 atomic fluorescence technique. Model 1130 is divided into pump and lysimeter modules, which is 160 mainly used for the collection and resolution of GOM. Model 1135 is the particle collection module that is mainly used to collect and analyze atmospheric Hg in the particulate state. During the actual 161 162 monitoring, considering the low air pressure on the Tibetan Plateau, we reduced the airflow of the pump module to 7.5 L/min (Swartzendruber et al., 2009; Zhang et al., 2015a; Zhang et al., 2016; 163 Lin et al., 2019; Lin et al., 2022) and the airflow in model 2537B was reduced to 1 L/min to ensure 164 that atmospheric Hg monitoring could be continuously performed. All monitoring data were 165 166 converted to concentrations under standard atmospheric pressure. The Tekran 2537B analyzer was

automatically calibrated every 23 hours using the instrument's internal Hg permeation source and
was calibrated before and after monitoring using an external Hg source. The same instrument setup
was used for QNNP and Nyingchi (Lin et al., 2019; Lin et al., 2022). The Tekran ambient Hg
analyzer has been described in detail in previous studies (Landis et al., 2002; Rutter et al. 2008; de
Foy et al. 2016).

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2.3 Meteorological data and other pollutant data

173 During the monitoring period, the Vantage Pro2 weather station (Davis Instruments, USA) 174 recorded local temperature (accuracy of 0.1°C), relative humidity (accuracy of 1%), wind speed 175 (accuracy of 0.1 m s⁻¹), wind direction (accuracy of 1°), barometric pressure (accuracy of 0.1 hPa), solar radiation (accuracy of 1 W m⁻²), and UV index (accuracy 0.1 MEDs). Hourly measurements 176 of PM_{2.5}, PM₁₀, SO₂, NO₂, O₃, and CO concentrations and the air quality index (AQI) were obtained 177 178 from monitoring stations hosted by the Ministry of Ecology and Environment of China and 179 published by the China Environmental Monitoring Center. The station was set up 10 km from the 180 atmospheric Hg monitoring station.

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2.4 Backward trajectory simulation and potential source analysis

182 To better understand the transport paths of atmospheric GEM, the Hybrid Single-Particle 183 Lagrangian Integrated Trajectory (HYSPLIT) model was used to calculate backward trajectories (Stein et al., 2015; Chai et al., 2017; Chai et al., 2016; Hurst and Davis, 2017; Lin et al., 2019). The 184 185 HYSPLIT model (https://www.arl.noaa.gov/hysplit/hysplit/) is a hybrid approach that combines 186 Lagrangian and Eulerian methods, which was developed by the National Oceanic and Atmospheric 187 Administration (NOAA) as a tool to explain the transport, dispersion, and deposition of particles in 188 the atmosphere. The backward trajectory simulation used Global Data Assimilation System (GDAS) 189 data with 1°x1° latitude and longitude horizontal spatial resolution and 23 vertical heights every 6 190 hours in this study. We examined the effect of arrival height on the trajectories using different arrival 191 heights (50 m, 100 m, 400 m, and 1,000 m) in December 2016 (Figure S2). The results showed that 192 the calculated trajectories of the air masses were almost the same when the arrival height was below 193 400 m. The trajectory arrival altitude was then set to 100 m a.g.l in this study. The trajectories were 194 computed every 6 hours with an inverse time of 120 hours. The trajectories could cover China, 195 Nepal, India, Pakistan, and most of West Asia. Here, we combined the backward trajectory with 196 real-time Hg monitoring concentrations to represent the trajectories of GEM concentrations. Cluster 197 analysis was performed after the trajectory calculation. Cluster analysis can indicate the main 198 trajectory of incoming pathways and the GEM concentration indicated by the incoming trajectories.

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2.5 Principal component analysis

200 Principal component analysis (PCA) is a data reduction method that allows a number of 201 measured variables to be categorized into several factors that represent the behavior of the entire 202 dataset (Jackson, 2005). In many previous Hg studies, PCA has been used to analyze the 203 relationships between Hg and multiple pollutants and meteorological variables (Brooks et al., 2010; 204 Cheng et al., 2012; Liu et al., 2007; Zhou et al., 2019; Lin et al., 2022). Prior to running the PCA, 205 all variables were normalized by the standard deviation. To check whether PCA was the appropriate 206 method for the dataset used in this study, Kaiser-Meyer-Olkin's measure of sampling adequacy 207 (MSA>0.5) and Bartlett's Test of sphericity (P<0.05) tests were performed during data analysis. 208 Total variance and rotated scree plots were used to determine the number of factors during the PCA 209 analysis, and components with variance ≥ 1.0 were retained. Variables with high factor loadings 210 (generally >0.5) were identified as potential sources of Hg in this study.

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3.1 Atmospheric Hg Monitoring in Lhasa

Results and Discussion

Atmospheric Hg monitoring in Lhasa comprised the subsequent Indian Summer Monsoon (S-213 214 ISM) and Westerly circulation (WEC) periods from August 29, 2016 to February 2, 2017. Figure 2 215 shows the variation in atmospheric Hg concentrations at the station during the monitoring period. 216 During the whole monitoring period, the mean concentrations of GEM, GOM, and PBM at the station were 2.26 \pm 1.97 ng m⁻³, 36.4 \pm 48.9 pg m⁻³, and 54.5 \pm 119.5 pg m⁻³ (mean concentration \pm 217 218 standard deviation), respectively. During the S-ISM period, the concentrations of GEM, GOM, and 219 PBM were 2.73 ± 1.48 ng m⁻³, 38.4 ± 62.7 pg m⁻³, and 59.1 ± 181.0 pg m⁻³, respectively. While during 220 the WEC period, the concentrations of GEM, GOM, and PBM were 2.11±2.09 ng m⁻³, 35.8±43.3 pg m⁻³, and 52.9±90.1 pg m⁻³, respectively. The GEM concentrations during the S-ISM period were 221 significantly higher than those during the WEC period (p < 0.01), while the mean concentrations of 222 223 GOM and PBM in the S-ISM period were slightly higher than those in the WEC period. Overall, 224 GEM concentrations showed a decreasing trend throughout the monitoring period, with the average

225 weekly concentration decreasing from 3.21 ng m⁻³ at the beginning of the monitoring period to 1.60 226 ng m⁻³ at the end of the monitoring period, which is contrast to previous studies, which showed that 227 atmospheric Hg concentrations in the Northern Hemisphere is low in summer and high in winter 228 (Horowitz et al., 2017; Jiskra et al., 2018). The different variation trends between Lhasa and the 229 whole Northern Hemisphere may be related to the special location of Lhasa. Unexpectedly high 230 concentrations were found at irregular intervals for all Hg species. The occurrence time of these 231 high concentrations was random, and high GEM concentrations did not always occur at the same 232 time as high GOM or PBM concentrations, indicating the complexity of the Hg sources of the 233 species. For GOM and PBM, relatively comparable trends between them may be related to similar 234 sources, transport, and transformation reactions in the atmosphere.



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Figure 2 Time series of GEM, GOM, and PBM concentrations over the sampling period. The





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In early September, GOM concentrations were generally higher than PBM concentrations. In the subsequent period, PBM concentrations were always higher than GOM concentrations, which may indicate that the sources and composition of pollutants at this time were not consistent with those in the latter period. GOM and PBM may undergo mutual transformation in the atmosphere, which may be related to temperature, humidity, and atmospheric composition(Rutter and Schauer, 2007; Rutter et al., 2008), and thus the concentration distributions of GOM and PBM may also be related to the changes of local climate and atmospheric composition from S-ISM to WEC periods.

246 The GOM and PBM concentrations were higher in November and December. Since GOM and PBM

are mainly from local emissions, the changes in their concentrations may indicate that there are

248 more local sources at this period. As Lhasa enters the heating season in November-December, and

- 249 there are more local religious activities at this time, there may be more local combustion sources.
- 250 Table 1 lists the concentrations of Hg and other pollutants during the monitoring period. PM_{2.5}
- 251 concentration increased significantly in the WEC1 period, indicating the presence of more

		Table 1 S	Statistics m	etrics of s	pecies F	Ig, mete	orologic	al factors,	and othe	er polluta	unts		
		GEM	рвм	MOS	Tamp	Ціт	Wind	Solar	00	NO	Č	DM 2.2	Ő
Period	Stat	MTD		MOD	remp	111n111	speed	radiation	5	1402	õ	1 1 1 2.5	202
		(ng m ⁻³)	(pg m ⁻³)	(pg m ⁻³)	(°C)	(%)	(m s ⁻¹)	(W m ⁻²)	(mg m ⁻³)	(µg m ⁻³) (µg m ⁻²	³) (μg m ⁻³) (μg m ⁻³
	Mean	2.73	59.08	38.39	14.42	61.45	1.70	212.60	0.39	26.44	72.18	16.26	4.19
	SD	1.48	181.38	62.85	3.70	16.96	1.41	313.90	0.18	19.36	28.68	13.12	2.08
S-ISM	Median	2.36	24.50	30.30	13.90	62.00	1.30	7.00	0.30	22.00	77.00	13.00	4.00
	Min	0.40	-11.44	1.89	5.90	14.00	0.00	00.00	0.10	1.00	1.00	1.00	1.00
	Max	18.87	2165.70	988.50	24.70	93.00	13.00	1290.00	1.70	110.00	133.00	81.00	21.00
	Mean	2.39	57.55	37.46	6.90	26.01	1.28	169.56	0.64	45.24	45.39	47.97	5.51
	SD	2.35	104.15	50.98	5.75	15.08	1.39	246.66	0.51	31.01	34.44	46.19	5.35
WEC1	Median	1.79	38.80	27.83	6.90	24.00	06.0	00.00	0.50	41.00	43.00	35.00	5.00
	Min	0.31	0.00	1.89	-6.20	1.00	0.00	00.00	0.10	2.00	1.00	1.00	1.00
	Max	48.93	1797.30	774.86	21.60	94.00	9.80	973.00	4.40	152.00	133.00	458.00	145.00
	Mean	1.47	42.82	32.05	0.22	25.59	1.97	123.61	0.56	26.78	65.22	23.73	4.49
	SD	1.12	45.39	17.14	3.98	14.67	1.85	186.98	0.30	22.61	26.30	18.77	2.61
WEC2	Median	1.17	33.77	27.50	0.20	24.00	1.30	00.0	0.50	19.00	71.00	19.00	4.00
	Min	0.33	0.00	9.91	-8.80	2.00	0.00	0.00	0.20	2.00	1.00	1.00	1.00
	Max	20.86	589.43	165.06	10.20	68.00	11.20	662.00	2.60	99.00	108.00	118.00	32.00

252 particulate matter during this period. This could explain the elevated concentrations of GOM and

253 PBM from November-December.

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Table 2 shows the distributions of atmospheric Hg concentrations in some provincial capitals in China and nearby monitoring stations from literature. In general, the GEM concentration in Lhasa is low among the provincial capitals in China. The GEM concentration in other provincial capitals

258 of China was approximately 3-10 ng m⁻³. Guiyang, Chongqing, and Lanzhou were the nearest 259 provincial capitals to Lhasa, with GEM concentrations reported in the literature, all located in 260 western China. Guiyang had a very high GEM concentration due to the presence of local Hg mines 261 (Liu et al., 2011; Yang et al., 2009). The GEM concentration in Chongqing was approximately three 262 times higher than that of Lhasa. The higher GEM concentration in Chongqing was likely due to its 263 proximity to the Hg-contaminated area and large population (Yang et al., 2009). Compared to 264 Lanzhou, another high-altitude city, the GEM concentration in Lhasa was approximately half that 265 of Lanzhou, which may be owing to the overall cleaner environment with fewer local pollution sources in Lhasa (Yin et al., 2020a). The GOM and PBM concentrations in other provincial capitals 266 267 were relatively less monitored. However, GOM concentrations in Lhasa were significantly higher 268 than those in cities with GOM monitoring, such as mega-cities like Beijing and Shanghai, and even 269 higher than those in Guiyang, where Hg mines are located. GOM concentrations in provincial capitals nationwide were mainly concentrated between 3-20 pg m⁻³, whereas the GOM 270 271 concentrations in Lhasa were approximately 2-10 times higher than the average concentration in provincial capitals. The high GOM concentration in Lhasa is likely due to its high altitude. Lhasa is 272 273 exposed to much higher solar radiation and has more ice surfaces than inland areas, which may have 274 contributed to the oxidation of GEM or the re-release of GOM deposited in snow and ice (Steffen 275 et al., 2008; Dommergue et al., 2003; Song et al., 2018). In contrast, the PBM concentration in Lhasa 276 was at a lower level, only somewhat higher than that in Hefei. The monitoring period in Lhasa was 277 mainly in winter when there were more particulate matter emissions than in summer owing to 278 heating combustion. The PM2.5 concentration in Lhasa was low throughout the monitoring period 279 (Table 1), which indicated that the local particulate matter emissions were low; this may be the main 280 reason for the low PBM concentration.

	- 1				GEM	GOM	PBM	GEM (local time	diurnal var ¢/GEM cone	iation centration)	
юсацоп	annae	type	region	Montoring period	(ng m ⁻³)	(pg m ⁻³)	(pg m ⁻³)	Peak (ng m ⁻³)	Valley (ng m ⁻³)	Variation (ng m ⁻³)	reterence
Lhasa	3600	City	Southwest	8/2016-2/2017	2.26±1.97	36.4±48.9	54.5±119.5				This study
Beijing	40	City	North china	12/2008-11/2009	3.22±1.74	10.1 ± 18.8	98.2±112.7				(Zhang et al., 2013)
Hefei	30	City	East china	7/2013-6/2014	4.07 ± 1.91	3.67±5.11	30.0 ± 100.3				(Hong et al., 2016)
Shanghai	4	City	East china	2014	4.19 ± 9.13	21 ± 100	197±877				(Duan et al., 2017)
Lanzhou	1525	City	Northwest	10/2016-10/2017	4.48±2.32						(Yin et al., 2020a)
Jinan	148	City	East china	10/2015 -7/2016	4.91 ± 3.66		451.9±433.4				(Li et al., 2017)
Chongqing	300	City	Southwest	2006-2007	6.74 ± 0.37						(Yang et al., 2009)
Nanjing	25	City	East china	2011	7.9±7.0						(Zhu et al., 2012)
Guiyang	1150	City	Southwest	8/2009-12/2009	9.72±10.2	35.7±43.9	368±676				(Liu et al., 2011)
Ev-K2, Nepal	5050	Remote		11/2011-4/2012	1.2 ± 0.2			18/1.30	6/1.10	0.20	(Gratz et al., 2013)
Nam Co, China	5300	Remote		11/2014-3/2015	1.33 ± 0.24						(Yin et al., 2018)
Waliguan, China	3816	Remote		9/2007-9/2008	1.98 ± 0.98	7.4±4.8	$19.4{\pm}18.1$	6/2.30	14/1.94	0.36	(Fu et al., 2012b)
Shangri-La, China	3580	Remote		11/2009-11/2010	2.51 ± 0.73	8.22±7.9	38.32±31.26	17/2.48	6/1.71	0.77	(Zhang et al., 2015a)
Gongga, China	1640	Remote		5/2005-6/2006	3.98			11/4.45	2/3.55	06.0	(Fu et al., 2008)
QNNP, China	4267	Remote		4/2016-8/2016	1.42 ± 0.37	21.4 ± 13.4	25.6 ± 19.1	6/2.04	13/1.11	0.93	(Lin et al., 2019)
Nyingchi, China	3263	Remote		3/2019-9/2019	1.01 ± 0.27	12.8±13.3	9.3±5.9	20/1.07	6/0.96	0.11	(Lin et al., 2022)

Table 2 Comparison of atmospheric H9 concentrations at some provincial capitals in China and some nearby monitoring stations

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282 Compared to nearby monitoring stations (Table 2), Hg species concentrations in Lhasa were 283 high. Namco station is the nearest station; its altitude is 4,730 m and the distance between the two 284 stations is approximately 120 km. The GEM concentration in Namco was 1.33±0.24 ng m⁻³, which 285 was only 59% of Lhasa. This is likely because the Namco region is sparsely populated with minimal 286 local pollution and is far from major Hg pollution sources (Yin et al., 2018). Compared to the QNNP 287 (Lin et al., 2019) on Mt. Everest, which is approximately 500 km apart, the GEM, GOM, and PBM 288 concentrations in Lhasa were approximately 1.6, 1.7, and 2.1 times higher, respectively. Our 289 previous studies demonstrated that the QNNP is influenced by transported air masses from the 290 Indian subcontinent, indicating that the concentration in Lhasa is high in the Tibetan Plateau. 291 Compared to another typical highland site, Nyingchi, Lhasa had much high levels of atmospheric 292 Hg species, which may be related to the vegetation uptake effects and strong wet deposition in 293 Nyingchi (Lin et al., 2022). Among the surrounding stations, only Mt. Gongga and Shangri-La 294 stations had higher GEM concentrations than Lhasa. The GEM concentrations reported at Mt. 295 Gongga station ranged from May 2005 to July 2006. Considering that the smelting activities near 296 this site were crude at that time and there were almost no air pollution control measures, the high 297 local GEM concentrations may be strongly affected by local smelting activities and fuel combustion 298 (Fu et al., 2008). In contrast, GEM concentrations in the Shangri-La region were mainly controlled by the monsoon, and Zhang et al. (2015a) suggested that all local GEM above 2.5 ng m⁻³ are 299 300 associated with the transport of dry air carrying domestic and foreign regional anthropogenic 301 emissions. Comparing these sites only for the monsoon period, the GEM concentration in Lhasa 302 was higher than that in the Shangri-La region. As for GOM and PBM, the concentrations at the 303 Lhasa station were much higher than those in the surrounding areas. The average GOM 304 concentration in the surrounding areas was approximately 10 pg m⁻³, which was only 27% of that 305 in Lhasa, and the average PBM concentration in the surrounding areas was approximately 28 pg m⁻ 306 ³, which was only 54% of that in Lhasa. Considering that GOM and PBM are mainly from local or 307 surrounding sources or atmospheric transport (Lindberg and Stratton, 1998; Seigneur et al., 2006; 308 Lynam et al., 2014), high GOM and PBM concentrations may indicate additional local sources of 309 Hg in Lhasa.

- 310 **3.**2
 - 3.2 Unexpected high concentration events in Lhasa

To investigate the reasons for the random high atmospheric Hg concentration events in Lhasa, typical high-value occurrence processes (defined as GEM concentrations above 10 ng m⁻³ monitored more than five times on a single day) were selected for analysis in the S-ISM and WEC periods, respectively. A total of seven high GEM concentration events were identified, of which, September 3, November 10, November 19, and December 9, 2016, were selected for analysis; September 18, October 3, 2016, and January 27, 2017 were omitted due to lack of meteorological data or Hg concentration data for the proximity date.

318 During the S-ISM period (Figure 2), there was a clear peak in Hg concentration on September 319 3, while the GEM, GOM, and PBM concentrations were approximately 1.6, 1.5, and 2.3 times the 320 average daily value, respectively. Comparing the two days before and after the high-concentration 321 event (Table 3, Figure 3. a), the concentrations of the three pollutants NO₂/PM_{2.5}/SO₂, were higher 322 on September 3. High GEM concentrations were accompanied by winds of 2.12 m/s from the 323 southwest, with NO₂ and SO₂ concentrations higher than usual as well as increased PBM 324 concentrations. NO₂ and SO₂ are typical combustion source pollutants, and the presence of PM_{2.5}, 325 and PBM may indicate more combustion sources in the day. Thus, it can be inferred that the elevated 326 Hg concentration event on September 3 may have originated mainly from a combustion event 327 nearby or further away.

328 During the WEC period, significantly high values were observed on November 10, November 329 19, and December 9. On November 10 (Figure 3.b), the increase in atmospheric Hg species 330 concentrations was accompanied by significant increases in CO/NO₂/PM₁₀/PM_{2.5}/SO₂ 331 concentrations. Two similar peaks in atmospheric Hg species concentrations were also observed 332 around November 10, with relatively lower peak concentrations. During these three events, the 333 concentrations of other pollutants were higher than usual, whereas wind speeds were extremely low 334 during the event periods. In addition, extremely high PBM concentrations (297.7±189.3 pg m⁻³, 335 maximum 621.2 pg m⁻³) were observed at midnight on November 12 which, considering the 336 extremely low wind speed and the presence of a low height nocturnal boundary layer, may indicate 337 that the high concentrations originated from local sources.

Table	3 Compara	ation of th	ne polluta	nt concer	itrations '	with the t	wo days l	before an	d after th	e high H	g concent	ration ev	ents
Iter	u	9.1-2	9.3	9.4-5	11.8-9	11.10	11.11- 12	11.17- 18	11.19	11.20- 21	12.7-8	12.9	12.10- 11
GEM	(ng m ⁻³)	2.72	4.63	3.03	1.47	3.16	2.96	2.15	6.87	2.66	2.76	5.43	3.24
PBM	(pg m ⁻³)	25.32	64.17	31.52	36.31	47.21	94.80	53.41	74.78	55.40	38.71	66.74	49.05
GOM	(pg m ⁻³)	40.95	63.84	42.55	32.76	39.48	42.03	35.29	119.39	44.86	43.84	69.65	43.75
Temp.	()°C)	16.61	15.86	15.32	7.06	7.76	8.80	4.20	4.49	4.94	3.13	2.88	2.77
Hum.	(%)	58.47	67.89	68.71	19.96	22.02	21.04	21.82	23.35	25.67	23.50	26.86	26.09
Wind	(m s ⁻¹)	2.37	1.46	1.49	1.47	0.93	1.17	1.05	0.71	0.75	0.88	0.89	0.67
Barometer	(hPa)	651.62	651.08	652.39	654.90	654.33	652.31	655.11	652.18	653.57	657.99	655.66	653.08
rainfall	(mm)	0.01	0.17	0.36	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
Solar Rad.	(W m ⁻²)	241.20	201.15	214.39	189.02	194.45	182.30	166.14	161.59	180.02	141.61	139.69	136.64
CO	(mg m ⁻³)	0.34	0.36	0.32	0.39	0.76	0.63	0.64	0.71	0.72	0.67	0.80	0.74
NO_2	$(\mu g m^{-3})$	20.44	26.00	21.77	41.38	57.64	66.48	54.58	53.58	55.27	48.87	55.33	56.23
O_3	$(\mu g m^{-3})$	85.73	77.42	72.56	59.27	57.43	49.46	34.73	34.79	29.51	31.37	28.38	26.81
PM_{10}	(µg m ⁻³)	33.90	32.78	26.95	80.71	130.07	155.44	107.98	119.79	146.29	125.82	137.74	143.13
$PM_{2.5}$	$(\mu g m^{-3})$	11.98	17.10	10.98	31.20	75.07	64.85	51.21	50.13	53.83	62.80	65.46	62.71
SO_2	(µg m ⁻³)	4.23	5.89	4.91	3.83	6.07	5.71	6.50	8.92	9.44	6.00	6.88	69.9

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On November 19 (Figure 3.c), there was a significant increase in all atmospheric Hg species concentrations. During the elevated period, the average GEM, GOM, and PBM concentrations were 17.94 ± 10.54 ng m⁻³, 302.5 ± 218.5 pg m⁻³, and 162.4 ± 54.0 pg m⁻³, respectively, which were 8.3, 5.7,

342 and 4.6 times higher than the average concentrations on November 17-18. However, only a slight increase in the peak CO/ PM_{2.5}/SO₂ concentration was observed during the event period. Therefore, 343 344 the increase in atmospheric Hg species concentrations may indicate that this event was unique, with certain sources of high Hg concentrations. The low wind speed and low nocturnal boundary layer 345 height both contributed to the accumulation of atmospheric Hg species, which also indicated that 346 347 the elevated Hg was from local sources. Similar to the event on November 19, sharp increases in 348 atmospheric Hg species concentrations were observed only on December 9 (Figure 3.d). The 349 atmospheric Hg species concentration increase events during the WEC period occurred at night and, given that it coincided with the winter heating period, local combustion sources may be the unique 350 351 sources that may contribute to the Hg concentration increase.

















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Figure 3 Comparison of the pollutant concentrations with those at the two days before and

after the high Hg concentration events. The dotted lines indicate the area where the Hg

concentrations are significantly elevated

356 **3.3 Diurnal variation of atmospheric Hg in Lhasa**

357 Figure 4.a shows the diurnal variation of atmospheric Hg in Lhasa during the monitoring period.

358 The mean GEM concentration during this period was 2.26±1.97 ng m⁻³, and the maximum 359 difference of hourly average GEM diurnal variation was 1.89 ng m⁻³. GEM concentration was low during the day; the lowest concentration of the day appeared from 14:00-18:00 (UTC+8). The GEM 360 concentration kept increasing and reached the peak at midnight. During the night, the GEM 361 concentration maintained in high value with little dissipation. Subsequently, the GEM concentration 362 363 decreased rapidly with the sunrise. Overall, the diurnal variation of GEM concentration was similar to CO, NO₂, and relative humidity, and opposite to O₃ and wind speed. The high GEM 364 365 concentrations at night can be attributed to the lower boundary layer height at night (average at 131 m a.g.l. during the observation period, between 19:00 and 07:00, UTC+8) and the lower wind speed 366 367 (Figure 4). Also, the average night temperature in Lhasa during the monitoring period was 5.8°C, 368 and the residents heating combustion may bring about the release of GEM. After sunrise, the 369 boundary layer height increased rapidly and the diffusion quickly occurred in the lower atmosphere. 370 During the increase of wind speed, airflows were carried from other regions with few populations, 371 which may lead to a decrease in GEM concentrations. For both GOM and PBM, the higher GOM 372 concentrations occurred during the day and higher PBM concentration occurred during the night. 373 No obvious increase occurred for GOM and PBM concentrations at the same time, which may 374 indicate that there is no local common external source, and the variation of GOM and PBM concentrations may come from the gas-particle redistribution process of atmospheric Hg(II). 375



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Figure 4 Diurnal variations of Hg species, concentrations of other pollutants, and meteorological information from S-ISM to WEC periods. The short horizontal line represents the concentration error range for each time period.

In particular, during the S-ISM period (Figure 4.b), the diurnal variation in GEM 380 381 concentrations fluctuated frequently, which indicates that the factors affecting the concentration 382 variation were very complicated. The mean concentration of GEM during this period was 2.73 ng 383 m⁻³, and the maximum difference of GEM diurnal variation was 1.40 ng m⁻³. September was the 384 ending period of the Indian summer monsoon, and the external transboundary transport of air masses 385 by the monsoon may have been weakened. Simultaneously, the average temperature in September 386 was high (14.4°C) without residents heating during the night, which may be the reason for the small 387 diurnal variation in GEM concentrations. Overall, GEM concentrations remained lower during the 388 day, increased, and remained high at night. No clear pattern of variation was observed in GOM and 389 PBM during this period.

390 During the WEC1 period, the diurnal variation in GEM concentrations was exceedingly large. The mean GEM value was 2.39 ng m⁻³ with great GEM diurnal variation value (3.01 ng m⁻³). The 391 392 diurnal trend was roughly the same as the whole monitoring period, but the lowest value was 1.06 ng m⁻³ and the highest value was 4.07 ng m⁻³. The increase in night concentrations may be related 393 394 to the beginning of the heating season, whereas the burning of yak dung, firewood, or other fuels 395 for residential heating may release GEM and PBM, and accumulate within the boundary layer at 396 night, leading to higher GEM concentrations (Rhode et al., 2007; Xiao et al., 2015; Chen et al., 397 2015). In contrast, low day values may be related to the wind field. Although the wind speed during 398 the WEC1 period was similar to that during the S-ISM period, westerly circulation was prevalent in 399 Lhasa during the WEC1 period, with air masses coming from the sparsely populated area with lower 400 GEM concentrations. The westerly winds carry away the locally accumulated GEM air masses in 401 Lhasa and bring in low GEM concentration air masses from west of Lhasa. During this period, the 402 diurnal variation in GOM concentrations was not clear.

403 The influence of wind flow was evident during the WEC2 period. During this period, GEM 404 concentrations in Lhasa were the lowest and close to 1.00 ng m^{-3} in the late afternoon when wind 405 speeds were the highest. Nighttime concentrations were lower in the WEC2 period than in the WEC1 period, probably because wind speeds were also high at night during the WEC2 period (19:00-4:00 UTC+8, mean wind speed 1.74 m/s). The heating season was still in progress during the WEC2 period, but GEM emissions from heating were likely carried away by the continuous flow of air masses. Therefore, the mean GEM concentration in Lhasa during the winter was likely strongly influenced by wind speed.

411 Compared to the diurnal variation patterns obtained from monitoring in other remote areas of 412 China (Figure S3), the diurnal variation of GEM in Lhasa is more unique, and a wide diurnal 413 variation in concentration was observed. Valleys of GEM concentrations in the afternoon were also 414 observed in Waliguan and Namco, but the concentration variation ranges were small in both areas. 415 This could be related to the regional characteristics of the three sites. Waliguan is located in a 416 mountainous area with no significant local anthropogenic sources, and is mainly influenced by 417 valley winds (Fu et al., 2012b). Namco is located in the central Tibetan region away from 418 anthropogenic sources. Although the wind speed was also high in the afternoon in Namco (Yin et 419 al., 2018), it is likely that the diurnal variation in atmospheric GEM concentrations was small 420 because the atmosphere was well-proportioned and there were no local sources; therefore, the 421 changes in wind speed might not affect the GEM concentration. In contrast, with the difference in 422 GEM concentrations between Lhasa and the transmitted external air mass, wind speed could 423 significantly influence the GEM concentrations in Lhasa. Meteorological conditions may have 424 played a more important role. We conjecture that the high nighttime concentrations in Lhasa may 425 originate mainly from local emissions, while the high wind speed and mixing of clean external air 426 masses in the afternoon reduce the local GEM concentrations.

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3.4 Analysis of factors affecting atmospheric Hg concentration in Lhasa

Overall, four principal components were obtained for each period, from S-ISM to WEC2, using PCA analyze, to analyze the relationships between Hg and multiple pollutants and meteorological variables. The Hg monitoring data, meteorological factors, and pollutant data in Lhasa were statistically and dimensionally reduced by PCA to analyze the relationships among them. The components related to Hg were selected and analyzed separately for each period, and the principal components were extracted (Table 4). According to the variable's loadings on each component, they were assigned as Special Hg-related factor, Local emission factor, and Wind factor.

Tentative Identification	Spee	cial Hg-re	lated	Local e	mission	W	ind
preiod	S-ISM	WEC1	WEC2	WEC1	WEC2	S-ISM	WEC2
GEM	0.87	0.63	0.9	0.44	0.3	-0.21	-0.13
PBM	0.96	0.79	0.63				
GOM	0.95	0.87	0.95	0.11	0.1		
Temp				-0.29	0.2		0.92
Hum		-0.19	0.17	-0.26	-0.18	-0.26	-0.75
Wind_Speed	-0.1	-0.14		-0.37		0.87	0.91
Rain							
Solar_Rad.				-0.21			
СО	0.17		0.11	0.87	0.69	-0.22	0.51
NO_2		0.13		0.92	0.96	-0.25	
O ₃			-0.36	-0.89	-0.83	0.53	
PM_{10}	-0.14			0.86	0.69		0.14
PM _{2.5}	-0.16		0.16	0.83	0.86		
SO_2	0.26	0.17	0.86	0.75			
Variance Explained	19.6	13.69	21.88	35.13	24.94	9.25	18.32

Table 4 PCA factor loadings (varimax rotated factor matrix) for Hg in Lhasa, China.

Note: variables with high factor loadings (> 0.5) were marked in bold. For readability, variables with very low factor loadings (< 0.1) are not presented.

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436 A special Hg-related factor (Factor 1) was assigned owing to the continuous high positive 437 loading of GEM, GOM, and PBM from the SISM to the WEC periods; only high positive loading of SO₂ was found in the WEC2 period. For this factor, there was no significant relationship with 438 439 any other meteorological factors or pollutants. The high positive loading of this factor for all three 440 Hg species and the low correlation with meteorological factors and other pollutants may indicate 441 that this is a specific source of Hg. As the short transmission distance of the GOM and PBM, the 442 special Hg-related source should be closer to Lhasa. However, no particular source of Hg around 443 Lhasa has been reported in the literature, so the source indicated by this factor remains unclear.

Factor 2 had slight positive relationship with GEM and GOM, high positive loading of CO/NO₂/PM₁₀/PM_{2.5}/SO₂, and high negative loading of O₃ during the WEC period. The high positive loadings of CO/NO₂/PM₁₀/PM_{2.5}/SO₂ may indicate that this factor is highly correlated with the combustion source. As O₃ concentration rapidly decreased after sunset (Figure 4. a), high negative loading of O₃ further indicated that this factor represented events during the nighttime with low O₃ concentrations. Since the WEC period overlapped with the heating season in Lhasa, Factor
2 may be strongly associated with heating combustion.

451 The wind factor (Factor 3) involves high positive loading of wind speed and low negative 452 loading of GEM concentration during the S-ISM and WEC2 periods. This factor reveals the 453 scavenging effect of wind on the local GEM in Lhasa. For high wind speed, the air masses with low 454 GEM concentrations in the surrounding area mixed with the air masses in Lhasa, leding to a reduction in the GEM concentration. Concurrently, some GEM in Lhasa was carried away from the 455 456 city by wind. At wind speed greater than 4m s⁻¹, the average GEM concentration in Lhasa was 457 1.31 ± 0.93 ng m⁻³, which is similar to the average concentration of Nam Co (1.33 ± 0.24 ng m⁻³) in the Tibetan hinterland. Factor 3 was consistent with the analysis of the wind effect on the diurnal 458 variation of Hg concentrations (Section 3.2), indicating the effect of strong winds on urban Hg 459 460 removal.

461

3.5 Atmospheric Hg source trajectories and potential source regions in the Lhasa area

462 Figure 4 shows the GEM backward trajectory paths from the S-ISM period to the WEC2 period 463 in Lhasa. During the S-ISM period, most trajectories (cluster 1, representative GEM concentration 464 of 2.66 ng m⁻³, 87.53% of the trajectory during this period, Table S1) originated from or passed 465 through the south of Lhasa. The source of the trajectory points directly to the Indian Ocean, likely 466 as these transported air masses are still subject to Indian monsoon action in September. According 467 to the UNEP Hg emission inventory (Unep, 2013), there are few anthropogenic emissions along this trajectory, indicating that the GEM may originate from the Indian Ocean or locally from Lhasa. 468 469 Clusters 2 and 3 may indicate the trajectories of air masses driven by westerly circulation, which 470 had a low proportion in the S-ISM period, with slightly different GEM concentrations from different 471 sources.

During the WEC1 period, the driving factor of the air mass gradually shifted from Indian monsoon to westerly circulation. Clusters 2 and 3 are trajectories driven by the higher-height westerly circulation during the WEC1 period with a higher proportion than in the S-ISM period. Cluster 1 came from the southwest of Lhasa, and the air mass moved along the Himalayas before entering the Tibetan Plateau and was transported to Lhasa. Compared to the S-ISM period, the GEM concentration of this trajectory decreased slightly, which may be related to the source of the air mass 478 and the areas it passes through. WEC2 showed little change in trajectory sources compared to WEC1, 479 but all trajectory concentrations decreased significantly. Both WEC1 and WEC2 were in winter, 480 and both had similar trajectories related to the driving wind field. Significantly decreasing GEM concentrations may suggest a local influence in Lhasa City. The local GEM in Lhasa consists of 481 background concentrations superimposed on local emissions, and the share of local emissions 482 483 decreases under better dispersion conditions (higher wind speeds) during the WEC2 period. The GEM concentration during the WEC2 period in Lhasa was only 0.16 ng m⁻³ higher than that during 484 485 the WEC period in the QNNP region $(1.31 \pm 0.42 \text{ ng m}^{-3})$ (Lin et al., 2019)).

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periods. The thickness of the line represents the ratio of the cluster in the time period, the background is the global Hg emission inventory developed by UNEP (2013a).

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3.6 Comparison of three typical sites on the Tibetan Plateau

491 Different levels and variation patterns of Hg species concentrations were observed in the
492 Qomolangma National Nature Preserve (QNNP, 4,276 m) (Lin et al., 2019), Nyingchi (SET, 3,263
493 m) (Lin et al., 2022), and Lhasa (3,650 m) areas, which are typical monsoon-influenced, canyon,
494 and urban areas on the Tibetan Plateau, respectively. Predictably, Hg species concentrations were

495 generally lower on the Tibetan Plateau, while background areas with few populations on the Plateau 496 were lower than those on the plains in mainland China, and cities on the Plateau were lower than 497 those on the plains. Although the transboundary transport of pollutants has received considerable 498 notice (Huang et al., 2016b; Wang et al., 2018; Lin et al., 2019; Zhu et al., 2019; Feng et al., 2019), 499 the comparison between QNNP and Lhasa indicates that the contribution of local anthropogenic 500 sources may be significant for atmospheric pollutant concentrations. The comparison also suggests 501 that atmospheric Hg emissions from urban residential life may be an important source that can be 502 valuable in inventory studies.

503 Vastly different patterns of Hg species concentrations were found in the QNNP and SET due 504 to differences in geography and vegetation cover. Atmospheric Hg transported to QNNP is subject 505 to a combination of monsoonal action and the pumping effect of glacial winds. However, 506 atmospheric Hg entering the SET area has a slow elevation increasing path, and abundant 507 precipitation and vegetation have a significant trapping effect on atmospheric Hg. While for Lhasa, 508 a direct effect of local wind fields on the accumulation or rapid removal of GEM concentrations 509 was found. This suggests that although global transport is important in the Hg cycle, the pollution 510 at each location is likely to be strongly influenced by the local environment, even in exceptionally 511 clean areas such as the Tibetan Plateau. However, there are still relatively few studies on GOM and 512 PBM in the Tibetan Plateau, and more comprehensive investigations are required on how changes 513 in Hg speciation transformation on snow and ice surfaces affect the environment, the effect of stratospheric intrusion on GOM concentrations (which is common on the Plateau), and the effect of 514 515 particulate matter on Hg(II) gas-particle partitioning, which would help to understand the Hg species 516 change on the Plateau and throughout the world.

517 **4.** Conclusions

Lhasa is the largest city on the Tibetan Plateau; thus, its atmospheric Hg concentrations represent the highest level of atmospheric Hg pollution in this area. Unexpectedly high concentrations of atmospheric Hg species were found in Lhasa. The GEM concentrations were higher than the Northern Hemisphere background concentrations, and the GOM and PBM concentrations were high among Chinese cities. Monitoring of atmospheric Hg in Lhasa showed that the mean concentrations of GEM, GOM, and PBM during the S-ISM period (2.73 ± 1.48 ng m⁻ ³, 38.4 ± 62.7 pg m⁻³, and 59.1 ± 181.0 pg m⁻³, respectively) were higher than those during the WEC period (2.11 ± 2.09 ng m⁻³, 35.8 ± 43.3 pg m⁻³, and 52.9 ± 90.1 pg m⁻³, respectively). Combined with the trajectory analysis, the high atmospheric Hg concentrations during the S-ISM phase may have originated from external long-range transport.

528 Analysis of the overall concentration changes revealed some irregular and sudden high 529 atmospheric Hg concentration events in Lhasa. Analysis of these events suggests that local sources 530 (such as combustion events) can cause severely elevated concentration events under low wind 531 speeds and the presence of a low-height nighttime boundary layer. Analysis of the diurnal variation 532 of concentrations confirmed that low wind speeds and a low height nocturnal boundary layer would 533 lead to the elevated local Hg concentrations. In contrast, higher wind speeds could rapidly remove 534 atmospheric Hg from Lhasa. PCA analysis of the influencing factors indicates that local sources, 535 especially special Hg-related sources, are important factors influencing the variability of 536 atmospheric Hg. The PCA analysis also indicated the important role of higher wind speeds in 537 reducing atmospheric Hg concentrations in the urban areas of Lhasa, likely owing to the large Hg 538 concentration difference between Lhasa and surrounding areas.

539 Up to this study, we have obtained atmospheric Hg monitoring data from four typical areas of 540 the Tibetan Plateau: Lhasa, QNNP, SET, and Namco. The atmospheric Hg concentrations in the 541 background areas were at or below the average GEM concentration of Northern Hemisphere, with 542 higher levels in the urban area of Lhasa. Factors such as long-range transport of atmospheric Hg, 543 effects of local meteorological conditions, local glaciers, etc., were considered in these studies. 544 Further monitoring of additional areas and regional simulations are required to confirm the 545 atmospheric Hg transport patterns and fluxes.

546

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- 552 Data availability. All the data presented in this paper can be made available for scientific

- 553 purposes upon request to the corresponding authors.
- Author contributions. HL,XW, YT, QZ and XY designed the research and performed field
- measurements. HL and YT performed the data analysis and model simulations. HL led the paper
- writing. LC,CY, ZC, QZ, SK, JL, JS and BF contributed to the scientific discussion and the paper
- 557 preparation.
- 558 **Competing interests.** The authors declare that they have no conflict of interest.
- 559

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