Speciated atmospheric mercury at Waliguan Global Atmospheric Watch station in the northeastern Tibetan Plateau: implication of dust related sources for particulate bound mercury

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1 Abstract

2 To understand the ambient levels and sources of atmospheric mercury (Hg) in the Tibetan 3 Plateau, a full-year continuous measurement of speciated atmospheric mercury was conducted at 4 Waliguan (WLG) Baseline Observatory (3816 m a.s.l.) from May 2012 to April 2013. Mean concentrations (±1SD) of gaseous elemental mercury (GEM), gaseous oxidized mercury (GOM) 5 6 and particulate bound mercury (PBM) during the whole study period were 1.90 ± 0.80 ng m⁻³, 12.0 7 \pm 10.6 pg m⁻³ and 65.4 \pm 63.2 pg m⁻³, respectively. Seasonal variations of GEM were very small, 8 while those of PBM were quite large with mean values being four times higher in cold (102.3 ± 66.7 9 pg m⁻³) than warm (22.8 \pm 14.6 pg m⁻³) season. Anthropogenic emissions to the east of Tibetan 10 Plateau contributed significantly to GEM pollution at WLG, while dust particles originated from 11 desert and Gobi regions in Xinjiang province and Tibetan Plateau to the west of WLG were 12 responsible to PBM pollution at WLG. This finding is also supported by the significant positive 13 correlation between daily PBM concentration and daily cumulative absorbing aerosol index (AAI) 14 encountered by air masses transported during the preceding two days.

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Keywords: Speciated atmospheric mercury, Particulate bound mercury, Anthropogenic mercury
 emissions, Dust related sources

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19 Introduction

20 Mercury (Hg) is a toxic pollutant of global concern due to its long lifetime in air, 21 bioaccumulation in aquatic system, and detrimental impacts on human and animal health. 22 Atmospheric Hg is operationally defined in three forms, i.e., gaseous elemental mercury (GEM), 23 gaseous oxidized mercury (GOM), and particulate bound mercury (PBM). These Hg species can be 24 transformed among each other through complex physical and chemical processes (Lyman et al., 25 2020; Selin, 2009). For example, GEM can be oxidized to form GOM, and GEM and/or GOM can 26 adsorb on atmospheric aerosols to form PBM (Ariya et al., 2015). GEM has a lifetime in air of 0.5-27 2 years while GOM and PBM only have a lifetime of hours to weeks (Ariya et al., 2015; Murphy et 28 al., 2006). Because of their different lifetimes, GEM can be transported globally via atmospheric 29 circulation whereas PBM is limited to regional transport (Pirrone et al., 2010). These Hg species 30 can be removed from the atmospheric through dry and wet deposition processes. Once deposited to 31 earth's surface, Hg can be converted to methylmercury by biological processes, which can cause 32 potential risks to ecological and human health (Jonsson et al., 2014; Wright et al., 2018). On the 33 other hand, Hg accumulated in soil and water bodies can be emitted into the atmosphere, which 34 plays an important role in the global atmospheric Hg cycle (Obrist et al., 2018; Wang et al., 2016).

36 Spatiotemporal variations of atmospheric Hg and Hg speciation fractions within the total Hg 37 are controlled by many factors, among which anthropogenic emissions are important ones (Driscoll 38 et al., 2013; Fu et al., 2012b). Global anthropogenic Hg emissions to the atmosphere were estimated to be 2224 Mg yr⁻¹, of which 82-87%, 10-18% and 3-4% were in the form of GEM, GOM and PBM, 39 40 respectively (AMAP/UNEP 2018). In China, concentrations of GEM and PBM were generally 41 elevated compare to the observations in Europe and America. The fractions of PBM in total 42 atmospheric Hg in urban areas of China were in the range of 5.2–17.2%, higher than those from 43 total anthropogenic Hg emissions (Fu et al., 2015; Zhang et al., 2015b). In addition, observations of 44 speciated atmospheric Hg in China in both urban and rural areas of China showed generally higher 45 PBM than GOM levels (Fu et al., 2015b), which is in contrast with the higher GOM than PBM 46 fractions in the total anthropogenic Hg emissions in China (Zhang et al., 2015b). These findings 47 indicate that additional emission sources and other physical and chemical processes contributed to 48 the elevated PBM concentrations in China. For example, natural sources, such as biomass burning 49 and dust related sources, and gas-particle partitioning can also produce PBM (Amos et al., 2012; 50 Obrist et al., 2008). The impact of these sources and processes on atmospheric PBM, although is 51 potentially important, has not been well investigated by previous studies.

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53 Speciated atmospheric Hg has been monitored in China in recent years, but observations in the 54 Tibetan Plateau region are very limited. The Tibetan Plateau, also known as the third pole of the 55 world with an average altitude of over 4000 m a.s.l., is an ideal place for assessing transport and transformation of atmospheric pollutants in China and other Asian regions (Chen et al., 2019; 56 57 Loewen et al., 2007). The Tibetan Plateau is surrounded by East Asia and South Asia, which are the 58 two most important source regions of atmospheric Hg in the world (Zhao et al., 2013). The 59 Taklamakan and Gobi deserts are located to the west and north, respectively, of the Tibetan Plateau. 60 Since the Tibetan Plateau region is strongly impacted by westerlies, Indian Summer Monsoon and 61 East Asia Summer Monsoon (Fig. 1), it is affected by air pollutants from anthropogenic and natural 62 source regions adjacent to the Tibetan Plateau (Che et al., 2011). Previous studies postulated that air 63 masses passing over the urban and industrial areas in western China and South Asia were important 64 sources of atmospheric GEM at Waliguan and Nam Co (4730 m a.s.l.) on the northeastern edge and 65 Midlands of the Tibetan Plateau, respectively (Fu et al., 2012a; Yin et al., 2018). At Shangri-La, 66 located on the southeastern edge of the Tibetan Plateau, the identified atmospheric GEM source regions were also located in Southeast Asia and mainland China (Zhang et al., 2015a). Speciated 67 68 atmospheric Hg measurement was only carried out during a warm season in Qomolangma Natural 69 Nature Preserve (4276 m a.s.l.) of Tibetan Plateau (Lin et al., 2019). Low GEM concentrations (means: 1.33 to 1.42 ng m⁻³), but elevated PBM concentrations (means: 25.6 to 49.0 pg m⁻³), were 70

observed in middle and southern Tibetan Plateau as compared to those in rural central and eastern China (Yin et al., 2018; Lin et al., 2019). To date, long-term observations of speciated atmospheric Hg in the Tibetan Plateau region are still lacking, limiting our capacity to fully understand the spatiotemporal patterns of Hg in this region and associated source regions and controlling factors.

In this study, one-year continuous monitoring of speciated atmospheric Hg was carried out at WLG in Tibetan Plateau region. Data were analyzed carefully for exploring the ambient levels, seasonal and diurnal patterns, and source regions of speciated atmospheric Hg in this region. Knowledge generated from this study is needed for establishing future emission control policies in order to preserve many sensitive ecosystems in this region.

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82 2 Materials and methods

83 **2.1 Measurement site**

84 The measurement site is situated at the summit of Mt. Waliguan located at northeastern edge 85 of the Tibetan Plateau in northwest China. It is a station known as Waliguan (WLG) Baseline 86 Observatory (100°54' E, 36°17' N, 3816 m a.s.l.) (Fig.1), which is the only station in inner Asia in 87 the Global Atmospheric Watch (GAW) program of World Meteorological Organization (WMO). 88 This area has a typical high plateau continental climate, and western winds dominant at the site (Fu 89 et al., 2012a; Okamoto and Tanimoto, 2016; Xu et al., 2018). WLG is mainly surrounded by the arid 90 and semi-arid grassland and desert lands. The population density is very low and industrial activities 91 sparsely distributed within 80 km around WLG. Anthropogenic Hg emissions in Qinghai province 92 is relatively low, and are mostly located to the east of WLG (Fu et al., 2015; Sun et al., 2020; Wu 93 et al., 2006). The Taklimakan Desert and Gobi Desert of Xinjiang province are located to the west 94 of WLG, and the Gobi Desert of Hexi Corridor and southern Inner Mongolia are located to the north 95 of WLG (Fig. 1).

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97 2.2 Sampling method

98 2.2.1 Measurements of speciated atmospheric mercury

High-temporal resolution measurements of GEM, GOM and PBM were carried out using the 2537B-1130-1135 Atmosphere Speciation Mercury Analysis System (Fig. 1, Tekran Inc., Toronto, Canada) from May 2012 to April 2013. The Tekran Model 2537B Mercury Vapour Analyzer provides continuous analysis of GEM in air at 0.1 ng m⁻³ detection limit. The instrument samples air and captures vapour phase Hg on the cartridges containing ultra-pure gold adsorbent media. The amalgamated Hg is thermally desorbed and detected using Cold Vapour Atomic Fluorescence Spectrometry (CVAFS). The Model 1135 Particulate Mercury Unit, together with the Model 1130 106 Mercury Speciation Unit, allows the Model 2537B Mercury Vapor Analyzer to simultaneously 107 monitor and differentiate between GEM, GOM and PBM (fine fraction, < 2.5 um) in ambient air. 108 KCl-coated annular denuders was installed in the specially designated location of Model 1130 109 Mercury Speciation Unit before the instrument starts running. The typical detection limits for the 110 GOM and PBM measurements during a 2-hour sampling duration are 2 pg m^{-3} , respectively 111 (http://www.tekran.com/files/Tekran-2537B-Unique-Features.r103.pdf). The instrument's 112 workflow is controlled by the controller which is capable of executing an automatic sampling and 113 analysis program (Fu et al., 2016; Lindberg et al., 2002). Due to the low air pressure at WLG, the 114 total sampling flow rate of the GOM and PBM was programed to be 6.6 lpm (referenced to standard temperature and pressure conditions). The Tekran 2537 was sampling GEM at a flow 115 116 rate of 0.6 lpm, while the Tekran 1130 pump module pulled additional air at 6 lpm. A 2-hour 117 duration was selected for GOM and PBM sampling, during which GEM is continuously 118 measured at a 5-minute interval.

119 This system has been used to monitor atmospheric Hg species worldwide, including the North 120 America Atmospheric Mercury Network (AMNet) and the Global Mercury Observation System 121 (GMOS) (Lan et al., 2012; Sprovieri et al., 2016). In this study, data QA/QC procedure followed 122 the GMOS Standard Operation Procedure and Data Quality Management (D'Amore et al., 2015). 123 Although KCl-coated annular denuders have been the most popular and widely applied method for 124 measuring ambient GOM, large analytical uncertainties in GOM may exist due to the trace level 125 and complicated chemical compounds of GOM which may not be fully collected by denuders (Ariya et al., 2015; Cheng and Zhang, 2017; Gustin et al., 2015; Gustin et al., 2019). Analysis and 126 127 discussions presented in this study are mostly focused on GEM and PBM, considering the larger 128 uncertainties in GOM than GEM and PBM.

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130 2.2.2 Meteorological data and backward trajectory calculation

131 Meteorological parameters, including air temperature (AT), relative humidity (RH), rainfall 132 (RF), wind direction (WD) and wind speed (WS) were obtained from the local weather station at 133 WLG. In order to identify the effect of long-range transport of Hg emissions on the distributions of 134 atmospheric Hg at WLG, backward trajectories arriving the site at 100 m above the ground were 135 calculated every 4 hours using the TrajStat software and gridded meteorological data from the Air 136 Resource Laboratory, National Oceanic and Atmospheric Administration (NOAA) (Wang et al., 137 2009). To investigate the source regions potentially influencing GEM and PBM concentrations at WLG, a weighing algorithm based on measured concentrations, known as the concentration 138 139 weighted trajectory (CWT) approach, was applied in this study. In this procedure, the CWT value indicates the source strength of a $0.5^{\circ} \times 0.5^{\circ}$ grid cell (CWT_{ij}) to the WLG and is defined as: 140

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$$C_{ij} = \frac{1}{\sum_{l=1}^{M} \tau_{ijl}} \sum_{l=1}^{M} C_l \tau_{ijl}$$

where C_{ij} is the average CWT value of speciated atmospheric Hg in the grid cell (*i*,*j*), C_l is the 4-h means of measured Hg concentration at WLG, τ_{ijl} is the number of trajectory endpoints in the grid cell (*i*,*j*) associated with the C_l sample, and *M* is the number of samples that have trajectory endpoints in grid cell (*i*,*j*). A point filter is applied as the final step of CWT to eliminate grid cells with few endpoints. Weighted concentration fields show concentration gradients across potential sources. This method helps determine the relative significance of potential source regions (Cheng et al., 2013; Zhang et al., 2016).

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150 **2.2.3 Ancillary parameters and analysis**

151 Anthropogenic emissions of GEM and PBM in $0.5^{\circ} \times 0.5^{\circ}$ grid cells in the studied domain 152 were obtained from the 2010 global emission dataset developed by the Arctic Monitoring and 153 Assessment program (AMAP) (AMAP/UNEP, 2013). Gridded monthly biomass burned areas at 154 0.25° spatial resolution were obtained from the fourth version of the Global Fire Emission Database 155 (GFED4) (Giglio et al., 2013). Absorbing Aerosol Index (AAI) constitutes one of the most useful 156 space-borne data sets, offering temporal and spatial information on UV absorbing aerosols (black 157 carbon, desert dust) distributions. Desert dust and biomass burning related aerosols are the dominant 158 aerosol types detected by the AAI, and AAI is therefore a useful parameter for qualitatively identifying the dust and biomass burning related sources. The AAI data are available on daily and 159 160 monthly basis at a spatial resolution of 1×1 degree. Generally, non-absorbing aerosols (e.g., sulfate 161 and sea-salt) yield negative AAI values, UV-absorbing aerosols (e.g., dust and smoke) yield positive 162 AAI values, and clouds yield near-zero values (Prospero et al., 2002). Such information can be used 163 for identifying distinct desert dust aerosol sources and analyzing dust and smoke transport patterns 164 (Chiapello et al., 1999; Kubilay et al., 2005; Moulin and Chiapello, 2004). A detailed description of 165 the AAI product is given in (Herman et al., 1997; Torres et al., 1998). In this study, Global monthly 166 gridded (1×1 degrees) AAI products during our study period were obtained from the Tropospheric Emission Monitoring Internet Service (TEMIS) (http://www.temis.nl/airpollution/absaai/). 167

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- 169 To study the effect of dust related sources on the variations in PBM concentration at WLG, we 170 calculated the daily cumulative AAI (Σ AAI) based on the 2-day backward trajectory and gridded 171 AAI data. Further analysis between the daily Σ AAI and mean PBM concentration were conducted 172 to assess the effect of dust related sources on the variations of PBM at WLG.
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174 **3 Results and discussion**

175 **3.1 Concentrations of GEM, GOM and PBM**

176 Time series of hourly speciated atmospheric Hg concentrations is shown in Fig. 2. Mean ± 1 sd (median) concentrations of GEM, GOM and PBM at WLG during the whole sampling campaign 177 were 1.90 ± 0.80 (1.59) ng m⁻³, 12.0 ± 10.6 (8.6) pg m⁻³ and 65.4 ± 63.2 (39.9) pg m⁻³, respectively. 178 179 Mean GEM level at WLG was relatively higher than the background levels in the Northern 180 Hemisphere (1.5-1.7 ng m⁻³) (Fu et al., 2015; Sprovieri et al., 2016). Mean GEM concentration at 181 WLG was relatively lower than that early observed in Mt. Gongga (mean = 3.98 ± 1.62 ng m⁻³, 1sd) 182 and Shangri-La (mean = 2.55 ± 0.73 ng m⁻³, 1sd) located on the eastern edge of the Tibetan Plateau in 2005 and 2009 respectively, but much higher than that later observed in Qomolangma Natural 183 Nature Preserve (mean = 1.42 ± 0.37 ng m⁻³, 1sd) and Nam Co (mean = 1.33 ± 0.24 ng m⁻³, 1sd) in 184 185 the inland Tibetan Plateau in 2016 and 2014 respectively (Fu et al., 2012b; Fu et al., 2009; Lin et 186 al., 2019; Yin et al., 2018; Zhang et al., 2015a). In general, atmospheric GEM levels in remote areas 187 are closely related to the regional atmospheric Hg budget. The inland Tibetan Plateau is sparsely 188 populated and with no large-scale industrial activities. However, Some monitoring sites on the 189 northeastern and eastern edges of the Tibetan Plateau, such as WLG, Mt. Gongga and Shangri-La, 190 are not too far away from the anthropogenic Hg source regions in middle and eastern China, and 191 thus were impacted by anthropogenic Hg emissions through long-range transport, which explained 192 the relatively higher GEM concentrations at these stations than inland Tibetan Plateau stations (Fu 193 et al., 2008; Zhang et al., 2015a). The impact of regional and long-range transport of Hg originated 194 from anthropogenic emissions on the elevated GEM level at WLG will be discussed in details in 195 section 3.2 and 3.3 below.

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197 Currently, there is a great debate on the measurement accuracy of GOM using KCl-coated 198 denuder, and a small load of Hg could also cause analytical uncertainties in Tekran-based GOM and PBM measurements (Ambrose, 2017). Therefore, GOM data in this study was only compared with 199 200 previously reported data collected using the same method. The mean GOM concentration at WLG 201 was slightly higher than those in rural areas of North America and China, but lower than those in 202 urban areas in China (Fu et al., 2012b; Zhang et al., 2016). GOM is mainly affected by local to 203 regional emission sources and atmospheric processes (Sheu and Mason, 2001). Since WLG is 204 isolated from primary anthropogenic sources, the relatively high level of GOM at WLG was 205 probably mainly caused by atmospheric processes. Intrusion of GOM enriched air from free 206 troposphere could be one reason, a phenomenon that has been reported in Qomolangma Natural Nature Preserve in the southern Tibetan Plateau (Lin et al., 2019). Additionally, GOM at WLG 207 generally showed relatively higher concentrations during daytime (Fig. S4), indicating in situ 208 209 photochemical production of GOM as another important mechanism causing high GOM levels at 210 WLG.

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212 PBM concentrations at WLG showed large variations with the maximum hourly value reaching 655 pg m⁻³. The overall mean PBM concentration at WLG (68.1 pg m⁻³) was significantly higher 213 (by up to ~5-40 times) than those reported for remote areas in the northern Hemisphere (Kim et al., 214 2012; Lan et al., 2012), but was similar to observations in the urban areas in China (Fu et al., 2012b; 215 216 Fu et al., 2015). Elevated PBM concentrations in Chinese urban areas were most likely caused by 217 strong local anthropogenic emissions. However, in the remote areas with few primary anthropogenic 218 emissions, long-range transport should be the major cause for highly elevated PBM concentrations. 219 PBM has an atmospheric residence time ranging from a few days to weeks and can undergo regional 220 transport (Seigneur et al., 2004; Zhang et al., 2019). Therefore, high PBM levels at WLG were 221 probably mainly caused by long-range transport of anthropogenic and natural emissions, which was 222 discussed in details in the following Sections.

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224 3.2 Seasonal and diurnal distributions of GEM, GOM and PBM

225 Daily values of GEM, GOM, PBM, AT, RH, WS, RF, and air water content were aggregated 226 into monthly average values to reveal seasonal variations during the study period (Table S1 and Fig. 227 3). In the discussion below, warm (May to October) and cold (November to April) seasons were 228 compared. Mean GEM level in the cold season (1.84 ng m⁻³) was relatively lower than that in the 229 warm season (1.95 ng m^{-3}) (Table S1), which was likely due to the strengthening westerlies 230 originated from or passing over regions with low anthropogenic emissions during the cold season 231 (Zhang et al., 2015b). However, elevated monthly mean GEM levels were observed from February 232 to April (Table S1, Fig. S1, Fig. S2), and many high GEM events were frequently observed in the 233 cold season (Fig. S3), likely due to the long-range transport from northern South Asia where has 234 been experiencing industrialization and urbanization and thus strong anthropogenic Hg emissions 235 (AMAP/UNEP, 2018; Chakraborty et al., 2013). Airflows originated from these areas had high 236 GEM concentrations and could be transported to WLG in the cold season (Lin et al., 2019; Yin et 237 al., 2018). In addition, higher than seasonal-average GEM concentrations were also observed during 238 July to September (Table S1, Fig. 3), which could be attributed to the strengthening East Asia 239 Summer Monsoon during the warm season. The prevailing wind from the east direction (Fig. 4, 240 Table S1) could transport GEM from eastern Qinghai and southern Gansu province of China to 241 WLG during the East Asian Summer Monsoon season.

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Unlike GEM, mean and median GOM and PBM values were both higher in the cold season
than in the warm season (Table S1, Fig. 3, Fig. S1, Fig. S2). Also, the high GOM and PBM events

245 occurred mainly in the cold months (Fig. S3). Lower GOM and PBM concentrations in the warm 246 season were probably attributed to the increasing removal processes of these water soluble Hg 247 species, and this is consistent with previous observations with wet Hg deposition fluxes peaked in 248 the warm rainy season (Cole et al., 2014). In addition, low RH in the cold season would be 249 conducive to the formation of GOM and PBM through atmospheric chemical and physical 250 transformations (Fain et al., 2009; Lin et al., 2019). Higher PBM concentrations at WLG were 251 frequently detected with westerly and northerly winds (Fig. 4), which were mainly from the desert 252 and Gobi areas of western Tibetan Plateau, Xinjiang, southern Gansu and southwestern Inner 253 magnolia, suggesting that desert dust related sources in these regions could be potential sources of 254 PBM at WLG in the cold season.

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256 No significant differences in GEM concentration were observed between daytime (7:00-19:00, 257 1.89 ± 1.01 ng m⁻³) and nighttime (20:00-06:00, 1.86 ± 0.91 ng m⁻³) at WLG (Fig. S4). This was also the case for PBM (80.0 ± 81.5 pg m⁻³ versus 75.0 ± 72.5 pg m⁻³). The diurnal pattern of PBM at 258 259 WLG was different from those observed in Nam Co, Qomolangma Natural Nature Preserve and Mt. 260 Gongga in the Tibetan Plateau where PBM generally peaked during daytime under valley breeze 261 condition (Fu et al., 2009; Lin et al., 2019; Yin et al., 2018). The above findings at WLG suggested 262 that local sources and in situ atmospheric transformations may only have minor impacts on PBM 263 concentration. Instead, atmospheric circulation over the Tibetan plateau and long range transport 264 from the other source regions should be the main factors controlling the diurnal and seasonal 265 variations of GEM and PBM concentrations at WLG. In contrast, mean concentration of GOM 266 during daytime $(17.2 \pm 16.5 \text{ pg m}^{-3})$ was 31.3% higher than that during night $(13.1 \pm 10.3 \text{ pg m}^{-3})$ 267 at WLG. Given that there was a lack of strong anthropogenic emissions around the station or in the 268 surrounding areas, such a daytime elevated GOM phenomenon should be likely attributed to the in 269 situ production of GOM via GEM oxidation during the daytime (Ariya et al., 2015; Fain et al., 2009). 270 Therefore, local meteorology and photochemical production could be important controlling factors 271 for the observed diurnal patterns of GOM at WLG.

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273 **3.3 Source identification of GEM and PBM**

During the whole study period, the prevailing winds at WLG were from southwestern quadrant (46.5%) mainly originated from and passing over Tibetan Plateau and southern Xinjiang under the control of the westerlies. Average GEM concentrations (1.58 to 1.91 ng m⁻³) associated with this wind sector were overall lower than those associated with other wind sectors (Fig. 4), suggesting the areas southwest of WLG were not important source regions of GEM at WLG. In contrast, GEM concentrations (means: 2.42 to 2.87 ng m⁻³) associated with northeast wind sector were highly 280 elevated. The northeast wind mainly came from the low-altitude regions in northwestern China with 281 many anthropogenic Hg sources, which could have contributed to GEM at WLG. In contrast to GEM, maximum PBM concentrations (means: 68.6 to 97.8 pg m⁻³) were associated with wind 282 283 sectors of southwestern and northwestern quadrants and lowest PBM concentrations (means: 49.6 to 63.3 pg m⁻³) were associated with wind sector of the eastern quadrants (Fig. 4). The southwest 284 285 and northwest winds were mainly originated from and passed over deserts and Gobi regions, 286 including the largest Taklimakan Desert in Asia. These areas are the main dust source regions in 287 middle and eastern China (Che et al., 2011; Chen et al., 2017), and therefore would be an important 288 source of PBM at WLG.

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290 To better understand the sources and long-range transport of atmospheric Hg at WLG, CWT 291 values for GEM and PBM were calculated and are shown in Fig. 5. Higher GEM CWT values were 292 mainly located in eastern Qinghai, southern Gansu, western Shanxi, and southwestern Inner 293 Mongolia of China and northern South Asia, whereas lower values were mainly located western 294 Qinghai, Xinjiang and Xizang provinces (Fig. 5a). By matching the gridded GEM CWT values with 295 the gridded anthropogenic GEM emissions in the study domain, we found GEM CWT values were significantly positively correlated with anthropogenic GEM emissions ($R^2 = 0.55$, p < 0.01, Fig. 6a). 296 297 This indicates GEM at WLG was mainly caused by long-range transport of anthropogenic GEM 298 emissions from industrial areas in western China, and this is overall consistent with the findings 299 discussed above that were based on wind dependence of GEM at WLG.

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301 Differing from the case of GEM, higher PBM CWT values were mainly located in southern 302 Xinjiang, western Qinghai and south-central Xizang provinces, whereas the regions to the east of 303 WLG, where many industrial sources were located, showed relatively lower PBM CWT values (Fig. 304 5b). In addition, gridded PBM CWT values showed a negative correlation with gridded 305 anthropogenic PBM emissions (Fig. 6b). These findings indicate that long-range transport of 306 anthropogenic PBM emission was unlikely the major sources of PBM at WLG. Instead, long-range 307 transport of dust particles originated from deserts and Gobi regions in western China, such as 308 Taklimakan desert, Qaidam desert and Badain Jaran desert (Fig. S5), is responsible for PBM at 309 WLG. These regions contain the major deserts and Gobi areas in East Asia and can release up to 25 310 million tons dust particles annually. Dusts from these regions could be transported to the northwestern, middle and even eastern China through the westerlies over the Tibetan plateau (Che 311 et al., 2011; Chen et al., 2017; Xuan et al., 2000). Previous studies showed that atmospheric PBM 312 concentrations (86.1-517 pg m⁻³) over the Taklimakan Desert are remarkably higher than those 313 314 observed from background sites in China and even comparable to those measured in most of the 315 Chinese metropolitan cities (Huang et al., 2020). We thus concluded that the dry airflows transported

316 the PBM-enriched dust aerosols from the desert and Gobi regions to WLG, and contributed

317 significantly to the elevated PBM concentrations at WLG.

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319 3.4 Impact of desert dust related sources on PBM

320 To evaluate the impact of dust related sources on the temporal variations of PBM concentration, 321 daily cumulative AAI (Σ AAI) encountered by air masses transported during the preceding two days 322 were calculated, as shown in Fig. 7 together with daily PBM concentrations. PBM concentrations 323 maintained at relatively low levels in warm months (May to September), increased since October, 324 and reached the highest levels in winter and early spring (December to March) (Table S1). Daily 325 \sum AAI showed negative values from June to October, but large positive values in winter and early spring (Table S1 and Fig. 7). A significant positive correlation ($r^2 = 0.31$, p < 0.01) was observed 326 327 between daily SAAI and daily PBM concentration (Fig. 7), indicating that the long-range transport 328 of dust and/or biomass burning related sources played an important role in the temporal variations 329 of PBM concentration at WLG. Biomass burning related sources were not likely the major causes 330 because the air masses ended at WLG were mainly originated from and passed over regions with 331 low biomass burning area (Fig. S5). PBM concentrations showed a negative correlation with CO 332 and BC concentrations, which are mainly emitted from the industrial and biomass burning activities 333 (Table S2). Hence, we conclude that the dust related sources were the dominant source of PBM at 334 WLG. Previous studies analyzing spatiotemporal patterns of atmospheric dust based on satellite 335 remote sensing generated dust aerosol index have shown the Taklimakan area as the dominant 336 source of dust episodes in Asia, especially in every spring season. Desert dust is a significant carrier 337 of atmospheric aerosol and PBM to the cryosphere of Western China and can also have global 338 impact through long-range transport (Huang et al., 2020; Zhang et al., 2008).

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340 Desert and Gobi areas are important sources of atmospheric particles. Global dust particle emissions were estimated to range from 500 to 5000 Tg yr⁻¹ with an average value of 1836±903 Tg 341 yr⁻¹. In China, the desert and Gobi dust particle emissions were estimated to range from 100 to 459 342 Tg yr⁻¹ with an average value of 242±120 Tg yr⁻¹ (Table S3). Hg content in total suspended particles 343 (TSP) from desert dust was averaged at 0.33 μ g g⁻¹ from existing studies (Table S3). Based on the 344 345 above numbers, total particulate bound mercury (Hg-TSP) emissions from desert dust related 346 sources were roughly estimated to be 606 ± 298 (range from 165 to 1650) Mg yr⁻¹ globally and 80 \pm 40 (range from 33 to 151) Mg yr⁻¹ in China (Table S3). Given that PM_{2.5} generally accounts for 347 40% of TSP in dust aerosols, PBM (referred as to mercury bound to fine particles, e.g., of diameter 348 <2.5 µm) emissions from desert dust related sources were 242.4 Mg yr⁻¹ globally and 32 Mg yr⁻¹ in 349

350 China. These values exceed the anthropogenic PBM emissions in the world (75 Mg yr⁻¹) and China

351 (16 Mg yr⁻¹), suggesting desert and Gobi areas as important sources of atmospheric PBM emissions

- on regional to global scales.
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354 Besides emissions from anthropogenic and dust related sources, gas-particle partitioning 355 between GOM and PBM also affect PBM level in the atmosphere. Thus, the intrusion of GOM-rich 356 air from free troposphere would also have an impact on PBM (Ariya et al., 2015; Lin et al., 2019; 357 Tsamalis et al., 2014). The PBM/GEM ratios at WLG were similar to those observed at 358 Qomolangma Natural Nature Preserve and Nam Co. in the Tibetan Plateau, but much higher than 359 those in Chinese urban and remote areas (Lin et al., 2019; Yin et al., 2018). On the other hand, the 360 PBM/GOM ratios at WLG were relatively lower than the values observed from the other two 361 Tibetan sites (Fig. 8). Generally, gas-particle partitioning of GOM and PBM is mainly controlled by 362 air temperature (Amos et al., 2012), however, no clear dependence of monthly PBM/GOM ratio on 363 monthly mean air temperature was observed, e.g., similar PBM/GOM ratios were observed between 364 the coldest months (December to February) and other seasons (Fig. 8). This indicates the elevated 365 PBM in winter and early spring at WLG were not likely caused by the enhanced gas-particle 366 partitioning of GOM under low air temperature. Besides, air masses travelling heights at WLG did 367 not show clear seasonal variations throughout the study period, indicating elevated PBM 368 concentrations at WLG in winter and the early spring were unlikely associated with intrusions of 369 free troposphere air masses.

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371 4 Conclusions

372 This study presented the first full-year continuous speciated Hg data set and identified potential 373 sources causing high GEM and PBM at WLG in the Tibetan Plateau. Mean GEM level at WLG was 374 slightly higher than the background level of GEM in the Northern Hemisphere. Mean PBM level at 375 WLG was much higher compared with the reported values in remote areas in the Northern 376 Hemisphere. Seasonal variations in GEM concentration indicated that Hg emissions from 377 anthropogenic source regions and long-rang transport played important roles on the high GEM 378 levels at WLG. High PBM concentrations at WLG were observed in cold season, which were mainly 379 caused by dust aerosol sources from the desert and Gobi areas. Analysis from CWT and ΣAAI 380 northern Xinjiang, eastern Qinghai, southern Gansu, southwestern Shaanxi, western Inner Mongolia 381 of China and northern South Asia could be the main source areas of GEM, while southern Xinjiang, 382 southwestern Inner Mongolia, northern Gansu, western Oinghai and Tibet of China were likely the source regions of PBM at WLG. Long-range transport of dust particles from desert and Gobi areas 383 384 contribute to the elevated PBM at WLG. The estimated PBM emissions from dust particles

- 385 suggested that dust from desert and Gobi areas are critical sources of PBM on regional to global
- 386 scales, which should be paid more attention in future studies.

387 Data Availability:

388 All the dataset used in this study can be found in Supporting information.

389 Author contribution:

- 390 H.Z., X.W.F, and X.B.F. initiated the project and designated the field experiments. H.Z, X.W.F.,
- B.Y., B.X.L., P.L., and G.P.Z. carried out the field observations. H.Z., and X.W.F. prepared the
 manuscript with contributions from all co-authors.
- 393

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399 Competing interests:

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

401 **Supporting information:**

- Supporting Information Figure S1-S5
- Supporting Information Table S1-S3

404 **Reference**

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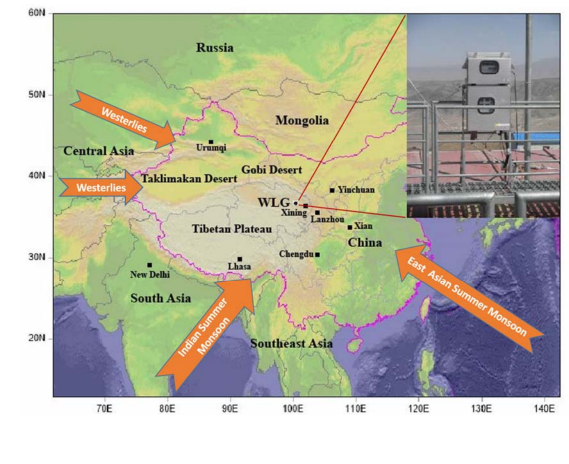
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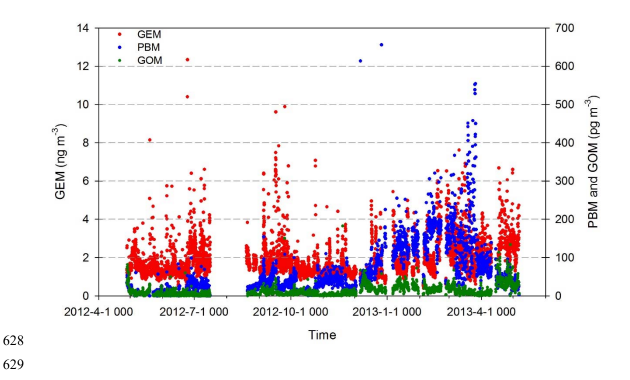
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- 592 Figure Captions
- Fig. 1: The map showing the location of WLG, and distributions of the deserts and cities aroundWLG.
- 595 Fig. 2: Time series of hourly GEM, GOM and PBM concentrations in ambient air at WLG.
- 596 Fig. 3: Monthly means of GEM, GOM and PBM at WLG during a full-year sampling period. Error
- 597 bars indicate the 1sd of monthly mean values.
- 598 Fig. 4: Frequency distribution of wind direction and wind-sector based mean GEM and PBM
- 599 concentrations during the study period.
- 600 Fig. 5: Identified source regions of atmospheric GEM and PBM at WLG during the study period.
- 601 (A) gridded $(0.5^{\circ} \times 0.5^{\circ})$ values for GEM, and (B) gridded $(0.5^{\circ} \times 0.5^{\circ})$ values for PBM. Gray line
- 602 enclosed regions indicate desert locations in China.
- 603 Fig. 6: Correlation between the simulated GEM or PBM CWT value and their respective
- anthropogenic emissions (AMAP/UNEP, 2018). The location of each gridded CWT value is
- 605 matched with that of anthropogenic emission. Gridded anthropogenic emissions are divided into 20
- 606 groups with equal number of grids, starting from the lowest to highest emission values.
- 607 Fig. 7: Variations in daily mean PBM concentration and daily cumulative Absorbing Aerosol Index
- 608 (AAI) during the proceeding two days at WLG.
- 609 Fig. 8: The monthly PBM/GEM and PBM/GOM ratio with air temperature and air mass traveling
- 610 height during a full-year sampling period. Column (a) is the mean PBM/GEM and PBM/GOM ratios
- 611 at Qomolangma Natural Nature Preserve and Nam Co in the inland Tibetan Plateau, column (b) is
- 612 the mean PBM/GEM and PBM/GOM ratios of Chinese cities, and column (c) is the mean
- 613 PBM/GEM and PBM/GOM ratios of Chinese remote areas.
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627 Fig. 2: Time series of hourly GEM, GOM and PBM concentrations in ambient air at WLG.

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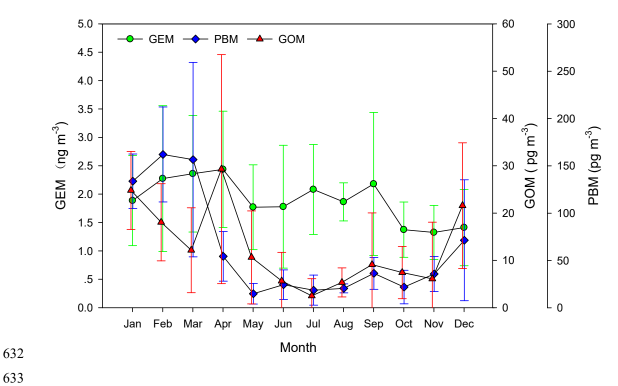
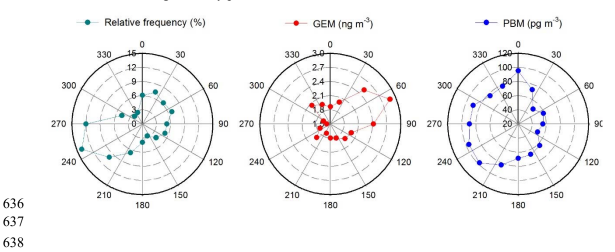
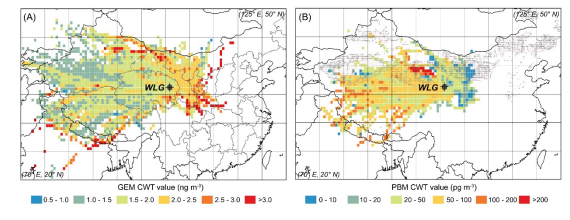


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- 639 Fig. 5: Identified source regions of atmospheric GEM and PBM at WLG during the study period.
- 640 (A) gridded $(0.5^{\circ} \times 0.5^{\circ})$ CWT values for GEM, and (B) gridded $(0.5^{\circ} \times 0.5^{\circ})$ CWT values for PBM.
- 641 Gray line enclosed regions indicate desert locations in China.





645 Fig. 6: Correlation between the simulated GEM or PBM CWT value and their respective 646 anthropogenic emissions (AMAP/UNEP, 2018). The location of each gridded CWT value is 647 matched with that of anthropogenic emission. Gridded anthropogenic emissions are divided into 20 648 groups with equal number of grids, starting from the lowest to highest emission values.

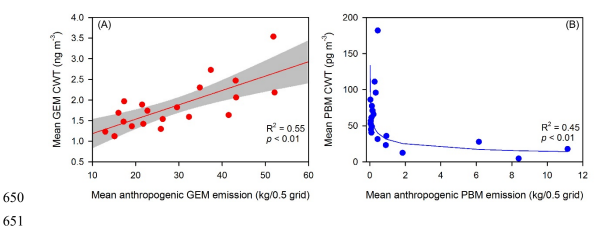


Fig. 7: Variations in daily mean PBM concentration and daily cumulative Absorbing Aerosol Index (AAI) during the preceding two days at WLG.

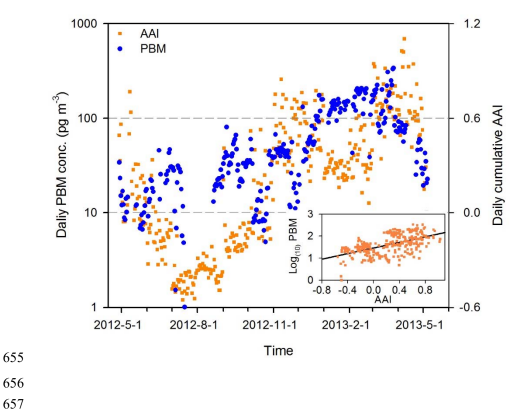


Fig. 8: The monthly PBM/GEM and PBM/GOM ratio with air temperature and air mass traveling
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at Qomolangma Natural Nature Preserve and Nam Co in the inland Tibetan Plateau, column (b) is
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