

Measurement report: Atmospheric mercury in a coastal city of Southeast China: inter-annual variations and influencing factors

Jiayan Shi^{1,2,4}, Yuping Chen^{1,2,3}, Lingling Xu^{1,2,*}, Youwei Hong^{1,2}, Mengren Li^{1,2}, Xiaolong Fan^{1,2}, Liqian Yin^{1,2}, Yanting Chen^{1,2}, Chen Yang^{1,2,3}, Gaojie Chen^{1,2,3}, Taotao Liu^{1,2,3}, Xiaoting Ji^{1,2,3}, Jinsheng Chen^{1,2,*}

¹ Center for Excellence in Regional Atmospheric Environment, Institute of Urban Environment, Chinese Academy of Sciences, Xiamen 361021, China

² Key Lab of Urban Environment and Health, Institute of Urban Environment, Chinese Academy of Sciences, Xiamen 361021, China

³ University of Chinese Academy Sciences, Beijing 100049, China

⁴ College of Resources and Environment, Fujian Agriculture and Forestry University, Fuzhou 350002, China

Correspondence: Lingling Xu (linglingxu@iue.ac.cn) and Jinsheng Chen (jschen@iue.ac.cn)

Abstract. Long-term monitoring of atmospheric mercury is an important part of the effective evaluation of the Minamata Convention on Mercury. Gaseous elemental mercury (GEM) along with conventional air pollutants and meteorological parameters were simultaneously observed in Xiamen city, Southeast China in January and July over the period 2012 – 2020. GEM concentrations in January highest in 2015 (4.47 ng m⁻³) and decreased to 2020 (3.93 ng m⁻³), while GEM concentrations in July were highest in 2017 (2.65 ng m⁻³) and lowest in 2020 (1.56 ng m⁻³). The temporal variation of GEM was typically characterized by higher concentrations in winter than in summer and in nighttime than in daytime. Bivariate polar plots and Concentrations-weighted trajectories (CWT) model were used to identify the source regions of GEM on a local and regional scale. The results indicate that the high GEM concentrations in January 2015 were likely due to a combination of high-level Hg emissions and adverse meteorological conditions. Generalized Additive Models (GAMs), a regression analysis method, were established and applied to investigate the influencing factors on the inter-annual variation of GEM. The factors, anthropogenic emissions, meteorological conditions and transportation explained $37.8 \pm 11.9\%$, $31.4 \pm 9.0\%$ and $30.8 \pm 9.9\%$ in average to the variation of GEM concentrations, respectively. There was a positive relationship of daily GEM concentrations with *T* and RH, mostly linking to natural surface emissions and Hg chemical transformations. The interpretation rate of anthropogenic emissions has significantly decreased since 2012, indicating the effectiveness of emission mitigation measures in reducing GEM concentrations in the study region.

Keywords: Inter-annual variation; Gaseous elemental mercury; GAMs; CWT analysis; Meteorology factors.

1 **1 Introduction**

2 Atmospheric mercury, because of its neurotoxicity, long persistence and high bioaccumulation, is
3 defined as a global pollutant that poses a threat to the health of the global population. Especially the
4 organic form of mercury, like methylmercury (MeHg), is associated with neurocognitive deficits in
5 human fetuses and cardiovascular effects in adults (Axelrad et al., 2007; Roman et al., 2011). In order to
6 protect human health and the environment from adverse effects of mercury, a legally-binding
7 international treaty Minamata Convention on Mercury was adopted in October 2013 and entered into
8 force in August 2017 (UN Environment, 2017). The Minamata Convention requires parties to reduce
9 mercury emissions and to assess the effectiveness of mitigation measures. China is one of the world's
10 largest mercury emitters as well as the signatory to the Minamata Convention. The annual atmospheric
11 mercury emissions in China were about 565 tons in 2015, accounting for about a quarter of global
12 anthropogenic mercury emissions (AMAP/UNEP, 2018). The predominant anthropogenic Hg emission
13 sources in China were industrial coal combustion, coal-fired power plants, nonferrous metal smelting,
14 and cement production (Zhang et al., 2015). The characteristics of Hg emissions such as large emission
15 amount and industrial production-dominated emissions made that China has great potential to reduce
16 mercury emissions through the implementation of the Minamata Convention on Mercury. A mercury
17 emissions inventory of China for 1978 – 2014 has reported that anthropogenic mercury emissions peaked
18 in 2011 and then showed a downward trend (Wu et al., 2016). In addition, China had reduced
19 anthropogenic mercury emissions by 127 tons from 2013 to 2017 (Liu K. et al., 2019). Among them,
20 mercury emissions from the coal-fired power plants fell from 105 tons in 2007 to 73 tons in 2015 (Zhang
21 et al., 2015; Liu et al., 2018). China has introduced a series of air cleaning measures including upgrading
22 precipitator devices, newly built desulfurization and denitrification devices and ultra-low emission
23 renovations since 2013, which led to synergistic removal of mercury (Liu K. et al., 2019). Strict
24 restrictions of mercury on the mining, production, utilization, import and export have also been imposed
25 in China since 2017 (<https://www.mee.gov.cn/>, last access: 17 May 2022).

26 Atmospheric mercury is usually classified into three categories according to the determination
27 technique: gaseous elemental mercury (GEM), gaseous oxidized mercury (GOM) and particulate bound

28 mercury (PBM). Because of its stability and volatility, GEM is the dominant form of atmospheric
29 mercury, accounting for up to 95%. GEM has an atmospheric residence time of 0.5 – 2 years and can
30 spread globally before being deposited to earth's surfaces (Schroeder and Munthe, 1998; Zhang et al.,
31 2013; Yuan et al., 2021). On the other hand, GOM and PBM have relatively high reactivity and dry/wet
32 deposition rates and are therefore easier to be removed from the atmosphere. Field observations are vital
33 to understand the long-term variation of Hg levels in the atmosphere. There are some global and regional
34 mercury monitoring networks around the world which provided a long-term monitoring result. For
35 example, a steep decline trend of GEM concentrations ($0.05 \text{ ng m}^{-3} \text{ yr}^{-1}$) was observed at Mace Head,
36 Ireland, from 2013 to 2018 (Custodio et al., 2020), while GEM data from Cape Point, South Africa,
37 showed a slight increase from 2007 to 2017 (Slemr et al., 2020). China has also carried out a series of
38 observational studies in the past two decades. These observation studies on GEM concentrations mostly
39 focused on 1 – 2 years, and few continuous GEM observation records over multiple years in China were
40 published (Fu et al., 2015). A three-year measurement at Chongming Island, East China, showed that the
41 annual GEM concentrations significantly decreased from 2.68 ng m^{-3} in 2014 to 1.60 ng m^{-3} in 2016, at
42 a rate of $-0.60 \pm 0.08 \text{ ng m}^{-3} \text{ yr}^{-1}$ (Tang et al., 2018). Whereas, a multi-year observation of GEM in
43 Guiyang, Southwest China, showed an increasing trend from 8.40 ng m^{-3} in 2002 to 10.2 ng m^{-3} in 2010,
44 and the increase mainly occurred during the cold season (Fu et al., 2015).

45 The variation of GEM concentrations is influenced by a variety of factors such as anthropogenic
46 emissions, meteorological conditions, as well as intra- and inter-regional transport (Tang et al., 2018; Liu
47 C. et al., 2019; Zhang et al., 2021). In previous studies, the impact of anthropogenic emissions changes
48 was often quantified by compiling emission inventories (Zhang et al., 2015; Wu et al., 2016; Liu K. et
49 al., 2019; Cai et al., 2020). The trajectory-based analysis method was frequently applied to analyze the
50 impact of regional transport (Tang et al., 2018; Wang et al., 2021). However, the results like the impacts
51 of anthropogenic emissions and regional transport derived from above different method systems could
52 not be comparable. Generalized additive models (GAMs), a regression analysis method, are data-driven
53 and able to incorporate non-linear relationships of air pollution with numerical and categorical variables,
54 which have been introduced into influencing factor identification in recent years (Wood and Augustin,
55 2002). The impact of local anthropogenic emissions, regional transport, and meteorological factors on

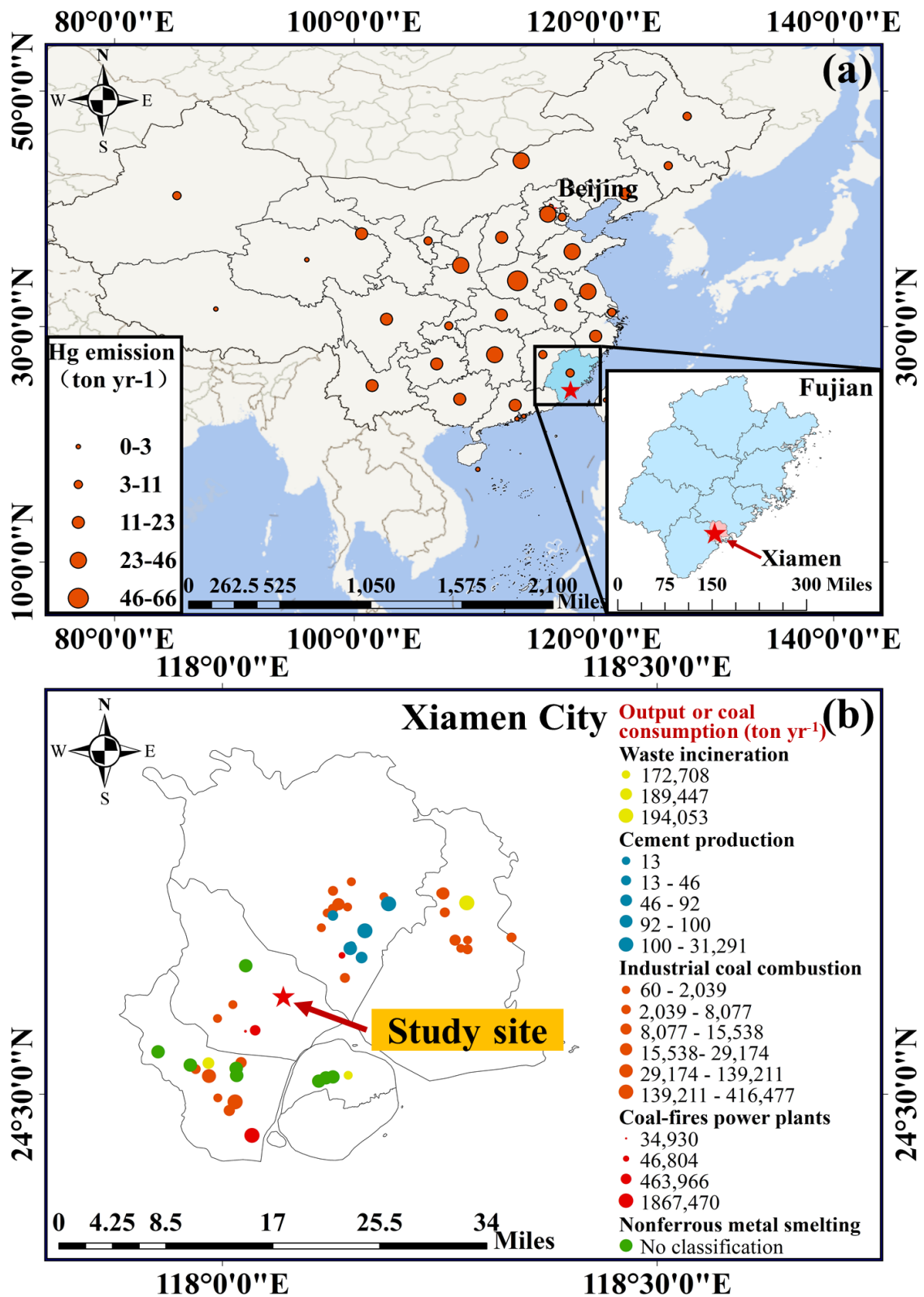
56 GEM concentrations in Nanjing, East China was quantified by using the GAMs (Zhang et al., 2021).
57 GAMs were also used to explain the decline of GEM concentrations in Beijing and the result showed
58 that reduction of anthropogenic mercury emissions, variation in meteorological conditions, and change
59 in globe background level explained 51.5%, 47.1% and 1.4% of the decrease in GEM concentrations,
60 respectively (Wu et al., 2020). It can be seen that GAMs are a promising tool to explore the effect of
61 factors like anthropogenic emissions and natural perturbations on GEM concentrations.

62 China has adopted aggressive atmospheric control measures in the last decade. Long-term GEM
63 observation was very necessary to investigate the variation of GEM levels and its influencing factors. In
64 this study, GEM, conventional pollutants and meteorological parameters were simultaneously observed
65 in Xiamen, a coastal city in Southeast China, in January (represents winter) and July (represents summer)
66 over the period 2012 – 2020. The main objectives of this study are: (1) to characterize the inter-annual,
67 seasonal and diurnal variations of GEM in Xiamen, a coastal city of Southeast China, (2) to identify the
68 source regions of GEM on a local and regional scale and their influence on annual concentrations of
69 GEM, (3) to investigate the influencing factors including anthropogenic emissions, regional transport
70 and meteorology on the inter-annual variation of GEM concentrations in the study region.

71 **2 Method**

72 **2.1 Site description**

73 The study site (Xiamen, 118°04'13''E, 24°36'52''N) is located in the Institute of Urban Environment,
74 Chinese Academy of Sciences in Jimei District of Xiamen City, Fujian Province, China (Fig. 1a). The
75 site was characterized by a typical subtropical monsoon climate, with the prevailing ocean monsoon in
76 summer and northerly or northeasterly air masses from the inland of China in winter. Industrial point
77 sources were mainly distributed to the northeast and the southwest of the study site (Fig. 1b). The
78 instruments were placed on the roof of a building (~80 m above the ground). The outdoor air inlet of the
79 sampling unit was located at 2 m above the rooftop of the building.



80
 81 Figure 1. (a) The location of study site in Xiamen City, Fujian Province, China and the regional distribution
 82 map of anthropogenic Hg emissions in China in 2014 (Wu et al., 2016). Note that the red dots represent the
 83 amount of mercury emitted by each province. (b) The distribution of local industrial point sources in Xiamen.
 84 Note that the colors of the dots represent different industrial categories, and the size represents a company's
 85 output or coal consumption (ton yr⁻¹).

86 2.2 Atmospheric mercury measurements

87 **Instrument operation:** Atmospheric mercury was measured by the Tekran 2537B/1130/1135 system
88 (Tekran Inc., Toronto, Canada). During the 1 h sampling period, the GOM and PBM were first collected
89 onto the KCl-coated quartz annular denuder and the quartz filter respectively, and GEM concentrations
90 were measured by a Tekran 2537B Hg vapor analyzer every 5 min at a sampling flow rate of 1 L min⁻¹.
91 The detection limit of GEM measurement was 0.06 ng m⁻³. In the following 1 h desorbing phase, the
92 PBM and GOM were sequentially desorbed and then quantified by the Tekran 2537B. The principle and
93 the routine maintenance of the equipment in detail have been described in the previous study (Xu et al.,
94 2015). The Tekran 2537B analyzer was calibrated automatically every 25h using the internal Hg
95 permeation source inside the instrument, and the accuracy of this permeation source was calibrated every
96 12 months with manual injection of Hg by a syringe from an external Hg source (Module 2505).

97 **Hg species selection:** GEM has a low chemical reactivity comparing to GOM and PBM in the
98 atmosphere. GEM concentrations are largely affected by factors like anthropogenic emissions and
99 atmospheric physical processes, which could be well represented by conventional pollutants like CO and
100 meteorological parameters like BLH. Whereas, GOM and PBM concentrations were strongly affected
101 by chemical transformation processes, which has no known suitable indicators. In addition, involving
102 chemical transformation effect as a main factor in the GAM models would make it more complicate.
103 Thus, we used the statistic method to explore the influencing factors on the more stable Hg species, GEM
104 in this study.

105 **Observation period selection:** This study was based on the GEM observation data of January and
106 July (representative of winter and summer, respectively) in 2012, 2013, 2015, 2017 and 2020. There was
107 a different time period of instrument malfunction among years. To ensure the study time periods were
108 consistent and the data could be comparable among years, we chosen the representative months of GEM
109 data in this study. Xiamen is located in the coastal region of Southeast China under the control of
110 subtropical oceanic monsoon, which has a significant distinction in meteorology between summer and
111 winter. According to the climate features in the study region, January and July well reflect the
112 characteristic of the winter and summer seasons, respectively. In addition, a whole year GEM
113 concentration observation in Xiamen supported that the GEM data in January can represent the GEM

114 characteristics of winter and July can represent summer (Xu et al., 2015). In order to match other
115 parameters, the time resolution of GEM concentrations was converted from the original 5 min to 2 h. For
116 most of the data (70%) during the study period, the proportion of GOM and PBM in TGM was less than
117 5%. Thus, GEM concentrations in this study were directly compared to TGM in the below analysis.

118 **2.3 Meteorological parameters and criteria air pollutants**

119 In this study, the conventional pollutants (including SO₂, NO₂, O₃, CO, PM₁₀, PM_{2.5}) and
120 meteorological parameters (including wind speed (WS), wind direction (WD), relative humidity (RH),
121 air temperature (*T*) and surface air pressure (SP)) were obtained from Xiamen air quality monitoring
122 station. Note that above pollutants concentrations and meteorological data were averaged into 2 h time
123 intervals. Other meteorological parameters: boundary layer height (BLH), downward UV radiation at the
124 surface (UVB) and low cloud cover (LCC) were obtained from the European Centre for Medium-Range
125 Weather Forecasts (ECMWF) reanalysis (<https://www.ecmwf.int>, last access: last access: 23 March
126 2022).

127 **2.4 Concentration weighted trajectory (CWT)**

128 Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model and Global Data
129 Assimilation System data (<https://www.arl.noaa.gov/>, last access:5 May 2022) were applied to calculate
130 the 72h backward air mass trajectories. The interval of backward trajectories was 2 h and the arrival
131 height was set as 500 m above the ground level.

132 Concentrations-weighted trajectories (CWTs) analysis is useful to identify pollution sources spatially
133 for long-term environment measurement (Cheng et al., 2013). The principal of the CWT method is that
134 each grid cell is assigned a weighted concentration by averaging the sample concentrations associated to
135 trajectories that cross that grid cell as follows:

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

136 Where C_{ij} is the average weighted concentration in the ij th cell, l is the index of the trajectory, M is
137 the total number of trajectories, C_l is GEM concentration observed on arrival of trajectory l , τ_{ijl} is the
138 time spent in the ij th cell by trajectory.

139 Based on the maximum distance travelled by the 72h back trajectories over the study period, a
140 geographical domain from 17° to 60 N and from 45° to 135° E (from 4° to 37° N and from 100° to 153°
141 E) was found to be suitable in January (July). This domain encompassed the locations of nearly all the
142 trajectory segment endpoints from all the back trajectories. The resolution for CWT was $1^\circ \times 1^\circ$ in this
143 study.

144 **2.5 Model development**

145 Generalized Additive Models, a regression analysis method, have been used to establish the
146 relationship between GEM and various variables, and to investigate the influencing factors on the inter-
147 annual trend of GEM concentrations (Gong et al., 2017). The models were run by the following steps:
148 parameter selection, model establishment, and model quality control.

149 **Parameter Selection:** 16 variables we obtained from site observation and web downloads generally
150 fell in four categories: anthropogenic emissions (SO_2 , NO_2 , O_3 , CO , $\text{PM}_{2.5}$, PM_{10}), surface meteorology
151 (T , RH , WS , WD , SP), high-altitude meteorology (BLH , UVB and LCC) and air transportation (24h-
152 Latitude and 24h-Longitude) (Table S1). All the variables were standardized by min-maximum method.
153 The normalized data eliminates the effects of differences in dimension and ranges of values between
154 indicators. The standardized variables were then screened using two methods: Statistical judgment and
155 the meaning of the variables. Firstly, the collinearity diagnosis method was adopted to make judgment
156 according to the variance inflation factor (VIF). PM_{10} , SO_2 , NO_2 , SP and UVB were rejected into the
157 model due to their high collinearity ($\text{VIF} > 5$). Secondly, we considered the meaning of the remaining
158 parameters. Parameters O_3 and $\text{PM}_{2.5}$ were easily rejected into the model. CO is mainly sourced from
159 anthropogenic emissions and has a long atmospheric residence time (compared to SO_2 and NO_2). In
160 addition, Hg emissions in Fujian provinces were dominated by combustion sources (Liu K. et al., 2019).
161 Hence, we used CO to represent anthropogenic Hg emissions. After determining the first parameter CO ,
162 we put the remaining parameters (WS , WD , T , RH , BLH , LCC , 24h-Latitude and 24h-Longitude) into
163 the model one by one. The performance of GAMs was judged according to Akaike Information Criterion
164 (AIC) and R^2 values. Specifically, as the parameters were successively added into the model, the AIC
165 decreased and R^2 increased. In this step, WS , WD and LCC were rejected. Based on this method, 6

166 variables including CO, RH, T , BLH, 24h-Latitude and 24h-Longitude were eventually selected into the
167 model (Table S2).

168 Considering that parameters of the same category might interact, we used interaction functions of
169 tensors. RH, T and BLH interaction was used to represent the meteorological factor. 24h-Latitude and
170 24h-Longitude interaction, which indicates the backward trajectory endpoint location during last 24 h
171 was used to represent the transportation factor. Given that the 6 selected variables passed the collinearity
172 test, the three factors anthropogenic Hg emissions represented by CO, meteorological factor represented
173 by RH, T , and BLH interaction, and transportation represented by 24h-Latitude and 24h-Longitude
174 interaction were considered to be independent of each other. Those selected variables were used in the
175 following model establishment.

176 **Model establishment:** GAMs were performed using R version 4.1.2 with the “mgcv” package. The
177 equation can be described as follows:

$$g(\mu) = f_1(x_1) + f_2(x_2) + \dots + f_k(x_k) + \varepsilon, \quad (3)$$

178 Where x_j ($j = 1, 2, 3, \dots, k$) are different meteorological predictors and f_j is a smooth function of the
179 predictors; ε is the residual; μ is the expected value of the response variable; and g is the link function
180 specifies the relationship between the non-linear formulation and the expected value. We used the
181 "identity link" function with a Gaussian distribution because the relationship between GEM and the three
182 factors conformed to a Gaussian distribution and the estimation of GAMs was considered unbiased. In
183 order to ensure the balance between under-fitting and overfitting of observation data, we used a penalized
184 cubic regression as a smooth function. In the running process of the model, the concentration contribution
185 of the smoothed independent variable (anthropogenic emissions, meteorological effects, and
186 transportation) to the dependent variable (GEM) was output and converted into contribution ratio, which
187 is helpful to determine the degree of each variable driving the prediction. To eliminate the effect of
188 seasonality on variance clustering, we used “seasons” as an input variable when building the model with
189 the whole dataset, and then run the model separately in summer and winter.

190 **Model Quality Control:** The accuracy of GAMs simulation was assessed using a 10-fold cross-
191 validation test. The principle of the test is dividing the whole dataset into ten subsets randomly, and in
192 each round of cross-validation, nine subsets are used to fit the model and the remaining one is predicted.

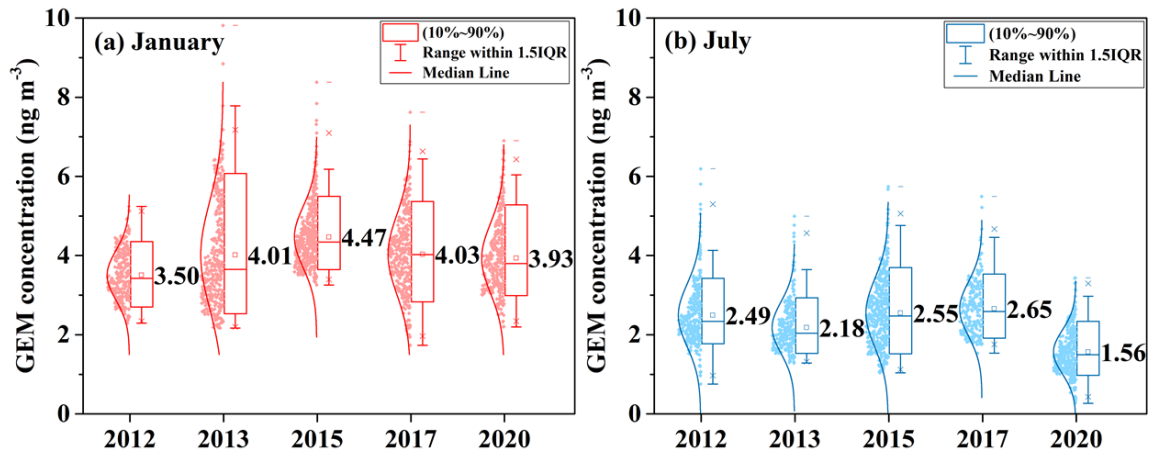
193 This process was repeated 10 times to ensure that every subset was tested. The 10-fold cross-validation
194 results showed a good coincidence between the GAMs and cross-validated results ($R^2 = 0.89$, Fig. S1),
195 demonstrating the reliability of the established model. In order to test the underlying assumptions of
196 homogeneity, normality and independence of GAMs to ensure the validity and accuracy of the model,
197 we used the following methods (Fig. S2): (1) Quantile-quantile (QQ) plots (Sample quantiles against
198 theoretical quantiles), (2) scatterplots (residuals against linear predictor), (3) histograms of the residuals
199 and (4) scatter plots (response against fitted value). The QQ-plot showed that the GAMs produced good
200 results around the average concentrations and the residuals showed a normal distribution.

201 **3 Results and discussion**

202 **3.1 Temporal variations of GEM concentrations**

203 **3.1.1 Inter-annual variation**

204 Monthly concentrations of GEM ranged from 3.50 to 4.47 ng m⁻³ in January and 1.56 to 2.65 ng m⁻³
205 in July during the whole study period, with mean values of 4.04 ± 1.01 ng m⁻³ and 2.29 ± 0.83 ng m⁻³,
206 respectively. The GEM concentrations in Xiamen were several times higher than the Hemisphere
207 background concentrations (~ 1.5 ng m⁻³) (Sprovieri et al., 2016; Diéguez et al., 2019). Comparisons of
208 TGM/GEM concentrations in Xiamen with other urban and rural areas in East Asia over the last decade
209 are shown in Table 1. The mean concentrations of GEM mostly fell in a range of 2 – 5 ng m⁻³ in East
210 Asia (except for some background sites). The winter GEM concentrations in Xiamen was slightly higher
211 than those measured at rural and background monitoring sites such as Changdao Island, Tibetan Plateau
212 region, and Mt. Ailaoshan (Zhang et al., 2016a; Yin et al., 2018; Wang et al., 2020), while lower than
213 those reported from inland urban sites like Lanzhou, Nanjing, and Shanghai (Zhu et al., 2012; Duan et
214 al., 2017; Yin et al., 2020). The summer GEM concentrations in Xiamen were relatively low, comparable
215 to those of Changdao Island, Yongheung Island and Fukuoka (Marumoto et al., 2015; Lee et al., 2016;
216 Wang et al., 2020).



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218

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Figure 2. Statistical summaries of gaseous elemental mercury (GEM) concentrations in Xiamen in (a) January and (b) July of the study years.

Table 1. Comparisons of TGM/GEM concentrations in Xiamen with other urban and rural areas in East Asia over the period 2010 – 2020.

Locations	Classification	Time	TGM/GEM (ng m ⁻³)	Winter	Summer	References
Xiamen, CN	Suburban (Coast)	03/2012–02/2013	3.50±1.21	/	/	(Xu et al., 2015)
Mt. Changbai, CN	Rural	07/2013–07/2014	1.68±0.47	/	/	(Liu C. et al., 2019)
Haidian, CN	Urban	02/2018–03/2018	2.77±0.91	/	/	(Wang et al., 2021)
Shanghai, CN	Suburban	06/2014–12/2014	4.19±9.13	5.5±6.6	3.8–5.3	(Duan et al., 2017)
Shanghai, CN	Suburban	12/2015–02/2016	2.77±1.36	2.88	2.87	(Qin et al., 2019)
Hefei, CN	Suburban	07/2013–06/2014	4.07±1.91	4.05 ± 1.81	4.08±1.99	(Hong Q. et al., 2016)
Ningbo, CN	Coastal	04/2011–04/2013	3.30±1.40	3.70	2.60	(Yu et al., 2015)
Ningbo, CN	Urban (Coastal)	07/2013–01/2014	3.26±1.63	/	/	(Hong Y. et al., 2016)
Ningbo, CN	Urban (Coastal)	12/2016–11/2017	2.44±0.95	2.62±1.05	2.26±0.78	(Yi et al., 2020)
Nanjing, CN	Urban	01/2011–12/2011	7.94±6.99	5.5±2.5	9.9±8.2	(Zhu et al., 2012)
Mt. Ailaoshan, CN	Rural	05/2011–05/2012	2.09±0.63	2.04±0.58	2.20±0.60	(Zhang H. et al., 2016)
Taoyuan city, Taiwan, CN	Suburban	10/2017–09/2018	2.61±6.47	2.19±0.75	2.27±1.67	(Sheu et al., 2019)
Taichung city, Taiwan, CN	Rural	10/2014–09/2015	1.19	/	/	(Fang et al., 2017)
Yongheung, Korea	Island	01/2013–08/2014	2.80±1.10	3.50–3.70	2.30±0.90	(Lee et al., 2016)
Gyodong Island, Korea	Rural	08/2015–09/2017	2.70±2.60	2.8±2.9	1.70±1.0	(Lee et al., 2019)
Fukuoka, Japan	Urban	06/2012–05/2013	2.33±0.49	2.31±0.44	2.37±0.58	(Marumoto et al., 2015)
Nam Co, Tibetan Plateau, CN	Plateau	01/2012–10/2014	1.33±0.24	1.14±0.18	1.50±0.20	(Yin et al., 2018)
Changdao Island, CN	Rural (Coastal)	10/2013–07/2015	2.52±0.82	2.87±1.16	2.25±0.51	(Wang et al., 2020)
Lanzhou, CN	Urban	10/2016–10/2017	4.48±2.32	5.06±2.45	4.45±2.10	(Yin et al., 2020)

221 The inter-annual variability of GEM concentrations in January and July are shown in Fig. 2. The GEM
222 concentrations in Xiamen showed a minor variation between years. Specifically, GEM concentrations in
223 January were lowest in 2012 (3.50 ng m⁻³), highest in 2015 (4.47 ng m⁻³), and comparable in the years of
224 2013, 2017 and 2020. Whereas GEM concentrations in July was highest in 2017 (2.65 ng m⁻³) and
225 significant lowest in 2020 (1.56 ng m⁻³). According to the published data, Wu et al. (2016) estimated
226 atmospheric Hg emissions in China decreasing from 547 tons in 2010 to 530 tons in 2014. The report
227 from AMAP/UNEP showed that the anthropogenic Hg emissions in China were 565.2 t in 2015 relative
228 to 575.2 t in 2010 (AMAP/UNEP, 2018). An inventory over the period 1978-2017 revealed that China's
229 anthropogenic Hg emission was highest in 2013 and then decreased until 2017 (Liu K. et al., 2019). It
230 could be expected that the anthropogenic Hg emissions in China had a downward trend over the period
231 2012-2020 and the peak emission was most likely to occur in 2012 to 2014 (Fig. S3,4). Recent studies
232 have indicated either a stable or a slight decreasing trend for GEM or TGM concentrations in Chinese
233 cities after 2013 when China has applied the vigorous measures to control air pollution (Qin et al., 2020;
234 Wu et al., 2020; Yin et al., 2020). For instance, it was reported that GEM concentrations at Chongming
235 Island in East China significantly decreased from 2014 to 2016, and the inflection point occurred before
236 2014 (Tang et al., 2018). Note, those measurements mostly lasted for 2 – 4 years. So far, the observations
237 of GEM concentrations over a long time period were scarce. Our result suggests the influencing factors
238 on the variation of GEM in East China would be complex over the last decade.

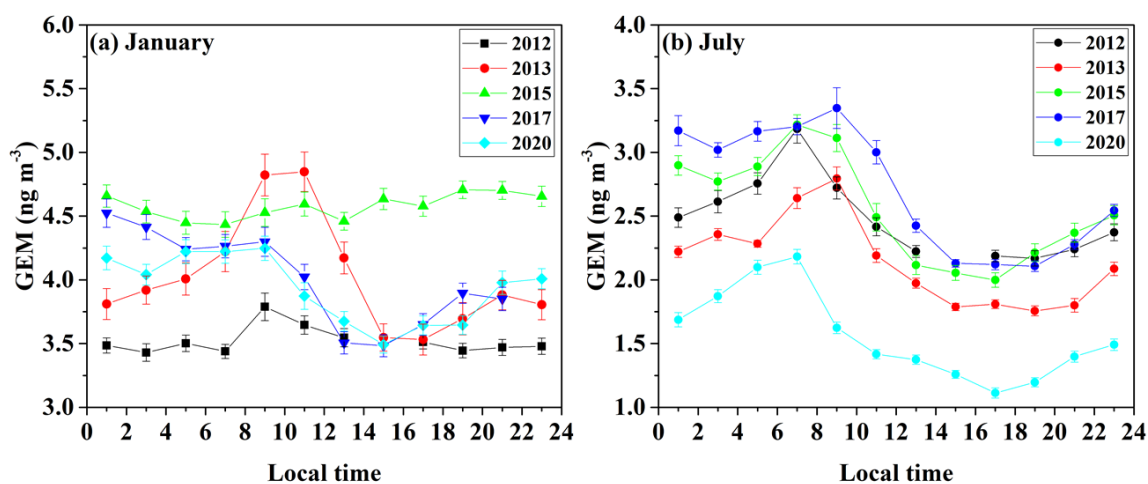
239 Coal combustion is one of the leading mercury sources in China (Wu et al., 2006). Table S3 summaries
240 the consumption of coal and the statistics of annual SO₂ and NO_x emissions in Fujian Province over the
241 period 2012 – 2020. The coal consumption in Fujian and China exhibited a similar variation, firstly
242 decreasing to a valley in 2016 and then showing an upward trend from 2016 to 2020 (Fig. S5). Whereas,
243 there was a decreasing trend for SO₂ and NO_x emissions during the whole study period, with a most
244 remarkable reduction of emissions in 2016. In 2013, the Chinese State council issued an air pollution
245 prevention and control action plan. Since then, plenty of emissions control measures, like accelerate the
246 elimination of backward production capacity, accelerate the promotion of central heating, upgrade and
247 building air pollution control devices, have been widely implemented in China. Previous studies have
248 found that mercury can be synergistically removed in the process of desulfurization and de-nitration

249 (Zhang Y. et al., 2016; Liu K. et al., 2019). The installation of selective catalytic reduction to control
250 nitrogen oxide emissions is often accompanied by the oxidation of GEM to GOM, and the combined
251 application of selective catalytic reduction and flue gas desulfurization could further reduce TGM
252 emissions to the atmosphere (Rallo et al., 2012). Nonetheless, the inconsistent inter-annual variation in
253 GEM concentrations and SO₂/NO_x emissions indicates that additional factors, like GEM emission
254 sources other than coal combustion and/or meteorological changes strongly affected the inter-annual
255 variation trend of GEM in the study region.

256 **3.1.2 Seasonal and diurnal patterns**

257 The GEM concentrations in Xiamen were approximately 1.41 – 2.52 times higher in winter than in
258 summer ($P < 0.001$, one-way ANOVA) over the study years. The similar seasonal variation was widely
259 observed in cities including Shanghai, Ningbo, Lanzhou, and Yongheung, as well as Gyodong Island,
260 and Changdao Island (Yu et al., 2015; Hong Y. et al., 2016; Lee et al., 2016; Duan et al., 2017; Lee et
261 al., 2019; Wang et al., 2020; Yi et al., 2020; Yin et al., 2020). However, a reverse seasonal variation with
262 higher GEM in summer than in winter was observed in Nanjing, Chongming Island, Mt. Ailaoshan and
263 Tibetan Plateau region (Zhu et al., 2012; Zhang H. et al., 2016; Tang et al., 2018; Yin et al., 2018). There
264 were many factors responsible for the seasonal variation of GEM in Xiamen. In terms of Hg emission
265 sources, local industrial emissions were relatively stable over the course of a year. The key factor with
266 seasonal changes was the increased usage of coal for heating which mainly occurred in northern China
267 in cold seasons. Although there was no coal consumption for heating in southern China, GEM is well
268 mixed due to its prolonged lifetime (0.5–2 years) (Schroeder and Munthe, 1998; Zhang et al., 2013).
269 Monsoons change the source–receptor relationships at observation sites, and thus affect the seasonal
270 variation of GEM concentrations (Fu et al., 2015; Liu C. et al., 2019). For Xiamen, air masses in winter
271 mainly originated from north directions which passed through numerous intensive anthropogenic GEM
272 emissions areas (Fig. 1a), while air mass in summer mainly originated from ocean with less GEM point
273 sources. Another important factor is that the mixing heights were reduced due to stable inversion layer
274 in winter. As a result, GEM diffused slowly and accumulated easily in the surface layer. In addition, for
275 the sites in the Northern Hemisphere, the greater removal of GEM by wet and dry deposition could also
276 lead to lower GEM concentrations in warmer seasons (Fu et al., 2008; Tang et al., 2018).

277 The diurnal variations of bihourly GEM concentrations were consistent among years (Fig. 3). In
 278 general, the GEM concentration peaked in the early morning, decreased to a valley in the afternoon, and
 279 then rose during the night. The diurnal pattern of GEM concentrations in January 2015 were gentle than
 280 other years of the same period, which might be related to the enhanced effect of air mass transport (Fu et
 281 al., 2012; Nguyen et al., 2022). The diurnal pattern of GEM concentrations in Xiamen is consistent with
 282 other urban sites like Guiyang, Hefei and Guangzhou (Feng et al., 2004; Chen et al., 2013; Fu et al.,
 283 2015). Previous studies often attributed diurnal variations of GEM to the effect of various anthropogenic
 284 emissions, photochemical oxidation and the diurnal variation of BLH (Hong Y. et al., 2016; Duan et al.,
 285 2017). The diurnal pattern of GEM in Xiamen was similar to those of SO₂, NO₂ and CO (Fig. S6a – c),
 286 reflecting the combined effects of common anthropogenic emissions and the diel fluctuation in
 287 meteorology. The decreasing trend of GEM from early morning to afternoon was likely related to the
 288 intensified turbulent mixing in the boundary after sunrise while the nighttime had the opposite condition
 289 (Fig. S6d). The photo-oxidation of GEM might contribute a small part to the diurnal variation of GEM,
 290 but it was not the dominant factor for the daytime GEM reduction due to its low reaction rate in the study
 291 region.



292
 293 **Figure 3. The diurnal trend of GEM concentrations in (a) January and (b) July over the study years. Note**
 294 **that the value of error bars has been reduced tenfold.**

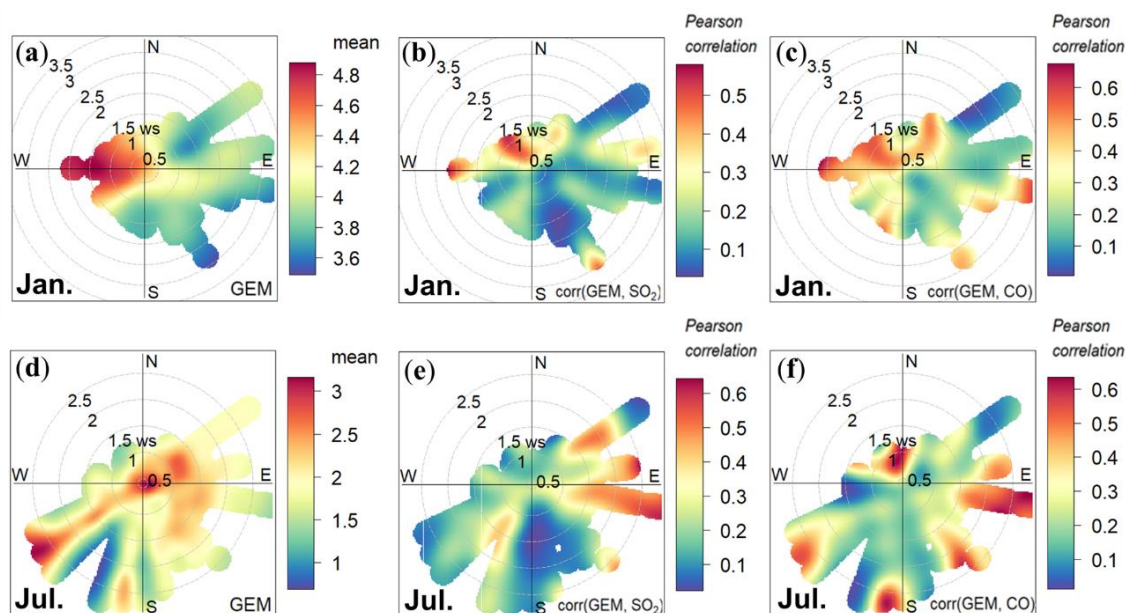
295 **3.2 Potential source regions of GEM**

296 **3.2.1 Local emissions**

297 According to the industrial Hg emissions inventory in China, the main Hg emissions industries
298 included industrial coal combustion, coal-fired power plants, non-ferrous metal smelting, cement
299 production, and waste incineration (Zhang et al., 2015; Wu et al., 2016). The relationships between the
300 spatial distribution of industrial sources and bivariate polar plots of GEM concentrations would shed
301 light on the influence of local anthropogenic point sources (within Xiamen city) on GEM concentrations.
302 As shown in Fig.4a, 4d, the elevated GEM concentrations in the bivariate polar plots of individual
303 seasons were highly concentrated, suggesting that the spatial distribution of main Hg point sources was
304 similar among years. However, the polar plot results of GEM concentrations were distinctly different
305 between seasons likely due to the shift in the dominant wind. In January, the elevated GEM
306 concentrations were associated with west wind with a low WS of $0.5 - 2.0 \text{ m s}^{-1}$ (Fig. 4a), which indicates
307 nearby emission sources. They were likely the industrial coal combustion, and non-ferrous metal
308 smelting upwind from the west of the monitoring site (Fig. 1b). The conventional pollutants are the good
309 indicators of primarily anthropogenic sources. In Xiamen city, the coal-fired power plants contributed
310 57.8% of local industrial SO_2 emissions and 62.4% of CO industrial emissions (Air pollution emissions
311 inventory of Xiamen city in 2016, unpublished report). A close correlation of GEM with SO_2 and CO in
312 westerly WD with low speeds (Fig. 4b, 4c) further supported above conclusion that the contributions of
313 local industrial sources to GEM.

314 In July, the elevated GEM concentrations occurred when winds came from the southwest with the WS
315 about $2.5 - 3 \text{ m s}^{-1}$ (Fig. 4d). As shown in Fig. 1b, there are many industrial clusters in the southwest
316 direction of the observation site including coal-fired power plants, industrial coal combustion and
317 nonferrous metal smelting. Accordingly, we suspected that the local industrial clusters upwind of the
318 southwest to the study site caused an evident increase in GEM concentrations. High WS of the southwest
319 wind likely weakened the correlation between GEM and SO_2 (Fig. 4e) while GEM and CO remained a
320 good correlation in the southwest wind due to their stable chemical properties (Fig. 4f). Another elevated
321 GEM concentration condition occurred when wind came from the east with a lower WS of $0 - 2 \text{ m s}^{-1}$
322 (Fig. 4d). Such low WS suggests a stagnant meteorological condition which was unfavorable for GEM

323 dispersion. In addition, GEM and SO₂ showed a good correlation in the case of the east wind with WS
 324 of 1 – 2 m s⁻¹. Hence, we could speculate that the upwind point sources, like industrial coal combustion
 325 and nonferrous metal smelting, as well as the adverse atmospheric diffusion conditions contributed to the
 326 increasing GEM concentrations. As shown in Table S4, GEM in Xiamen was overall positively correlated
 327 with SO₂, NO₂, CO and PM_{2.5}, but the correlation coefficient fluctuated remarkably among years. In
 328 addition, the inter-annual variation of GEM concentrations was not coincided with those of SO₂ or NO_x
 329 emissions as mentioned above. Thus, the contribution of local Hg point sources to atmospheric GEM
 330 varied among year and it could not fully explain the inter-annual variation of GEM concentrations in the
 331 study region.



332
 333 **Figure 4. Bivariate polar plots of GEM concentrations (a, d), the correlation coefficients of GEM with SO₂ (b,**
 334 **e) and CO (c, f) as a function of WS and WD in January and July during the whole study period. Note that**
 335 **GEM in ng m⁻³, wind speed in m s⁻¹, wind direction in °.**

336 3.2.2 Long distance migration

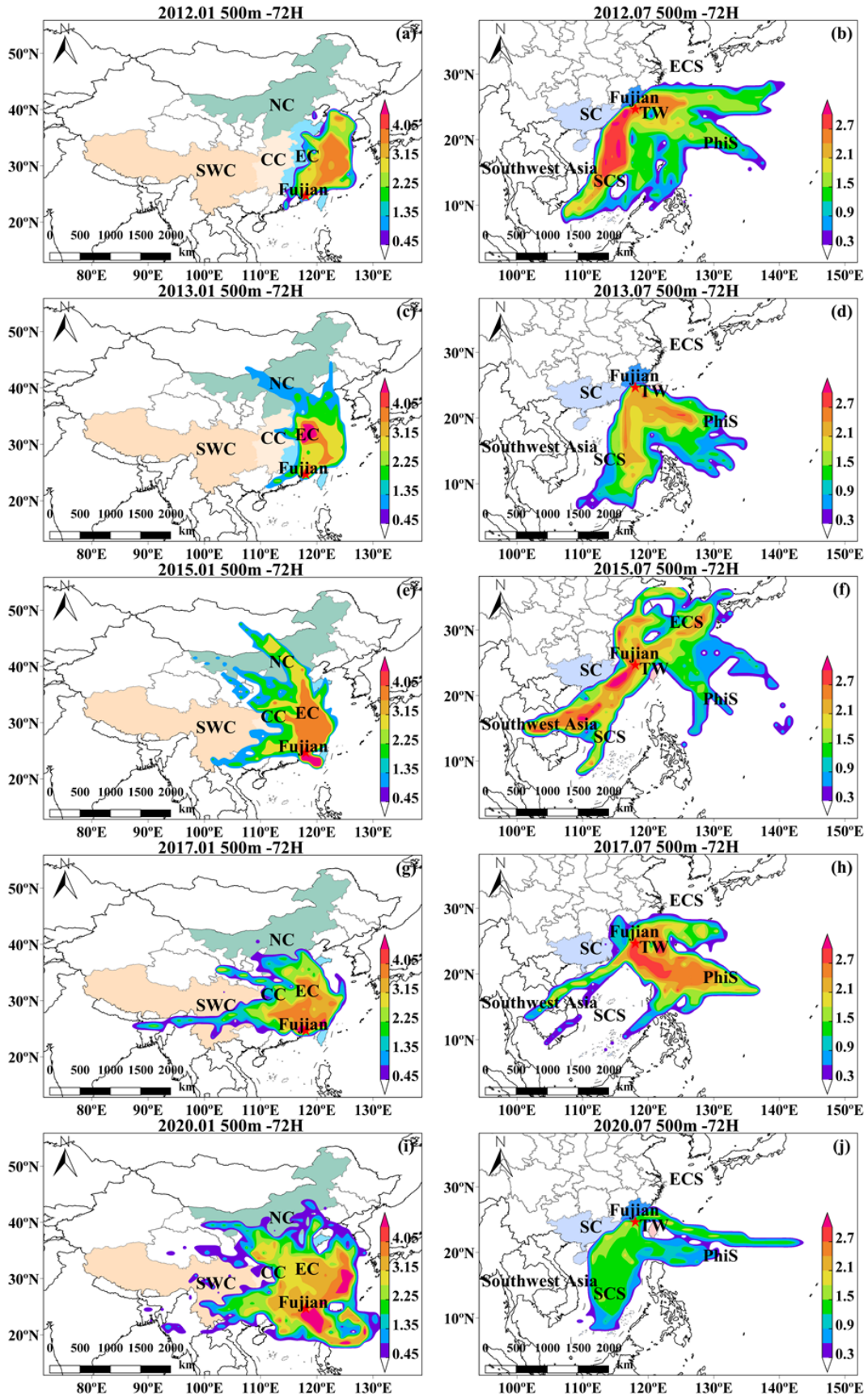
337 The result of CWT analysis based on HYSPLIT backward air mass trajectories are shown in Fig. 5.
 338 On the whole, the GEM concentrations in Xiamen were under an influence of continental air masses in
 339 January and a mixing influence of continental and marine air masses in July. To describe it better, the
 340 potential source regions of GEM were marked as Fujian province, East China (EC), Southwest China
 341 (SWC), North China (NC), Central China (CC), South China Sea (SCS), Philippines Sea and Taiwan

342 Strait (PhiS+TW), as well as Philippines Sea and East China Sea (PhiS+ECS) (The details of region
343 definition are presented in Supporting Information).

344 It can be seen that the higher weights of CWT mostly dominated in Fujian province and EC in January.
345 The results could reasonably be expected because anthropogenic Hg emissions in East China was
346 extremely large due to the dense industries (Zhang et al., 2015). The potential source regions for GEM
347 in Xiamen also included ECS in January 2012, CC in January 2017, and PhiS+TW in January 2020. The
348 North China Plain region was one of the heaviest mercury polluted area in China (Zhang et al., 2015),
349 but to our surprise, it did not become an important source region for Xiamen GEM. This is most likely
350 due to the effect of transmission distances, while GEM concentrations decreased with long transport
351 distances. In contrast, short air mass trajectories, like from the Fujian province, indicated a relatively
352 stagnant air condition which is not conducive to the diffusion of GEM (Zhang et al., 2021). The weights
353 of CWT in January 2015 were higher than those of other study years and the region of high CWT values
354 spread over Fujian province and EC. Previous studies have also reported the high levels of GEM in 2015
355 among urban Beijing, Changdao Island, rural Shanghai and Chongming Island (Tang et al., 2018; Qin et
356 al., 2020; Wang et al., 2020; Wu et al., 2020), which suggests a heavy GEM pollution on a large regional
357 scale during 2015. The high GEM concentrations in January 2015 was likely due to a combination of
358 high-level Hg emissions and adverse meteorological conditions. The annual atmospheric mercury
359 emissions in China were about 565 tons in 2015, which was roughly 20% higher than those in 2010
360 (AMAP/UNEP, 2018). According to anthropogenic mercury emissions inventory in China during 1978
361 – 2017, mercury emissions might peak around 2013, and remained high in 2015 (Figure S2). In addition,
362 an adverse effect of meteorological conditions due to extreme 2015 – 2016 El Niño event might also
363 result in an increase of GEM concentrations in 2015 (Nguyen et al., 2022).

364 The dominant clean marine air masses helped explain the lower concentration of GEM in July than in
365 January. As shown in Fig. 5, the dominant source region for GEM in Xiamen in July mostly were SCS
366 and PhiS+TW. The evasion of GEM from the ocean to the atmosphere was reported to be an important
367 natural source (accounting for 50%) (Mason, 2008). When under the control of marine air masses, the
368 ocean emissions would become a significant source for GEM in the study region. The CWT values were
369 extremely high in SCS in July 2012 and in PhiS+TW in July 2017, while the CWT values were elevated

370 in SCS and ECS in July 2015. The relatively high GEM concentrations for Xiamen in July 2017 was
371 likely associated with the contribution of extremely high GEM from PhiS+TW. The elevated GEM
372 concentrations for Xiamen in July 2015 passed more closely through Southeast Asia where the intense
373 biomass burning often occurred (Friedli et al., 2009; Sheu et al., 2013; Liu et al., 2016). A previous study
374 on Hainan Island also pointed to the possibility of long-range transport of GEM from Southeast Asia to
375 South China (Liu et al., 2016).



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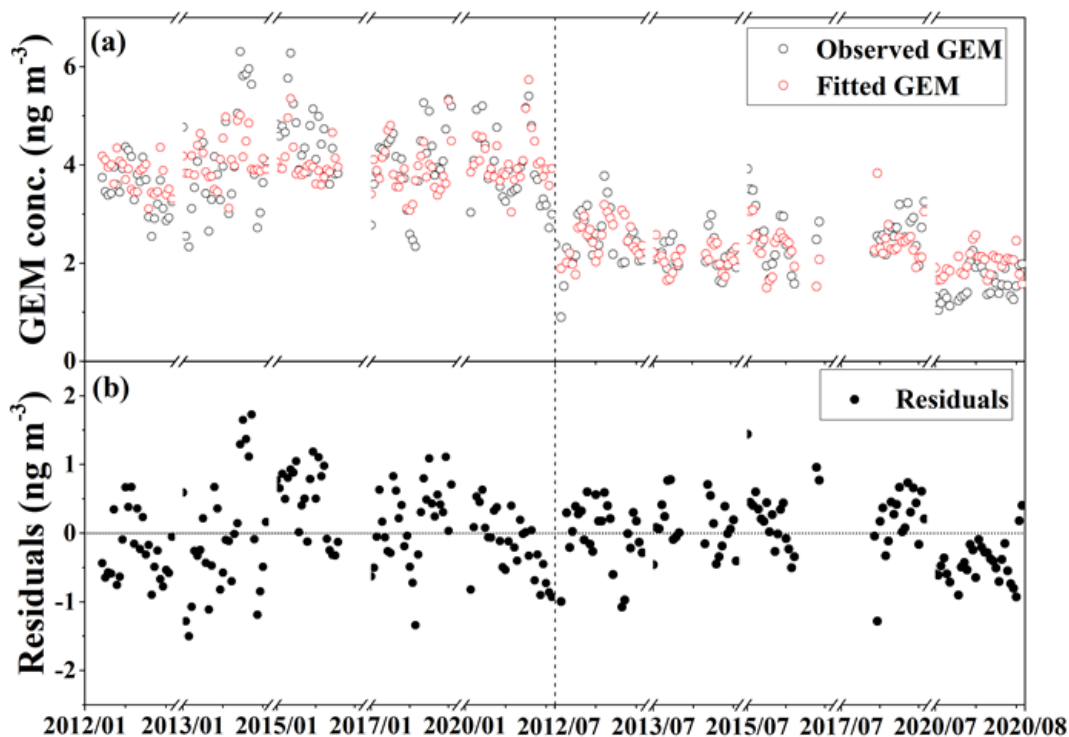
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Figure 5. Concentration weighted trajectory (CWT) in January (a, c, e, g, i) and July (b, d, f, h, j) during the study years. The legend represents the weighted GEM concentrations (ng m^{-3}).

379 **3.3 Factors affecting GEM concentrations**

380 **3.3.1 Model evaluation**

381 GAMS were applied to investigate the influencing factors on the inter-annual variation of GEM in this
382 study. The fitted (observed) GEM concentrations derived from the GAMS were approximately $3.98 \pm$
383 0.47 ng m^{-3} ($4.00 \pm 0.84 \text{ ng m}^{-3}$) in January and $2.25 \pm 0.41 \text{ ng m}^{-3}$ ($2.23 \pm 0.61 \text{ ng m}^{-3}$) in July, reflecting
384 that the model approximates the concentration of GEM to the mean. The observed and fitted GEM
385 concentrations showed a good consistency in time series (Fig. 6a) and the residuals were normally
386 distributed (Fig. 6b). The R^2 of the observed and fitted GEM concentrations was 0.75 (Fig. S7) and the
387 variance interpretation rate was 75.8%. In previous studies using GAMS to quantify impact factors on air
388 pollutants, the R^2 was generally between 0.35 and 0.86 (Gong et al., 2018; Li et al., 2019; Wu et al., 2020;
389 Wu et al., 2021; Zhang et al., 2021). The fitted result of the GAMS in our study falls in this range.

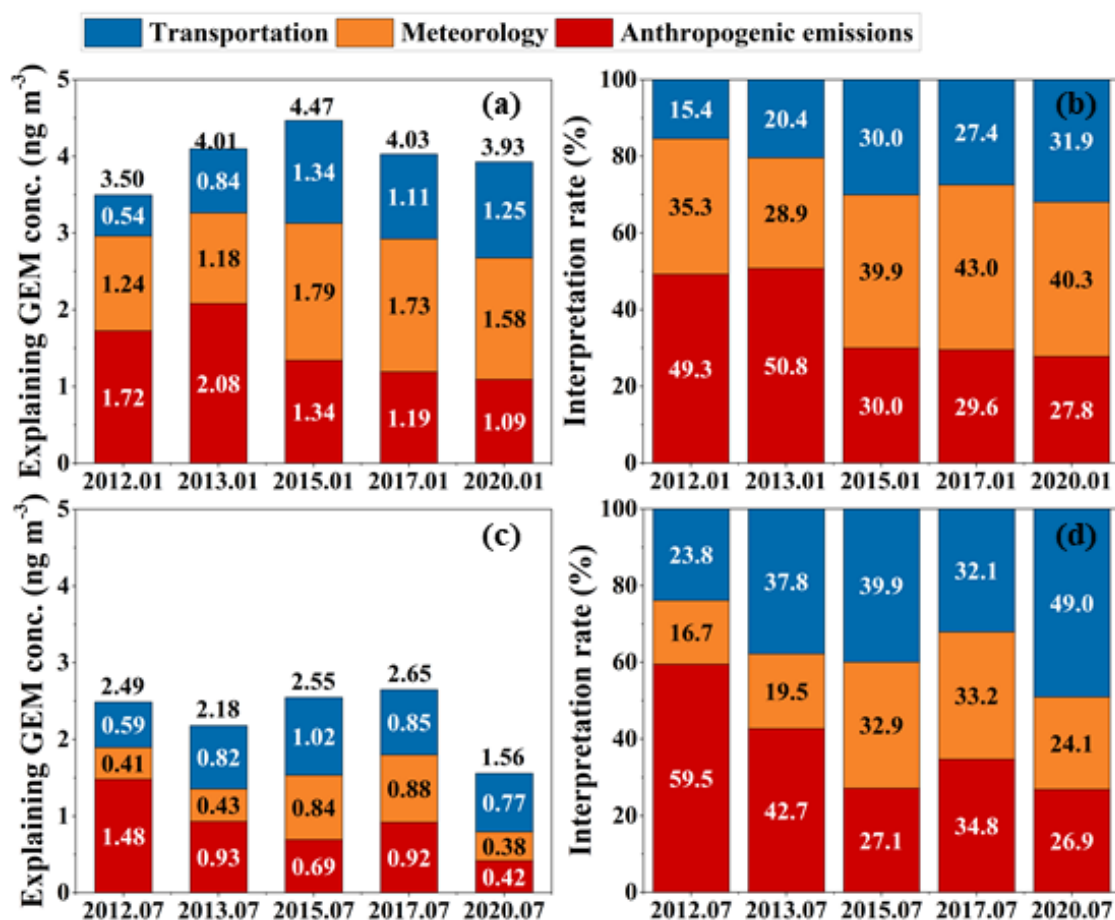


390
391 **Figure 6. Time series of (a) observed and fitted GEM concentrations in January and July and (b) residual**
392 **distribution during the whole study period.**

393 3.3.2 Inter-annual variations of factor contributions

394 The 6 screened variables represented the factors of anthropogenic emissions (CO), meteorology (T,
395 RH, and BLH interaction) and transportation (24h-Latitude and 24h-Longitude interaction). The
396 contributions of the three factors to the variation of GEM concentrations in January and July of the study
397 years are shown in Fig. 7. The inter-annual interpretation rate of anthropogenic emissions varied from
398 27.8% to 50.8% in January and from 26.9% to 59.5% in July, with a mean value of $37.8 \pm 11.9\%$ during
399 the whole study period. The interpretation rate of the emission factor to the variation of GEM
400 concentrations was pronouncedly high in 2012 and decreased overall to 2020. China has adopted a series
401 of desulphurization and denitrification measures for air pollution control during the period 2010 – 2015.
402 The capacity of desulphurization and denitrification units in China had reached 99% and 92% of the total
403 installed capacity of coal power plants, and 1.6×10^8 kw had been upgraded to achieve ultra-low emissions
404 by 2015. In addition, strict restrictions of mercury on the mining, production, use, import and export were
405 also be imposed in China since 2017 (<http://www.mee.gov.cn>, last access: 17 May 2022). We observed
406 a positive relationship between GEM and the representative variable CO in most of the CO observed
407 range (Fig. 8a, b). That is, GEM concentrations basically decreased with the reduction of CO which
408 stands for anthropogenic emissions. The most significant reduction in GEM concentrations explained by
409 anthropogenic emissions was 0.55 ng m^{-3} from July 2012 to July 2013 and 0.74 ng m^{-3} from January
410 2013 to January 2015. The results reveal that the vigorous air pollution control measures in China were
411 very effective reducing GEM concentrations in the study region.

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Figure 7. The explaining concentration (ng m⁻³) (a, c) and interpretation rate (%) (b, d) of the three factors to the variation of GEM concentrations during the study years.

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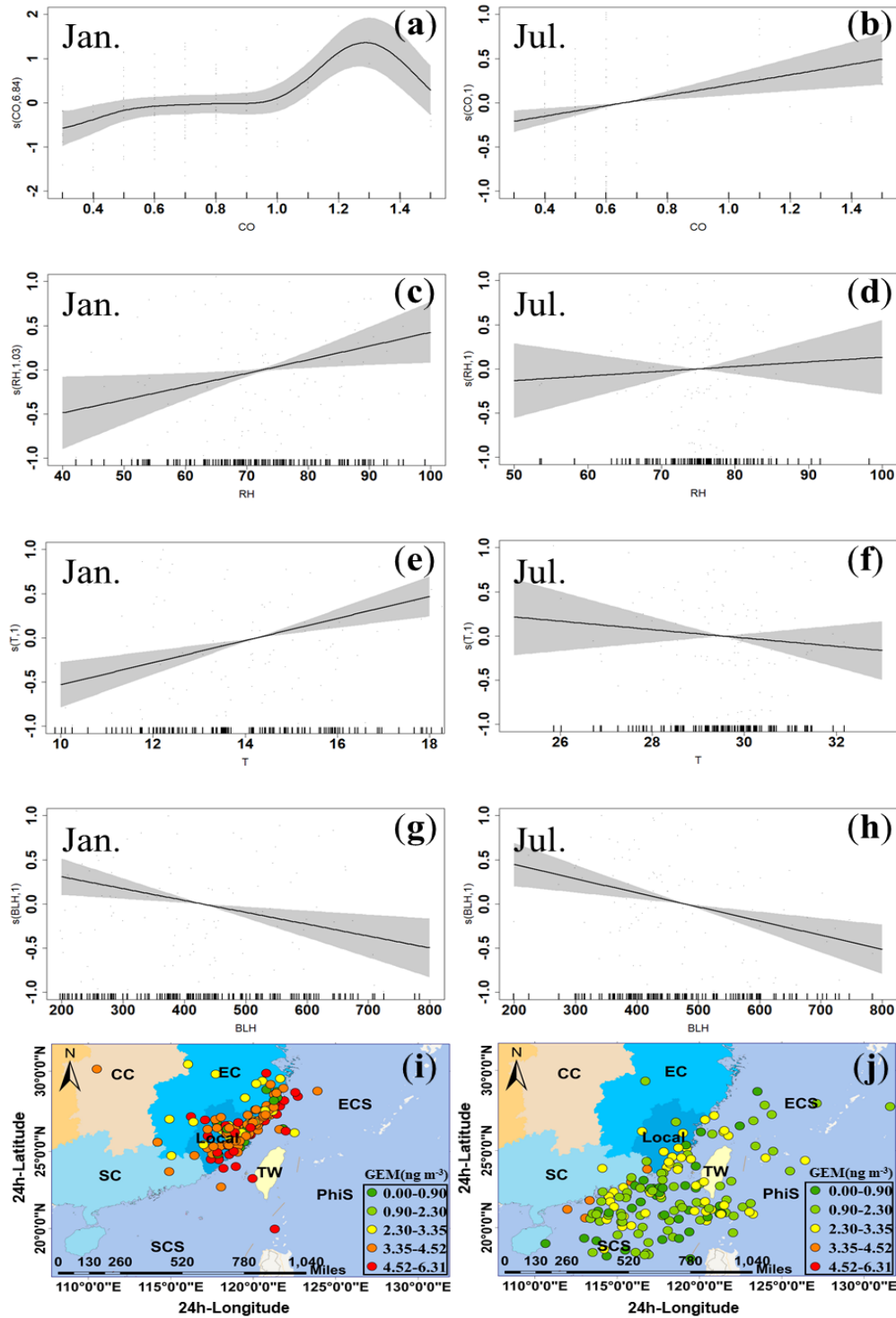
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Meteorology was another important factor which explained, on average, $31.4 \pm 9.0\%$ to the variation of GEM concentrations during the whole study period. The inter-annual interpretation rate of meteorology largely varied from 28.9% to 43.0% in January and from 16.7% to 33.2% in July. The largest part of GEM concentrations explained by meteorological factor occurred in January 2015 and July 2017, which were 1.79 ng m⁻³ and 0.88 ng m⁻³, respectively. The meteorological factor was represented by RH, *T* and BLH interaction, which influenced GEM variations by several pathways. One is that the meteorological factor drives the Hg chemical transformation (Ariya et al., 2015). In this study, GEM concentrations linearly increased with the increasing RH (Fig.8c). The raise in RH might enhance the liquid phase reduction of process that convert reactive Hg to GEM (Horowitz et al., 2017; Saiz-Lopez et al., 2018; Huang et al., 2019). Another important role is that the meteorological factor is highly related to the Hg emissions from natural surfaces. According to Fig. 8e, f, GEM concentrations increased with the raise of temperature in January, possibly because the increase of temperature would promote the Hg

428 emissions from natural surface (Horowitz et al., 2017; Qin et al., 2019). The meteorological factor also
429 affects the physical state of the atmosphere. A higher BLH likely indicated good diffusion conditions
430 and was more conducive to the reduction of GEM concentrations (Fig. 8g, h). A case in January 18 - 28,
431 2013 shown that the daily GEM concentrations increased with the increase of T and RH as well as the
432 decrease of BLH (Fig. S8). The case also implies that increased temperature likely accelerating natural
433 surface emissions and high RH being favorable for the liquid phase reduction of reactive Hg to GEM
434 contributed to GEM in the atmosphere.



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Figure 8. Spline of GEM to the chosen variables, (a, b) CO (mg m⁻³) in January and July, (c, d) RH (%) in January and July, (e, f) Temperature (°C) in January and July, (g, h) BLH (m) in January and July, (i, j) 24h-Longitude and 24h-Latitude in January and July. The grey background around the line is 95% confidence bounds for the response. The short lines on x axes show the distribution of data points. The number in the bracket of ordinate title is the estimated degree of freedom. The colored dots in figure (i, j) represents the interactive influence of 24h backward trajectories latitude and longitude coordinates.

442 Regional transportation explained a comparable proportion of GEM concentrations with the
443 meteorological factor, which made up $30.8 \pm 9.9\%$ of total variation during the whole study period.
444 During the period of 2012 – 2020, the interpretation rate of transportation displayed an increasing
445 variation from 15.4% to 31.9% in January and a slightly large variation between 23.8% and 49.0% in
446 July. The endpoint of 24 h backward trajectories and corresponding GEM concentrations are shown in
447 Fig. 8i, j. The high GEM concentrations mostly presented in the areas with 24h-Longitude of $116^\circ -$
448 124°E and 24h-Latitude of $24^\circ - 30^\circ\text{N}$. This is consistent with the backward trajectory results in section
449 3.3.2 that the main source regions of GEM were Fujian province and East China. The interpretation rate
450 of transportation increased significantly among years, indicating the growing importance of
451 transportation from high Hg emission zone to low emission zone in the background of national
452 anthropogenic emissions reduction.

453 **4 Conclusions**

454 Long-term observation of GEM concentrations along with conventional pollutants and meteorological
455 parameters were conducted in Xiamen city, Southeast China. GEM concentrations showed no distinct
456 variation trends over the period 2012 – 2020. GEM concentrations in January were highest in 2015 (4.47
457 ng m^{-3}) and decreased to 2020 (3.93 ng m^{-3}), while GEM concentrations in July were highest in 2017
458 (2.65 ng m^{-3}) and lowest in 2020 (1.56 ng m^{-3}). The temporal variation of GEM was characterized by
459 higher values in winter than in summer and in nighttime than in daytime, which is consistent with those
460 in most urban cities.

461 Upwind point sources within Xiamen city contributed to the elevated GEM concentrations in the study
462 site. Nevertheless, the inter-annual variation trend of GEM was not consistent with those of local SO_2
463 and NO_x emissions, suggesting that anthropogenic emissions might be not the dominant influencing
464 factor for GEM variation. The CWT results showed that the pronounced high GEM concentrations in
465 winter 2015 was a regional phenomenon, which was likely due to a combination of high-level Hg
466 emissions and adverse meteorological conditions.

467 The three factors, anthropogenic emissions, meteorological conditions and transportation, which were
468 represented by CO, (RH, T and BLH) interaction, and (24h-Longitude, 24h-Latitude) interaction,

469 explained $37.8 \pm 11.9\%$, $31.4 \pm 9.0\%$ and $30.8 \pm 9.9\%$ to the variation of GEM concentrations during the
470 whole study period, respectively. There was a positive relationship of GEM with T and RH, suggesting
471 that the meteorological factor influenced GEM variations by several pathways, possibly by affecting the
472 emissions from natural surface and the liquid phase reduction of reactive Hg to GEM. The interpretation
473 rate of transportation and meteorology on GEM variations displayed an increasing trend. In contrast,
474 anthropogenic emissions showed a decreasing interpretation rate since 2012, indicating the effectiveness
475 of emission mitigation measures in reducing GEM concentrations in the study region.

476 **Data availability.** A dataset for this paper can be accessed at <https://doi.org/10.5281/zenodo.6573605>
477 (Shi et al., 2022). High-altitude meteorology parameters can be acquired from the European Centre for
478 Medium-Range Weather Forecasts (ECMWF) reanalysis (<https://www.ecmwf.int>, last access: last access:
479 23 March 2022), Gridded meteorological data are available from the Global Data Assimilation System
480 (<https://www.arl.noaa.gov/>, last access:5 May 2022), The details are also available upon request from the
481 corresponding author.

482

483 **Author contributions.** JS and LX designed this study and analysis the data. YuC, LX, YH, ML, XF,
484 YaC, CY GC, LT, JX and JC were involved in the scientific discussion and offered valuable suggestions
485 for modifications. JS and LX wrote the manuscript. YuC and JC helped revise the manuscript. and LY
486 managed finances. All authors reviewed the paper.

487

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

489

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492

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498

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