

1 **Atmospheric Speciated Mercury Concentrations on an Island between China and Korea:**  
2 **Sources and Transport Pathways**

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27 **Abstract**

28 As a global pollutant, mercury (Hg) is of particular concern in East Asia where anthropogenic  
29 emissions are the largest. In this study, speciated Hg concentrations were measured on Yongheung  
30 Island, the western most island in Korea, located between China and the Korean mainland to identify  
31 the importance of local and regional Hg sources. Various tools including correlations with other  
32 pollutants, conditional probability function, and back-trajectory based analysis consistently indicated  
33 that Korean sources were important for gaseous oxidized mercury (GOM) whereas, for total gaseous  
34 mercury (TGM) and particulate bound mercury (PBM), regional transport were also important. A  
35 trajectory cluster based approach considering both Hg concentration and the fraction of time each  
36 cluster was impacting the site was developed to quantify the effect of Korean sources and out-of-  
37 Korean source. This analysis suggests that contributions from out-of-Korean sources were similar to  
38 Korean sources for TGM whereas Korean sources contributed slightly more to the concentration  
39 variations of GOM and PBM compared to out-of-Korean sources. The ratio of GOM/PBM decreased  
40 when the site was impacted by regional transport, suggesting that this ratio may be a useful tool for  
41 identifying the relative significance of local sources vs. regional transport. The secondary formation  
42 of PBM through gas-particle partitioning with GOM was found to be important at low temperatures  
43 and high relative humidity.

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45 **Keywords** : *TGM, GOM, PBM, regional transport, gas-particle partitioning, cluster analysis*

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47 **1. Introduction**

48

49 Mercury (Hg) is the only metal that exists as a liquid at standard conditions (US EPA, 1997) which  
50 results in it having a significant vapor pressure and presence in the atmosphere. In the atmosphere, Hg  
51 generally does not constitute a direct public health risk at the level of exposure usually found (Driscoll  
52 et al, 2007). However, once Hg is deposited into aquatic systems, it can be transformed into methyl-  
53 mercury (MeHg) which is very toxic and readily bioaccumulates through the food web (Mason et al.,

54 1995). Many studies show that one of the major sources of MeHg in aquatic and terrestrial system is  
55 atmospheric deposition of inorganic Hg (Landis and Keeler, 2002, Mason et al., 1997). Fish  
56 consumption has been considered to be the major exposure pathway of Hg for humans (Mergler et al.,  
57 2007; UNEP, 2013). In Korea, You et al. (2012) showed that MeHg concentrations in blood were  
58 affected by fish consumption as well as by gender difference. However, rice consumption was also  
59 found to be the predominant pathway of MeHg exposure for the inhabitants residing in a highly  
60 contaminated area of China (Zhang et al., 2010).

61 Atmospheric mercury exists in three major inorganic forms, including gaseous elemental mercury  
62 (GEM,  $\text{Hg}^0$ ), gaseous oxidized mercury (GOM,  $\text{Hg}^{2+}$ ) and particulate bound mercury (PBM,  $\text{Hg}(p)$ ).  
63 The sum of the GEM and GOM is often called as total gaseous mercury (TGM). Due to its high water  
64 solubility and deposition velocity, GOM has short atmospheric residence times (~day) and,  
65 consequently, its ambient concentration is mainly affected by local sources. Besides the anthropogenic  
66 sources, the free troposphere has been identified as an important GOM source (Huang and Gustin,  
67 2012; Weiss-Penzias et al., 2009; Timonen et al., 2013). In contrast, GEM, which comprises more  
68 than 95 percent of the total Hg in ambient air, can be transported long distances because it is relatively  
69 inert and has low water solubility and deposition velocity (Lin and Pehkonen, 1999). The residence  
70 time of PBM is dependent on the size of associated particles, but generally, it has been assumed to be  
71 a few days (Fang et al., 2012, Zhang et al., 2001). Measurements of GOM and PBM are challenging  
72 and uncertain due to their extremely low concentrations and complex chemical reactivity, and because  
73 their chemical forms are not actually known (Pirrone et al., 2013). In most studies, GOM and PBM  
74 have operational definitions for the mercury species collected by a KCl coated denuder and by a  
75 quartz filter downstream of a KCl denuder, respectively. It is typically assumed that GOM comprises  
76  $\text{HgCl}_2$ ,  $\text{HgBr}_2$ ,  $\text{HgO}$ ,  $\text{Hg}(\text{NO}_3)_2$ , and  $\text{HgSO}_4$ . However, the sampling method including the use of a  
77 KCl denuder has been shown to be subject to interferences from ozone, water vapor and possibly  
78 other compounds (Lyman et al. 2010; Talbot et al., 2011; Jaffe et al., 2014; Finley et al., 2013; Gustin  
79 et al., 2013; Huang et al., 2013; McClure et al., 2014), and it also should be noted that the different  
80  $\text{Hg}(\text{II})$  compounds have different collection efficiencies by the KCl coated denuder (Gustin et al.,

81 2015).

82 In the atmosphere, Hg species can be interconverted through various redox reactions. It is known  
83 that GOM can be produced by homogeneous and heterogeneous reactions of GEM with O<sub>3</sub>, OH, and  
84 Br/BrO (Hedgecock and Pirrone, 2004; Obrist et al., 2011; Subir et al., 2011), but there is no  
85 consensus on which oxidants are most important under which environmental conditions. GEM can  
86 also be formed through reduction of GOM predominantly in cloud water (Subir et al. 2011, 2012).  
87 GOM can also be converted to PBM through gas-particle partitioning, with the partition coefficient,  
88 K<sub>p</sub>, inversely correlated with temperature and positively correlated with particle surface area (Lyman  
89 and Keeler, 2005, Rutter and Schauer, 2007a,b; Liu et al., 2010). Since GEM makes up the bulk of the  
90 total Hg in ambient air its formation through reduction processes of divalent Hg may not be important.  
91 However, the secondary formations of GOM through the oxidation of Hg<sup>0</sup> followed by gas-particle  
92 partitioning formation of PBM can contribute significantly to their ambient concentrations.

93 The region of largest anthropogenic Hg emissions is East and Southeast Asia, contributing 39.7%  
94 of the total global anthropogenic emissions (UNEP, 2013). In Korea, atmospheric Hg emissions have  
95 generally decreased since 1990 (Kim et al., 2010). However, Hg levels in Korea are likely to be  
96 highly susceptible to Chinese emissions because China alone accounts for about one third of the  
97 global total (UNEP, 2013) and Korea is situated just east (and generally downwind) of China.  
98 According to the recent studies, Hg concentrations in blood of Koreans are more than 4~8 times  
99 higher than those found in US and Germany, and approximately 26% of Koreans have higher blood  
100 mercury concentrations than a USA guideline level (<http://envhealth.nier.go.kr>), indicating that there  
101 is an urgent need to identify the Hg sources and pathways controlling Hg concentrations in Korea.

102 This study was designed to identify the contribution of various Hg sources including direct  
103 emissions from anthropogenic and natural sources and indirect secondary formation processes to  
104 atmospheric Hg concentrations in Korea. In order to achieve these objectives, Hg concentrations were  
105 measured in the western most island in Korea, located in between eastern China and the Korean  
106 mainland, so that, depending on wind patterns, the effects of Chinese and Korean Hg emissions could  
107 be evaluated. Previously, our group qualitatively evaluated the impact of local Korean sources and

108 regional Chinese sources on TGM concentrations at the same sampling site (Lee et al. 2014).  
109 However, that work was unable to identify the effect of sources on Hg levels in Korea because only  
110 TGM was measured whereas all three Hg species are needed since they have very different physical  
111 and chemical properties. In this study, the importance of sources and pathways were both qualitatively  
112 and quantitatively evaluated using all three Hg species' concentrations measured throughout the  
113 extended sampling period.

114

## 115 **2. Materials and methods**

### 116 **2.1 Site description**

117 TGM, GOM and PBM were measured on the roof of a three-story building on Yongheung Island  
118 (YI), the westernmost island in Korea (Fig. 1). YI is a small island located about 15~20 km west from  
119 mainland Korea with a population of 5,815. The Yongheung Coal-fired Power Plant (YCPP), located  
120 approximately 4.5 km southwest of the sampling site (Fig 1c), emits about 0.11 ton yr<sup>-1</sup> of Hg. To the  
121 east of the sampling site, industrial (the Incheon industrial complex shown as light violet color in Fig  
122 1b and Fig 1c) and metropolitan (Seoul shown as a pink color in Fig 1b and Fig 1c) areas are located  
123 in mainland Korea, and, in the southern direction, there are three large coal fired power plants (Fig 1b).  
124 The Hg emission rate of anthropogenic sources in Korea was estimated to be 8.04 ton yr<sup>-1</sup> in 2010,  
125 with cement production being the largest source type (AMAP/UNEP, 2013).

126

### 127 **2.2 Sampling and analysis**

128 From January 2013 to August 2014, three atmospheric mercury species: TGM (GEM+GOM), GOM,  
129 and PBM ( $\leq 2.5 \mu\text{m}$ ) were measured during eight intensive sampling periods (Table 1). TGM  
130 concentrations were measured every 5 min using a mercury vapor analyzer (Tekran 2537B). This  
131 instrument contains two gold cartridges which collect and thermally desorb Hg alternately. Desorbed  
132 Hg is quantified using cold vapor atomic fluorescence spectrometry (CVAFS). Outdoor air at a flow  
133 rate of 1.0 L min<sup>-1</sup> was transported through a 3-m-long heated sampling line (1/4" OD Teflon) into the  
134 analyzer. There is a possibility that some of GOM may be sorbed in the line upstream of the gold traps

135 although the sampling line and inlet were maintained at 50°C to prevent the GOM sorption. However,  
136 it is typically assumed that Tekran 2537 can collect and analyze TGM in most of studies (Temme et al.,  
137 2002; Gustin et al., 2013; Han et al., 2014; Zhang et al., 2015). The Tekran 2537B was automatically  
138 calibrated daily using an internal permeation source. Manual injections were also used to evaluate  
139 these automated calibrations before each sampling campaign using a saturated mercury vapor  
140 standard. The relative percent difference between manual injection and automated calibration was  
141 <2%. Five-point manual calibration was also performed by injecting Hg vapor into the sampling line  
142 two times during the study period. The R<sup>2</sup> ranged from 0.9991 to 0.9997 between mass injected and  
143 Tekran reported area, and the average relative percent difference between the mass injected and the  
144 mass calculated was 5.5%. The method detection limit (0.04 ng m<sup>-3</sup>) was calculated as three times the  
145 standard deviation obtained after injecting 1 pg of the mercury vapor seven times. The recovery rate  
146 (96 ± 3%) was obtained by directly injecting Hg vapor into the sampling line between the sample inlet  
147 and the Tekran 2537B in a zero-air stream.

148 GOM and PBM were collected manually using an annular denuder coated with KCl followed by a  
149 quartz filter, respectively, at a flow rate of 10 L min<sup>-1</sup>. To identify any diurnal variations, all samples  
150 were separately collected during the daytime (07:00-19:00) and nighttime (19:00-07:00) except during  
151 the 7<sup>th</sup> sampling period when they were measured every 2 hrs. The sampling system including an  
152 elutriator, an impactor, a KCl-coated denuder and a filter pack was housed in a custom-made sampling  
153 box maintained at 45°C to prevent hydrolysis of KCl. After sampling, the denuder and quartz filter  
154 were thermally desorbed using a tube furnace at 525°C and 900°C, respectively, to convert Hg<sup>2+</sup> to  
155 Hg<sup>0</sup> in a carrier gas of zero air. The heated air was then transported into a Tekran 2537B for  
156 quantification. Field blanks for GOM and PBM were collected once for each sampling period, and  
157 their average values were 0.23 ± 0.12 pg m<sup>-3</sup> and 0.25 ± 0.09 pg m<sup>-3</sup>, respectively.

158 The sampling methods used in this study are currently the most accepted methods for the  
159 measurement of atmospheric GOM and PBM, however there are many studies reporting that these  
160 methods are subject to interferences from ozone, water vapor and possibly other compounds (Lyman  
161 et al. 2010; Talbot et al., 2011; Jeff et al., 2014; Finley et al., 2013; Gustin et al., 2013; Huang et al.,

162 2013; McClure et al., 2014) although recent side-by-side measurements with two Tekran systems  
163 showed good agreement and no impact from added ozone and increasing relative humidity (Edgerton,  
164 personal communication). Also, it should be noted that the concentrations of PBM measured during  
165 12-hrs of sampling time (all sampling periods except in the 7<sup>th</sup>) may have been biased due to Hg loss  
166 from filters over the long sampling period; however for model development any loss of PBM is  
167 assumed to be the same for each sampling period. Therefore, it should be noted that the GOM and  
168 PBM measurements reported in this study may be somewhat biased even though, at present, it is not  
169 possible to quantify the magnitude of these uncertainties.

170 Meteorological data including temperature, wind speed, wind direction, relative humidity and solar  
171 radiation were also measured every 5 min at the sampling site using a meteorological tower (DAVIS  
172 Inc weather station, Vintage Pro2<sup>TM</sup>).

173 Hourly concentrations of SO<sub>2</sub>, NO<sub>2</sub>, CO, O<sub>3</sub> and PM<sub>10</sub> were obtained from the national air quality  
174 (NAQ) monitoring station (<http://www.airkorea.or.kr/>) located approximately 8 km east from the  
175 sampling site. These concentrations were compared with those measured at another national air  
176 quality monitoring station located approximately 24 km west of the Hg sampling site, and there were  
177 no statistical differences between sites (p-value < 0.001), indicating that the spatial distribution of  
178 these pollutants was relatively uniform across the area

179

### 180 **2.3. Backward trajectory and Cluster analysis**

181 Three-day backward trajectories were calculated using the NOAA HYSPLIT 4.7 with GDAS  
182 (Global Data Assimilation System) meteorological data which supplies 3-hour, global 1 degree  
183 latitude-longitude datasets of the pressure surface. Hourly 3-day back-trajectories were calculated for  
184 each hour of sampling, and the arrival heights of both 200 m and 500 m were used to describe the  
185 local and the regional transport meteorological pattern. Since GOM and PBM were measured for 12hr  
186 for most time periods, hourly trajectories were matched to the 12hr-averaged GOM and PBM  
187 concentration; therefore, in total 12 back-trajectories represented one averaged GOM or PBM  
188 concentrations. Although there have been many studies using different time scales for measurements

189 of pollutants and for meteorological data (Amato and Hopke, 2012; Galindo et al., 2011; Kim et al.,  
190 2007) this mismatch might increase the uncertainty for the trajectory-based approaches.

191 The backward trajectories were clustered into groups with similar transport patterns using NOAA  
192 HYSPLIT 4.7. This method minimizes the intra-cluster differences among trajectories while  
193 maximizing the inter-cluster differences. The clustering of trajectories is based on the total spatial  
194 variance (TSV) method. TSV is the sum of all the cluster spatial variances (SPVAR) which is the sum  
195 of the squared distances between the endpoints of the cluster's component trajectories and the mean of  
196 the trajectories in that cluster. In this study, five clusters were chosen based on a large increase in TSV  
197 for larger clusters (Fig. 4S), as described in Draxler et al. (2014) and Kelly et al. (2012). A more  
198 detailed description of the clustering process can be obtained in Draxler et al. (2014).

199

## 200 **2.5. Conditional Probability Function (CPF)**

201 The conditional probability that a given concentration from given wind direction will exceed a  
202 predetermined threshold criterion, was calculated using the following equation.

$$203 \quad \text{CPF}_{\Delta\theta} = \frac{m_{\Delta\theta}}{n_{\Delta\theta}} \quad (\text{Eq.1})$$

204 where  $m_{\Delta\theta}$  is the number of occurrences from wind sector  $\Delta\theta$  where the concentration is higher than a  
205 criterion value, and  $n_{\Delta\theta}$  is the total number of occurrence from this wind sector. In this study, 16  
206 sectors were used ( $\Delta\theta = 22.5^\circ$ ), and calm winds were excluded from the calculation because of  
207 isotropic behavior of the wind vane for such conditions. For TGM, two threshold criteria of the upper  
208 10 and 25 percentile were chosen while only the upper 25 percentile was used for GOM and PBM  
209 concentrations due to the smaller number of samples. The 1hr-averaged wind direction (WD) data  
210 were used for 12hr-averaged GOM and PBM concentrations for most of periods, so that in total 12  
211 WS and WD were used for one averaged GOM and PBM concentrations to create CPF. It should be  
212 noted that there are some unquantifiable uncertainties derived from using different time resolutions  
213 between measurements of GOM and PBM and WD data although overall trajectories did not diverge  
214 significantly at this sampling site for most sampling periods.

215



## 216 2.6. Potential Source Contribution Function (PSCF)

217 The PSCF model counts each trajectory segment endpoint that terminates within given grid cell. A  
218 high PSCF value signifies a potential source location. The PSCF value was calculated as:

$$219 \text{ PSCF value} = \frac{P[B_{ij}]}{P[A_{ij}]} = \frac{m_{ij}}{n_{ij}} \quad (\text{Eq.2})$$

220 If  $N$  is the total number of trajectory segment endpoints over the study period and if  $n$  segment  
221 trajectory endpoints fall into the  $ij$ th cell, the probability of this event ( $P[A_{ij}]$ ) is calculated by  $n_{ij}/N$ . If  
222  $m_{ij}$  is the number of segment endpoints in the same  $ij$ th cell when the concentrations are higher than a  
223 criterion value, the probability of this high concentration event,  $B_{ij}$ , is given by  $P[B_{ij}] = m_{ij}/N$ . The  
224 criterion value was the top 25% concentration and the cell size of  $0.5^\circ$  by  $0.5^\circ$  was used for tracing  
225 sources. To reduce the uncertainty in a grid cell with a small number of endpoints, an arbitrary weight  
226 function  $W_{ij}$  was applied when the number of the endpoints in a particular cell was less than three  
227 times the average number of endpoints ( $N_{ave}$ ) for all cells (Fu et al., 2011, Han et al., 2007, Polissar et  
228 al., 2001a, 2001b).

$$229 W_{ij} = \begin{pmatrix} 1.0 & N_{ij} > 3N_{ave} \\ 0.70 & 3N_{ave} > N_{ij} > 1.5N_{ave} \\ 0.40 & 1.5N_{ave} > N_{ij} > N_{ave} \\ 0.20 & N_{ave} > N_{ij} \end{pmatrix} \quad (\text{Eq. 3})$$

230 In this study, trajectories were hourly calculated

231

## 232 3. Results and discussion

### 233 3.1. General trends of three Hg species

234 To maintain the consistency of the sampling duration, the 12hr-averaged GOM and PBM  
235 concentrations were used for the 7<sup>th</sup> sampling period to identify the general trends of Hg species. The  
236 average TGM, GOM, and PBM concentrations were  $2.8 \pm 1.1 \text{ ng m}^{-3}$ ,  $8.3 \pm 9.7 \text{ pg m}^{-3}$ , and  $10.9 \pm$   
237  $11.2 \text{ pg m}^{-3}$ , respectively (Table 1). Since the GOM concentration was much lower than TGM the  
238 reported TGM concentration can be considered a good approximation of the GEM concentration.  
239 TGM varied from 0.1 to  $18.8 \text{ ng m}^{-3}$ ; the highest concentration was observed around 2 am on March  
240 18, 2014 (Fig. 2). GOM and PBM concentrations peaked at  $50.9 \text{ pg m}^{-3}$  during the daytime on March

241 19, 2014 and  $43.7 \text{ pg m}^{-3}$  during daytime on January 22, 2013, respectively (Fig. 2). The various Hg  
242 species did not follow similar concentration patterns although PBM was statistically significantly  
243 correlated with TGM (Pearson correlation coefficient,  $r= 0.235$ ,  $p\text{-value}= 0.03$ ).

244 When the data were grouped into three categories including the first (Apr., 2013, May, 2013, Mar.,  
245 2014, May, 2014), the second (Aug, 2013, Aug., 2014.), and the third (Jan., 2013, Feb., 2013) periods,  
246 both TGM (ANOVA/Tukey test,  $p\text{-value}<0.001$ ) and PBM ( $p\text{-value}=0.024$ , Kruska-Wallis test) had  
247 the highest concentrations in cold period ((Jan., 2013, Feb., 2013) while there was no statistical  
248 difference in GOM concentrations among different categories ( $p\text{-value}= 0.288$ , Kruskal-Wallis test)  
249 (Fig. 3). Observed TGM concentrations were substantially lower than those measured in a suburban  
250 and remote site in China and metropolitan areas of Korea (Seoul), but higher than at most North  
251 American sites and at a rural site of Korea (Chuncheon) (Table 2). GOM and PBM concentrations  
252 were in between those typically found at urban locations and at a rural site in Korea and were much  
253 lower than those typically measured in China.

254 The TGM concentration varied diurnally, generally showing morning maximums (07:00-12:00) and  
255 minimums during the nighttime. In urban areas, TGM concentrations are typically higher during the  
256 nighttime due to a combination of decreased GEM loss by daytime oxidation, increased use of  
257 household heating systems and decreased mixing heights at night (Kim et al., 2012; Han et al., 2014).  
258 In contrast daytime peaks have been observed in rural and remote areas, likely due to increased  
259 volatilized of  $\text{Hg}^0$  from natural sources (Choi et al., 2008; Cheng et al., 2014). Overall these results  
260 suggest that TGM concentrations at this site are elevated due to the proximity of regional sources and  
261 daily variations are controlled by natural emissions from the ocean and soil surfaces.

262 GOM concentrations were the highest in spring ( $10.7 \pm 10.1 \text{ pg m}^{-3}$ ) and the lowest in summer  
263 ( $6.2 \pm 4.9 \text{ pg m}^{-3}$ ) with statistically insignificant seasonal variation (Fig. 3). The lack of a GOM  
264 seasonal variation for could be an indicator of insignificant secondary formation through  
265 photochemical oxidation reactions, but it might be also be due to the small sample numbers and/or  
266 relatively long sampling duration (12hr). PBM concentrations did have statistically significant  
267 seasonal variations with the highest average concentration in winter ( $17.8 \pm 16.7 \text{ pg m}^{-3}$ ) and the lowest

268 average concentration in summer ( $5.8 \pm 4.1 \text{ pg m}^{-3}$ ) (Fig. 3). Higher PBM concentrations in winter  
269 were likely caused by increased biomass burning and residential heating, decreased removal from the  
270 atmosphere due to the lower precipitation depth, and/or lower temperatures which favor partitioning  
271 to the aerosol phase. Previous studies also often observed the highest PBM concentrations in winter  
272 (Mao et al., 2012; Amos et al., 2012; Lan et al., 2012).

273 In Korea emissions of PBM from anthropogenic sources are much smaller than gaseous emissions  
274 (the proportion of GEM, GOM, and PBM released are 64.4%, 28.8%, and 6.8%)(Kim et al., 2010).  
275 The fact that PBM concentrations are similar to GOM even though significantly less PBM is released  
276 suggests that a significant portion of atmospheric PBM may be due to secondary formation through  
277 gas-particle partitioning. This process is characterized by a partition coefficient,  $K_p$ , which is inversely  
278 correlated with temperature (Rutter and Schauer, 2007), possibly causing the distinct seasonal  
279 variation in PBM concentrations at the sampling site.

280 The relationship between  $K_p$ , defined as:

$$281 \quad K_p = \frac{PBM/PM}{Hg_{gas}} \quad (\text{Eq. 4})$$

282 where PM represents the particle mass, and  $Hg_{gas}$  is the concentration of gaseous Hg and relative  
283 humidity (RH) was examined (PM<sub>10</sub> concentration was used for PM in Eq. 4 in this study). RH was  
284 included since in recent studies  $K_p$  was found to increase at high relative humidity in colder seasons  
285 (Lyman and Keeler, 2005; Liu et al., 2010). Note that the sampling site was located in a coastal area  
286 with generally high RH.

287 Some previous studies suggested that all gaseous mercury species including  $Hg^0$  may deposit on  
288 particles (Xiu et al., 2005; 2009); however, others suggested that the gas-particle partitioning of GOM  
289 occurred but assumed that the adsorption of  $Hg^0$  on particles was negligible due to its high vapor  
290 pressure (Amos et al., 2012; Rutter and Schauer, 2007). Consistent with this hypothesis, we found a  
291 statistically significant multiple linear relationship between  $K_p$  with temperature and relative humidity  
292 (Fig. 4):

$$293 \quad \text{Log}(K_p) = -2.518 - 0.036(T) + 0.017(RH) \quad (\text{Eq. 5})$$

294 where T and RH indicate atmospheric temperature ( $^{\circ}\text{C}$ ) and relative humidity (%), respectively. The

295 multiple linear equation fit the data well ( $R^2=0.29$ ,  $R=0.54$ ,  $p\text{-value}<0.001$ ), and both variables of  
 296 temperature and relative humidity were statistically significant ( $p\text{-value}<0.001$ ). The partial  
 297 correlation coefficients were  $-0.389$  for temperature ( $p\text{-value}<0.001$ ) and  $0.375$  for RH ( $p\text{-}$   
 298  $\text{value}=0.001$ ). When each of the temperature and the RH was used as a single independent variable  
 299 the  $\log(K_p)$  regression equation was still significant with a Pearson correlation coefficient of  $-0.42$   
 300 ( $R^2=0.18$ ,  $p\text{-value}<0.001$ ) and  $0.39$  ( $R^2= 0.15$ ,  $p\text{-value}< 0.001$ ), somewhat lower than that from the  
 301 multiple regression (Fig. 1S). Amos et al. (2012) found an empirical gas-particle partitioning  
 302 relationship between  $K_p$  and T using the Hg data obtained from five monitoring sites in United States  
 303 and Canada and two laboratory experiments, and average equation is :

$$\text{Log}(1/K_p) = (10 \pm 1) - (2500 \pm 300)/T \quad (\text{Eq. 6})$$

304 where T is in Kelvin unit. When a partial linear regression was performed with T in this study similar  
 305 equation was derived:  $\text{Log}(1/K_p) = 13.5 - 3362.7/T$ . In Amos et al. (2012), the coefficients,  $\beta$  and  $y_0$ ,  
 306 ranged from  $-1600$  to  $-3300$  and  $6$  to  $13$ , respectively, and  $R^2$  ranged from  $0.16$  to  $0.57$  in different  
 307 monitoring sites. Rutter and Schauer (2007a) also determined the relationship between  $\text{Log}(1/K_p)$  for  
 308 urban aerosols with inverse temperature and found the slope and intercept were  $-4250 \pm 480$  and  $10 \pm 2$ ,  
 309 respectively, with  $R^2$  of  $0.77$ . Another study using datasets from 10 AMNet sites located in North  
 310 America found  $R^2$  ranged between  $0.04$  and  $0.53$ . In summary the values derived in this study fall  
 311 between those reported by other studies. Somewhat lower  $R^2$  in this study is probably caused by  
 312 smaller number of samples, different composition of aerosol, different GOM species, and/or longer  
 313 sampling duration.

315 Han et al. (2014) also found a significant multiple linear relationship between the ratio of  
 316 PBM/GOM with temperature and relative humidity at a rural site ( $R^2= 0.613$ ,  $\beta$  for T=  $-0.774$ ,  $\beta$  for  
 317 RH=  $0.33$ ) but not at an urban site. The lower correlation coefficient and the beta values found in this  
 318 study compared to those from Han et al. (2014) is probably due to the greater impact of primary  
 319 anthropogenic sources around the sampling site. A large anthropogenic source near the sampling site  
 320 is likely to weaken the relative contribution of gas-particle partitioning (secondary formation) to the  
 321 variations of ambient PBM and GOM concentration because both Hg species can be also strongly

322 affected by the primary source emission. In the study of Amos et al. (2012), the highest  $R^2$  was shown  
323 in Experimental Lakes Area of Canada and Reno, NV where no large anthropogenic sources of Hg is  
324 located.

325 In previous researches, GOM concentration measured using KCl denuder is subject to interferences  
326 under the conditions of high ozone and relative humidity (Gustin et al., 2012; Huang et al., 2013;  
327 Gustin et al., 2015) although it is currently the most accepted method. To evaluate the possible error  
328 on interpreting result,  $K_p$  was re-calculated with re-calculated GOM concentration using the empirical  
329 equation suggested by McClure et al. (2014) ( $RH=0.63 \text{ GOM loss \%} + 18.1$ ). Since McClure et al.  
330 (2014) suggested this equation at RH of 21 to 62%, the GOM concentrations collected only when RH  
331 was from 20% to 65% were re-calculated. The multi-linear relationship was compared between re-  
332 calculated  $K_p$  and original  $K_p$ , and the results were very similar to each other (Fig. 2S).

333

### 334 **3.2. Tracing sources of Hg species**

335 Correlations between Hg and other pollutant concentrations are often used to identify sources. For  
336 example good correlations with  $\text{SO}_2$  and CO typically indicate the impact of coal combustion (Pirrone  
337 et al., 1996; Han et al., 2014), and a strong correlation between Hg and CO has often been used as an  
338 indicator for regional transport because both pollutants have similar sources and do not easily  
339 decompose by reaction and undergo deposition during transport (Weiss-Penzias et al., 2003, 2006;  
340 Kim et al., 2009) although a few recent studies showed the significant bromine-induced oxidation of  
341 GEM in the mid-latitude marine boundary layer as well as in the polar atmosphere (Ariya, 201; Obrist  
342 et al., 2011). A good correlation between Hg and  $\text{NO}_2$  suggests the site is being impacted by local  
343 sources because the lifetime of  $\text{NO}_2$  is relatively short compared with that of CO (Seinfeld and Pandis,  
344 2006). In this study TGM concentrations were well correlated with  $\text{SO}_2$ , CO, and  $\text{PM}_{10}$  concentrations  
345 but not with  $\text{NO}_2$  concentrations (Table 3), indicating that regional transport of TGM emitted from  
346 coal combustion was impacting the site throughout much of the sampling period. PBM concentrations  
347 also had a statistically significant relationship with TGM and CO suggesting regional transport is also  
348 important for PBM, but GOM was not correlated with any other pollutant suggesting it is impacted to

349 a greater extend by local sources (see additional discussion below).

350 CPF plot shows that the top 25% TGM concentrations were associated with winds from the NNW  
351 and eastern direction, pointing towards northeastern China and inland Korean sources; however, when  
352 the criterion was set to the top 10% the winds from NNW became less important and the sources  
353 located in southern and eastern areas from the sampling site were identified as an important source  
354 direction (Fig. 5). The CPF plot for GOM is significantly different from the one for PBM. High PBM  
355 concentrations were associated with northern winds while GOM concentrations were enhanced during  
356 southeastern winds.

357 These results suggest that for PBM regional transport from Chinese and North Korea sources were  
358 more important than Korean sources; in contrast coal fired power plants located in the southern  
359 direction rather than regional transport impacted GOM concentration. It should be noted that this  
360 result is in apparent conflict with the finding that there was no relationship between GOM and SO<sub>2</sub>  
361 concentrations. Total SO<sub>2</sub> emissions from power plants in China (18.6 Tg yr<sup>-1</sup>) are much larger than in  
362 Korea (0.09 Tg yr<sup>-1</sup>), and SO<sub>2</sub> emission rates per capita and per area in China also greatly surpass  
363 those in Korea (Lu et al., 2010). Much larger SO<sub>2</sub> emissions in China raise the background SO<sub>2</sub>  
364 concentration in the region and may mask any correlation between GOM and SO<sub>2</sub> even if coal fired  
365 power plants located in the southerly direction from the sampling site impacted GOM concentrations.  
366 In support of this hypothesis the pollution rose indicates that high SO<sub>2</sub> concentrations are associated  
367 with westerly winds while high GOM concentrations are associated with southerly winds. It should be  
368 noted that only the top 25% of GOM and PBM concentrations were used as the criteria for the CPF  
369 plot because the number of samples for both species were significantly less than for TGM due to their  
370 longer sampling duration (12hr).

371 Among the eight sampling periods, the second period (April, 2013) had the highest TGM, PBM and  
372 the second highest GOM average concentration, and SO<sub>2</sub>, NO<sub>2</sub>, CO, and PM10 were also quite high  
373 (Table 1). During this period, TGM was statistically well correlated with SO<sub>2</sub> (r= 0.55), NO<sub>2</sub> (r=0.56),  
374 and CO (r=0.36), with the highest Pearson correlation coefficient with NO<sub>2</sub>, the characteristic local  
375 pollutant. In addition, the CPF plot (for TGM) and the back-trajectories were also associated with

376 easterly winds transporting air masses from major Korean urban areas, supporting the previous  
377 suggestion that inland sources enhanced all three Hg concentrations during the second sampling  
378 period.

379 In contrast the fifth sampling period had the lowest GOM, PBM and the second lowest TGM  
380 concentrations (Table 1). Note however that the TGM concentrations for the first couple of days  
381 reached approximately  $5 \text{ ng m}^{-3}$  and gradually decreased to about  $1 \text{ ng m}^{-3}$  during the last days of  
382 sampling (Fig. 2), indicating that there was likely two different sources affecting Hg concentrations  
383 during this period. Back-trajectories associated with high TGM concentrations passed through  
384 northeastern China, North Korea and the industrial/metropolitan areas of Korea before arriving at the  
385 sampling site whereas trajectories during the low TGM concentrations spent long periods within the  
386 ocean boundary layer. Although Hg can be emitted from ocean surface (Han et al., 2007; UNEP, 2013)  
387 heavy rain and low solar radiation occurring during the last two days of this period probably inhibited  
388 emissions of Hg from the ocean surface.

389

### 390 **3.2.1 GOM/PBM ratio**

391 According to the CPF results, the winds from NW and NE of the sampling site were responsible for  
392 the elevated PBM concentrations while easterly winds pointing towards inland Korea were associated  
393 with increased GOM concentrations (Fig. 5). The finding that regional transport of TGM and PBM to  
394 the site is important is supported by their significant correlation with CO (Table 3). In order to  
395 identify the relative importance of local sources relative to regional transport, the ratio of GOM/PBM  
396 was used as an indicator because the atmospheric residence time of GOM is widely regarded to be  
397 shorter than that of PBM (even though there is no consensus on what specific chemical forms are  
398 collected by KCl-coated denuders). The GOM/PBM ratio should be higher if local sources are more  
399 important, and the GOM/PBM ratio is likely to decrease as regional transport becomes more  
400 important. In this study the GOM/PBM ratios were categorized into three groups: low (0~2), middle  
401 (2~8), and high (>8) and the frequency of wind direction was compared (Fig. 6). The result clearly  
402 indicates that the southerly and southeasterly winds were associated with high GOM/PBM ratios and

403 that the westerly and northerly winds indicative of regional transport from China prevailed at lower  
404 GOM/PBM ratios. There was a weak negative correlation between the ratio of GOM/PBM and CO  
405 concentration at a significance level of 0.1 ( $p$ -value= 0.089) (Fig. S3), supporting the assertion that the  
406 GOM/PBM ratio decreased with the increased effect of regional transport. Lynam and Keeler (2005)  
407 also found that high GOM/PBM ratio was observed with influences from local sources and low  
408 GOM/PBM ratios appeared with influence from regional sources in Detroit. In Korea, Kim et al.  
409 (2009) found a significant increase in the PBM/GOM ratio during high PM<sub>2.5</sub> concentration events  
410 caused by regional transport from China.

411 The reciprocal of this ratio (i.e. GOM/PBM) was used to calculate  $K_p$  which indicates that the  
412 secondary formation of PBM through gas-particle partitioning was became more important as the  
413 significance of regional transport increased, suggesting that secondary production was favored when  
414 air underwent regional transport, as has been shown previously (Lynam and Keeler, 2005). It should  
415 be noted that there are uncertainties related to different aerosol composition, particle surface area  
416 variations, and/or temperature change which can affect both  $K_p$  and the GOM/PBM ratio.

417

### 418 **3.2.2 PSCF results**

419 In order to locate potential source areas in more detail, PSCF was used. For TGM, potential sources  
420 were located in Liaoning, Shandong, and Henan provinces of China along with the southern area of  
421 Korea (Fig. 7). Liaoning province, where large non-ferrous smelters are situated, is the province with  
422 the largest Hg emissions in China; Shandong and Henan provinces are also large Hg emission areas,  
423 emitting about 30-40 ton yr<sup>-1</sup> (Fu et al., 2012) in part due to a large lead smelter (Wang et al, 2014),  
424 and biomass burning (Huang et al., 2011).

425 The probable source areas of PBM identified by PSCF were similar to those for TGM, indicating  
426 that both Chinese and inland Korean sources enhanced PBM concentrations, with the exception of  
427 metropolitan (Seoul) and industrial (Incheon) areas located in northwestern South Korea which  
428 emerged as more prominent source areas for PBM than for TGM (Fig. 7). Only Korean sources  
429 including metropolitan (Seoul) and the industrial areas in southern Korea were identified as probable



430 source areas for GOM (Fig. 7); regional transport of GOM from China was not important. The Yellow  
431 Sea between China and Korea was also associated with high PSCF values, possibly indicating the  
432 shipping ports located on the western coast of Korea as important source areas. However, it should be  
433 noted that it might be a trailing effect derived by relatively short sampling duration. A trailing effect is  
434 often observed, especially with a limited number of measurements or short sampling period, since  
435 PSCF evenly distributes weight along the path of trajectories so that PSCF results often identify areas  
436 upwind and downwind of real sources as a source area (Han et al., 2007). However, it should be also  
437 noted that the marine boundary layer provides good conditions for active Hg oxidation reactions due  
438 to an abundance of oxidants (Auzmendi-Murua et al., 2014); therefore, the possibility of areas over  
439 the ocean being a GOM source should not be excluded.

440 It should be noted that different temporal resolutions for trajectories (hourly) and concentrations  
441 (every 12hr) were used for GOM and PBM. Since trajectory directions can significantly change over  
442 the course of 12hrs there is a possibility that some source areas could be misidentified, especially for  
443 more distant regional sources. However, upon investigation it was determined that over the 12 hr  
444 sampling periods the trajectories did not diverge significantly at this sampling site for most sampling  
445 periods.

446

### 447 **3.2.3 Source attribution based on cluster analysis**

448 In an effort to quantify the contribution of national and foreign sources to the measured Hg  
449 concentrations the back trajectories were grouped into five clusters using the trajectory cluster  
450 analysis feature of HYSPLIT. Among the five clusters, clusters 1 and 5 represent trajectories  
451 originating from outside (South) Korea whereas the trajectories grouped in the cluster 4 originated  
452 and passed through the (South) Korean peninsula (Fig. 8). Clusters 2 and 3 contain trajectories from  
453 China and the Korean peninsula, but cluster 2 was more associated with Liaoning province and North  
454 Korea while cluster 3 originated more from Shandong and Henan provinces. Clusters 1 through 5  
455 contributed 12%, 31%, 26%, 20%, and 11% of the total time, respectively, and the associated  
456 concentrations with each cluster are shown in Table 4. Concentration ranges of three Hg species for

457 each cluster were shown as the box-and-whisker plots in the supplementary file (Fig. 5S). When  
 458 considering that cluster 4 is associated with the local transport from inland Korea and the cluster 1 and  
 459 5 are associated with the regional transport from outside of (South) Korea, the maximum and 75<sup>th</sup>  
 460 percentile values as well as the arithmetic average are higher in cluster 4 for GOM and PBM than  
 461 those in the clusters 1 and 5 (Fig. 5S).

462 The TGM concentration was the highest for cluster 5; however, GOM and PBM concentrations had  
 463 the lowest averages for this cluster. Cluster 5 contains the back-trajectories originating from Mongolia  
 464 and Russia and passing through northeastern China before arriving at the sampling site, which  
 465 suggests regional transport was important for this cluster (Fig. 8). Average CO concentrations were  
 466 pretty similar for all clusters, but it was the second highest for cluster 5 (cluster 2 was highest). The  
 467 highest total average GOM and PBM concentrations were associated with cluster 4 which includes  
 468 trajectories distributed over the Korean peninsula, suggesting that Korean sources were responsible  
 469 for the enhanced GOM and PBM concentrations. For cluster 4, the highest Pearson correlation  
 470 coefficient between GOM and PBM concentrations ( $r=0.721$ ) was observed, indicating that both Hg  
 471 species were emitted from similar sources. For other clusters, there were no statistically significant  
 472 correlations between GOM and PBM except for cluster 2 ( $r=0.209$ ,  $p\text{-value}<0.001$ ). In addition, both  
 473 average  $\text{NO}_2$  concentration ( $19.4 \pm 14.9$  ppb) and the correlation coefficient between  $\text{NO}_2$  and TGM  
 474 ( $r= 0.688$ ) were the highest for cluster 4, supporting the finding of impact from Korean sources.

475 In order to consider both Hg concentration and the fraction of time for each cluster, the following  
 476 equation was used to quantify the effect of Korean and out-of- Korean sources to the Hg concentration  
 477 at the receptor site.

$$478 \quad \text{source contribution of cluster, } i = \frac{\left(\frac{N_i}{N_{total}}\right) \times C_j}{\sum_{i=1}^n \left\{ \left(\frac{N_i}{N_{total}}\right) \times C_j \right\}} \quad (\text{Eq.6})$$

479 where  $(N_i/N_{total})$  indicates the percentage of time associated with the cluster,  $i$ ,  $i$  is the cluster number,  
 480  $n$  is the number of clusters (equal to five in this study), and  $C_j$  indicates the average Hg concentration  
 481 associated with the cluster,  $i$ . Compared to the other clusters, the source contributions of clusters 1 and  
 482 5, which represent regional transport, were relatively low for all Hg species (Table 4). Cluster 4

483 contributed more significantly, especially for GOM and PBM, indicating the importance of Korean  
484 sources. The source contribution of cluster 2 was the highest for PBM compared to other Hg species,  
485 suggesting that North Korean sources were an important contributor to the high PBM concentrations  
486 measured, likely due to coal and biomass burning in North Korea (Kim et al., 2013; NI, 2001; NI,  
487 2003).

488 In order to quantify the contribution of Korean vs. out-of-Korean sources (note that “Korean”  
489 means “South Korean” throughout the manuscript), the source contributions of the clusters were used.  
490 Clusters 1 and 5 were used to represent the effect of sources outside of Korea and the cluster 4 was  
491 used to indicate the effect of sources in Korea. Since clusters 2 and 3 contain mixed trajectories from  
492 Korea and out-of-Korea their contribution was divided evenly between in and out of Korea. The  
493 results indicate that the sources in Korea and outside Korea contributed about 50% each to the  
494 concentration variation of TGM measured at the sample site during the sampling period while the  
495 Korean sources affected GOM and PBM more significantly, accounting for approximately 52.3% and  
496 53.4%, respectively (Table 4). These results augment the CPF and PSCF results which only use  
497 concentrations that are in the top 25<sup>th</sup> percentile. While CPF and PSCF found that for high  
498 concentration events Korean sources were most important for GOM while for TGM and PBM  
499 regional transport from China and North Korea were also important, the cluster based approach  
500 suggests that for all 3 species Korean and out-of-Korean sources contributed approximately 50% each  
501 to the concentration variations seen by the site. When the geometric mean concentrations were used  
502 for each cluster a similar result was obtained for relative contributions as the results using arithmetic  
503 mean concentrations.

504 It should be noted that errors always exist in calculating trajectories, causing uncertainties in all  
505 trajectory-based approaches. Trajectory errors vary considerably from case to case; Stohl (1998)  
506 suggested uncertainties might be 20% of the distance travelled by trajectories while Draxler (1996)  
507 found that the final error was about 10% of the travel distance.

508

#### 509 **4. Conclusion and Implications**

510 This study was initiated to identify the sources affecting speciated mercury concentrations  
511 measured on an island located between mainland Korea and Eastern China. Various tools were used  
512 to locate and quantify the sources, including correlations with other pollutants, CPF, and the back-  
513 trajectory based analysis (PSCF and cluster analysis). The results consistently show that Korean  
514 sources are most important for GOM while for other Hg species (TGM and PBM) regional transport  
515 from China and North Korea were also important. Existing methods including PSCF and CPF are able  
516 to locate the source direction and areas, but do not consider the frequency of the wind directions  
517 which can affect the long-term concentrations at the receptor site. For example, if the Hg  
518 concentration is high with easterly winds both CPF and PSCF identify the eastern areas as important  
519 source areas even if, in fact, winds are rarely blowing from east. In this work, it is true that sources  
520 located in the eastern direction from the sampling site are likely to be important for enhancing Hg  
521 concentrations, but based only on CPF and PSCF results it cannot be said that their contribution to the  
522 concentration variations at the site is also high.

523 Do address this problem a new approach that considers both the cluster frequency and the Hg  
524 concentration associated with each cluster was used to quantify the source contribution at the  
525 sampling site. On average, contributions from out-of-Korean sources were similar to Korean sources  
526 for TGM whereas Korean sources contributed slightly more to the concentration variations of GOM  
527 and PBM compared to the out-of-Korean sources. However, in general, conclusions using this  
528 approach are more uncertain when the concentration ranges are similar between clusters. Additional  
529 work is needed with this approach to determine if a different statistic (other than mean) would provide  
530 better results when there are not distinct concentration differences between clusters. In addition,  
531 uncertainties exist in the source attribution approach based on cluster analysis because the trajectories  
532 inevitably overlap between different clusters since the cluster analysis accounts for both variations in  
533 transport speed and direction simultaneously. Nevertheless, this new approach can augment existing  
534 methods including CPF and PSCF to help identify source contributions to the concentration variations  
535 at the sampling site.

536 The ratio of GOM/PBM proved to be a useful tool for identifying the relative significance of local

537 sources vs. regional transport. The GOM/PBM ratio decreased as the effect of regional transport  
538 increased and vice versa since GOM has a shorter atmospheric residence time than PBM. The  $K_p$   
539 calculated using the reciprocal of the GOM/PBM ratio was negatively correlated with atmospheric  
540 temperature and positively correlated with relative humidity, suggesting that the secondary formation  
541 of PBM was an important source of atmospheric PBM concentration at low temperature and high  
542 relative humidity. This result also suggests that the secondary formation of PBM becomes more  
543 favored when the air undergoes regional transport rather than local transport.

544

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550

#### 551 **Author Contributions**

552 The work presented here was carried out in collaboration between all authors. Gang S. Lee analyzed  
553 data and wrote the paper. Pyung R. Kim performed the experiments and interpreted the results. Young  
554 J. Han defined the research theme, interpreted the results, and wrote the paper. Yong S. Seo, Seung.  
555 M. Yi, and Thomas M. Holsen also interpreted the results and approved the final paper.

556

#### 557 **Conflicts of Interest**

558 The authors declare no conflict of interest.

559

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855 **Table list**

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882 **Figure list**

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910 trajectory and contribution for each cluster.

Table 1 Summarized concentrations of speciated Hg and other typical pollutants for each sampling period

Sampling periods	TGM (ng m <sup>-3</sup> )	GOM (pg m <sup>-3</sup> )	PBM (pg m <sup>-3</sup> )	SO <sub>2</sub> (ppb)	NO <sub>2</sub> (ppb)	CO (ppm)	O <sub>3</sub> (ppb)	PM <sub>10</sub> (μg m <sup>-3</sup> )
1 <sup>st</sup> : 2013.01.17 ~ 01.23	3.5 ± 0.8	5.8 ± 8.8	17.0 ± 16.5	5.8 ± 4.2	26.7 ± 12.4	0.5 ± 0.2	21.7 ± 13.5	55.9 ± 38.5
2 <sup>nd</sup> : 2013.02.25 ~ 03.01	3.7 ± 0.9	13.2 ± 14.8	19.5 ± 19.6	7.3 ± 3.0	28.8 ± 19.1	0.7 ± 0.2	28.3 ± 14.7	83.6 ± 28.6
3 <sup>rd</sup> : 2013.04.08 ~ 04.13	2.1 ± 0.4	4.3 ± 5.6	15.6 ± 13.8	4.9 ± 1.7	8.6 ± 3.3	0.4 ± 0.1	46.8 ± 6.0	45.8 ± 21.4
4 <sup>th</sup> : 2013.05.20 ~ 05.25	2.8 ± 1.0	4.2 ± 5.9	6.7 ± 7.3	5.7 ± 2.5	17.2 ± 5.9	0.5 ± 0.1	36.2 ± 19.4	51.9 ± 21.6
5 <sup>th</sup> : 2013.08.19 ~ 08.24	2.3 ± 0.9	3.2 ± 3.0	5.4 ± 4.6	4.7 ± 1.6	10.2 ± 3.8	0.5 ± 0.1	40.9 ± 29.6	34.2 ± 13.7
6 <sup>th</sup> : 2014.03.17 ~ 03.21	2.6 ± 1.2	12.8 ± 15.7	7.3 ± 3.5	7.5 ± 0.1	32.9 ± 75.1	0.4 ± 0.2	39.0 ± 12.5	66.7 ± 44.4
7 <sup>th</sup> : 2014.05.26 ~ 05.31	2.8 ± 0.7	13.5 ± 9.0	9.8 ± 12.2	5.8 ± 2.3	12.4 ± 5.6	0.5 ± 0.1	85.5 ± 23.3	124.5 ± 44.9
8 <sup>th</sup> : 2014.08.19 ~ 08.23	2.4 ± 1.1	10.7 ± 3.7	6.3 ± 3.6	3.4 ± 1.2	8.7 ± 6.0	0.4 ± 0.1	38.7 ± 16.5	32.0 ± 22.3
Average	2.8 ± 1.1	8.3 ± 9.7	10.9 ± 11.2	5.6 ± 5.0	18.2 ± 28.7	0.5 ± 0.2	42.1 ± 25.8	61.9 ± 42.8

Table 2. Comparisons of measured Hg concentrations with those reported in other studies.

Country	Site	Remarks	Year	TGM (ng m <sup>-3</sup> )	GOM (pg m <sup>-3</sup> )	PBM (pg m <sup>-3</sup> )	Reference
Korea	Seoul	Urban	2005-2006	3.22 ± 2.10	27.2 ± 19.3	23.9 ± 19.6	Kim et al., 2009
	Seoul	Urban	2006-2009	3.72 ± 2.96	11.3 ± 9.5	13.4 ± 12.0	Han et al., 2014
	Chuncheon	Rural	2006-2009	2.12 ± 1.47	2.7 ± 2.7	3.7 ± 5.7	Han et al., 2014
	<b>Yongheung</b>	<b>Island</b>	<b>2013-2014</b>	<b>2.8 ± 1.1</b>	<b>9.8 ± 9.9</b>	<b>10.6 ± 12.0</b>	<b>This study</b>
China	Guiyang	Urban	2009	9.7 ± 10.2	35.7 ± 43.9	368.0 ± 676.0	Fu et al., 2011
	Xiamen	Suburban	2012-2013	3.5	61.05	174.41	Xu et al., 2015
	Mt. Changbai	Remote	2005-2006	3.58 ± 1.78	65 ± 84	77 ± 136	Wan et al., 2009a,b
USA	Chicago	Urban	2007	2.5 ± 1.5	17 ± 87	9 ± 20	Gratz et al., 2013
	Reno, Nevada	Suburban	2007-2009	2.0 ± 0.7	18 ± 22	7 ± 7	Lyman and Gustin, 2009
	Rochester, NY	Urban	2008-2009	1.49	4.08	6.57	Huang et al., 2010
Canada	Nova Scotia	Urban	2010-2011	1.67 ± 1.01	2.07 ± 3.35	2.32 ± 3.09	Cheng et al., 2014
	Ontario	Rural	2006-2007	1.17	15.10	16.40	Baya and Van Heyst, 2010

Table 3. Correlation coefficients and p-values (in parenthesis) for speciated Hg with other pollutants for the whole sampling period. Correlation coefficients with an asterisk indicate a statistically significant relationship at  $\alpha = 0.05$ .

	<b>TGM</b>	<b>GOM</b>	<b>SO<sub>2</sub></b>	<b>NO<sub>2</sub></b>	<b>CO</b>	<b>O<sub>3</sub></b>	<b>PM<sub>10</sub></b>
<b>TGM</b>	-	-0.132 (0.233)	<b>0.115*</b> (0.001)	0.063 (0.074)	<b>0.571*</b> (<0.001)	-0.055 (0.115)	<b>0.401**</b> (<0.001)
<b>GOM</b>	-0.132 (0.233)	-	0.025 (0.822)	0.022 (0.846)	-0.149 (0.180)	0.143 (0.197)	0.128 (0.248)
<b>PBM</b>	<b>0.235*</b> (0.030)	0.021 (0.855)	-0.006 (0.954)	0.008 (0.941)	<b>0.215*</b> (0.048)	0.029 (0.794)	0.139 (0.206)

Table 4. Estimated contribution of Korean and out-of-Korean sources on variations of speciated Hg concentration.

\*: TGM is shown in  $\text{ng m}^{-3}$  while for both GOM and PBM the units are  $\text{pg m}^{-3}$ .

Cluster	Cluster frequency (%)	Average concentration *			Source contribution (%)			Korean (%)			Out-of-Korean (%)		
		TGM	GOM	PBM	TGM	GOM	PBM	TGM	GOM	PBM	TGM	GOM	PBM
<b>1</b>	12	2.2	6.9	7.7	9.6	10.5	8.6				9.6	10.5	8.6
<b>2</b>	31	2.8	8.3	12.6	31.5	32.5	36.5	15.7	16.3	18.2	15.7	16.3	18.2
<b>3</b>	26	2.9	8.1	10.0	27.3	26.6	24.3	13.7	13.3	12.1	13.7	13.3	12.1
<b>4</b>	20	2.6	9.0	12.3	18.9	22.8	23.0	18.9	22.8	23.0			
<b>5</b>	11	3.2	5.5	7.4	12.8	7.6	7.6				12.8	7.6	7.6
<b>Korean</b>								<b>48.3</b>	<b>52.3</b>	<b>53.4</b>			
<b>Out-of-Korean</b>											<b>51.7</b>	<b>47.7</b>	<b>46.6</b>

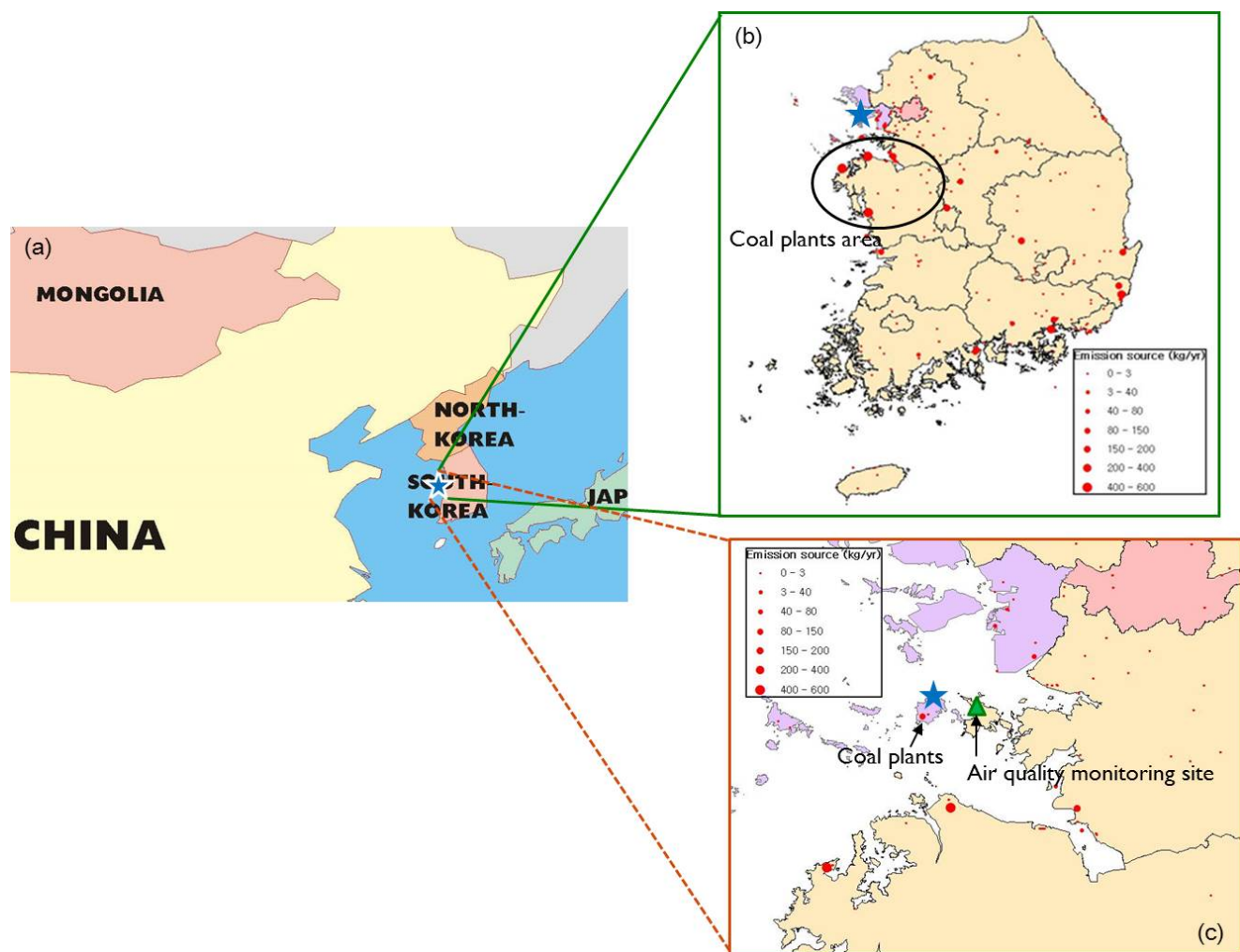


Figure 1. (a) The sampling site in Yongheung Island (the star mark). (b) Anthropogenic mercury emission sources in Korea. Blue star indicates the sampling site, and the green circle indicates the area where major Korean coal-fired power plants are located. (c) The enlarged image of the area near the sampling site.

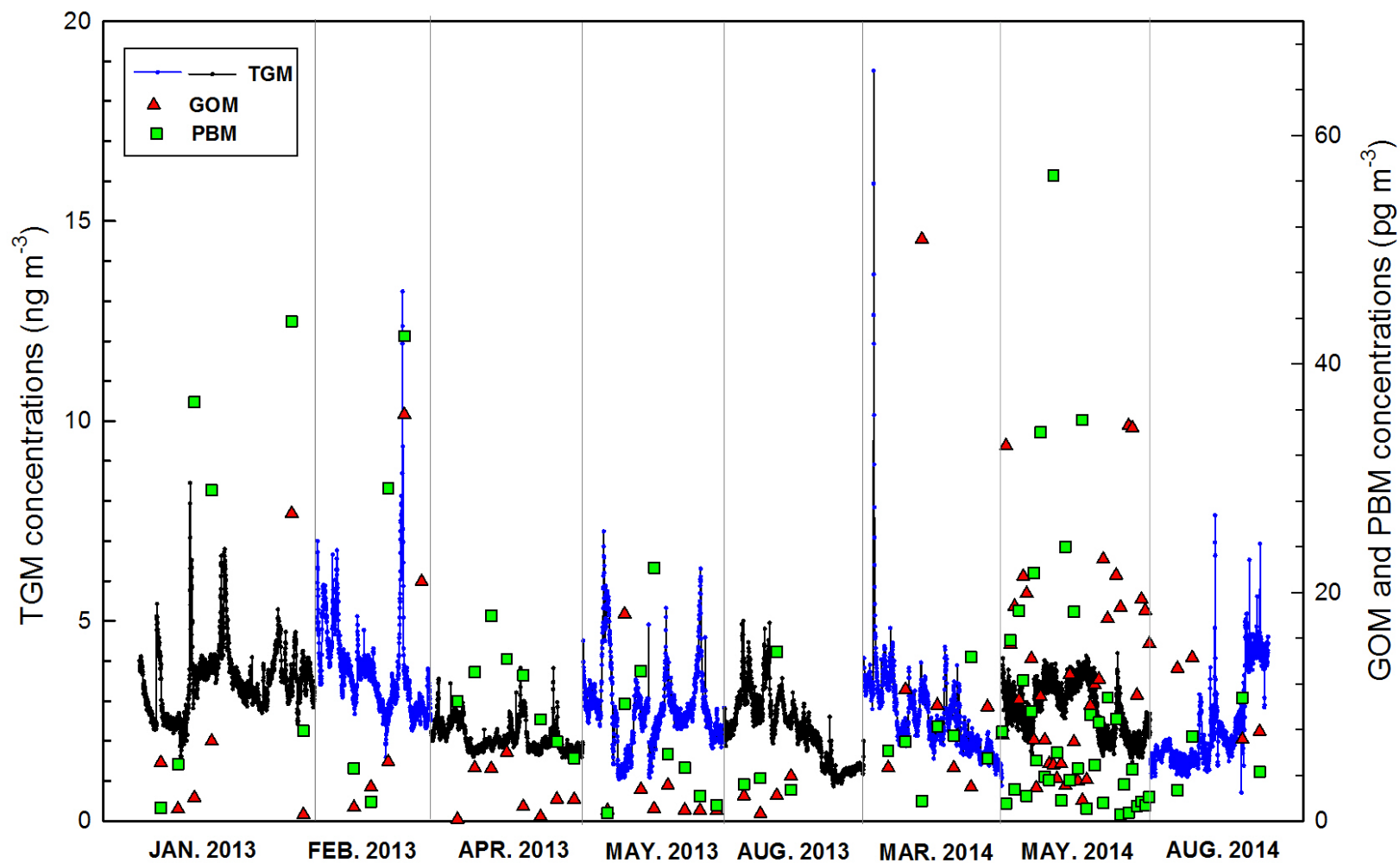


Figure 2. TGM, GOM, and PBM concentrations measured during the eight sampling periods. TGM was measured every 5 minutes while GOM and PBM were measured during 12hrs except for the 2hr measurements during May, 2014.



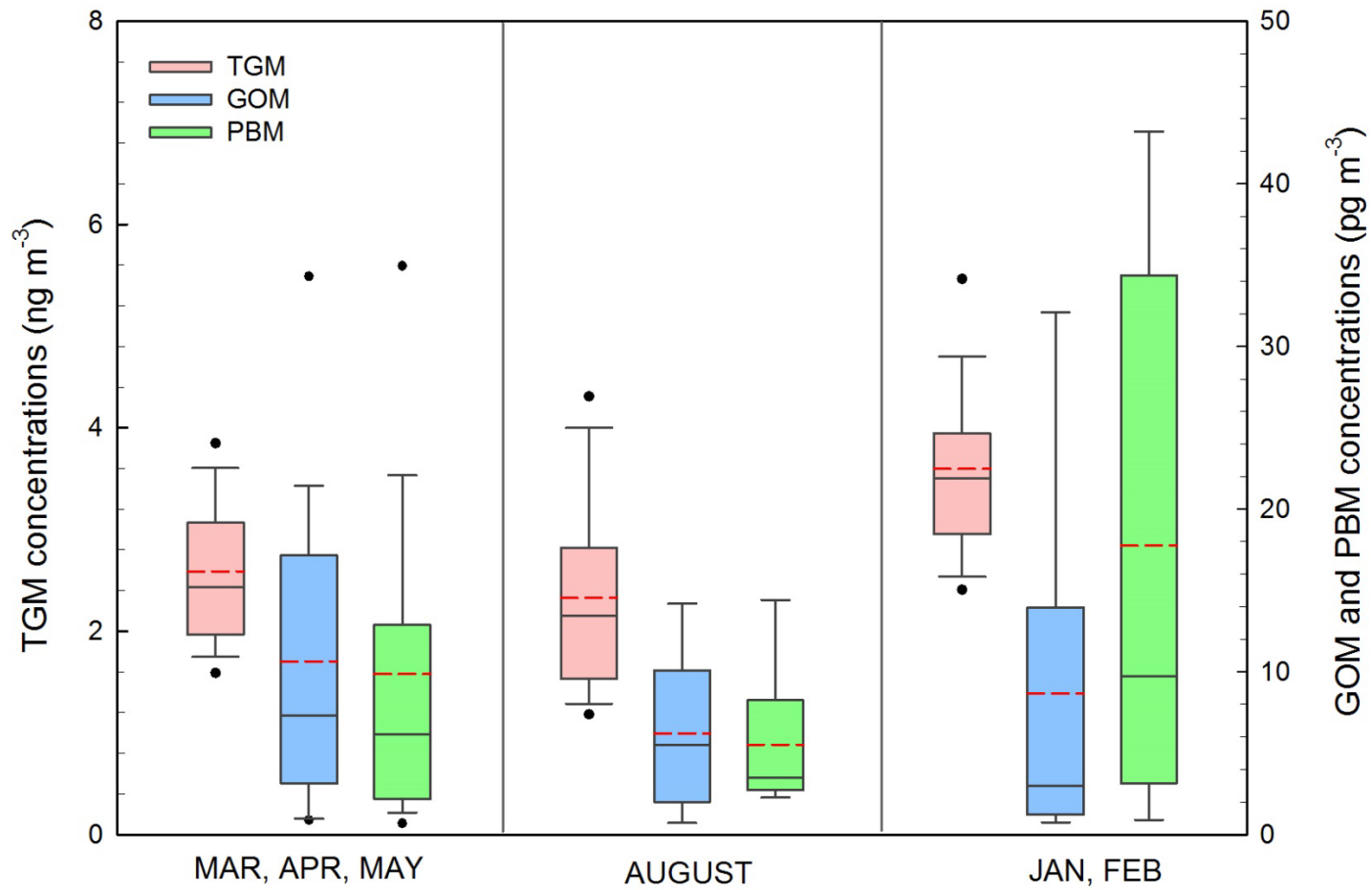


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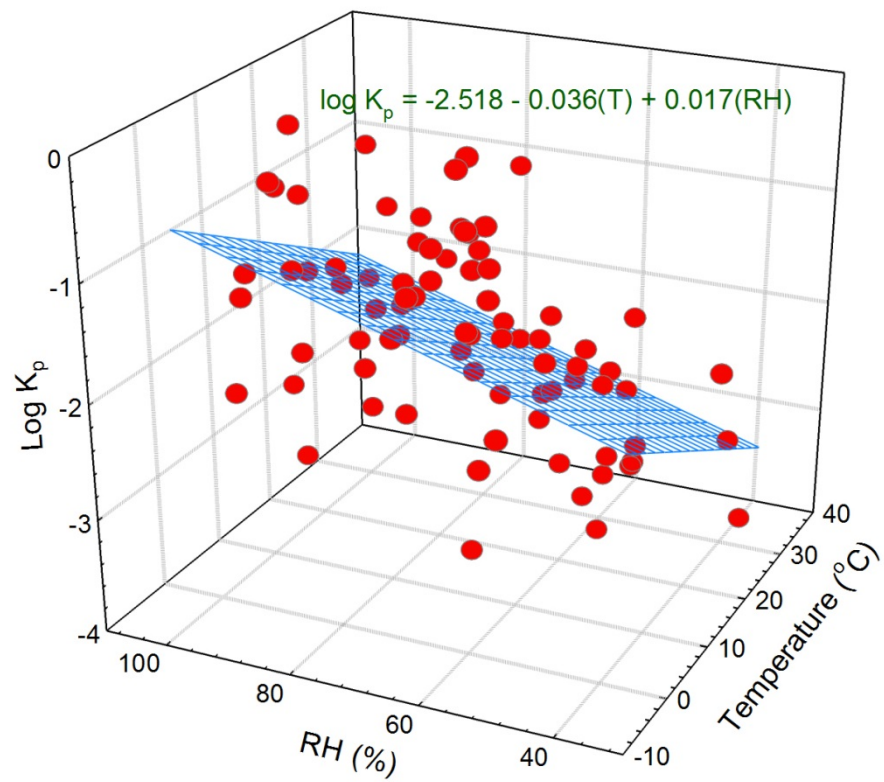


Figure 4. The gas-particle partitioning coefficient,  $K_p$ , related to atmospheric temperature and relative humidity (RH) (n=81).



Figure 5. CPF plots for TGM using the top 25% (left upper panel) and the top 10% (right upper panel) as a criterion, and for GOM (left bottom panel) and for PBM (right bottom panel). For both GOM and PBM, the criterion of the top 25% concentration was used.

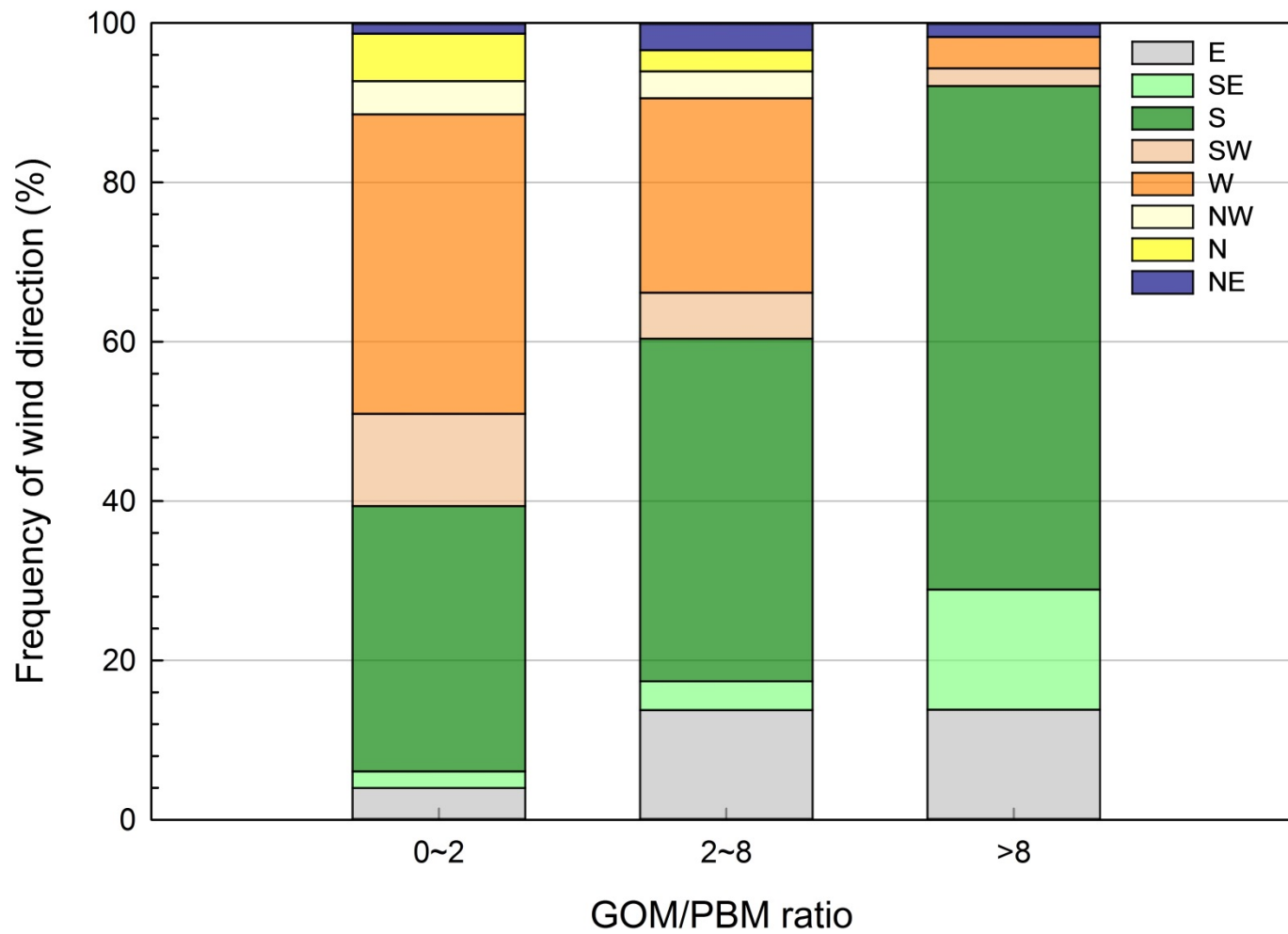


Figure 6. Frequency of wind direction with different GOM/PBM ratios. Southerly and easterly winds prevailed for the samples with high GOM/PBM ratio whereas the percentage of westerly winds increased as the GOM/PBM ratio decreased.

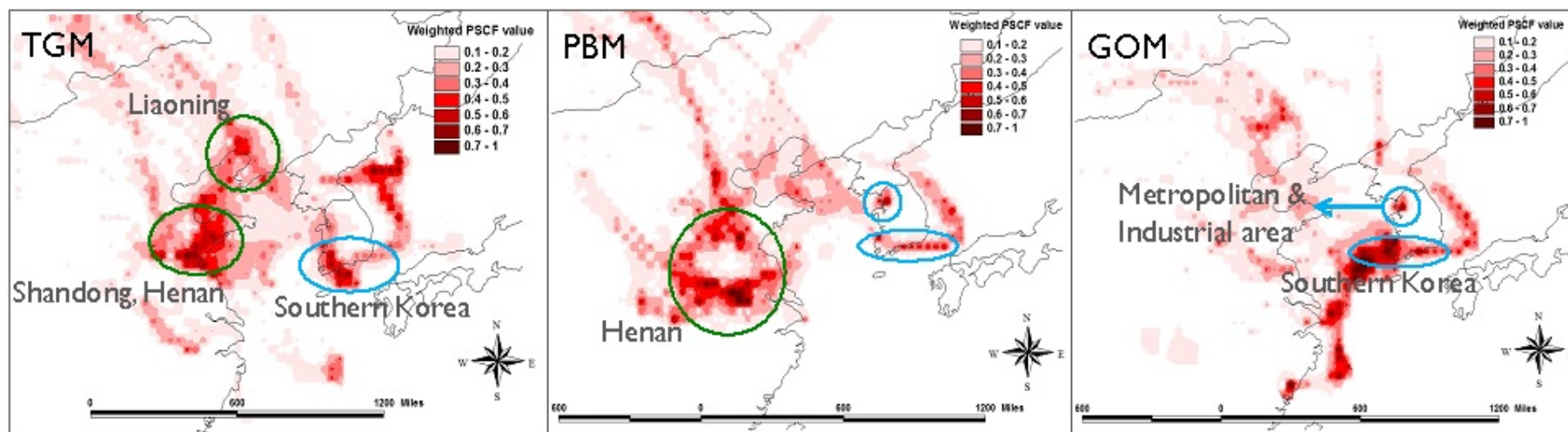


Figure 7. PSCF results for TGM (left), PBM (middle), and GOM (right) using the top 25% of concentrations as criteria.

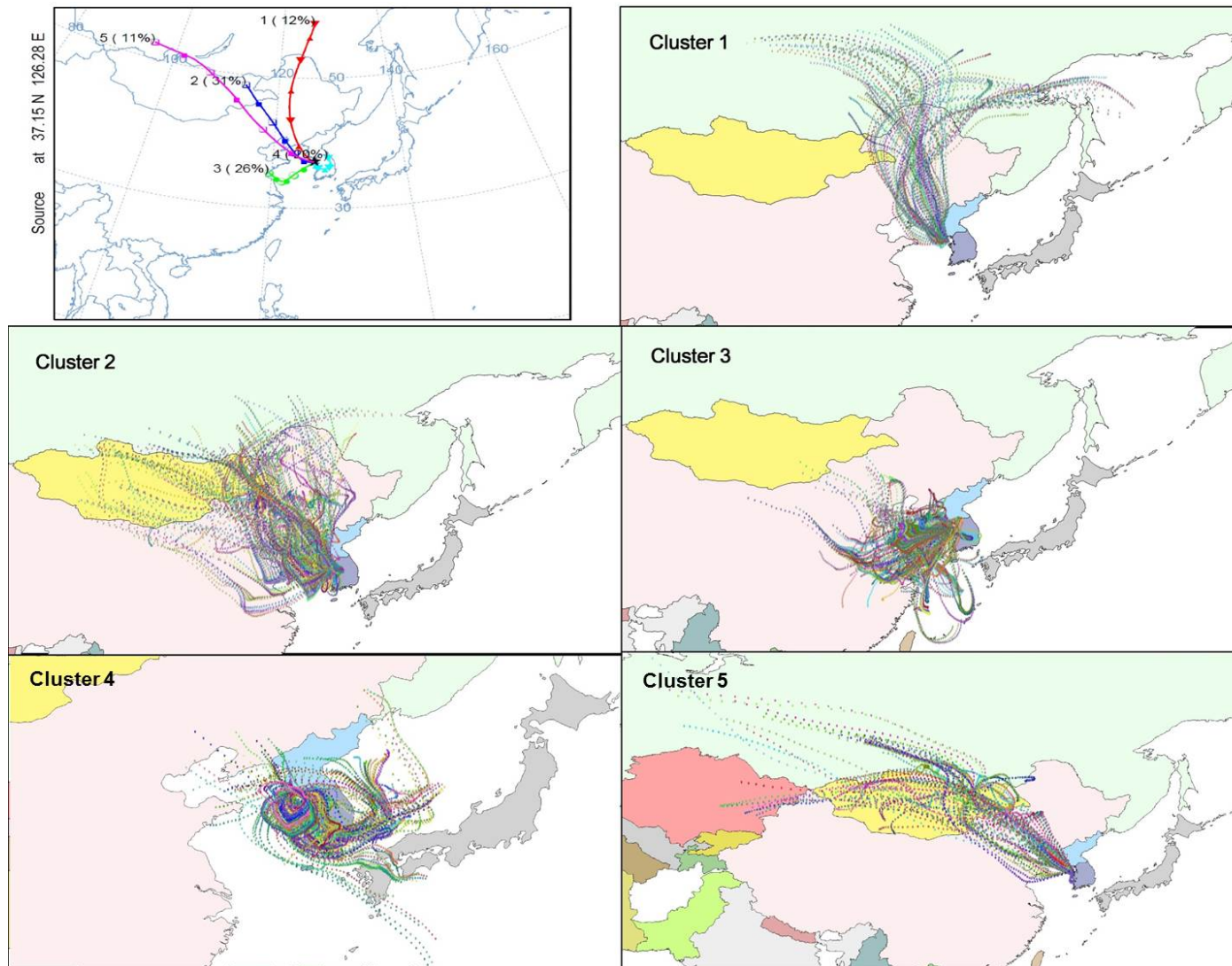


Figure 8. Back-trajectories for clusters 1 through 5. The left top panel indicates the mean back-trajectory and contribution for each cluster.