1 Atmospheric mercury deposition over the land

surfaces and the associated uncertainties in

observations and simulations: a critical review

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- 10 **Abstract.** One of the most important processes in the global mercury (Hg)
- biogeochemical cycling is the deposition of atmospheric Hg, including gaseous
- elemental mercury (GEM), gaseous oxidized mercury (GOM), and particulate-bound
- mercury (PBM), to the land surfaces. Results of wet, dry, and forest Hg deposition
- from global observation networks, individual monitoring studies, and observation-
- based simulations have been reviewed in this study. Uncertainties in the observation
- and simulation of global speciated atmospheric Hg deposition to the land surfaces
- 17 have been systemically estimated based on assessment of commonly used observation
- methods, campaign results for comparison of different methods, model evaluation
- with observation data, and sensitivity analysis for model parameterization. The
- 20 uncertainties of GOM and PBM dry deposition measurements come from the
- 21 interference of unwanted Hg forms or incomplete capture of targeted Hg forms, while
- that of GEM dry deposition observation originates from the lack of standardized
- 23 experimental system and operating procedure. The large biases in the measurements
- of GOM and PBM concentration and the high sensitivities of key parameters in
- resistance models lead to high uncertainties in GOM and PBM dry deposition
- simulation. Non-precipitation Hg wet deposition could play a crucial role in alpine
- and coastal regions, and its high uncertainties in both observation and simulation
- affect the overall uncertainties of Hg wet deposition. The overall uncertainties in the
- observation and simulation of the total global Hg deposition were estimated to be
- $\pm (25-50)$ % and $\pm (45-70)$ %, respectively, with the largest contributions from dry

- deposition. According to the results from uncertainty analysis, future research needs
- were recommended, among which global Hg dry deposition network, unified methods
- for GOM and PBM dry deposition measurements, quantitative methods for GOM
- speciation, campaigns for comprehensive forest Hg behavior, and more efforts on
- long-term Hg deposition monitoring in Asia are the top priorities.

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1 Introduction

- Mercury (Hg) is a global pollutant, characterized by its neurotoxicity, persistency and
- 39 bioaccumulation effect. It undergoes regional or global long-range transport via
- 40 atmospheric circulation, deposition to local or remote areas, methylation in
- ecosystems, and accumulation through food chain, posing high risks to human health
- and the environment (Obrist et al., 2018). Hg in the atmosphere has three major
- forms: gaseous elemental mercury (GEM), gaseous oxidized mercury (GOM), and
- particulate-bound mercury (PBM). The sum of GEM and GOM is called total gaseous
- 45 mercury (TGM), and the sum of GOM and PBM is also known as reactive mercury
- 46 (RM). GEM is the predominant form of atmospheric Hg (>90 %) with a long
- 47 residence time of several months to over one year due to its chemical inertness and
- low solubility. GOM used to be estimated to account for less than 1 % of atmospheric
- 49 Hg, which is easily scavenged by wet deposition, resulting in a short residence time of
- 50 hours to days (Schroeder and Munthe, 1998; Lindberg et al., 2007). However, recent
- studies (Lyman et al., 2010; Gustin et al., 2013; McClure et al., 2014; Gustin et al.,
- 52 2015) showed that there could be a significant underestimation of GOM due to the
- low capture efficiency of the KCl denuder method adopted by most observation sites
- in the presence of ozone and moisture. PBM (<10 % of atmospheric Hg) stays in the
- air for days to several weeks depending on particle size before scavenged by dry or
- wet deposition (Schroeder and Munthe, 1998; Lindberg et al., 2007; Ci et al., 2012;
- 57 Fu et al., 2012; Zhang et al., 2016a).
- 58 Deposition is one of the most important processes in global Hg cycling, leading to
- the sink of atmospheric Hg (Obrist et al., 2018). According to the Global Mercury
- Assessment 2018 (UN Environment, 2019), the annual Hg deposition to the land
- surfaces including freshwater is estimated to be 3600 t. Atmospheric Hg deposition
- can be broadly divided into wet and dry deposition. Hg wet deposition is mostly in the
- form of precipitation (rain, snow, etc.), with non-negligible contribution from non-

- precipitation forms (cloud, fog, dew, frost, etc.). Hg dry deposition is highly related to
- 65 the underlying surfaces, including forest canopies, grasslands, wetlands, agricultural
- 66 fields, deserts, background non-vegetated soils, contaminated sites, etc. (Zhang et al.,
- 67 2009). Forest canopy is regarded as an important sink of atmospheric Hg for its
- special forms of deposition, litterfall and throughfall (Gustin et al., 2008). Litterfall is
- a form of indirect Hg dry deposition through foliar uptake of atmospheric Hg, and
- throughfall includes wet-deposited Hg above the canopy and a portion of dry-
- deposited Hg washed off from the canopy (Wright et al., 2016). Hg deposition
- through litterfall has recently been drawn much attention to by the study of Wang et
- al. (2016a). The sum of litterfall and throughfall is regarded as the total Hg deposition
- 74 in forest canopies.
- Significant efforts have been made in the past decade on quantifying atmospheric
- 76 Hg deposition through both direct observations and model simulations, especially on
- dry deposition (Lyman et al., 2009; Zhang et al., 2009; Holmes et al., 2011; Lai et al.,
- 78 2011; Castro et al., 2012; Gustin et al., 2012; Peterson et al., 2012; L. Zhang et al.,
- 79 2012; Fang et al., 2013; Sather et al., 2013; Lynam et al., 2014; Sather et al., 2014;
- Huang and Gustin, 2015a; Weiss-Penzias et al., 2016a; Zhang et al., 2016b; Hall et al.,
- 2017; Sprovieri et al., 2017). Yet large uncertainties still exist due to limitations of the
- current methods for Hg deposition measurements and modeling (Gustin et al., 2015).
- The purpose of this paper is to give an overview of the uncertainties in the observation
- and simulation of global speciated atmospheric Hg deposition over the land surfaces.
- 85 In this paper, we investigated results from observations and simulations of global Hg
- deposition, reviewed methods adopted for Hg deposition measurements and modeling,
- 87 estimated the uncertainties of different methods for different Hg deposition forms, and
- summarized the overall uncertainty level of the global Hg deposition.

2 Observation-based estimation of global Hg deposition

90 2.1 Wet deposition

- 91 Precipitation is the major form of Hg wet deposition. There have been several
- observation networks of Hg wet deposition through precipitation. The Global Mercury
- Observation System (GMOS) is so far the only global scale network covering the
- northern hemisphere, the tropics, and the southern hemisphere (Sprovieri et al., 2017).
- The Mercury Deposition Network (MDN) of the National Atmospheric Deposition

- 96 Program (NADP) in North America is the earliest continental scale network
- specifically for Hg deposition (Prestbo and Gay, 2009; Weiss-Penzias et al., 2016a).
- 98 Hg wet deposition is also monitored in the European Monitoring and Evaluation
- 99 Programme (EMEP) for Europe (Tørseth et al., 2012; Bieser et al., 2014). A new
- Asia–Pacific Mercury Monitoring Network (APMMN) has recently been established
- 101 (Sheu et al., 2019).
- Sprovieri et al. (2017) reported a 5-year record (2011–2015) of Hg wet deposition
- at 17 selected GMOS monitoring sites, which provided a global baseline of the Hg
- wet deposition flux including regions in the southern hemisphere and tropical areas.
- The annual averages (multiple year ranges) of Hg wet deposition in the northern
- hemisphere, the tropics, and the southern hemisphere were 2.9 (0.2–6.7), 4.7 (2.4–
- 107 7.0), and 1.9 (0.3–3.3) μ g m⁻² yr⁻¹, respectively. The MDN network has a much
- longer history dating back to the 1990s. Weiss-Penzias et al. (2016a) analyzed records
- from 19 sites in the United States (U.S.) and Canada between 1997 and 2013, and
- discovered trends of Hg concentration in wet deposition, with the early time period
- 111 (1998–2007) producing a significantly negative trend ($-1.5\pm0.2~\%~yr^{-1}$) and the late
- time period (2008–2013) a flat slope (not significant). Therefore, the MDN data of
- 136 sites for the time period of 2008–2015 (http://nadp.slh.wisc.edu/mdn) were used
- in Figure 1 to represent the recent background Hg wet deposition level in North
- America. Fu et al. (2016a) summarized wet deposition measurements from 7
- monitoring sites in China. The annual Hg wet deposition fluxes at 6 rural sites were
- averagely 4.8 µg m⁻² yr⁻¹, while the annual flux at an urban site was as high as 12.6
- 118 $\mu g m^{-2} yr^{-1}$.
- Figure 1 summarizes the global distribution of the observed Hg wet deposition
- fluxes based on results from both these global or regional networks and individual
- studies. Overall, East Asia has the highest wet deposition flux (averagely 16.1 µg m⁻²
- 122 yr⁻¹), especially in the southern part of China where the RM concentration level is
- relatively high (Fu et al., 2008; Guo et al., 2008; Wang et al., 2009; Fu et al., 2010a;
- 2010b; Ahn et al., 2011; Huang et al., 2012b; Seo et al., 2012; Huang et al., 2013a;
- Sheu and Lin, 2013; Marumoto and Matsuyama, 2014; Xu et al., 2014; Zhu et al.,
- 2014; Huang et al., 2015; Zhao et al., 2015; Han et al., 2016; Fu et al., 2016a; Ma et
- al., 2016; Nguyen et al., 2016; Qin et al., 2016; Sommar et at., 2016; Cheng et al.,
- 2017; Travnikov et al., 2017; Chen et al., 2018; Lu and Liu, 2018). North America has

an average Hg wet deposition flux of 9.1 µg m⁻² yr⁻¹, and exhibits a descending 129 spatial profile from the southeastern part to the northwestern part, which is consistent 130 with the distribution of the atmospheric Hg concentration (L. Zhang et al., 2012; 131 Gichuki and Mason, 2014; Lynam et al., 2017). Europe has the lowest Hg wet 132 deposition level (averagely 3.4 μg m⁻² yr⁻¹) according to the available observation and 133 simulation data (Connan et al., 2013; Bieser et al., 2014; Siudek et al., 2016). 134 Observation data for the tropics and the southern hemisphere are scarce with large 135 uncertainties (Wetang'ula, 2011; Gichuki and Manson, 2013; Sprovieri et al., 2017). 136 The one exceptional tropical site with a wet deposition flux of 16.8 µg m⁻² yr⁻¹ is in 137 Kenya while the other sites in the tropics are all in Mexico (Wetang'ula, 2011; Hansen 138 and Gay, 2013). The two sites in the southern hemisphere with annual precipitation of 139 over 4000 mm are in Australia and have wet deposition fluxes of 29.1 and 18.2 $\mu g \ m^{-2}$ 140 yr⁻¹, respectively (Dutt et al., 2009). Seen from the bottom part of Figure 1, Hg wet 141 deposition flux is not significantly correlated with elevation. 142 Studies on non-precipitation Hg wet deposition (e.g., cloud, fog, dew, and frost) are 143 very limited so far. Fog or cloud Hg deposition is not yet considered in the global Hg 144 wet deposition observation network. However, studies (Stankwitz et al., 2012; Weiss-145 Penzias et al., 2016b; Gerson et al., 2017) have shown that cloud and fog water have 146 higher Hg concentration than rain water in the same region, and cloud and fog could 147 have a remarkable contribution to Hg wet deposition in high-elevation forests and 148 near-water surfaces. Stankwitz et al. (2012) and Gerson et al. (2017) found the 149 average cloud Hg deposition fluxes of two North American montane forests to be 7.4 150 and 4.3 µg m⁻² during the research periods, respectively, equivalent to rainfall Hg 151 deposition. In California coastline, fog Hg deposition, with only 2 % volume 152 153 proportion, accounts for 13 % of the total wet deposition (Weiss-Penzias et al., 2016b). Converse et al. (2014) found the annual dew and frost Hg deposition at a 154 high-elevation meadow in the U.S. to be about 0.12 μg m⁻² yr⁻¹, 2–3 orders of 155 magnitude smaller than wet deposition through precipitation. More standardized 156 methods are in urgent need for non-precipitation Hg wet deposition measurements. 157 2.2 **Dry deposition** 158 Figure 2 shows the global distribution of the GOM, PBM and GEM dry deposition 159 fluxes from observation-based estimation, either direct observation of dry deposition 160

or simulation based on Hg concentration observation. The global Hg dry deposition

network is very immature compared to the wet deposition network due to the 162 inconsistency in methods for estimation. GOM dry deposition fluxes were either 163 measured by the surrogate surface methods or simulated based on GOM concentration 164 measurements. PBM dry deposition fluxes were mainly estimated from the 165 measurements of total or size-resolved PBM concentrations. GEM dry deposition 166 fluxes were measured by different types of methods, including the surrogate surface 167 methods, the enclosure methods, and the micrometeorological methods. 168 Wright et al. (2016) presented an overview of GOM and PBM dry deposition. In 169 170 their work, the observation or simulation years for nearly one third of the reviewed studies were earlier than 2005, and only studies conducted in North America and Asia 171 were summarized. Therefore, this study included more studies carried out in recent 172 years and limited the observation or simulation year to be no earlier than 2005. Also, 173 studies in Europe and China were summarized in this study. As shown in Figure 2, 174 most studies on GOM dry deposition were conducted in North America and Europe, 175 among which direct observations of GOM dry deposition are mainly from North 176 America (Lyman et al., 2007; Lyman et al., 2009; Weiss-Penzias et al., 2011; Lombard 177 et al., 2011; Castro et al., 2012; Gustin et al., 2012; Peterson et al., 2012; L. Zhang et 178 179 al., 2012; Sather et al., 2013; Bieser et al., 2014; Sather et al., 2014; Wright et al., 2014; Huang and Guatin, 2015a; Enrico et al., 2016; Han et al., 2016; Zhang et al., 180 181 2016b; Huang et al., 2017). Regardless of the estimating methods, the average GOM dry deposition flux in North America (6.4 μg m⁻² yr⁻¹) is higher than in Europe (3.0 182 $\mu g \ m^{-2} \ yr^{-1}$). There have been very few studies on GOM dry deposition in Asia. A 183 significant correlation (R²=0.532, p<0.01) was found between the elevation and the 184 GOM dry deposition flux (see Figure 3), which could be due to higher GOM 185 concentrations at higher elevation and stronger atmospheric turbulence (Huang and 186 Gustin, 2015a). Nevertheless, significant discrepancies were found between the GOM 187 dry deposition fluxes from direct observations and from model simulations based on 188 measurements of GOM concentrations (see Figure 4). Results from size-resolved 189 PBM analysis and PBM dry deposition models show that East Asia has a much higher 190 average of PBM dry deposition flux (45.3 µg m⁻² yr⁻¹) than North America (1.1 µg 191 m⁻² yr⁻¹) (Fang et al., 2012a; Fang et al., 2012b; Zhu et al., 2014; Zhang et al., 2015; 192 Huang et al., 2016; Guo et al., 2017). 193 Zhu et al. (2016) reviewed the air-surface exchange of GEM. The observation years 194

195	for most of the reviewed studies were earlier than 2005. Since GEM concentrations
196	decreased significantly from early 1990s to 2005 in most regions in the world (Y.
197	Zhang et al., 2016), this study included more recent studies and limited the
198	observation or simulation year to be no earlier than 2005. The average GEM dry
199	deposition is lower in Europe (4.3±8.1 µg m ⁻² yr ⁻¹) while higher in North America
200	with more variation (5.2 \pm 15.5 μg m ⁻² yr ⁻¹) (Castelle et al., 2009; Baya and Heyst,
201	2010; Converse et al., 2010; Miller et al., 2011). The four Asian sites using all show
202	negative values, indicating the role of East Asia as a net emission source rather than a
203	net deposition sink (Luo et al., 2016; Ci et al., 2016; Yu et al., 2018). However, the
204	GEM dry deposition observations in Asia are still very limited.
205	Hg dry deposition is highly related to the underlying surfaces. Figure 5 exhibits the
206	dry deposition fluxes of GOM, PBM and GEM for different terrestrial surface types.
207	As shown in Figure 5a, high GOM dry deposition levels were found for grasslands
208	(mainly alpine meadows) and savannas. This is probably because of the enhanced Hg
209	oxidation process at high elevations with more halogen free radicals or more intensive
210	solar radiations (Huang and Gustin, 2015a). Urban areas also have high GOM dry
211	deposition fluxes due to high GOM concentrations. The low GOM dry deposition
212	fluxes on moist surfaces (near-water surfaces and croplands) might be partially
213	because of fog and dew scavenging (Malcolm and Keeler. 2002; Zhang et al., 2009).
214	The PBM dry deposition flux is high on surfaces with high human activities (urban
215	areas and croplands) and low in vegetative areas, implying the heavier PM pollution
216	in urban and rural areas than in remote areas (Figure 5b). Short-term observation of
217	GEM dry deposition shows high fluctuation. Therefore, we summarized model
218	estimations and one annual observation dataset (L. Zhang et al., 2012; Bieser et al.,
219	2014; Zhang et al., 2016b; Enrico et al., 2016), and found that the GEM dry
220	deposition does not only depend on GEM concentration, but also on the air-soil Hg
221	exchange compensation point (Luo et al., 2016). Regarding the annual air-surface Hg
222	exchange, instead of an important natural source, forests tend to be a net sink of
223	atmospheric Hg (Figure 5c).
224	2.3 Forest deposition

Hg deposition in forests is mainly in the forms of litterfall and throughfall. Wright et al. (2016) also made an extensive review of litterfall and throughfall Hg deposition. Wang et al. (2016a) made a comprehensive assessment of the global Hg deposition

- 228 through litterfall, and found litterfall Hg deposition an important input to terrestrial
- forest ecosystems (1180±710 Mg yr⁻¹). Not many new studies on forest Hg deposition
- have been reported since then. Therefore, here we only briefly introduce the spatial
- distribution of forest Hg deposition. South America was estimated to bear the highest
- 232 litterfall Hg deposition (65.8±57.5 μg m⁻² yr⁻¹) around the world (Teixeira et al.,
- 2012; Buch et al., 2015; Fostier et al., 2015; Teixeira et al., 2017; Fragoso et al., 2018;
- Shen et al., 2019). There have been numerous forest Hg deposition studies in the
- recent decade in East Asia with the second highest average litterfall Hg deposition
- 236 flux $(35.5\pm27.7 \,\mu g \,m^{-2} \,yr^{-1})$ (Wan et al., 2009; Wang et al., 2009; Fu et al., 2010a; Fu
- et al., 2010b; Gong et al., 2014; Luo et al., 2016; Ma et al., 2015; Han et al., 2016; Fu
- et al., 2016a; Ma et al., 2016; Wang et al., 2016b; Zhou et al., 2016; Zhou et al.,
- 239 2017). Lower levels of litterfall Hg deposition fluxes were found in North America
- 240 $(12.3\pm4.9 \,\mu\text{g m}^{-2} \,\text{yr}^{-1})$ and Europe $(14.4\pm5.8 \,\mu\text{g m}^{-2} \,\text{yr}^{-1})$ (Larssen et al., 2008; Obrist
- et al., 2009; Fisher and Wolfe, 2012; Juillerat et al., 2012; Obrist et al., 2012; Risch et
- al., 2012; Benoit et al., 2013; Navrátil et al., 2014; Gerson et al., 2017; Risch et al.,
- 2017; Risch and Kenski, 2018). Throughfall Hg deposition is another important way
- for Hg input in forests, Wright et al. (2016) summarized previous studies and reported
- the median throughfall Hg deposition to be 49.0, 16.3 and 7.0 μ g m⁻² yr⁻¹ in Asia,
- Europe and North America, respectively. Large discrepancies in Asian co-located
- comparisons between rainfall and throughfall Hg depositions (32.9±18.9 and 13.3±8.6
- 248 μg m⁻² yr⁻¹, respectively) could indicate a high dry deposition level in Asian forests
- 249 (Wan et al., 2009; Wang et al., 2009; Fu et al., 2010a; Fu et al., 2010b; Luo et al.,
- 2016; Ma et al., 2015; Han et al., 2016; Fu et al., 2016a; Ma et al., 2016; Wang et al.,
- 251 2016b; Zhou et al., 2016).

252 3 Uncertainties in Hg deposition observation

3.1 Uncertainties in the measurements of Hg wet deposition

3.1.1 Measurements of Hg wet deposition through precipitation

- 255 Hg wet deposition through precipitation, mostly rainfall, is easier to measure than dry
- deposition and usually more reliable. The rainfall Hg wet deposition flux is calculated
- 257 as follows (Zhao et al., 2018):

$$F_{\text{wet,rainfall}} = \sum_{i=1}^{n} C_i \cdot D_i$$
 (1)

where $F_{\text{wet,rainfall}}$ is the total rainfall Hg wet deposition flux; n is the number of precipitation events during a certain period; C_i is the total Hg concentration in rainwater during Event i; and D_i is the precipitation depth of Event i. As shown in Eq. (1), the overall uncertainty in rainfall Hg wet deposition originates from both the analytical methods of Hg concentration in rainwater and the measurements of precipitation depth.

Both manual and automatic precipitation sample collectors were used in previous 265 studies (Fu et al., 2010a; Gratz and Keeler, 2011; Marumoto and Matsuyama, 2014; 266 267 Zhu et al., 2014; Brunke et al., 2016; Chen et al., 2018). Automatic precipitation sample collectors cover the lid automatically when it is not raining to prevent 268 potential contamination, while manual collectors require manually placing collectors 269 before precipitation events and retrieving them after events. The measurements of 270 precipitation volume by samplers have non-negligible uncertainties (Wetherbee, 271 2017). The relative standard deviations (RSDs) of daily and annual precipitation depth 272 measurements in MDN were estimated to be 15 % and 10 %, respectively (Wetherbee 273 et al., 2005). The event-based sampling volume biases of two types of samplers used 274

276 The total Hg concentration in rainwater samples is usually analyzed by oxidation, purge and trap, and cold vapor atomic fluorescence spectrometry (CVAFS) following 277 278 EPA Method 1631. GMOS reported the ongoing precision recovery (OPR) for every 12 samples to be generally within 93–109 % (Sprovieri et al., 2017). The relative 279 280 percentage difference (RPD) for MDN precipitation Hg analysis is generally within 10 % according to inter-laboratory comparisons (Wetherbee and Martin, 2018). For 281 282 individual studies (Fu et al., 2010a; Huang et al., 2015; Zhao et al., 2018), the RSD is also generally less than 10 %. 283

in APMMN were estimated to be up to 11–18 % (Sheu et al., 2019).

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The overall relative uncertainty of the precipitation Hg wet deposition flux was calculated to be approximately $\pm (15-20)$ % using the following equation:

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$$\delta_F(\text{wet}) = \frac{U_F(\text{wet})}{F_{\text{wet}}} = \sqrt{\left(\frac{U_C}{C}\right)^2 + \left(\frac{U_D}{D}\right)^2} = \sqrt{\delta_C^2 + \delta_D^2}$$
 (2)

where δ_F (wet) and U_F (wet) are the relative and absolute uncertainties of Hg wet deposition flux, respectively; δ_C and U_C are the relative and absolute uncertainties of the total Hg concentration in precipitation water, respectively; and δ_D and U_D are the relative and absolute uncertainties of the precipitation depth, respectively.

3.1.2 Measurements of Hg wet deposition through cloud, fog, dew and frost

- Non-precipitation Hg wet deposition, e.g., cloud, fog, dew and frost, could account for
- a notable proportion of the total wet deposition in montane, coastal, arid, and semi-
- arid areas (Lawson et al., 2003; Sheu and Lin, 2011; Stankwitz et al., 2012; Blackwell
- and Driscoll, 2015b). Quantifying Hg in cloud or fog helps better understand the
- impact of long-range transport and local sources on global Hg cycling (Malcolm et al.,
- 297 2003). The non-precipitation Hg deposition flux is calculated as follows:

$$F_{\text{wet,non-precipitation}} = \sum_{j=1}^{m} C_j \cdot D_j$$
 (3)

- where $F_{\text{wet,non-precipitation}}$ is the non-precipitation Hg deposition flux; m is the number
- of non-precipitation wet deposition events during a certain period; C_i is the total Hg
- concentration in non-precipitation wet deposition water during Event j; and D_j is the
- non-precipitation wet deposition depth of Event *j*.

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- Both active and passive collectors have been used to collect cloud or fog water
- 304 (Lawson et al., 2003; Malcolm et al., 2003; Kim et al., 2006; Sheu and Lin, 2011;
- Schwab et al., 2016; Weiss-Penzias et al., 2018). The major uncertainty lies in the
- deposition depth. The deposition depth of cloud, fog, dew or frost is usually modeled
- based on meteorology (Converse et al., 2014; Katata, 2014). The fog deposition depth
- can be measured by standard fog collectors (SFC). The uncertainty of fog deposition
- depth measurements is mainly from the collecting efficiency of SFC depending on the
- wind speed, wind direction, or mesh types (Weiss-Penzias et al., 2016b; Fernandez et
- al., 2018). Montecinos et al. (2018) evaluated the collection efficiency of SFC to be
- 312 up to 37 %. Therefore, there is extremely large uncertainty in the measurements of the
- fog deposition depth. Based on the fog deposition studies (Weiss-Penzias et al.,
- 2016b; Fernandez et al., 2018; Montecinos et al., 2018), the overall uncertainty of
- non-precipitation Hg deposition flux observation is estimated to be $\pm (200-300)$ %.
- Note that the true uncertainty range is not symmetric about the mean because some of
- the underlying variables are lognormally distributed (Streets et al., 2005). A better
- interpretation of " \pm (200–300) %" might be "within a factor of 3–4".

3.2 Uncertainties in the measurements of Hg dry deposition

- Direct measurements of the Hg dry deposition flux is technically challenging, large
- uncertainties still exist in quantify Hg dry deposition accurately (Wright et al., 2016).
- Three major categories of methods for direct Hg dry deposition measurements are the

surrogate surface methods, the enclosure methods, and the micrometeorological methods (Zhang et al., 2009; Huang et al., 2014).

3.2.1 Measurements of RM (GOM and PBM) dry deposition

Most of the RM dry deposition measurements used the surrogate surface methods
(Huang et al., 2014; Wright et al., 2016). The micrometeorological methods and the
enclosure methods were also adopted in some studies (Poissant et al., 2004; Zhang et
al., 2005; Skov et al., 2006), but not widely used due to the high uncertainties in the
measurements of GOM and PBM concentrations using the Tekran system. For the
surrogate surface methods, the RM dry deposition flux is determined using the
following equation (Huang et al., 2014):

$$F_{\rm dry,SS} = \frac{M}{A \cdot t} \tag{4}$$

where $F_{\text{dry,SS}}$ is the Hg dry deposition flux using the surrogate surface methods; M is the total Hg amount collected on the material during the sampling period; A is the surface area of the collection material; and t is the exposure time.

Different surrogate surfaces were used to measure different RM forms. Mounts with cation-exchange membranes (CEMs) are widely used for GOM dry deposition measurements (Lyman et al., 2007; Lyman et al., 2009; Castro et al., 2012; Huang et al., 2012a; Peterson et al., 2012; Sather et al., 2013). The down-facing aerodynamic mount with CEM is considered to be the most reliable deployment for GOM dry deposition measurements so far (Lyman et al., 2009; Huang et al., 2014). Knife-edge surrogate surface (KSS) samplers with quartz fiber filter (QFFs) and dry deposition plates (DDPs) with overhead projection films were deployed for PBM dry deposition measurements (Lai et al., 2011; Fang et al., 2012b; Fang et al., 2013). However, these samplers are not well verified to reflect the deposition velocity of PBM, and hence not widely accepted. KCl-coated QFFs were used to measure the total RM (GOM+PBM) dry deposition, but failed to capture GOM efficiently (Lyman et al., 2009; Lai et al., 2011).

The uncertainties of RM dry deposition mainly come from the capture efficiency of sampling surface, the turbulent condition near the surface, and the analysis of the membrane. CEMs exhibited a GOM capture rate of 51–107 % in an active sampling system (Huang and Gustin, 2015b). The CEM mounts designed to measure only GOM dry deposition capture part of fine PBM (Lyman et al., 2009; Huang et al.,

2014), while the KSS samplers with QFFs designed to measure only PBM dry 355 356 deposition may also collect part of GOM (Rutter and Schauer, 2007; Gustin et al., 2015). Based on the RM concentration measurements and the surrogate surface 357 method evaluations, the GOM concentration related uncertainty is estimated to be 358 ±50 % (Lyman et al., 2009; Lyman et al., 2010; Gustin et al., 2012; Fang et al., 2013; 359 Zhang et al., 2013; Huang et al., 2014). The design of the sampler (e.g., the sampler 360 orientation, the shape of the sampler, variation in turbulence, low surface resistances, 361 passivation, etc.) leads to the surface capture efficiency related uncertainty which is 362 363 about ±50 % for GOM (Lyman et al., 2009; Lai et al., 2011; Huang et al., 2012a). The overall uncertainty in surface capture efficiency could decline to about ± 30 % at 364 annual level. Calculating based on the method described by Eq. (2), the overall 365 uncertainty of GOM dry deposition observation is $\pm (60-70)$ %. There is not enough 366 information to quantify the overall uncertainty of PBM dry deposition observation in 367 a similar way. However, its uncertainty is usually considered to be higher than that of 368 GOM dry deposition measurements. Based on the distribution of daily samples in the 369 study of Fang et al. (2012b), the overall uncertainty of PBM dry deposition 370 measurements is assumed to be $\pm (80-100)$ %. 371 3.2.2 Measurements of GEM dry deposition 372 373 GEM has a low dry deposition velocity due to its mild activity, high volatility and low water solubility, and deposited GEM could re-emit into the atmosphere (Bullock et al., 374 2008; Fu et al., 2016b). Various methods have been applied to studies on air-surface 375 GEM exchange, among which the enclosure methods and the micrometeorological 376 methods were most commonly used (Zhang et al., 2009; Agnan et al., 2016; Zhu et al., 377 2016; Yu et al., 2018). Both Agnan et al. (2016) and Zhu et al. (2016) have presented 378 comprehensive reviews on air–surface GEM exchange and introduced the two types 379 of methods for measurements. The uncertainty of air-surface GEM exchange flux 380 381 using the micrometeorological methods were estimated to be up to ± 30 % (Meyers et al., 1996; Lindberg and Meyers, 2001; Fritsche et al., 2008; Sommer et al., 2013a; 382 Zhu et al., 2015b). However, Zhu et al. (2016) summarized existing air–surface GEM 383 384 exchange studies and found that the mean flux using micrometeorological methods is higher than using DFCs by a factor of 2. Agnan et al. (2016) found the uncertainty of 385 GEM flux to be in the range of -180 % to +120 %. Therefore, the overall uncertainty 386 of GEM dry deposition observation is estimated to be $\pm (100-200)$ %. 387

3.3 Uncertainties in the measurements of Hg deposition in forests

- In forest ecosystems, the presence of canopy changes the form of Hg deposition. The
- sum of litterfall and throughfall is more commonly used to represent the total Hg
- deposition in forests (Wang et al., 2016a; Wright et al., 2016).

3.3.1 Litterfall Hg deposition measurements

- Litterfall Hg deposition includes the dry and wet deposited Hg on leaves and bark as
- well as the captured Hg emitted from the soil (Blackwell and Driscoll, 2015a; Wright
- et al., 2016). Litterfall Hg deposition flux is calculated as follows (Fisher and Wolfe,
- 396 2012):

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$$F_{\text{litterfall}} = \frac{E_A \cdot C_l \cdot M_l}{A \cdot t} \tag{5}$$

- where $F_{\text{litterfall}}$ is the litterfall Hg deposition flux; E_A is the litterfall trap area
- expansion factor (note: leaves outside the area above the trap could fall into the trap
- due to horizontal air fluctuation); C_l is the Hg mass concentration in litterfall; M_l is the
- 401 total dry weight of litterfall; A is the litterfall trap area; and t is the sampling time.
- Litterfall samples are collected during the leaf-growing or -falling seasons with
- litterfall traps or collectors (Fisher and Wolfe, 2012). Total litterfall consists of leaves
- and needles, woody material such as twigs and bark, and reproductive bodies such as
- flowers, seeds, fruits, and nuts (Meier et al., 2006; Risch et al., 2012). The total litter
- 406 mass collected by different samplers could cause a RSD of 16 % (Risch et al., 2012)
- and Risch et al., 2017). The Hg content in litterfall can be determined by thermal
- decomposition, amalgamation, and cold vapor atomic absorption spectrophotometry
- 409 (CVAAS) following EPA Method 7473 (Richardson and Friedland, 2015; Fu et al.,
- 2016a; Zhou et al., 2017; Risch et al., 2017). Alternatively, the litterfall samples can
- be digested and analyzed following EPA Method 1631E (Fu et al., 2010a; Fisher and
- Wolfe, 2012). The uncertainty in litterfall Hg content analysis is about $\pm 7 \%$
- according to the Litterfall Mercury Monitoring Network developed by NADP (Risch
- et al., 2017) and individual studies (Benoit et al., 2013; Ma et al., 2015; Zhou et al.,
- 415 2016; Gerson et al., 2017).
- Therefore, the event-based uncertainty of litterfall Hg deposition observation is
- estimated to be ± 18 % based on Eq. (2). The Litterfall Mercury Monitoring Network
- and many other studies only collect litterfall during the falling season each year,
- which will cause some underestimation. Moreover, based on the assumption that the

- 420 total Hg concentration in litterfall is linearly accumulated during the growing season,
- some studies estimated litterfall Hg concentration by multiplying a scale factor, which
- may cause extra uncertainty (Bushey et al., 2008; Poissant et al., 2008; Fu et al.,
- 423 2010a; Gong et al., 2014). Taking this into consideration, the overall uncertainty of
- litterfall Hg deposition observation is estimated to be $\pm (20-30)$ %.

425 3.3.2 Throughfall Hg deposition measurements

- Throughfall Hg deposition includes the wet-deposited Hg passing through the canopy
- and a portion of dry-deposited Hg washed off from the canopy (Blackwell and
- Driscoll, 2015a; Wright et al., 2016). Throughfall Hg deposition flux is calculated as
- follows (Fisher and Wolfe, 2012):

$$F_{\text{throughfall}} = \frac{E_A \cdot C_t \cdot V_t}{A \cdot t} \tag{6}$$

- where $F_{\text{throughfall}}$ is the throughfall Hg deposition flux; E_A is the throughfall funnel
- area expansion factor; C_t is the Hg mass concentration in throughfall; V_t is the total
- volume of throughfall; *A* is the throughfall funnel area; and *t* is the sampling time.
- Throughfall under canopy is usually collected using a passive bulk throughfall
- collector with a funnel connected a bottle for water storage (Wang et al., 2009; Fisher
- and Wolfe, 2012; Åkerblom et al., 2015) or collected as open-field rain collection if
- the environmental condition permits (Choi et al., 2008; Fu et al., 2010a; Fu et al.,
- 438 2010b; Han et al., 2016). Attention should be paid to potential litterfall contamination
- and cloud or fog deposition influence at high elevation sites if the collector is not
- sheathed (Fisher and Wolfe, 2012; Wright et al., 2016). Throughfall samples are
- usually analyzed following EPA Method 1631E (Fisher and Wolfe, 2012). Therefore,
- throughfall Hg deposition should have a similar uncertainty as rainfall Hg deposition.
- Considering the possible interference for throughfall sample collection, the overall
- uncertainty of throughfall Hg deposition observation is estimated as $\pm (20-30)$ %.

4 Uncertainties in Hg deposition simulation

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4.1 Uncertainties in models for Hg wet deposition

4.1.1 Model for precipitation Hg wet deposition

- 448 Hg wet deposition through precipitation is an important process in global or regional
- chemical transport models (CTMs), such as GEOS-Chem and CMAQ-Hg (Lin et al.,
- 450 2010; Y. Zhang et al., 2012; Bieser et al., 2014; J. Zhu et al., 2015; Horowitz et al.,

2017). As shown in Eq. (1), precipitation Hg wet deposition is the product of the total 451 Hg concentration in rainwater and the precipitation depth. In CTMs, the precipitation 452 Hg concentration contains more uncertain factors. Hg in rainwater originates from the 453 scavenging of GOM and PBM in both free troposphere and boundary layer. Based on 454 previous modeling work for Hg wet deposition in the United States using GEOS-455 Chem (Selin and Jacob, 2008), GOM and PBM contributed 89 % and 11 % to the total 456 Hg wet deposition, respectively, and 60% of the GOM induced wet deposition 457 originated from scavenging in the free troposphere. Seo et al. (2012) and Cheng et al. 458 459 (2015) also reported higher scavenging coefficient for GOM than for PBM. Therefore, Hg redox chemistry in the free troposphere, aqueous phase Hg speciation, aqueous 460 phase sorption, and the scavenging process tend to be the dominant sources of 461 uncertainties (Lin et al., 2006; Lin et al., 2007; Cheng et al., 2015). 462 In the simulation of Hg wet deposition by the GEOS-Chem model, the uncertainty 463 of precipitation depth is usually within ± 10 % because it is based on assimilated 464 meteorological observations from the Goddard Earth Observing System (GEOS) 465 instead of meteorological models (Y. Zhang et al., 2012). Y. Zhang et al. (2012) 466 conducted a nested-grid simulation of Hg over North America using GEOS-Chem, 467 468 and reported the normalized bias of the annual Hg wet deposition flux to be ranging from -14% to +27% comparing to the MDN observations. Horowitz et al. (2017) 469 470 used GEOS-Chem to reproduce observed Hg wet deposition fluxes over North America, Europe, and China and also got low bias (0–30 %). The CMAQ-Hg model 471 exhibits a higher uncertainty level because the precipitation depth is simulated by 472 meteorological models (e.g., MM5 or WRF) and its uncertainty has a strong impact 473 on model prediction on Hg wet deposition (Lin et al., 2006). In the study of Bullock et 474 al. (2009), the precipitation simulated by MM5 was averagely 12% greater than 475 observed and the CMAQ simulation of Hg wet deposition was averagely about 15% 476 above the MDN observations. However, different boundary conditions could cause a 477 25% difference (Bullock et al., 2009). Holloway et al. (2012) found that the CMAQ-478 Hg model underestimated wet deposition by 21 % on an annual basis and showed 479 average errors of 55 %. Based on the comparison between observed and modeled 480 results and the sensitivity of key parameters, the overall uncertainty of precipitation 481 Hg wet deposition simulation is estimated to be $\pm (30-50)$ % depending on the adopted 482 models. 483

4.1.2 Model for non-precipitation Hg wet deposition

- Non-precipitation Hg wet deposition simulation has never been considered in CTMs,
- but performed in some individual studies with Hg concentration data for cloud, fog,
- dew or frost samples (Ritchie et al., 2006; Converse et al., 2014; Blackwell and
- Driscoll, 2015b). Non-precipitation deposition depth can be estimated using resistance
- 489 models, analytical models or sophisticated atmosphere-soil-vegetation models. Katata
- 490 (2014) reviewed different types of models for fog deposition estimation, and found
- 491 the four most sensitive factors to be canopy homogeneity, droplet size spectra, droplet
- capture efficiency, and canopy structure. Since fog is the most important form of non-
- 493 precipitation deposition, the overall uncertainty in the simulation of non-precipitation
- 494 Hg wet deposition is estimated to be $\pm (200-300)$ % or a factor of 3–4 based on the
- sensitivity analysis in the study of Katata (2014).

4.2 Uncertainties in models for Hg dry deposition

- 497 Hg dry deposition flux is proportional to the corresponding Hg concentration (Zhang
- 498 et al., 2009):

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$$F_{\text{dry}} = v_d \cdot C_z \tag{7}$$

- where F_{dry} is the Hg dry deposition flux; C_z is the Hg concentration at reference
- height z; and v_d is the dry deposition velocity.
- In this part, the uncertainties of speciated Hg concentration measurements were
- first discussed, followed by the uncertainty analyses of Hg dry deposition models.

4.2.1 Uncertainties in speciated Hg concentration measurements

- Although many new methods and apparatus have been or are being developed to
- better determine speciated Hg concentrations in ambient air, up to now the Tekran
- 507 2537/1130/1135 system is still the most widely used commercial instrument for
- continuous measurements of speciated Hg (Gustin et al., 2015). Regional and global
- monitoring networks such as Atmospheric Mercury Network (AMNet) and GMOS
- 510 have all been using the Tekran systems and developed systematic quality assurance
- and quality control (QA/QC) protocols to assure data quality (Obrist et al., 2018).
- Therefore, this section is mainly to assess the uncertainties of the Tekran system.
- Tekran 2537 uses a pair of gold trap cartridges (A/B) to capture GEM in order to
- achieve continuous observation and to reduce the uncertainty of GEM measurements.
- The standard operating procedure (SOP) of GMOS for the determination of GEM

516 requires the RPD of the average of five consecutive A trap concentrations and five consecutive B trap concentrations to be less than 10 % (Sprovieri et al., 2017). In field 517 comparisons held by EMEP, the RSD from Tekran measurements are also generally 518 within 10 % (Aas, 2006). However, in the Reno Atmospheric Mercury 519 Intercomparison eXperiment (RAMIX) campaign, the RPD between two co-located 520 Tekran systems was as high as 25–35 % (Gustin et al., 2013). This was possibly 521 related to other factors, such as the configuration of the manifold, which could be 522 occasional or systemic. Therefore, the overall uncertainty of GEM concentration 523 524 measurements by the Tekran system is estimated to be $\pm (10-30)$ %. Tekran 1130 uses a KCl-coated denuder to pre-concentrate GOM, and the collected 525 GOM is then thermally desorbed at 500 °C and converted to GEM for quantification. 526 A number of studies have reported the significant interference of ozone and humidity 527 on the GOM capture rate of the denuder (Lyman et al., 2010; Jaffe et al., 2014; 528 McClure et al., 2014; Gustin et al., 2015). McClure et al., (2014) found that the KCl-529 530 coated denuder only captures 20–54 % HgBr₂ in the ambient air under the influence 531 of humidity and ozone. Huang et al. (2013) compared denuder- and membrane-based methods, and reported that the KCl-coated denuder only captures 27-60 % of the 532 533 GOM measured by CEMs. Discrepancy with a factor of 2–3 at times was found between the Tekran system and other new methods in the RAMIX campaign (Gustin 534 535 et al., 2013). Cheng and Zhang (2017) developed a numerical method to assess the uncertainty of GOM measurements, and estimated the GOM concentrations measured 536 537 at 13 AMNet sites to be underestimated by a factor of 1.3 to more than 2. Gustin et al. (2015) reported that the capture efficiency ratio of CEMs over the denuder method for 538 five major GOM compounds ranges from 1.6 to 12.6. Recent studies (Huang and 539 Gustin, 2015a; Huang et al., 2017) applied a correction factor of 3 for Tekran GOM 540 data when modeling dry deposition flux. Therefore, the overall uncertainty of the 541 GOM concentration measured by the Tekran system is estimated to be ± 200 % or 542 within a factor of 3. It should be noted that the correction factor of 3 is not universally 543 applicable. Different humidity levels or ozone concentrations lead to a significant 544 change in underestimation. Different chemical forms of GOM also have different KCl 545 capture efficiencies. Therefore, accurate quantification methods for measuring the 546 total and chemically speciated GOM concentration are in urgent needs. 547 Tekran 1135 uses a quartz filter downstream the KCl denuder to collect PM_{2.5}, and 548 the collected fine particles are then thermally desorbed at 800 °C at a pyrolyzer and 549

converted to GEM for the quantification of PBM, or rather PBM_{2.5}. The uncertainties

in PBM concentration measurements have not been systemically assessed so far.

Gustin et al. (2015) pointed out that breakthrough of GOM from the upstream denuder

could result in the retention of GOM on the quartz filter and induce consequent PBM

overestimation. The RAMIX campaign showed that the RSD of PBM measurements

was 70–100 % when the Tekran systems were free standing (Gustin et al., 2013).

Coarse PBM is neglected in Tekran measurements with an impactor removing all

coarse particles. However, based on the estimation of Zhang et al. (2016b), about

558 30 % of PBM could be on coarse particles. Regarding the limited evidence from

previous studies, the overall uncertainty of the PBM concentration measured by the

Tekran system is estimated to be ± 100 % or a factor of 2.

4.2.2 Resistance model for GOM dry deposition

Based on Eq. (7), the dry deposition velocity (v_d) is the key parameter in the

determination of Hg dry deposition flux. It can be estimated using a resistance model

564 (Zhang et al., 2002; Zhang et al., 2003):

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$$v_d = \frac{1}{R_a + R_b + R_c} \tag{8}$$

where R_a is the aerodynamic resistance depending on the meteorological conditions

and the land use category; R_b is the quasi-laminar resistance, a function of friction

velocity and the molecular diffusivity of each chemical species (Zhang et al., 2002);

and R_c is the canopy resistance which can be further parameterized as follows:

$$R_c = \left(\frac{1 - W_{st}}{R_{ct} + R_{rr}} + \frac{1}{R_{rr}}\right)^{-1} \tag{9}$$

where W_{st} is the fraction of stomatal blocking under wet conditions; R_{st} is the

stomatal resistance; R_m is the mesophyll resistance; and R_{ns} is the non-stomatal

resistance which is comprised of in-canopy, soil, and cuticle resistances. Cuticle and

soil resistances for GOM are scaled to those of SO₂ and O₃ by the following equation:

$$R_{x,\text{GOM}} = \left(\frac{\alpha_{\text{GOM}}}{R_{x,\text{SO}_2}} + \frac{\beta_{\text{GOM}}}{R_{x,\text{O}_2}}\right)^{-1}$$
(10)

where R_x is the cuticle or soil resistance; α and β are two scaling parameters (Zhang

et al., 2003; L. Zhang et al., 2012). Among the numerous parameters in the resistance

578 model the two scaling factors for the non-stomatal resistance components regarding

579 the solubility and reactivity of the chemical species are the most sensitive ones. The

- values for HNO₃ ($\alpha=\beta=10$) used to be applied in the model for GOM (Marsik et al.,
- 581 2007; Castro et al., 2012; L. Zhang et al., 2012). However, some other studies found
- the values for HONO ($\alpha=\beta=2$) are probably more suitable for GOM due to equivalent
- effective Henry's Law constants (H^*) between HONO and HgCl₂ (Lyman et al.,
- 584 2007). Huang and Gustin (2015a) indicated that no single value could be used to
- calculate GOM dry deposition due to the unknown GOM compounds. Various values
- for the two scaling parameters ($\alpha=\beta=2, 5, 7$ and 10) were used in Huang et al. (2017)
- to identify dominant GOM deposition species.
- The uncertainties of R_a and R_b are estimated to be generally small, within the range
- of ± 30 % (Zhang et al., 2003; Huang et al., 2012a), while the uncertainty of R_c usually
- has a larger impact, especially through the selection of α and β . Lyman et al. (2007)
- changed the values of α and β from 2 to 10, and found a 120% enhancement of v_d .
- With a correction factor of 3 for the GOM concentration measured by Tekran, Huang
- and Gustin (2015a) got similar modeled and measured GOM dry deposition values
- with bias of up to ± 100 %. Huang et al. (2017) also applied the correction factor of 3,
- tested different values of α and β , and found the bias of GOM dry deposition
- simulation to be up to a factor of 2.5. As discussed above, the overall uncertainty of
- the GOM concentration measured by Tekran is within a factor of 3. If the GOM dry
- deposition simulation is directly based on the Tekran GOM data, its uncertainty level
- would be much higher than a factor of 3. However, recent studies (Huang et al., 2014;
- Huang and Gustin, 2015a; Huang et al., 2017) have used a correction factor of 3 for
- 601 GOM concentration data which offsets the uncertainty of GOM dry deposition.
- Therefore, the overall uncertainty in GOM dry deposition simulation is estimated to
- 603 be a factor of 2.5–4 or $\pm (150-300)$ %.

4.2.3 Resistance model for PBM dry deposition

- For PBM dry deposition, resistance models regarding both fine and coarse particles
- are more and more widely applied based on the theory that v_d for atmospheric
- particles strongly depends on particle size (Dastoor and Larocque, 2004; Zhang et al.,
- 2009; Zhang and He, 2014). Many independent studies (Fang et al., 2012b; Zhu et al.,
- 609 2014) showed that Hg in coarse particles constitutes a large mass fraction of the total
- PBM, which was previously neglected. PBM measured by Tekran 2537/1130/1135
- only considers fine particles. Taking coarse particles into consideration, the total PBM
- dry deposition is calculated as follows (Zhang et al., 2016b):

$$F_{\text{dry,PBM}} = C_f \left(v_f + \frac{f}{1 - f} v_c \right) \tag{11}$$

where $F_{\text{dry,PBM}}$ is the total PBM dry deposition flux; C_f is the mass concentration of PBM in fine particles; v_f and v_c are the dry deposition velocities of PBM for fine and coarse particles, respectively; and f is the mass fraction of PBM in coarse particles. v_f and v_c can be calculated using the following equation (Zhang et al., 2001):

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$$v_x = v_g + \frac{1}{R_a + R_s}$$
 (12)

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where v_x is v_f or v_c ; v_g is the gravitational settling velocity; R_a is the aerodynamic

620 resistance; and R_s is the surface resistance which can be parameterized as a function of collection efficiencies from Brownian diffusion, impaction, and interception 621 mechanisms (L. Zhang et al., 2012; Zhang et al., 2016b). Zhang and He (2014) have 622 developed an easier bulk algorithm based on the v_x scheme of Zhang et al. (2001) to 623 make this model more widely applicable in monitoring networks. 624 Zhang et al. (2001) conducted a model comparison with two PBM dry deposition 625 schemes, and the results showed that the differences between models are generally 626 within the range of 20 %. However, recent studies found the proportion of coarse 627 particles plays a crucial role in the evaluation of PBM dry deposition velocity (Zhang 628 et al., 2016b). Zhang et al. (2016b) assumed that 30 % of the total PBM is on coarse 629 particles, and found that 44 % PBM deposition was caused by coarse particle 630 631 deposition. We tested the model used by Zhang et al. (2016b), and found a 2-fold change when we increased the coarse PBM proportion from 30 % to 50%. In other 632 words, the uncertainty of the PBM deposition velocity could be about $\pm (60-100)$ %. 633 As discussed above, the overall uncertainty of the PBM concentration measured by 634 Tekran is about ± 100 %. Considering both aspects and applying the calculation 635 method based on Eq. (2), the overall PBM uncertainty in GOM dry deposition 636 simulation is estimated to be $\pm (120-150)$ %. 637

4.2.4 Bidirectional model for GEM dry deposition

GEM dry deposition can also be calculated using the resistance model with different parameters. However, the re-emission and natural emission of GEM must be taken into consideration. Net GEM dry deposition is estimated from the difference between the estimated unidirectional deposition flux and the modeled total re-emission plus natural emission in the resistance model (L. Zhang et al., 2012).

A bidirectional air-surface exchange model modified from the resistance model is more and more recommended in recent years (Zhang et al., 2009; Bash, 2010; Wang et al., 2014; Zhang et al., 2016b; Zhu et al., 2016). In the bidirectional scheme, the GEM dry deposition flux can be calculated as follows (Zhang et al., 2009):

$$F_{\text{dry,GEM}} = \frac{\chi_a - \chi_c}{R_a + R_b} \tag{13}$$

$$\chi_{c} = \left(\frac{\chi_{a}}{R_{a} + R_{b}} + \frac{\chi_{st}}{R_{st} + R_{m}} + \frac{\chi_{g}}{R_{ac} + R_{g}}\right) \left(\frac{1}{R_{a} + R_{b}} + \frac{1}{R_{st} + R_{m}} + \frac{1}{R_{ac} + R_{g}} + \frac{1}{R_{cut}}\right)^{-1} (14)$$

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where $F_{\text{dry},\text{GEM}}$ is the net GEM dry deposition flux; χ_a is the GEM concentration at a

reference height; R_a , R_b , R_{st} , R_m , R_{ac} , R_g and R_{cut} are aerodynamic, quasi-laminar, 651 stomatal, mesophyll, in-canopy aerodynamic, ground surface and cuticle resistances, 652 respectively (Zhang et al., 2016b); and χ_{st} and χ_g are canopy, stomatal and ground 653 surface compensation points, respectively. Based on observations on different land use 654 categories, Wright and Zhang (2015) have proposed a range of χ_{st} and χ_{g} . 655 The studies of L. Zhang et al. (2012) and Zhang et al. (2016b) have shown the great 656 657 importance of the previously neglected GEM dry deposition. Due to the presence of natural and re-emission of GEM, the net GEM dry deposition has a higher uncertainty 658 659 level than GOM and PBM dry deposition. Although both the studies of L. Zhang et al. (2012) and Zhang et al. (2016b) reported the uncertainty of net GEM dry deposition to 660 661 be averagely about a factor of 2, there were many exceptions (over a factor of 2–5) according to L. Zhang et al. (2012), especially when the net GEM dry deposition 662 fluxes were at low level. Based on the above concern and the sensitivity analysis 663 conducted in the study of Zhang et al. (2016b), the overall uncertainty of the net GEM 664 665 dry deposition simulation is within a factor of 2 or ±100 % when GEM dominates the total Hg dry deposition, while it could be as high as a factor of 5 or ± 400 % when 666

4.3 Uncertainties in models for forest Hg deposition

GOM+PBM dominate the total dry deposition.

The study of Wang et al. (2016a) is to date the only modeling study for litterfall Hg deposition. Monte Carlo simulation was adopted to assess the global Hg deposition through litterfall based on the measured litterfall Hg concentrations and the global litterfall biomass distribution. The estimated global annual Hg deposition through litterfall was reported to be 1180 t with a relative uncertainty of ± 60 %. At the site level comparison, the difference is within a factor of 2. Therefore, the overall

675 uncertainty of litterfall Hg deposition is estimated to be $\pm (60-100)$ %. There is no modeling study on throughfall Hg deposition so far. Consequently, we can only use 676 the overall uncertainty of wet and dry deposition simulation to represent throughfall, 677 which will be discussed in the next section. 678

Summary of uncertainties in Hg deposition to terrestrial surfaces

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Based on the review work above, the overall uncertainties of wet, dry, and forest Hg 680 deposition can be calculated using the following equation: 681

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$$\delta_{A+B} = \frac{U_{A+B}}{F_{A+B}} = \frac{\sqrt{U_A^2 + U_B^2}}{F_{A+B}} = \frac{\sqrt{F_{A+B}^2 P_A^2 \delta_A^2 + F_{A+B}^2 P_B^2 \delta_B^2}}{F_{A+B}} = \sqrt{P_A^2 \delta_A^2 + P_B^2 \delta_B^2}$$
(15)

where δ_A , δ_B , and δ_{A+B} are the relative uncertainties of Part A, Part B, and the total deposition flux, respectively; U_A , U_B , and U_{A+B} are the absolute uncertainties of them, 684 respectively; F_{A+B} is the total deposition flux; and P_A and P_B are the proportions of 685 Part A and Part B deposition fluxes, respectively. 686 Table 1 summarizes the previously estimated relative uncertainties for wet, dry, and 687 forest Hg deposition fluxes. Although the uncertainty of precipitation Hg deposition 688 flux is low, the uncertainty of non-precipitation Hg deposition has been neglected. 689 690 Due to the condensation effect, non-precipitation deposition could contribute equivalent or even larger proportion to Hg wet deposition than rainfall (Stankwitz et 691 al., 2012; Blackwell and Driscoll, 2015b; Weiss-Penzias et al., 2016b; Gerson et al., 692 693 2017). Considering the global area of hotspot regions for cloud, fog, dew, and frost, such as alpine and coastal regions, the overall contribution of non-precipitation 694 695 deposition to Hg wet deposition is approximately 5–10 %. Given the high uncertainty level of non-precipitation Hg deposition, the overall uncertainties in the observation 696

and simulation of global Hg wet deposition are estimated to be $\pm (20-35)$ % and $\pm (30-$ 55) %, respectively. Hg dry deposition has a much larger uncertainty level than wet deposition from both observation and simulation perspectives. High GOM deposition fluxes were exhibited in North America, while high PBM deposition fluxes were found in East Asia (Wright et al., 2016). Based on the global observation and simulation data (Wright et al., 2016; Zhang et al., 2016b), the ratio of global GOM dry deposition over PBM dry deposition could be in the range of 1:1 to 3:1, and the ratio of global GEM dry deposition over RM (GOM+PBM) dry deposition could be in the range of

- 1:9 to 9:1. Therefore, the overall uncertainties in the observation and simulation of
- global Hg dry deposition are estimated to be $\pm (50-90)$ % and $\pm (90-130)$ %,
- 708 respectively.
- 709 Without studies specifically on throughfall deposition modeling, the uncertainty of
- throughfall Hg deposition simulation has been estimated based on the uncertainties of
- both wet and dry deposition simulation, and turned out to be about $\pm (50-90)$ %.
- Studies on both litterfall and throughfall Hg deposition (Larssen et al., 2008; Navrátil
- et al., 2014; Luo et al., 2016; Ma et al., 2015; Fu et al., 2016a; Wang et al., 2016a;
- Gerson et al., 2017) showed that the relative contributions of litterfall and throughfall
- could be in the range of 2:3 to 4:1. Accordingly, the overall uncertainties in the
- observation and simulation of global forest Hg deposition are estimated to be $\pm (15-$
- 717 25) % and $\pm (40-70)$ %, respectively.
- Based on global and regional modeling studies (Selin and Jacob, 2008; Wang et al.,
- 719 2016a; UN Environment, 2019), the relative contributions of wet, dry, and litterfall
- Hg deposition are estimated to be approximately 1:2:1. With the previously estimated
- 721 uncertainty ranges for wet, dry, and litterfall deposition, the overall uncertainties in
- the observation and simulation of global total Hg deposition are calculated to be
- $\pm (25-50)$ % and $\pm (45-70)$ %, respectively. It should be noted that the low overall
- uncertainty for observation can only be achieved when Hg deposition networks are
- 725 established worldwide.

6 Implications and future research needs

- With a big effort of literature review, this study has estimated the uncertainties in the
- observation and simulation of global Hg deposition to the land surfaces through
- 729 different pathways. The implications from the comprehensive uncertainty analysis and
- 730 the derivative research needs in the future are as follows:
- 731 (1) The observation methods for both wet and forest Hg deposition fluxes have low
- uncertainty levels. Although large uncertainties still exist in the methods for Hg dry
- deposition measurements, the overall uncertainty in global Hg deposition observation
- can be as low as $\pm (25-50)$ %. Optimized surrogate surfaces and DFCs are economic
- approaches for RM and GEM measurements, respectively, and could be useful
- methods for the global dry deposition network.
- 737 (2) Methods with high time resolution for the accurate measurements of GOM and
- 738 PBM concentrations are in urgent needs. On account of the GOM dry deposition

velocity, the chemical form of GOM also plays a crucial role. Different model parameterizations should be applied for different GOM species. Therefore, quantification methods for measuring different GOM species need to be developed to improve the simulation of GOM dry deposition flux.

- (3) More comparisons between observation and simulation of the GEM dry deposition flux should be conducted to improve model parameterization. Moreover, the GEM deposition process is complicated in forests. It is useful to measure the above-canopy apparent deposition flux, the under-canopy dry deposition flux, the litterfall deposition flux, and the throughfall deposition flux at the same site to get a more comprehensive understanding of the process.
- (4) Non-precipitation Hg wet deposition has been neglected in the global monitoring networks and modeling studies. Cloud, fog, or even dew and frost Hg deposition could be quite important in hotspot regions, such as alpine and coastal areas. It could be enriched in aqueous Hg and affect other deposition processes, or in other words, change the overall Hg residence time. Extremely large uncertainties still exist in both observation and simulation of non-precipitation Hg wet deposition. More standardized sampling methods are required for long-term observation of non-precipitation Hg wet deposition.
- (5) Asia has the highest atmospheric Hg concentration level. However, the Hg deposition studies in Asia are still quite limited. The Hg wet deposition network in Asia is not as mature as in North America and Europe, and there are only a few scattered studies on dry deposition in East Asia. The Hg wet and dry deposition processes in Asia could be quite different from those in North America and Europe because of the high atmospheric Hg and high PM condition in Asia.

Data availability. Data presented in this study were all generated from published literature and are available from the original researchers. Data in this research are available in the Supplement Information.

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1430 Table Captions

Table 1. Summary of relative uncertainties of different types of Hg deposition to

terrestrial surfaces.

Table 1. Summary of relative uncertainties of different types of Hg deposition to terrestrial surfaces.

Type of Hg deposition	Relative uncertainty in observation (%)	Relative uncertainty in simulation (%)
Wet deposition	±(20–35)	±(30-55)
Precipitation	±(15–20)	±(30–50)
Cloud, fog, dew, and frost	±(200-300)	±(200–300)
Dry deposition	$\pm (50-90)$	±(90-130)
GOM dry deposition	$\pm (60-70)$	$\pm (150-300)$
PBM dry deposition	±(80–100)	$\pm (120-150)$
GEM dry deposition	±(100–200)	±(100-400)
Forest deposition	±(15–25)	±(40-70)
Litterfall	±(20–30)	±(60–100)
Throughfall	±(20–30)	±(50–90)
Overall	±(25–50)	±(45-70)

Figure Captions 1438 Figure 1. Global distribution of the observed Hg wet deposition fluxes by observation 1439 networks around the world ($\mu g m^{-2} vr^{-1}$). 1440 Figure 2. Global distribution of the (a) GOM, (b) PBM, and (c) GEM dry deposition 1441 fluxes (µg m⁻² yr⁻¹) from observation-based estimation. 1442 Figure 3. Relationship between the elevation and the GOM dry deposition flux. 1443 1444 Figure 4. Comparison between the GOM dry deposition fluxes from direct observations and from model simulations based on measurements of GOM 1445 concentrations. The numbers in brackets stand for the numbers of samples. 1446 **Figure 5.** Dry deposition fluxes (cyan columns with black bars as standard deviations) 1447 of (a) GOM, (b) PBM and (c) GEM for different terrestrial surface types. "Water" 1448 stands for the terrestrial surfaces near water. The numbers in brackets stand for the 1449 numbers of samples. 1450 1451

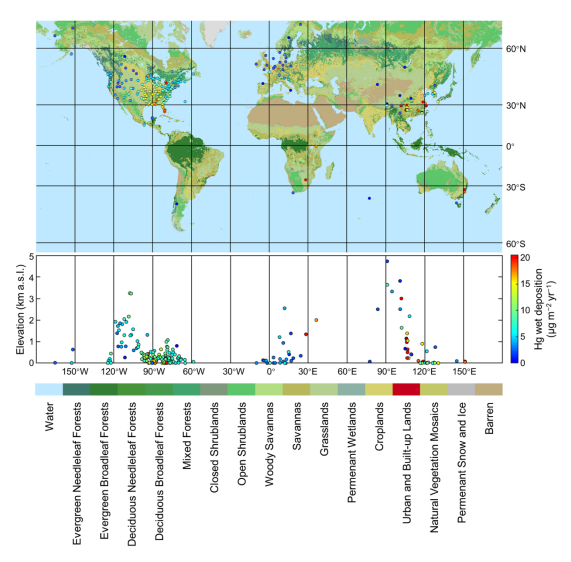


Figure 1. Global distribution of the observed Hg wet deposition fluxes by observation networks around the world ($\mu g \ m^{-2} \ yr^{-1}$).

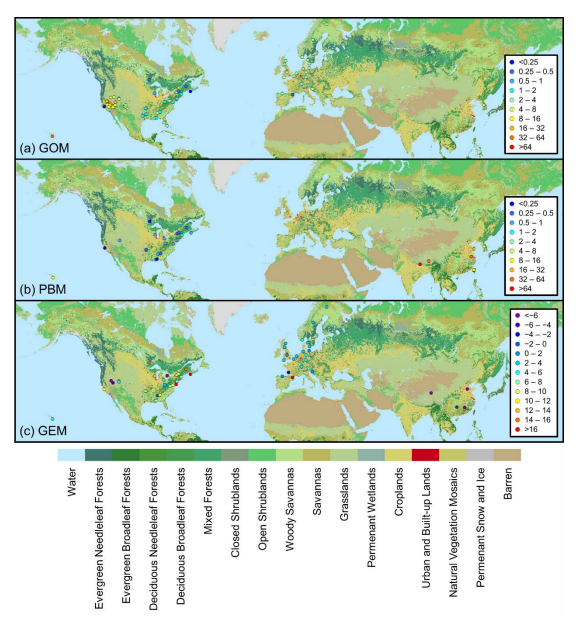


Figure 2. Global distribution of the (a) GOM, (b) PBM, and (c) GEM dry deposition fluxes ($\mu g \ m^{-2} \ yr^{-1}$) from observation-based estimation.

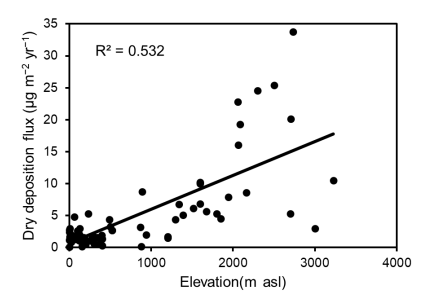


Figure 3. Relationship between the elevation and the GOM dry deposition flux.

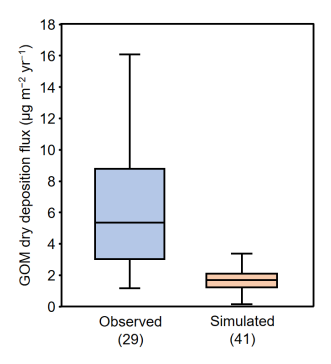


Figure 4. Comparison between the GOM dry deposition fluxes from direct observations and from model simulations based on measurements of GOM concentrations. The numbers in brackets stand for the numbers of samples.

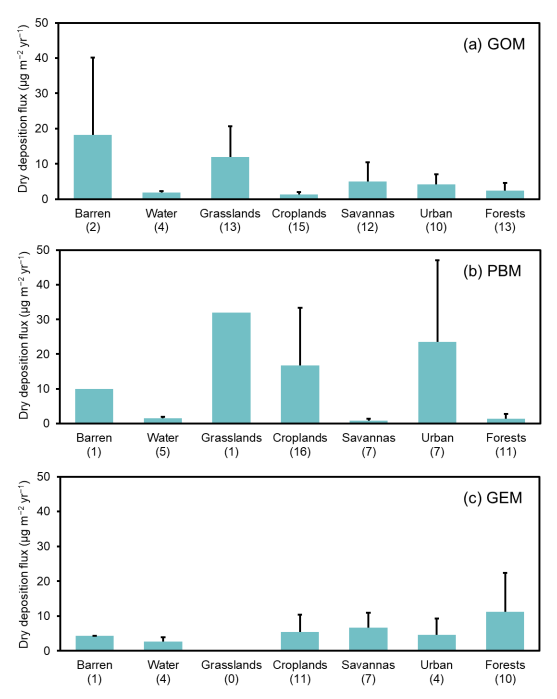


Figure 5. Dry deposition fluxes (cyan columns with black bars as standard deviations) of (a) GOM, (b) PBM and (c) GEM for different terrestrial surface types. "Water" stands for the terrestrial surfaces near water. The numbers in brackets stand for the numbers of samples.