



Observations of atmospheric mercury in China: a critical review

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Received: 28 January 2015 – Published in Atmos. Chem. Phys. Discuss.: 22 April 2015

Revised: 11 August 2015 – Accepted: 11 August 2015 – Published: 24 August 2015

Abstract. China presently contributes the largest amount of anthropogenic mercury (Hg) emission into the atmosphere in the world. Over the past decade, numerous studies have been conducted to characterize the concentration and forms of atmospheric Hg in China, which provide insights into the spatial and temporal distributions of atmospheric Hg through ground-based measurements at widely diverse geographical locations and during cruise and flight campaigns. In this paper, we present a comprehensive review of the state of understanding in atmospheric Hg in China. Gaseous elemental mercury (GEM) and particulate-bound mercury (PBM) measured at the remote sites in China are substantially elevated compared to the background values in the Northern Hemisphere. In Chinese urban areas, the highly elevated GEM, PBM and gaseous oxidized mercury (GOM) were mainly derived from local anthropogenic Hg emissions, whereas regional anthropogenic emissions and long-range transport from domestic source regions are the primary causes of the elevated GEM and PBM concentrations at remote sites. Using 7–9 years of continuous observations at a remote site and an urban site, a slight increase in atmospheric GEM ($2.4\text{--}2.5\% \text{ yr}^{-1}$) was identified (paired samples test: $p < 0.01$), which is in agreement with the increasing domestic anthropogenic emissions. Anthropogenic GEM emission quantity in China estimated through the observed GEM / CO concentration ratios ranged from 632 to 1138 t annually over the past decade, 2–3 times larger than published values using emission activity data. Modeling results and filed measurements show dry deposition is the predominant process for removing Hg from the atmosphere, 2.5–9.0 times larger than wet deposition, due to the elevated atmospheric GEM and

PBM concentrations that facilitate dry deposition to terrestrial landscapes. Further studies to reconcile the observed and simulated Hg concentrations, to understand the impact of domestic emission reduction on Hg concentration and deposition and to delineate the role of Hg emission and deposition of China in the global Hg biogeochemical cycle, are needed.

1 Introduction

Mercury (Hg) is a toxic air pollutant that exists in the atmosphere. There are three operationally defined Hg forms: gaseous elemental mercury (GEM), particulate-bound mercury (PBM), and gaseous oxidized mercury (GOM). The sum of GEM and GOM is known as total gaseous mercury (TGM). Because of its mild reactivity, high volatility, low dry deposition velocity and water solubility, GEM is the most abundant form of Hg in the atmosphere. GEM has an atmospheric residence time of 0.5–2 years and can spread globally before being deposited to earth's surfaces (Schroeder and Munthe, 1998). Rapid transformations of GEM to GOM and PBM have been observed in the polar regions, upper free troposphere, lower stratosphere, and marine boundary layer, indicating a much shorter residence time of GEM in these environments (Schroeder et al., 1998; Hedgecock and Pirrone, 2003; Lyman and Jaffe, 2012; Timonen et al., 2013; Murphy et al., 1998). On the other hand, GOM and PBM have much higher dry deposition velocity and water solubility, and can be readily removed from the atmosphere. The persistence of GEM makes Hg one of the most widely distributed pollutants in the atmosphere.

Both anthropogenic and natural sources emit GEM into the atmosphere, while anthropogenic sources also emit GOM and PBM (Pirrone et al., 2010; Pacyna et al., 2010). Due to the anthropogenic emission and recycling of deposited mercury, the budget of atmospheric Hg has been increasing by a factor of 3 since the onset of the industrial revolution (Mason and Sheu, 2002). Over the past 2 decades, advancement in emission control measures has greatly reduced anthropogenic Hg emissions and led to a decrease in atmospheric TGM concentrations in Europe and North America (Slemr and Scheel, 1998; Slemr et al., 2003; Cole et al., 2013). The global anthropogenic emissions, however, have remained relatively unchanged during the same period, probably caused by the increase in anthropogenic emissions in Asia because of the increasing energy use and lower degree of emission control (AMAP/UNEP, 2013; L. Zhang et al., 2015). China is the world's strongest source region of anthropogenic Hg emission. The annual anthropogenic Hg emissions were in the range from 500 to 1000 t (Streets et al., 2005; Wu et al., 2006; Liang et al., 2013), approximately 1/3 of the global anthropogenic emissions. GEM emission fluxes from terrestrial landscapes in China were also found to be significantly higher than those observed in Europe and North America (Fu et al., 2015).

There is a need to assess the source, transport, transformation and deposition of atmospheric Hg in China, which is critical in understanding the global biogeochemical cycle of Hg. In the past ~ 15 years, significant understanding has been obtained regarding atmospheric Hg studies in China. The knowledge is crucial to resolve the spatial distribution of atmospheric Hg, constrain the Hg emission inventory, assess Hg transport, and evaluate the roles of China emissions in the regional- and global-scale Hg biogeochemical cycles. In our previous study (Fu et al., 2012c), we synthesized the observations of atmospheric Hg in China before 2011. Over the past 4–5 years, many advances have been achieved in atmospheric Hg studies in China. In this paper, we comprehensively review the atmospheric Hg data in China, discuss spatial and temporal distributions and estimate the Hg emissions in China based on the observations. A new analysis regarding the long-term trend of atmospheric GEM is also presented. The trend helps explain the discrepancy between decreasing atmospheric GEM concentrations in Europe and North America and increasing global anthropogenic emissions. The implications and future research needs for further understanding of atmospheric Hg in China are also presented.

2 Observational approaches

2.1 Atmospheric total gaseous mercury (TGM) and gaseous elemental mercury (GEM)

Most TGM and GEM measurements in China were made using the commercial model 2537 A/B automated ambient mercury vapor analyzer (Tekran, Canada). This analyzer has a detection limit of $\sim 0.1 \text{ ng m}^{-3}$ as specified by the Tekran Instrument Corporation. There have been debates as to whether Tekran model 2537 measures TGM or only GEM. Under the typical conditions of the marine boundary layer (presence of sea salt aerosols and humid air), GOM is not likely to bypass the sampling line with a soda lime trap and an unheated Teflon sampling tube, and therefore the measurement is GEM (Fu et al., 2010b; Moore et al., 2013). For other instances, the measured atmospheric Hg vapor likely represents TGM. GEM concentration in China is approximately 2 orders of magnitude higher than GOM concentration. Therefore, the measured TGM concentration is often analytically indistinguishable from GEM concentration given the magnitude of the detection limit. In this review, we refer the atmospheric TGM and GEM observations in China to GEM for clarity and consistency.

The RA-915 +Hg analyzer (Lumex, Russia) has also been used for monitoring of GEM in studies at sites with elevated concentrations (Yang et al., 2009; Ci et al., 2011a, b). This analyzer can measure GEM at a very short interval of 1 s and has a detection limit from 0.3 to 1.0 ng m^{-3} (Ci et al., 2011a). Previous studies revealed that the RA-915 +Hg analyzer has a good agreement with the traditional gold trap/CVAFS (cold-vapor atomic fluorescence spectrometry) method and the Tekran 2537 automated analyzer for GEM measurements in urban areas, with GEM concentrations ranging from 1 to 100 ng m^{-3} (Kim et al., 2006; Fu et al., 2011b). However, further field intercomparisons are needed for validating the RA-915 +Hg analyzer's performance at remote and/or high-altitude sites. A manual method that collects GEM on gold-coated beads or sand traps followed by thermal desorption and detection of CVAFS has also been used (Fang et al., 2004; Xiang and Liu, 2008), which compares well with automated GEM analyzers (Ebinghaus et al., 1999).

2.2 Atmospheric gaseous oxidized mercury (GOM) and particulate bounded mercury (PBM)

There are currently no standard methods for measurements of GOM and PBM due to the unknown chemical composition of GOM and PBM as well as a lack of reliable sampling and calibration techniques. Therefore, the measurements of the two atmospheric Hg forms (especially GOM) should be regarded as a qualitative indication (Gustin and Jaffe, 2010; Gustin et al., 2015). Several studies utilized the Tekran 2537/1130/1135 automated speciation analyzer system for GOM and PBM measurements (Fu et al., 2011a;

Zhang et al., 2013; Xu et al., 2015). This system has been described in detail elsewhere (Landis et al., 2002; Lindberg et al., 2002; Lan et al., 2012). Coarse particles with an aerodynamic size of $>2.5\ \mu\text{m}$ are firstly removed by a glass inertial impactor inlet prior to entering the system. Therefore, the PBM measured by this system represents Hg bounded with fine ($<2.5\ \mu\text{m}$) particulates. The detection limits of GOM and PBM have been reported to be $0.5\text{--}6\ \text{pg m}^{-3}$, depending on sampling duration (Landis et al., 2002; Poissant et al., 2005). There are uncertainties about GOM and PBM measurements using the Tekran system, including a lack of calibration standards for GOM and PBM, incomplete collection and loss of GOM and PBM during sampling (Jaffe et al., 2014; Gustin et al., 2015). Recent field and laboratory intercomparisons suggested that the GOM measured by the Tekran system may be biased low by a factor of 1.6–10.6 times compared to recently developed methods using the Detector for Oxidized Hg Species (DOHGS), nylon membranes and cation exchange membranes, and the magnitude of bias can be influenced by the forms of GOM, ozone and relative humidity (Lyman et al., 2010; Huang et al., 2013; Gustin et al., 2013; Ambrose et al., 2013; McClure et al., 2014).

A manual method similar to the Tekran 2537/1130/1135 system has also been deployed (Fu et al., 2012a; Zhang et al., 2015a). This method collects GOM and PBM onto a KCl-coated quartz annular denuder and a quartz fiber filter in sequence. After sampling, the annual denuder and quartz filter (transported manually into a quartz tube) are flushed with zero air, followed by thermal desorption that converts GOM and PBM into GEM for CVAFS detection. The detection limits of GOM and PBM based on 3 times the standard deviations of blanks are 1.2 and $4.2\ \text{pg m}^{-3}$, respectively (Fu et al., 2012a). GOM measurement using the manual method is also subjected to the analytical uncertainties associated with the Tekran automated speciation system. Hence, GOM data presented in this paper probably underestimate ambient GOM concentrations in China.

Total particulate-bounded mercury (TPM) has also been measured in China. In these studies, airborne particles were collected onto a glass and/or quartz fiber filter without any upstream particulate collection devices. Fu et al. (2008b) and Xu et al. (2013) used a thermal desorption coupled with detection by a Tekran 2537 analyzer and a Lumex RA-915 +Hg analyzer, while others used acid digestion and CVAFS detection (Fang et al., 2004; Z.W. Wang et al., 2006; Xiu et al., 2009). The thermal desorption method yields relatively lower ($\sim 30\%$) values compared to the acid digestion method, likely due to the matrix interferences that reduce the collection efficiency of Hg onto gold pre-concentration traps (Lynam and Keeler, 2002).

2.3 Wet deposition

Wet deposition flux of atmospheric Hg can be estimated via periodic or event-based measurements of Hg concentration in

rainwater and rainfall depth. Previous studies in China used both a bulk collector (Fu et al., 2008b; Guo et al., 2008; Wan et al., 2009a; Fu et al., 2010a) and a wet-only collector (Fu et al., 2010b; Liu et al., 2011; Wang et al., 2012; Zhu et al., 2014; Zhou et al., 2013). The bulk collector tends to overestimate the wet deposition flux due to the dry deposition of TPM and GOM in the sampling period. Such artifacts may be lowered with a short sampling duration (e.g., <1 week) or at remote sites where TPM and GOM concentration is low (Landis and Keeler, 1997). Given that TPM concentrations in China are generally elevated (Sect. 3.1), previous observations using bulk collectors likely overestimated wet deposition fluxes.

2.4 Atmospheric dry deposition

Dry deposition is an important removal pathway of atmospheric Hg. Dry deposition velocities for atmospheric Hg vary significantly with the physical and chemical forms of Hg, meteorological conditions, and surface characteristics (Gustin and Jaffe, 2010). Zhang et al. (2009) reviewed the dry deposition velocities of different atmospheric Hg forms. Using the measured concentrations of atmospheric Hg species and modeled dry deposition velocities, Zhang et al. (2012) estimated the speciated and total Hg dry deposition fluxes in eastern and central North America. The result suggested that dry deposition flux of Hg is comparable to the wet deposition flux in North America.

Given the levels of GEM, GOM and PBM being highly elevated (more details in Sect. 3.1), dry deposition of Hg may be more important in China than in North America. Studies of dry deposition flux in China are limited due to the challenges in accurately measuring speciated Hg concentrations and dry deposition velocities. Modeling approaches have been attempted to estimate dry deposition fluxes (Lin et al., 2010; L. Wang et al., 2014). However, there may exist significant uncertainties in the estimates because the simulated Hg concentrations are significantly different from the observed concentrations of speciated atmospheric Hg.

Dry deposition flux of Hg in forest ecosystems in China has also been investigated (Wang et al., 2009; Fu et al., 2010b, a). These studies estimated deposition fluxes using data of throughfall and litterfall, which provide a reasonable estimate of total dry deposition in forest ecosystems (St Louis et al., 2001; Zhang et al., 2012). However, due to the incomplete understanding of the sources of Hg in litterfall, air–foliage Hg exchange and dry deposition to forest soil, these estimates should be considered approximate.

3 Results and discussion

3.1 Atmospheric Hg observed at ground-based sites

3.1.1 GEM concentrations, spatial distribution and its relationship with anthropogenic Hg emissions

Table 1 shows the statistical summary of speciated atmospheric Hg concentrations and associated site information. Averaged GEM concentrations at remote sites were in the range of 1.60–5.07 ng m⁻³ (mean = 2.86 ± 0.95 ng m⁻³, *n* = 12), generally elevated compared to those observed in North America, Europe, South Africa and South America (Table 1). The mean GEM concentrations at 11 of the 12 remote sites in China exceeded the background concentration of GEM (1.5–1.7 ng m⁻³) in the Northern Hemisphere (Lindberg et al., 2007). Clear spatial variations of GEM concentrations were observed at the remote sites (Fig. 1). In general, GEM concentrations in northeastern, northwestern, and southwestern China (Mt. Changbai, Mt. Waliguan, and Mt. Ailao, means = 1.60–2.09 ng m⁻³) were lower (*t* test, *p* < 0.01) than those (2.31–5.07 ng m⁻³) in northern, southern, and eastern China (e.g., Chengshantou, Chongming Island, Mt. Leigong, Miyun, Mt. Damei, and Mt. Dinghu). This pattern is similar to the spatial distribution of anthropogenic Hg⁰ emissions (Fig. 1). Most of the Hg emission sources in China are located in northern, central, eastern, and southern China. Anthropogenic GEM emissions in northeastern (Jilin and Heilongjiang provinces), western (Xinjiang, Xizang, and Qinghai provinces), and southwestern China (mainly in Yunnan Province) are lower (Fig. 1). The mean GEM concentrations at remote sites increased with the regional GEM emissions surrounding the sampling sites (Fig. 2), suggesting the influence of anthropogenic GEM emissions on the distribution of atmospheric Hg in remote areas.

Mean GEM concentrations at urban sites ranged from 2.50 to 28.6 ng m⁻³ (mean = 9.20 ± 7.56 ng m⁻³, *n* = 13), significantly higher than the GEM concentrations measured at remote sites (*t* test, *p* < 0.01). In contrast to remote sites, no significant positive correlations were observed between mean GEM concentrations at urban sites and regional GEM emissions surrounding the sampling sites (*p* > 0.05 for all correlations between GEM and emissions within 1° × 1°, 3° × 3° and 5° × 5° grid cells, Fig. 2). Additionally, the mean GEM concentrations at urban sites were significantly different from that at remote sites (*t* test, *p* < 0.01), whereas the anthropogenic GEM emissions within 1° × 1° (*t* test, *p* = 0.06), 3° × 3° (*t* test, *p* = 0.11), and 5° × 5° (*t* test, *p* = 0.19) grid cells surrounding the urban sites and remote sites were not significantly different. We conclude that anthropogenic GEM emissions in the vicinity of the sampling sites play an important role for the highly elevated GEM concentrations at urban sites. Urban areas in China are densely populated and heavily industrialized. The high energy demand and the presence of other large point sources (e.g.,

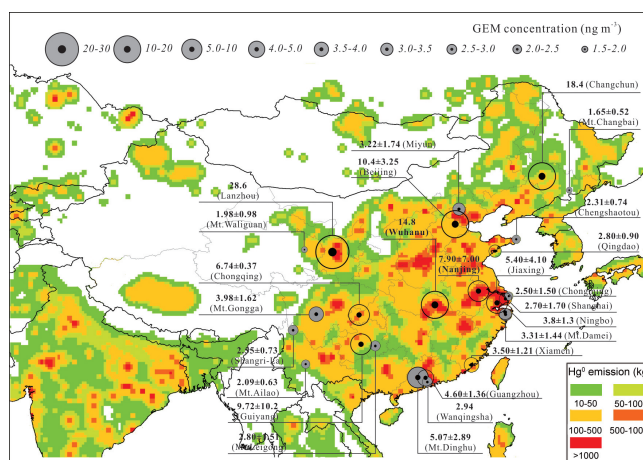


Figure 1. A map showing the locations and atmospheric GEM concentrations observed at ground-based sites in China (gray circles and hollow circles indicate remote and urban sites, respectively). GEM concentrations are indicated by the size of the circles. Data sets were from literatures in Table 1) and 0.5° × 0.5° gridded anthropogenic Hg⁰ emissions in Asia (AMAP/UNEP, 2013).

iron and steel production, non-ferrous metals smelting, cement production) contribute to the large Hg⁰ emissions (Fu et al., 2011a; Zhu et al., 2012; Chen et al., 2013). For example, the annual anthropogenic Hg emissions in Guiyang and Nanjing reached 1.75 t and 4.26 t, respectively (Zhu et al., 2012; Fu and Feng, 2015). Large emissions of GEM from natural surfaces in the urban areas may also play a role for the elevated GEM concentrations (Feng et al., 2005; D. Y. Wang et al., 2006; Zhu et al., 2012). There is a clear spatial distribution pattern of GEM at urban sites (Fig. 1). The mean GEM concentrations in the six coastal cities including Qingdao, Shanghai, Ningbo, Xiamen, Guangzhou and Jiaxing were in the range of 2.70–5.4 ng m⁻³, significantly lower (*t* test, *p* < 0.01) than that (6.74 to 28.6 ng m⁻³) in other inland cities, probably caused by the influence of clean marine air.

3.1.2 TPM and PBM concentrations

TPM/PBM concentrations at remote and urban sites of China ranged from 18.9 to 154 pg m⁻³ (mean = 51.8 pg m⁻³, *n* = 8) and from 141 to 1180 pg m⁻³ (mean = 530 pg m⁻³, *n* = 9), respectively, significantly higher (several to dozens of times) than the observations in North America and Europe (Table 1). The elevated TPM/PBM concentrations in urban areas were most likely due to local anthropogenic emissions (Z. W. Wang et al., 2006; Xiu et al., 2005; Fu et al., 2011a). PBM has an atmospheric residence time ranging from a few hours to a few weeks and can undergo regional transport and further influence PBM at the remote sites (Sheu and Mason, 2001). A positive correlation exists between PBM and GEM concentrations observed at remote and urban sites in China

Table 1. Atmospheric Hg concentrations at ground-based stations in China and other regions worldwide.

Site	Location			Type	Study period	GEM (ng m ⁻³)	PBM/TPM (pg m ⁻³)	GOM (pg m ⁻³)	Reference
	Lon	Lat	Altitude (m a.s.l.)						
Mt. Changbai	128.112° E	42.402° N	740	Remote forest	10/2008–10/2010 07/2013–07/2014	1.60 ± 0.51 1.73 ± 0.48	18.9 ± 15.6	5.7 ± 6.8	Fu et al. (2012b) (Fu et al., 2014)
Mt. Waliguan	100.898° E	36.287° N	3816	Remote grassland	09/2007–09/2008	1.98 ± 0.98	19.4 ± 18.0	7.4 ± 4.8	Fu et al. (2012a)
Mt. Ailao	101.017° E	24.533° N	2450	Remote forest	05/2011–05/2012	2.09 ± 0.63	31.3 ± 28.0	2.2 ± 2.3	Zhang et al. (2015b)
Chengshantou	122.68° E	37.38° N	30	Remote coast	07&10/2007, 01&04/2009	2.31 ± 0.74			Ci et al. (2011a)
Chongming Island	121.908° E	31.522° N	11	Remote coast	09–12/2009	2.50 ± 1.50			Dou et al. (2013)
Shangri-La	99.733° E	28.017° N	3580	Remote forest	11/2009–10/2010	2.55 ± 2.73	37.8 ± 31.0	7.9 ± 7.9	Zhang et al. (2015a)
Mt. Leigong	108.2° E	26.39° N	2178	Remote forest	05/2008–05/2009	2.80 ± 1.51			Fu et al. (2010b)
Wanqingsha	113.55° E	22.7° N	3	Remote coast	11–12/2009	2.94			Li et al. (2011)
Miyun	116.775° E	40.481° N	220	Remote forest	12/2008–11/2009	3.22 ± 1.94	98.2 ± 113	10.1 ± 18.8	Zhang et al. (2013)
Mt. Damei	121.565° E	29.632° N	550	Remote forest	04/2011–04/2013	3.31 ± 1.44	154 ± 104	6.3 ± 3.9	Yu et al. (2015)
Mt. Gongga	102.117° E	29.649° N	1640	Remote forest	05/2005–07/2007	3.98 ± 1.62	30.7 ± 32.0*	6.2 ± 3.9	Fu et al. (2008a); Fu et al. (2008b)
Mt. Dinghu	112.549° E	23.164° N	700	Remote forest	09/2009–04/2010	5.07 ± 2.89			Chen et al. (2013)
Mt. Jiuxian	118.11° E	25.71° N	1700	Remote forest	11/2010, 01&04&08/2011		24.0 ± 14.6		Xu et al. (2013)
Shanghai	121.54° E	31.23° N	19	Urban	08–09/2009 07/2004–04/2006	2.70 ± 1.70	560 ± 220*		Friedli et al. (2011) Xiu et al. (2009)
Qingdao	120.5° E	36.16° N	40	Urban	01/2013	2.80 ± 0.90	245 ± 174*		Zhang et al. (2014)
Xiamen	118.05° E	24.60° N	7	Urban	03/2012–02/2013	3.50 ± 1.61	174 ± 280	61 ± 69	Xu et al. (2015)
Ningbo	121.544° E	29.867° N	10	Urban	10/2007–01/2008	3.79 ± 1.29			Nguyen et al. (2011)
Guangzhou	113.355° E	23.124° N	60	Urban	11/2010–10/2011	4.60 ± 1.60			Chen et al. (2013)
Jiaxing	120.7° E	30.833° N	10	Urban	09/2005	5.40 ± 4.10			Wang et al. (2007)
Chongqing	106.5° E	29.6° N	350	Urban	08/2006–09/2007	6.74 ± 0.37			Yang et al. (2009)
Nanjing	118.78° E	32.05° N	100	Urban	01–12/2011 06/2011–02/2012	7.90 ± 7.00	1100 ± 570*		Zhu et al. (2012) Zhu et al. (2014)
Guiyang	106.72° E	26.57° N	1040	Urban	11/2011–11/2002 12/2009–11/2010	8.40 ± 4.87 10.2 ± 7.06			Feng et al. (2004) Fu and Feng (2015)
Beijing	116.392° E	38.898° N	48	Urban	08–12/2009 02&09/1998 01–12/2006	9.72 ± 10.2 10.4 ± 3.25	368 ± 276 272 573 ± 551*	35.7 ± 43.9	Fu et al. (2011a) Liu et al. (2002) Schleicher et al. (2015)
Wuhan	114.3° E	30.6° N	20	Urban	–/2002	14.8			Xiang and Liu (2008)
Changchun	125.319° E	43.824° N	270	Urban	–/2001	18.4	276*		Fang et al. (2004)
Lanzhou	103.79° E	36.067° N	1540	Urban	–/2004	28.6			Su et al. (2007)
Southeastern coastal cities				Urban	04&07&10&12/1994 11/2010, 01&04&08/2011		955* 141 ± 128		Duan and Yang (1995) Xu et al. (2013)
Suriname, South America	56.983W	5.933° N		Remote	03–07/2007	1.40			Müller et al. (2012)
Cape Point, South Africa	18.483° E	34.35° S	230	Remote	01–12/2009	0.87			Slemr et al. (2011)
Cape Grim, Australia	144.683° E	40.683° S		Remote	2011–02/2013	0.85–0.96			Slemr et al. (2015)
North America				Urban		1.60–4.50	2.5–25.4	6.9–37.2	Peterson et al. (2009); Song et al. (2009); Engle et al. (2010); Liu et al. (2010); Brooks et al. (2010); Lan et al. (2012)
				Remote		1.32–1.66	1.6–13.7	0.5–5.6	Lan et al. (2012); Cole et al. (2013)
Europe				Urban		1.9–3.4	12.5	2.5	Dommergue et al. (2002); Li et al. (2008)
				Remote		1.40–1.93	3.0–32.2	9.1–26.5	Slemr and Scheel (1998); Lee et al. (1998); Munthe et al. (2003); Kock et al. (2005)
Antarctic				Remote		0.23–1.20	116–344	12–224	Dommergue et al. (2010)

(PBM/TPM: * Indicates TPM (total particulate-bound mercury) and the rest indicate PBM (particulate-bound mercury on particles with an aerodynamic diameter < 2.5 µm.)

(Fig. 3), indicating PBM and GEM shared common emission sources. Size-fractionation of particulate Hg in Beijing, Nanjing, and several coastal cities of southeastern China has been investigated. The studies revealed that PBM (the fraction associated with particles having a diameter of < 2.5 µm) constituted 40–70 % of TPM. Given that fine particles are removed from the atmosphere less efficiently than large particles (Zhang et al., 2001), it is possible that the fraction of PBM in TPM in remote areas is higher than that in urban areas.

The fraction (1.0–17.2 %) of TPM/PBM in total atmospheric Hg (THg, defined as the sum of TGM and TPM/PBM) in the urban and remote areas of China was larger than that (0.1–1.2 %) in North America and Europe (Table 1). The fraction of TPM in THg at the urban sites of northern and eastern China was particularly high (5.2–17.2 % as compared to the overall mean fraction at 2.5 %), and higher than the emission ratios of TPM/THg from most anthropogenic emission sources (L. Zhang et al., 2015). These results imply that secondary formation of TPM via

gas-particle partitioning TGM is an important source of atmospheric TPM in urban areas of China. Atmospheric particulate matter (PM) pollution is a special concern in China. The concentrations of PM_{2.5} in eastern and northern China are the highest in the world (van Donkelaar et al., 2010). Elevated atmospheric PM probably facilitates the formation of TPM in the atmosphere, which can explain the large fraction of TPM. The two highest PBM/THg ratios at remote sites were observed at Mt. Damei (4.4 %) and Miyun (3.0 %); both are located in the major anthropogenic source regions in eastern China (Fig. 1). The PBM/THg ratios (1.0–1.5 %) at the other four remote sites in northeastern, northwestern, and southwestern China are relatively lower. This is consistent with the spatial distribution of anthropogenic Hg emissions and atmospheric PM pollution in China (van Donkelaar et al., 2010; AMAP/UNEP, 2013). PBM is more readily removed from the atmosphere than GEM. Therefore, the PBM/THg ratios are expected to be higher near emission sources.

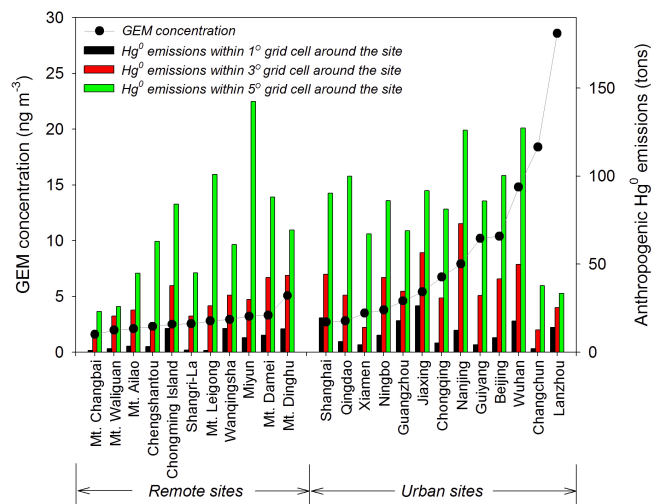


Figure 2. Atmospheric GEM concentrations and total anthropogenic Hg^0 emissions within the $1^\circ \times 1^\circ$, $3^\circ \times 3^\circ$, and $5^\circ \times 5^\circ$ grid cells surrounding the ground-based sites in mainland China.

3.1.3 GOM concentrations

GOM concentrations in urban areas have been measured in Guiyang (southwestern China) and Xiamen (southeastern China). Mean concentrations of GOM in Guiyang and Xiamen were 35.7 and 61.0 pg m^{-3} , respectively (Table 1), higher than most observations in the urban areas of North America and Europe (typically $< 15 \text{ pg m}^{-3}$), except in St. Louis, Missouri, USA (GOM mean = 37.2 pg m^{-3}), and Reno, Nevada, USA (GOM mean = 26.0 pg m^{-3} ; Peterson et al., 2009; Engle et al., 2010). The former was heavily impacted by nearby sources and the latter was influenced by in situ production and mixing of GOM from the free troposphere (Peterson et al., 2009; Engle et al., 2010). In China, anthropogenic emission is the dominant factor contributing to the elevated GOM concentrations in urban areas. Transient high GOM concentrations were often associated with plumes from a coal-fired power plant and a cement factory in Guiyang (Fu et al., 2011a).

Mean GOM concentrations at seven remote sites of China ranged from 2.2 to 10.1 pg m^{-3} (mean = $6.6 \pm 2.4 \text{ pg m}^{-3}$), significantly lower than those observed in urban areas (t test, $p < 0.01$) and comparable to the values in Europe and North America (t test, $p = 0.21$; Table 1). Sources of GOM at remote sites include local emissions, in situ photochemical production, and intrusion of GOM-enriched air from the free troposphere. At the remote sites in eastern and northern China, the local sources (such as domestic heating in small settlements) may be responsible for the elevated GOM concentrations. The impact of long-range transport from industrial and urbanized areas is rare. GOM has a much greater dry deposition velocity ($0.1\text{--}7.6 \text{ cm s}^{-1}$) than GEM ($0.01\text{--}0.3 \text{ cm s}^{-1}$) and PBM ($0.02\text{--}2.1 \text{ cm s}^{-1}$; Zhang et al., 2009), and there-

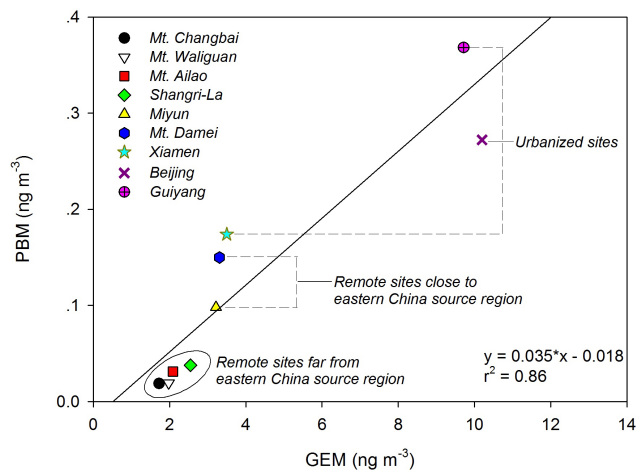


Figure 3. Correlation between GEM and PBM ($\text{Hg-P}_{2.5}$) concentrations in remote and urban areas of China.

fore it has a much shorter atmospheric residence time and limited long-range transport (Lindberg and Stratton, 1998). However, under low air humidity and high wind speed, the possibility of regional transport of GOM cannot be ruled out. For example, the observation at Mt. Waliguan, northeastern Tibetan Plateau, found many high-GOM events were related to air plumes originating from industrial and urbanized centers that are about 90 km to the east of the sampling site (Fu et al., 2012a). Several studies observed a remarkable diel pattern of GOM with higher concentrations during daytime that cannot be explained by emissions (Fu et al., 2012a; Zhang et al., 2013). Although not as significant as in the polar region, free troposphere and marine boundary layer (Lindberg et al., 2002; Swartzendruber et al., 2006; Timonen et al., 2013), in situ GOM production may be an important source of GOM in remote areas of China. There is currently no evidence suggesting the mixing of GOM-enriched air from the upper troposphere/lower stratosphere in China because most of the sampling sites are located in low-altitude areas or the site characteristics do not favor intrusion of an air mass from the upper atmosphere.

3.2 Atmospheric Hg in the marine boundary layer

The first measurement of GEM in the marine boundary layer near China was made onboard a ship during a cruise campaign in the South China Sea (SCS) in August 2007 (Fu et al., 2010c). The SCS is surrounded by China, Vietnam and Laos in the Indo-China Peninsula, and the Philippines. The GEM concentration displayed a spatial pattern with concentrations ($1.29\text{--}1.94 \text{ ng m}^{-3}$) in the open sea close to the background values ($1.5\text{--}1.7 \text{ ng m}^{-3}$) in the Northern Hemisphere and with elevated concentrations ($2.5\text{--}4.71 \text{ ng m}^{-3}$) at locations close to mainland China (Fig. 4). Most elevated GEM events were associated with air masses from southern China and the Indo-China Peninsula (Fu et al., 2010c). Atmospheric

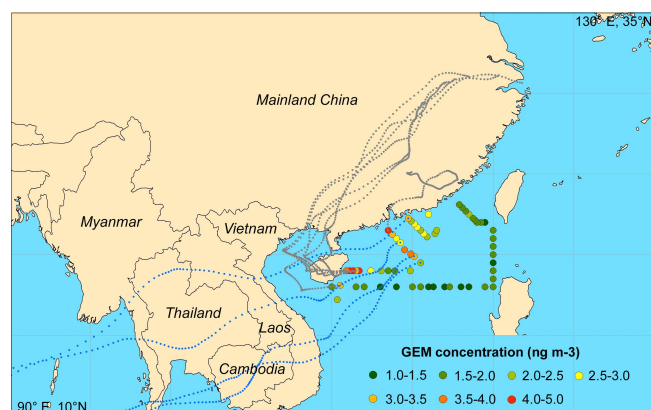


Figure 4. Spatial distribution of GEM concentrations and backwards trajectories in the marine boundary layer of the South China Sea (Fu et al., 2010c).

Hg emissions from mainland China have been investigated extensively (Streets et al., 2005; Wu et al., 2006; Shetty et al., 2008). However, Hg emissions in the Indo-China Peninsula are not well understood. The mean GEM concentration in Da Nang, central eastern Vietnam, was reported to be up to 3.86 ng m^{-3} , suggesting significant anthropogenic Hg emissions in the Indo-China Peninsula (Sheu et al., 2013). Outflows of Hg from mainland China and the Indo-China Peninsula to the SCS vary with the monsoonal climate (Fu et al., 2010c). The cruise measurements by Fu et al. (2010c) were conducted in summer and the observations do not suggest impact of Hg outflows from China and the Indo-China Peninsula on the open sea of the SCS because of the southeastern monsoonal winds. However, as the northwestern monsoonal winds become predominant in winter and spring, the impact of Hg emissions in China and the Indo-China Peninsula into the SCS is expected to increase. For example, the mean GEM concentration in Dongsha Island of the SCS in spring (March–April, 2008) was 56 % higher than that in summer (August, 2008; Sheu et al., 2013).

GEM in the marine boundary layer was also investigated in the Bohai Sea and Yellow Sea from 9 to 18 July 2010 (Ci et al., 2011b). A mean GEM concentration of $2.61 \pm 0.50 \text{ ng m}^{-3}$ was reported, similar to the values observed in the SCS (Fu et al., 2010c) and significantly higher than those measured in the Atlantic Ocean, North Sea, Baltic Sea, Adriatic Sea, and Mediterranean Sea (Leermakers et al., 1997; Wangberg et al., 2001; Temme et al., 2003; Gardfeldt et al., 2003; Sprovieri and Pirrone, 2008). A GEM concentration gradient from the Chinese coast to the eastern Bohai Sea and Yellow Sea was observed (Ci et al., 2011b). Elevated GEM concentrations were mostly related to the outflows from China (Ci et al., 2011b). GEM concentrations were slightly higher in the Yellow Sea ($1.50\text{--}2.50 \text{ ng m}^{-3}$) compared to the nearby western Korea coast ($1.40\text{--}2.00 \text{ ng m}^{-3}$; Nguyen et al., 2011). GEM concentrations in the marine

boundary layer of the East China Sea (ECS) were measured onboard during the cruise campaign from Shanghai, China, to the Antarctic (Xia et al., 2010). A mean concentration of $2.32 \pm 0.49 \text{ ng m}^{-3}$ in the ECS and the coast of Japan and Korea was observed (Xia et al., 2010). These studies show that anthropogenic Hg emissions in China have a strong impact on the observed GEM concentration in marine areas east of China.

3.3 Atmospheric Hg in the free troposphere and lower stratosphere

GEM concentrations in the upper troposphere/lower stratosphere (10–12 km above sea level) over mainland China were measured onboard a passenger aircraft during dozens of CARIBIC flights from Frankfurt, Germany, to Guangzhou, China (Slemr et al., 2009, 2014; Lai et al., 2011). The GEM concentrations were in the range of $0.5\text{--}1.5 \text{ ng m}^{-3}$ for most cases, except near Guangzhou, China, where a GEM concentration exceeding 2.0 ng m^{-3} was observed. Aircraft measurements of GEM over the South China Sea (SCS) in flights from Guangzhou, China, to Manila, Philippines (Lai et al., 2011), showed a concentration range of $0.8\text{--}1.8 \text{ ng m}^{-3}$. These GEM levels are comparable to the high-altitude observations made over southern Europe, western Africa, Atlantic Ocean, and western South America (Slemr et al., 2009, 2014) as well as North America and the North Pacific Ocean (Talbot et al., 2007; Lyman and Jaffe, 2012). Concentrations of ozone, halogen species and other oxidants in the upper troposphere/lower stratosphere are highly elevated and can facilitate depletion of GEM (Slemr et al., 2009; Lyman and Jaffe, 2012; Talbot et al., 2007). Also, the upper troposphere/lower stratosphere has high horizontal wind speed and weak vertical mixing. Hence, the GEM concentrations measured at high altitude over China were similar to the global level of GEM concentrations in the upper troposphere/lower stratosphere.

Nevertheless, several high-GEM events in the upper troposphere/lower stratosphere over mainland China were observed (Slemr et al., 2009). These events were accompanied by elevated CO , aerosols, NO_y , and CH_3CN concentrations transported from the planetary boundary layer over southern China (Slemr et al., 2009). In a transport event in July 2007, the GEM concentrations in the upper troposphere/lower stratosphere were elevated to $2.0\text{--}4.0 \text{ ng m}^{-3}$. This event was impacted by anthropogenic sources in southern China via strong convective processes (Lai et al., 2011). The emission plumes from mainland China on GEM were also detected in the free troposphere during the flights over East Asia (Friedli et al., 2004), which reported several strong GEM pollution events with concentrations exceeding 2.5 ng m^{-3} (up to 6.3 ng m^{-3} ; Friedli et al., 2004).

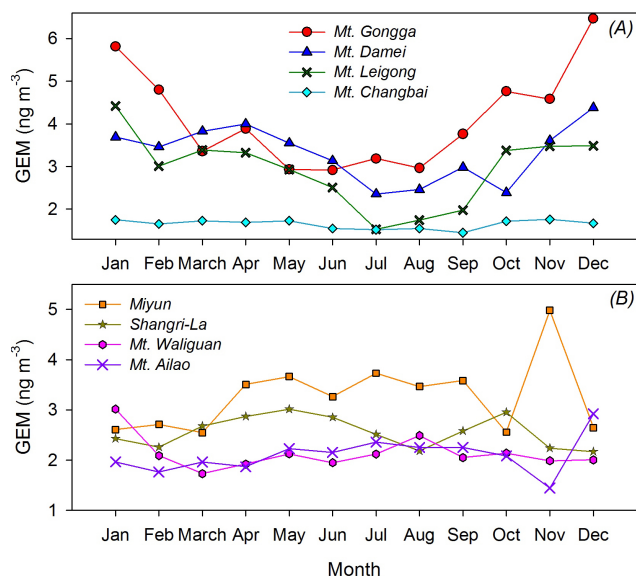


Figure 5. Monthly variations of GEM concentrations at remote sites of China.

3.4 Seasonal variations of GEM and the influence of monsoon

Figure 5 shows the seasonal variations of GEM concentration at ground-based remote sites in China. The variations can be classified into two distinct types (Fig. 5a and b). The mean GEM concentrations at Mt. Gongga, Mt. Damei, Mt. Leigong, and Mt. Changbai in cold seasons (from October to March) were significantly higher (t test: $p < 0.05$ for all) than that in warm seasons (from April to September). On the other hand, the means of GEM at Miyun, Shangri-la, Mt. Ailao, and Mt. Waliguan in warm seasons were relatively higher (2–17%) than that in cold seasons, although the differences were not significant (t test: $p = 0.06$ – 0.66). Seasonal variations of GEM concentrations in urban areas of China were also different among sites. In Guiyang (southwestern China) and Xiamen (southeastern China), monthly mean GEM concentrations were significantly higher (t test: $p < 0.05$ for both) in cold seasons than that in warm seasons (Feng et al., 2004; Xu et al., 2015). In contrast, monthly mean GEM concentrations in Nanjing (eastern China) were found to be $\sim 60\%$ higher in warm seasons than cold seasons (t test: $p < 0.05$; Zhu et al., 2012). In Guangzhou (southern China), the lowest concentration occurred in summer and the maximum was observed in spring ($\sim 22\%$ higher; Chen et al., 2013).

Several hypotheses have been made to explain the seasonal variations of GEM in China, including seasonal changes in anthropogenic GEM emissions and natural emissions. The seasonal emission changes mainly resulted from coal combustion for urban and residential heating during cold seasons. This source lacks emission control devices and releases large amounts of Hg capped by a shallow planetary bound-

ary layer, leading to elevated GEM concentrations at some of the monitoring sites in cold seasons (Feng et al., 2004; Fu et al., 2008a, 2010b). GEM emission from natural surfaces is influenced by many factors (Gustin et al., 2008; Zhang et al., 2009). Due to increasing solar radiation and soil/air temperature, elevated GEM emission fluxes from soil and water in warm seasons were frequently observed (Fu et al., 2012c), which may partially explain the higher GEM concentrations in Nanjing in warm seasons. However, emissions from domestic heating during winter could not explain the lower winter GEM concentrations observed at Miyun, Shangri-La, Mt. Waliguan and Mt. Ailao (Fig. 5b), and the higher natural emissions in summer could not explain the lower summer GEM concentrations measured at Mt. Gongga, Mt. Damei, Mt. Leigong, and Mt. Changbai (Fig. 5a). This implies other not-yet-understood factors that played a role in the observed GEM seasonal variation.

The northerly winter monsoon (northwestern monsoon) and East Asian summer monsoon (southwestern and southeastern monsoon) strongly influence the seasonal winds in East Asia (An, 2000). Summer winds in eastern China mainly originate from the Indian Ocean and the western Pacific Ocean and then move northwards from southern China to northern China, while winter winds mainly originate from Siberia and western Asia and then move southwards from northern China to southern and eastern China (Fig. 6). Given the spatial patterns of anthropogenic GEM emissions in China and East Asia (Fig. 1), monsoonal winds can change the source–receptor relationships at observational sites and subsequently the seasonal GEM trends. Here we calculated the anthropogenic GEM sources impacted factors (ASIF, a conditional possibility of GEM transport from regional source regions to a receptor) for the ground-based sites in China (Fig. 7). The sites located in western, southwestern, and northern China yielded higher ASIF values in the warm season because GEM source regions were up-wind under the influence of the East Asian summer monsoon. Nanjing (eastern China) also showed a larger ASIF in the warm season because easterly winds passed through the Yangtze River delta (one of the most important GEM source regions in China, Fig. 1) prior to arriving in Nanjing. The sites in southern, eastern, and northeastern China showed higher ASIF values in the cold season due to transport of anthropogenic emissions of GEM from the north and west by the northerly winter monsoon. As shown in Fig. 7, the variations of ASIF values in the cold and warm seasons at remote and urban sites of China were in a good agreement with the variations of atmospheric GEM concentrations (except at Mt. Gongga). Monsoonal winds do not have a noticeable influence in the Mt. Gongga area due to the mountainous barrier (the elevation of the summit of Mt. Gongga is 7556 m a.s.l.). Under such circumstances, other factors, including local and regional GEM emissions, atmospheric photochemical processes, mixing layer depth, and meteorologi-

cal conditions played a more important role in the observed seasonal variation.

Seasonal variations of TPM/PBM mostly showed lower concentrations in summer and higher concentrations (up to 1 order of magnitude higher) in winter and fall (Z.W. Wang et al., 2006; Fu et al., 2008b; Zhu et al., 2014; Xu et al., 2015; Xiu et al., 2009; Zhang et al., 2013). The higher TPM/PBM in winter was likely caused by direct PBM emissions, formation of secondary particulate mercury via gas-particle partitioning and a lack of wet scavenging processes (Z. W. Wang et al., 2006; Fu et al., 2008b; Zhu et al., 2014). Data regarding the seasonal variation of GOM concentrations were sparse. The observations at Mt. Gongga (southwestern China) and Miyun (northern China) showed lower GOM concentrations in winter, opposite to the seasonal patterns of GEM (Fu et al., 2008b; Zhang et al., 2013). Possible explanations include stronger gas-particle partitioning of GOM at lower air temperature and higher PM concentrations in winter, as well as faster photochemical production of GOM in warm seasons (Fu et al., 2008b; Zhang et al., 2013).

3.5 Diel variations of atmospheric Hg

Figure 8 displays the diel variations of GEM at the remote sites in China. Most observations showed elevated concentrations during daytime except at Mt. Waliguan and Miyun. Such a diel pattern highlights the impact of regional anthropogenic Hg emissions. Most of these remote sampling sites are located at high altitude far away from major source areas, and therefore were influenced by the diel transition of mountain valley breezes that brought Hg-polluted air to the sampling sites during daytime (Fu et al., 2008a, 2010b; Chen et al., 2013). The higher concentrations during nighttime at Mt. Waliguan were mainly related to downward intrusion of free troposphere airflow originating from industrialized and urbanized areas via long-range transport (Fu et al., 2012a). For urban sites, diel trends of GEM in Guiyang, Hefei, Xiamen, and Guangzhou were similar, with higher concentrations at night due to the accumulation of Hg emitted from local sources in the shallow nocturnal boundary layer (Feng et al., 2004; Chen et al., 2013; Li, 2012). The diel trend of GEM in Nanjing exhibited higher concentrations from early morning to noon, which was attributable to natural emissions and downward mixing of air masses from aloft (Zhu et al., 2012).

Elevated PBM concentrations were observed at night in Guiyang, Mt. Waliguan, and Miyun. The most pronounced diel variation of PBM was observed in Guiyang, with the peak nighttime hourly PBM concentration observed about 9 times higher than the minimum observed during daytime, suggesting the accumulation of local PBM emissions at night (Fu et al., 2011a). The nighttime elevated PBM at Miyun was related to the formation of a temperature inversion layer (Zhang et al., 2013), while that at Mt. Waliguan was caused

by downward intrusion of PBM-enriched air originating from regional industrialized and urbanized areas (Fu et al., 2012a).

GOM observations in China showed increasing concentrations during daytime, coinciding with increasing solar radiation and atmospheric oxidant concentrations (Fu et al., 2008b, 2011a; Zhang et al., 2013; Zhang et al., 2015a). The diel trend of GOM is more pronounced at remote sites where the impact of anthropogenic emissions was weaker and in warm seasons when levels of atmospheric oxidants and solar radiation were high (Fu et al., 2008b; Zhang et al., 2013). This result is consistent with the observations in North America and Europe (Munthe et al., 2003; Lan et al., 2012). Both anthropogenic sources and photochemical oxidation can give atmospheric GOM. Photochemical oxidation produces approximately 8000–15 000 t of atmospheric GOM annually (Selin et al., 2007; Driscoll et al., 2013), about 10–20 times greater than the quantity of anthropogenic emissions. Although this process mainly occurred in the free troposphere, stratosphere, polar regions, and marine boundary layer (Lindberg et al., 2002; Swartzendruber et al., 2006; Fain et al., 2009; Lyman and Jaffe, 2012; Timonen et al., 2013), the diel pattern of GOM at the remote sites in China suggests that photochemical oxidation may also play an important role in the formation of GOM in the continental boundary layer.

3.6 Long-term trend of GEM

Gradual decreases in GEM concentrations in South Africa, Europe, and North America have been well documented (Slemr and Scheel, 1998; Slemr et al., 2003, 2011; Ebinghaus et al., 2002; Cole et al., 2013). Slemr et al. (2011) showed that GEM concentrations decreased from ~ 1.35 in 1996 to ~ 0.9 ng m^{-3} in 2008 at Cape Point, South Africa, and from ~ 1.75 in 1996 to ~ 1.4 ng m^{-3} in 2009 at Mace Head, Europe (a decrease of $0.024\text{--}0.035$ $\text{ng m}^{-3} \text{yr}^{-1}$ or $1.4\text{--}2.7\%$ yr^{-1}). The 10-year trends of GEM concentrations at five ground-based sites in the Arctic and Canada also showed a decreasing trend at a rate of $0.013\text{--}0.035$ $\text{ng m}^{-3} \text{yr}^{-1}$ ($0.9\text{--}2.2\%$ yr^{-1} ; Cole et al., 2013). Such a decrease is consistent with the decrease in anthropogenic Hg emissions in Europe and North America (Pacyna et al., 2006; Streets et al., 2011; AMAP/UNEP, 2013). The available GEM data in China are not sufficient to conclude a long-term concentration trend. However, a preliminary assessment can be obtained using monitoring data that spanned up to 7–9 years. For example, Fu and Feng (2015) found that the annual mean GEM concentration in Guiyang, southwestern China, increased from 8.40 ng m^{-3} (geomean = 7.46 ng m^{-3}) in 2002 to 10.2 ng m^{-3} (geomean = 8.88 ng m^{-3}) in 2010. Assuming a linear increasing rate, a mean annual rate of 0.16 $\text{ng m}^{-3} \text{yr}^{-1}$ (2.5% yr^{-1}) was found. The increase in GEM in Guiyang was mainly observed during cold seasons (paired samples test: $p < 0.05$ during the cold season vs. $p = 0.82$ during the warm season), when the impact of long-

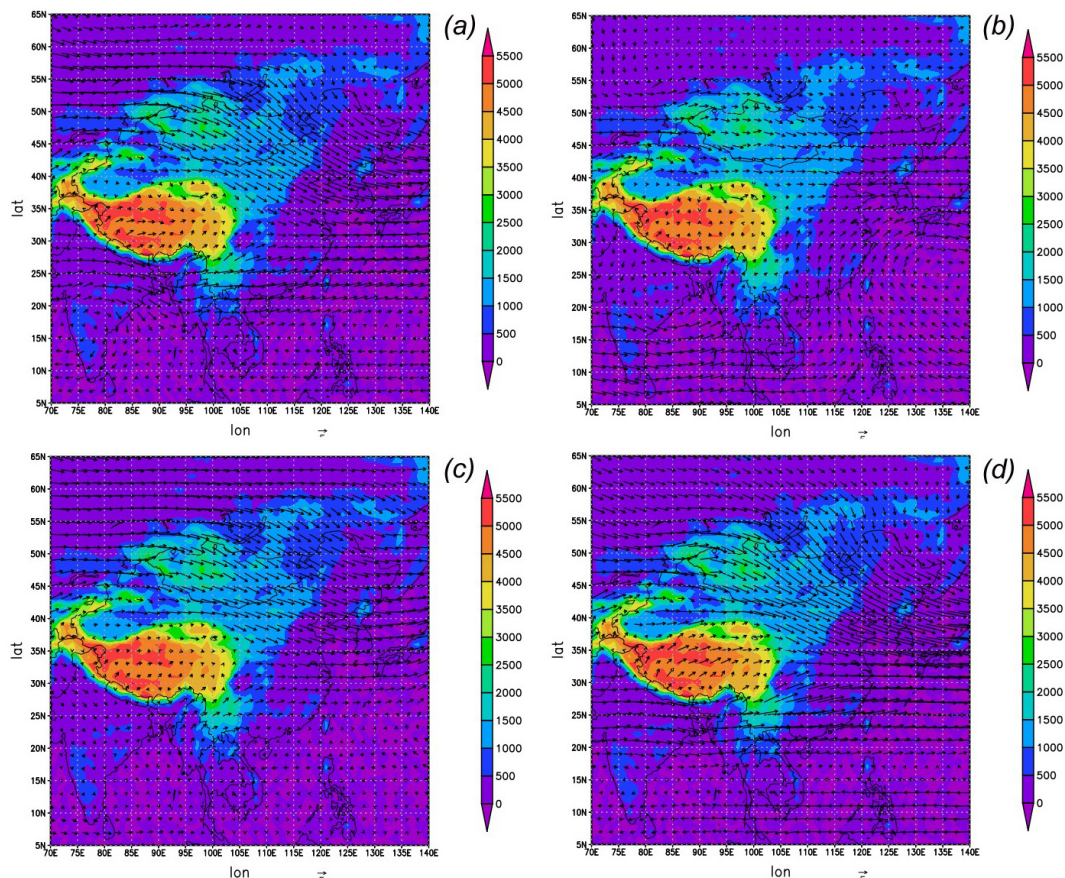


Figure 6. Wind field at 650 hPa and geopotential height from 2011 to 2013: (a) spring, (b) summer, (c) fall and (d) winter. It was calculated using the Grid Analysis and Display System (GADS) and gridded meteorological data ($1.0^\circ \times 1.0^\circ$) from the NCEP FNL Operational Global Analysis system.

range transport from major source regions of China played a more important role. The measurements at Mt. Changbai, northeastern China (unpublished data presented in the 12th ICMGP in Jeju, Korea, 2015) showed that the annual mean GEM concentrations were increasing during the 2009–2013 period with a mean of $1.75 \pm 0.09 \text{ ng m}^{-3}$ in 2013 compared to a mean of $1.55 \pm 0.31 \text{ ng m}^{-3}$ in 2009 (paired samples test: $p < 0.01$), corresponding to an annual increase of $0.04 \text{ ng m}^{-3} \text{ yr}^{-1}$ ($2.4 \% \text{ yr}^{-1}$; Fig. 9). The increase in atmospheric GEM in Guiyang and Mt. Changbai is consistent with the increasing anthropogenic Hg emissions ($4.2 \% \text{ per year}$) in China (L. Zhang et al., 2015). After 2013, GEM concentrations at Mt. Changbai showed a slight decrease in 2014 and 2015. The long-term trend should be further evaluated using continuous monitoring data.

3.7 Inferences on Hg emissions in China

Field measurements of atmospheric Hg have been applied to constrain Hg emissions. Pan et al. (2007) used a GEM concentration measured onboard aircrafts and an inverse modeling method to estimate Hg emissions in China, which es-

timated an annual GEM emission of 1140 t in China, ~ 3.4 times greater than the GEM emission estimated using activity statistics in China for 1999 (340 t). Using GEM observations at 27 remote sites globally (of which 4 sites are in mainland China) and GEOS-Chem inverse modeling, Song et al. (2015) predicted a total GEM emission of 1180–2030 t (median = 1550 t) from Asia, of which 650–1770 t (median = 1210 t) were from anthropogenic sources. Estimates of GEM emission in China have also been made using correlation analysis using observed GEM and trace gas concentrations. This approach is based on the measured ratios of GEM to trace gases in plumes originating from China and verified inventories of trace gases in China. Table 2 summarizes the observed GEM / CO ratios in China. The GEM / CO correlation slopes varied slightly in different years without systematic changes. The variations of GEM / CO correlation slopes observed in different years and at the receptors may partially reflect spatial and seasonal patterns. Using published GEM / CO correlation slopes and CO emissions in China, the GEM emissions in China over the past decade were estimated to range from 632 to 1138 t (Table 2), consistent with the re-

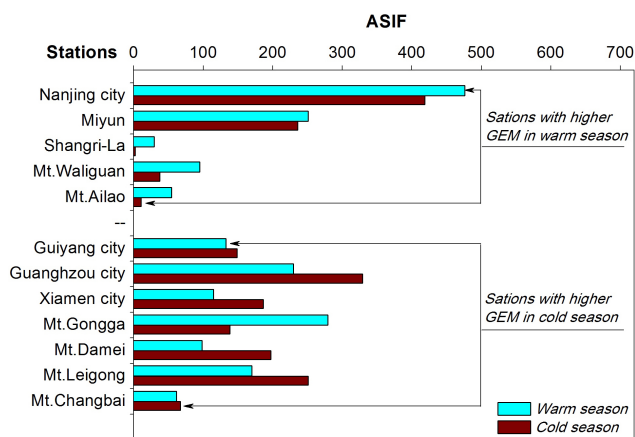


Figure 7. Dimensionless anthropogenic Hg^0 sources impacted factor (ASIF) at Chinese ground-based stations in cold and warm seasons. ASIF is defined as $\text{ASIF} = \frac{\sum N_{ij} \times \text{Hg}_{ij}^0}{\sum N_{ij}}$, where N_{ij} is the number of trajectory endpoints in a $0.5^\circ \times 0.5^\circ$ grid cell (i, j) during the sampling period, and Hg_{ij}^0 is the total anthropogenic Hg^0 emissions in a $0.5^\circ \times 0.5^\circ$ grid cell (i, j). The source region domain contains 20×20 grid cells at a spatial resolution of $0.5^\circ \times 0.5^\circ$. The ground-based stations (receptors) are located in the center of the domain. The 5-day backward trajectory that ended at a height of ~ 1000 m a.g.l. was utilized for the calculation of ASIF.

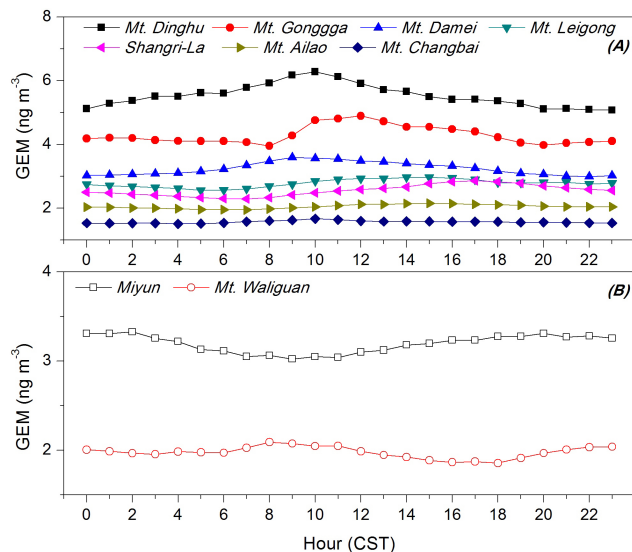


Figure 8. Diel trends of GEM concentrations at the remote sites in China. The diel trends at the remote sites in China were calculated on the basis of at least 1 year of continuous measurements (the time schedule for each of the sites was shown in Table 1).

sults derived from inverse modeling (Pan et al., 2006; Song et al., 2015). An overall increase in GEM emission from 2001 to 2009 was evident because of the increasing GEM/CO correlation slopes and CO emissions. CO emissions in China are not available from 2011 to 2012 and the estimated GEM

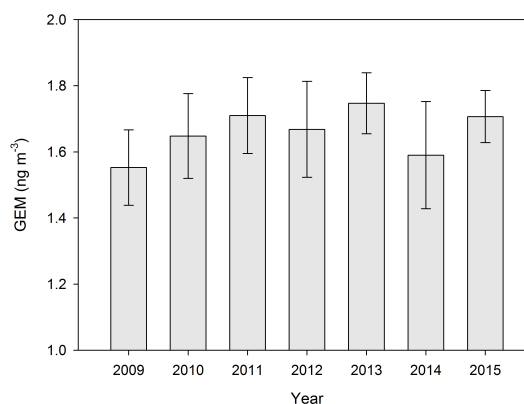


Figure 9. Annual mean GEM concentrations at Mt. Changbai in northeastern China.

emissions in the 2 years were based on CO emissions in 2010 (Tohjima et al., 2014), which could result in an underestimate of GEM. Considering the agreement between inverse modeling and correlation estimates, it is likely that GEM emissions in China are larger than the values estimated by activity statistics ($312\text{--}430 \text{ t yr}^{-1}$; AMAP/UNEP, 2013; L. Zhang et al., 2015). Recent field measurements of Hg emission factors for large point sources have reduced the uncertainties in Hg emission estimates (Wang et al., 2014). However, knowledge gaps in the emission estimates in China remain for the source categories whose emission factors have not been verified. Hg emissions related to the production, consumption and recycling of Hg in artisanal mercury and gold production, polyvinyl chloride (PVC) production, mercury batteries, mercury sphygmomanometers, and mercury thermometers, are potentially large Hg sources in China but have not been well evaluated (Li et al., 2009; Ren et al., 2014; Minshwaner et al., 2010). Other reasons causing earlier underestimation include natural emissions and conversion of GOM and/or PBM to GEM in the stack or in the plume during transit (Jaffe et al., 2005).

3.8 Long-range transport of atmospheric Hg as a result of Hg emissions and monsoon climate

Anthropogenic Hg emissions in China are mainly from industrial and urban sources (D. Y. Wang et al., 2014). These sources not only cause highly elevated GEM concentrations locally, but also impact the GEM concentrations in remote areas via long-range transport. Fig. 10 shows the source regions of GEM at Mt. Waliguan (northwestern China), Mt. Changbai (northeastern China), Mt. Leigong (southwestern China) and Mt. Damei (eastern China) indicated by the potential source contribution function (PSCF). The source regions of Mt. Waliguan were industrialized and urbanized areas in northwestern China 100–400 km to the east of the sampling site and northern India (Fu et al., 2012a). The source regions of Mt. Changbai were the megacities of Beijing, Tianjin, and

Table 2. GEM / CO ratios in air plumes and estimated GEM emissions in China.

Source region	Year	Receptor	GEM / CO ratio (pg m ⁻³ /ppb)	CO emission (×10 ⁶ t yr ⁻¹); Tohjima et al. (2014)	Estimated GEM emission (t yr ⁻¹)	Reference
Eastern, northern, and northeastern China	2001	ACE-Asia flight	6.8 ± 1.2 (<i>n</i> = 16)	140	763	Friedli et al. (2004); Pan et al. (2006)
Eastern China	2004	Hedo station, Japan and MBO, USA	4.4 ± 1.4 (<i>n</i> = 10)	179	632	Jaffe et al. (2005); Weiss-Penzias et al. (2007)
Eastern and southern China	2005	MBO, USA and CARIBIC flights	7.1 ± 1.5 (<i>n</i> = 3)	182	1036	Weiss-Penzias et al. (2007); Slemr et al. (2014)
Northwestern, southwestern, and southern China	2006	CARIBIC flights	7.6 ± 6.8 (<i>n</i> = 16)	176	1073	Slemr et al. (2009)
Northwestern, southwestern, and southern China	2007	CARIBIC flights	8.4 ± 3.2 (<i>n</i> = 9)	169	1138	Slemr et al. (2009); Sheu et al. (2010)
Northwestern China	2009	Mt. Waliguan, China	9.4 ± 3.5 (<i>n</i> = 20)	150	1130	Fu et al. (2015)
Southwestern and southern China	2011	Mt. Ailao, China	6.4 ± 6.1 (<i>n</i> = 8)	159	812	Fu et al. (2015)
Southwestern and southern China	2012	Mt. Ailao, China	6.2 ± 2.0 (<i>n</i> = 7)	159	791	Fu et al. (2015)

Shijiazhuang (Fu et al., 2012b), similar to the source regions of Miyun, northern China (Zhang et al., 2013). The source regions of Mt. Leigong included Chongqing, Guiyang, and Kunming (three major cities in southwestern China) as well as northern Myanmar, where forest biomass burning released a large quantity of air pollutants to the atmosphere (Streets et al., 2003). The source regions of GEM at Mt. Damei were central Anhui Province and western Jiangshu Province, both being important industrial regions in eastern China (Yu et al., 2015). These source regions corresponded to the locations of Hg emission sources in China and other Asian countries (Fig. 1). These results elucidated that the dominant transport routes of GEM are highly related to dominant wind directions governed by the monsoons. On the other hand, local emission sources played a more predominant role in urban areas (Fu et al., 2011a; Zhu et al., 2012).

Air outflows from mainland China may also impact GEM concentrations in other regions. For instance, measurements of atmospheric GEM in Seoul, Korea, showed that approximately 79 % of the high-GEM events were related to air masses originating from China (Choi et al., 2009). Flight measurements in the free troposphere over the Asia Pacific region also showed evidence of GEM outflow (Friedli et al., 2004). GEM emitted in China and East Asian countries can undergo intercontinental transport as suggested by the GEM / CO ratio measured at Mt. Bachelor Observatory (MBO) in the Pacific Northwest of the USA (Jaffe et al., 2005; Weiss-Penzias et al., 2006; Strode et al., 2008; Timonen et al., 2013). The trans-Pacific transport of Hg mainly occurred in spring and episodically resulted in a 10–40 % increase in GEM at the monitoring sites in North America (Jaffe et al., 2005; Weiss-Penzias et al., 2006; Durnford et al., 2010).

3.9 Atmospheric Hg deposition in China

The deposition fluxes of THg and methyl mercury (MeHg) via precipitation, throughfall, and litterfall measured in China are presented in Table 3. The mean THg and MeHg concentrations in rainwater collected by wet-only precipitation collectors at remote sites ranged from 3.0 to 4.8 ng L⁻¹ (mean = 4.0 ng L⁻¹, *n* = 4) and from 0.03 to 0.04 ng L⁻¹ (mean = 0.04 ng L⁻¹, *n* = 2), respectively. Such aqueous THg and MeHg concentrations were much higher in urban areas, with mean values in the ranges of 13.3–52.9 ng L⁻¹ (mean = 31.2 ng L⁻¹, *n* = 5) and 0.05–0.31 ng L⁻¹ (mean = 0.18 ng L⁻¹, *n* = 2), respectively. The THg and MeHg concentrations in rainwater collected by bulk precipitation collectors at remote sites of China were much higher than those using wet-only precipitation collectors (Table 3). Some of the bulk precipitation samples were collected at locations close to anthropogenic Hg sources and on a monthly basis (Guo et al., 2008; Fu et al., 2008b; Wan et al., 2009b), and therefore dry deposition of TPM and GOM can also contribute to the measured wet deposition. Wet-only deposition fluxes of THg and MeHg were 1.8–7.0 μg m⁻² yr⁻¹ (mean = 5.1 μg m⁻² yr⁻¹, *n* = 4) and 0.01–0.06 μg m⁻² yr⁻¹ (mean = 0.04 μg m⁻² yr⁻¹, *n* = 2) at remote sites, and 13.4–56.5 μg m⁻² yr⁻¹ (mean = 31.9 μg m⁻² yr⁻¹, *n* = 4) and 0.05–0.28 μg m⁻² yr⁻¹ (mean = 0.16 μg m⁻² yr⁻¹, *n* = 2) at urban sites, respectively (Table 3). Wet deposition fluxes of THg and MeHg at urban sites in China were higher compared to those in North America and Europe, but wet deposition fluxes of THg at remote sites were in the lower range of those observed in North America and Europe (Table 3).

Significant positive correlations were observed between rainwater THg concentrations and PBM concentrations ($R^2 = 0.66$, $p < 0.05$, $N = 7$) and between rainwater THg concentrations and GOM concentrations ($R^2 = 0.78$,

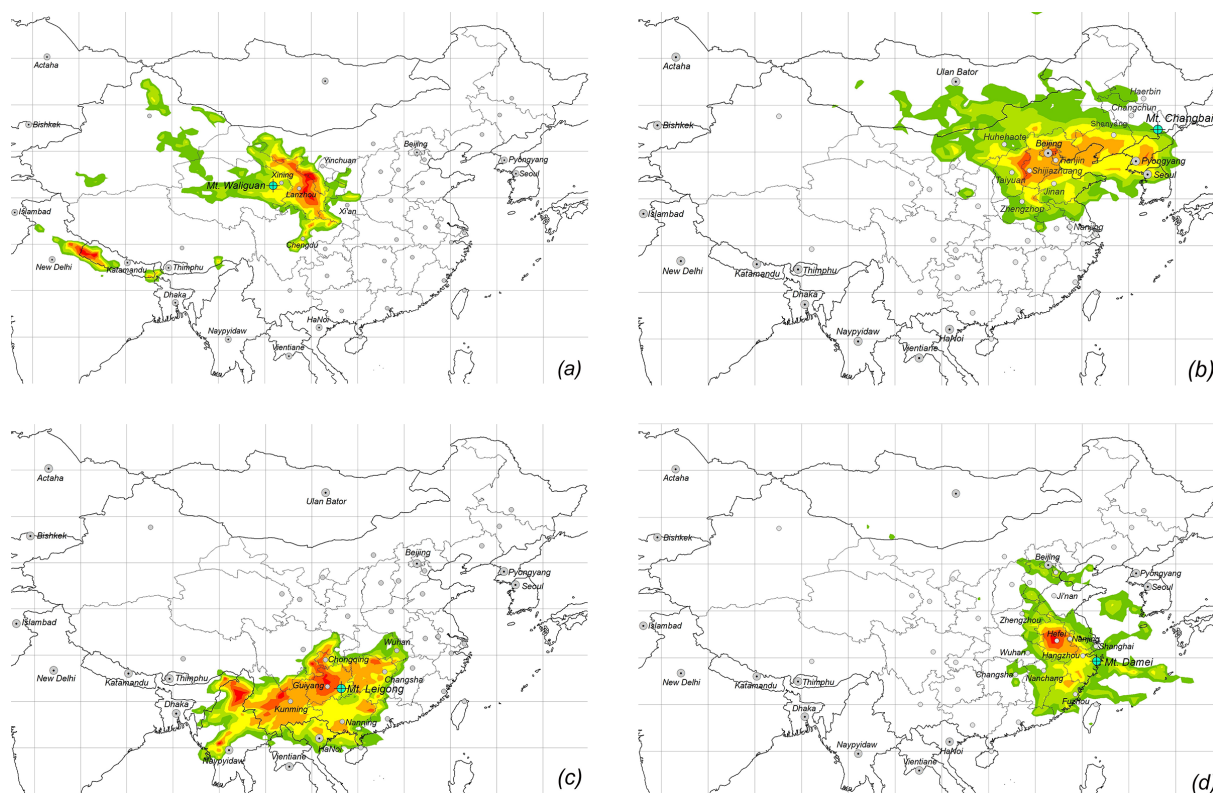


Figure 10. Potential source regions of GEM at (a) Mt. Waliguan, (b) Mt. Changbai, (c) Mt. Leigong, and (d) Mt. Damei (Fu et al., 2010b; Fu et al., 2012a; Fu et al., 2012b; Yu et al., 2015).

$p < 0.05$, $N = 6$), resulting in the positive correlations between wet deposition fluxes and PBM ($R^2 = 0.69$, $p < 0.05$, $N = 7$) and GOM concentrations ($R^2 = 0.92$, $p < 0.01$, $N = 6$). This indicates that washout of PBM and GOM during rain events contributes to Hg wet deposition in China. Such a contribution is particularly important in urban areas where PBM and GOM concentrations are highly elevated. In remote areas of China, however, washout of elevated atmospheric PBM does not seem to drive a notable increase in Hg wet deposition flux. This is likely because of the low washout rate of PBM during rain events (Lee et al., 2001; Seigneur et al., 2004). Also, most of the remote sites were at high altitude with low-level clouds, which reduced the contribution of Hg washout.

Study of dry deposition of Hg is of special importance for constructing the regional and global mass budget of Hg. Lin et al. (2010) simulated the dry deposition of Hg in East Asia using the CMAQ-Hg model. The results demonstrated a remarkable spatial distribution of dry deposition fluxes ranging from 10 to $150 \mu\text{g m}^{-2} \text{yr}^{-1}$. The greatest deposition fluxes were found to be related to emission sources in the major urban areas of China (Lin et al., 2010). Dry deposition of Hg was also estimated in forest areas of China by subtracting wet deposition flux measured in open areas from the sum of deposition via throughfall and lit-

terfall (Table 3). Using this approach, annual deposition flux in Tieshanping forest near the megacity of Chongqing was estimated to be $262 \mu\text{g m}^{-2} \text{yr}^{-1}$ (Wang et al., 2009), which is of the same order of magnitude compared to the model-predicted deposition. At the remote Mt. Gongga and Mt. Leigong forest in southwestern China, annual dry deposition fluxes reached 66.5 and $43.9 \mu\text{g m}^{-2} \text{yr}^{-1}$, respectively (Fu et al., 2010a, 2010b). Such dry deposition fluxes were 1.8–13.6 times the reported dry deposition fluxes in North America (range 13.3 – $34.0 \mu\text{g m}^{-2} \text{yr}^{-1}$, mean = $19.2 \mu\text{g m}^{-2} \text{yr}^{-1}$, $n = 5$) and Europe (range 25.3 – $36.0 \mu\text{g m}^{-2} \text{yr}^{-1}$, mean = $29.6 \mu\text{g m}^{-2} \text{yr}^{-1}$, $n = 5$) (Lee et al., 2000; Grigal et al., 2000; Rea et al., 2001; St Louis et al., 2001; Lindberg et al., 2007). The much greater dry deposition of Hg in China was likely due to the elevated GEM and PBM concentrations that enhance the accumulation of Hg in leaves and on foliage surfaces (St Louis et al., 2001; Frescholtz et al., 2003). Dry deposition contributes 72–90 % of total Hg deposition in forest areas of China, consistent with the results predicted by modeling studies that dry deposition is the predominant removal pathway of atmospheric Hg (Lin et al., 2010; L. Wang et al., 2014).

Table 3. Hg concentrations and deposition fluxes in precipitation, throughfall, and litterfall in China and other regions.

Site	Location			Type	Study period	Samples	Hg concentration		Deposition flux		References	
	Lon	Lat	Altitude (m a.s.l)				(ng L ⁻¹ or ng g ⁻¹)		($\mu\text{g m}^{-2}\text{ yr}^{-1}$)			
							THg	MeHg	THg	MeHg		
Mt. Ailao, Yunnan	101.02	24.53	2500	Remote	06/2011–05/2012	Precipitation Litterfall	3.0 54.0		5.4 71.2		Zhou et al. (2013)	
Mt. Leigong, Guizhou	108.20	26.39	2178	Remote	05/2008–05/2009	Precipitation Throughfall Litterfall	4.0 8.9 91.0	0.04 0.1 0.48	6.1 10.5 39.5	0.06 0.12 0.28		Fu et al. (2010b)
Mt. Damei, Zhejiang	121.57	29.63	550	Remote	08/2012–07/2013	Precipitation Litterfall	4.1 46.6		7.0 26.0			Lang (2014)
Nam Co, Tibet	90.99	30.77	4730	Remote	07/2009–07/2011	Precipitation	4.8	0.03	1.75	0.01		Huang et al. (2012)
Mt. Gongga ¹ , Sichuan	102.12	29.65	1640	Remote	01–12/2006	Precipitation*	9.9		9.1			Fu et al. (2008b)
Mt. Gongga ² , Sichuan	101.93	29.58	3000	Remote	05/2005–04/2005	Precipitation* Throughfall Litterfall	14.2 40.2 35.7	0.16 0.3	26.1 57.1 35.5	0.30 0.43		Fu et al. (2010a)
Mt. Changbai, Jilin	128.47	42.40	750	Remote	08/2005–07/2006	Precipitation*	13.4		8.4			Wan et al. (2009a)
Puding, Guizhou	105.80	26.37	1145	Remote	08/2005–07/2006	Precipitation*	20.6	0.18	24.8	0.22		Guo et al. (2008)
Hongjiadu, Guizhou	105.85	26.88	1130	Remote	08/2005–07/2006	Precipitation*	39.4	0.18	34.7	0.16		Guo et al. (2008)
Yinzidu, Guizhou	106.12	26.57	1088	Remote	08/2005–07/2006	Precipitation*	35.7	0.18	38.1	0.19		Guo et al. (2008)
Dongfeng, Guizhou	106.13	26.85	970	Remote	08/2005–07/2006	Precipitation*	37.4	0.20	36.3	0.19		Guo et al. (2008)
Wujiangdu, Guizhou	106.77	27.32		Remote	08/2005–07/2006	Precipitation*	57.1	0.25	39.6	0.17		Guo et al. (2008)
Guiyang	106.72	26.57	1040	Urban	07–09/2008	Precipitation	13.3	0.05	13.4	0.05		Liu et al. (2011)
Xiamen	118.31	24.60	50	Urban	07/2013–02/2014	Precipitation	26.6		30.4			Wu (2014)
Chongqing				Urban	06/2010–06/2011	Precipitation	30.7	0.31	28.7	0.28		Wang et al. (2012); Y. M. Wang et al. (2014)
Tieshanping, Chongqing	104.68	29.63	500	Urban	03/2005–03/2006	Precipitation Throughfall Litterfall	32.3 69.7 105		29.0 71.3 220			Wang et al. (2009)
Nanjing	118.78	32.05	100	Urban	06/2011–02/2012	Precipitation	52.9		56.5			Zhu et al. (2014)
North America				Urban		Precipitation	7.8–15.0		8.4–17.2			Mason et al. (2000); Keeler et al. (2005); Prestbo and Gay (2009)
				Remote		Precipitation Throughfall Litterfall	4.9–22.4 6.6–20.7 32.0–57.0	0.12 0.22 0.06–0.55	2.5–21.5 1.6–12.0 7.2–15.0	0.09 0.09 0.01–0.24		Rea et al. (2001); St Louis et al. (2001); Demers et al. (2007); Choi et al. (2008); Prestbo and Gay (2009); Fisher and Wolfe (2012)
Europe				Remote		Precipitation Throughfall Litterfall	11.9–18.0 22.8–29.0 28.0–68.0		7.0–36.0 15.0–39.0 4.0–32.5			Munthe et al. (1995); Lee et al. (2000); Schwesig and Matzner (2000); Wangberg et al. (2007); Larssen et al. (2008)

(Precipitation: * indicates bulk precipitation and the rest indicate wet-only precipitation. Mt. Gongga: ¹ elevation of the sampling site was 1600 m above sea level.

² Elevation of the sampling site was 3000 m above sea level).

3.10 Limitations, implications and research needs

Although the methods used in quantifying mercury concentrations and forms have significant uncertainties as discussed in Sect. 2.2, the relative magnitudes of GEM and PBM concentrations measured using similar analytical protocols (Tekran speciation method) at remote and urban sites in China were substantially higher than the background concentrations in the Northern Hemisphere, which provide evidence of the influence of anthropogenic and natural Hg emissions in China and neighboring countries. Many issues and knowledge gaps exist and deserve further study.

First, there is no coordinated observational network for atmospheric Hg in China. In North America and Europe, much effort has been made to establish global or continental observational networks including the Global Mercury Observation System (GMOS), American Atmospheric Mercury Network (AMNet), Canadian Atmospheric Mercury Network (CAMNet) and Mercury Deposition Network (MDN). The network

observations have been integrated with modeling studies, leading to improved understanding of Hg fate and transport.

The geographical differences in atmospheric compositions (e.g., different levels of air pollutants) provide an opportunity to investigate Hg behavior in air under different atmospheric conditions. Air pollution has become a major problem in China. Previous studies have shown that the concentrations of aerosols, sulfur dioxide (SO₂), nitrogen dioxide (NO₂) and ozone in the atmosphere were higher in China relative to other regions in the Northern Hemisphere (Burrows et al., 1999; Savage et al., 2004; van Donkelaar et al., 2010). These pollutants play important roles in the physicochemical processes in the atmosphere and can affect the fate of atmospheric Hg. Previous studies have shown that gas-particle partitioning of atmospheric GOM and heterogeneous oxidation of GEM at aerosol surfaces facilitate PBM formation (Lindberg et al., 2002; Sprovieri et al., 2005; Amos et al., 2012), which may deplete GEM and GOM and contribute to the elevated PBM concentrations in China. However, there were a few studies that provided detailed assessments of the

effect of aerosol and other air pollutants on the transformation and wet and dry deposition of atmospheric Hg in China.

GEM concentrations in China are highly elevated and can enhance the uptake of GEM by vegetation. Such uptake may represent an important sink of atmospheric Hg in China. Deposition fluxes through litterfall and throughfall at several forest sites in China were found to be much higher than magnitudes in North America and Europe. There is a growing amount of evidence that Hg found in vegetation biomass does not likely result from the uptake in the root zone followed by upward translocation to stem and leaves (Frescholtz et al., 2003; Cui et al., 2014), which points to the importance of atmospheric uptake in Hg accumulation in vegetative biomass. We therefore hypothesize that the high Hg deposition through litterfall and throughfall is a result of the elevated atmospheric GEM and PBM concentrations that drive the uptake and accumulation of Hg in vegetative biomass, and that Hg uptake by vegetation is a major sink for removing different Hg forms from the atmosphere in China. One area of data deficiency is the lack of deposition data over other vegetation coverages such as grass and crop. Garland and cropland are dominant terrestrial ecosystems in China that cover 3.9×10^6 and 1.6×10^6 km² of land area, respectively. It is therefore needed to quantify their roles in the removal of atmospheric Hg in China.

As a persistent air pollutant, the outflow of GEM in China will have an impact on the global Hg concentration and deposition. Signals of Hg emissions in China have been detected at receptors in Korea, Japan, the Pacific Northwest and western North America (Weiss-Penzias et al., 2007; Swartzendruber et al., 2008; Jaffe et al., 2005; Nguyen et al., 2010). Model results also suggest that Hg emissions in Asia enhanced Hg concentration and deposition elsewhere (Seigneur et al., 2004; Durnford et al., 2010). As more measurement data in China become available, further research efforts should be made to reconcile the observational and model results. Although earlier modeling studies have utilized Hg observations in China and other Asian regions to construct the mass budget for estimate of Hg outflow from East Asia, there is a systematic discrepancy in the magnitude and distribution of concentration and dry deposition between the observational and model results (Lin et al., 2010; L. Wang et al., 2014; Song et al., 2015). Such mismatch may be caused by a combination of reasons including the uncertain Hg emission estimate and the incomplete understanding of physical and chemical transformation, wet and dry deposition of atmospheric Hg in China. Atmospheric composition in China is different from that of a typical airshed because of the heavy emission loading of PM, SO₂, NO_x, and ozone precursors, which may result in complex atmospheric processes that are not yet understood. Also, Hg dry deposition to vegetation was enhanced due to the generally elevated GEM concentrations, and such an enhanced uptake process has not been implemented in atmospheric Hg models. More studies should focus on the transformation and removal of at-

mospheric Hg in China, which will help understand the impact of Hg emission in China on global Hg pollution.

Continuous, long-term observations of atmospheric Hg in China should be carried out to assess the temporal change of the atmospheric Hg burden. Preliminary assessment of GEM concentrations in China points to an increase corresponding to the increase in the regional emission inventory. This finding is in contrast to most observations in remote sites in the Northern Hemisphere (Slemr et al., 2011; Cole et al., 2013). More data that allow analysis of the long-term trend of GEM in China and other Asian countries are needed because (1) the long-term trend of GEM concentrations may not share a similar pattern throughout the Northern Hemisphere, and (2) regional emission influence may exceed the influence of atmospheric circulation of global emissions. In addition, the observed long-term trend should be carefully verified with regional and global modeling assessment to understand the role of Hg emission in China in the global background of atmospheric Hg. Furthermore, the impact of ongoing efforts of Hg emission reduction in China caused by the co-benefits of air pollution control devices as well as targeted Hg emission control, in response to the adoption of the Minamata Convention on Mercury, should be thoroughly investigated to better understand the associated benefit to global Hg biogeochemical cycling.

4 Conclusions

As the largest atmospheric Hg source region in the world, much attention has been paid to understanding the characteristics and behavior of atmospheric Hg in China. This paper provides an integrated review of these understandings presented in peer-reviewed literature. GEM and PBM have been found to be substantially elevated in both remote and urban areas of China compared to those observed in North America and Europe. A strong spatial variation in GEM and PBM concentrations was observed, with high concentrations related to regional anthropogenic Hg emissions in central and eastern China. Annual GEM emission in China was estimated to be 632–1138 t over the past decade using reported GEM/CO correlation slopes and emissions of CO in China, agreeing with the results of reverse modeling but higher than published values estimated from emission activity statistics. Existing field measurements suggested an increase in GEM concentrations at a remote site and an urban site in China over recent years, opposite to reported long-term trend in Europe and North America. The increasing trend in China is possibly caused by the increase in anthropogenic Hg emissions in the past decade, and indicates that the influence of regional emissions on GEM levels in China exceed global emission influence. GEM concentrations in China displayed two distinct seasonal patterns driven by monsoonal climate that governs the transport of anthropogenic Hg emissions in China and in neighboring countries. Wet deposition fluxes of Hg in remote

areas of China were not significantly higher than the values observed in the rest of the world, while the dry deposition as measured through litterfall and by model simulation was highly elevated compared to those observed in Europe and North America. This indicates that dry deposition of Hg is likely the dominant pathway for removing atmospheric Hg in China. However, further studies are needed to better quantify dry deposition of atmospheric Hg and to improve our understanding of the atmospheric Hg budget in China, as well as to elucidate the impact of Hg emission changes on global Hg pollution.

Acknowledgements. This work was funded by the National 973 Program of China (2013CB430003) and the National Science Foundation of China (41473025, 41273145, 41428301). We also acknowledge Lei Zhang, Shuxiao Wang, Tijian Wang, and Laiguo Chen for providing the diel and monthly mean atmospheric gaseous mercury concentrations at Miyun, Nanjing, Mt. Dinghu, and Guangzhou.

Edited by: R. Ebinghaus

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