

Five-year records of ~~Total~~ Mercury wet Deposition flux at GMOS sites in the Northern and Southern Hemispheres

Francesca Sprovieri¹, Nicola Pirrone², Mariantonia Bencardino¹, Francesco D'Amore¹, Helene Angot^{3,4}, Carlo Barbante^{17,10}, Ernst-Günther Brunke⁵, Flor Arcega-Cabrera¹⁵, Warren Cairns¹⁷, Sara Comero⁸, María del Carmen Diéguez⁷, Aurélien Dommergue^{3,4}, Ralf Ebinghaus⁶, Xin Bin Feng¹², Xuewu Fu¹², Patricia Elizabeth Garcia⁷, Bernd Manfred Gawlik⁸, Ulla Hageström⁹, Katarina Hansson⁹, Milena Horvat¹¹, Jože Kotnik¹¹, Casper Labuschagne⁵, Olivier Magand^{4,3}, Lynwill Martin⁵, Nikolay Mashyanov¹³, Thumeka Mkololo⁵, John Munthe⁹, Vladimir Obolkin¹⁶, Martha Ramirez Islas¹⁴, Fabrizio Sena⁸, Vernon Somerset⁵, Pia Spandow⁹, Massimiliano Varde^{1,17}, Chavon Walters⁵, Ingvar Wängberg⁹, Andreas Weigelt⁶, Xu Yang¹², and Hui Zhang¹²

¹CNR Institute of Atmospheric Pollution Research, Rende, Italy

²CNR Institute of Atmospheric Pollution Research, Rome, Italy

³Univ. Grenoble Alpes, Laboratoire de Glaciologie et Géophysique de l'Environnement, Grenoble, France

⁴CNRS, Laboratoire de Glaciologie et Géophysique de l'Environnement, Grenoble, France

⁵Cape Point GAW Station, Climate and Environ. Research & Monitoring, South African Weather Service

⁶Helmholtz-Zentrum Geesthacht, Germany

⁷INIBIOMA-CONICET-UNComa, Bariloche, Argentina

⁸Joint Research Centre, Italy

⁹IVL, Swedish Environmental Research Inst. Ltd., Sweden

¹⁰University Ca' Foscari of Venice, Italy

¹¹Jožef Stefan Institute, Ljubljana, Slovenia

¹²State Key Laboratory of Environmental Geochemistry, Inst. of Geochemistry, Chinese Academy of Sciences

¹³St. Petersburg State University, Russia

¹⁴Instituto Nacional de Ecología y Cambio Climático (INECC), Mexico

¹⁵Universidad Nacional Autónoma de México (UNAM), Unidad de Química, Sisal, Mexico

¹⁶Limnological Institute SB RAS, Irkutsk, Russia

¹⁷CNR Institute for the Dynamics of Environmental Processes, Venice, Italy

Correspondence to: Francesca Sprovieri (f.sprovieri@iia.cnr.it)

Abstract. The atmospheric deposition of mercury (Hg) occurs via several mechanisms including dry and wet scavenging by precipitation events. In an effort to understand the atmospheric cycling and seasonal depositional characteristics of Hg, wet deposition samples were collected for approximately five years at 17 selected GMOS monitoring sites located in the Northern and Southern Hemispheres in the framework of the Global Mercury Observation System (GMOS) project. Total mercury (THg) exhibited annual and seasonal patterns in Hg wet deposition samples. Inter-annual differences in total wet deposition are mostly linked with precipitation volume, with the greatest deposition flux occurring in the wettest years. This data set provides a new insight into baseline concentrations of THg concentrations in precipitation worldwide, particularly in regions, such as the Southern Hemisphere and tropical areas where wet deposition as well as atmospheric Hg species were not investigated before, opening the way for future and additional simultaneous measurements across the GMOS network as well as new findings in future modeling studies.

1 Introduction

Mercury (Hg) is a persistent pollutant of global concern due to its toxicity and its capacity to bioaccumulate aquatic food chains with serious consequences on human and wildlife health (Driscoll et al., 2013). Long-range atmospheric transport is the main pathway for contamination of remote ecosystems, therefore atmospheric deposition is the primary indicator for the understanding of its impact on aquatic and terrestrial ecosystems (Schroeder and Munthe, 1998; Lindberg et al., 2002). Hg exists in the atmosphere mainly in three operationally defined forms: gaseous elemental mercury (GEM), oxidized gaseous mercury (GOM), and particulate bound mercury (PBM). Globally, GEM is the predominant form whereas GOM and PBM are thought to be rapidly dry deposited and wet scavenged by precipitation (Lindberg et al., 2007). Due to the current Currently, Hg dry deposition is often estimated by models using ambient concentrations of Hg measurements and meteorological parameters due to the lack of existing direct and accurate measurements of Hg dry deposition (Gustin et al., 2012; Zhang et al., 2012); (Gustin et al., 2012; Zhang et al., 2012), therefore the investigation of Hg fluxes to terrestrial and aquatic surfaces in different part of the world are mainly performed by wet deposition measurements (Gratz et al., 2009; Feng et al., 2009). Hg wet deposition represents the air-to-surface flux in precipitation (Lindberg et al., 2007). Previous studies suggested that the magnitude of Hg wet deposition varies geographically and seasonally due to climatic conditions, atmospheric chemistry, and human influences i.e. emissions of Hg from anthropogenic sources (Vanarsdale et al., 2005; Selin and Jacob, 2008; Prestbo and Gay, 2009). Current annual atmospheric deposition of Hg has been estimated to be 3200 Mg y^{-1} deposited on land and 3700 Mg y^{-1} into oceans (Mason et al., 2012). The preindustrial deposition rate has been estimated to be 1000 Mg y^{-1} deposited on land and 2500 Mg y^{-1} into oceans (Selin, 2009). Developed countries in North America and Europe have reduced their anthropogenic Hg use and emissions (Hylander, 2001), but Hg use and emission are still occurring widely around the world (Pacyna et al., 2010; Pirrone et al., 2010). In North America seasonal patterns in wet deposition are observed in both depositional flux and concentration THg concentrations in precipitation and Hg wet deposition amounts have been observed with the highest values in the summer and lowest values in winter (Pacyna et al., 2010; Mason et al., 2000); (Keeler et al., 2005; Choi et al., 2008; Prestbo and Gay, 2009); (Mason et al., 2000; Keeler et al., 2005; Choi et al., 2008; Prestbo and Gay, 2009; Pacyna et al., 2010). Explanations for this observation include more effective Hg scavenging by rain compared to snow (Keeler et al., 2005; Selin and Jacob, 2008), and a greater availability of soluble Hg due to convective transport in summer events (Keeler et al., 2005; Strode et al., 2007, 2008). Geographic differences in Hg wet deposition may be explained in part by the proximity to atmospheric sources. Results from the National Atmospheric Deposition Program's (NADP) Mercury Deposition Network (MDN) sites in the Northeastern United States exhibit a geographic trend with southern and coastal sites receiving higher Hg concentrations and depositional in precipitation and wet deposition fluxes (Vanarsdale et al., 2005; Prestbo and Gay, 2009) due to their location nearer to the East coast megalopolis and downwind of anthropogenic emission sources such as coal burning power plants and waste incinerators. In addition, gaseous evasion of Hg from marine waters is a significant global source of atmospheric Hg and GEM which throughout active oxidation processes may also contribute to elevated depositional fluxes in coastal regions (Mason and Sheu, 2002). A similar pattern exists in northern Europe with a clear gradient in atmospheric concentrations and deposition (Munthe et al., 2003) (Munthe et al., 2003, 2007; Sprovieri et al., 2016). Hg wet deposition data are therefore im-

Table 1. Key information on Station locations that are part of the 17-GMOS monitoring network and general characteristics of the sites (i.e., code, name, country, latitude, longitude, elevation), including the years of sampling as well as the type of monitoring stations in respect to the Hg measurements carried out as speciated (M) or not (S). (M/S or S/M = change of the site from Master to Secondary (or reverse)). In bold, external GMOS partners are indicated

	Code	Name	Country	Lat	Lon	Elev. Elev. (m a.s.l.)	Collector Collector-Type	years Type* of sampling	Type*	
Northern Hemisphere	H1	NY	Ny-lesund	Norway	78.90	11.88	+212	bulk-modified IVL-bulk	M 2012-2015	M
	2	PAL	Pallas	Finland	68,00	24,24	340	bulk-modified IVL-bulk	2011-2014	S
	3	RAO	Råö	Sweden	57,39	11,91	5	bulk-modified IVL-bulk	2011-2014	M
	4	MHE	Mace Head	Ireland	53,33	-9,91	5	wet-only	2012-2014	S
	5	LIS	Listvyanka	Russia	51,85	104,89	670	wet-only	2012-2013	S
	6	CMA	Col Margherita	Italy	46,37	11,79	2545	bulk-modified IVL-bulk	2014	S
	7	ISK	Iskrba	Slovenia	45,56	14,86	520	wet-only	2011-2015	M
	8	MCH	Mt. Changbai	China	42,40	128,11	741	wet-only	2011-2014	M/S
	9	LON	Longobucco	Italy	39,39	16,61	1379	wet-only	2012-2013	M
	10	MWA	Mt. Waliguan	China	36,29	100,90	3816	wet-only	2012-2014	M
	11	MAL	Mt. Ailao	China	24,54	101,03	2503	wet-only	2011-2014	S/M
Tropics	12	SIS	Sisal	Mexico	21,16	-90,05	7	wet-only	2013-2014	S
	13	CST	Celestún	Mexico	20,86	-90,38	3	wet-only	2012-2013	S
Southern Hemisphere	14	AMS	Amsterdam Island	TAAF	-37,80	77,55	70	wet-only	2013-2014	M
	15	CPT	Cape Point	South Africa	-34,35	18,49	230	wet-only	2011-2015	S
	16	CGR	Cape Grim	Australia	-40,68	144,69	94	bulk-modified IVL-bulk	2013-2015	S
	17	BAR	Bariuloche	Argentina	-41,13	-71,42	801	wet-only	2014-2015	M

* M=Master; S= Secondary

portant for verifying atmospheric models, understanding the biogeochemical cycling of Hg on a regional/global scale, and investigating ecosystem impacts. Regional monitoring networks with properly chosen monitoring sites can provide accurate estimates of wet deposition at regional scales. Long-term Hg wet deposition measurements exist at many locations within already established regional network, such as in the United States as part of the MDN or in Europe as part of the EMEP program European Monitoring and Evaluation Programme (EMEP); however, before the establishment of the global-Hg network by the GMOS Global Mercury Observation System (GMOS) on global scale, long-term measurements of ambient Hg concentrations and measurements of Hg wet deposition fluxes were lacking (Lindberg et al., 2007; Selin, 2009; Zhang and Wright, 2009) in several regions of the world. Although a number of monitoring stations have been in fact established to better understand the impact of Hg wet deposition on ecosystems in many countries in the Northern Hemisphere (Wängberg et al., 2007; Prestbo and Gay, 2009; Sanei et al., 2010) several regions of the world (i.e., regions which are becoming increasingly impacted by anthropogenic activities in general), and prevalently the Tropical zone and the Southern Hemisphere, were particularly lacking in wet deposition data available, in terms of concentrations and deposition Hg fluxes.

To address this concern, seasonal and annual variations of Hg wet deposition and concentration at 17 ground-based sites in the Northern and Southern Hemispheres were monitored as a part of GMOS (www.gmos.eu). Here an overview of the seasonal/annual Hg wet deposition patterns across the 17 sites, is presented, briefly examining meteorological/climatological conditions, as well as indicators of anthropogenic air mass sources and/or atmospheric chemical conditions in relation to Hg wet deposition results observed. This study is the first multi-year comparison of Hg wet deposition worldwide and provides insights into annual and seasonal variations, as well as spatial gradient in Hg deposition patterns.

dance with the measurement practice adopted in well-established regional monitoring networks and based on the most recent literature (Brown et al., 2010a, b; Steffen et al., 2012; Gay et al., 2013). For THg in precipitation an ad-hoc Standard Operating Procedure has been developed and adopted within the network, and furthermore the management of the measurement program at most of the GMOS sites consisting in analysis of all precipitation samples, cleaning procedures, distribution of the sample bottles to all sites, have been performed by three reference laboratories (IVL, Sweden; CNR-IIA, Italy, and IJS, Slovenia) whereas the precipitation samples related to some other GMOS sites, in Russia (~~Listvyanka~~)[Listvyanka (LIS)], in China (~~(Mt. Walinguan (MWA), Mt. Ailao (MAL), and Mt. Changbai)~~(MCH)], and in South Africa (~~Cape Point~~)[Cape Point (CPT)] have been analyzed by local laboratories. The analytical performance and the QA/QC of the analysis carried out by the reference laboratories as well as by the local laboratories were confirmed by the results achieved during International Inter-comparison exercises for Hg in water (i.e., Brooks Rand Instruments Inter-laboratory Comparison Study). GMOS sites predominantly collected bi-weekly samples. However, considering the spatial distribution and the diversity of meteorological parameters and conditions characterizing the monitoring sites locations, the sampling frequency was sometime different across the sites. THg concentrations in precipitation samples, refrigerated and kept in the dark before the analysis (to avoid photo-induced reduction of the Hg in the precipitation sample), were determined according to the U.S. EPA Method 1631 (version E) (1631, 2002): each sample was first oxidized by BrCl (0.5 mL/100 mL sample), followed by neutralization with hydroxylamine hydrochloride ($NH_2OH \cdot HCl$). Stannous chloride ($SnCl_2$) was then added to the sample to reduce $Hg_{(aq)}^{2+}$ to $Hg_{(g)}^0$ which was quantified by Cold Vapor Atomic Fluorescence Spectrometry (CVAFS) using a Tekran Mercury Analysis System Model 2600 (Tekran Inc. Corporation, Canada). Working Hg standards solutions were obtained from a Standard Reference Material (SRM) produced by accredited laboratory (ISO/IEC 17025). Calibration standards were analyzed in the range from 0.2 to 100 ng/L (Recovery 93-109%). The standard curve was used within the coefficient of determination (r^2) greater than 0.998 (linear). Initial (IPR) and ongoing precision and recovery (OPR) solutions (5 ppt) were analyzed prior to the analysis of samples and again after every 12 samples (Recovery 91-103%). These values were within the quality control acceptance criteria for performance in the EPA Method 1631e. The method detection limit (MDL; 40 CFR 136, Appendix B) for Hg has been determined to be 0.02 ng/L. The minimum level of quantification (ML) has been established as 0.05 ng/L for THg. The QA/QC of the analysis were obtained using replicates, method blanks, field blanks, initial/ongoing precision recovery (IPR/OPR) standards, matrix spikes and certified reference materials (CRMs) with different certified Hg concentrations. Method and field blanks were always below the respective MDL, indicating minimal contamination during sampling, transport, and treatment for this study. Additionally, the sampling train materials [i.e., fluorinated polyethylene (FLPE) bottles, cylindrical glass funnels, Teflon adapters along with the glass capillary S-shaped tubes (to prevent loss of mercury from the sample) etc.] were thoroughly acid-cleaned and rinsed with ultra-pure water in the Hg laboratory before and after sampling steps, and randomly tested for Hg concentrations; they were always below the MDL. All of these materials have been triple-bagged in zip-type plastic bags to keep them clean prior to use in the field. The results of “blanks” analysis allowed us to exclude possible contamination of all samples during different steps.

2.3 Hg wet deposition flux calculation

Considering the geographical distribution of the 17 sites located at different latitude and longitude, and therefore, under different meteorological and climatologically conditions, the precipitation was not collected over an entire year at each station due to limited amount of precipitation samples occurring during specific periods (i.e., dry seasons). Therefore, Hg flux was necessarily estimated based on the volume-weighted mean (VWM) concentration and the annual total precipitation amount collected at each site. The annual THg wet deposition flux can be approximated by the following equation:

$$F_W = C_{Hg_x} \sum_{i=1}^{i=n} P^i / 1000$$

where F_W is the annual THg wet deposition flux ($\mu g m^{-2} yr^{-1}$), and C_{Hg_x} is the volume-weighted mean (VWM) concentration of THg ($ng L^{-1}$). P^i (mm ; $1 mm = 1 L m^{-2}$) represents the precipitation amount associated to each wet deposition sample.

In addition, considering also the variability of the rainy samples frequency collected at each sites (see Tables S3 and S4) the rainfall amounts as well as THg wet deposition flux weighted data at each site have been calculated taking into account for each single sample the associated sampling time normalizing the weighted data with 15-days sampling reference as prescribed within the GMOS-Standard Operating Procedure (SOP) for determination of total Hg in precipitation adopted in the GMOS network (Munthe et al., 2011)

3 Hg wet deposition patterns and inter-annual variability

~~Annual THg wet deposition flux ($\mu g m^{-2} yr^{-1}$) during 2011–2015 at the 17 GMOS sites~~

The annual variations in THg concentration and wet deposition recorded at all 17 monitoring GMOS sites are summarized in Tables ~~?? and ??~~. Tables ?? and ?? S1 and S2. Both Tables list the monitoring sites according to their latitude and for each site, rain amounts collected, the number of the sampling days as well as the annual wet deposition flux ~~and average THg wet deposition flux calculated for each year in~~ the period 2011-2015. ~~The latter was calculated taking into account the number of sampling days at each site for each sampler~~ rainfall amounts as well as THg wet deposition flux weighted data have been normalized with 15-days sampling reference as described in the section above. Annual THg wet deposition fluxes calculated at each site according to their latitude are shown in Figure 1. The Hg deposition at each site tends to vary from year to year, but to a different degree at different locations. It is well known that the magnitude of Hg wet deposition varies geographically and seasonally due to different meteorological and climatic conditions, atmospheric chemistry, and anthropogenic influences (Vanarsdale et al., 2005; Selin and Jacob, 2008; Prestbo and Gay, 2009). Therefore, considering the ~~10-11~~ sites distributed in the Northern Hemisphere, the discussion of the results will be separately related to the seven European sites (ruling out the discussion on the data related to LIS site, due to the lower number of samples collected over the sampling period, thus not enough representative for such conclusion) and the three Chinese sites (see Tables ~~?? and ??~~ S1, S2, S3 and S4) as well as those located in the tropical area and the sites distributed in the Southern Hemisphere. Considering the THg wet deposition from 2012 to 2014 at the European sites, there appears to be a geographical trend with an increase in Hg deposition from north ~~([Arctic area, i.e., Ny Alesund, Pallas etc.])~~ (NYA Norway, Pallas (PAL), Finland etc.) to south in the Northern Hemisphere ~~([i.e., Rao (RAO), Sweden, Mace Head, Listvyanka (MHE), Ireland, LIS, Col Margherita, Longobucco (CMA),~~

[Italy, Longobucco \(LON\), Italy](#)]. At the Chinese sites as well as at lower latitude (i.e., Tropical area and Southern Hemisphere) no [north-south](#) spatial trend has been observed. However, it is important to point out that the sites in the Southern Hemisphere are limited in number compared to those in the Northern Hemisphere and the data coverage is less complete for each year considered. This makes detailed evaluation of spatial trends at the southern sites difficult. In addition, apart from [Cape Point \(CPT\)](#), no historical records of THg deposition exist for the new stations established in the GMOS project.

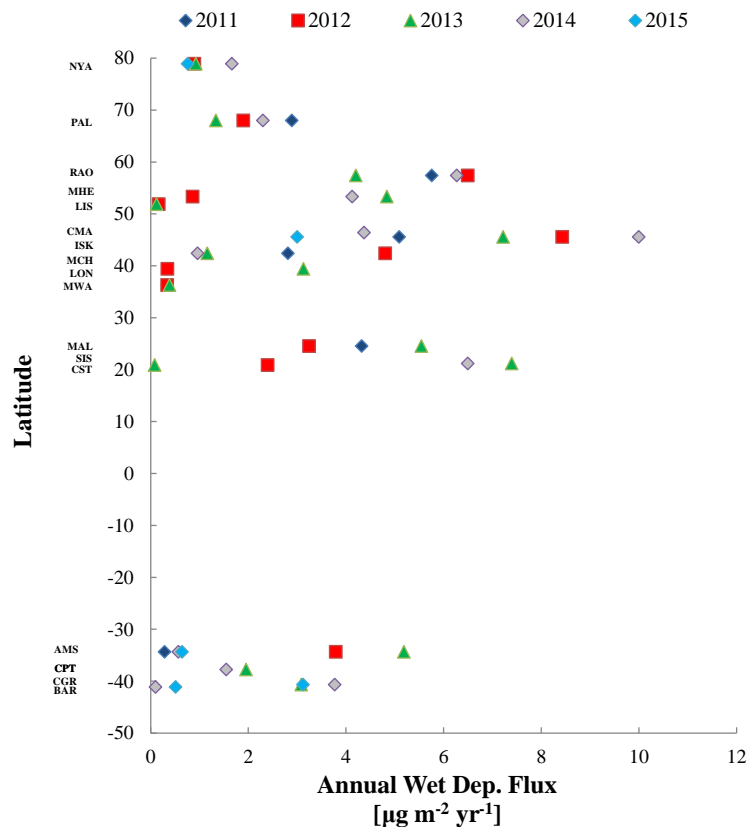


Figure 1. Average THg wet deposition Flux ($\mu\text{gm}^{-2}\text{d}^{-1}$) calculated during 2011-2015. Scatterplot reporting the Annual Wet Deposition Fluxes vs Latitude, observed at the 17 GMOS sites during 2011-2015 years.

The geographical trend observed at the European stations with higher deposition of Hg in southern sites than in the north is in line with emission patterns with the main source areas in central and eastern Europe. The present data in combination

with ground-based atmospheric Hg measurements performed within the GMOS project during 2012 - 2015 period indicate that these findings are in good agreement with the geographical distribution of atmospheric Hg with a downward gradient from the Northern to the Southern Hemisphere (Sprovieri et al., 2016). ~~Figure 1 shows~~ Figures 1 and 2 show from 2012 to 2014 (the period with more data coverage) a general increasing of THg wet deposition from ~~Ny Alesund station (Norway) to Iskrba~~ NYA station to Iskrba (ISK), Slovenia; this finding is particularly evident during the 2013 ~~for sites at lower latitudes (i.e., Mace Head, Ireland, Col Margherita and Longobucco, Italy).~~ This patterns and 2014 (Figure 2) event if the pattern is not apparent for ~~other sites such as Listvyanka (Russia) LON site in 2013 and for MHE and CMA for 2014~~ indicating the influence of other ~~emission sources parameters and/or atmospheric transport pathways.~~ In ~~order to compare~~ Figure 2 are reported the THg wet deposition ~~at all sites and look for a confirmed geographical trend in Europe, average wet deposition values were calculated ($ngm^{-2}d^{-1}$) normalizing the calculations on the effective number of sampling days. The results are shown in Figure 2. Comparing annual average wet deposition flux as is shown in Figure 1, and considering for example the 2013 period common to most of European sites, all measurements performed in the Northern Hemisphere, apart Col Margherita, where data is missing for that period, generally fits into a clear south to north decreasing trend. Deposition fluxes calculated on annual basis taking into account the annual precipitation amounts recorded at each site. It well known that wet deposition~~ of atmospheric Hg at any given location is influenced by factors such as: (a) atmospheric Hg concentration depending upon the local, regional and global sources; (b) site location in relation to the predominant wind direction in relation to the source areas; (c) precipitation amount which removes Hg from the atmosphere, ~~and~~ (d) type of precipitation (rain or snow), (e) length of precipitation events which affect Hg concentrations-

~~Precipitation amounts collected at all GMOS sites during 2011-2015~~
~~In particular,~~ (f) height and thickness of the precipitating cloud layer in the atmosphere, and the degree of convection involved, (g) and at least but not less important than the others, the oxidizing capacity of the atmosphere which can be the dominant factor particularly in remote/polar areas. Hg concentrations appear to be higher at the beginning of a precipitation event (i.e., rain or snow), and lower at the end of a precipitation event (Keeler et al., 2005; Gratz et al., 2009; Prestbo and Gay, 2009; Chen et al., 2014). This is most evident during periods of prolonged precipitation (i.e., over a period of several days). It is obvious therefore that the Hg deposition obtained at some sites, ~~is should be~~ strongly influenced by the precipitation amounts. ~~The In particular the~~ annual deposition amounts during the 2011-2015 period ~~is reported in Figure 3 which shows show~~ the influence of the precipitation amount on Hg deposition between, for example, ~~Rao (Sweden) site and Pallas (Finland) site~~ RAO site and PAL site (Figure 2). The THg wet deposition fluxes recorded during 2011, 2012, 2013, and 2014 were respectively $5.8 \mu gm^{-2}y^{-1}$, $6.5 \mu gm^{-2}y^{-1}$, $4.2 \mu gm^{-2}y^{-1}$, and $6.3 \mu gm^{-2}y^{-1}$ at ~~Rao~~ RAO site. This is more than two times higher than at Pallas during the same years ($2.9 \mu gm^{-2}y^{-1}$, $1.9 \mu gm^{-2}y^{-1}$, $1.3 \mu gm^{-2}y^{-1}$ and $2.3 \mu gm^{-2}y^{-1}$), and since the precipitation amounts are also a factor of two higher at ~~Rao~~ RAO site in comparison to ~~Pallas~~ PAL, the Hg deposition results seem to be consistent with this increase in the south compared to the northern sites. These findings also confirmed the results obtained by (Munthe et al., 2007) during an assessment on available Hg data in precipitation carried out from 1996 to 2002 at five Scandinavian EMEP monitoring stations, and among them also at ~~Rao and Pallas~~ RAO and PAL GMOS sites. (Munthe et al., 2007) highlights, in fact, that the highest annual Hg wet deposition and yearly averaged THg concentrations in precipitation

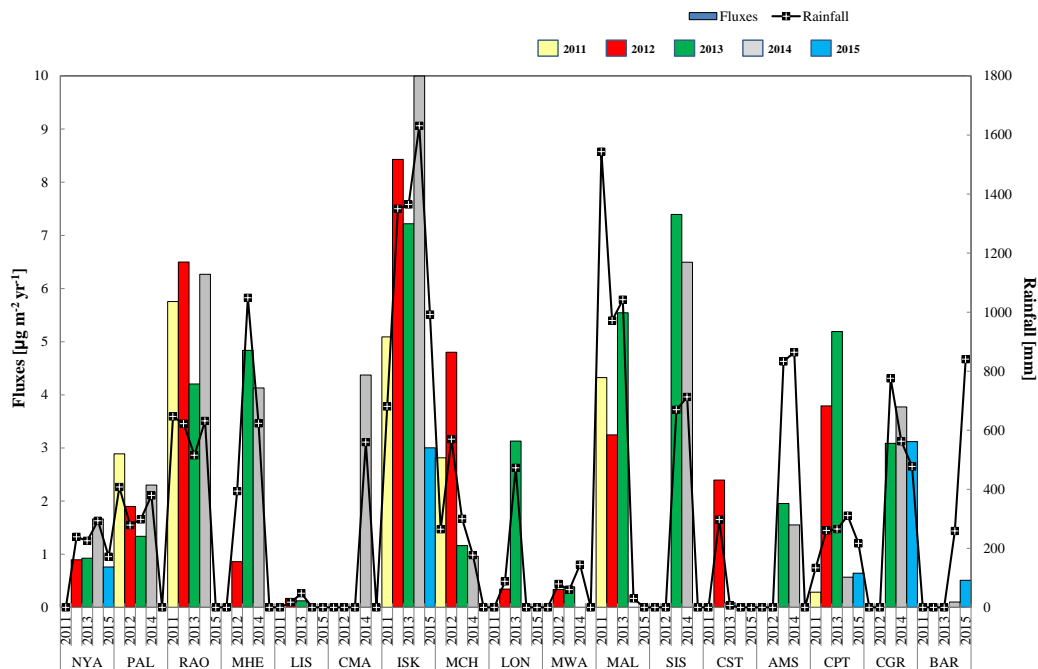


Figure 2. Annual Wet Deposition Fluxes and the corresponding cumulative precipitation amounts (Rainfall), observed at the 17 GMOS sites during 2011-2015 years.

have been recorded at the southern Scandinavian coastal sites where the highest average annual deposition amounts also occurred. The annually based THg wet deposition flux ($\mu\text{g m}^{-2}\text{y}^{-1}$) calculated, conversely, at Mt. Changbai, Mt. Walinguan and Mt. Ailao MCH, MWA and MAL show no significant geographical trend with high variability and notable differences in concentrations among the sites during the same period. These stations are all remote sites in China, and considering the 2012, 2013, and 2014 period which is the most representative in terms of number of samples recorded, it is possible to see (Figure 2) that the averaged THg wet deposition fluxes ($\text{ng m}^{-2}\text{d}^{-1}$) in remote areas of China were not significantly higher than the values observed at the rest of the GMOS sites (i.e., ISK, MHE, RAO) (Figure 2). At the sites located at lower latitude and Southern Hemisphere the relationship between precipitation amount and deposition was not as evident as in the Northern hemisphere. At the Sisal monitoring station (SIS), a coastal site of the Tropical area located on the Yucatan peninsula (Gulf of Mexico), the 2013 annual wet THg deposition flux was $67.34 \pm 7.4 \mu\text{g m}^{-2}\text{y}^{-1}$ and the average wet Hg deposition flux was $20.5 \text{ ng m}^{-2}\text{d}^{-1}$ whereas the rainfall amounts was 669.6 mm which is lower than the rainfall recorded at the remote southern sites, such as Amsterdam Island (833.6 ± 833.2 mm rainfall), southern Indian Ocean, and Cape Grim (CGR), Australia (775.6 mm rainfall) where

the annual wet Hg deposition flux recorded were considerably lower at 1.95 and 3.1 $\mu\text{gm}^{-2}\text{y}^{-1}$, respectively, and the average wet Hg deposition flux as well at 7.2 and 10.6 $\mu\text{gm}^{-2}\text{y}^{-1}$, respectively (see Tables ?? and ??S1 and S2). The 2013 and 2014 annual wet deposition flux recorded at SIS are comparable or higher than those observed at most GMOS sites in the Northern and Southern Hemisphere (Tables ?? and ??S1 and S2). Because of the Hg deposition at any given location is dependent upon both THg concentrations (which has a geographical component) in precipitation and precipitation amounts (Munthe et al., 2007), the results obtained across the sites located from the Tropical area to the Southern Hemisphere highlighted that in this case, the geographical component in terms of local meteorology and local emission sources, has had a higher influence on the THg results. During the sampling period SIS was typically influenced by air masses originated from Atlantic Ocean coming from east-south-east, but crossing the Caribbean Islands and/or Central/South America with occasional air masses coming from east-north-east mostly during the winter period crossing the south of Florida and Caribbean Archipelago prior to arrive at the monitoring site (Sena et al., 2015; Sprovieri et al., 2016). Very few Hg deposition measurements have been performed at tropical latitudes (Hansen and Gay, 2013; Shanley et al., 2008);(Shanley et al., 2015). (Shanley et al., 2015) in a study over seven years (2005-2012) on Hg wet deposition at Puerto Rico (Caribbean Archipelago, US) highlighted that despite receiving prevailing unpolluted air off the Atlantic Ocean from northeasterly trade winds, wet Hg deposition recorded at the site was about 30% higher than that observed in Florida and the Gulf Coast, which in turn, are the highest deposition areas in the U.S., and thus greater than at all other MDN sites. The wet Hg deposition map from the MDN, in fact, shows a general pattern of relatively low deposition over the western U.S. ($\sim 2 - 5\mu\text{gm}^{-2}\text{y}^{-1}$) and higher in the eastern U.S. (6-15 $\mu\text{gm}^{-2}\text{y}^{-1}$) due to increasing precipitation and location of important anthropogenic Hg sources. In addition, in the Eastern U.S. a north-south latitudinal gradient exists in wet Hg loading, with wet deposition reaching a maximum in the SE U.S. over Florida (Prestbo and Gay, 2009; Selin, 2014). Despite its unpolluted, tropical setting, Puerto Rico seems to fit as a southern extension to a latitudinal gradient of increasing Hg deposition from north to south in the eastern U.S. (Shanley et al., 2015). The high wet Hg deposition at SIS can be directly linked to the meteo-climatic conditions and pressure systems typical of the tropics. The higher THg wet deposition observed at latitudes lower than south of Florida and or Mexico, such as Puerto Rico (27.9 $\mu\text{gm}^{-2}\text{y}^{-1}$) an unpolluted tropical site crossed often by air masses detected at SIS prevalently in summer and fall and few in winter, also suggests that frequent high convective clouds in this subtropical region likely access the reservoir of oxidized Hg species in the upper free troposphere (Guentzel et al., 2001; Driscoll et al., 2013; Nair et al., 2013). (Shanley et al., 2015) found that the high Hg deposition was not correlated to GOM at ground level but to the maximum height of rain detected within clouds (obtained from the echo tops using the NOAA-NEXRAD radar station) suggesting that droplets in high convective cloud tops scavenged GOM from above the mixing layer (Shanley et al. (2015) and references therein). Numerous studies suggest in fact that the upper free troposphere holds a large pool of GOM that has been oxidized from the global Hg pool (Driscoll et al., 2013; Swartzendruber et al., 2006; Weiss-Penzias et al., 2009) and that frequent high convective clouds occurring in tropical regions, particularly closer to the Equator, scavenge GOM by precipitation being readily soluble (Lindberg et al., 2007; Selin and Jacob, 2008; Holmes et al., 2010). Closer to the equator, the Hadley cell structure indeed gives way to the Intertropical Convergence Zone (ICT), and the atmospheric circulation there may affect upper-atmosphere Hg levels. The few measurements in the Northern-Hemisphere tropics, such as SIS, generally indicate lower Hg fluxes than those measured

at lower tropical latitude probably due to fewer convective rain events with clouds that reach the upper atmosphere (Shanley et al. (2015) and references therein). The higher annual wet Hg deposition observed at SIS compared to the other GMOS sites could be also due to a contribution of air masses crossing areas with discrete anthropogenic emission sources, particularly in late spring and summer, such as the metropolitan area of San Juan and/or minor industrial plants in Fajardo and Antille Islands, and/or from air masses crossing, particularly in winter, several coal power plants and waste incinerations in the southern United States and southern Florida (Latysh and Wetherbee, 2007). In addition, also legal and/or illegal gold mining activities which are widespread (Veiga et al., 2006; Sprovieri et al., 2016) in the southern regions of the Yucatan peninsula (i.e., Nicaragua; Guatemala, etc.) could contribute to the Hg wet deposition at SIS.

The southern sites, AMS, CPT, CGR, and Bariloche (BAR), Argentina are more remote compared to SIS. AMS is a very small island located in the southern Indian Ocean where atmospheric Hg concentrations recorded during the same period were remarkably steady with annual median of $1.03 \pm 0.10 \text{ ngm}^{-3}$ and lower than those recorded at the Tropical sites (Angot et al., 2014); (Sprovieri et al., 2016) but slightly higher than annual averages and medians recorded at ~~Cape Grim~~ CGR in 2013 (Slemr et al., 2014). Both AMS and ~~Cape Grim~~ CGR for most of the time receive clean marine air masses (Slemr et al., 2014; Angot et al., 2014). Previous studies (Mason and Sheu, 2002; Sprovieri et al., 2003; Holmes et al., 2009; Sprovieri et al., 2010b, a) analyzed atmospheric observations of GOM from Mediterranean, Pacific and Atlantic cruises in terms of Hg chemistry and deposition in the marine atmosphere, and suggested that elevated levels of halogen atoms, and in particular of Br in the marine boundary layer (MBL) are an important source of GOM from oxidation of GEM, that more readily deposited throughout sea-salt aerosols followed by aerosol deposition. GEM evasion from marine waters therefore, could represent a significant source of atmospheric Hg which contributes to depositional fluxes in marine regions (Mason and Sheu, 2002), such as ~~Amsterdam Island, and Cape Grim~~ AMS, and CGR. In 2013, among the Southern sites, the highest annual ~~and average~~ THg wet deposition flux have been recorded at CPT ($5.2 \mu\text{gm}^{-2}\text{y}^{-1}$ ~~and $37.1 \text{ ngm}^{-2}\text{d}^{-1}$~~) ~~which also~~ which also showed the lowest both ~~deposition~~ precipitation amount (264.9 mm) and the number of sampling days (Tables ~~?? and ??~~ S1 and S2) compared to AMS (with annual wet deposition flux of $1.95 \mu\text{gm}^{-2}\text{y}^{-1}$ ~~and $7.2 \text{ ngm}^{-2}\text{d}^{-1}$~~ , considering a rainfall of 833.2 mm) and CGR (with wet deposition flux of $3.1 \mu\text{gm}^{-2}\text{y}^{-1}$ ~~and $10.6 \text{ ngm}^{-2}\text{d}^{-1}$~~ , considering a rainfall of 775.6 mm). These findings have not been observed at CPT in 2014 with the lowest annual wet deposition flux ($0.57 \mu\text{gm}^{-2}\text{y}^{-1}$) and comparable precipitation amounts and number of sampling days of the year before (see Tables ~~?? and ??~~).

~~Seasonal distribution of rainfall amounts, at the European GMOS sites from 2011 to 2015~~

~~Seasonal distribution of volume-weighted THg concentration in precipitation at the European GMOS sites from 2011 to 2015~~

~~Seasonal distribution of THg wet deposition flux at the European GMOS sites from 2011 to 2015~~

~~Seasonal distribution of THg wet deposition flux averaged on the number of sampling days, at the European GMOS sites from 2011 to 2015~~

S1 and S2). CPT is situated on the southern tip of South Africa (Sprovieri et al., 2016; Brunke et al., 2016), and during the wetter season (May till October) normally precipitation increased due to the passage of cold fronts moving from West to East (Brunke et al., 2016). ~~(Brunke et al., 2004)~~ In a previous study by Brunke et al. (2004) it was highlighted that CPT receives

~~clean marine air most of the time whereas~~ continental and polluted air masses ~~are observed at the site~~ more frequently during the winter period with air masses advected to the station from north to north-western (Rautenbach and Smith, 2001; Brunke et al., 2004) region where the Gauteng and Mpumalanga provinces are located. These south African areas represent the major anthropogenic Hg sources with former mine dumps from gold mining and large coal-burning power stations (Dabrowski et al., 2008). Therefore, in the first instance, the highest annual average THg wet deposition flux observed at CPT in 2013 compared to the other southern sites which received more precipitation amounts than the CPT site ~~seem to could~~ be prevalently influenced by regional/large scale emission sources during the sampling period. Measurements of atmospheric Hg deposition in ~~Bariloche (BAR), Argentina BAR~~ have been carried out for the first time from 2014 till 2015. BAR site has been established inside a well protected natural reserve in Northern Patagonia, on the shore of Gutierrez River at south-east of the Nahuel Huapi lake. GEM records at BAR station resemble background concentrations comparable to levels found in Antarctica and other remote locations of the South Hemisphere with annual mean GEM concentrations of $0.9 \pm 0.14 \text{ ngm}^{-3}$ (Diéguez et al., 2015; Sprovieri et al., 2016). The annual THg wet deposition flux calculated at BAR in 2014 was very low ($0.1 \mu\text{gm}^{-2}\text{yr}^{-1}$), however, it is necessary to point out that the number of samples carried out during the year was scarce ($n = 91$), therefore, ~~the average wet deposition flux value ($1.1 \text{ ngm}^{-2}\text{d}^{-1}$) obtained is~~ less representative than that recorded in 2015 (~~$3.0 \text{ ngm}^{-2}\text{d}^{-1}$~~) and calculated over a number of sampling days of nearly 50% of the year. The 2015 THg wet deposition flux was $0.5 \mu\text{gm}^{-2}\text{yr}^{-1}$ ~~and an average wet deposition flux of $3.0 \text{ ngm}^{-2}\text{d}^{-1}$~~ which is lower than those recorded at most of the other southern GMOS sites ~~with a comparable number of sampling days and, conversely, more close to the value observed in the Arctic, at Ny Alesund station ($4.2 \text{ ngm}^{-2}\text{d}^{-1}$).~~

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20 4 Seasonal patterns and Influence of meteorological conditions on Hg wet deposition

4.1 European Stations

In this study, seasons are delineated according to the ~~metereological~~ meteorological definition. Since THg wet deposition flux depends on the total precipitation amount and the concentration of total Hg in that precipitation, the seasonal cycles of both these parameters are shown along with the cycles of Hg wet deposition in Figures ~~??, ??, ?? and ??~~. Seasonal 3, 4 and 5. In particular, Figures 3 and 4 show that seasonal trends of THg in precipitation are clearly evident at all sites, with increased Hg concentrations and deposition observed during spring and summer months at most of them, implying a significant dependence on meteorological conditions throughout the years. The seasonal variability in Hg concentrations and Hg deposition has been reported in previous studies in North America (Hoyer et al., 1995; Landis and Keeler, 1997) and Europe (Iverfeldt, 1991; Munthe et al., 2007). The warm month maximum in seasonal THg wet deposition is predominant at most European GMOS sites (Figure 4), except at ~~Mace Head (MHE) and Longobueco (LON) MHE~~ where the maximum THg wet deposition occurs during the winter ~~and the fall seasons, respectively~~. However, the patterns of THg concentrations and precipitation amounts reveal that at most of the sites, the seasonal THg wet deposition maximum corresponds to the maximum in precipitation amounts collected, except at ~~Ny Alesund (NYA), Iskrba (ISK) and LON-NYA, ISK and LON~~ (Figure 5). Therefore, the

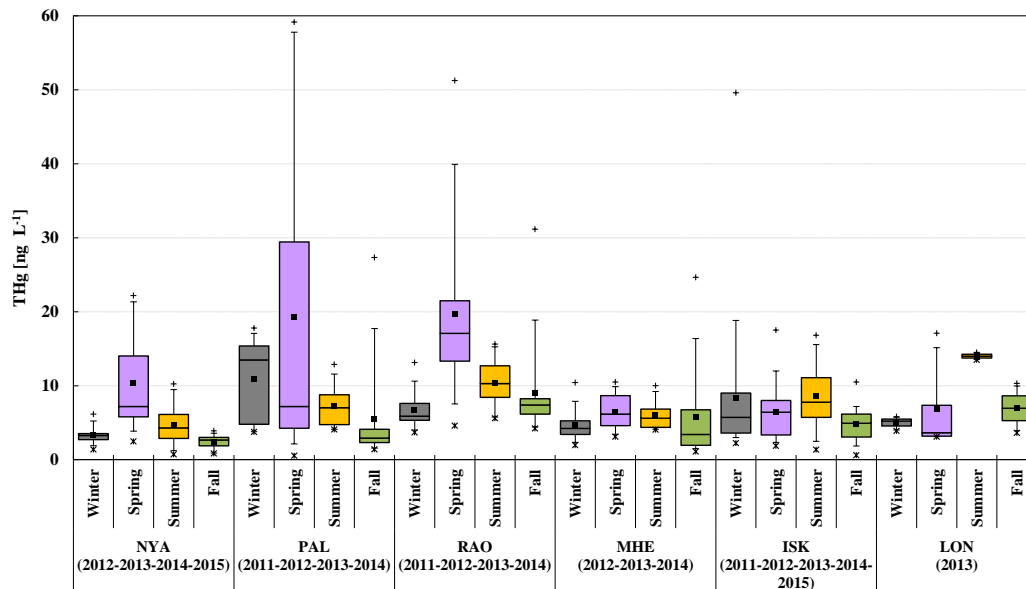


Figure 3. Seasonal distribution of volume-weighted THg concentration in precipitation at the European GMOS sites from 2011 to 2015. Each box includes the median (midline), mean (■), 25th and 75th percentiles (box edges), 5th and 95th percentiles (whiskers), minimum (*) and maximum (+).

dominant factor in determining the Hg wet deposition loading recorded at all the European sites was generally related to the amounts of precipitation collected. Hg concentrations in rainfall at NYA peaked in spring, and decreased through the summer, in fall and winter seasons (Figure ??3). Rainfall mean were fairly equally distributed in all seasons except the winter season (Figure 5). Thus, wet Hg loading was highest in spring, intermediate in winter and summer and lowest in fall (Figures ?? and ??4 and 5). High levels of soluble species could in general be due to direct enhanced atmospheric oxidation of GEM to GOM, which occurs in regions with high concentrations of oxidants such as polar regions during springtime (where AMDEs occur, such as NYA). At Pallas (PAL), PAL Hg concentrations in rainfall increased through the winter, peaking in spring, and decreased through the summer and fall (Figure 3). Rainfall was not fairly equally distributed in all seasons but lowest values were recorded during winter and spring and highest rainfall was observed in summer followed by a decreasing during the fall season (Figure 5). Thus, wet Hg loading was highest in summer, intermediate in fall, and lowest in winter and spring (Figures ?? and ??).

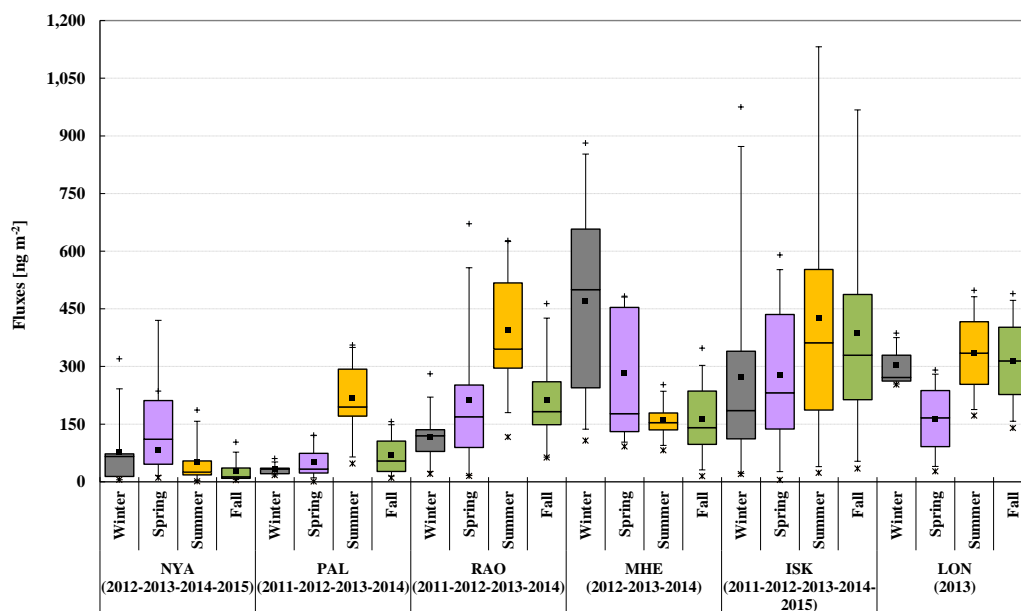


Figure 4. Seasonal distribution of time sampling flux-weighted (by 15 days reference) at the European GMOS sites from 2011 to 2015. Each box includes the median (midline), mean (■), 25th and 75th percentiles (box edges), 5th and 95th percentiles (whiskers), minimum (*) and maximum (+).

Seasonal distribution of rainfall amounts, at the three Chinese GMOS sites from 2011 to 2014

Seasonal distribution of volume-weighted THg concentration in precipitation at the three Chinese GMOS sites from 2011 to 2014

Seasonal distribution of THg wet deposition flux at the three Chinese GMOS sites from 2011 to 2014

5 Seasonal distribution of THg wet deposition flux averaged on the number of sampling days, at the three Chinese GMOS sites from 2011 to 2014

10 Similar rainfall (Figure 4). Similar behavior was observed at RAO, where Hg concentrations in rainfall peaked in spring, and decreased in fall and winter through the summer season. Therefore, wet Hg loading was highest in summer and the lowest in winter with intermediate values in spring and fall. At MHE, Hg concentrations in rainfall increased through the winter, peaked in spring, and decreased through the summer and fall seasons. Rainfall mean was fairly equally distributed in all seasons except the winter season. Thus, (Figure 4), whereas at MHE, wet Hg loading was highest in winter, intermediate in spring and

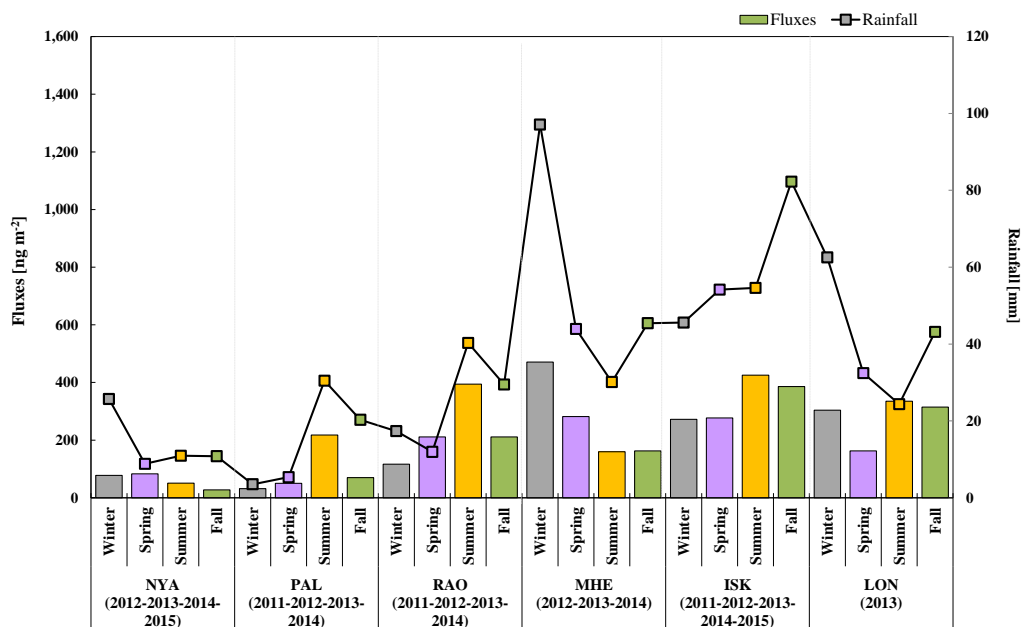


Figure 5. Seasonal mean values distribution of sampling-weighted (by 15 days reference) fluxes and rainfall, at the European GMOS sites from 2011 to 2015.

summer when also highest rainfall amounts have been recorded, and the lowest in fall (Figures ?? and ??4 and 5). At ISK, Hg concentrations in rainfall increased from the winter, and wet Hg loading peaked in summer through spring, and decreased in fall. Rainfall mean was fairly equally distributed in spring and summer seasons except the winter season which shows the lowest rainfall whereas they peaked in fall season. Thus, wet Hg loading increased from the winter, peaked in summer through spring, and decreased in fall, following the same behavior of Hg concentrations in rainfall. (Figure ?? and winter, respectively (Figures 3 and 4), whereas rainfall was highest in fall and lowest in winter (Figure 5).

LON shows highest seasonal THg wet deposition in autumn-summer and the lowest during spring. In this latter case, it is necessary to point out that these results are related to one year (2013) in contrast to the other sites in which all precipitation samples were grouped and analyzed season by season for a period of three to five years. Among the European sites the highest THg wet deposition have been recorded at the remote RAO and PAL stations during the more photochemically active summer months, whereas lower amounts were found in deposited in the colder months. In addition, rainfall amount during summer seems to be identified as the overriding factor controlling wet Hg loading at these sites. The lowest concentrations and total wet deposition were seen in winter months at most of sites. The

seasonal pattern in the atmospheric Hg, with highest precipitation concentrations and wet deposition typically seen in summer and lowest concentrations and wet deposition in winter, was believed partly to be the result of increased convection and mixing during the warmer summer months which can increase the ability of the air to transport Hg over longer distances, leading to greater precipitation amounts that remove Hg from the atmosphere. This may also indicate the role of precipitation type in the amount of Hg wet deposition, as rain may have a greater capacity to scavenge and hold different forms of Hg than snow. Higher Hg deposition, typically observed during the warmer months, was likely could be the result of a mix of meteorological, source emission, and atmospheric chemistry influences. For example, it is widely known that the concentrations of oxidants such as ozone, OH radicals, and acids that oxidize GEM to GOM are higher during warmer months and would lead to elevated concentrations of oxidized species (Schroeder and Munthe, 1998; Lin and Pehkonen, 1999). Scavenging of soluble oxidized Hg species has also been considered to be more efficient in summertime precipitation events than in winter due to differences in the cloud microphysical processing between rain and frozen precipitation (Hoyer et al., 1995).

~~Seasonal distribution of rainfall amounts, at the tropical GMOS site (Sisal, Mexico) in 2013 and 2014~~

~~Seasonal distribution of volume-weighted THg concentration in precipitation, at the tropical GMOS site (Sisal, Mexico) in 2013 and 2014~~

~~Seasonal distribution of THg wet deposition flux, at the tropical GMOS site (Sisal, Mexico) in 2013 and 2014~~

~~Seasonal distribution of THg wet deposition flux averaged on the number of sampling days, at the tropical GMOS site (Sisal, Mexico) in 2013 and 2014~~

4.2 Chinese Stations

China has been regarded as one of the largest atmospheric Hg emission sources region in the world (Streets et al., 2005; Wu et al., 2006). However, limited monitoring sites and data are available to understand Hg deposition patterns in China. Few previous measurements of THg deposition in China have been conducted in remote areas like Mt. Fanjing (Xiao et al., 1998), Mt. Leigong (Fu et al., 2010), Wujiang River basin (Guo et al., 2008), and Mt. Gongga (Fu et al., 2008, 2010) in southwestern China, as well as at ~~Mt. Changbai (MCH)~~ (Wan et al., 2009) in northeastern China. In order to evaluate the spatial and temporal distribution of THg at the three GMOS Asian stations, all measurements performed from 2011 to 2014 at ~~Mt. Changbai (MCH), Mt. Walinguan (MWA), and Mt. Ailao (MAL)~~ MCH, MWA, and MAL were grouped by season and by site (Figures ~~??, ??, ?? and ??~~ 6, 7 and 8). Seasonal variations of THg in precipitation were observed at the three Chinese sites (Figure ~~??~~ 6). The results obtained during the sampling period were similar to the seasonal variations of THg in precipitation in other Chinese regions, such as Wujiang River Basin, Guizhou, China, but in contrast to the observations in North America (Landis et al., 2002), Adirondacks (Choi et al., 2008) and Great Lakes region (Hall et al., 2005), which found increased THg concentration during summer months (Prestbo and Gay, 2009). Geographic differences in Hg wet deposition worldwide may be explained in part by the proximity to atmospheric sources and regional difference in anthropogenic emission sources. Atmospheric Hg species, in particular, GEM and PBM have been found to be substantially increased over recent years in both remote and urban areas of China, especially in central and eastern China, compared to those observed in North America and Europe which reported opposite long-term trends (Fu et al., 2015). The increasing trend in China is possibly caused by the

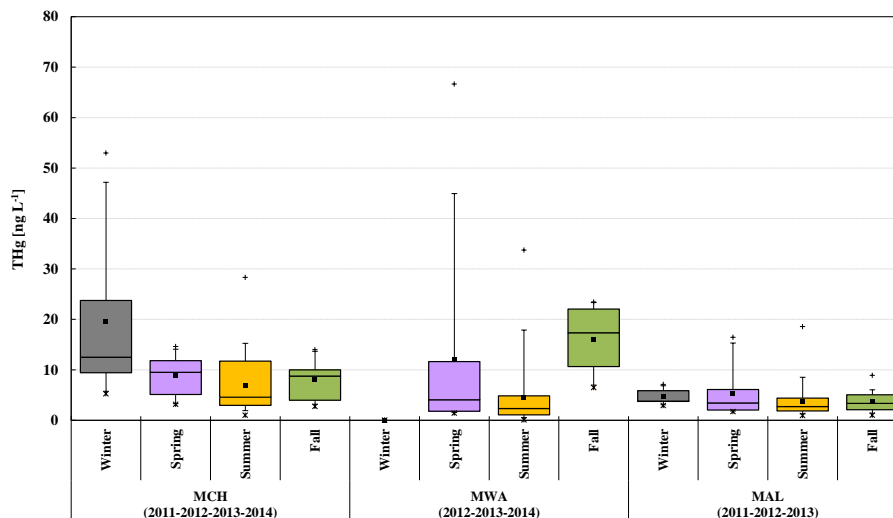


Figure 6. Seasonal distribution of volume-weighted THg concentration in precipitation at the three Chinese GMOS sites from 2011 to 2014. Each box includes the median (midline), mean (■), 25th and 75th percentiles (box edges), 5th and 95th percentiles (whiskers), minimum (*) and maximum (+).

increase in anthropogenic Hg emissions in the past decade, and indicates that the influence of regional emissions on Hg levels in China exceed global emission influence ((Lindberg et al., 2007) and references therein). The seasonal variation of weighted THg concentration observed in precipitation with highest value in winter and lowest in summer (see Figures ??, ??, ?? and ??6), could be attributed in a first instance, to lower rainy-rain amounts collected in winter (Figure 8). The results obtained at the three Chinese sites show in fact that the THg concentrations varied with rain amount. In particular, at MCH, THg concentrations slightly increased in autumn, peaked during the winter season, and decreased during spring and summer when the lowest values were recorded. The reverse trend has been observed in precipitation amount ~~through the seasons. Average with the highest value observed in summer and the lowest in winter~~ (Figure 8). THg wet deposition trend ($ngm^{-2}d^{-1}$) is comparable with that of the precipitation amount, with values of THg flux increased from winter, ~~through spring,~~ and peaked in summer (Figure 8). Ruling out the winter season at MWA during which very few rainy samples have been collected, thus not representative for the present discussion, weighted THg concentrations peaked in fall ~~and decreased during spring~~ with lowest values in ~~summer period~~ spring. Therefore, on average wet Hg loading was highest in spring, ~~intermediate in fall~~ and lowest in summer. ~~The positive or negative correlation between THg concentrations and the precipitation amount has not~~

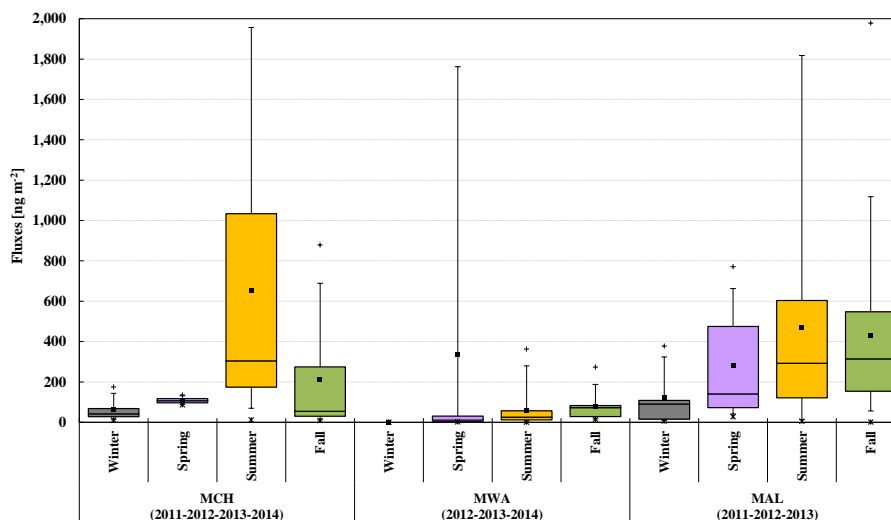


Figure 7. Seasonal distribution of sampling-weighted (by 15 days reference) fluxes, at the three Chinese GMOS sites from 2011 to 2014. Each box includes the median (midline), mean (■), 25th and 75th percentiles (box edges), 5th and 95th percentiles (whiskers), minimum (*) and maximum (+).

been obviously observed at MAL where At MAL the rainy samples show a fairly seasonal variability during all seasons with lowest average rainfall in winter and the highest in fall, whereas summer (Figure 8), while THg concentrations showed high values in winter and lowest in fall, and wet Hg loading was highest in summer, intermediate in fall and spring and the and lowest values were recorded in winter. (Fu et al., 2015) Fu et al. (2015) highlight significant positive correlations between rainwater THg concentrations and PBM and GOM concentrations, resulting in positive correlations between wet deposition fluxes and PBM and GOM concentrations. This has been explained by the authors with the washout process of PBM and GOM during rain events which could contribute to enhance Hg wet deposition in China, particularly in urban areas where PBM and GOM concentrations are much higher. Wet deposition is in fact, commonly distinguished in terms of in-cloud and below-cloud washout and involves oxidized mercury forms (GOM, PBM). Gaseous Hg^0 does not undergo direct scavenging by precipitation because of its low solubility, but it can be washed out indirectly through dissolution and oxidation in cloud water. In remote areas of China, however, washout of elevated atmospheric PBM does not seem to drive a notable increase in Hg wet deposition flux, probably due to the low washout rate of PBM during rain events at high altitude monitoring sites, such as MAL and MWA where low-level clouds reduced the contribution of Hg washout (Lee et al., 2001; Seigneur et al., 2004).

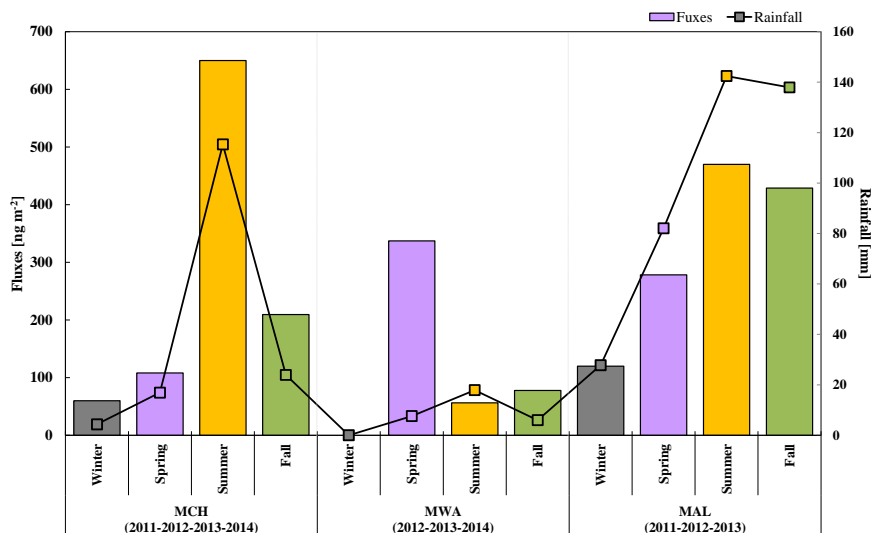


Figure 8. Seasonal mean values distribution of sampling-weighted (by 15 days reference) fluxes and rainfall, at the three Chinese GMOS sites from 2011 to 2014.

(Guo et al., 2008) in a previous study in Guizhou on Hg in precipitation also pointed out that maximum THg concentrations in rainy samples during winter-cold seasons may be related to coal burning in domestic activities. Similar conclusions have also been reported in a study performed by Wang et al. (2012) (Wang et al., 2012) at three Chinese sites (urban, residential and near-remote sites) in Chongqing province from 2010 to 2011, where they also found a high correlation between THg and particulate Hg (PBM) concentrations, suggesting that THg concentration in precipitation may be influenced by the PBM concentration. Additionally, comparable seasonal behavior of Hg concentrations in precipitation with our results have been also observed, but with annual mean THg concentrations (ngL⁻¹) significantly higher than those observed at MCH, MWA, and MAL sites which are located in remote Chinese areas. The seasonal pattern in deposition flux observed at the remote MCH, MAL, and MWA are comparable with those observed at remote sites of Europe and North America (Choi et al., 2008; Mason et al., 2000; Keeler et al., 2005; Sanei et al., 2010; Lombard et al., 2011), with maximum values during warmer months (Figures ??, ??, ?? and ??). It was suggested by (Keeler et al., 2005) and (Mason et al., 2000) that this annual maximum was mainly due to more effective scavenging by rain in summer than by snow in the cold season (Sorensen et al., 1994; Mason et al., 2000; Keeler et al., 2005; Selin and Jacob, 2008). Mercury Hg is not incorporated into cold cloud precipitation as efficiently as in warm cloud precipitation (Landis et al., 2002). Other explanations for this observation have been addressed by the authors

including a greater availability of soluble Hg due to convective transport in summer events (Guentzel et al., 2001; Keeler et al., 2005), and a summer increase in Hg-containing soil derived particles in the atmosphere (Sorensen et al., 1994).

4.2.1 Tropical Station: SISAL, Mexico

~~Seasonal distribution of rainfall amounts, at the four GMOS sites in the Southern Hemisphere from 2012 to 2015~~

5 ~~Seasonal distribution of volume-weighted THg concentration in precipitation, at the four GMOS sites in the Southern Hemisphere from 2012 to 2015~~

~~Seasonal distribution of THg wet deposition flux, at the four GMOS sites in the Southern Hemisphere from 2012 to 2015~~

~~Seasonal distribution of THg wet deposition flux averaged on the number of sampling days, at the four GMOS sites in the Southern Hemisphere from 2012 to 2015~~

10 Hg deposition measurements are rare in tropical latitudes, with very few scientific publications in the past decade (Shanley et al. (2015) and references therein). The tropics are a particularly important region regarding global atmospheric chemistry. Due to intense ultraviolet radiation and high water vapor concentrations, high OH concentrations oxidize inorganic and organic gases, and induce an efficient removal from the atmosphere of the oxidized products (Shanley et al. (2015) and references therein). Strong convective events in the tropical regions leads to huge volumes of air being drawn out of the sub-cloud
15 layer with the resultant chemical composition of the precipitation coming from the capture of gases and small particles by the liquid phases of cloud and rain. Hg deposition measurements started in Mexico at Celestùn station (CST) in 2012 (see Table 1), but after a short time period of sampling, the monitoring station changed the location with SIS, therefore, we refer the discussion to the SIS data related to both 2013 and 2014 years during which sufficient precipitation samples have been recorded ([Figures 9, 10 and 11](#)). Despite receiving unpolluted air off the Atlantic Ocean from northeasterly and southeasterly
20 trade winds, during most of the years (Sena et al., 2015), the site recorded higher wet Hg deposition fluxes during summer and fall compared to those observed during the other seasons ([Figure 10](#)). The SIS high Hg deposition rates, comparable to other sites in the Northern Hemisphere, such as the Chinese sites (i.e., [MWA/MCH](#)) or European sites (i.e., ISK) that sometimes are also impacted by anthropogenic emissions, are driven in part by high rainfall events more intense during summer and fall, and less during winter and spring period. The high wet Hg deposition flux at this site suggests that other tropical areas
25 may be hotspots for Hg deposition as well. A number of studies have suggested that this could be due to higher precipitation and the scavenging ratios from the global pool in the sub-tropical free troposphere where high concentrations of oxidized Hg species exist (Guentzel et al., 2001; Seigneur et al., 2004; Selin and Jacob, 2008). These findings were also highlighted in previous studies in south of Florida and the Gulf of Mexico coastal areas confirming that local and regional Hg emissions play only a minor role on wet Hg deposition (Guentzel et al., 2001; Sillman et al., 2013) suggesting that the primary source of
30 scavenged oxidized Hg could be the global pool. Weather patterns in SIS exhibit a seasonality [in](#) annual rainfall, with highest rainfall from June/July through October/November. Summer tropical waves and systems characterized by deep convection and low pressure produced greater rainfall. During summer and fall, the site indeed receives rainfall from deep convection associated with tropical waves embedded in the prevailing easterly airflow. THg concentrations were higher in low volume samples. With larger storms Hg concentrations were diluted, this means that rainout of Hg was maximum (the decreasing of

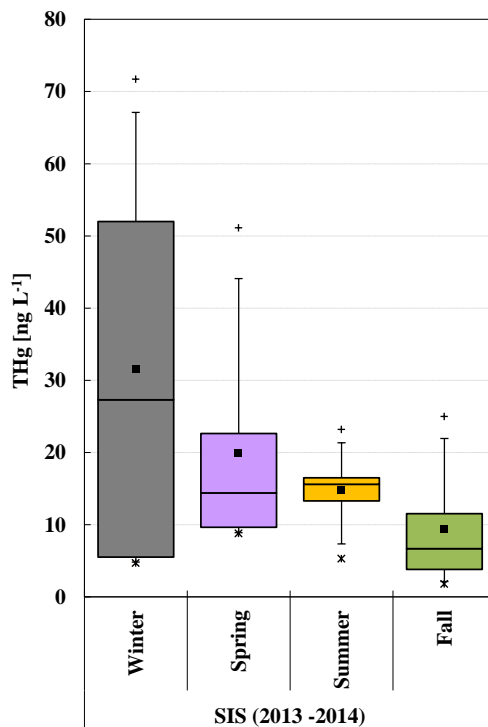


Figure 9. Seasonal distribution of volume-weighted THg concentration in precipitation at the tropical GMOS site (Sisal, Mexico) in 2013 and 2014. Each box includes the median (midline), mean (■), 25th and 75th percentiles (box edges), 5th and 95th percentiles (whiskers), minimum (*) and maximum (+).

Hg concentrations with the increasing of the rainfall depth). Weighted THg concentrations in rainfall (ngL^{-1}) increased from the fall, peaked in winter, and decreased through the spring and summer. On average terms (Figure 9), THg in wet deposition was highest in summer, intermediate in fall, and lowest in spring and winter (Figures ??, ??, ?? and ?? Figure 10). The higher summer Hg deposition flux is not driven by higher Hg concentrations in rainfall since the highest Hg concentrations in rain samples occurred in winter (Figures ??, ??, ?? and ??). Different mechanisms leading to enhanced Hg concentrations in rain during the winter including greater anthropogenic emissions are probably associated with higher use of fossil flues-fuels in

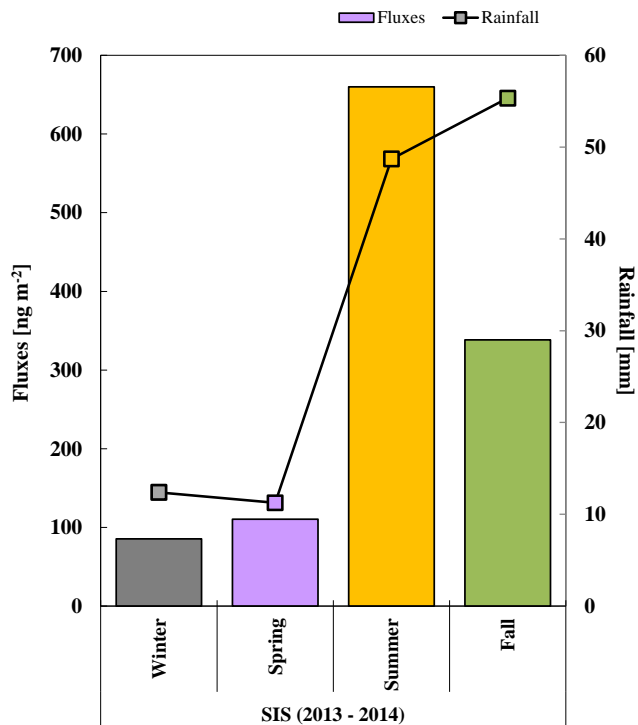


Figure 10. [Seasonal distribution of sampling-weighted \(by 15 days reference\) fluxes, at the tropical GMOS site \(Sisal, Mexico\) in 2013 and 2014. Each box includes the median \(midline\), mean \(■\), 25th and 75th percentiles \(box edges\), 5th and 95th percentiles \(whiskers\), minimum \(*\) and maximum \(+\).](#)

power plants during the cold season. As reported in Section 3 relating to the annual wet deposition patterns, the THg wet deposition observed at SIS could also be influenced by air masses crossing particularly in winter the southern ~~Unite~~ [United States](#) and southern Florida where several coal power plants and waste ~~incinerations~~ [incinerators](#) (Latysh and Wetherbee, 2007) are located. The high wet deposition of Hg during the rainy seasons (May/June to October/November), in contrast, could be due to more efficient scavenging processes of reactive gaseous mercury from the free troposphere by tall convective thunderstorms, and the concentration of GOM by the sea breeze effect, where the diurnal alternation of onshore and offshore winds can lead to

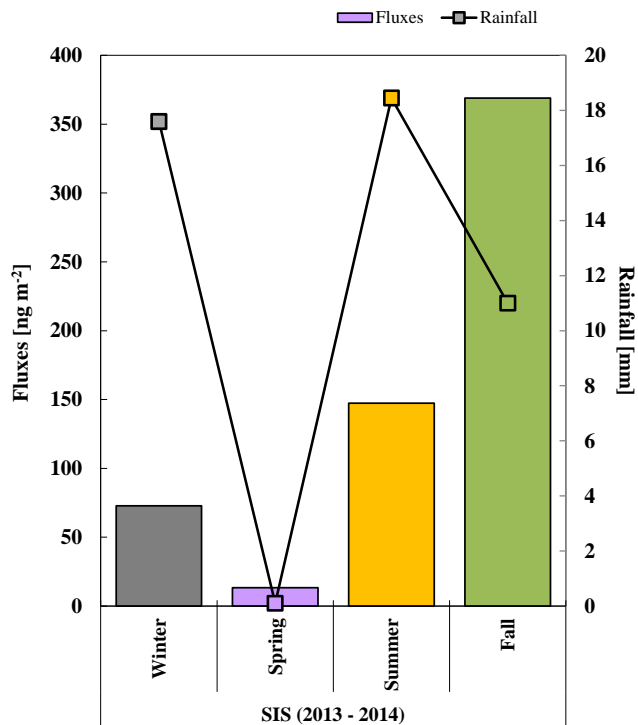


Figure 11. [Seasonal mean values distribution of sampling-weighted \(by 15 days reference\) fluxes and rainfall, at the tropical GMOS site \(Sisal, Mexico\) in 2013 and 2014.](#)

a buildup of pollutants in the air mass. Greater information on Hg deposition and cycling is needed in tropical regions, where populations are more likely to be exposed to Hg through fish consumption and artisanal gold mining activity.

4.2.2 Southern Hemisphere Stations

In remote areas far from any local sources, atmospheric deposition has been recognized as the main source of Hg to the ocean
 5 (Lindberg et al., 2007; Pirrone et al., 2008). Hg can then be reemitted back to the atmosphere via gas exchange, and modeling studies suggest that reemission from oceans is a major contributor to atmospheric concentrations of GEM, particularly in

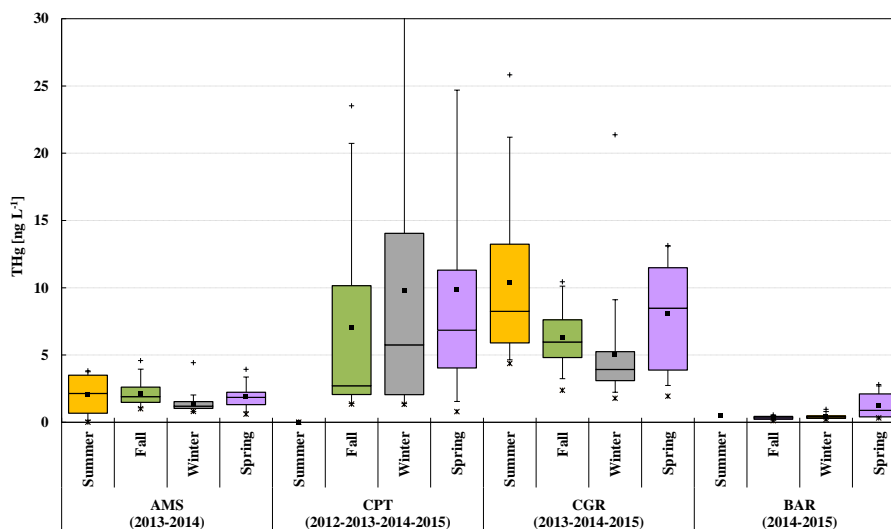


Figure 12. Seasonal distribution of volume-weighted THg concentration in precipitation, at the four GMOS sites in the Southern Hemisphere from 2012 to 2015. Each box includes the median (midline), mean (■), 25th and 75th percentiles (box edges), 5th and 95th percentiles (whiskers), minimum (*) and maximum (+).

the Southern Hemisphere where oceans were shown to contribute more than half of the surface atmospheric concentration ((Strode et al., 2007) and references therein). In the Southern Hemisphere we considered the four monitoring sites, Amsterdam Island (AMS), southern Indian Ocean AMS, CPT, South Africa, Cape Grim (CGR), Australia, and Bariloche (BAR), Argentina CGR, and BAR, which recorded a representative number of samples over the 2012-2015 period. Figures ??, ??, ?? and ??

5 12 and 13 show the box plots related to rainfall, THg concentrations in precipitation as well as wet deposition flux of Hg recorded whereas Figure 14 shows the mean values of rainfall amounts with the corresponding mean values of Hg fluxes at the four southern sites. An NSA-171 (Eigenbrodt) collector was set up at AMS at the beginning of the 2013. The GMOS site experiences a mild oceanic climate with monthly median air temperature ranged from 11 °C in austral winter to 17 °C in austral summer and frequent presence of clouds (Sciare et al., 2009). In 2013 and 2014 AMS displays the highest precipitation

10 amounts-a season variation of the precipitation amounts with the highest values collected during the warmer seasons (spring and summer)(Fig. ??, ??). Also winter season (Figure 14). On the contrary, the THg wet deposition flux patterns follow the same trend observed for the rainfall highlighting that the main factor driving the flux seems to be the amount of rain collected (Fig. ??). The THg fluxes pattern seems to be in agreement with the results of atmospheric Hg speciation measurements

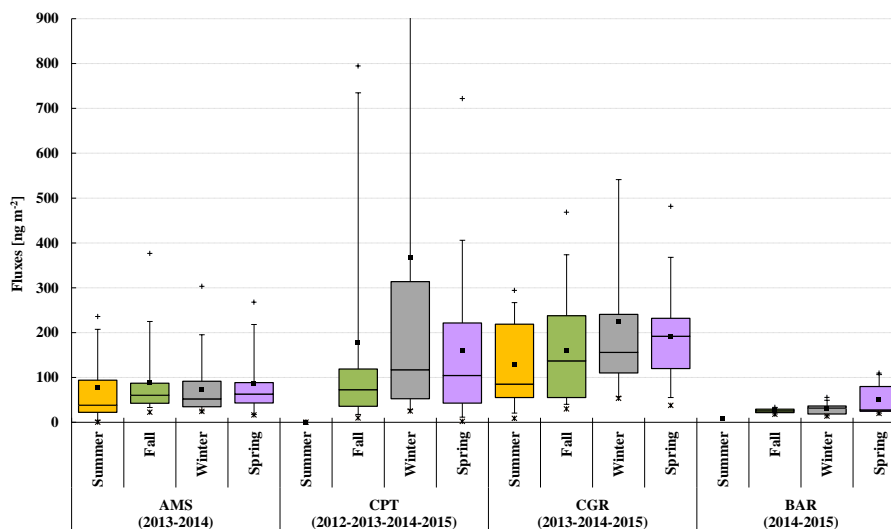


Figure 13. Seasonal distribution of sampling-weighted (by 15 days reference) fluxes, at the four GMOS sites in the Southern Hemisphere from 2012 to 2015. Each box includes the median (midline), mean (■), 25th and 75th percentiles (box edges), 5th and 95th percentiles (whiskers), minimum (*) and maximum (+).

carried out during the same period at AMS, and in particular with the GOM seasonal pattern observed since January 2012 by (Angot et al., 2014) that highlighted a higher frequency of GOM events between December and March (summer). However, additional and integrated measurements in ambient air and rainwater samples to improve our understanding of deposition processes and oxidation mechanisms should be addressed. The didn't show a similar variation throughout the seasons as well as the THg concentrations in precipitation samples (Figures 12 and 13). At CPT the variation of Hg concentrations in precipitation and Hg wet deposition fluxes driven by the precipitation amounts collected at AMS occurred also at CPT where, where apart the dry summer season, Hg concentrations concentration in precipitation, Hg wet deposition fluxes as well as the precipitation amounts, followed the same trend during the rainy season (May till October), with a maximum in wintertime for all the parameters recorded. CPT experiences a Mediterranean-type climate that is characterized by rather dry summers comprising moderate temperatures. The austral autumn to spring season normally experience increased precipitation due to the passage of cold fronts moving from West to East, therefore, CPT generally receives clean marine air from the Atlantic Ocean whereas continental and polluted air masses are observed at the site more frequently, mainly during the winter period (Brunke et al., 2004, 2016), due to the prevailing air masses from the north to northwestern sector (Rautenbach and Smith, 2001; Brunke

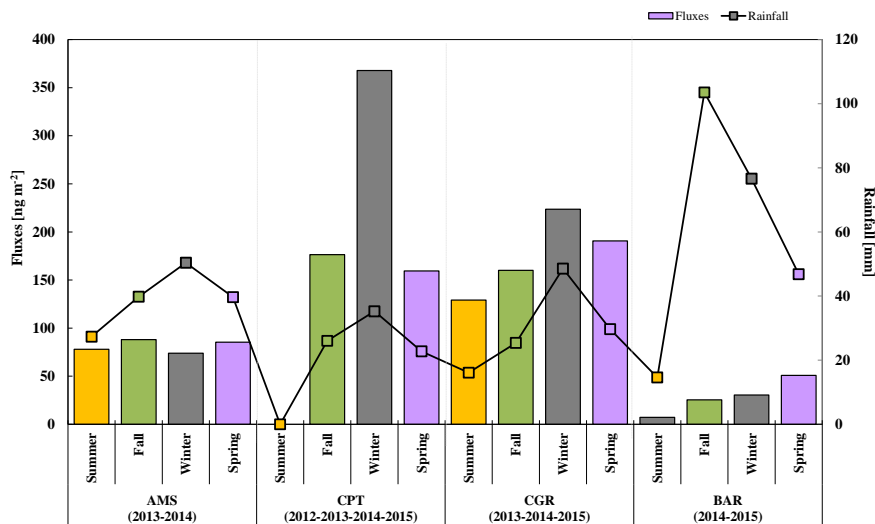


Figure 14. [Seasonal mean values distribution of sampling-weighted \(by 15 days reference\) fluxes and rainfall, at the four GMOS sites in the Southern Hemisphere from 2012 to 2015.](#)

et al., 2004). The highest THg concentrations and wet deposition fluxes recorded during the winter season could be due also to the contribution of polluted air masses crossing Cape Town metropolitan area before arriving at the stations. However, [in a previous a more recent study on GEM concentrations and THg in precipitation carried out by Brunke et al. \(2016\) over a period of seven years \(2007-2013\) by \(Brunke et al., 2016\)](#) highlighted that GEM, THg, CO and ²²²Rn levels within the urban-marine events observed at CPT did not substantially differ from those seen in the marine rain episodes, concluding that no significant local anthropogenic influences were detected on THg concentrations. Conversely, a significant positive correlation was found CPT between GEM and THg concentrations, and with the Southern Oscillation Index (SOI), suggesting that both GEM and THg concentrations are primarily influenced by large scale meteorology which in turn controls Hg emission sources in terms, for example, of enhanced sea surface temperature that could increase large scale droughts leading to a raised biomass burning (Brunke et al., 2016).

Measurements of atmospheric Hg deposition in Australia have never been reported before [\(?\) \(Jardine and Bunn, 2010\)](#). From 2013 till 2015, at [Cape Grim GAW Station \(CGR\) CGR GAW Station](#), located on the north-western coast of Tasmania, Australia, highest value in rainfall have been observed during winter an lowest in summer, whereas Hg concentrations peaked in summer and dropped to lowest values in winter (see [Fig. ??, ??, ?? and ??](#)). [The trend of Hg wet-deposition fluxes conversely seems](#)

to be driven by the precipitation amounts even if a small seasonal variability of Hg loading was displayed. ~~Figure 12 and 14~~). Indeed, an increase in precipitation volume results in ~~an increase of the Hg deposition flux. This is accompanied by~~ a decrease in Hg concentrations in rain, probably due to the dilution of the washout loading (Prestbo and Gay, 2009). This means that any ~~changes change~~ in meteorological conditions, especially precipitation, complicate the interpretation of GMOS observations at different latitude and might mask any trends due to ~~change changes~~ in Hg emissions. The trend of Hg wet deposition fluxes shows a seasonal variability with highest values in spring and lowest in cold seasons. At BAR the highest precipitation amounts in 2014 and 2015 were collected during the fall and winter seasons and decreased in spring when the highest THg concentrations occurred (see ~~Fig. ??, ??, ?? and ??~~ Figures 12 and 14). Therefore, the seasonal THg wet deposition peaked in spring and decreased during the cold seasons (Figure 13). It is necessary to point out, however, that in ~~both 2014 at BAR no and~~ 2015 at BAR very few samples have been recorded in fall and summer ~~as well as in 2015, during the same seasons the number of sampling days was very low particularly in summer~~ (Tables S1, S2, S3 and S4). This means that further measurements and studies are needed to draw any conclusion and improve our understanding of deposition processes and oxidation mechanisms in this region. There are very few previous observations of Hg wet deposition in the Southern Hemisphere, and this makes difficult any comparison of data recorded during GMOS. The results observed at the four southern GMOS sites highlighted that the magnitude of wet deposition is affected by two main factors: amount of precipitation and the THg concentration in precipitation influenced by soluble Hg species (oxidized Hg) in the atmosphere. High levels of soluble species could in general be due to direct anthropogenic emissions of Hg oxidized species or by enhanced atmospheric oxidation of GEM to GOM, which occurs in regions with high concentrations of oxidants such as southern locations (where more solar radiation occurs) or polar regions during springtime (where AMDEs occur).

20 5 Conclusions

Mercury deposition measurements are critical for constructing an accurate global Hg budget and to model the benefits or consequences of changes in Hg emissions, for example, as proscribed by the Minamata Convention. ~~Early models of The scarce availability of long term wet Hg deposition had few measurements data for calibration or validation, and tended to overestimate of models could give uncertainties in modeling applications to assess~~ the influence of local emission sources. A synthesis of ~~all~~ available Hg measurements in precipitation from ~~GMOS network selected GMOS ground-based sites~~ is presented, including trends and seasonal cycles. ~~These Wet deposition samples were collected for approximately five years, from 2011 to 2015, at 17 selected GMOS monitoring sites located in the Northern and Southern Hemispheres, as well as in the Tropical area. In the Northern Hemisphere and specifically at the European stations a geographical trend with an increase in THg wet deposition from north to south have been observed. These findings are in good agreement with the geographical distribution of atmospheric Hg data obtained during the same period within the GMOS network with a downward gradient from the Northern to the Southern Hemisphere. At the other GMOS monitoring sites in the Northern Hemisphere (i.e., Chinese sites) as well as at lower latitude (i.e., Tropical area and Southern Hemisphere) no north-south spatial trend has conversely been observed. Annual and seasonal patterns in Hg wet deposition are clearly evident at all GMOS sites, implying a significant~~

dependence on meteorological conditions throughout the years. Most of ground-based sites report, in particular, Hg deposition strongly influenced by the precipitation amounts. In the Northern Hemisphere, inter-annual differences in THg wet deposition are mostly linked with precipitation volume, with the greatest deposition flux occurring in the wettest years, whereas at the sites located at lower latitude and Southern Hemisphere the relationship between precipitation amount and deposition was not as evident as in the North. It is however to point out the need to expand the global network particularly in the Tropics and Southern Hemisphere regions in order to provide more information throughout long-term monitoring activities. As a start point of a global network, these results provide a set of data for modeling applications to fully understand THg wet deposition patterns as well as the transformation and deposition mechanisms of atmospheric Hg. With broad geographic coverage including mostly background and remote sites ~~with few~~ as well as local or regional sources, GMOS's observation network gives important insights to modeling applications to evaluate future Hg trends and its fate and transport on global scale. The results on THg wet deposition carried out in this study open the way for new avenues in future modeling studies as well as highlight the need of additional and integrated measurements in ambient air and rainwater samples to improve our understanding of deposition processes and oxidation mechanisms. These new observations in fact, give scientists and modelers some insight into baseline concentrations of THg concentrations in precipitation and depositional fluxes especially in the tropical area, and in the Southern Hemisphere where wet deposition as well as atmospheric Hg species were not investigated before. Greater information on Hg deposition and cycling is obviously needed in these regions. Moving forward, in addition to continued monitoring GMOS sites, integration with other ground-based monitoring sites at strategic locations along with integrations with atmospheric Hg species and other key oxidants, identification of the compounds making up GOM and PBM_{2.5} continue to be needed. Knowledge of these exact chemical species would also lead to improved understanding of the chemistry and wet and dry deposition processes of oxidized Hg ~~specie~~ species in different air masses. These and other uncertainties are the subject of ongoing research. The magnitude of Hg dry deposition is to date uncertain, especially dry deposition of GEM, and few measurements are available to constrain model estimates. Further measurements of dry deposition, especially in locations where wet deposition measurements are available, would dramatically improve scientific understanding of the Hg cycle. Wet deposition measurements worldwide would assist modelers in constraining the atmospheric Hg budget on global scale, as would additional direct measurements of dry deposition across the GMOS network.

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Interactive comment on “Five-year records of Total Mercury Deposition flux at GMOS sites in the Northern and Southern Hemispheres”

by Francesca Sprovieri et al.

First of all, we thank the three reviewers for their effort and useful suggestions reported for the manuscript on mercury wet deposition flux performed at the GMOS sites distributed worldwide. We completed the revision of the manuscript according to comments provided by reviewers taking into account the important input and corrections they highlighted.

We appreciate very much their valuable comments for improving the readability and interpretation of the manuscript. We think that after this review, our manuscript has been now improved. Below we report point by point our detailed responses to the comments for each Reviewer. Thank you very much once more.

Anonymous Referee #1

1. Title: Commonly, “total deposition flux” means sum of wet and dry deposition. There is no discussion about Hg dry deposition in the paper. So, probably, “Five-year records of wet mercury deposition flux....” would be more proper title for the paper.

Reply: We thank the reviewer and the other two reviewers (that highlighted the same comment) for pointing out an inaccurate title of the manuscript. We agree with you, therefore, following your input we revised the title according to. Please, see the Title of the revised version of the manuscript.

2. Page 2, line 19. “.....in both depositional flux and concentration with the highest values...”
Page 2, line 25. “ ...coastal sites receiving higher Hg concentrations and depositional Fluxes....” Concentrations in air or in precipitation are mentions here? Please, specify to avoid misleading.

Reply: We thank this reviewer for pointing out an inaccurate statement in the original text. We have made thorough revision in the revised version of the paper to eliminate and/or correct such an inaccurate statement. Please, see page 2, lines 20-23 and 27-31 in the revised version of the manuscript. Thank you.

3. Page 2, lines 27-29. “.....gaseous evasion of Hg from marine waters is a significant global source of atmospheric Hg and may also contribute to elevated depositional fluxes in coastal regions.....” Wet deposition is mostly comprised precipitation removal of highly soluble oxidized Hg. Significant Hg evasion from the ocean, which is in poorly soluble elemental form, does not necessarily mean elevated deposition. Oxidation of GEM to GOM is essential.

Reply: Yes, that’s right. We agree with you in respect to the oxidation processes of GEM to GOM which are essential within the wet deposition mechanisms to remove the highly soluble form of Hg from the atmosphere. Therefore, in order to make the sentence more clear and accurate, we revised the sentence. Please, see the revised version of the manuscript at page 2, lines 31-33. Thank you.

4. Page 2, line 34. “.....the EMEP program the GMOS”...These acronyms which appear for the first time require explanation and references.

Reply: Done, Thank you. We explained these acronyms in the text, and reported the references. Please, see now at page 3, lines 5 and 6.

5. Page 2, line 33 – Page 3, line 1. “Long-term Hg wet deposition measurements exist at many locations within the United States as part of the MDN or in Europe as part of the EMEP program; however, before the establishment of the global Hg network by the GMOS, long-term of ambient Hg concentrations and measurements of Hg wet deposition fluxes were lacking” The second part of the sentence contradicts the first part.

Reply: Yes, Thank you for your comment. We revised these sentences according to in order to make clear what they mean. Please, see the revised text at page 3, lines 3-13.

6. Page 6, Section 2.3. What were the criteria of data coverage for calculation of annual and seasonal mean values? As it follows from Tables 2-3 and Figs. 1-3 the annual data are not available for all the stations for all years.

Reply: We appreciate very much the comment of the reviewer regarding the criteria of data coverage for calculation of annual and seasonal mean values. Regarding the data, unfortunately, we did not gain a full coverage as well as samples collected with an homogenous time frequency. For that concerning data coverage, the GMOS wet deposition samples have been carried out with irregularity due to technical troubles with stuff and some problems in situ. Otherwise, since many GMOS stations provided wet deposition data recorded for the first time at their locations, we believe that, even if with a partial coverage, they could be helpful in making possible a global picture of the issue under study. To overcome the irregularity in time-sampling frequency we have done a normalization of both rainfall amounts and Hg fluxes in respect to the ideal time-sampling period that is equal to 15 days, as previously established in our GMOS Standard Operating Procedures. Concerning the Tables 2 and 3 and the Figures 1-3, they have been replaced and re-organized in order to accommodate all comments and suggestions made on data coverage criteria and related calculation as well as on precision of measurements data etc., made also by the other two reviewers. Please, see Tables 2 and 3 that are replaced with Table S1 and Table S2 reported within the supplementary material added to the revised version of the manuscript. The supplementary material added also includes additional two new tables (Table S3 and Table S4) reporting useful information about measurements and data calculation. Figure 1 has been replaced with a scatter plot as suggested by the reviewer n.3 and Figure 2 and Figure 3 have been merged in a “new” Figure 2 in order to make clear the relationship of wet deposition fluxes and the rainfall amounts at each site for each year, and we have also switched x/y axes according to the reviewer n.3 and enlarged the bars to make easier reading the Figure itself. Please, see the revised version of the manuscript along with the supplementary material included. Thank you.

7. Figs. 1-3. The bars at the diagrams are very thin that makes difficult reading the figures.

Reply: Regarding Figure 1-3, according to your comment, and in order to accommodate also the comment of the third referee (see the comment of Mark Cohen Number 9 on Figure 1 within the “Specific Comments” section, and several his suggestion reported earlier related to the need to

combine fluxes vs. precip. amounts), Figure 1 has been replaced with the new Figure 1 (scatter plot), and Figure 2 and Figure 3 have been merged in an unique Figure 2 in which we reported the rainfall vs Fluxes. In addition, as you suggested, in the new Figure 2, the bars were enlarged in order to make more clear and easy to read them. Please, see the new Figures 1 at page 7 and 2 at page 9 in the revised version of the manuscript. We hope that the revised Figures (1-3) will meet the comments and suggestion of both the reviewers. Thank you for your input.

8. Page 8, lines 9-12. “Deposition of atmospheric Hg at any given location is influenced by factors such as : :” Since Hg deposition mostly consists of scavenging oxidized Hg forms (GOM or PBM) the list of factors should also include the oxidizing capacity of the atmosphere. The oxidation chemistry can be a dominating factor for Hg deposition at least in remote regions.

Reply: Yes, valid comment related to the oxidizing capacity of the atmosphere as another important factor. We revised these sentences including this important factor in the text. Please, see now at page 8, lines 14-22. Thank you.

9. Page 10, lines 3. Probably, the units of the average wet Hg deposition flux should be $\text{ng m}^{-2} \text{d}^{-1}$ instead of $\text{ug m}^{-2} \text{y}^{-2}$.

Reply: We thanks this reviewer for this important point. According also to the other two Referees regarding the precision of the measurements as well as the method firstly adopted to calculate the deposition fluxes, based on the exact sampling days at each site for each year, (Please, see also the comments n.2 & 3 of the second referee and related our replies as well as the comment number 6 in “specific comments” of the third referee and related our reply), we decided to revise all calculation taking into account the sampling frequency at each site for each year, which unfortunately was not constant across the sites. For this reason we decided to normalize all rainfall and fluxes data taking into account the two-weeks (15 days) as reference period based on the standard operating procedure (SOP) for THg in precipitation adopted within GMOS network. Therefore, the units that firstly referred to the number of days during the first analysis of the results, have been ruled out the text according to the revised calculation performed. Please, see the revised text of the manuscript. Thank you.

10. Page 11, line 21. “...showed the lowest both deposition amount (264.9 mm) and...” Should it be read as “precipitation amount”?

Reply: Yes, that’s right. We corrected it, thank you. Please, see at page 11, lines 21-22.

11. Pages 27-28. The Conclusions are too general and does not contain any particular findings on spatial trends and seasonal variation, factors affecting Hg wet deposition etc.

Reply: The conclusions have been integrated and reorganized according to your suggestion and comment. Please, see the revised version of the manuscript at the section “Conclusion”, pages 27 and 28. Thank you.

12. There are also numerous typos throughout the text which need spell-checking.

Reply: Yes, thank you very much for highlighting this. We revised the manuscript to correct typos throughout the text. Please, see the corrections done within the whole new revised version of the manuscript which is now improved. Thank you once more.

Anonymous Referee # 2

1. The paper deals only by wet deposition. What about dry deposition? The title should be changed then to "...Hg wet deposition fluxes..."

Reply: Yes, we revised the Title according to. Please, see the revised version of the manuscript.

2. P6 - What is the uncertainty of Hg wet deposition flux calculations?

Reply: As reported within the manuscript, because of different meteorological and climatologically conditions of the sites, the precipitation was not collected over an entire year at each station due to limited amount of precipitation samples occurring during specific periods (i.e., dry seasons). Therefore, all flux calculations reported herein used rain depth determined from the bottle catch and the uncertainty of Hg wet deposition flux calculations is therefore strictly linked to the precipitation volume of each sample. Weighed sample aliquots (50–100 mL) are pretreated following the standard procedure reported within the manuscript. Average blank values was determined for each analytical run and subtracted to determine sample Hg concentrations. Average analytical uncertainty for Hg in precipitation has been calculated with the relative standard deviation (RDS), where the RDS is the standard deviation of the three replicate analysis of each sample. Low precipitation volume samples (<1.5 ml) have very high uncertainty and a very low impact on the volume-weighted mean concentration, and were therefore not reported. The bi-weekly precipitation volume-weighted mean (VWM) concentration was determined using data only from samples considered valid. In addition, all data regarding rainfall amounts and Hg fluxes have been normalized in respect to the ideal time-sampling period that is equal to 15 days, as previously established in our GMOS Standard Operating Procedures. Both rainfall and Hg fluxes have been reported and discussed within the paper by the box and whisker plots, showing the variability of the available samples collected at each site and each season of observation. Furthermore, in Table S3 and S4 of the Supplemental Material, we also provided, on annual basis, the number of single collected sample with related basic statistics (min, Max, mean and St. Dev.). Thank you for your comment.

3. The data in the manuscript should be provided according to the precision of measurements.

Reply: Thank you for your comment. We think that this comment is strictly linked to the previous comment reported above (n. 2). We appreciate very much the reviewer for pointing out this important issue. Please, see our reply reported above as well as the revised version of the manuscript which now also includes a supplementary material document. The supplementary material added to the revised manuscript also includes additional two new tables (Table S3 and Table S4) reporting useful information and statistics about measurements and data calculation. Please, see the revised version of the manuscript and the supplementary material included. Thank you.

4. Although the Hg source identification is not the subject of the present paper it would be interesting to make a comparison with the study performed by Sun et al. ES&T, 2014. There are several parts in the manuscript where sources of Hg are discussed, which could be further supported by stable isotope analysis especially in China and their relation to global emissions (P20).

Reply: We appreciate very much the suggestion of the reviewer regarding the work performed by Sun et al. (2014), therefore, to accommodate the above comments, we focused our attention on the results and interesting study performed on Hg stable isotope signatures of coal deposits worldwide and historical coal combustion emission. At the same time, as the reviewer pointed out the identification of Hg source is not the subject of the present paper, since in this study, we limit our discussions to assess Hg wet deposition fluxes at some GMOS site (including regions where Hg measurements in air and deposition have not performed before, such as Tropics and Southern Hemisphere) and their seasonal and inter-annual variation reporting somewhere possible impact/effects on data recorded from mixing of different emission sources based on previous discussion and interpretation published in the literature at several GMOS sites part of them already established before the GMOS network and published historical series of Hg data in ambient air and precipitation. Therefore, according to the above discussion (and discussion in the text) and due to the lack of direct evidence and/or experiments on Hg isotope fractionation during atmospheric Hg transformation and deposition in this study to better identify Hg emission sources, we found difficult to start and include here a discussion on this topic. This also to avoid just include a and/or few sentences which in contrast should require more investigation and studies on this important and very interesting issue. Therefore, we thanks to this reviewer to highlight this issue and will take into account the suggestion for a future work on Hg precipitation data and the influence that the contributions from different sources may be better explained throughout Hg isotope fractionation factors and composition which should be updated whenever more data are available also for modeling applications. Thank you once more for your suggestion and comments.

5. The spell-checking is needed throughout the paper – several typos present.

Reply: Yes, Thank you. Following your input and the input also by the other two reviewers on this issue, we revised throughout the manuscript the several typos and corrected them according to. Please, see the revised manuscript.

Referee # 3 (Mark Cohen)

Specific Comments

- 1) The title is misleading, suggesting that both dry and wet deposition is being reported. Perhaps the title could be reworded to be something like the following: “Five-year records of mercury wet deposition flux at GMOS sites in the Northern and Southern Hemispheres”

Reply: Yes, thank you. We revised the Title according to your suggestion as well as the other two reviewers. Please, see the new Title of the manuscript. Thank you.

- 2) Page 2, Lines 8-11. Could mention that dry deposition is often estimated via models using measurements of ambient concentrations of mercury and meteorological parameters.

Reply: Yes, that's right. Following your suggestion, we integrated the sentences according to. Please see at page 2, lines 8-12.

- 3) Page 2, Lines 29-30. What is the gradient in northern Europe?

Reply: The north-south gradient in atmospheric mercury concentrations in northern Europe has been discussed in a manuscript published by Wangberg et al. (2001) (Atmospheric Environment Journal) which reports a summary of the results obtained in the framework of two joint MAMCS-MOE EU-funded projects, and in particular, the above paper reports the results from the MOE (Mercury Over Europe) project, in which has been discussed the north-south gradient for TGM with the highest values in the south considering measurements of TGM performed at six monitoring sites in North Europe, and confirming these findings reported in previous studies at the same sites (i.e., Smolke et al., 1999). In the manuscript Sprovieri et al. (2016) this has also been deeply discussed in the framework of the GMOS global network, thus considering several monitoring sites and confirming the gradient north-south, but extending the discussion from the Northern Hemisphere to the Southern Hemisphere, throughout the Tropical areas. Please, see Sprovieri et al. (2016) published in ACP journal. In this paper, in fact, there is a whole Section (Section 4.2) dedicated to the Northern – Southern hemispherical gradients in which we calculated the probability density functions (PDFs) of the 2013 and 2014 data and related histograms following the Scott rule, and, considering the mean (X) of the experimental measures for the northern (XN), southern (XS) and tropical (XT) groups and the confidence intervals evaluated from the *Student t test* among them, we observed a clear gradient of GEM concentrations from the Northern to the Southern Hemisphere (XN > XT > XS). The spatial gradient observed from northern to southern regions is also highlighted in two Figures (5 and 6), which also report the statistical monthly distribution of GEM values at all GMOS sites in the Northern and Southern hemispheres as well as in the tropical area. The gradients calculated are reported within the following table:

Northern (XN), Southern (XS) and Tropical (XT)	Mean X (2013-2014)	Δ ng m ⁻³
XN	1.54	0.31
XT	1.24	0.25
XS	0.99	0.56

For major details, please, see the paper Sprovieri et al. (2016), ACP. Thank you for your comment.

- 4) Page 3, Lines 7-8; and page 5, Lines 6-8. Why were these particular 17 sites chosen out of the 43 monitoring stations worldwide? Why were some sites excluded?

Reply: We thanks the reviewer for point out this important issue. The GMOS network to date consists of 43 monitoring sites globally distributed, and it has been established within the EU-funded project "GMOS" which start at the end of 2010. Several sites managed by the GMOS partners received funds to establish their own site during the development of the project, therefore, not all sites started together all Hg measurements in ambient air and precipitation, therefore, we chosen the most representative sites looking at their location (i.e., southern hemisphere, tropical

sites etc.) and their data coverage in order to have a number scientifically representative of results to discuss the data in different regions of the world. Obviously, this is a first tentative to discuss all together data from sites “globally” distributed taking into account that there are yet several difficulties linked to different meteorological conditions, emission sources and so on. Therefore, it well known for us that additional sites covering different regions and further investigations need to better understand the mercury fate and transport. GMOS network is ongoing and other additional sites are adding to the global network, therefore, we hope to collect in the next future more information and valid data to be discuss along with modeling applications for providing new insights in the mercury chemistry on regional and global scale. Thank you once more for your comment and suggestion.

5) Table 1. Several questions and suggestions to consider:

- (a) “Elev.” – could give units (m-asl) in the table. The units are given later in the text, but for clarity, could be included in the table.
- (b) The Sampling frequency could be included, e.g., 2-weeks for some sites, etc.
- (c) What is the meaning of sites listed as “M/S” and “S/M”?
- (d) If the site is a member of a national/regional network, this network could be listed.
- (e) The years of data collection could be noted for each site (e.g., 2013-2014, etc.)
- (f) In my opinion, would be very helpful to show the sites on a global map, perhaps with insets with close-ups as needed for clarity (e.g., for Europe)

Reply: We thanks the reviewer for point out several suggestions related to Table 1. We including them (point: a, c, d, and e) according to. Regarding point (f) we included a map named as Figure 1S within the supplementary material document added to the manuscript. Please, see the revised Table 1 in the revised version of the manuscript at page 4 of the revised version of the paper, as well as the supplementary material. Regarding the point (b) on the sampling frequency for each site, we included within the supplementary material two new tables (S3 and S4) in which we reported the exact number of the total yearly sampling days as well as the total number of single samples for each site. From these detailed information, it is possible to draw the sampling frequency related to each site for each year. Unfortunately the sampling frequency was not constant across the sites. For this reason we decided to normalize all rainfall and fluxes data taking into account the two-weeks (15 days) as reference period based on the standard operating procedure (SOP) for THg in precipitation adopted within GMOS network. Thank you very much for your input and suggestions.

- 6) Table 2, and associated text. It is not clear to me what “n days” data mean, and why it would be less than ~365 days per year. At some points in the manuscript, it seems that it might be being implied that if there was no precipitation during a given period (e.g., page 6, lines 20-21), then that period would not be reported as being a day of sampling for that year? But I don’t think that this is what you mean. My understanding of wet deposition samples is that the collector is in the field for a certain period (e.g., 2 weeks) and any precipitation that falls during this period is collected. So, in the usual case, if the site is operational, then the sampling generally occurs for the entire year, i.e., ~365 days. There may be some sampling periods where no precipitation is collected, but this is still a “sample” to be counted in the

number of sampling days for that year. So, it would be helpful to clarify what is meant for each site, for each year, when the number of sampling days is less than 365. Was the site “closed” for the non-sampling days, i.e., the collector was not being operated? And if this is the case, and since there are seasonal patterns to precipitation and mercury wet deposition, it is not clear to me that normalizing the measured deposition by the number of sampling days is a reasonable approach. In other words, the periods when the sampler was “on” would not necessarily be representative of the “average”. I’m not sure if it’s really useful to present data for fragments of years, given the seasonality, and given that the dates of collection are not given. In my opinion, it might be best just to give the data for a site when an entire year of samples was collected (or at least most of the year).

Reply: We appreciate very much the reviewer for pointing out this important issue. We think that this comment is strictly linked to the previous comment reported above (Referee 1, comment n.6). Therefore, please, see also our related reply to the comment n.6) as well as our reply to your comment reported above (n.5). The “n days” means the number of days during which each station was able to collect wet deposition samples. Unfortunately, GMOS wet deposition samples have been carried out with irregularity due to technical troubles with stuff and some problems in situ mainly found at those stations that began their Hg monitoring within the GMOS project. Therefore, when we had less than 365 days per year it means that, even if the corresponding station was “open”, the collector was not being operated. Otherwise, since many GMOS stations provided wet deposition data recorded for the first time at their locations, we believe that, even if with a partial coverage, they could be helpful in making possible a global picture of the issue under study. With this in mind, in this revised version of the paper we also tried to improve the presentation of data by normalizing them in respect of the ideal time-sampling period that is equal to 15 days, as previously established in our GMOS Standard Operating Procedures. In this way, the box and whisker plots, now report a consistent picture showing the variability of the wet deposition samples available on seasonal basis. Furthermore, in this revised version of the paper, we included additional information regarding the variability on annual basis of the wet deposition fluxes available at each GMOS site (See Tables S3 and S4 in the Supplemental Material).

- 7) Page 5, Lines 14-15: Additional description could be given in the text regarding the “bulk-modified” sites, e.g., at least a few sentences describing the sampling protocol at these sites. E.g., what does “bulk-modified” mean? Also, should be noted that bulk-collection sites collect some dry deposition.

Reply: We thanks the reviewer for point out this issue. We integrated this part of the manuscript following this comment (please, see the revised version of the manuscript at pages 4 and 5, lines 10 and 1- 4, respectively) including also the reference in which a detailed description of the bulk sampler used within the network is reported and its equivalence with the wet-only collectors. In addition, we replaced in Table 1 the “Bulk-modify” with the “IVL-Bulk” sampler which is the correct definition of the samplers that the sites reported in Table 1 used. Thank you for your comment.

- 8) Page 5, Lines 28-30. As noted above in comments on Table 1, it would be helpful to give the sampling frequency of each site.

Reply: Yes, thank you. Please, see our reply to your previous comment n.5 and the additional Tables reported within the supplementary material document added to the manuscript.

9) Figure 1. Several comments/suggestions:

(a) Figure is too small to read easily. One suggestion would be to switch the x/y axes, i.e., put the sites along the x-axis on the bottom, and the flux on the y-axis. And then, use the whole width of the page, so that the data can be more easily distinguished. Another suggestion might be to use symbols rather than bars.

(b) In the text, you refer to European sites extensively, and it would be helpful if these were grouped in the Figure. I know that you ordered the sites by latitude. But, in my opinion, you refer so many times to the “European sites” and refer to trends, etc., that it is really inconvenient to have to filter out the Chinese sites, etc.

(c) Could consider showing a scatter plot of deposition flux vs. latitude instead, or in addition.

Reply: We thanks the reviewer for detailed comments and suggestions. We considered at the end firstly the suggestion (c) replacing the old Figure 1 with a scatter plot of deposition flux vs latitude (please, see new Figure 1 at page 7); in addition, we also reported the deposition flux and the rainfall as an unique figure (see Figure 2 at page 9) in order to make more clear and simplify the interpretation/explain the results and avoid constantly going back and forth between the separate flux and the precipitation plots, as you also suggested within your comment reported later (n.16). Moreover, in order to improve and to make easier the Figure 2 reported in this part of the manuscript, we followed also your suggestion (a) switching the x/y axes, and enlarging the bars so that the data can be more easily distinguished. In addition, taking into account the comment n. 16 of the reviewer reported below and elsewhere, we re-organized all the Figure(s) related to both Section 3 and Section 4, including in each sub-section a Figure reporting the deposition flux as a function of precipitation amount according to. Thank you once more for your input and effort.

10) Page 7, Lines 4-5. Seems that there are 11 sites in the Northern Hemisphere, rather than 10? And you discuss the European sites and Chinese sites extensively, but not the Russian site LIS. Why is this? Also, here you say 7 European sites, but Figures 4-7 for European sites show only 6 sites. Could mention at some point why is the CMA site in Italy is not included in Figures 4-7.

Reply: We thanks to the reviewer for point out the number of sites in the Northern Hemisphere. There was a mistake in the text. In the revised manuscript we corrected the number of the sites at page 6, line 26, replacing 10 with 11 sites. We didn't discuss the results related to the Russian site because of the number of rainy samples was unfortunately inconsistent for any discussion or conclusion also from statistical point of view. Regarding CMA we didn't included within the seasonal variation graphics because for this site we have available few samples related to only two seasons and thus not covering a whole year (see Tables S1, S2, S3, S4). Therefore, we decided to rule out the site from the discussion of the re-organized Figures 4-7. The Figures 4-7, now became Figure 3, 4 and 5. Please, see the revised Figures at pages 13, 14 and 15, respectively, included within the revised manuscript. Thank you.

11) Page 7, Lines 7-9. The trend is not that clear in Figure 1, partly because the Russian and Chinese sites are interspersed in the Figure with the European sites. As noted above, I understand that you've listed the sites according to latitude, but ultimately, I think might be

clearer if you group by region first, given that the discussion is predominantly carried out by region. Also, the LON site does not seem to fit the European trend noted.

Reply: We thanks to the reviewer for this suggestion. However, as reported in our reply above, Figure 1 has been replaced with the new Figure 1 which consists in a scatter plot as suggested by this reviewer leaving the x/y axes as the original Figure 1 in order to make, in our opinion, more clear for the readers the general trend of the data observed according to the latitude. In addition, the re-organization of the Figures (firstly Figs 1-3, and now Figure 1 and Figure 2) made following your previous comments and the suggestion repeated later, could be from our point of view, a good compromise to better understand the discussion presented here and elsewhere not penalizing the general trend observed across the sites, particularly in the Northern Hemisphere. The Figures have been reorganized taking into account the precipitation amounts for each site and for each year in corresponding with the wet deposition fluxes as suggested. Please, see the new Figures 1 and 2 at pages 7 and 9, respectively. In addition, even according to your suggestion, in the Figure 2 we switched the x/y axes and enlarged the bars which are more clearly visible and in vertical position. Please, see our reply to your comment and suggestion n.9. We hope now that these changes could help to make more clear the discussion and interpretation of the results. Regarding LON site, we corrected this sentence in agreement with you. Please, see at page 8, lines 6-8 in the revised version of the manuscript. Thank you for your input and suggestions.

12) Page 7, Line 10. ... no north-south spatial trend has been observed.

Reply: Yes, that's right. We integrated the sentence according to. Please, see at page 7, lines 1-2, thank you.

13) Page 8, Lines 9-12. Wet deposition of atmospheric Hg at any given location...

Reply: Corrected, please, see the revised version of the manuscript at page 8, line 14. Thank you.

14) Page 8, Lines 9-12. Wet deposition also depends on the type of precipitation (e.g., snow vs. rain), and the height and thickness of the precipitating cloud layer in the atmosphere, and the degree of convection involved. These are included at several points later in the document, but when I read this at this point, it seemed like important factors were being left out.

Reply: Yes, that's right. We integrated these factors according to along with other important factors suggested by the Referee 1. Please, see the revised manuscript at page 8, lines 15-22. Thank you.

15) Figure 3. As noted at several points in the manuscript, the relative proportion of snow vs. rain (or frozen vs. liquid) precipitation can be an important factor in interpreting the wet deposition data. Are there any site-specific data on this could be shown in Figure 3, or in a different figure?

Reply: Yes, we agree with you. Unfortunately, we have no info/data from the sites on this. They didn't make this classification of each sample recorded. This also probably due to some logistic difficulties sometimes to reach the remote sites.

16) Page 9, Lines 1-9. The idea that more wet Hg deposition occurs with more precipitation is mentioned here and at several other points in the document. I think it might be really useful

to show Figure(s) that show the deposition flux as a function of precipitation amount. This might be easier to interpret/explain than constantly going back and forth between the separate flux and the precipitation plots.

Reply: We agree with your suggestion. We re-organized all the Figure(s) related to both Section 3 and Section 4, including in each sub-section a Figure reporting the deposition flux as a function of precipitation amount according to. Please, see also our reply to the comment n. 9 in which we listed the revisions made in both manuscript and Figures. Please, see the revised manuscript and the new Figures reported in the Sections 3 and 4. Thank you.

17) Page 11, Lines 26-34. Here, you present arguments that suggest that the relatively high Hg wet deposition at CPT is due at least in part to contributions from local and regional sources. But then on page 26, lines 15-17, you cite a study that purportedly concluded that no significant local anthropogenic influences were found in Hg concentrations at CPT. How can these conflicting situations be reconciled?

Reply: We thanks the reviewer for highlighting this important point. In this part of the text we reported some research and results obtained at CPT in the past by the authors cited in the paper, and in particular we referred our results of Hg wet deposition observed in 2013 to a study performed by Brunke et al. in 2014, and assuming as first instance that these high Hg concentrations could be due to influences from anthropogenic sources considering the origin of air masses prevailing at CPT during specific period of the year (Please, see the revised version of the text at page 10, lines 5-8). However, as pointed out at page 26 in the revised manuscript, lines 2-10, we also reported the results obtained in a more recent work by the same authors (Brunke et al., 2016) where, throughout additional measurements performed, they concluded that during the same period of the year the high Hg concentrations recorded at CPT are probably related to other processes occurring more than to significant local anthropogenic emission sources, and highlighting the positive correlation they found between GEM/THg concentrations, and the Southern Oscillation Index (SOI) giving such conclusion:... *“suggesting that both GEM and THg concentrations are primarily influenced by large scale meteorology which in turn controls Hg emission sources in terms, for example, of enhanced sea surface temperature that could increase large scale droughts leading to a raised biomass burning”*... In addition, we have made thorough revision in the paper to correct inaccurate words to make clear the sentences. Please, see the revised version of the manuscript related to this issue at both page 10 and page 26. Thank you once more.

18) Figures 4-5-6-7. Are the box-whisker plots showing statistics for the sample-by-sample distributions for each season? If so, then it would definitely be important to know the sampling frequency. Might be useful to state what the boxes mean (25%, 50%, 75%?), and what the whiskers mean (5%, 95%?)

Reply: We thanks the reviewer for highlighting this important point. Regarding the sampling frequency, please, see our reply to your comments n. 5, 6 and 8 and the related information reported within the new Tables (S3 and S4) in the supplementary material added to the manuscript from which is possible to understand the sampling frequency related to each site. In addition, following your suggestion, we reported what the boxes mean and the whiskers mean within the caption of the Figures. Figures 4-5-6-7 as well as the others following them included within Section 4, have been replaced/re-organized following your comment and suggestion reported elsewhere, therefore, the

new Figures have also a new numeration (Figures 4-5-6-7 are now 3, 4 and 5) and the other accordingly. Please, see the Figures and captions within Section 3 and Section 4. Thank you.

19) Figure 5. Seems like could reduce y-axis to 0-60 ng/lit to show data more clearly.

Reply: yes, we agree with you and we revised the old Figure 5 (now Figure 3) according to. Please, see the revised version of the manuscript. Thank you.

20) Figures 4-5-6-7: Again, maybe could add a figure that shows flux as a function of precip... This might be very illuminating.

Reply: Done. Please, see our reply to your comment n. 16 and suggestion reported elsewhere on this issue. Please, see the revised version of the Figures included within the manuscript. Thank you.

21) Figures -5-6-7: Why is CMA not included as a European station? I guess because not enough data?

Reply: Yes, that's right, the coverage related to this site is not enough consistent to drawn interesting conclusion and/or discussion. Please, see also our reply to your comment n. 10 on this issue. Thank you.

22) Figures 4-5-6-7: Would be useful somehow to show degree of solid vs. liquid precip, e.g., in Figure 4, if these data were available.

Reply: Unfortunately, we have no data and detailed information from the sites on this. Please, see also our reply to the above comment n.15.

23) Page 13, Line 5: Why were only 91 days sampled at the site? Was this because there was no precipitation, or was this because the site was simply not operated during that time?

Reply: Yes, the last one. The BAR site started the THg measurements with a delay in respect to most of the GMOS sites considered in the paper. Please, see also our reply to your comment n.4 about our chosen of the GMOS sites among the 43 which constitute to date the network and our reply to the comment n.6 related to the first Referee about the "data coverage". Thank you for your comment.

24) Page 15, Lines 3-11. Here, and in some other places in the document, it seems that you are just restating the information that can be clearly seen in the Figures. The manuscript is pretty long, and perhaps some efficiency could be obtained by omitting at least some of this type of reiteration?

Reply: Yes, we agree with you. We tried to optimize/summarize through the manuscript the sentences related to what is clearly showed in the Figures according to. Please, see the revised version of the manuscript as examples at page 13, lines 8-11; page 14, lines 7-11, page 15, lines 1-7 and so on. Thank you.

25) Page 15, Lines 16-17. As mentioned earlier in other contexts, it might be really helpful here to show a graph of flux vs. precipitation amount.

Reply: yes, thanks. Please see our previous reply to your comment reported earlier in other context and the revised version of the manuscript and Figures. Thank you once more.

26) Page 17, Line 6. What emissions are larger in the warmer months?

Reply: In this part of the paper we want only to point out that during such period of the year, in this case, during the warmer months, the concomitance of meteorological conditions (such as high temperature, higher solar radiations etc.) along with other existing conditions and characterizing a such site (such as local emission sources) could enhance the effects on Hg chemistry that in other situation/conditions (for example during the cold months) not occurs or not occurs with the same “intensity”. We reported, in fact, in this part of the manuscript the example of some oxidants (i.e., O₃, OH radicals etc.) that under conditions of high temperature, higher solar radiation typical of warmer months, show higher concentrations and the photo-oxidation processes are enhanced. These conditions/parameters all together could give enhanced oxidized mercury species in the atmosphere for the conversion of GEM to GOM. In addition, we have made thorough revision in the revised version of the paper to replace “was likely” with “could be” being the first an inaccurate word which could give unclear the sentences (i.e., page 16, line 6). Please, see the revised version of the manuscript. Thank you for your comment.

27) Page 19, Lines 2-3. Perhaps too much to ask, but would it be possible to show maps of emissions in relation to the sites?

Reply: We thanks the reviewer for his effort and several input and suggestion that we try to strictly follow because we strongly believe that they are improving the quality of the paper. Anyway, as the reviewer reported above in one of his comment, the manuscript is pretty long, although we tried to synthesize and when it was possible delete several sentences, therefore, considering also the number of Figures and new Tables included partially in a supplementary material document added to the manuscript with further and useful information, we prefer not added additional map or figures. This also considering that some comments reported in the paper on this issue are related to previous studies already published in the literature, therefore, we think that inserting the references in the text this could be sufficient to understand the discussion reported in our manuscript. In addition, we would like to point out that in the same special issue there are other some papers (i.e., De Simone et al.) where modeling application and measurements performed at some of the same GMOS sites are discussed along with useful emission maps. Thank you once more.

28) Page 21, Lines 1-2. Sorry to be repetitive, but again, could may be show a graph of flux vs. precipitation amount.

Reply: To accommodate the above comments of this reviewer, we have revised the Figures as also reported earlier in our replies to the comments on this issue . Please see our previous reply and the revised version of the manuscript. Thank you.

29) Page 22, Lines 10-15. Not sure what you mean by “washout”. Are you referring to below-cloud scavenging of PBM by falling precipitation? Perhaps you could explain a bit more about the phenomena that you are describing here.

Reply: yes, we refer to the below-cloud scavenging of PBM by falling precipitation. Following your suggestion, we explain a little bit about the phenomena. Please, see the revised version of the manuscript at page 18, lines 8-11. Thank you.

30) Page 26, Lines 1-3. Here, and at a few other points, you note patterns in relation to GOM or other measurements. Would a graphic be useful here to show the relationship, e.g., GOM vs. volume-weighted-mean concentration in precip?

Reply: Thank you for your suggestion. In the original version of the manuscript, we didn't include additional graphic and or details on Hg measurements in air to avoid to report and/or repeat some issue and results reported within other recent manuscripts and/or included within the same special issue (such as Sprovieri et al. where atmospheric Hg results obtained within the GMOS network have been discussed and presented), even if has been often highlighted the need of additional investigation on the relationship of atmospheric Hg speciation measurements vs Hg in precipitation. Anyway, in the revised version of the manuscript, we rewrote and/or deleted some sentences according to the new calculation performed for rainfall amounts and wet deposition fluxes normalizing the weighted data on each sample at each site with 2-weeks reference time as described earlier. The new analysis for AMS data samples gave us different results on the seasonal trend of the precipitation amounts and THg wet deposition flux patterns. These findings make the previous instance on the relationship between GOM vs Hg precipitation data no longer valid. Therefore, we delete the sentence related to the possible correspondence in this case among atmospheric and precipitation data. We are grateful to the reviewer for this and for his previous suggestions which have prompted us to make a more accurate and scientifically objective analysis than that previously made taking into account only the number of sampling days. Please, see the revised version of the manuscript at page 24, lines 9-13, and page 25, lines 1-5. Thank you.

31) Page 27, Lines 30-31. You state that early models tended to overestimate the influence of local emissions sources. This may or may not have been true, for one or more models, but I feel you'd need to cite a lot of different papers really make this statement. To me, seems like an overly provocative statement, and one that is not really needed for the paper? The general idea that observations are critical for model evaluation is certainly valid, but I don't think you can (or need to) make this sweeping statement about "early models". Indeed, Sunderland et al (2016) have recently pointed out that "early models" may have significantly underestimated the influence of local emissions sources! Sunderland, E. M., C. T. Driscoll, J. K. Hammitt, P. Grandjean, J. S. Evans, J. D. Blum, C. Y. Chen, D. C. Evers, D. A. Jaffe, R. P. Mason, S. Goho and W. Jacobs (2016). Benefits of Regulating Hazardous Air Pollutants from Coal and Oil Fired Utilities in the United States. *Environmental Science & Technology* 50(5): 2117-2120.

Reply: Thank you for your comment. We absolutely agree with you regarding the sweeping statement reported in this part of the paper. We revised this section taking into account what recently Sunderland et al. (2016) pointed out. Please, see the revised version of the manuscript at page 27, lines 22-24 within the Section "Conclusions". Thank you once more.

32) Page 27, Line 31. Is this really all available GMOS wet dep data, or just the data from selected sites for selected years? Also, are the GMOS wet dep data (and other data?) available? Perhaps this could be mentioned?

Reply: yes, we agree with you, thank you. In this paper we referred only to selected monitoring sites which provided Hg data in precipitation during the development of the GMOS project. Please, see the revised sentence in the new text at page 27, lines 25-28.

However, we would like to point out that Hg measurements across the GMOS network are ongoing, including wet deposition samples, thus the number of the ground-based sites is growing, and the data from them as well. The data coming from the GMOS network are available upon request and protected by a policy document available for the scientific community (i.e., it can be downloaded from the GMOS web page (www.gmos.eu), please, see the following link: http://www.gmos.eu/public/GMOS-Governance_Data_Policy_rev160705.pdf).

33) Page 28, Line 1. Having data a “remote” sites with few local or regional sources is important, for sure, but having data a sites that are influenced by local and regional sources are also important for better understanding of Hg atmospheric fate and transport (and model evaluation), etc.

Reply: Yes, thank you, we agree with you. We revised the statement according to. Please, see at page 28, lines 8-10. Thank you.

Technical Corrections

1) Page 3, Line 1: ... long-term measurements of ambient Hg concentrations and measurements of Hg wet deposition fluxes were lacking...

Reply: Yes, corrected, thank you. See at page 3, lines 6-8.

2) Table 2. There is a vertical line in the top of the table (see clip below, with red circle), that I think should be removed.

Reply: Yes, corrected, thank you. Please, see the supplementary material added to the manuscript where, now Table 2 becomes Table S1.

		2011					2012				
		Annual Wet Dep. Flux [$\mu\text{g m}^{-2} \text{ yr}^{-1}$]	Rainfall [mm]	ndays [d]	Weighted HgT [ng L^{-1}]	Aver Wet Dep. Flux [$\text{ng m}^{-2} \text{ d}^{-1}$]	Annual Wet Dep. Flux [$\mu\text{g m}^{-2} \text{ yr}^{-1}$]	Rainfall [mm]	ndays [d]	Weighted HgT [ng L^{-1}]	Aver Wet Dep. Flux [$\text{ng m}^{-2} \text{ d}^{-1}$]
Northern Hemisphere	NYA	-	-	-	-	-	0.9	238.6	350	3.8	2.6
	PAL	2.9	407.4	363	7.1	8.0	1.9	278.6	332	6.8	5.7
	RAO	5.8	646.6	364	8.9	15.8	6.5	621.8	366	10.4	17.8
	MHE	-	-	-	-	-	0.9	393.7	113	2.2	7.6
	LIS	-	-	-	-	-	0.2	17.4	18	9.7	9.4
	CMA	-	-	-	-	-	-	-	-	-	-
	ISK	5.1	680.2	224	7.5	22.7	8.4	1349.7	363	6.2	23.2
	MCH	2.8	264.6	119	10.6	23.6	4.8	569.4	228	8.4	21.1
	LON	-	-	-	-	-	0.3	88.2	19	3.9	18.1
	MWA	-	-	-	-	-	0.3	79.5	127	4.3	2.7
MAL	4.3	1543.2	222	2.8	19.5	3.2	971.5	202	3.3	16.1	
Tropics	SIS	-	-	-	-	-	-	-	-	-	-
	CST	-	-	-	-	-	7.4	797.1	155	8.1	15.5
Southern Hemisphere	AMS	-	-	-	-	-	-	-	-	-	-
	CPT	0.3	133.5	119	2.1	2.4	3.8	260.3	147	14.6	25.8
	CGR	-	-	-	-	-	-	-	-	-	-
	BAR	-	-	-	-	-	-	-	-	-	-

3) Table 3. I think “uom” refers to “units of measurement”, but maybe clearer just to put the units, or spell out “units of measurement. Better yet to include the units directly in the table.

Reply: Yes, we followed your suggestion including the units directly in the revised Table 3 (now Table S2 in the supplement material document). Please, see Table S2. Thank you.

- 4) Page 6, Lines 31-32: ...the number of the sampling days as well as the annual wet deposition flux and average THg wet deposition flux calculated for each year in the period 2011-2015.

Reply: Thank you, the sentence has been corrected according to and integrated on the basis of new calculation performed. Please, see at page 6, lines 18-22.

- 5) Page 6, Line 32: As noted above, it is really unclear how valid any of the partial-year data are, given that it is unclear if the missing data are from rainy or dry seasons, etc.

Reply: Please, see our reply on this issue to your comment reported above within the “Specific Comments” section and the revised version of the Section 3 and 4 of the manuscript. Thank you for your input and suggestion.

- 6) Page 8, Line 17. ... during the 2011-2015 period are is reported in Figure 3

Reply: Yes, that’s right. However, this sentence in the manuscript has been changed. Please, the revised version at page 8, line 26-28. Thank you.

- 7) Page 11, Lines 19-24. Seem like sometimes you refer to sites using the 3-letter abbreviation, and sometimes you refer the sites using the full name of the site. Since the graphics all use the 3-letter abbreviation, maybe better to just use these in the text throughout. Could give the full name the first time it was mentioned, with the abbrev in parentheses, and then just use the abbreviation from then on?

Reply: Yes, thank you. We followed your suggestion. Please see the revised version of the manuscript.

- 8) Page 13, Line 13. “meteorological” is misspelled.

Reply: Corrected. Please, see at page 12, line 22. Thank you.

- 9) Page 20, Line 5. “rain” not “rainy”

Reply: Corrected, thank you. Please, see at page 17, line 4 of the revised manuscript.

- 10) Page 22, Line 5. ... The positive or negative correlation between THg concentrations and the precipitation amount has not been obviously observed at MAL where the rainy samples shows a fairly seasonal variability, during all seasons with lowest average rainfall in winter and the highest in fall...

Reply: The sentence has been integrated and corrected, thank you. Please, see at page 17, line 14 and page 18, lines 1-4 of the revised manuscript.

- 11) Page 23, Line 18. ... exhibit a seasonality in annual rainfall, ...

Reply: Corrected, thank you. Please, see at page 20, line 30 of the revised manuscript.

12) Page 24, Line 10. “fuels” not “flues”

Reply: Corrected, thank you. Please, see at page 21, line 6 of the revised manuscript.

13) Page 24, Line 12. “United States”

Reply: Corrected, thank you. Please, see at page 22, line 2 of the revised manuscript.

14) Page 24, Line 12. “waste incinerators”

Reply: Corrected, thank you. Please, see at page 22, line 3 of the revised manuscript.