Five-year records of mercury wet deposition flux at GMOS sites in the Northern and Southern hemispheres

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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 5 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. Interannual 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 in 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). Currently, Hg dry deposition is often estimated by models, using measured ambient concentrations of Hg and meteorological parameters, due to the lack of existing direct and accurate measurement methodologies (Sexauer Gustin et al., 2012; Zhang et al., 2012). Therefore, the investigation of Hg fluxes to terrestrial and aquatic surfaces in different parts of the world are often based on wet deposition measurements (Gratz et al., 2009; Fu 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 yr\(^{-1}\) deposited on land and 3700 Mg yr\(^{-1}\) into oceans (Mason et al., 2012). The preindustrial deposition rate has been estimated to be 1000 Mg yr\(^{-1}\) deposited on land and 2500 Mg yr\(^{-1}\) 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 emissions are still occurring widely around the world (Pacyna et al., 2010; Pirrone et al., 2010). In North America, seasonal patterns in THg concentrations in precipitation and Hg wet deposition amounts have been observed with the highest values in the summer and lowest values in the winter (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 in precipitation and wet deposition fluxes (Vanarsdale et al., 2005; Prestbo and Gay, 2009) due to their location closer 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 GEM which, throughout active oxidation processes, may also contribute to elevated deposition 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, 2007; Sprovieri et al., 2016). Hg wet deposition data are therefore important 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 European Monitoring and Evaluation Programme (EMEP); however, before the establishment of the Hg network by the Global Mercury Observation System (GMOS) on a 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 (Winberg et al., 2007; Prestbo and Gay, 2009; Sanei et al., 2010), 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 (http://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 multiyear comparison of Hg wet deposition worldwide and provides insights into annual and seasonal variations, as well as a spatial gradient in Hg deposition patterns.
Table 1. Station locations that are part of the GMOS 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 with respect to the Hg measurements carried out as speciated (M) or not (S). M/S or S/M indicates change of the site from master to secondary or vice versa. External GMOS partners are indicated in bold.

<table>
<thead>
<tr>
<th>Code</th>
<th>Name</th>
<th>Country</th>
<th>Lat</th>
<th>Long</th>
<th>Elev. (m a.s.l.)</th>
<th>Collector type</th>
<th>Years of sampling</th>
<th>Type*</th>
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<tbody>
<tr>
<td>1 NYA</td>
<td>Ny-Ålesund</td>
<td>Norway</td>
<td>78</td>
<td>90</td>
<td>11, 88</td>
<td>IVL bulk</td>
<td>2012–2015</td>
<td>M</td>
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<td>2 PAL</td>
<td>Pallas</td>
<td>Finland</td>
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<td>24, 24</td>
<td>IVL bulk</td>
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<td>3 RAO</td>
<td>Råö</td>
<td>Sweden</td>
<td>57</td>
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<td>11, 91</td>
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<tr>
<td>4 MHE</td>
<td>Mace Head</td>
<td>Ireland</td>
<td>53</td>
<td>33</td>
<td>–9, 91</td>
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<td>5 LIS</td>
<td>Listvyanka</td>
<td>Russia</td>
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<td>85</td>
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<td>6 CMA</td>
<td>Col Margherita</td>
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<td>11, 79</td>
<td>wet only</td>
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<td>7 ISK</td>
<td>Iskrba</td>
<td>Slovenia</td>
<td>45</td>
<td>56</td>
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<td>2011–2015</td>
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<td>8 MCH</td>
<td>Mt. Changbai</td>
<td>China</td>
<td>42</td>
<td>40</td>
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<td>IVL bulk</td>
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<td>9 LON</td>
<td>Longobucco</td>
<td>Italy</td>
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<td>39</td>
<td>16, 61</td>
<td>wet only</td>
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<td>Mt. Walliguan</td>
<td>China</td>
<td>36</td>
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<td>12 SIS</td>
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<td>Mexico</td>
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<td>13 CST</td>
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<td>14 AMS</td>
<td>Amsterdam Island</td>
<td>TAAF</td>
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<td>wet only</td>
<td>2013–2014</td>
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<td>Cape Point</td>
<td>South Africa</td>
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<td>Cape Grim</td>
<td>Australia</td>
<td>–40</td>
<td>68</td>
<td>144, 69</td>
<td>IVL bulk</td>
<td>2013–2015</td>
<td>S</td>
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<tr>
<td>17 BAR</td>
<td>Bariloche</td>
<td>Argentina</td>
<td>–41</td>
<td>13</td>
<td>–71, 42</td>
<td>wet only</td>
<td>2014–2015</td>
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*T = master; S = secondary. TAAF indicates Terres Australes et Antarctiques Françaises.

2 Experimental

2.1 GMOS ground-based monitoring sites

The global Hg monitoring network has been established in the framework of the GMOS and presented in Sprovieri et al. (2016). It has been developed by integrating previously ongoing ground-based Hg monitoring stations as part of regional networks with those established as part of GMOS also in regions of the world where atmospheric Hg measurements were previously limited. To date, the GMOS network consists of 43 monitoring stations worldwide distributed and located in climatically diverse regions, including polar areas (Sprovieri et al., 2016). In the present study, we refer the discussion on Hg wet deposition to a representative number of 17 ground-based sites distributed in the Northern and Southern hemispheres (Fig. S1 in the Supplement). Table 1 provides key information on the 17 monitoring sites such as their location (country, coordinates, etc.), elevation (m a.s.l.), and type of monitoring stations (master and secondary sites) with respect to the atmospheric Hg measurements performed (Hg speciation and total gaseous mercury – TGM/GEM measurements, respectively) along with THg wet deposition sampling.

2.2 Sample collection, analytical procedure, and QA/QC

Precipitation samples were collected across the sites primarily using wet-only collectors, (i.e., N-CON MDN or the Eigenbrodt NSA 171 wet-only samplers). Where necessary, due to site constraints or operator availability, few GMOS sites (Table 1) alternatively collected bulk precipitation samples using the Swedish Environmental Research Institute (IVL) bulk sampler (Chazin et al., 1995; Munthe et al., 2011; UNI, 2010). The detailed description of the Swedish bulk sampler is reported elsewhere (Chazin et al., 1995; Munthe et al., 2011; UNI, 2010), where it also has been highlighted that utilizing clean handling and analysis technique is equivalent to using a wet-only sampler in remote areas, and/or areas not subject to large amounts of anthropogenic emission sources, as is the case in most of the GMOS ground-based sites. Within GMOS, special attention was paid with respect to protocol harmonization, data quality collection, and data management in order to assure a full comparability of site-specific observational data sets. During the imple-
mentation stage of the GMOS global network, harmonized standard operating procedures (SOPs) as well as common quality assurance/quality control (QA/QC) protocols have been addressed (Munthe et al., 2011) in accordance 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. Furthermore, the management of the measurement program at most of the GMOS sites consisting of analysis of all precipitation samples, cleaning procedures, and distribution of the sample bottles to all sites, has been performed by three reference laboratories (IVL, Sweden; CNR-IIA (CNR Institute of Atmospheric Pollution Research), Italy; and IJS (Jožef Stefan Institute), Slovenia), whereas the precipitation samples related to some other GMOS sites, such as in Russia – Listvyanka (LIS); in China – Mt. Waliguan (MWA), Mt. Ailao (MAL), and Mt. Changbai (MCH); and in South Africa – 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 intercomparison exercises for Hg in water (i.e., Brooks Rand Instruments Inter-laboratory Comparison Study). GMOS sites predominantly collected biweekly samples. However, considering the spatial distribution and the diversity of meteorological parameters and conditions characterizing the monitoring site locations, the sampling frequency was sometimes different across the sites. THg concentrations in precipitation samples, re-frigerated 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 US EPA Method 1631 (version E) (1631 U. E. M., 2002); each sample was first oxidized by BrCl (0.5 mL/100 mL sample), followed by neutralization with hydroxylamine hydrochloride (NH₂OH·HCl). Stannous chloride (SnCl₂) was then added to the sample to reduce Hg²⁺ to Hg⁰ 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⁻¹ (recovery 93–109 %). The standard curve was used within the coefficient of determination (r²) 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 (Code of Federal Regulations) 136, Appendix B) for Hg has been determined to be 0.02 ng L⁻¹. The minimum level of quantification (ML) has been established as 0.05 ng L⁻¹ 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 (CRM) 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 (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 climatological 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 precipitation amount collected at each site during the times that the Hg sampler was operating. This means that the THg wet deposition flux has been approximately calculated by the following equation:

\[ F_W = C_{Hg} \sum_{i=1}^{n} P^i \times 1/1000, \]

where \( F_W \) is the THg wet deposition flux (μg m⁻² yr⁻¹) and \( C_{Hg} \) is the VWM concentration of THg (ng L⁻¹). \( P^i \) (mm; 1 mm = 1 L m⁻²) represents the precipitation amount associated with each wet deposition sample.

It is also necessary point out that the “precipitation” and “rainfall” terms throughout this work have been interchangeably used and, further, rainfall included all forms of precipitation (i.e., rain, frozen). The frozen precipitation has been considered as “liquid rain equivalent”. In addition, to overcome the irregularity in time-sampling frequency of the rainy samples collected at each site (see Tables S3 and S4) the rainfall amounts as well as THg wet deposition flux weighted data have been normalized with respect to the ideal time-sampling period, equal to 15 days, as previously established in our GMOS standard operating procedures (Munthe et al., 2011). In this way, all data resulted comparable and harmonized.
3 Hg wet deposition patterns and interannual variability

The annual variations in THg concentration and wet deposition recorded at all 17 monitoring GMOS sites are summarized in Tables S1 and S2. Both tables list the monitoring sites according to their latitude and rain amounts collected, the number of the sampling days, as well as the annual wet deposition flux calculated for each site for each year in the period 2011–2015. The rainfall amounts as well as THg wet deposition flux weighted data have been normalized with a 15-day sampling reference as described in the section above. Annual THg wet deposition fluxes calculated at each site according to their latitude are shown in Fig. 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 11 sites distributed in the Northern Hemisphere, the discussion of the results will be separately related to the 7 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 and thus not representative enough for such a conclusion) and the 3 Chinese sites (see Tables S1–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-Ålesund – NYA, Norway; Pallas – PAL, Finland) to south in the Northern Hemisphere (i.e., Råå – RAO, Sweden; Mace Head – MHE, Ireland; Listvyanka – LIS, Russia; Col Margherita – 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 CPT, no historical records of THg deposition exist for the new stations established in the GMOS project.

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). Figures 1 and 2 show, from 2012 to 2014 (the period with more data coverage), a general increasing of THg wet deposition from the NYA station to Iskrba (ISK), Slovenia; this finding is particularly evident during the 2013 and 2014 (Fig. 2) event if the pattern is not apparent for the LON site in 2013 and for MHE and CMA for 2014, indicating the influence of other parameters and/or atmospheric transport pathways. In Fig. 2, the THg wet deposition fluxes calculated on an annual basis are reported, 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; (d) type of precipitation (rain or snow); (e) length of precipitation events which affect Hg concentrations; (f) height and thickness of the precipitating cloud layer in the atmosphere and the degree of convection involved; and (g) 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 should be strongly influenced by the precipitation amounts. In particular, the annual deposition amounts during the 2011–2015 period show the influence of the precipitation amount on Hg deposition between, for example, the RAO site and the PAL site (Fig. 2). The THg wet deposition fluxes recorded during 2011–2014 were, respectively, 5.8, 6.5, 4.2, and 6.3 µg m$^{-2}$ yr$^{-1}$ at the RAO site. This is more than 2 times higher than at Pallas during the same years (2.9, 1.9, 1.3, and 2.3 µg m$^{-2}$ yr$^{-1}$), and since the precipitation amounts are also a factor of 2 higher at the RAO site in comparison to 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 the 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 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 (µg m$^{-2}$ yr$^{-1}$) calculated, conversely, at MCH, MWA, and MAL shows 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 2012–2014 period, which is the most representative in terms of number of samples recorded, it is possible to see that the averaged THg wet deposition fluxes (ng m$^{-2}$ day$^{-1}$) 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) (Fig. 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 Yucatán Peninsula (Gulf of Mexico), the 2013 annual wet THg deposition flux was 7.4 µg m$^{-2}$ yr$^{-1}$, whereas the rainfall amount was 669.6 mm, which is lower than the rainfall recorded at the remote southern sites, such as Amsterdam Island (AMS) (833.2 mm rainfall), southern Indian Ocean, and Cape Grim (CGR), Australia (775.6 mm rainfall), where the annual wet Hg deposition flux recorded was considerably lower at 1.95 and 3.1 µg m$^{-2}$ yr$^{-1}$, respectively (see Tables S1 and S2). The 2013 and 2014 annual wet deposition fluxes recorded at SIS are comparable or higher than those observed at most GMOS sites in the Northern and Southern hemispheres (Tables S1 and S2). Because of the Hg deposition at any given location is dependent upon both THg concentrations (which have 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 emis-
sion sources has had a higher influence on the THg results. During the sampling period, SIS was typically influenced by air masses originating from the Atlantic Ocean coming from east–southeast, but crossing the Caribbean islands and/or Central/South America with occasional air masses coming from east–northeast mostly during the winter period and crossing the south of Florida and Caribbean archipelago prior to arriving 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, 2015). Shanley et al. (2015), in a study over 7 years (2005–2012) on Hg wet deposition in 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 US, and thus was 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 US ($\sim$2–5 µg m$^{-2}$ yr$^{-1}$) and higher in the eastern US (6–15 µg m$^{-2}$ yr$^{-1}$) due to increasing precipitation and location of important anthropogenic Hg sources. In addition, in the eastern US, a north–south latitudinal gradient exists in wet Hg loading, with wet deposition reaching a maximum in the southeastern US 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 US (Shanley et al., 2015). The high wet Hg deposition at SIS can be directly linked to the meteorological 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 in Puerto Rico (27.9 µg m$^{-2}$ yr$^{-1}$), an unpolluted tropical site crossed often by air masses detected at SIS prevalently in summer and fall and less so 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 with GOM at ground level but with 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 at 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 the Antilles 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 Yucatán Peninsula (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 an annual median of 1.03 ± 0.10 ng m$^{-3}$ 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 CGR in 2013 (Slemr et al., 2014). Both AMS and 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; Holmes et al., 2009; Sprovieri et al., 2003, 2010a, b) 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 bromine (Br) in the marine boundary layer (MBL) are an important source of GOM from oxidation of GEM that are 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 AMS and CGR. In 2013, among the southern sites, the highest annual THg wet deposition fluxes has been recorded at CPT (5.2 µg m$^{-2}$ yr$^{-1}$) which also showed both the lowest precipitation amount (264.9 mm) and the number of sampling days (Tables S1 and S2) compared to AMS (with an annual wet deposition flux of 1.95 µg m$^{-2}$ yr$^{-1}$, considering a rainfall of 833.2 mm) and CGR (with wet deposition flux of 3.1 µg m$^{-2}$ yr$^{-1}$, 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 µg m$^{-2}$ yr$^{-1}$) and comparable precipitation amounts and number of sampling days.
of the year before (see Tables 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–October) normally precipitation increased due to the passage of cold fronts moving from west to east (Brunke et al., 2016). In a previous study by Brunke et al. (2004), it was highlighted that CPT receives continental and polluted air masses more frequently during the winter period with air masses advected to the station from north to northwestern (Rautenbach and Smith, 2001; Brunke et al., 2004) regions 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, could be prevalently influenced by regional/large-scale emission sources during the sampling period. Measurements of atmospheric Hg deposition in BAR have been carried out for the first time from 2014 to 2015. The BAR site has been established inside a well-protected natural reserve in northern Patagonia, on the shore of the Gutierrez River, southeast of the Nahuel Huapi lake. GEM records at the BAR station resemble background concentrations comparable to levels found in Antarctica and other remote locations of the Southern Hemisphere with annual mean GEM concentrations of 0.9 ± 0.14 ng m⁻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 µg m⁻² yr⁻¹); however, it is necessary to point out that the number of samples carried out during the year was scarce (n = 91) and therefore less representative than that recorded in 2015 and calculated over a number of sampling days of nearly 50 % of the year. The 2015 THg wet deposition flux was 0.5 µg m⁻² yr⁻¹, which is lower than those recorded at most of the other southern GMOS sites.

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 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 Figs. 3–5.

In particular, Figs. 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 (Fig. 4) except at
Figure 4. Seasonal distribution of sampling-weighted fluxes (by 15-day 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 (+).

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

MHE, where the maximum THg wet deposition occurs during the winter. 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 NYA, ISK, and LON (Fig. 5). Therefore, the 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, fall, and winter seasons (Fig. 3). Rainfall was fairly equally distributed in all seasons except the winter season (Fig. 5). Thus, wet Hg loading was highest in spring, intermediate in winter and summer, and lowest in fall (Figs. 4 and 5). High levels of soluble species could in general be due to direct enhanced atmospheric oxidation of GEM to GOM, which oc-
curs in regions with high concentrations of oxidants, such as in polar regions during springtime (where atmospheric mercury depletion events – AMDEs – such as NYA occur). At PAL, Hg concentrations in rainfall increased through the winter, peaking in spring, and decreased through the summer and fall (Fig. 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 (Fig. 5). Thus, wet Hg loading was highest in summer and lowest in winter (Fig. 4). Similar behavior was observed at RAO (Fig. 4), whereas at MHE, wet Hg loading was highest in winter, when also the highest rainfall amounts have been recorded, and the lowest in fall (Figs. 4 and 5). At ISK, Hg concentrations in rainfall and wet Hg loading peaked in summer and decreased in fall and winter, respectively (Figs. 3 and 4), whereas rainfall was highest in fall and lowest in winter (Fig. 5). LON shows highest seasonal THg wet deposition in 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 pretreatment samples were grouped and analyzed season by season for a period of 3 to 5 years. Among the European sites, the highest THg wet deposition has been recorded at the remote RAO and PAL stations during the more photochemically active summer months, whereas lower amounts were found to be 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, 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 Pekkonen, 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).

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 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 MCH, MWA, and MAL were grouped by season and by site (Figs. 6–8). Seasonal variations of THg in precipitation were observed at the three Chinese sites (Fig. 6). The results obtained during the sampling period were similar to the seasonal variations of THg in precipitation in other Chinese regions, such as in the Wujiang River basin, Guizhou, China, but in contrast to the observations in North America (Landis et al., 2002), the Adirondacks (Choi et al., 2008), and the 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 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 the highest value in winter and lowest in summer (see Fig. 6), could be attributed in a first instance to lower rain amounts collected in winter (Fig. 8). The results obtained at the three Chinese sites show in fact that the THg concentrations varied with rainfall 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 the precipitation amount, with the highest value observed in summer and the lowest in winter (Fig. 8). THg wet deposition trend is comparable to that of the precipitation amount, with values of THg flux increasing from winter and peaking in summer (Fig. 8). Ruling out the winter season at MWA during which very few rainy samples have been collected (and is thus not representative for the present discussion) weighted THg concentra-
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 (+).

Figure 7. Seasonal distribution of sampling-weighted (by 15-day 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 (+).

Concentrations peaked in fall with lowest values in spring. Therefore, on average, wet Hg loading was highest in spring and lowest in summer. At MAL, the rainy samples show a fairly seasonal variability during all seasons, with the lowest rainfall in winter and the highest in summer (Fig. 8), while THg concentrations showed high values in winter and lowest in fall, and wet Hg loading was highest in summer, and lowest values were recorded in winter. 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 enhancing 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 at-
Figure 8. Seasonal mean values distribution of sampling-weighted (by 15-day reference) fluxes and rainfall at the three Chinese GMOS sites from 2011 to 2014.

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). 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 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) at three Chinese sites (urban, residential and near-remote sites) in the 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 (ng L\(^{-1}\)) significantly higher than those observed at the 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 in 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. 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). Hg is not incorporated into cold cloud precipitation as efficiently as it is into warm cloud precipitation (Landis et al., 2002). Other explanations for this observation have also 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

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 with regard to 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 lead to huge volumes of air being drawn out of the subcloud 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 Celestún 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 during which sufficient precipitation samples have been recorded (Figs. 9–11). Despite receiving unpolluted air off the Atlantic Ocean from northeasterly and southeasterly 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 (Fig. 10). The SIS high Hg deposition rates, comparable to other sites in the Northern Hemisphere, such as the Chinese sites (i.e., MCH) or European sites (i.e., ISK) that sometimes are also impacted by anthropogenic emissions, are driven in part by high rainfall events more intensely during summer and fall, and less during winter and spring periods. The high wet Hg deposition flux at this site suggests that other tropical areas 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 subtropical 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 the south of Florida and the Gulf of Mexico coastal areas, confirming that local and regional Hg emissions play only a minor role in wet Hg deposition (Guentzel et al., 2001; Sillman et al., 2013) and suggesting that the primary source of 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 Hg concentrations with the increasing of the rainfall depth). Weighted THg concentrations in rainfall (ng L$^{-1}$) peaked in winter and decreased through the spring and summer (Fig. 9). THg in wet deposition was highest in summer and lowest in spring and winter (Fig. 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. Different mechanisms leading to enhanced Hg concentrations in rain during the winter, including greater anthropogenic emissions, are probably associated with higher use...
of fossil fuels in power plants during the cold season. As reported in Sect. 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 United States and southern Florida where several coal power plants and waste 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 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 (Lindberg et al., 2007; Pirrone et al., 2008). Hg can then be re-emitted back to the atmosphere via gas exchange, and modeling studies suggest that re-emission from oceans is a major contributor to atmospheric concentrations of GEM, particularly in 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, AMS, CPT, CGR, and BAR, which recorded a representative number of samples over the 2012–2015 period. Figures 12 and 13 show the box plots related to rainfall, THg concentrations in precipitation, as well as wet deposition flux of Hg recorded, whereas Fig. 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 experienced a mild oceanic climate with monthly median air temperature ranging 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 displayed a seasonal variation of the precipitation amounts, with the highest values collected during the winter season (Fig. 14). On the contrary, the THg wet deposition flux patterns did not show a similar variation throughout the seasons as well as the THg concentrations in precipitation samples (Figs. 12 and 13). At CPT, the Hg concentrations in precipitation, Hg wet deposition fluxes as well as the precipitation amounts, followed the same trend during the rainy season (May–October), with a maximum in winter 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 experiences 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 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 the Cape Town metropolitan area before arriving at the stations. However, a more recent study on GEM concentrations and THg in precipitation carried out by Brunke et al. (2016) over a period of 7 years (2007–2013) 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 at 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 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 to 2015, at the CGR GAW station, located on the northwestern coast of Tasmania, Australia, highest value
in rainfall have been observed during winter and lowest in summer, whereas Hg concentrations peaked in summer and dropped to lowest values in winter (see Figs. 12 and 14). Indeed, an increase in precipitation volume results in a decrease in Hg concentrations in rain, probably due to the dilution of the washout loading (Prestbo and Gay, 2009). This means that any change in meteorological conditions, especially precipitation, complicates the interpretation of GMOS observations at different latitudes and might mask any trends due to 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 Figs. 12 and 14). Therefore, the seasonal THg wet deposition peaked in spring and decreased during the cold seasons (Fig. 13). It is necessary to point out, however, that in both 2014 and 2015 at BAR very few samples have been recorded in fall and summer (Tables S1–S4). This means that further measurements and studies are needed to draw any conclusion and improve our understanding of deposition processes and oxidation mech-
anisms 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).

5 Conclusions

Mercury deposition measurements are critical for constructing an accurate global Hg budget and modeling the benefits or consequences of changes in Hg emissions, for example, as prescribed by the Minamata Convention. The scarce availability of long-term wet Hg deposition data for calibration or validation of models could cause uncertainties in modeling applications to assess the influence of local emission sources. A synthesis of available Hg measurements in precipitation from selected GMOS ground-based sites is presented, including trends and seasonal cycles. Wet deposition samples were collected for approximately 5 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 has 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 those 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 the ground-based sites report, in particular, Hg deposition strongly influenced by the precipitation amounts. In the Northern Hemisphere, interannual 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 in the Southern Hemisphere the relationship between precipitation amount and deposition was not as evident as in the north. It is however necessary 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 starting 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 background and remote sites as well as local or regional sources, GMOS’s observation network gives important insights to modeling applications to evaluate future Hg trends and their fate and transport on a global scale. The results of THg wet deposition carried out in this study open the way for new avenues.
in future modeling studies as well as highlight the need for 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, and 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 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 a global scale, as would additional direct measurements of dry deposition across the GMOS network.

6 Data availability

Mercury data discussed in this paper are reported within the GMOS central database and are available upon request at http://sdi.iia.cnr.it/geoint/publicpage/GMOS.

The Supplement related to this article is available online at doi:10.5194/acp-17-2689-2017-supplement.

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

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References


Diéguez, M., García, P., and Sprovieri, F.: Atmospheric mercury fluxes in North Patagonia: first continuous records of the EMMA station (Global Mercury Observation System, Bartolome, Argentina), in: 12th International Conference on Mercury as a


