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Four years (2011–2015) of total gaseous mercury measurements from the Cape Verde Atmospheric Observatory

Katie A. Read¹, Luis M. Neves², Lucy J. Carpenter¹, Alastair C. Lewis¹, Zoe F. Fleming³, and John Kentisbeer⁴

¹National Centre for Atmospheric Science (NCAS), Department of Chemistry, University of York, York, YO10 5DD, UK
²Instituto Nacional de Meteorologia Geofísica (INMG), Delegação de São Vicente, Monte, CP15, Mindelo, Republic of Cape Verde
³National Centre for Atmospheric Science (NCAS), University of Leicester, Leicester, LE1 7RH, UK
⁴Centre for Ecology and Hydrology (CEH), Bush Estate, Penicuik, Midlothian, EH26 0QB, UK

Correspondence to: Katie A. Read (katie.read@york.ac.uk)

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Abstract. Mercury is a chemical with widespread anthropogenic emissions that is known to be highly toxic to humans, ecosystems and wildlife. Global anthropogenic emissions are around 20% higher than natural emissions and the amount of mercury released into the atmosphere has increased since the industrial revolution. In 2005 the European Union and the United States adopted measures to reduce mercury use, in part to offset the impacts of increasing emissions in industrialising countries. The changing regional emissions of mercury have impacts on a range of spatial scales. Here we report 4 years (December 2011–December 2015) of total gaseous mercury (TGM) measurements at the Cape Verde Observatory (CVO), a global WMO-GAW station located in the subtropical remote marine boundary layer. Observed total gaseous mercury concentrations were between 1.03 and 1.33 ng m⁻³ (10th, 90th percentiles), close to expectations based on previous interhemispheric gradient measurements. We observe a decreasing trend in TGM (−0.05 ± 0.04 ng m⁻³ yr⁻¹, −4.2 % ± 3.3 % yr⁻¹) over the 4 years consistent with the reported decrease of mercury concentrations in North Atlantic surface waters and reductions in anthropogenic emissions. The decrease was more visible in the summer (July–September) than in the winter (December–February), when measurements were impacted by air from the African continent and Sahara/Sahel regions. African air masses were also associated with the highest and most variable TGM concentrations. We suggest that the less pronounced downward trend inclination in African air may be attributed to poorly controlled anthropogenic sources such as artisanal and small-scale gold mining (ASGM) in West Africa.

1 Introduction

Mercury is present in the atmosphere in three main forms: gaseous elemental mercury Hg⁰, which is the most common form in the gas phase; oxidised mercury Hg II (GOM or RGM); and Hg bound to particulate matter (PBM). Total gaseous mercury (TGM) is the combined measurement of Hg⁰ (or gaseous elemental mercury (GEM)) + RGM, with Hg⁰ typically contributing around 90–99% of the total Hg or TGM.

Anthropogenic sources of mercury account for around 30% of the total amount and include emissions from coal burning, mining, cement production, oil refining, and waste incineration. One-third of the anthropogenic emissions are thought to come from deliberate biomass burning with Africa as the single largest continental source; therefore, in this region there could be an influence from Sahel African biomass burning during the months of November through to February (Roberts et al., 2009; De Simone et al., 2015). Hg⁰ reacts slowly with atmospheric oxidants with a global lifetime of around 6–8 months (Selin et al., 2007; Holmes et al., 2010), and so can be transported to remote regions. When oxidised to less volatile Hg II, it can be deposited either through wet deposition processes (precipitation scavenging) or by surface uptake (Sexauer Gustin et al., 2012; Schroeder and Munthe, 2010).
Hg\textsuperscript{0} also undergoes slow dry deposition through air–surface exchange with both terrestrial and aquatic surfaces (Zhang et al., 2009; Wang et al., 2016). Once deposited, transformation to highly toxic species such as the neurotoxic methylmercury allows bioaccumulation in food chains and poses a health risk to humans and a damaging effect to ecosystems (US EPA, 1997). Previously deposited mercury can also be reduced back to Hg\textsuperscript{0} through the natural weathering of mercury-containing rocks, geothermal activity, or from volcanic activity, and then re-emitted back to the atmosphere (Smith et al., 2008; Qureshi, 2011).

Reactions of Hg\textsuperscript{0} to Hg\textsuperscript{II} with the hydroxyl radical (OH) and ozone (O\textsubscript{3}) were historically accepted as the dominant photochemical oxidation mechanisms (Bergan and Rodhe, 2001; Lin et al., 2006; Seigneur et al., 2006; Selin et al., 2007; Ponpanprasert et al., 2008). Recent work has suggested that there may be significant other oxidants such as atomic halogens (Holmes et al., 2010; Wang et al., 2014) and more complex two-step oxidation schemes, which include further reactions with NO\textsubscript{2} and HO\textsubscript{2}; however, the kinetics are highly uncertain (Goodsite et al., 2004; Holmes et al., 2010). Heterogeneous oxidation in clouds may also contribute but is not experimentally proven (Ariya et al., 2009; Calvert and Lindberg, 2005).

Strode et al. (2007) estimated that 36\% of all mercury emissions in the Northern Hemisphere come from the ocean both through primary emission (ocean upwelling and mercury-containing rocks) and from re-emission of previously deposited mercury (as Hg\textsuperscript{II}), but this increases to 55\% as one moves into the Southern Hemisphere (Strode et al., 2007). The major anthropogenic source affecting the remote marine boundary layer is likely to be long-range transport of Hg\textsuperscript{0} from combustion (smelting, waste incineration, chemical plants) rather than from Hg\textsuperscript{II}, which is more likely to deposit regionally due to its relatively short lifetime of 4.8 h (Zhang et al., 2012). Other industrial sources for Hg include artisanal and small-scale gold mining (ASGM), which are known to occur in West Africa (Telmer and Velga, 2009; UNEP, 2013) and will likely regionally influence the measurements described here. For the 2013 UNEP global assessment, ASGM emission data were compiled from field and industry reports but with an uncertainty of ca. ±43\% due to the multitude and varying nature of ASGM sites. In recent years, global emissions from ASGM, and in particular the proportion of global emissions attributed to South America and sub-Saharan Africa, appear to be increasing; however, this assumption may be due to improved reporting (Muntean et al., 2014). The majority of global anthropogenic emissions of Hg to the atmosphere in 2010 are associated with ASGM (37\%), with one-third thought to be from sub-Saharan Africa (UNEP, 2013).

A community strategy developed by the EU was adopted in 2005 and listed 20 actions to reduce mercury emissions, cut mercury supply and demand, and protect people against exposure. This strategy had a strong focus on the need to take a global approach and included actions relating to multilateral negotiations for the conclusion of a legally binding convention on mercury (http://ec.europa.eu/environment/chemicals/mercury/strategy_en.htm). The UNEP Global Mercury partnership led by the US Environmental Protection Agency took a similar approach (http://web.unep.org/chemicalsandwaste/global-mercury-partnership) and these initiatives formed the basis of the Minamata Convention on Mercury, which was agreed in 2013 and is a global treaty to protect human health and the environment from the adverse effects of mercury (http://www.mercuryconvention.org/).

It has been a source of contradiction that in the Northern Hemisphere, while both measured atmospheric Hg concentrations and wet deposition fluxes have been decreasing since 1990 (Soerensen et al., 2012) and 1996–2013 (Slenczka et al., 2011; Weigelt et al., 2015); global Hg emissions during this period were calculated to be increasing (Pacyna et al., 2010; Streets et al., 2011). Very recently, however, Zhang et al. (2016), using a revised inventory and the global model GEOS-CHEM, have shown that global Hg emissions may also be decreasing. They suggest that a large discrepancy in the emissions data was from locally deposited mercury close to coal-fired utilities. It is thought that this source has declined more rapidly than was previously predicted due to shifts in mercury speciation from air pollution control technology targeted at SO\textsubscript{2} and NO\textsubscript{2} (Zhang et al., 2016). Flue gas desulphurisation (FGD) – which controls SO\textsubscript{2} emissions – washes out Hg\textsuperscript{II}, whilst selective catalytic reduction (SCR) to control NO\textsubscript{x} emissions also oxidises Hg\textsuperscript{0} to Hg\textsuperscript{II}. The effects of FGD, in addition to the recent phase-out of Hg from commercial products (UNEP Minamata Convention on Mercury) and lower global estimates from small-scale gold mining, serve to explain the globally decreasing atmospheric concentrations in the model. Zhang et al. (2016) also found that the larger emission decreases observed in North America and Europe generally offset the increases from other major polluted regions, e.g. from coal-fired utilities in East Asia (Pacyna et al., 2010; Pirrone et al., 2013).

Using data from ship cruises, Soerensen et al. (2012) observed a significant decreasing trend of atmospheric mercury concentrations over the North Atlantic of −0.046 ng m\textsuperscript{−3} yr\textsuperscript{−1} (−2.5 \% yr\textsuperscript{−1}), with smaller trends at more southern latitudes (Soerensen et al., 2012). They suggest that this decline is due to decreasing oceanic evasion driven by declining subsurface water Hg\textsuperscript{0} concentrations (−5.7 \% yr\textsuperscript{−1} since 1999; Mason et al., 2012).

Here we report 4 years (December 2011–December 2015) of TGM measurements at the Cape Verde Observatory (CVO), a clean marine background station located in the subtropical Atlantic. The measurements presented here are part of the EU Global Mercury Observation System (GMOS) network. The GMOS network of sites was established in 2011 with the aim of addressing known gaps in the spatial and temporal measurement of mercury, as well as improving knowl-
edge of Hg speciation. The data are being used to validate regional and global scale atmospheric Hg models in order to improve understanding of global Hg transport, deposition, and re-emission as well as providing a contribution to future international policy development and implementation (www.gmos.eu).

2 Experimental

The CVO was established in 2006 as a multilateral project between the UK, Germany and Republic of Cape Verde. Long-term atmospheric measurements include reactive trace gases including ozone, carbon monoxide, nitrogen oxides and volatile organic compounds (National Centre for Atmospheric Sciences (NCAS), University of York, UK), long-lived greenhouse gases (Max Planck Institute (MPI), Jena, Germany), and physical and chemical characterisation of aerosol (Leibniz Institute for Tropospheric Research (TROPOS), Leipzig, Germany). Details of the measurements and characteristics of the station can be found in Carpenter et al. (2010).

The CVO is positioned on the northeastern side of São Vicente (16.85°N, 24.87°W), one of 10 islands in the Cape Verde archipelago (Fig. 1). The island is of volcanic origin and the CVO is situated 50 m from the coastline. The climate is warm (mean annual air temperature is 24.0 °C ± 2.0 °C) and dry with extremely low annual rainfall (< 200 mm), which occurs mostly during the rainy season of July–November. The site receives air masses from the north-easterly trade winds for 95% of the time, which have travelled typically for five days over the ocean. Research flights carried out over the CVO in summer 2007 established that the boundary layer is well mixed (Read et al., 2008). There is no coastal shelf to the island at this location point, and the CVO conditions are considered to be representative of the North Atlantic open ocean boundary layer. Radiosonde and ceilometer data show there is no diurnal pattern evident in boundary layer heights, which suggests no systematic difference between daytime and night-time entrainment rates (Carpenter et al., 2010).

Air is sampled from the main laboratory glass manifold (10 m height of inlet, 2 in. diameter, residence time 4 s) and then through a 2 m length of 1/4 in. Teflon tubing and a particulate filter which is changed every 2 months. The entire inlet is heated. A Tekran 2537B analyser (Tekran Inc., Toronto, Canada) was used for the TGM measurements and is described in detail elsewhere (Steffen et al., 2014) and so only a brief summary is presented here. The analytical principle collects the TGM onto gold traps with subsequent thermal desorption and detection by atomic fluorescence spectroscopy (λ = 253.7 nm; Bloom and Fitzgerald, 1988). It is, however, likely that the measurement at this site is of GEM rather than TGM since RGM is lost very easily to any salt deposits in the inlet lines and filters. Samples of 5 L volume are obtained every 5 min (1 L min−1 flow rate) with a detection limit of around 0.1 ng m−3, using a dual-trap set-up. Concentrations in ng m−3 are reported at a standard pressure of 1013 hPa and a standard temperature of 273.14 K. Cali-
In calibrations are performed every 72 h using an internal mercury permeation source which injects a known amount of Hg(0) into mercury-free zero air (using a TEKRAN zero air filter, part no: 90-25360-00). The calibration consists of a zero and a span on each channel. The effective span was 19.08 ng m\(^{-3}\) for a sample volume of 5 L. The permeation rate was externally validated using manual injections of saturated mercury vapour taken from a Tekran 2505 mercury vapour calibration unit and after 5 years found to be within \(\sim 3.6\%\) of the instrumental set point. The detection limit of the instrument was 0.1 ng m\(^{-3}\).

Instruments to make trace gas and meteorological measurements are provided by the Atmospheric Measurement Facility (AMF), which is part of NCAS. Ozone measurements were made using a UV photometric analyser (Thermo Electron Corporation). The instrument had a detection limit of 0.05 ppb and a precision of < 1 ppbV. Carbon monoxide data presented here were measured using a vacuum UV fluorescence technique (Aerolaser 5001). It was sensitive to 1 ppbV and linear up to 100 ppm. The accuracy of the measurements was < 2 ppbV. In 2012 the Global Atmospheric Watch (GAW) audited these measurements and the report can be found at http://www.wmo.int/pages/prog/arep/gaw/documents/CVO_2012.pdf. Nitrogen oxide measurements were made using a low-detection, high-accuracy (sensitivity 0.3 and 0.35 pptV and accuracy of 5.5 and 5.9 % for NO and NO\(_2\) respectively) chemiluminescence analyser (AQD Inc.). The meteorological measurements presented here were made using a Campbell Scientific automatic weather station. For more information about the present NCAS instrumentation at the CVO refer to https://www.ncas.ac.uk/index.php/en/the-facility-amf/291-amf-main-category/cvao/cvao-amf-instruments/1557-cvao-amf-instruments.

Four years of data are presented here obtained between 5 December 2011 and 5 December 2015. In calculating annual statistics, we have used data from 1 December to 30 November. The data were quality-controlled using the central GMOS-Data Quality Management (G-DQM) system (Cinnirella et al., 2014; D’Amore et al., 2015). The G-DQM allows harmonisation of data across the network and is able to acquire and process data in near-real time, allowing immediate diagnosis of issues. It was developed using harmonised standard operating procedures, which had been established over many years by European and Canadian monitoring networks, together with recent literature (Brown et al., 2010; Gay et al., 2013; Steffen et al., 2012). An additional filter has been applied to the data presented here to exclude periods when the relative humidity was higher than 90 %, as the data were prone to increased uncertainties due to water condensing in the instrument. Instrument issues led to some significant data gaps; a lamp failure caused major data gaps between July and August 2012 and May and June 2014, whilst a pump failure caused downtime between October 2012 and January 2013.

3 Results and discussion

3.1 Statistics and seasonal cycles

The mean TGM concentration over 2011–2015 was 1.191 ± 0.128 ng m\(^{-3}\) and the 4-year time series is shown in Fig. 2. Sprovieri et al. (2016) showed that the CVO measurements (site referred to as CAL rather than CVO) fit well within the north–south gradient of TGM data. Other sites of reference, which receive background air similar in origin to CVO, include Mace Head, Ireland; Nieuw Nickerie, Suriname; and Cape Point, South Africa (Table 1). The remoteness of the CVO is reflected in the small variability of the TGM measurements, compared to other sites.

The data shown in Table 1 illustrate the dominating effect of emissions from the Northern Hemisphere compared to the Southern Hemisphere, with Mace Head (53° 20’ N, 9° 54’ W) TGM concentrations averaging 7–9 % higher than those ob-
Figure 3. Seasonal cycle of TGM at CVO. The bars represent the standard deviation of the monthly averages.

Table 1. Average TGM concentrations and standard deviation statistics from comparable sites in 2013 and 2014. Data from Sprovieri et al. (2016).

<table>
<thead>
<tr>
<th>Site (latitude, longitude)</th>
<th>Mean ± standard deviation for 2013 (ng m$^{-3}$)</th>
<th>Mean ± standard deviation for 2014 (ng m$^{-3}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mace Head, Ireland (53°20' N, 9°54' W)</td>
<td>1.46 ± 0.17</td>
<td>1.41 ± 0.14</td>
</tr>
<tr>
<td>Calhau, Rep. of Cape Verde (16°51' N, 24°52' W)</td>
<td>1.22 ± 0.14</td>
<td>1.20 ± 0.09</td>
</tr>
<tr>
<td>Nieuw Nickerie, Suriname (5°56' S, 56°59' W)</td>
<td>1.13 ± 0.42</td>
<td>1.28 ± 0.46</td>
</tr>
<tr>
<td>Cape Point, South Africa (33°56' S, 18°28' E)</td>
<td>1.03 ± 0.11</td>
<td>1.09 ± 0.12</td>
</tr>
</tbody>
</table>

served at Cape Point. The site at Nieuw Nickerie experiences 10% higher concentrations in the air arriving from the north compared to the south (Müller et al., 2012) and is additionally impacted by emissions from biomass burning and gold mining from South America (Sprovieri et al., 2010). Comparisons to ship-borne field campaigns in the Atlantic made between 1977 and 2001 (Sprovieri et al., 2010) show that the data from CVO are more comparable with southern Atlantic conditions than the northern Atlantic.

The CVO TGM monthly mean data show a weak seasonal cycle (1.289 ± 0.134 ng m$^{-3}$ December maximum, 1.130 ± 0.128 ng m$^{-3}$ June minimum, Fig. 3) with generally higher concentrations in winter and lower in summer. This cycle is similar, both in shape and magnitude, to that observed within subtropical maritime air masses at Mace Head, which is shallower than for other air masses (Weigelt et al., 2015) and generally not so defined as that of other remote sites in the Northern Hemisphere (Temme et al., 2007; Holmes et al., 2010). Selin et al. (2007) show that the mean seasonal amplitude of 12 northern mid-latitude sites between the maximum in January (winter) and minimum August (summer) is 0.19 ng m$^{-3}$, compared to the CVO amplitude of 0.14 ng m$^{-3}$ (December–June).

A smaller seasonal cycle may mean that O$_3$ plays a more dominant role in the oxidation of Hg$^0$ compared to other oxidants such as OH (Temme et al., 2007; Selin et al., 2007; Holmes et al., 2010). The equatorial nature of the CVO site means that solar irradiance and water vapour are high year-round (Carpenter et al., 2010; Whalley et al., 2010). This may lead to a less pronounced change in oxidation capacity between summer and winter when compared to sites at higher latitudes.

An influence of air masses from the Southern Hemisphere without any pronounced seasonal variation (Slemr et al., 2015) may be another reason for the smaller amplitude in the seasonal cycle at the CVO. However, air mass back trajectory analyses show that the CVO receives very little air representative of the Southern Hemisphere (~1.3% of all data, Fig. S1, Supplement). Further, the highest frequency of southerly air masses arriving at the CVO occurs during August and September, which would serve to increase the mercury seasonal cycle amplitude rather than reduce it.
Anthropogenic emissions of mercury affecting the Atlantic region include emissions from coal combustion, which tend to have maximum impact in February–March due to a dominance of air from continental regions such as North America. This is also observed in the seasonal distribution of anthropogenic combustion tracers such as carbon monoxide (Selin et al., 2007; Weigelt et al., 2015; Read et al., 2009). Ocean emissions of Hg\(^0\) from the reduction of Hg\(^{II}\) to Hg\(^0\), driven by increased biological production are at a maximum in June in the NH but December in the SH (Strode et al., 2007). The seasonal trend may also be affected by meteorological differences in seasonal circulation patterns and cycles in boundary layer heights, clouds, precipitation, and dry deposition characteristics (Dastoor and Larocque, 2004; Selin et al., 2007).

### 3.2 Four-year trends

The Theil–Sen function (Theil, 1950; Sen, 1968) was used to evaluate the 4-year dataset inclination based on monthly TGM medians by season. The results are shown in Fig. 4. In this function the slopes between all \(x, y\) pairs are calculated and the Theil–Sen estimate is the median of all these slopes. This analysis was performed using the Openair package in R (Carslaw and Ropkins, 2012). The advantage of using this function is that it gives accurate confidence intervals and is resistant to outliers. Statistics used for this plot can be found in Table S1 in the Supplement.

Over 4 years the data show a weak downward inclination \((-0.042 \pm 0.04\) ng m\(^{-3}\) yr\(^{-1}\), \(p < 0.01\) significance level). This decrease is more significant in the data collected during the Cape Verdean summer \((-0.079 \pm 0.054\) ng m\(^{-3}\) yr\(^{-1}\), \(p < 0.001\) in June–August) than in the winter \((-0.009 \pm 0.179\) ng m\(^{-3}\) yr\(^{-1}\), \(p < 0.1\) in December–February). Previous studies have shown a stronger decreasing trend in air that has been influenced by anthropogenic emissions, for which there is more ready detection of the impact of regulation (Selin et al., 2007; Cole et al., 2014; Weigelt et al., 2015; Zhang et al., 2016).

The seasonal trends calculated here imply that there are differences in the sources of mercury that affect the winter months compared to the summer months, potentially with a smaller decline in emissions over this winter period. This may be because CVO measures Hg coming from the same source region throughout the year but that the emission from that source has not declined as much in winter as it has in summer, e.g. from residential burning. Alternatively the difference could be explained by a difference in air mass between seasons bringing air from different sources that have experienced different trends in emissions over the years. We consider this latter scenario to be the more likely explanation, since air masses originating from continental Africa, which may be influenced by ASGM or biomass burning, frequently reach the CVO in winter (Carpenter et al., 2010), but are more rare in summer. A further alternative explanation for the difference in trends between seasons would be a change in global oxidant concentrations (such as OH) and that this effect had a seasonal dependence, but there is no evidence to support this from studies that estimate OH fields (Hartmann et al., 2013).

In order to understand better the drivers of the TGM behaviour, observations were classified according to the origin and pathways of air masses arriving at the CVO over a 10-day period using the UK Met Office NAME dispersion model in passive tracer mode (Ryall et al., 2001). The air mass classifications have been used previously for evaluating the source
Figure 5. (a) Boundary definition of the eight geographical regions: Coastal African, Europe, Sahara, Sahel, North America, Atlantic, South America, and South Africa. (b) Trajectory frequency maps for each of the seven air mass types using HYSPLIT trajectories and Openair: AAC – Atlantic and African coastal; AM – Atlantic marine; NAA – North American and Atlantic; NCA – North American and coastal African; EUR – European (with minimal African influence); AFR – African (with minimal European influence); and EUR/AFR – European and African.

Figure 6. Histograms of observed TGM classified by air mass for the full 2011–2015 dataset.
Table 2. Statistics for the individual air mass classified data.

<table>
<thead>
<tr>
<th>Mass</th>
<th>Mean ± 1σ standard deviation (25th–75th percentiles, number of points) ng m⁻³</th>
<th>% time the site receives air mass</th>
</tr>
</thead>
<tbody>
<tr>
<td>AM</td>
<td>1.14 ± 0.11 (1.08–1.22, 432)</td>
<td>6 %</td>
</tr>
<tr>
<td>AAC</td>
<td>1.15 ± 0.12 (1.08–1.24, 1633)</td>
<td>24 %</td>
</tr>
<tr>
<td>NAA</td>
<td>1.21 ± 0.08 (1.17–1.27, 449)</td>
<td>7 %</td>
</tr>
<tr>
<td>NCA</td>
<td>1.20 ± 0.10 (1.15–1.27, 975)</td>
<td>15 %</td>
</tr>
<tr>
<td>EUR</td>
<td>1.18 ± 0.13 (1.08–1.26, 1161)</td>
<td>17 %</td>
</tr>
<tr>
<td>AFR</td>
<td>1.23 ± 0.16 (1.14–1.29, 1448)</td>
<td>22 %</td>
</tr>
<tr>
<td>EUR/AFR</td>
<td>1.22 ± 0.11 (1.16–1.30, 586)</td>
<td>9 %</td>
</tr>
</tbody>
</table>

Figure 7. A correlation matrix separated by season to show the correlation between pairs of data, using the corPlot function in Openair (Carslaw and Ropkins, 2012). The ellipses are visual representations of a scatter plot. The colour scale highlights the strength of the correlation (red being the strongest and blue the weakest), and the number is the $r^2$ of the data. The data were daily averaged before correlating to remove any bias from diurnal variability. The order the variables appear is due to their similarity with one another, through hierarchical cluster analysis.

Biomass burning, of both anthropogenic and biogenic origins, is prevalent in Africa. In Northern Hemisphere Africa, burning occurs primarily in the Sahel, moving from the northern to the southern Sahel between November and February (Roberts et al., 2009). From Fig. 5b it would appear that there are few trajectories which originate from this region; however, an influence from biomass burning could be one explanation for the variable, and sometimes higher, mercury concentrations within AFR air masses. Previous studies have found a relationship between TGM and carbon monoxide during such episodes (Slemr et al., 2006; Brunke...
Figure 8. (a) Time series of TGM and carbon monoxide. The plot is normalised by dividing by the compounds’ mean value, and the grey dots indicate when the air was characterised as AFR. The shaded periods correspond to the 10-day back trajectories in (b). Clockwise from top left: 15 December 2013, 03:00 GMT; 17 December 2013, 18:00; 23 December 2013, 03:00; and 20 December 2013, 00:00.

et al., 2012). Figure 7 shows a correlation analysis using a matrix method between pairs of data at the CVO for AFR air, separated by season. O₃ and CO show a strong positive correlation particularly in spring, but also in summer and winter, consistent with their shared pollution sources and with CO being a precursor of O₃ over long transport times. Higher wind speeds tend to be associated with air masses that have travelled further (from continental regions) and therefore have undergone greater photochemical production, which might explain the positive correlation of O₃ with wind speed. In spring a strong positive correlation of TGM with O₃ suggests a shared anthropogenic source when these concentrations are at their seasonal high in the Northern Hemisphere. The lack of correlation of O₃ with CO in autumn, however, suggests a more localised source for CO, perhaps from biomass burning, but this is not reflected in a positive relationship with the TGM concentrations. Instead, TGM has a strong correlation with NOₓ (NO + NO₂), which suggests that the TGM concentrations are influenced by anthropogenic pollution sources within closer proximity, for example from pollution emitted from cities on the coast of West Africa. This may also explain the high variability in the TGM concentrations observed in the air classified as AAC, which has also travelled over the African coast; however, the concentrations in that air mass are lower due to longer time spent over the ocean. Air classified as AFR and EUR/AFR shows the highest levels of TGM and these may be better explained by an additional source further in-land. If sources of mercury from ASGM from West Africa have any impact on these measurements and trends, then this may be reflected in a weaker correlation with CO in autumn and winter. This is because we speculate that the activity might
be more commonly carried out in the dry season (November to April) when crops cannot be grown; however, there is little evidence to support this. There is no data for TGM in AFR in summer.

We next consider episodes when TGM concentrations were enhanced, to investigate the potential influence of biomass burning on the measurements. On two occasions TGM exceeded 1.7 ng m\(^{-3}\) (\(\sim 0.5 \text{ ng m}^{-3}\) higher than the mean levels detailed in Table 1). The first period was during December 2013; Fig. 8a shows normalised CO and TGM concentrations during this month. The shaded periods correspond to AFR trajectories, shown in detail in Fig. 8b. The figure shows that, for this period of relatively elevated concentrations in AFR classified air, there is no significant correlation with CO, and that the trajectories do not originate over the biomass-burning region of the Sahel. Thus, biomass burning appears not to be a contributor to the high TGM concentrations during this period.

A similar analysis was performed for the period 19 September 2015 until the 19 October 2015 and the corresponding plots and trajectories are shown in Fig. 9. In this case the period of elevated concentrations is shorter, with the episode lasting around a week. From the trajectories the air may have been influenced by air from the biomass region in Sahel Africa, which is at its most northern location in the month of October (Roberts et al., 2009). However, CO was not elevated during the period of peak [TGM] suggesting that biomass burning was not the source.

It has been previously established that West Africa is an important source region for ASGM activity (Telmer and Velga, 2009) but it is difficult to determine whether West Africa is a growing source of emissions since data have been limited and are subject to large uncertainty. It is likely, however, that the ASGM emissions are less regulated than anthropogenic emissions from coal combustion, ferrous/non-ferrous metal and cement production from Europe and the US (UNEP, 2013), and so is less likely to be decreasing in source strength.

A further analysis using the Theil–Sen function was performed to evaluate trends in the 4-year dataset within individual air masses, based on seasonal TGM medians (Fig. 10). The overall data (top left panel) show a decrease of \(-0.051 \text{ ng m}^{-3} \text{ yr}^{-1}\) (95% confidence interval of \(-0.1\) to \(-0.022 \text{ ng m}^{-3} \text{ yr}^{-1}\)), as shown in the green text. Over the 4-year period the concentration changes were (in \(\text{ng m}^{-3} \text{ yr}^{-1}\)) \(-0.046 \pm 0.037\), \(-0.021 \pm 0.043\), \(-0.052 \pm 0.060\), \(-0.061 \pm 0.044\), \(-0.058 \pm 0.039\), \(-0.051 \pm 0.043\), \(-0.077 \pm 0.034\), for AAC, AFR, AM, EUR, EUR/AFR, NAA, and NCA respectively (see Table S3 in the Supplement for the number of points used to derive
From an analysis of trajectories it was found that the AFR air was within the Sahel region outlined earlier for only around 1–2 % of the time and mostly in October when the burning is at its most northern point (Fig. 5b). In order to evaluate whether these air masses biased the decrease in AFR air, a sensitivity analysis was performed on the Theil–Sen analysis with the October data removed. The data flagged as southerly were also removed. The updated concentration change was \(-0.022 \pm 0.036 \text{ng m}^{-3} \text{yr}^{-1}\); thus, the filtering had essentially no effect on the AFR trend.

Weigelt et al. (2015) observed an annual decrease in TGM concentrations at Mace Head (53°20'N, 9°54'W; 10 m a.s.l.) of between \(-0.021\) and \(-0.023 \text{ng m}^{-3} \text{yr}^{-1}\) between 1996 and 2013, close to the upper confidence levels shown here (Weigelt et al., 2015). Within subtropical maritime classified air masses (air from south of \(28°\)N, west of \(10°\)W) the decrease was \(-0.016 \pm 0.002 \text{ng m}^{-3} \text{yr}^{-1}\). Further south at Cape Point, the trend for 17 years (1996–2013) was reported as \(-0.018 \text{ng m}^{-3} \text{yr}^{-1}\) (95 % significance interval of \(-0.035\) to \(-0.013 \text{ng m}^{-3} \text{yr}^{-1}\)). These concentration decreases are lower than the global calculation using the GEOS-CHEM model for the northern Atlantic of \(-0.04 \text{ng m}^{-3} \text{yr}^{-1}\) (Soerensen et al., 2012) but may not take into account the very recent decreases in emissions (Weigelt et al., 2015; Soerensen et al., 2012). The overall concentration decrease (2011–2015) observed here of \(-0.05 \text{ng m}^{-3} \text{yr}^{-1}\) is more similar to the GEOS-CHEM study.

4 Conclusions

We report a 4-year decreasing trend in total gaseous mercury (TGM) concentrations over the subtropical North Atlantic of \(-0.05 \pm 0.04 \text{ng m}^{-3} \text{yr}^{-1}\), a rate of decline that is in agreement with a GEOS-CHEM analysis for the northern Atlantic (Soerensen et al., 2012). A downward trend in concentration was observed in six out of seven different air mass types, all associated broadly with long-range transport of air from the US and Europe over the North Atlantic Ocean to the measurement location. The smallest and least significant downward trend (\(-0.02 \pm 0.03 \text{ng m}^{-3} \text{yr}^{-1}\)) was observed in air that was influenced by West Africa, where emissions are less understood and may well be static or possibly still increasing. The UNEP Global Mercury Assessment report in 2013 suggested that more work is needed to improve emissions estimates for West African sources, including field measurements around ASGM sites.

Data availability. The data presented here are freely available at the Centre for Environmental Data Analysis (CEDA) at http://catalogue.ceda.ac.uk/uuid/0ae5eb7ce3ad4885a7223dd7b69f4db6 (Read, 2010). Other levels of data are available within the GMOS central database upon request at http://sdi.iiia.cnr.it/geoint/publicpage/GMOS/gmos_historical.zul (GMOS Database, 2014).

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Figure 10. 4-year TGM trends using a Theil–Sen function based on seasonal TGM medians separated by air mass. The data for all air masses are shown in the top left panel with the rest of the panels showing the data separated by the seven air mass classifications. The green text shows the slope estimate and the 95 % confidence intervals are in brackets. In each case the solid red line shows the trend estimate and the dashed red lines show the 95 % confidence intervals for the trend based on resampling methods.
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Competing interests. The authors declare that they have no conflict of interest.

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