

Reply to Comments from: Anonymous (Referee #1)

This is a straightforward paper that provides details on trends in measured atmospheric mercury (Hg) at Cape Point and focuses on the changing trend in the last decade compared to the prior measurements. The paper concludes that the trends found at Cape Point are consistent with what is being found at other locations where long term measurements are being made, either of atmospheric gaseous mercury or wet deposition. I find no flaws in the paper and it is a straightforward analysis of the data which shows a clear trend. It adds to the dataset on this and supports an interesting global trend. My comment relates mostly to the paper and how it is written. Given the straightforward nature of the information I would combine the results and discussions together because as the paper is now written it is repetitive and too long. Also, there is no need for a conclusion as the results are straightforward. I would say that the paper could be reduced by 2 pages if the authors do this. I would make for a better read. The analytical methods are well known and the authors have previously published and it is clear the data is of high quality.

Reply: *We followed the recommendations of both reviewers to shorten the paper by merging the sections “Results” and “Discussion” in the revised manuscript.*

Reply to Comments from: Anonymous (Referee #2)

1. This paper analyzed the long-term trends of gaseous elemental mercury (GEM) concentrations at Cape Point between September 1995 - December 2005 and since March 2007 until June 2015. The paper focuses on the changing trend sign between 2004 and 2007. The authors concluded that the trend at Cape Point is qualitatively consistent with the trend changes in other observation sites and suggests a change in worldwide mercury emissions. Overall, the analysis is very straightforward and the paper is clearly written. However, I found the analysis might be a bit overly simplified and too qualitative. Some specific comments are given below. First, I can see that the measurement data at Cape Point is part of the GMOS and is an important dataset. But it is hard for me to see what the novelty of this paper is because the authors always stated that the results of this study are consistent with previous studies.

Reply: *The paper reports a reversal of trend of atmospheric mercury at Cape Point from a downward trend in the years 1995 – 2004 to an upward trend since March 2007 and tries to provide an explanation for it. Changing downward trends have been reported for atmospheric mercury concentrations and mercury wet deposition in the northern hemisphere, but Cape Point data is the only data set long enough to establish trends in the southern hemisphere.*

2. The comparison with emission inventory is quite weak. The authors tried to link the increase trend of GEM levels measured at Cape Point during 2007-2015 with the study of Zhang et al (2016), which indicated that the global mercury emissions had increased very slightly during 2000-2010, that is, by +5%. This is farfetched because the emissions given by Zhang et al. and the measurements are different years.

Reply: *We are aware of temporal mismatch between the trend since March 2007 until June 2015 and the emission estimates of Zhang et al. (2016) until 2010 and the problems with general attribution of trends in concentrations to trends in emissions. To ascribe the trend of mercury concentrations to the trends of emissions one needs a) emission estimates made with the same methodology over the years and b) emission estimates whose period matches the period of the monitoring. Emission estimates which fulfil both conditions are difficult to get. The more recent emission estimates tend to use improved methodology, i.e. are not consistent over the time. Because of methodical differences emission estimates by different authors are not consistent*

and cannot be compared. Emission inventories are also usually made with a delay of several years when the statistical input data are available. We are not experts on emissions inventories and have thus to rely on published work. To the best of our knowledge the estimate of worldwide emissions by Zhang et al. (2016) for 2010 is the most recent one which fulfil both conditions.

3. Line 30-33: The trend change is qualitatively consistent with the trend changes in GEM concentrations observed at Mace Head, Ireland, and in mercury wet deposition over North America suggesting a change in worldwide mercury emissions. This is a very important selling point for this paper. However, I wonder if this can be applied to worldwide emission. And more importantly, this should be supported by the data of Hg emissions or at least the drivers for Hg emission change. However, there is almost no analysis or discussions on the changes of Hg emissions in the paper.

Reply: *We have added a paragraph discussing the trends in worldwide mercury emission inventories. Lines 252-261 in the revised manuscript.*

4. Seasonal changes of biomass burning shall be discussed in more details and a more quantitative way if possible.

Reply: *We used CO measurements as a tracer for different seasonal trends of biomass burning but the result was inconclusive. As mentioned in the text, CO concentrations are highly variable and thus need longer periods to establish significant trends. Also mentioned in the text are large differences between estimates of mercury emissions from different authors.*

5. Line 259-262: This statement needs support of detail analyses.

Reply: *The hypothesis of biomass burning being the cause of different seasonal trends is discussed in the paragraph after line 262.*

6. Finally, the authors repeated some sentences in a few places of the paper. For example, Line 83-85 “According to Zhang et al. (2016) the worldwide anthropogenic emissions decreased from 2890 Mg yr⁻¹ in 1990 to 2160 Mg yr⁻¹ in 2000 and increased slightly 84 to 2280 Mg yr⁻¹ in 2010 “ was repeated in Line 250-252, and stated again in the conclusions. The downward trend during 1995-2004 was repeated for many times. The conclusions repeated the statements

in results and discussions. And there are many sentences in the abstract same as that in conclusions. Therefore, the authors shall short the paper by deleting the repetitive sentences.

Reply: *We followed the recommendations of both reviewers to shorten the paper by merging the sections “Results” and “Discussion”.*

1 **Trend of atmospheric mercury concentrations at Cape Point for 1995 – 2004**
2 **and since 2007**

3

4 Lynwill G. Martin¹, Casper Labuschagne¹, Ernst-Günther Brunke¹,- Andreas Weigelt^{2*}, Ralf Ebinghaus²,
5 Franz Slemr³

6

7 ¹South African Weather Service c/o CSIR, P.O.Box 320, Stellenbosch 7599, South Africa

8 ²Helmholtz-Zentrum Geesthacht (HZG), Institute of Coastal Research, Max-Planck-Strasse 1, D-21502
9 Geesthacht, Germany,

10 ³Max-Planck-Institute for Chemistry, Hahn-Meitner-Weg 1, D-55128 Mainz, Germany

11

12 *now at: Federal Maritime and Hydrographic Agency (BSH), [D-22589](#) Hamburg, Germany

13

14

15 Lynwill Martin: lynwill.martin@weathersa.co.za

16 Casper Labuschagne: casper.labuschagne@weathersa.co.za

17 Ernst-Günther Brunke: egbrunke@gmail.com

18 Andreas Weigelt: andreas.weigelt@bsh.de

19 Ralf Ebinghaus: ralf.ebinghaus@hzg.de

20 Franz Slemr: franz.slemr@mpic.de

21

22

23 **Abstract**

24

25 Long-term measurements of gaseous elemental mercury (GEM) concentrations at Cape Point, South
26 Africa, reveal a downward trend between September 1995 and December 2005 and an upward one
27 since March 2007 until June 2015 implying a change in trend sign between 2004 and 2007. The trend
28 change is qualitatively consistent with the trend changes in GEM concentrations observed at Mace
29 Head, Ireland, and in mercury wet deposition over North America suggesting a change in worldwide
30 mercury emissions.

31

32 Seasonally resolved trends suggest a modulation of the overall trend by regional processes. The trends
33 in absolute terms (downward in 1995 – 2004 and upward in 2007 – 2015) are the highest in austral
34 spring (SON) coinciding with the peak in emissions from biomass burning in South America and
35 southern Africa. The influence of trends in biomass burning is further supported by a biennial variation
36 in GEM concentration found here and an ENSO signature in GEM concentrations reported recently.

37

38

39 **Introduction**

40

41 Mercury and especially methyl mercury which bio-accumulates in the aquatic nutritional chain are
42 harmful to humans and animals (e.g. Mergler et al., 2007; Scheuhammer et al., 2007; Selin, 2009; and
43 references therein). Mercury, released into the environment by natural processes and by
44 anthropogenic activities, cycles between the atmosphere, water, and land reservoirs (Selin et al.,
45 2008). In the atmosphere, mercury occurs mostly as gaseous elemental mercury (GEM) which with an
46 atmospheric lifetime of 0.5 – 1 yr can be transported over large distances (Lindberg et al., 2007).
47 Mercury is thus a pollutant of global importance and as such on the priority list of several international
48 agreements and conventions dealing with environmental protection and human health, including the
49 United Nations Environment Program (UNEP) Minamata convention on mercury
50 (www.mercuryconvention.org).

51

52 Because of fast mixing processes in the atmosphere, monitoring of tropospheric mercury
53 concentrations and of its deposition will thus be the most straightforward way to verify the decrease
54 of mercury emissions expected from the implementation of the Minamata convention. Regular
55 monitoring of atmospheric mercury started in the mid-1990s with the establishment of mercury
56 monitoring networks in North America (Temme et al., 2007; Prestbo and Gay, 2009; Gay et al., 2013).
57 Until 2010 only a few long-term mercury observations have been reported from other regions of the
58 northern hemisphere and hardly any from the southern hemisphere (Sprovieri et al., 2010). The Global
59 Mercury Observation System (GMOS, www.gmos.eu) was established in 2010 to extend the mercury
60 monitoring network, especially in the southern hemisphere (Sprovieri et al., 2016).

61

62 Decreasing atmospheric mercury concentrations and wet mercury deposition have been reported for
63 most sites in the northern hemisphere (Temme et al., 2007; Prestbo and Gay, 2009; Ebinghaus et al.,
64 2011; Gay et al., 2013). At Cape Point, the only site in the southern hemisphere with a long-term record
65 exceeding a decade, decreasing mercury concentrations were also observed between 1996 and 2004
66 (Slemr et al., 2008). The worldwide decreasing trend has been at odds with increasing mercury
67 emissions in most inventories (Muntean et al., 2014, and references therein). Soerensen et al. (2012)
68 thought that decreasing mercury concentrations in sea water of the North Atlantic were responsible
69 for the decrease, at least in the northern hemisphere. The most recent inventories, however, attribute
70 the decrease of atmospheric mercury concentrations to a decrease in mercury emissions since 1990
71 (Zhang et al., 2016). The decrease in mercury emissions was attributed to the decrease of emissions
72 from commercial products, changing speciation of emission from coal-fired power plants, and to the
73 improved estimate of mercury emissions from artisanal mining. According to Zhang et al. (2016) the
74 worldwide anthropogenic emissions decreased from 2890 Mg yr⁻¹ in 1990 to 2160 Mg yr⁻¹ in 2000 and
75 increased slightly to 2280 Mg yr⁻¹ in 2010.

76

77 In the first approximation, the observed trends in atmospheric mercury should follow these changes.
78 There is indeed some recent evidence that the downward trend in the northern hemisphere is slowing
79 or even turning upwards (Weigelt et al., 2015; Weiss-Penzias et al., 2016). Here we report and analyse
80 the trends of atmospheric mercury concentrations at the GAW station Cape Point between 1995 and
81 2004 and since March 2007 until June 2015.

82

83 **Experimental**

84

85 The Cape Point site (CPT, 34° 21'S, 18° 29'E) is operated as one of the Global Atmospheric Watch (GAW)
86 baseline monitoring observatories of the World Meteorological Organization (WMO). The station is
87 located on the southern tip of the Cape Peninsula within the Cape Point National Park on top of a peak
88 230 m above sea level and about 60 km south from Cape Town. The station has been in operation since
89 the end of the 1970s and its current continuous measurement portfolio includes Hg, CO, O₃, CH₄, N₂O,
90 ²²²Rn, CO₂, several halocarbons, particles, and meteorological parameters. The station receives clean
91 marine air masses for most of the time. Occasional events with continental and polluted air can easily
92 be filtered out using a combination of CO and ²²²Rn measurements (Brunke et al., 2004). Based on the
93 ²²²Rn ≤ 250 mBq m⁻³ criterion about 35% of the data are classified annually as baseline.

94

95 Gaseous elemental mercury (GEM) was measured by a manual amalgamation technique (Slemr et al.,
96 2008) between September 1995 and December 2004 and by the automated Tekran 2537B instrument
97 (Tekran Inc., Toronto, Canada) since March 2007. Typically, ~ 13 measurements per month were made
98 using the manual technique, each covering 3 h sampling time. The manual technique was compared
99 with the Tekran technique in an international intercomparison (Ebinghaus et al., 1999) and provided
100 comparable results.

101

102 Since March 2007 GEM was measured using an automated dual channel, single amalgamation, cold
103 vapor atomic fluorescence analyzer (Tekran-Analyzer Model 2537 A or B, Tekran Inc., Toronto,
104 Canada). The instrument utilized two gold cartridges. While one is adsorbing mercury during a sampling
105 period, the other is being thermally desorbed using argon as a carrier gas. Mercury is detected using
106 cold vapor atomic fluorescence spectroscopy (CVAFS). The functions of the cartridges are then
107 interchanged, allowing continuous sampling of the incoming air stream. Operation and calibration of
108 the instruments follow established and standardized procedures of the GMOS (Global Mercury
109 Observation System, www.gmos.eu) project. The instrument was run with 15 min sampling frequency
110 while 30 min averages were used for the data analysis. All mercury concentrations reported here are
111 given in ng m⁻³ at 273.14 K and 1013 hPa.

112

113 The Mann-Kendal test for trend detection and an estimate of Sen's slope were made using the program
114 by Salmi et al. (2002).

115

116 **Results and discussion**

117

118 **1. Trend**

119

120 The upper panel of Figure 1 shows monthly average GEM concentrations calculated from all data since
121 March 2007 until June 2015 and in the lower panel monthly average GEM concentrations were
122 calculated from baseline data, i.e. GEM concentrations measured at ^{222}Rn concentration $\leq 250 \text{ mBq m}^{-3}$.
123 The slope of the least square fit of all data ($0.0222 \pm 0.0032 \text{ ng m}^{-3} \text{ yr}^{-1}$, $n=99$) is not significantly
124 different from the slope calculated from the baseline data only ($0.0219 \pm 0.0032 \text{ ng m}^{-3} \text{ yr}^{-1}$). Sen's
125 slope and trend significance for all ($0.0210 \text{ ng m}^{-3} \text{ yr}^{-1}$) and baseline ($0.0208 \text{ ng m}^{-3} \text{ yr}^{-1}$) data are listed
126 in Table 1. Sen's slopes tend to be somewhat lower than the slopes from the least square fits but they
127 are in agreement within their 95% uncertainty range. All trends are highly significant, i.e. at level \geq
128 99.9%. The results are essentially the same whether monthly median or monthly average
129 concentrations are used. This shows that the trend is robust and not influenced by occasional pollution
130 or depletion events.

131

132 ~~The trends were calculated for different seasons (austral fall – March, April, May; winter – June, July,~~
133 ~~August; spring – September, October, November; and summer – December, January, February) for the~~
134 ~~period since March 2007 until June 2015 from all and baseline data. These are listed in Table 1.~~
135 ~~Although the 95% uncertainty ranges of seasonal Sen's slopes overlap, the least square fit slopes for~~
136 ~~different seasons are statistically different at $> 99\%$ significance level. Irrespective of whether monthly~~
137 ~~averages or medians or least square fit or Sen's slope are used, a consistent picture emerges with~~
138 ~~upward trends where the slopes decrease in the following order: austral spring (SON) $>$ summer (DJF)~~
139 ~~$>$ winter (JJA) $>$ fall (MAM). (Brunke et al., 2010, 2012).~~

140

141 For comparison we also calculated the trends for the manually measured GEM concentrations during
142 the period September 1995 – December 2004 periods. These data have an annual coverage of only
143 about 300 hours per year, i.e. about 3% in contrast to the Tekran measurements since 2007 where the
144 coverage was nearly 100%. Baseline data were not filtered out from this data set because a) on average
145 only 13 measurements were available per month and b) ^{222}Rn was measured only since March 1999

146 and cannot thus be used as criterion for the whole period. In Table 2 we list the trends calculated from
147 the least square fit of the monthly medians. Monthly averages provide qualitatively the same trends
148 with lower significance, because of their larger sensitivity to extreme GEM concentrations. The trend
149 of all monthly medians of $-0.0176 \pm 0.0027 \text{ ng m}^{-3} \text{ year}^{-1}$ is somewhat higher than $-0.015 \pm 0.003 \text{ ng m}^{-3}$
150 year^{-1} (Slemr et al., 2008) calculated from the 1996 and 1999 – 2004 annual averages but within the
151 uncertainty of both calculations.

152

153 ~~Seasonal trends for the 1995 – 2004 period are all downward and their slopes are decreasing in the~~
154 ~~following order: austral fall > summer > winter > spring (note the negative sign of the slopes). The~~
155 ~~difference between fall and summer as well as between winter and spring is not significant. In absolute~~
156 ~~terms the slope during austral autumn (MAM) is the smallest and for spring (SON) is the highest for~~
157 ~~both the 1995 – 2004 and 2007 – 2015 data sets.~~

158

159 ~~2. Seasonal variation~~

160

161 ~~For analysis of seasonal variation we detrended the monthly averages using their least square fits. The~~
162 ~~detrended monthly averages were then averaged according to months. Figure 2a shows the seasonal~~
163 ~~variation of relative monthly averages with their respective standard error. No systematical seasonal~~
164 ~~variation is apparent in this plot. We noted, however, a two-year periodicity in the monthly averages.~~
165 ~~Figure 2b shows the monthly averages of the detrended monthly values for a 2-year period. Despite the~~
166 ~~somewhat higher standard errors of the monthly averages (number of averaged months for biennial~~
167 ~~variation being only half of those for the seasonal variation), the monthly averages vary between 0.95~~
168 ~~and 1.05 as do the monthly averages for the seasonal variation (Figure 1a). Taken collectively, however,~~
169 ~~the relative GEM concentrations during the second year are significantly (>99.9%) higher than those in~~
170 ~~the first year. This is a clear sign of a biennial variation of GEM concentrations at Cape Point.~~

171

172 ~~Discussion~~

173

174 ~~Tropospheric biennial oscillations (TBO) in tropospheric temperature, pressure, wind field, monsoon,~~
175 ~~etc. has been previously reported in the literature (e.g. Meehl, 1997, Meehl and Arblaster, 2001, 2002,~~
176 ~~Zheng and Liang, 2005). Meehl and Arblaster (2001) also report that TBO with roughly a 2 – 3 years~~

177 ~~period encompasses most ENSO years with their well known biennial tendency. Slemr et al. (2016)~~
178 ~~analysed mercury data from Cape Point in South Africa, Mace Head in Ireland, and from CARIBIC~~
179 ~~measurements in the upper troposphere and found an ENSO signature in all these data sets. Thus the~~
180 ~~finding of biennial variation of GEM concentrations at Cape Point is consistent with the ENSO influence.~~

181
182 All GEM concentrations show an upward trend of $0.0210 (0.0127 - 0.0284) \text{ ng m}^{-3} \text{ year}^{-1}$ between
183 March 2007 and June 2015. This trend is almost identical with $0.0208 (0.0141 - 0.0280) \text{ ng m}^{-3} \text{ year}^{-1}$
184 when only baseline (i.e. GEM concentrations at ^{222}Rn concentrations $\leq 250 \text{ mBq m}^{-3}$) are considered.
185 Occasional pollution and depletion events (Brunke et al., 2010; 2012) thus do not influence the trend
186 of the all data set. A decreasing trend of $-0.015 \pm 0.003 \text{ ng m}^{-3} \text{ yr}^{-1}$ was derived from annual medians
187 of all GEM concentrations at Cape Point measured manually during the years 1996 and 1999 – 2004
188 (Slemr et al., 2008). These data were obtained by a manual technique and have an annual coverage of
189 only about 300 hours per year, i.e. about 3% in contrast to the Tekran measurements since 2007 where
190 the coverage was nearly 100%. Here we derive a downward trend of $-0.0176 \pm 0.0027 \text{ ng m}^{-3} \text{ year}^{-1}$
191 from the least square fit of the monthly medians since September 1995 until December 2004. Despite
192 the different temporal coverage (because of incomplete data set, only the 1996 and 1999 – 2004
193 annual medians were used by Slemr et al. (2008)), both trends are in good agreement. Because of the
194 small data coverage and ^{222}Rn data available only since 1999 we were not able to filter out the baseline
195 data for the whole 1995 – 2004 period and to determine their trend separately.

196
197 The upward trend after March 2007 and the downward trend between 1995 and 2004 were measured
198 by different techniques: the former one with a Tekran instrument and the latter one using the manual
199 technique. For reasons outside of our control we could not operate both techniques side by side for a
200 reasonable length of time. Although the measurements by both techniques agreed well during an
201 international field intercomparison (Ebinghaus et al., 1999), we do not claim here that they are
202 comparable without an extended intercomparison of both techniques at Cape Point. Assuming internal
203 consistency of each of the data sets, it is however obvious that the decreasing trend between 1995
204 and 2004 turned to an increasing one since 2007 implying that the turning point was located between
205 2004 and 2007.

206
207 The changing trend reversal at Cape Point is the most pronounced but not the only signevidence that
208 the hemispheric trends in mercury concentrations are changing. An analysis of 1996 – 2013 data from

209 Mace Head, classified according to the geographical origin of the air masses, showed a) that the
210 downward trend of mercury concentration in air masses originating from over the Atlantic Ocean south
211 of 28°N is substantially lower than for all other classes originating north of 28°N and b) that all
212 downward trends for air masses originating from north of 28°N are decelerating (Weigelt et al., 2015).
213 The apparent inconsistency that no decelerating trend for air masses from south of 28°N was found
214 can be explained by the fact that the changes of a smaller trend are likely to be more difficult to detect.
215 Weiss-Penzias et al. (2016) recently reported that the wet mercury deposition was decreasing at 53%
216 of the sites in the U.S. and Canada and was increasing at none of the sites over the period 1997 – 2013.
217 Over the period 2008-2013, however, the mercury wet deposition was decreasing only at 6% of the
218 sites but was increasing at 30% of the sites. Thus the sign change of the trend at Cape Point somewhere
219 between 2004 and 2007 is just one more indication that trends in atmospheric mercury concentrations
220 are changing world-wide.

221

222 Trends in mercury concentrations and mercury wet deposition are most likely related to changes in
223 worldwide emission (Pacyna et al., 2016). Most anthropogenic emission inventories show nearly
224 constant or increasing anthropogenic emissions between 1990 and 2010 (Wilson et al., 2010; Streets
225 et al., 2011; Muntean et al., 2014) which is inconsistent with the worldwide decreasing trend in
226 atmospheric mercury concentrations and mercury wet deposition over this period. This inconsistency
227 has been explained by decreasing emissions from North Atlantic Ocean due to reduced mercury
228 concentrations in subsurface ocean water (Soerensen et al., 2012) and more recently by a substantial
229 reduction of mercury emissions from coal fired power plants and from commercial products between
230 1990 and 2000 (Zhang et al., 2016). The most recent inventory by Zhang et al. (2016) estimated that
231 the worldwide anthropogenic emissions decreased from 2890 Mg yr⁻¹ in 1990 to 2160 Mg yr⁻¹ in 2000
232 and increased slightly to 2280 Mg yr⁻¹ in 2010. To the best of our knowledge no more recent emission
233 estimates have been published so far (the emission estimates are always delayed by several years
234 needed for the collection of the underlying statistical data). Since the potential to reduce emissions
235 from the commercial products and from coal fired power plants was largely exhausted between 1990
236 and 2000 a further increase of worldwide mercury emissions between 2010 and 2015, mostly from
237 increasing coal burning and artisanal small scale gold mining can be expected.

238

239 Seasonally resolved trends may provide some information about the processes influencing the trends
240 at Cape Point. For the period 1995 – 2005 we find the smallest downward trend (-0.0132 ng m⁻³ year⁻¹)
241 in The trends were calculated for different seasons (austral fall (MAM) and the largest one (-0.0198 ng

242 $\text{m}^{-3}\text{-year}^{-1}$) in austral March, April, May; winter – June, July, August; spring (SON). In the – September,
243 October, November; and summer – December, January, February) for the period since March 2007 –
244 until June 2015 data the lowest upward trend is found for austral fall (MAM, around $0.010 \text{ ng m}^{-3}\text{-year}^{-1}$)
245 from all and the highest in baseline data. These are listed in Table 1. Although the 95% uncertainty
246 ranges of seasonal Sen's slopes overlap, the least square fit slopes for different seasons are statistically
247 different at > 99% significance level. Irrespective of whether monthly averages or medians or least
248 square fit or Sen's slope are used, a consistent picture emerges with upward trends where the slopes
249 decrease in the following order: austral spring (SON, $\sim 0.037 \text{ ng m}^{-3}\text{-year}^{-1}$.) > summer (DJF) > winter
250 (JJA) > fall (MAM). Seasonal trends for the 1995 – 2004 period shown in Table 2 are all downward and
251 their slopes are decreasing in the following order: austral fall > summer > winter > spring (note the
252 negative sign of the slopes). The difference between fall and summer as well as between winter and
253 spring is not significant. In absolute terms the slope during austral autumn (MAM) is the smallest and
254 for spring (SON) is the highest for both the 1995 – 2004 and 2007 – 2015 data sets.

255
256 The difference in seasonal GEM trends may could originate from the seasonal trends of GEM emissions
257 or from climatological shifts in regional transport patterns or both.

258
259 Hg emissions from coal fired power plants, the largest anthropogenic Hg source, tend to be nearly
260 constant over the year (Rotty, 1997). On the contrary, biomass burning is a highly seasonal
261 phenomenon with maximum emissions during August - September both in southern America and
262 southern Africa (Duncan et al., 2003; van der Werff et al., 2006). Taking into account a delay by ~ 3
263 months due to intrahemispherical air mixing time, the October - November coincide with the maximum
264 absolute seasonal trends: an upward one for 2007 – 2015 and a downward one for the 1995 – 2004
265 periods. Biomass burning emission inventories suggest a small decrease in CO emissions from Africa
266 and more pronounced one from South America between 1997 and 2004, but differences between
267 different inventories render it very uncertain (Granier et al., 2011). As the emission estimates by
268 Granier et al. (2011) end in 2010, no trend in emissions from biomass burning in 2007 – 2015 period
269 can be given. Nonetheless, the ambient Cape Point CO data has shown a measurable decrease during
270 2003 till 2014 (Tohir et al., 2015). We tried to calculate seasonal trends of baseline CO mixing ratios
271 for 1995 – 2004 and 2007 – June 2015 periods but none of the trends was significant. The 1995 – 2004
272 and 2007 – June 2015 periods are probably too short to reveal trends in CO data obscured by strong
273 seasonal and interannual variations. Nevertheless, the ENSO signature both in Hg and CO data from
274 Cape Point, Mace Head, and CARIBIC was found to be consistent, within large uncertainty margins,

275 with emissions from biomass burning (Slemr et al., 2016). In summary, seasonal variations of emissions
276 from biomass burning in southern Africa and America as well as ENSO signature are consistent with a
277 hypothesis of emissions from biomass burning as a major driving force behind the different seasonal
278 trends as seen in the Cape Point data.

279

280 Seasonal variation of mercury concentrations was also investigated. For this we detrended the monthly
281 averages using their least square fits. The detrended monthly averages were then averaged according
282 to months. Figure 2a shows the seasonal variation of relative monthly averages with their respective
283 standard error. No systematical seasonal variation is apparent in this plot. We noted, however, a two-
284 year periodicity in the monthly averages. Figure 2b shows the monthly averages of the detrended
285 monthly values for a 2 year period. Despite the somewhat higher standard errors of the monthly
286 averages (number of averaged months for biennial variation being only half of those for the seasonal
287 variation), the monthly averages vary between 0.95 and 1.05 as do the monthly averages for the
288 seasonal variation (Figure 1a). Taken collectively, however, the relative GEM concentrations during the
289 second year are significantly (>99.9%) higher than those in the first year. This is a clear sign of a biennial
290 variation of GEM concentrations at Cape Point.

291

292 Tropospheric biennial oscillations (TBO) in tropospheric temperature, pressure, wind field, monsoon,
293 etc. has been previously reported in the literature (e.g. Meehl, 1997, Meehl and Arblaster, 2001, 2002,
294 Zheng and Liang, 2005). Meehl and Arblaster (2001) also report that TBO with roughly a 2 – 3 years
295 period encompasses most ENSO years with their well-known biennial tendency. Slemr et al. (2016)
296 analysed mercury data from Cape Point in South Africa, Mace Head in Ireland, and from CARIBIC
297 measurements in the upper troposphere and found an ENSO signature in all these data sets. Thus the
298 finding of biennial variation of GEM concentrations at Cape Point is consistent with the ENSO influence.

299

300 **Conclusions**

301

302 We report here an increasingupward trend for mercury concentrations at Cape Point for the period
303 2007 – 2015. As mercury concentrations at Cape Point decreased over the period 19961995 – 2004 we
304 conclude that the trend must have changed sign between 2004 and 2007. Such a change is qualitatively
305 consistent with the trend changes observed in atmospheric mercury concentrations at Mace Head in

306 the Northern Hemisphere (Weigelt et al., 2015) and in mercury wet deposition at sites in North America
307 (Weiss-Penzias et al., 2016). Combining all this evidence it seems that the worldwide mercury
308 emissions are now increasing, after a decade or two of decreasing emissions. This finding is consistent
309 with the temporal development of mercury emissions in the most recent mercury inventory (Zhang et
310 al., 2016).

311

312 For both periods, 1995 – 2004 and 2007 – 2015, seasonally resolved trends were different in different
313 seasons. We believe that the observed trends of GEM concentrations at Cape Point result from the
314 trend of worldwide mercury emissions and are additionally modulated by regional influences. During
315 1995 – 2004 the highest downward trend was observed in austral spring (SON) and winter (JJA). For
316 the 2007 – 2015 period the highest upward trend was found in austral spring. Hg emissions from
317 biomass burning in South America and southern Africa both peak in August and September (Duncan et
318 al., 2003, van der Werff et al., 2006). Although the trend of these emissions is uncertain because of
319 differences between different emission inventories (Granier et al., 2013), it can produce different
320 trends in different seasons. Biennial variation of the GEM concentrations at Cape Point, reported here,
321 suggest that climatological changes of transport patterns can also play a role in seasonally different
322 trends. The detection of the ENSO signature in GEM concentrations at Cape Point (Slemr et al., 2016)
323 is consistent with the influence of both emissions from biomass burning and changing regional
324 transport patterns.

325

326

327 **Acknowledgment**

328

329 The GEM measurements made at Cape Point have been supported by the South African Weather
330 Service and have also received financial support from the Global Mercury Observing System (GMOS),
331 a European Community funded FP7 project (ENV.2010.4.1.3-2). We are grateful to Danie van der
332 Spuy for the general maintenance of the Tekran analyser at Cape Point.

333

334

335 **References**

336

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479 **Tables**

480

481 Table 1: Sen's slopes calculated from monthly GEM averages of all and baseline (i.e. $^{222}\text{Rn} \leq 250 \text{ mBq}$
 482 m^{-3}) data for March 2007 – June 2015.

Data	Sen's slope [$\text{ng m}^{-3} \text{ year}^{-1}$]	n	Significance [%]	Range at 95% signif. level [$\text{ng m}^{-3} \text{ year}^{-1}$]
All data	0.0210	99	>99.98	0.0127 – 0.0284
All Baseline	0.0208	97	>99.98	0.0141 – 0.0280
Fall (MAM, all data)	0.0089	27	95.99	-0.0009 – 0.0198
Fall (MAM, baseline)	0.0108	27	98.78	0.0018 – 0.0223
Winter (JJA, all data)	0.0153	25	99.29	0.0025 – 0.0294
Winter (JJA, baseline)	0.0152	25	98.68	0.0020 – 0.0287
Spring (SON, all data)	0.0375	24	99.74	0.0142 – 0.0556
Spring (SON, baseline)	0.0361	24	99.84	0.0160 – 0.0563
Summer (DJF, all data)	0.0287	23	99.87	0.0119 – 0.0440
Summer (DJF, baseline)	0.0269	21	99.79	0.0020 – 0.0287

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485 Table 2: Least square fit of monthly median of all GEM concentrations for September 1995 – December
 486 2004.

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Data	Slope [$\text{ng m}^{-3} \text{ year}^{-1}$]	n	Signif. level [%]
All data	-0.0176 ± 0.0027	94	>99.9
Fall (MAM)	-0.0132 ± 0.0052	23	>95
Winter (JJA)	-0.0189 ± 0.0049	23	>99.9
Spring (SON)	-0.0198 ± 0.0038	24	>99.9
Summer (DJF)	-0.0154 ± 0.0065	24	>95

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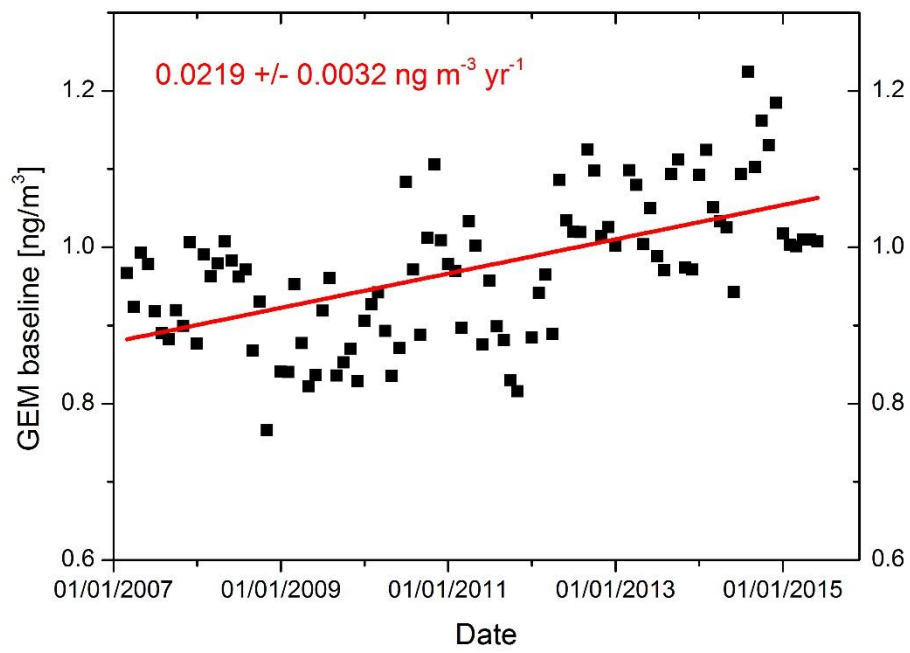
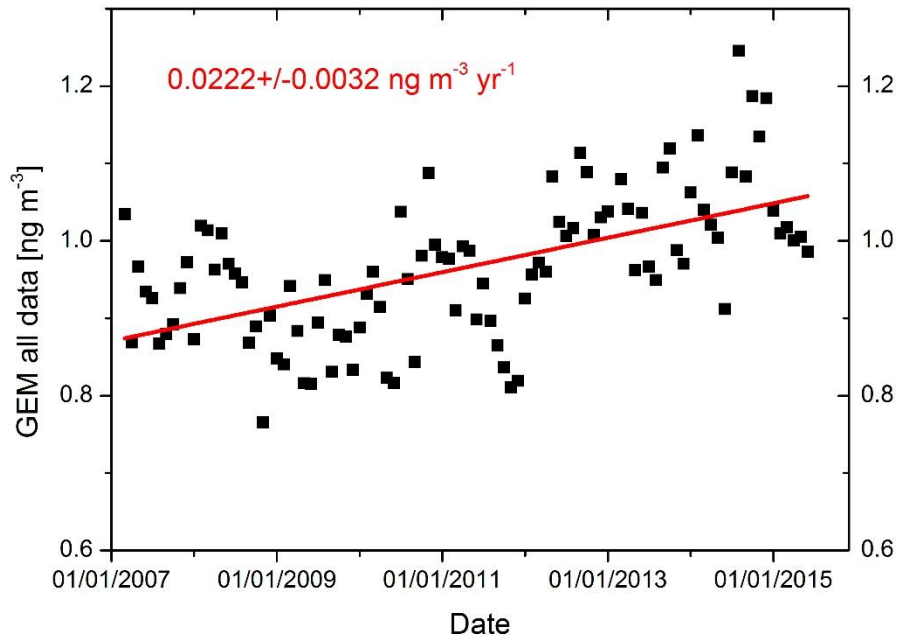
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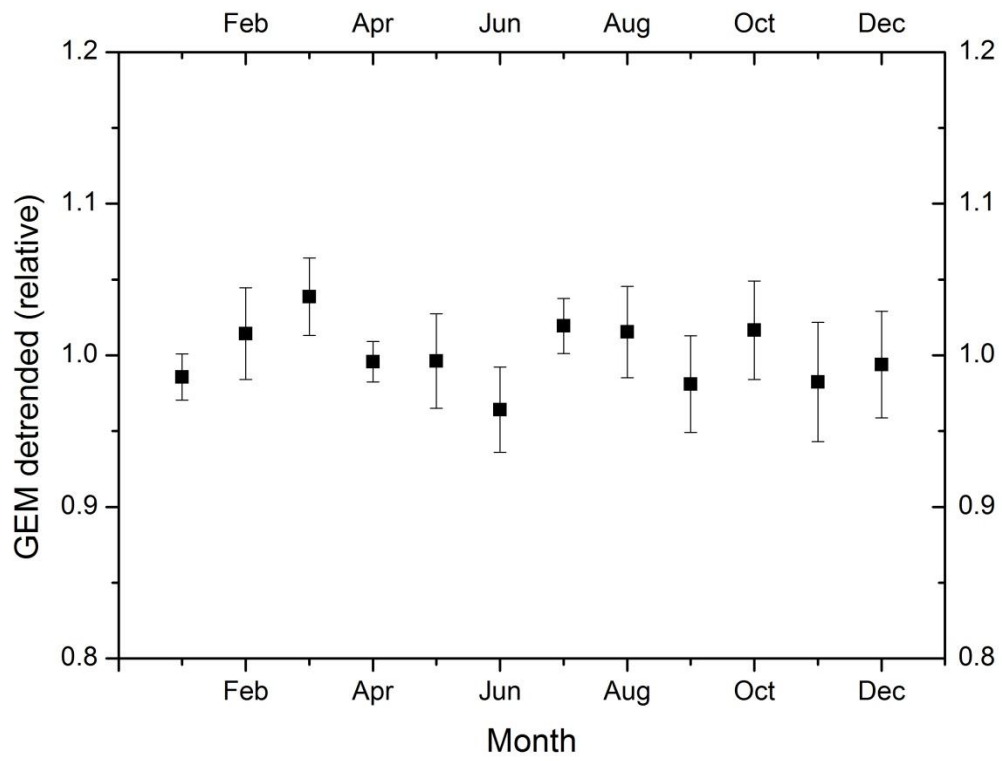
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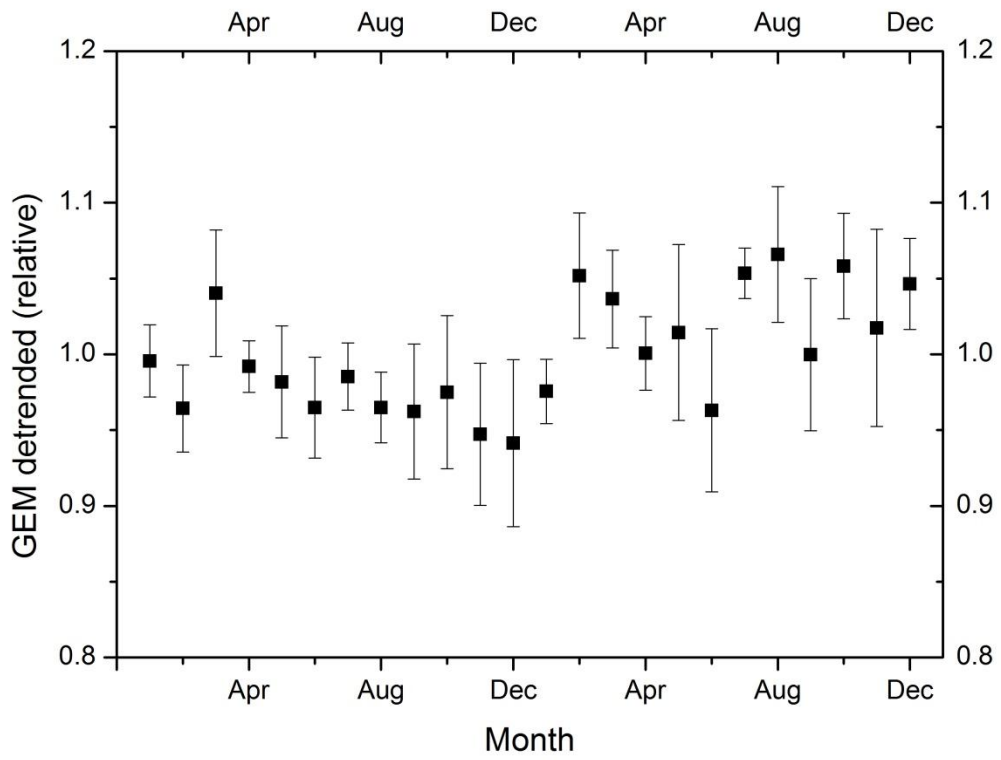


501 Figure 1: Monthly average GEM concentrations and their least square fit: upper panel all data, lower
502 panel baseline data (i.e. only GEM concentrations at ^{222}Rn concentrations $\leq 250 \text{ mBq m}^{-3}$).

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509 Figure 2: Seasonal (upper panel) and biennial (lower panel) variation of detrended monthly averages.

510 The error bars denote the standard error of the monthly average.

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