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Comment

Interactive comment on “Correlation slopes of GEM / CO, GEM / CO₂, and GEM / CH₄ and estimated mercury emissions in China, South Asia, Indochinese Peninsula, and Central Asia derived from observations in northwest and southwest China” by X. W. Fu et al.

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Received and published: 15 December 2014

Reponse to Anonymous Referee #2:

Firstly, we would like to acknowledge the anonymous reviewer for dictating the time to read our original manuscript and provide valuable suggestions. These suggestions are very helpful and constructive. We have made careful revision in the revised manuscript

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based on the reviewer's recommendations.

The authors present measurements of GEM concentrations made at three remote sites in China. Using concurrently measured CO, CO₂, and CH₄ mixing ratios they calculate GEM/CO, GEM/CO₂, and GEM/CH₄ concentration ratios for a number of pollution events. The authors compare these ratios with ratios reported by others for China and some other areas. The origin of the individual pollution events are attributed to four major source areas using backward trajectories. The transport from the source area to the measurement sites takes usually only a few days and thus chemical reactions of all measured species can be neglected. The measured GEM/CO, GEM/CO₂, and GEM/CH₄ concentration ratios then represent the emission ratios of the source area. Using the CO, CO₂, and CH₄ emission inventories for the source areas, the authors then calculate the GEM emissions for these areas. The authors find that the GEM emissions for all areas are substantially higher than the anthropogenic emissions from inventories.

The method is scientifically sound, the assumptions on which it is based were discussed by Jaffe et al. (2005) and Brunke et al. (2012) – see references in the paper. The paper is generally well organized and well written and should be published in a final version. However, the authors should address several weaknesses in the final version:

Major comment 1: The discussion is difficult to follow because the measured ratios in ng m⁻³ ppb⁻¹ or ng m⁻³ ppm⁻¹ are compared with emission ratios in t t⁻¹ (in the text though not in the Table 2). Using molar or mass ratios for both the measured and inventory ratios would help. Please use uniform units.

Reply: We agree that using different units in the manuscript made the manuscript somewhat difficult to read. Most of the previous studies used the units of pg m⁻³/ppb and pg m⁻³/ppm for correlation slopes of GEM/CO, GEM/CH₄ and GEM/CO₂. To be consistent with these previous studies while maintaining the readability of the paper,

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we have provided the conversions of $\text{pg m}^{-3}/\text{ppb}$ and $\text{pg m}^{-3}/\text{ppm}$ to g/g in the revised manuscript. The added information can be found in section 3.2, Table 1 and Table 2.

Major comment 2: One possible source of mismatch between observed ratios and ratios from inventories is ignored. The authors state that different circulation patterns (e.g. monsoon) transport the pollution to the measurement sites only during certain seasons. Because the GEM, CO, CO₂, and CH₄ emissions have all their specific seasonal variation (e.g. wetlands as a source of CH₄ peak in summer whereas CO and CO₂ emissions from residential heating peak in winter), their ratios will vary with season. Thus comparing annual emission ratios from inventories with observed emission ratios in a certain seasons can introduce a seasonal mismatch. Such mismatch could perhaps explain the overestimation of GEM emissions from GEM/CH₄ emission ratio and CH₄ annual emission inventory. The proper solution would be to calculate the regional emissions for the seasons for which the emission ratios were observed and using this as a basis for the estimation of GEM emissions instead of annual emissions. This would need a temporally resolved CO, CO₂, CH₄ emission inventories. This approach would of course provide only seasonal GEM emissions. If the temporally resolved CO, CO₂, and CH₄ emissions are not available the authors should at least mention this caveat.

Reply: This is an important point. Due to the different sources between GEM and CO, CH₄, and CO₂, the correlation slopes for GEM/CO, GEM/CH₄ and GEM/CO₂ vary temporally and spatially. We recognize that the correlation slopes cannot be observed uniformly throughout seasons and across the studied regions. This could be an inherent limitation for the correlation method. We introduced this artifact in many sections in the revised manuscript such as in line 402-407 on page 13 and line 539-552 on page 17. We believe that these revisions improve the discussion of the manuscript.

Major comment 3: The authors have data at disposal from which also CO/CO₂, CH₄/CO₂ and CH₄/CO ratios of the events could be calculated and compared for consistency with the official CO, CO₂, and CH₄ inventories. This could provide an insight

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into the seasonal mismatch mentioned above and perhaps also answer the question about the reliability of CH₄ emissions.

Reply: This is an excellent suggestion. We have added the correlation slopes for CO/CO₂, CH₄/CO₂ and CH₄/CO for the studied regions in the revised manuscript (line 342-355 on page 11, Table 1 and Table 2). The addition of the information helps better understand the discrepancy between observed correlation slopes of GEM/CO, GEM/CH₄ and GEM/CO₂ and estimated emission ratios. For example, we found the CO/CO₂ correlation slopes were consistent with anthropogenic emission inventories whereas observed CH₄/CO₂ and CH₄/CO were much lower than anthropogenic inventories, indicating anthropogenic inventories on CO and CO₂ may be more accurate than CH₄. This is in a good agreement with the assessment of uncertainties in CH₄, CO, and CO₂ inventories and also help explain why we got higher estimated emissions on the basis of GEM/CH₄ slopes. However, we did not go further to study the seasonal patterns because we have not obtained a sufficient number of correlation slopes in each season and correlation slopes in different season may be related to air masses for different regions. The temporal aspect of correlation slopes may also introduce uncertainties.

Major comment 4: The ranges of the GEM emissions calculated in this paper for different regions are rather narrow probably because the uncertainties of CO, CO₂, and CH₄ emissions were not considered. It should be mentioned that the inventories of CO₂, CO, and CH₄ in EDGARv3.0 are themselves uncertain by 10%, 50%, and 50%, respectively (Olivier et al., RIVM Bilthoven, RIVM Report #773301 001, NOP Report #410200051, 2001). Including the CO₂, CO and CH₄ emissions uncertainties of the used inventory would make many differences in the discussion insignificant.

Reply: We agree with this comment. The effect of uncertainties of CO, CO₂, and CH₄ emissions was discussed in the revised manuscript in line 527-536 on page 16.

Major comment 5: The calculated GEM emissions should be compared with the latest

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EDGARv4 gridded mercury inventory (Muntean et al., Sci. Total Environ. 494-495, 337-350, 2014).

Reply: The study by Muntean et al. (2014) is the most updated emission inventories. We have added this study into the manuscript.

I think that the authors in their final version of the paper should also address some minor problems listed below:

Minor points 1: Section 2, “Experimental”: The authors investigated pollution events lasting 8-24 hours (page 24994). What criteria were used to find out a “pollution event”?

Reply: These events were defined in line 204 and line 213-217 on page 7.

Minor points 2: Page 24987, line 13: There are 4 regions but only 3 GEM/CO2 ratios listed here.

Reply: There are some errors in the original manuscript, and we corrected them already (line 28-29 on page 2).

Minor points 3: Page 24987, line 27: Even different but collocated sources of GEM and CH4 will provide a GEM/CH4 emission ratio for a certain area. If chemical reactions during the transport can be neglected the measured GEM/CH4 emission ratio should correspond to the respective ratio of GEM and CH4 emissions from the area inventories. “Fewer common emission sources” thus should not be a problem but perhaps the different seasonal variations of GEM and CH4 emissions. The overestimation of CH4 emissions seems thus to be the primary cause for the overestimated GEM emissions.

Reply: This comment helped us better understand the correlation method. We deleted our previous discussions and talked the effect of overestimation of CH4 emissions on the discrepancy (line 458-468 on page 14).

Minor points 4: Page 24992: What are the standard conditions for mercury concentrations given in the paper? 1013 hPa and 273.16 K or something else? Please state

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explicitly.

Reply: We added the information in line 160-161 on page 6.

Minor points 5: Page 24993, line 1: “ensure” instead on “insure”

Reply: This word was changed.

Minor points 6: Page 24994, correlation analysis: It is not clear to me whether the correlations were made using the normal least-square fit or a bivariate correlation such as described by Cantrell (Atmos. Chem. Phys. 8, 5477, 2008). Both methods provide the same R or R² but different slopes, i.e. emission ratios. The problem with the least square fit is that the slope of e.g. GEM/CO is not equal to 1/slope CO/GEM which is physically incorrect. The normal least square fit assumes measurement uncertainty only in y whereas bivariate correlation needs uncertainties both in x and y. Bivariate correlation would be the correct method. If the authors used the normal least-square fit they should recalculate the slopes using the program provided with the paper by Cantrell (2008).

Reply: We thank the reviewer for point this out and would like to clarify it. The correlations were calculated based on SPSS using orthogonal least-square correlation. We also checked the slope and 1/slope for the observations. We found the difference in general fell in the range of 10% (may be due to the fact of significant correlations between GEM and CO, CH₄, and CO₂). Therefore, we did not change the method in the revised manuscript.

Minor points 7: Section 3.2: The authors may refer also to Hg/CO, Hg/CO₂, and Hg/CH₄ emission ratios published in a recent paper by Slemr et al. (Atmosphere 5, 342, 2014).

Reply: The papers by Slemr et al., (2009 and 2014) have been used in the revised manuscript.

Minor points 8: Section 3.3: Artisanal gold mining is not mentioned in this section. Ac-C10185

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According to the new EDGARv4 gridded mercury inventories (Muntean et al., Sci. Total Environ. 494-495, 337-350, 2014) this should be a very important source, especially for estimating the GEM emissions for South Asia, Indochinese Peninsula, Central Asia, and possibly even for China. This could be another explanation for the difference between the GEM emissions from emission ratios and GEM inventory emissions.

Reply: The contributions of artisanal gold mining is added in the revised manuscript in line 383-387 on page 12.

Minor points 9: Section 3.4: A map showing the four areas (China, South Asia, Indochinese Peninsula, Central Asia) of pollution origin for which the CO, CO₂, and CH₄ emissions from the inventories were calculated is necessary. Were the seasonally resolved CO, CO₂, and CH₄ emissions used or only annual emissions? CO/CO₂, CH₄/CO₂, and CH₄/CO emission ratios can be calculated from the correlations of available data. Are they consistent with the inventory ratios? If not what might be the reason and what does it mean for the calculated GEM emissions?

Reply: Figure 1 and Table 2 show the annual emissions of GEM, CO, CH₄, and CO₂. The CO/CO₂, CH₄/CO₂, and CH₄/CO emission ratios were calculated in Table 2 and compared with observed CO/CO₂, CH₄/CO₂, and CH₄/CO correlation slopes (line 342-355 on page 11). These discussions helped explain the uncertainties of CO, CH₄, and CO₂ emissions in the studies regions.

Minor points 10: Table 3: The estimated GEM emissions should also be compared with the emissions from the latest EDGARv4 gridded mercury emission inventory (Muntean et al., 2014).

Reply: The most updated inventory by Muntean et al. (2014) was added in Table 3.

Minor points 11: Figure 7: What is “warm season”?

Reply: We defined the “warm season” and “cold season” in the revised manuscript.

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Please also note the supplement to this comment:

<http://www.atmos-chem-phys-discuss.net/14/C10180/2014/acpd-14-C10180-2014-supplement.pdf>

Interactive comment on Atmos. Chem. Phys. Discuss., 14, 24985, 2014.

ACPD

14, C10180–C10187,
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