Can any currently identified chemistry rationalize the RAMIX results?

As we noted in the manuscript attempts to rationalize the discrepancies in the RAMIX dataset using any currently identified chemistry are extremely problematic. This is particularly the case for the reactions of GEM with NO$_3$ and O$_3$ that are suggested by Ambrose et al. (2013) as a potential mechanism for the high RGM concentrations measured by the DOGHS instrument. We have discussed these reactions in detail (Hynes et al., 2009) and we would suggest that more recent work confirms our conclusions. Any hypothesized oxidation chemistry for GEM is constrained by the fact that the HgO molecule is very weakly bound (Shepler and Peterson, 2003). Taking the binding energy of 17 kJ/mole calculated by Shepler and Peterson (2003) makes an abstraction reaction of Hg(0) with NO$_3$ endoergic by 195 kJ/mol.

$$\text{Hg} + \text{NO}_3 \rightarrow \text{HgO} + \text{NO}_2 \quad \Delta H^0 = 195 \text{ kJ/mol}$$

Recent work by Dibble et al. (2012) calculated a binding energy of 21 kJ/mol for an Hg-NO$_3$ adduct, suggesting that any such adduct would be too short lived to undergo further reaction.

Measurements of the reaction of Hg(0) with ozone have recently been reported by Rutter et al. (2012). They reported two experiments in which they monitored the decay of Hg(0) in a large excess of ozone.

Fig 1a) [O$_3$]: 1.2 x 10$^{13}$ molecules cm$^{-3}$, [Hg(0)]: 10.7 ng m$^{-3}$, 3.2 x 10$^7$ atoms cm$^{-3}$

Fig 1b) [O$_3$]: 6.5 x 10$^{12}$ molecules cm$^{-3}$, [Hg(0)]: 75.5 ng m$^{-3}$, 2.3 x 10$^8$ atoms cm$^{-3}$

If the reaction can be treated as a simple gas phase bimolecular reaction with reactants proceeding to products, i.e. any reverse reaction can be ignored, and one reactant is in large excess then the reaction proceeds under pseudo-first
order conditions. Since the ozone concentration is in large excess it remains essentially constant, hence the GEM concentration should decay exponentially and a plot of ln[Hg(0)] vs time should be linear and give a pseudo-first order decay rate, k’. A plot of k’ vs the excess reactant, ozone, should then give a bimolecular rate coefficient. Typically, if a reaction shows good pseudo-first order behavior the decaying reactant will show an exponential decay for 3 1/e times. This means that it will decay exponentially until the concentration is ~5% of the initial concentration. Deviations from this behavior are a clear indication that it is not possible to treat the reaction as a simple bimolecular process. In the Rutter et al. (2012) study the authors noted that “After the first 1.4 h a divergence from the model was observed in which the net oxidation was decreased with respect to the model predictions. Data collected after this point became inconsistent and tended to show a decreased net oxidation, or even reduction-like behavior, at a variety of experimental times. Such behavior was also observed after 1.4 h in the GEM-ozone experiments. No definitive explanation is available, but all data collected after 1.4 h are considered unreliable due to potential artifacts.” The model referred to here is simply pseudo-first order behavior for the GEM-ozone experiments. Rutter et al. (2012) obtained rate coefficients based on GEM concentrations that had decayed to only 98% and 96% of their original concentration and then acknowledge that the decays deviate from pseudo-first order behavior. This is a very clear indication that the ozone-GEM reaction cannot be treated as pseudo-first order. It seems clear that ozone and mercury react slowly on surfaces and we have observed such a reaction in our laboratory. A slow wall reaction of ozone with mercury could explain the slow decay rates initially observed by Rutter et al. (2012). There is no evidence that heterogeneously mediated oxidation of ozone occurs in the atmosphere but if it does a much more complicated formulism would be required to treat the process as described by Poschl et al. (2007).
Supplementary References:
Supplementary Figures

SI Figure 1: 22 hour sampling period from September 1st and 2nd. Comparison of the UM (red line) and UNR (green line) Tekrans with the UM 2P-LIF (black line) concentrations. The UM 2P-LIF signal was calibrated from the UM Tekran concentrations at the beginning of hour 13.
SI Figure 2: This is the same dataset as in Fig. 1 with an expanded concentration scale focusing on ambient measurements.
SI Figure 3: 22 hour sampling period from September 1st and 2nd. Comparison of the UM (black line) and UNR (green line) Tekrans with the UM 2P-LIF (black line) concentrations. The UM Tekran and UM 2P-LIF instruments were calibrated by the UNR Tekran concentration at the beginning of hour 35. In the case of the UM Tekran the full 22 hours of concentration measurements were multiplied by a constant factor that made the UNR and UM Tekran concentrations the same at the beginning of hour 35.
SI Figure 4: September 14 measurements 8-10.45 am. The background subtracted 2P-LIF signals from the ambient channel (black) and pyrolyzed channel (red) are shown. The gaps correspond to times when the laser was blocked to check power and background. The means and 1 standard deviation are shown.
SI Figure 5: September 14 measurements 8-10.45 am. The means of the ambient channel (black) and pyrolyzed channel (red) are shown. The error bars show both 2 standard errors (thicker line) and 2 standard deviations.
SI Figure 6: September 14 measurements 8-10.45 am. The RGM concentration obtained from the difference between the pyrolyzed and ambient Hg(0) concentrations is shown. The error bars show 2 standard errors.
SI Figure 7: September 14 measurements 12-2pm. The background subtracted 2P-LIF signals from the ambient channel (black) and pyrolyzed channel (red) are shown. The gaps correspond to times when the laser was blocked to check power and background. The means and 1 standard deviation are shown.
SI Figure 8: September 14 measurements 12-2pm. The means of the ambient channel (black) and pyrolyzed channel (red) are shown. The error bars show both 2 standard errors (thicker line) and 2 standard deviations.
SI Figure 9: September 14 measurements 12-2pm. The RGM concentration obtained from the difference between the pyrolyzed and ambient Hg(0) concentrations is shown. The error bars show 2 standard errors.
SI Figure 10: September 14 measurements: hour 8-19. The means of the 2P-LIF measurements are shown. The x-axis error bar shows the sample time. The y-error bar is 2SE of the mean. The measurements for the UNR speciation units spec 1 and spec 2 show the sum of gaseous and particle bound oxidized mercury. The values of Spec 2 using the correction suggested by Gustin et al. (2013) are also shown. The spike concentrations were reported by the UW manifold team. The DOHGS concentrations are 5 minute averages.
SI Figure 11. Fig. 15 (Manuscript) data plotted with an expanded concentration scale
SI Figure 12: September 13 measurements: hour 8-14. The means of the 2P-LIF measurements are shown. The x-axis error bar shows the sample time. The y-error bar is 2SE of the mean. The measurements for the UNR speciation units spec 1 and spec 2 show the sum of gaseous and particle bound oxidized mercury. The values of Spec 2 using the correction suggested by Gustin et al. (2013) are also shown. The DOHGS concentrations are hourly averages.
SI Fig 13. September 15th ambient measurements. Comparison of RGM as measured by the 2P-LIF pyrolysis instrument, UNR Spec 2, and our manual denuder measurement. The UW DOHGS and Spec1 systems were sampling from the RAMIX manifold with continuous HgBr$_2$ spiking during this period.
SI Figure 14: September 16th KCl manual denuder measurements. The temporal decomposition profiles (TDP) for the tandem denuder pair, D4 and D8 are shown.
SI Figure 15: September 16th KCl manual denuder measurements. The temporal decomposition profiles (TDP) for the tandem denuder pair, D5 and D6 are shown.
SI Figure 16: September 16th KCl manual denuder measurements. The temporal decomposition profiles (TDP) for the tandem denuder pair, D7 and D9 are shown.

147.61 pg/m³ (D7)
5.21 pg/m³ (D9)