Reply to Referee #2

We appreciate the very comprehensive review of the manuscript. Based on the line numbers that the referee refers to, it appears that it is the original submission was used as the basis for review, rather than the revised submission that it on the website.

To facilitate our response we have set the reviewers comments in red and our response in black type.

General Comments:

(1) In several areas of the paper (e.g., lines 63-64; 144-147) the authors incorrectly include particulate bound mercury (PBM) into their definition of total gaseous mercury (TGM). In the literature TGM is generally used to describe Hg0 + divalent reactive gaseous mercury (RGM). In fact most ambient instruments that quantify TGM (e.g., Tekran Instruments Corporation Model 2537) use an integrated Teflon filter to exclude all particulate matter, and most monitoring networks include an integrated Teflon filter at the inlet of their sampling line to minimize gas/particle interactions in the sampling line that has been shown to create problems in reliably quantifying ambient gaseous mercury species (see below). If the system is pyrolyzing all ambient mercury species to Hg0 for detection, then perhaps coin a new operationally defined term such as total atmospheric mercury (TAM) to avoid confusion.
We appreciate the reviewer picking this up. We have replaced most of the “total gaseous mercury (TGM)” with total mercury (TM) defined as the sum of gaseous elemental and oxidized mercury plus particulate bound mercury. In section 3.2 TGM is the appropriate term.

(2) The authors need to provide additional details on their sampling configuration and calibration procedures (QA/QC) in the methods section to provide readers and reviewers the basic information necessary to inform and judge the implementation of their 2P-LIF system.

We provided a very detailed description of the instrument as deployed at RAMIX including the sampling and calibration procedures in a recent paper in Atmospheric Measurement Techniques, cited as Bauer et al. (2014).


The first sentences of section 2.2 have been rewritten to emphasize this:

“Bauer et al. (2002, 2003, 2014) provide a description of the operating principles of the 2P-LIF instrument. Bauer et al. (2014) provide a detailed description of the instrument deployed at RAMIX including the sampling configurations, data
processing, calibration and linearity tests together with examples of experimental data.”

(3) Though not a problem with this paper per se, the design and implementation of the RAMIX study manifold system described by Finley et al. (2013) for this work is problematic and the results described in this paper must be viewed through this lens. The overarching issue was the decision to allow ambient particulate matter (PM1) into the manifold. When the stated objective was to evaluate the efficacy of different measurement methodologies to quantify Hg0 and RGM the logical first step would have been to evaluate these gaseous species first without the complication of spiked gases interacting with ambient particles in the manifold. The curious choice of constructing the manifold using highly porous PFA Teflon tubing also creates multiple potential problems (i) absorption/desorption of spiked gas concentrations following diffusion gradients, (ii) non-conducting material combined with high (187 LPM) flow rate and low relative humidity could lead to electrostatic collection of PM on the internal manifold walls providing additional surfaces for gas phase adsorption, and (iii) poor conduction of heat applied to the external surface of the tubing at such a high manifold flow rate (Finley et al. reported using eight thermocouples to measure external temperature down the length of the manifold but did not report any measuring internal surface or air temperature). Using a blower to maintain flow through the manifold also added a reported 15% uncertainty in the spike concentrations (Finley et al., 2013). All these issues lead to relatively low reported average transmission efficiencies for Hg0 (92%), HgBr2 (76%), and O3 (93%) even under controlled laboratory conditions (Finley et al.,
The range for Hg0 transmission in the manifold was reported to be 71-101% by Prestbo (2014). If the spiked mass of target gases into the manifold were not conserved through the system, then definitive evaluation of the analytical instruments sampling from the manifold were compromised. I agree with the authors that 15% probably underestimates the overall uncertainty for Hg0 (Lines 205-207) and that the system should be characterized as “a semi-quantitative delivery system” (Lines 210-211). If the manifold cannot quantitatively and reproducibly transmit a relatively inert gas like Hg0 there is very little chance of reproducibly transmitting HgBr2.

We have modified “Section 3.1 RAMIX Manifold” because it may give the impression that “The range for Hg0 transmission in the manifold was reported to be 71-101% by Prestbo (2014),” as inferred by the reviewer. This was not reported by Prestbo but rather is in the Finley et al. paper.

In regards to the issue of particulate matter the Finley et al. paper also contains the following:

“A Teflon coated cyclone inlet (URG 2000-30EA), also heated to 115 °C, was attached to the manifold. The inlet gave a particle size cut of approximately 1 μm at the expected manifold flow rate of 185–230 LPM (manufacturer communication).” This indicates that only the submicron fraction of ambient particulate matter was introduced into the manifold. However we appreciate the comments and note that in “Section 5.0 Future Mercury Intercomparisons:” we discuss the need for
sampling in an “unreactive” configuration that would eliminate any issues associated with particulate matter.

We write:

Line 750: Based on the RAMIX results it should consist of a period of direct ambient sampling and then manifold sampling in both reactive and unreactive configurations. For example an unreactive configuration would consist of Hg(0) and oxidized mercury in an N₂ diluent eliminating any possibility of manifold reactions and offering the possibility of obtaining a manifold blank response.

(4) This reviewer agrees with the authors that previous RAMIX study papers invoke and discuss mercury oxidation and reduction chemistry that is not supported in either the theoretical or applied literature. Controlling the physical adsorption of gaseous species onto manifold surfaces and aerosols is the logical first direction for which to find answers.

Specific Comments:

(1) Introduction:

a. Lines 49-50: Sprovieri et al., 2016 Atmos. Chem. Phys. Discuss., doi:10.5194/acp-2016-466 is a more contemporary discussion of global background Hg0 concentrations in the Northern and Southern hemispheres from the global GMOS network.
We had replaced Sprovieri et al. (2010) with Slemr et al., 2011, in the revised submission. We have added Sprovieri et al. (2016) as an additional reference.

b. Lines 63-64: PBM is not part of TGM.
Modified as described above

(2) Experimental:

As indicated above, the authors need to supplement the QA/QC information.
See reply to point 2) in general comments.

i. Lines 122-126: The authors discuss calibrating the 2P-LIF system using a Tekran 2537B as a secondary transfer standard, and that unlike their previous field work the unit was turned off for one week for transportation to the study site. This discussion implies to the reader that there was some kind of additional uncertainty in the calibration due to the Tekran 2537B unit being powered down. As long as the authors powered up the unit and had argon purge gas flowing for 2-3 days prior to use the permeation system should have re-equilibrated and had no impact on their 2P-LIF system calibration. I suggest the authors clarify the circumstances of the Tekran 2537B operation status. Manual standard addition injections from the Tekran Model 2505 primary calibration source (that the authors indicated they had on site – Line 180) should have been conducted to verify the stability/accuracy of
the 2537B instrument perm tube system prior to initiation of the experiment. The authors should report results for any QA/QC injections.

In the initial period of operation we saw instability in the response factors from calibrations and the overall response between calibrations. This is discussed below. However we cannot clearly identify these problems as being due to the fact that the Tekran 2537 was shipped and went without power for one week and we have removed that statement from section 2.2. Section 3.2 now includes the statement: “Problems with instability in the UM Tekran may have been associated with the use of an external pump to supplement the internal Tekran pump, or with the fact that the UM instrument had been powered down for almost one week and relocated to a site at a significantly different ambient pressure.”

The Tekran 2537 was calibrated using the internal permeation source. The Tekran Model 2505 primary calibration source was at a different location at the University of Nevada, Reno. This was the location of the LIF system that was used for denuder analysis. In retrospect it is evident that we should have performed additional QA/QC procedures but we would note that at this period, the start of the intercomparison we were primarily focused on assembling two completely different LIF systems, the 2P-LIF system at the RAMIX site and the LIF system for denuder analysis that was located in a laboratory at UNR. We have modified the text in “3.2 UM Tekran Performance “ to discuss this as detailed below.

ii. Lines 122-126: Clarify if the 25’ sampling line was heated and shielded from the sun.
The sampling line was not heated and was not shielded from the sun and this is now noted in the manuscript.

iii. Lines 130-131: If the Tekran 2537B was not able to pull 2 LPM with the additional load of a 25’ sampling line it is not clear why the authors simply did not reduce the flow rate set point to 1.5 LPM (the manufacturer recommended flow rate). Adding a supplemental pump to the instrument exhaust and increasing the instruments internal vacuum can impact the permeation tube system performance creating uncertainty in the instruments reported values. The additional vacuum may also have played a role in the permeation tube “malfunction” described by the authors in Lines 230-231. Was the Tekran 2537B checked with manual standard addition injections from the Tekran Model 2505 primary calibration source in the external pump configuration?

There is an error in the manuscript here and in Bauer et al. (2014) and this is now corrected. The Tekran was indeed sampling at 1.5 LPM but it was necessary to add a supplemental pump to be able to sample at this flow. We were unaware that this might present potential problems. As noted above the Tekran Model 2505 primary calibration source was at a different location at the University of Nevada, Reno and it was not used to check the Tekran 2537 calibration.

iv. Lines 133-134: The authors indicate that the 2P-LIF system cannot detect RGM, but expressed concern about “deposition of RGM on the sampling lines followed by
heterogeneous reduction to GEM”. Were any actions taken to filter out RGM from the sampling line (at the manifold port) while allowing the GEM to pass to the instrument like incorporation of a soda and lime trap? Allowing HgBr2 into a Tekran 2537B can have long term contamination effects on the internal components (filter packs, tubing, connectors, and valves).

No actions were taken to filter out RGM from the sample line. This is now explicitly stated in the manuscript.

v. Lines 148-150: Was a second pump used to maintain flow through the sampling line not being actively sampled by the 2P-LIF system during the TOM difference experiment? Otherwise air in the sampling line void volume not being actively sampled would stagnate and not represent the correct temporal sample duration.

Both lines were continuously sampled at 10 L/min and the flow to the fluorescence cell was switched between the lines. This is now explicitly stated in the manuscript.

vi. Lines 152-169: How often were the KCl-coated manual annular denuder cleaned and recoated/conditioned?

The denuders were cleaned and recoated prior to the RAMIX deployment. Prior to sampling, the denuders were cleaned by heating to 500 °C and then bagged and taken to the sampling site. This is now explicitly stated in the text.
vii. Lines 231-234: The authors discuss comparison of the UM and UNR Tekran 2537 instruments and bring up the point again about the power down of their 2537B instrument. The authors should clarify how long the system was allowed to stabilize prior to the first perm tube calibration, and how often thereafter it was recalibrated. Was an independent QA auditor part of the study plan to ensure traceability across research group instruments? See below

b. Line 146: Again PBM is not part of TGM.

Modified as described above


Corrected
d. Line 165: Not heating the KCl-coated annular denuders during sampling can be problematic since the hydrophilic KCl coating will tend to absorb water vapor and can (i) interfere with RGM collection, and (ii) provide surfaces for heterogeneous reactions.

We would note that the humidity in Reno was low during the RAMIX experiment.

(3) Results:
a. Lines 231-236: In the absence of an independent auditor or standard addition injections to validate the respective instruments perm tube emission rates, it is not
possible to definitively establish the reason for the observed differences between the UM and UNR instruments. Based on the described behavior it could be related to contamination of one or both of the instruments with HgBr2 or an unstable permeation tube system. It would be useful for the authors to discuss the observed behavior as a function of the timing of HgBr2 spiking.

b. Lines 300-301: The authors point out in this discussion that the UM 2537B was systematically reporting a higher Hg0 value than the UNR instrument. If the authors believe the divergence between 2537 instruments was due to the UM instrument being turned off for shipment, then the UM instrument would be reporting lower values. This would be due to the fact that the amount of mercury being emitted by the unequilibrated perm tube system would be higher than expected during the calibration since excess Hg0 accumulated on the walls of the perm tube oven would be slowly desorbing – resulting in lower reported ambient concentrations.

See below

c. Line 399: The authors are discussing Fig. 7 in this discussion and then state “In addition, it is clear that the DOHGS system show very different temporal profiles of TOM.” I suspect the authors should reference Fig. 9 here since I do not see the DOHGS concentrations presented in Fig. 7.

This has been clarified.

d. Line 413: I do not see the DOHGS concentrations presented in SI Figure 6.

The DOHGS concentrations are shown in Fig. 10. The text has been modified.
e. Lines 422-423: I do not see the DOHGS concentrations presented in SI Figures 7-9 as referenced in the text.

As noted above, the DOHGS concentrations are shown in Fig. 10. The text has been modified.

f. Lines 447-455: It is unclear what the authors take home message for Section 3.5.3 discussion. It is also unclear why the Spec2 data are shown in SI Fig. 13 since it was sampling off the manifold while all the other measurements are from the trailer roof.

SI Fig. 13 shows both Spec 1 and Spec 2 data. Spec 2 was sampling ambient air but Spec 1 was sampling from the manifold which was spiked with HgBr₂. The Spec 1 data should not have been included and it has been removed.

g. Lines 489-530: Landis et al. 2002 documents the quantitative transport of HgCl₂ through the manual denuder elutriator/impactor inlet when properly heated.

See response below to i)

h. Lines 490-492: The authors implementation of the manual denuders method described by Landis et al., 2002 deviated in two ways (i) not using the suggested
elutriator/impactor inlet to remove large aerosols which may be retained by the denuder causing positive artifacts (the potential contamination of their RGM denuder sample by PBM is later discussed by the authors), and (ii) not heating the denuder system.

Now reads “As we describe above, our use of manual denuders was similar to that described by Landis et al. (2002) with the exception that we did not incorporate the integrated elutriator/acceleration jet and impactor/coupler on the denuder inlet and the denuders were not heated.”

i) Lines 493-494: Feng et al., 2004 reference cited by the authors to imply potential loss of RGM by the inlet elutriator/impactor inlet of the manual denuder system does not support their statement. The Feng et al. paper does not use or even mention this system or RGM loss in their particulate mercury methods paper. They used a method described by Lu et al., 1998 that does not use an inlet.

This reference has been corrected to Feng et al. (2003).


The abstract includes the following:

“It is shown that, under the experimental conditions examined, KCl-coated annular quartz denuders designed for ambient reactive gaseous mercury (RGM) collection could quantitatively collect HgCl2. It is also demonstrated that the impactors used to remove coarse airborne particulate matter could lead to a loss of up to one
third of the HgCl$_2$ in the gas stream.”

The authors cite Landis et al. (2002) but do not state that the impactors and denuders were heated.

Our text has been modified as follows:

“Landis et al. (2002) suggest that HgCl$_2$ is quantitatively transported through the manual denuder elutriator/impactor inlet when properly heated. In later work Feng et al. (2003) reported that such impactors could reduce the efficiency of RGM collection although in that work there is no reference to the temperature of the impactor. In this work no type of particle filtering was used on the inlets.”

j. Lines 517-522: Manual versus automated denuder and denuder in series experiments described here were previously conducted and presented in Landis et al., 2002. No significance difference between manual and automated systems, and no significant breakthrough from the first denuder. This previous work should be cited.

The Feng et al. (2003) also included laboratory experiments on manual denuders in series. Both works are now cited.

k. Lines 579-582: The critical review of the experimental design here is warranted and should go further to include recommendations to (i) improve the manifold design and sampling port configurations as previously discussed, (ii) include an independent auditor, and (iii) removal of PM from the gas phase experiments.
We have not tried to provide a detailed recommendation on the future manifold design. The reference to the GASIE campaign that included oversight by an independent panel consisting of three atmospheric scientists none of whom were involved in SO$_2$ research and our suggestion that:

“We would suggest that a future mercury intercomparison should be blind with independent oversight.”

Imply the inclusion of independent auditors.

As we have noted above the RAMIX manifold did incorporate a Teflon coated cyclone inlet.

i. Lines 122-126: The authors discuss calibrating the 2P-LIF system using a Tekran 2537B as a secondary transfer standard, ........

We have provided a single response to the comments related to the operation of the UM Tekran

vii. Lines 231-234: The authors discuss comparison of the UM and UNR Tekran 2537 instruments ................

(3) Results:

a. Lines 231-236: In the absence of an independent auditor or standard addition injections .................

b. Lines 300-301: The authors point out in this discussion that the UM 2537B was systematically reporting a higher Hg0 value than the UNR instrument. .................
As we have noted above the start of the intercomparison we were primarily focused on assembling two completely different LIF systems, the 2P-LIF system at the RAMIX site and the LIF system for denuder analysis that was located in a laboratory at UNR. We should also note that we are not Tekran experts and the UM Tekran was not part of the primary intercomparison, in retrospect it is evident that we should have performed additional QA/QC procedures on the Tekran. We believe the key point here is that it was not reasonable for Gustin et al. to use this data to conclude that by some unidentified mechanism oxidized mercury was being converted to Hg(0) and that the UM Tekran was measuring total mercury. It is also noteworthy that the offsets shown in Fig. 1 occurred prior to the start of the manifold spikes of HgBr₂ and cannot be associated with the elevated levels of HgBr₂ that were introduced into the manifold on Sept. 5th. We have now modified section 2.2 and added a section in Supplementary Information giving more details about the calibration and response instabilities in the UM Tekran.

We have modified Section 2.2 as follows:

**“3.2 UM Tekran Performance**

In evaluating the first week of the UM RAMIX measurements it became clear that there was some non-linearity in the relative responses of the 2P-LIF and UM Tekran systems and that better agreement was obtained by referencing the Hg(0) concentration to the UNR Tekran. Gustin et al., (2013) concluded that the UNR Tekran, based on the inlet configuration, only measured Hg(0) and they suggested
that the UM system, due to the long sampling line, was measuring total gaseous mercury (TGM). We compared the manifold Hg(0) readings from the UM and UNR Tekrans over the first 260 hours in which we took measurements. The absolute concentration difference relative to the UNR instrument is shown in Figure 1. Hour zero corresponds to 9 am on August 26th when we started measurements and hour 260 corresponds to midnight on September 5th. Over the first 24 hours the UM Tekran is offset by ~0.5 ng m⁻³ and then jumps to ~2 ng m⁻³ at hour 30 on August 27th with the difference decreasing over the next week of measurements in an almost linear fashion. Over most of this period the UW Tekran did not report Hg(0) measurements other than a small set of measurements on August 28th that are offset by ~0.5 ng m⁻³ relative to the UNR Tekran. It can be seen that by hour 250 on September 5th all three instruments had converged. After this period the agreement between the UW, UNR and UM Tekrans was good until September 8th, when the UM instrument became contaminated after a malfunction of our permeation oven, requiring replacement with a backup Tekran 2537A unit. Both the absolute response and the response factor, i.e. the calibration factor of the UM Tekran were somewhat unstable during this period and additional details are provided in the Supplementary Information. Our focus during this initial period of the intercomparison was on the two laser / sampling systems that were being set up. In retrospect we can acknowledge that greater attention should have been paid to quality assurance with the UM Tekran. We conclude that the difference between the UM and UNR instruments is an experimental artifact. Problems with instability in the UM Tekran may have been associated with the use of an external pump to
supplement the internal Tekran pump, or with the fact that the UM instrument had been powered down for almost one week and relocated to a site at a significantly different ambient pressure. It is also noteworthy that the offsets shown in Fig. 1 occurred prior to the start of the manifold spikes of HgBr$_2$ and cannot be associated with the elevated levels of HgBr$_2$ that were introduced into the manifold on Sept. 5$^{th}$. The observations cannot, in our view, be indicative of any type of chemistry within the manifold, nor can it be indicative of the UM instrument measuring TGM rather than Hg(0). “

The Supplementary Information now contains the following:

Initial operation of the UM Tekran.
The UM Tekran was powered up after arrival at the RAMIX site and calibrated after 24 hours and again after another 24 hours. The response factors were consistent at $\sim 6 \times 10^7$. During this period the Tekran was sampling ambient air and was not connected to the RAMIX manifold which, as noted in the main text, was below ambient pressure. After connection to the manifold via 25 ft of 0.25 inch tubing an external pump was connected to allow the instrument to maintain a 1.5 L min$^{-1}$ sampling rate. At hour 72 on August 29$^{th}$ the Tekran was recalibrated and the response factor dropped to $\sim 4.3 \times 10^7$. At the next recalibration on September 2$^{nd}$ at hour 175 the response factor increased to $\sim 6 \times 10^7$ and then on September 7$^{th}$ decreased to $\sim 4.5 \times 10^7$. In addition to the instability in the response factors there was some instability in overall response that was not related to calibration. After sampling began from the manifold at hour zero, which corresponds to 9 am on August 26$^{th}$, the UM Tekran was offset by $\sim +0.5$ ng m$^{-3}$ with respect to the UNR
Tekran and this offset increased to ~ +2 ng m\(^{-3}\) at hour 30 on August 27\(^{th}\). This increase in offset occurred after the UM system was disconnected from the manifold and flushed with the \(N_2\) blowoff from a liquid nitrogen tank that gave good zeros but would have increased the pressure in the system to ambient pressure. The large offset is caused by the UM instrument reading high and the UNR instrument reading low and then over the next 45 hours the instruments converge. This instability occurred prior to any manifold spiking with \(HgBr_2\). As noted in Section 3.2 of the manuscript, our focus during this initial period of the intercomparison was on the two laser systems that were being set up. We are unable to identify the reasons for these problems which may have been associated with the use of an external pump to supplement the internal Tekran pump, or with the fact that the instrument had been powered down for almost one week and relocated to a site at a significantly different ambient pressure. In retrospect we can acknowledge that greater attention should have been paid to quality assurance with the UM Tekran. We conclude that the difference between the UM and UNR instruments is an experimental artifact.
In-situ and Denuder Based Measurements of Elemental and Reactive Gaseous Mercury with Analysis by Laser-Induced Fluorescence. Results from the Reno Atmospheric Mercury Intercomparison Experiment.

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Abstract

The University of Miami (UM) deployed a sequential two photon laser-induced fluorescence (2P-LIF) instrument for the in-situ measurement of gaseous elemental mercury, Hg(0), during the Reno Atmospheric Mercury Intercomparison Experiment (RAMIX) campaign. A number of extended sampling experiments, typically lasting 6-8 hours but on one occasion extending to ~24 hours, were conducted allowing the 2P-LIF measurements of Hg(0) concentrations to be compared with two independently operated instruments using gold amalgamation sampling coupled with Cold Vapor Atomic Fluorescence Spectroscopic (CVAFS) analysis. At the highest temporal resolution, ~5 minute samples, the three instruments measured concentrations that agreed to within 10-25%. Measurements of total gaseous mercury (TGM) were made by using pyrolysis to convert total oxidized mercury (TOM) to Hg(0). TOM was then obtained by difference. Variability in the ambient Hg(0) concentration limited our ability to measure sensitivity for measurement of ambient TOM using this approach. In addition, manually sampled KCl coated annular denuders were deployed and analyzed using thermal dissociation coupled with single photon LIF detection of Hg(0). The TOM measurements obtained were normally consistent with KCl denuder measurements obtained with two Tekran speciation systems and with the manual KCl denuder measurements but with very large uncertainty. They were typically lower than measurements reported by the University of Washington (UW) Detector for Oxidized Hg Species (DOHGS) system. The ability of the 2P-LIF pyrolysis system to measure TGM was demonstrated during one of the manifold HgBr₂ spikes but the results did not agree well with those reported by the DOHGS system. The limitations of the RAMIX experiment and potential improvements that should be implemented in any future mercury instrument intercomparison are discussed. We suggest that instrumental artifacts make a substantial contribution to the discrepancies in the
reported measurements over the course of the RAMIX campaign. This suggests that caution should be used in drawing significant implications for the atmospheric cycling of mercury, from the RAMIX results.

1.0 Introduction:

The environmental and health impacts of mercury pollution are well recognized with impacts on human health and broader environmental concerns (U.S. EPA, 2000; UNEP, 2013; Mergler et al., 2007; Diez, 2009; Scheuhammer et al., 2007). There have been extensive reviews of global emissions, measurements and biogeochemical cycling of mercury, (Mason, 2009; Streets et al., 2011; Pirrone et al., 2009; Lindberg et al., 2007, Ebinghaus et al., 2009; Sprovieri et al., 2010; Selin, 2009) The concerns associated with the mercury problem have resulted in attempts to regulate and control emissions at both national and international levels. The latest attempt in the United States is incorporated in the Mercury and Air Toxics Standards (Houyoux, and Strum, 2011; US EPA, 2013) and international efforts by the United Nations Environment Program have led to the Minamata Convention on Mercury, a global legally binding treaty on mercury controls (UNEP, 2008; UNEP, 2013; UNEP, 2014).

There is a reasonable consensus on typical background concentrations of atmospheric mercury, which are extremely low. Currently typical concentrations range from 1.2–1.4 ng m⁻³ in the Northern Hemisphere and 0.9–1.2 ng m⁻³ in the Southern Hemisphere and appear to be decreasing (Slemr et al., 2011; Sprovieri et al., 2016) [1 ng m⁻³ is ~ 3x10⁶ atoms cm⁻³ or ~ 120 ppq (parts per quadrillion)] (Slemr,)]. Until recently it has been accepted that most of the mercury found in the boundary layer is elemental mercury, Hg(0) (Lindberg et al., 2007). Oxidized or reactive gaseous mercury (RGM), normally assumed to be in the Hg(II) oxidation state, has not been chemically identified and is thought to constitute a very small fraction of the total mercury concentration although recent work (Gustin et al., 2013; Ambrose et al., 2013) challenges this view. Our overall understanding of the atmospheric chemistry of mercury and the detailed elementary chemical reactions that oxidize Hg(0) is poor (Lin et al., 2006; Hynes et al., 2009; Subir et al., 2012) and the uncertainty of both the chemical identity and measurements of speciated oxidized mercury places few constraints on models. Atmospheric measurements of mercury represent a significant challenge in ultra-trace analytical chemistry and the issues associated with current techniques have been discussed by Gustin and Jaffe (2010). We have developed a laser-based sensor for the detection of Hg(0) using sequential two-photon laser-induced fluorescence (2P-LIF) (Bauer et al., 2002; Bauer et al. 2014). The instrument is capable of fast, in-situ, measurement of Hg(0) at ambient levels. By incorporating pyrolysis to convert RGM and particulate mercury to Hg(0) it is possible to measure total gas phase mercury (TGM mercury, i.e. the sum of Hg(0) plus gas phase and particulate bound oxidized mercury) and hence to measure total oxidized mercury (TOM, i.e. the sum of gas phase and particulate bound oxidized mercury) by difference. The Reno Atmospheric Mercury Inter-comparison Experiment (RAMIX) offered an opportunity to deploy the 2P-LIF instrument as part of an informal field intercomparison at the University of Nevada Agricultural Experiment Station (Gustin et al., 2013; Ambrose et al., 2013; Finley et al., 2013).
RAMIX was an attempt to inter-compare new Hg measurement systems with two Tekran 2537/1130/1135 systems. This is the instrumentation that is currently in use for the overwhelming majority of atmospheric Hg measurements. Participants included the University of Washington (UW), University of Houston (UH), Desert Research Institute (DRI), University of Nevada Reno (UNR) and the University of Miami (UM).

The specific goals for the project were:

1. Compare ambient measurements of gaseous elemental mercury, Hg(0), gaseous oxidized mercury (RGM) and particulate bound mercury (PBM) by multiple groups for 4 weeks.
2. Examine the response of all systems to spikes of Hg(0) and HgBr₂.
3. Examine the response of all systems to Hg(0) in the presence of the potentially interfering compounds ozone and water vapor.
4. Analyze the data to quantify the level of agreement and the results of interference and calibration tests for each measurement system.

In practice the instrument operated by UH only measured Hg(0) for the first week of the campaign and the cavity ring down spectroscopy (CRDS) instrument deployed by DRI did not produce any data. Hence RAMIX was primarily an intercomparison of the UM 2P-LIF instrument, the UW Detector for Oxidized Hg Species (DOHGS) that is based on two Tekran 2537 instruments, and a Tekran 2537 and two 2537/1130/1135 speciation systems deployed by UNR. Under these circumstances we were not able to compare 2P-LIF measurements made at high temporal resolution with the CRDS instrument. It did allow us to compare the 2P-LIF sensor with independently operated instruments that use preconcentration on gold coupled with analysis by CVAFS and to examine potential interference effects. Our focus here is to compare the short term variation in GEM on the timescale that the CVAFS instruments operate, ~ 5 minute samples, and examine the ability of the different instruments to capture this variation. In addition, we made measurements of TCM and hence TOM by difference and also employed manual denuder measurements to attempt to measure RGM directly. In prior publications, Gustin et al. (2013) and Ambrose et al. (2013) provide their interpretation of the RAMIX results and their conclusions have very significant implications for our understanding of atmospheric mercury chemistry. In this work we offer a contrasting view with different conclusions.

2.0 Experimental

2.1 RAMIX Intercomparison. A detailed description of the RAMIX location and the local meteorology was provided by Gustin et al. (2013). The original RAMIX proposal included participation from Tekran Corporation to build and test a field-deployed, high-flow sampling manifold that could be reliably spiked with 10-100 parts per quadrillion of RGM. Tekran proposed to supply both GOM and GEMHg(0) spiking using independent generators that were traceable to NIST standards and would be independent of the detection systems being evaluated. However, due to time constraints Tekran believed that it was unlikely that the manifold and ultra-trace spiking system could be manufactured and fully tested to their standards, so they declined to participate in RAMIX. (Prestbo, 2016). Instead, the UW group stepped in to supply and operate the sampling manifold and spiking system and the details of its characterization are
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provided in Finley et al. (2013). During the RAMIX campaign the 2P-LIF instrument sampled on 18 days, typically sampling for between 4 and 6 hours. The longest period of continuous sampling lasted for 26 hours and occurred on September 1st and 2nd. Over this 18 day period we sampled from the RAMIX manifold and, in addition, at the end of the campaign we sampled ambient air independently and also attempted to measure TOM by pyrolyzing the sample air and measuring the difference between Hg(0) and TGM. We also sampled RGM using KCl coated annular denuders using LIIF for real-time analysis.

### 2.2 The 2P-LIF system

Bauer et al. (2002, 2003, 2014) provide a description of the operating principles of the 2P-LIF instrument. Bauer et al. (2014) provide a detailed description of the 2P-LIF instrument deployed at RAMIX including the operating principles, sampling configurations, data processing, calibration and linearity tests and together with examples of experimental data. In summary, the system uses sequential two-photon excitation of two atomic transitions in Hg(0) followed by detection of blue shifted LIF. The instrumental configuration at RAMIX utilized an initial excitation of the Hg 63P1 - 61S0 transition at 253.7 nm, followed by excitation to the 71S0 level via the 71S0 - 63P1 transition at 407.8 nm. Both radiative decay and collisional energy transfer produce population in the 61P1 level. Blue shifted fluorescence was then observed on the strong 61P1 - 61S0 transition at 184.9 nm using a solar blind photomultiplier tube (PMT). By using a solar blind tube that is insensitive to laser scatter at the excitation wavelengths very high sensitivity is possible. The use of narrowband excitation of two atomic transitions followed by detection of laser-induced fluorescence at a third wavelength precludes the detection of any species other than Hg(0). The 2P-LIF instrument requires calibration, so Hg(0) was also measured with a Tekran 2537B using its internal permeation source as an absolute calibration. In prior field campaigns we have been able to transport the Tekran with it remaining powered on which is important in maintaining the stability of the permeation oven. This was not possible during the move from Miami to Reno and so the Tekran was powered down for about one week prior to the start of measurements. We sampled from the RAMIX manifold, which was below ambient pressure, through ~25 ft of ¼ in Teflon tubing. No filter was placed on the sampling line to attempt to remove ambient RGM or the HgBr2 spikes that were periodically added to the sample flow. The sampling line was not heated and was not shielded from the sun. The original RAMIX plan called for all instruments to be located close to the manifold for optimal sampling. Unfortunately the positioning of the trailers at the actual site precluded this and forced us to use a long sampling line. As a result, the internal pump on our Tekran was not able to draw the 21.5 SLPM required for sampling and an auxiliary pump was placed on the Tekran exhaust to boost the flow.

Under atmospheric conditions the 2P-LIF instrument cannot detect RGM so, in principle, this does not need to be removed from the sample gas. However, deposition of RGM on the sampling lines followed by heterogeneous reduction to GEM could produce measurement artifacts. The limit of detection for Hg(0) during RAMIX was ~30 pg m⁻³ for a 10 s or 100 shot average.

### 2.3 Measurements of TGM and TOM
We attempted to use the 2P-LIF instrument to measure \( \text{TGTM} \) and hence \( \text{TOM} \) by difference. Although we have routinely used this approach to convert \( \text{HgCl}_2 \) and \( \text{HgBr}_2 \) to \( \text{Hg}(0) \) in the laboratory, this was our first attempt to measure total oxidized mercury at ambient concentrations. A second sampling line was attached to the RAMIX manifold and a pyrolyzer was located directly at the manifold sampling port. The pyrolyzer consisted of an ~0.6 cm o.d. quartz tube, 15 cm in length and partially filled with quartz wool. Wrapped Nichrome wire encompassed an 8 cm section of tube that was heated until the quartz began to glow. The high temperature inside the pyrolyzer reduces both \( \text{RGM} \) and particulate mercury in the manifold air to \( \text{Hg}(0) \), which is then monitored by 2P- LIF and gives the sum of oxidized (both gaseous and particulate) and elemental mercury, i.e. \( \text{TGTM} \). Directly sampling from the manifold and measuring ambient \( \text{Hg}(0) \) then allows the concentration of \( \text{TOM} \) to be calculated as the difference between the two signals. 

Both lines were continuously sampled at 10 L/min and the flow to the fluorescence cell was switched between the pyrolyzed and unpyrolyzed sample lines in, typically, 5 min intervals to attempt to track fluctuations in \( [\text{Hg}(0)] \) that would obscure the relatively small signal increase attributable to \( \text{TOM} \).

### 2.4 Manual Denuder Sampling of RGM

We conducted manual denuder sampling on seven afternoons during the RAMIX campaign to attempt to quantify total RGM. We sampled using both KCl coated annular denuders and uncoated tubular denuders that were then analyzed using programmable thermal dissociation (Ernest et al., 2013). In both cases we monitored the \( \text{Hg}(0) \) that evolved during RGM decomposition, in real time using single photon LIF. Only the annular denuder results are presented here. The use of denuder sampling coupled with thermal dissociation has been described by Landis et al. (2003) and is used in the Tekran Model 1130 Mercury Speciation Units deployed during RAMIX. Air is pulled through a KCl coated annular denuder which captures RGM but transmits elemental and particulate mercury. After a period of sampling, typically one hour, the denuder is flushed with zero grade air and the denuder is heated to 500 °C. The RGM is thermally decomposed producing elemental mercury that desorbs from the denuder surface and is then captured and analyzed by a Tekran 2537. The KCl coated annular denuders used here were manufactured by URG Corporation and were identical to those described by Landis et al for manual sampling. They were located on top of one of the RAMIX instrument trailers a few feet from the entrance to the RAMIX manifold inlet. The denuders sampled at 10 SLPM, they were not heated and the integrated elutriator/acceleration jet and impactor/coupler described by Landis et al. and incorporated in the Model 1100 speciation unit were not placed on the denuder inlet. Hence no type of particle filtering was used on the inlets. The denuders were cleaned and recoated prior to the RAMIX deployment. Prior to sampling, the denuders were cleaned by heating to 500 °C and then bagged and taken to the sampling site. After a period of sampling that varied from ~1 to 4 hours, the denuders were capped, placed in sealed plastic bags, and transported to the analysis lab at the University of Nevada, Reno. On most of the sampling days a single denuder was opened and then immediately bagged serving as a field blank. On the final two days of sampling, denuders were sampled in pairs, i.e with two denuders connected inline so that the front denuder
sampled RGM and the rear denuder served as a blank and monitor of bleed-through of RGM. The blank concentrations are typically low as shown in Table 1; however on September 10th the blank shows a very high value that is indicative of significant contamination at some point during the cleaning or sampling process. For the analysis, a flow of He passed through the denuders and then into a fluorescence cell where any Hg(0) in the flow was detected by LIF. The LIF was monitored by two PMT’s set to different gains to increase the dynamic range of the detection system. Prior to the analysis, a known amount of mercury was injected into the flow through a septum using a transfer syringe. The syringe sampled from a Tekran Model 2505 Mercury Vapor Primary Calibration Unit. Without disrupting the gas flow the denuder was then placed in a clamshell tube furnace that had been preheated to 500°C. The evolution of the Hg(0) was monitored for, typically, 5-10 minutes and after the LIF signal had returned to baseline a second calibration injection was performed. A frequency doubled, Nd-Yag pumped dye laser was used to excite the Hg(0) 6^3P_1-6^3S_0 transition at 253.7 nm and resonance LIF was observed at the same wavelength. In this approach, the detection PMT detects both LIF and laser scatter, hence sensitivity is limited by the ratio of intensity of the LIF signal to the laser scatter. Since the 6^3P_1 level is efficiently quenched by both O_2 and N_2 (Breckenridge and Unemoto, 2007) the thermal analysis was performed in He buffer gas to achieve good detection sensitivity. The excitation beam then passed through a reference cell that contained a steady flow of Hg(0) from a permeation source. The LIF signal from the reference cell served to confirm that the laser output was stable.

3.0 Results:

3.1 RAMIX Manifold

As noted above, the RAMIX manifold had to be constructed and tested by the UW group under tight time constraints, and details of its characterization are provided in Finley et al. (2013). A critique of the manifold performance has been presented by Prestbo (2014) and we detail some key issues here. The manifold deployed at RAMIX was a different size than the prototype tested in the laboratory. The laboratory manifold showed very large variation in calculated transmission efficiencies of GEM Hg(0) after spiking with a permeation source. Recoveries from 71-101% were reported by Finley et al. for short-term spikes. The GEM authors speculate that this was associated with rapid changes in ambient Hg(0) but provide no measurements to support this. The Hg(0) source used for spiking was gravimetrically calibrated by the manufacturer but was not used at the calibration temperature requiring the output to be calibrated by a Tekran 2537B. After the equipment was moved to the RAMIX site the permeation tube output increased. The authors also acknowledge a significant uncertainty (± 15%) in the RAMIX manifold flow measurements that were required to calculate spike concentrations; hence this is the minimum uncertainty in calculated spike concentrations.

We find that several independent measurements of GEM Hg(0) spikes differ by as much as 30% from the value calculated by the manifold operators suggesting that (± 15%) underestimates the uncertainty. Because of these considerations we believe the RAMIX manifold is best treated as a semi-quantitative delivery system and that was not well characterized. We do not feel it is appropriate to
characterize “recoveries” as Gustin et al. (2013) have done because of the large uncertainty in Hg(0) spike concentrations. Rather, it is most useful to focus on sampling periods when multiple independent instruments show reasonable agreement.

3.2 UM Tekran Performance

In evaluating the first week of the UM RAMIX measurements it became clear that there was some non-linearity in the relative responses of the 2P-LIF and UM Tekran systems and that better agreement was obtained by referencing the Hg(0) concentration to the UNR Tekran. Gustin et al., (2013) concluded that the UNR Tekran, based on the inlet configuration, only measured Hg(0) and they suggested that the UM system, due to the long sampling line, was measuring TGM. We compared the manifold Hg(0) readings from the UM and UNR Tekrans over the first 260 hours in which we took measurements. The absolute concentration difference relative to the UNR instrument is shown in Figure 1. Hour zero corresponds to 9 am on August 26th when we started measurements and hour 260 corresponds to midnight on September 5th. Over the first 24 hours the UM Tekran is offset by ~0.5 ng m$^{-3}$ and the absolute offset jumps to ~2 ng m$^{-3}$ at hour 30 on August 27th with the difference decreasing over the next week of measurements in an almost linear fashion. Over most of this period the UW Tekran did not report Hg(0) measurements other than a small set of measurements on August 28th that are offset by ~0.5 ng m$^{-3}$ relative to the UNR Tekran. It can be seen that by hour 250 on September 5th all three instruments had converged.

After this period the agreement between the UW, UNR and UM Tekrans was good until September 8th, when the UM instrument became contaminated after a malfunction of our external permeation oven, requiring replacement with a backup Tekran 2537A unit. Both the absolute response and the response factor, i.e. the calibration factor of the UM Tekran were somewhat unstable during this period and additional details are provided in the Supplementary Information. Our focus during this initial period of the intercomparison was on the two laser systems that were being set up. In retrospect we can acknowledge that greater attention should have been paid to quality assurance with the UM Tekran. We conclude that the difference between the UM and UNR instruments is likely to be an experimental artifact possibly Problems with instability in the UM Tekran may have been associated with the use of an external pump to supplement the internal Tekran pump, or with the fact that the UM instrument had been powered down for almost one week and relocated to a site at a significantly different ambient pressure. The initial abrupt change to a large offset followed by the linear decrease over 300 hours offsets shown in Fig. 1 occurred prior to the start of the manifold spikes of HgBr$_2$ and cannot be associated with the elevated levels of HgBr$_2$ that were introduced into the manifold on Sept. 5th. The differences between the instruments cannot, in our view, be indicative of any type of chemistry within the manifold or sampling lines, nor can it be indicative of the UM instrument measuring TGM rather than Hg(0).

3.3 2P-LIF Measurements
The absolute Hg(0) concentrations reported for the 2P-LIF measurements typically use a single 10-minute section of Tekran concentration data to calibrate the 2P-LIF signal and place it on an absolute concentration scale. The complete time series of measurements then gives a long-term comparison of the 2P-LIF and Tekran instrumentation with the absolute 2P-LIF concentrations based on the single 10-minute calibration point.

### 3.3.1 September 5th

This was the first occasion on which the three independent Tekran 2537 instruments and the 2P-LIF system reported simultaneous measurements. The 2P-LIF system sampled from the RAMIX manifold for approximately 6.5 hours from ~10:30 am to 5 pm. Over the course of the sampling period there were two spikes of Hg(0) lasting one and two hours, respectively. The UW manifold team reported an initial 10 am Hg(0) spike concentration of 26.5 ng m\(^{-3}\) dropping to 24.4 ng m\(^{-3}\) over the course of the one hour spike. The two hour spike that began at 1 pm was reported to be ~12.4 ng m\(^{-3}\) dropping to 10.5 ng m\(^{-3}\) over the course of two hours. The ambient airflow in the manifold was spiked with HgBr\(_2\) for the whole of this sampling period and the reported level of the HgBr\(_2\) spike varied between 0.6-0.7 ng m\(^{-3}\). The levels of HgBr\(_2\) measured by the DOHGS instrument were consistent with this but the concentrations reported by the UNR speciation units were considerably lower and with a significant discrepancy between the two speciation units. Figure 2a shows the sequence of Hg(0) measurements from the UNR, UW and UM Tekrans together with the 5 minute averages of the 2P-LIF signal. The 2P-LIF instrument began manifold measurements in the middle of the initial 10 am Hg(0) spike and is scaled to the concentration at this time which all three Tekrans measured as ~22.5 ng m\(^{-3}\). The three Tekrans agree to better than 5% during both of the manifold spikes and, based on a pre-spike ambient concentration of 2 ng m\(^{-3}\) it suggests that the initial spike concentration was ~20.5 ng m\(^{-3}\). This suggests that the reported spike concentration was ~25-30% larger than the actual concentration introduced into the manifold. Fig. 2b shows an expanded concentration scale to highlight the nominally ambient measurements. There is some suggestion that it took some time for the spike to be completely removed, particularly after the second spike. At the completion of the second spike all the instruments drop to ambient but the UNR instrument sees two Hg(0) "pulses". Interestingly these show up with greatly reduced amplitudes in the UW and UM Tekran signals and also in the 2P-LIF signal. Figure 3 shows the % difference of the other instruments relative to the UM Tekran and over most of the sampling period the agreement between all the measurements is better than 10% over an ~7 hour period with 5 minute sampling resolution. This indicates that the 2P-LIF instrument is capable of stable operation over an extended time period with any drifts being corrected by normalization to the reference cell. Well calibrated independently operated Tekrans should be capable of agreement to better than 5% based on tests performed by the manufacturer and this level of agreement is achieved during subsets of the sampling period. It is not clear if the deviations that are observed, particularly the large deviations seen by the UNR Tekran after the second spike are related to presence of elevated levels of HgBr\(_2\) or other issues related to manifold operation. The fact that all the instruments observed these Hg(0)
pulses suggests that the artifact may be related to a process in the manifold rather than in in the UNR sampling line. However the significant differences in the magnitude of Hg(0) pulses observed by the different instruments are difficult to rationalize.

3.3.2 September 1st and 2nd

The UM and UNR systems sampled simultaneously for a 22 hour period offering an opportunity to compare the instruments over an extended sampling period. This sampling also occurred prior to any of the manifold spikes that introduced substantial concentrations of HgBr₂ into the manifold and sampling lines. Unfortunately, the UW instrument did not report any measurements during this sampling period. The UM system sampled for 26 hours and the complete dataset is described elsewhere, (Bauer et al. 2014). This includes a detailed analysis of the short-term, i.e. 1-10 seconds, variation in the Hg(0) concentration and the ability of the 2P-LIF system to capture this. Here we focus on the simultaneous sampling period and the variability that should be resolvable by both of the Tekrans and the 2P-LIF instruments. SI Figure 1 shows the 24 hour sampling period with the 2P-LIF signal calibrated by the UM Tekran concentration at the beginning of hour 13 (i.e 1 pm on September 1st) and the corresponding measurements from the UNR Tekran. SI Figure 2 shows the same data with an expanded y-axis to highlight the variation in the ambient measurements. All three instruments track each other quite well over the first 10 hours and then measure a nocturnal increase in Hg(0) which shows greater medium term variability in the concentration. The 2P-LIF concentrations are approximately 20% greater than the Tekran measurements during this period. At hour 33 (i.e. 9 am on September 2nd) there was a manifold spike with a reported concentration of 12.9 ng m⁻³ dropping to 11.9 ng m⁻³ over the course of one hour. The UNR Tekran is ~6% lower, the UM Tekran is ~20% lower and the 2P-LIF ~22% higher than the calculated spike concentration. SI Figure 3 shows the same measurement set but with all instruments normalized to the second manifold spike at hour 33. Figure 4 shows an expanded y-axis, the concentration scale, focusing on the ambient concentration measurements. It is apparent that we now see better agreement between the 2P-LIF and the UNR Tekran but that the UM Tekran lies systematically higher than the UNR Tekran. Figure 5 shows a three hour subset of the measurements corresponding to 5-8 am on the morning of September 2nd. The variation between the instruments is greater than 5% and the short term variations in the Hg(0) concentration vary between the three instruments. Using either calibration approach we see that all instruments capture both the nocturnal increase in Hg(0) concentration and the greater variability in the signal but that there are differences in the amplitude of the variability.

3.3.3 Hg(0) Intercomparison Conclusions

Almost all of the measurements of atmospheric concentrations of Hg(0) have been made with CVAFS instrumentation and the majority of those measurements have utilized the Tekran 2537. This work provides the first extensive comparison of the Tekran 2537 with an instrument that is capable of fast in-situ detection of Hg(0) using a completely different measurement technique. Measurements over two extended
sampling periods show substantial agreement between the 2P-LIF and Tekran measurements and suggest that all the instruments are primarily measuring the same species. Intercomparison precision of better than 25% was achievable over an extended sampling period and precision of better than 10% was achieved for subsets of the sampling period. As we discuss below it is difficult to determine the extent to which interferences from GOM GGM contribute to the differences observed.

3.4 Interference Tests.

As noted above, one component of the initial RAMIX proposal was an examination of the response of the various sensors to potential interfering compounds HgBr₂, O₃ and H₂O. An analysis of the 2P-LIF detection approach suggests that, at the spike levels employed during the RAMIX campaign, neither HgBr₂ nor O₃ should have any interference effects. Changes in the concentration of H₂O do affect the 2P-LIF signal because H₂O absorbs the 2P-LIF fluorescence signal and may quench the fluorescence. In addition, O₂ also absorbs the 2P-LIF signal and quenches fluorescence thus a change in the O₂ concentration will affect the linearity of the response. We have presented a detailed discussion of these effects (Bauer et al., 2014) including an examination of two types of interferences that have been observed in LIF sensors applied in atmospheric and combustion environments and concluded that these are not potential problems in 2P-LIF measurements of atmospheric Hg(0). As we have noted previously (Bauer et al., 2014), condensation in our sampling lines can produce artifacts in Hg(0) concentration measurements. Because of the low humidity in Reno it was not necessary to use any type of cold trap during ambient measurements but we did use a trap during manifold spikes of H₂O so our measurements do not address this as a potential interference.

3.4.1 O₃ Interference Tests.

On September 7th an ozone interference test was conducted by simultaneously spiking the sampling manifold with a high concentrations of Hg(0) and ozone. The spike in Hg(0) lasted from 9am to 7:30 pm and there were two ozone spikes, each of two hours duration. A comparison of the UM, UW and UNR Tekrans and the 2P-LIF signal is shown in Figure 6. The UW Tekran only measured for a portion of this period but agrees reasonably well with the other Tekrans. The 2P-LIF signal was calibrated by the UM Tekran reading during the initial Hg(0) spike at hour 9.30. The 2P-LIF signal was online for 6 minutes at the beginning of the first ozone spike and then went offline for ~40 minutes for instrument adjustments. When the 2P-LIF came back online the magnitude of the normalized signal was low relative to the Tekrans. At hour 13 all three instruments converge and agree well over the course of the second spike. The magnitude of the 2P-LIF signal could have been affected adversely by the adjustments but any reduction in signal should have been compensated by a corresponding change in the reference cell. The elevated levels of ozone were introduced into the manifold by UV irradiation of O₂ and adding the O₂/O₃ gas mixture directly into the manifold produced a reported ~8% relative increase of O₂ levels in the manifold mixing ratio. As we note above this additional O₂ would absorb some of the 2P-LIF signal but this would be a very small effect. The enhanced quenching by O₂ is more difficult to assess but cannot explain the discrepancy.
between the Tekrans and the 2P-LIF signal. In addition the agreement during the second ozone spike was good. One possible explanation is that the increase in the O₃ mixing ratio was larger than calculated for the first spike. A second series of O₃ spikes were conducted on September 13th when we were attempting to measure total gaseous mercury using pyrolysis as described below. The 2P-LIF measurements switched on a five-minute cycle between a pyrolyzed line that would have decomposed all the ozone in the sample and a line containing the ambient air spiked with ozone. There was no difference in the 2P-LIF signal from the two sampling channels again suggesting that O₃ has no interference effects.

The changes in the Hg(0) concentration measurements shown in Figure 6 track the predicted changes in calculated spike concentration. However the calculated spike concentration is concentrations, which are also shown are 20-40% higher than the actual measurements obtained made by the UM Tekran.

3.5 Measurements of TGM, TM and TOM

We made attempts to use the 2P-LIF instrument to measure TGM and hence TOM by difference by sampling through two manifold lines. A pyrolyzer was located at the manifold on one of the sampling lines to measure TGM. The other sampling line measured ambient Hg(0). TOM was calculated from the difference in the TGM and Hg(0) concentrations and in this sampling configuration the limit of detection for TOM depends on the short term variability in ambient Hg(0) which is significant and shows a diurnal variation. The pyrolysis system was set up and tested on September 12. Manifold sampling was conducted on the 13th and 14th and sampling from the trailer roof occurred on the 15th. We calculated the means of the pyrolysis and ambient channel concentrations, and the difference which gives the TOM concentration. We also calculated the standard deviations and standard errors (SE) and used these errors to calculate in quadrature the 2SE uncertainty in the derived TOM concentration. However, as discussed below, the errors in the means do not appear to capture the full variability in Hg(0), particularly at shorter sampling times.

3.5.1 September 14th

Our most extensive sampling took place on the 14th and we were able to sample for three ~ 2 hour periods between 9 am and 8 pm. On this day there were multiple manifold spikes of HgBr₂ and also an Hg(0) spike and we have made a detailed analysis of the data for each sampling period.

The third sampling period which included a large HgBr₂ spike provided the only definitive opportunity to demonstrate the capability of 2P-LIF coupled with pyrolysis to measure oxidized mercury. The third sampling period began at ~ hour 17.3 during a manifold HgBr₂ spike that began at hour 17. A short Hg(0) spike was also introduced at hour 18. Fig. 7 shows the 2P-LIF signals from the ambient and pyrolyzed sampling lines together with the means and 1 standard deviation. The UM Tekran was offline at this time and so the 2P-LIF concentrations are calibrated by the concentrations reported by the UNR Tekran at the beginning of the Hg(0) spike which are also shown. Both the UNR Tekran and UW Tekran report very similar Hg(0) concentrations during the Hg(0) spike. Both systems report an Hg(0)
concentration of 6.7 ng m⁻³ at the beginning of the spike which, since the pre-spike concentration was ~1.9 ng m⁻³, corresponds to a spike concentration of 4.8 ng m⁻³. This is lower than the calculated spike concentration of 6.1 ng m⁻³ reported by the manifold operators and suggests that the calculated spike was ~27% higher than the actual spike concentration introduced into the manifold. Fig. 8 shows the means of each set of ambient and pyrolyzed measurements together with the 2σ variation and 2SE of the mean. Fig. 9 shows the TOM concentrations calculated from the difference together with 2SE in the TOM concentration. The reported spike concentrations and DOHGS measurements are also shown. During the initial sampling period between ~17.3-17.8 hours the 2P-LIF pyrolysis measurements do not show evidence for an HgBr₂ spike. Taking the difference between the ambient and pyrolyzed measurements during this period we obtain

\[ [\text{TOM}] = 0.05 \pm 0.05 \text{ ng m}^{-3}. \]

Shortly before the introduction of the Hg(0) spike we see clear evidence for an increase in the Hg(0) concentration in the pyrolysis sample relative to the ambient sample. We speculate that the manifold adjustments that were made to introduce the additional Hg(0) spike produced either a change in the flow or some other change in the manifold conditions that allowed the HgBr₂ spike to reach our pyrolyzer, which, as mentioned above, was located at the manifold. This difference between the two 2P-LIF signals is clearly evident by inspection of Fig. 7. The Fig. 9 shows that the TOM concentration which should consist almost exclusively of HgBr₂ is significantly larger than both the reported HgBr₂ spike concentration and the concentrations reported by the DOHGS system which are in perfect agreement. Taking the difference between the ambient and pyrolyzed measurements for hour 18.02-18.35 we obtain

\[ [\text{TOM}] = 1.20 \pm 0.17 \text{ ng m}^{-3} \text{ with } 2\text{SE uncertainty.} \]

It is important to note again that the calculated Hg(0) spike concentration is 27% larger than the measured concentration. This large difference is most likely due to errors in the flows or the permeation source output but it suggests significant uncertainty that little confidence can be placed in the calculated concentration of the HgBr₂ spike. In addition, it is clear that the DOHGS measurements show a different temporal profile of TOM. The DOHGS system reports TOM concentrations that agree almost exactly with the calculated spike concentration, at the beginning of the spike period and drop to a very low background level that is below the detection limit at the end of the reported spike period. In contrast, the 2P-LIF measurements do not show an increased TOM concentration until shortly before the introduction of the Hg(0) spike and they take ~20 minutes to drop to background levels. The UNR speciation systems sample for 1 hour and this is followed by a 1 hour analysis period so they produce a single hourly average every two hours. During this period the UNR speciation system Spec1 sampled for ~20 minutes during the spike period and then for a further 40 minutes. Spec2 was sampling ambient air outside the manifold.

SI Figure 4 shows the 7σ average of the 2P-LIF signal from the ambient and pyrolysis sample lines for the first sampling period 8-10.45 hours together with the mean and 1 standard deviation (1σ) variation in the 2P-LIF signals. SI 5 shows the means together with the 2σ variation and 2SE of the mean. It is clear that there is significant short term variability in the ambient Hg(0) concentration. SI Fig. 6 shows the TOM concentrations calculated from the difference between the pyrolyzed and ambient channels together with the calculated 2SE in the TOM concentration. The reported spike concentration and DOHGS concentration...
measurements are also shown. If we take the means of the 2P-LIF ambient and pyrolysis measurements during the reported spike period we obtain: ambient: 2.06±0.05 ng m⁻³ and pyrolyzed: 2.21±0.03 ng m⁻³ giving a TOM concentration of 0.145±0.05 ng m⁻³. The 2P-LIF measurements are consistent with the detection of TOM but they are much lower than the calculated spike and DOHGS measurements shown in Fig. 10.

SI Figs. 7-9 show the corresponding plots for the second sampling period from ~ 12.2-14 hours. The alternating sampling between the ambient and pyrolysis channels is more even and SI Fig. 7 shows that there is still variability in ambient Hg(0). The means of all the samples give: ambient: 1.72±0.02 ng m⁻³, pyrolyzed: 1.70 ± 0.02 ng m⁻³. If we take the subset of measurements that coincide with the reported spike we obtain: ambient: 1.79±0.02 ng m⁻³ pyrolyzed 1.77±0.02 ng m⁻³. In this case, the 2P-LIF measurements do not detect HgBr₂ and are not consistent with the reported spike or DOHGS measurements.

SI Figs. 10 and 11 show the averages of the TOM concentrations from the 2P-LIF system together with the measurements from the UNR speciation systems, the reported spike concentrations and 5 min DOHGS concentrations. During this sampling period Spec1 sampled from the RAMIX manifold while Spec2 sampled ambient air outside the manifold. Gustin et al.(2013) detail problems with the response of the Spec2 system and applied a 70% correction that is also shown as “Spec2 corrected”. Because both the DOHGS and 2P-LIF pyrolysis systems are expected to measure the sum of gaseous (RGM) and particulate (PBM) oxidized mercury we have plotted the sum of the RGM and PBM concentrations from the speciation systems. They are plotted at the mid-point of the 1 hour sampling period.

Over most of the measurement period the 2P-LIF pyrolysis and Spec1 measurements are consistent and lower than the DOHGS measurements. The exception is the large spike in TOM seen by the 2P-LIF system at hour 18. The spike occurred during the initial portion of Spec1 sampling and, although it measures an increase in RGM relative to Spec2, the magnitude is not consistent with the 2P-LIF pyrolysis observations.

3.5.2 September 13 th

September 13 th was the first day we were able to sample with the pyrolysis system and we sampled over a period of 5 hours. The only manifold spike during this period was an O₃ spike at 1pm that lasted one hour so the speciation instruments were attempting to measure ambient RGM. SI Figure 12 shows averages of TOM concentrations as measured by the 2P-LIF pyrolysis system together with the hourly averages as measured by DOHGS and UNR speciation instruments. The x-axis error shows the duration of the 2P-LIF measurements together with 2SE y-axis error bars. Two of the averages of the 2P-LIF measurement give a physically unrealistic negative concentration suggesting that combining the 2SE errors in the means of the ambient and pyrolyzed channels underestimates the uncertainty in the TOM measurement.

3.5.3 September 15 th

On September 15 th we sampled from the trailer roof using the same sampling lines and again alternating between the pyrolyzed and unpyrolyzed channels. SI Figure 13 shows the averages of the 2P-
LIF signal from the ambient and pyrolysis channels together with the concentrations measured by the Spec2 system that was sampling ambient air outside the manifold. The concentration obtained from the UM denuder samples described below are also shown. The UW DOHGS and Spec1 systems were sampling from the RAMIX manifold with continuous HgBr₂ spiking during this period. We see some evidence for measurable RGM in the first hour of the measurements and this is not seen by Spec 2. Later measurements show no evidence for measurable RGM concentrations.

3.6 Limits of 2P-LIF detection of TOM

As we have noted above, the limit of our detection of TOM depends on the short term variability in the ambient Hg(0) concentration because we use a single fluorescence cell and switch between pyrolysis and ambient channels. We have attempted to give an estimate of the uncertainty by taking two standard errors of the means and combining the errors in quadrature to get an estimate of the uncertainty in the TOM concentration. If the mean of the ambient Hg(0) concentration is not fluctuating significantly on the timescale of channel switching this approach should give an accurate estimate of the uncertainty in TOM.

In fact our Hg(0) observations show that the fluctuations in the Hg(0) concentration show a significant diurnal variation, with large fluctuations at night, decreasing over the course of morning hours and being smallest in the afternoon. This can be seen in the long term sampling from September 1st and 2nd and in the observations from September 14th. The observation of statistically significant but physically unrealistic negative TOM concentrations on September 13th may be explained by this. Such an artifact could be produced by contamination in the Teflon valve switching system that alternates the flow to the fluorescence cell. This type of contamination should produce a constant bias that is not actually observed. It appears that the short term variability in Hg(0) concentration produces a small bias in some cases that is not averaged out by switching between the ambient and pyrolyzed channels. For example on September 13th the initial sample period of 1.2 hours gives an RGM concentration of 0.06±0.10 ng m⁻³ while two shorter sampling periods at hour 10.5 (36 min sample) and 13.5 (12 min sample) give 0.15±0.09 ng m⁻³. Our results suggest that the use of single detection channel with switching between ambient and pyrolyzed samples is not adequate to resolve the small concentration differences that are necessary to be able to monitor ambient TOM. It is necessary to set up two detection systems, one continuously monitoring ambient Hg(0) and the other continuously monitoring a pyrolyzed sample stream giving TGM to get the precision necessary to monitor ambient TOM. Over most of the measurement periods our results are consistent with the lower TOM values reported by the UNR speciation instruments although there is a large uncertainty in the concentrations that is actually difficult to quantify. In addition, it is important to emphasize that this was our first attempt to use the pyrolysis approach to attempt to measure TOM. It is possible that the pyrolyzer was not working efficiently on September 13th. The results from September 14th are more difficult to rationalize. The 2P-LIF pyrolysis system has the sensitivity to detect the much higher values of RGM reported by the DOHGS system and the reported spike concentrations of HgBr₂. At higher concentrations, as shown in Fig. 9, the 2P-LIF system can monitor HgBr₂ with ~10 minute time resolution. Our results,
however, cannot be reconciled with those reported by the DOHGS system or the spike concentrations reported by the UW manifold team.

3.7 Manual Denuder Measurements:

As we describe above, our use of manual denuders was similar to that described by Landis et al. (2002) with the exception that we did not incorporate the integrated elutriator/acceleration jet and impactor/coupler on the denuder inlet, and the denuders were not heated. Landis et al. (2002) suggest that HgCl$_2$ is quantitatively transported through the manual denuder elutriator/impactor inlet when properly heated. In later work Feng et al. (2004) suggested that such impactors could reduce the efficiency of RGM collection. Hence although in that work there is no reference to the temperature of the impactor. In this work no type of particle filtering was used on the inlets. In addition, we used single photon LIF to monitor the evolution of Hg(0) in real-time as the RGM decomposed on the hot denuder surface during oven analysis. The analysis was carried out in He buffer gas and the Hg(0) concentration was calibrated by manual injections. The first series of measurements, i.e. September 6-14$^\text{th}$ involved single denuder sampling. On the 15 and 16$^\text{th}$ we employed tandem sampling with two denuders in series to assess the extent of RGM “bleedthrough”. We used two sets of denuders on the 15$^\text{th}$ and four sets of denuders on the 16$^\text{th}$. Fig. 10 shows the raw data for a denuder analysis showing the preheat Hg(0) calibration injections and the temporal profile of the Hg(0) LIF signal for one of the September 16$^\text{th}$ samples, denuder 1. The two traces correspond to the two monitoring PMTs set at different gains to increase the dynamic range of the measurements. Fig. 11 shows the calibrated profile for the same denuder together with the “blank” i.e. the trailing denuder. The complete set of manual denuder data together with corresponding values for the UNR speciation units that are closest in sampling time are shown in Table 1. Sampling occurred on denuders 1, 4, 6 and 7. The “trailing” denuders which we have treated as blanks, are denuders 3, 5, 8 and 9. The advantage of monitoring the RGM decomposition in real-time is shown in the September 16$^\text{th}$ data. The temporal decomposition profiles (TDP) for three of the denuders shown in Fig 11 and SI Figures 14 and 15 show reasonable agreement both in absolute concentration of Hg(0) and the time for decomposition to occur. The fourth denuder sample, SI Fig. 16, is a factor of 4-5 higher in concentration and decomposes on a longer time scale with significant structure in the TDP. Comparing the TDPs for all eight denuders it is clear that the TDP for denuder 7, which shows the anomalously high value, is very different from the TDPs for the other three sample denuders. We believe that this TDP is associated with particulate mercury that has impacted on the denuder wall and decomposes on a slower timescale giving a very different temporal profile from RGM that was deposited on the denuder wall. SI Table 1 shows the values of RGM obtained from denuder analysis together with an indication of impact from a PBM component. We have also included measurements from the UNR speciation systems that overlap with, or are close to, the times when our measurements were made. We draw several conclusions from the measurements. The values we obtain from simultaneous measurements that are not influenced by the presence of PBM agree reasonably well with each other, are broadly consistent with the values reported by
the Tekran speciation systems and are typically much lower than the values from the UW DOHGS system. Two sets of tandem denuder measurements from September 15 and 16 indicate that there is not a significant level of “bleedthrough” onto the trailing denuders. This suggests that the large differences between the DOHGS system and the UNR speciation systems are not due to specific problems with the RAMIX manifold or the speciation systems deployed at RAMIX even though Spec 2 was not functioning properly as documented by Gustin et al. (2013). The tandem sampling also demonstrates that any denuder artifact is not a result of some type of “bleedthrough” artifact that is preventing RGM from being quantitatively captured by the first denuder. These results are consistent with prior work by Landis et al. (2002) and Feng et al. (2003). It is also noteworthy that the manually sampled denuders were at ambient temperature in contrast to the speciation denuders that are held at 50 C. Hence the absolute sampling humidities are similar but the relative humidities are very different. Finally, we suggest that there is value in monitoring RGM decomposition in real time as diagnostic of particulate impact—when utilizing the annular denuders without the impactor inlet designed to remove coarse particulate matter that may be retained due to gravitational settling.

4.0 Implications of RAMIX results.

We think a realistic assessment of the RAMIX results is imperative because the interpretation of the RAMIX data and the conclusions presented by Gustin et al. (2013) and Ambrose et al. (2013) have enormous implications for both our understanding of current experimental approaches to atmospheric sampling of mercury species and to the chemistry itself. Speciation systems using KCl denuder sampling are widely used in mercury monitoring networks worldwide to measure RGM concentrations and the Gustin et al. (2013) and Ambrose et al. (2013) papers suggests these results greatly underestimate RGM concentrations with no clear way to assess the degree of bias.

4.1 Intercomparison of Hg(0)

The assessment of the Hg(0) measurements is a little different in the two manuscripts with Ambrose et al. (2013), noting that “comparisons between the DOHGS and participating Hg instruments demonstrate good agreement for GEM” where GEM refers to Hg(0), and they found a mean spike recovery of 86% for the DOHGS measurements of $\text{GEM Hg}(0)$, based on comparisons between measured and calculated spike concentrations. Gustin et al. (2013) suggest that the UM Tekran agreed well with measurements of TM reported by the DOHGS system and they “hypothesize that the long exposed Teflon line connected to the UM Tekran unit provided a setting that promoted conversion of RM to GEM, or that RM was transported efficiently through this line and quantified by the Tekran system. The latter seems unlikely given the system configuration…”, where RM refers to reactive mercury. As we note above, we believe that the best explanation for discrepancies between the UM and UNR Tekrans is an experimental issue with the UM Tekran response during the initial period of sampling. We would suggest that data from September 5th, one of the few occasions when data from multiple instruments agreed over an extended period is not compatible with either transmission or inline reduction of RGM in our sampling line. What is also significant from this data is the very large discrepancy between the spike concentrations as measured
independently by three different Tekran systems and confirmed by the relative response of the 2P-LIF measurements and the calculated spike concentration. The discrepancy, on the order of 25-30%, is larger than the manifold uncertainties suggested by Finley et al. (2013). We note other examples of the measured Hg(0) spikes being significantly lower than the calculated concentrations. In prior work we have shown that both the Tekran and 2P-LIF systems show excellent agreement over more than 3 orders of magnitude in concentration when monitoring the variation in Hg(0) in an N$_2$ diluent. It is to be expected therefore that the “recovery” of high concentration spikes should show good agreement between the different instruments as observed in the September 5th data. The difference between the observations and the calculated manifold spike concentrations is, we would suggest, a reflection of the significant uncertainty in the calculated manifold spike concentration and is not a reflection of reactive chemistry removing Hg(0). In addition, random uncertainties in the flow calculations should not produce a consistently low bias relative to the calculated spike concentrations. As we note above in section 3.1 Ambrose et al. report an increase in the output of their Hg(0) permeation tube after the move to the RAMIX site but this assumes that their Tekran calibration is accurate. The results are consistent with their Tekran measuring too high an output from the permeation device. This is significant if the same Tekran is being used to calibrate the output of the HgBr$_2$.

A more difficult issue is the question of resolving the differences in the temporal variation of ambient Hg(0) at the 5 minute timescale as captured by the different instruments. The Tekran systems should be in agreement with a precision of better than 5% and the 2P-LIF system, with a much faster temporal resolution and detection limit, should be capable of matching this. The differences here are not consistently associated with a single instrument with, for example, the 2P-LIF having some systematic offset with respect to the CVAFS systems. The extent to which the larger (i.e. larger than 5%) observed discrepancy which ranged from 10% to 25% is a result of interferences or simply a reflection of instrument precision is difficult to assess. We note again that the UM instruments had to sample through a very long sampling line and we expect that oxidized mercury is deposited on the sampling line. However it is not possible to assess the extent to which oxidized mercury is reduced back to its elemental form introducing small artifacts. As we suggest below, an intercomparison of instrument response to variation in Hg(0) concentrations in a pure N$_2$ diluent with the Hg(0) concentration varying between 1-3 ng m$^{-3}$ would provide a definitive baseline measurement of the instrument intercomparison precision and accuracy. We suggest that such a measurement is a critical component of any future intercomparison of mercury instrumentation.

4.2 Comparison of Total Oxidized Mercury

To the best of our knowledge RAMIX is the only experiment that has measured ambient TOM using multiple independent techniques. It should again be emphasized that the TOM measurements using pyrolysis with 2P-LIF detection were the first attempt to perform such measurements and the use of a single channel detection system introduced large uncertainties into the measurements. The very large discrepancies between the measurements of TOM reported by the DOHGS system, the Tekran speciation systems and the limited number of 2P-LIF pyrolyzer measurements are the most problematic aspect of the
RAMIX measurement suite. Work prior to RAMIX and suggested a potential ozone and/or humidity interference in the operation of KCl coated annular denuders and a number of studies since have also reported such an effect (Lyman et al., 2010; McClure et al., 2014). Typically however the differences between the RAMIX measurements are large and are not germane to the differences between the DOHGS and 2P-LIF pyrolyzer measurements. The SI Figures give an example of the differences between the DOHGS measurements and the denuder and 2P-LIF measurements. Ambrose et al. (2013) note that the DOHGS measurements were, on average, 3.5 times larger than those reported by the Spec1 system and summarize the comparison with denuder measurements as follows: “These comparisons demonstrate that the DOHGS instrument usually measured RM concentrations that were much higher than, and weakly correlated with those measured by the Tekran Hg speciation systems, both in ambient air and during HgBr$_2$ spiking tests.” The discrepancy of a factor of 3.5 is an average value but, for example, examining the September 14 data at ~5 am the DOHGS system is measuring in excess of 500 pg m$^{-3}$ compared with ~20 pg m$^{-3}$ measured by the speciation systems, a factor of 25 difference. At this point the Hg(0) concentration was ~ 3 ng m$^{-3}$ so based on the DOHGS measurements oxidized mercury is ~ 15% of the total mercury concentration. A recent study by McClure et al. (2014) provided a quantitative assessment of the extent to which ozone and humidity impact the recovery of HgBr$_2$ on KCl recovery. They note that although they provide a recovery equation to compare with other studies, they do not recommend use of this equation to correct ambient data until more calibration results become available. In Fig 12, we show the ozone concentration and absolute humidity for a 35 hour sampling period on September 13$^{th}$ and 14$^{th}$ that included two ozone spikes and only sampled ambient TOM. Fig 13 shows the expected denuder recovery based on the formula determined by McClure et al. which varies between a typical value of ~70% dropping to ~50% during the ozone spikes. The figure also shows the reported recoveries i.e. the ratio of RGM as measured by either the UNR speciation systems or the 2P-LIF system divided by the value reported by the DOHGS system. These values are typically much lower than those predicted by the McClure recovery expression. In addition, on September 13$^{th}$ and for most of the 14$^{th}$ the 2P-LIF pyrolysis system sees little or no evidence for high spike concentrations of HgBr$_2$ but records levels that fluctuate around those reported by the speciation systems. The one exception is the spike at hour 18 on September 14$^{th}$. We suggest that the ability of the 2P-LIF pyrolysis system to monitor large spike concentrations is shown by the measurements during the September 14$^{th}$ HgBr$_2$ spike at hour 18. The evidence for an enhancement in the pyrolyzed sample stream is observable in the raw 7s averaged data and becomes clear taking 5 minute averages. The absolute value of the pyrolyzed enhancement is obtained relative to the concentration of the Hg(0) during the spike taken from the measurements by the UNR Tekran that are in excellent agreement with the DOHGS Hg(0) values. The 2P-LIF measurements show a significantly larger HgBr$_2$ concentration and a different temporal profile compared with the DOHGS instrument. In particular, it is very difficult to rationalize the difference between the 2P-LIF and DOHGS systems during the first hour of the spike. We would suggest it is difficult to make the case that both instruments are measuring the same species. It is clear that the 2P-LIF pyrolyzer is operating efficiently based on the clear observation of
TOM at the end of the spike. We again note that the 2P-LIF system is not sensitive to TOM. It is important to note that the DOHGS instrument requires an inline RGM scrubber to remove RGM before the measurement of Hg(0). This inline scrubber utilizes deposition on uncoated quartz wool and the results of Ambrose et al. (2013) imply that while uncoated quartz captures RGM efficiently in the presence of O₃, quartz with a KCl coating promotes efficient reduction to Hg(0).

It is also reasonable to question the extent to which the Tekran speciation systems operated at RAMIX reflect the performance of these systems when normally operated under recommended protocols. As noted above, the operation of the RAMIX manifold and the Tekran speciation systems has been questioned by Prestbo (2014). In our view the two most significant issues are the performance of the two 2537 mercury analyzers associated with each speciation system and the reduced sampling rate. The performance of the two 2537 units is detailed in Gustin et al. (2013) and, as they noted, there was a significant response in each instrument. Examination of Fig SI 6 of Gustin et al. (2013) shows the relative responses of the two instruments and, using concentrations up to 25 ng m⁻³ i.e. manifold spikes, they list a regression of 0.72 [Hg(0)] + 0.08 whereas for the non-spike data they obtain 0.62[Hg(0)] + 0.25. Their Table SI 5 lists the regression including spikes as 0.7 (±0.01) + 0.2, with all concentrations expressed in ng m⁻³. When considering the use of these analyzers to monitor oxidized mercury the important factor to consider is the loading on the gold cartridge. Table SI 3 lists the mean RGM concentrations from manifold sampling as 52 pg m⁻³ for SPEC1 and 56 pg m⁻³ for SPEC2. For a 1 hour sample at 4 L min⁻¹ this corresponds to a cartridge loading of 13 pg. This is similar to the cartridge loading for sampling a concentration of 0.6 ng m⁻³ at 4 L min⁻¹ for 5 minutes. If we examine Fig SI 6 of Gustin et al. (2013) we see that the regression analyses are based on higher concentrations than 0.6 ng m⁻³, i.e. higher cartridge loadings. At concentrations of 0.6 ng m⁻³ the ratio of SPEC2:SPEC1 obtained from these regressions would be 1.05, 0.85 and 1.06 depending on which regression formula is used. We should note that based on Table SI 6 the median RGM concentrations in manifold sampling were 41 and 46 pg m⁻³. The RGM concentrations for free standing sampling were even lower with means of 26 and 19 pg m⁻³ and medians of 23 and 14 pg m⁻³ for SPEC1 and SPEC2 respectively. For concentrations below 40 pg m⁻³ the cartridge loading drops below 10 pg and in addition, the Tekran 2537 integration routine becomes significant. Swartendruber et al. (2009) reported issues with the standard integration routine and note that below cartridge loadings of 10 pg the internal integration routine produces a low bias in the Hg(0) concentration. They recommend downloading the raw data, i.e. PMT output and integrating offline. This issue has recently been discussed by Slemr et al. (2016) in a reanalysis of data from the CARIBIC program. This compounds the problem of correcting the bias between SPEC1 and SPEC2. Because the speciation instruments were sampling at 4 L/min rather than the recommended 10 L/min a large number the measurements made by the speciation systems are based on uncorrected cartridge loadings of less than 10 pg m⁻³. Based on the above we caution against drawing significant conclusions based on differences between SPEC1 and the corrected SPEC2. These differences are the basis of the conclusions of Gustin et al. (2013) that “On the basis of collective assessment of the data, we hypothesize that reactions forming
RM (reactive mercury) were occurring in the manifold” (Gustin et al. (2013) abstract). Later they state
switched between instruments on September 9. Prior to switching the slope for the equation comparing
GOM as measured by Spec 1 versus Spec 2 adjusted was 1.7 ($r^2=0.57$, $p<0.05$, $n=76$) after switching this
was 1.2 ($r^2=0.62$, $p<0.05$, $n=42$). This indicates that although there may have been some systematic bias
between denuders SPEC 2 adjusted consistently measured more GOM than SPEC 1. We hypothesize that
this trend is due to production of RM in the manifold (discussed later)." If reactions in the manifold were
producing RM then this production would surely have resulted in the DOHGS measuring artificially high,
i.e. higher than ambient, concentrations of oxidized mercury. However, the paper by Ambrose et al. (2013)
(written by a subset of the authors of Gustin et al.(2013)) makes no mention of manifold production of
oxidized mercury. In fact Ambrose et al. (2013) state, in the supplementary information to their paper, “The
same two denuders, prepared by the same operator, were used in the Tekran® Hg speciation systems from
2 to 13 September. The denuders were switched between Spec. 1 and Spec. 2 on 9 September. From 2 to 9
September, the Spec. 1-GOM/Spec. 2-GOM linear regression slope was 1.7 ($r^2 = 0.57; p < 0.05; n = 76$);
from 9 to 13 September the Spec. 1-GOM/Spec. 2-GOM slope was 1.2 ($r^2 = 0.62; p < 0.05; n = 42$). These
results suggest that the precisions of the GOM measurements made with Spec. 1 and Spec. 2 were limited
largely by inconsistent denuder performance."

The oxidized mercury concentrations presented by Ambrose et al. (2013) for the RAMIX
measurements suggests a well-defined diurnal profile that peaks at night. It is important to note that the
error bars on this profile (Figure 3 of Ambrose et al.) are one standard error rather than one standard
deviation. The standard deviations, which actually give an indication of the range of concentrations
measured show much larger errors indicating significant day to day variation in these profiles.

Nevertheless, the measurements show much larger oxidized mercury concentrations than the speciation
systems and the very limited number of 2P-LIF measurements. As we note below, there is no known or
hypothesized chemistry that can reasonably explain the large RGM concentrations seen by the DOHGS
instrument. Both Gustin et al. (2013) and Ambrose et al. (2013) draw some conclusions about the chemistry
of mercury that have significant implications for atmospheric cycling. Gustin et al. suggest in their abstract
that “On the basis of collective assessment of the data, we hypothesize that reactions forming RM were
occurring in the manifold.” Later in a section on “Implications” they conclude “The lack of recovery of the
HgBr$_2$ spike suggests manifold reactions were removing this form before reaching the instruments.” The
residence time in the RAMIX manifold was on the order of 1s depending on sampling point and there is no
known chemistry that can account for oxidation of Hg(0) or reduction of RGM on this timescale. We would
suggest that the most reasonable explanation of the discrepancies between the various RAMIX
measurements includes both instrumental artifacts and an incomplete characterization of the RAMIX
manifold. If fast gas-phase chemistry is producing or removing RGM in the RAMIX manifold the same
chemistry must be operative in the atmosphere as a whole and this requires that we completely revise our
current understanding of mercury chemistry. The discrepancies between the DOHGS and speciation
systems are further indication that artifacts are associated with KCl denuder sampling under ambient conditions but we would suggest that RAMIX does not constitute an independent verification of the DOHGS performance and that the 2P-LIF measurements raise questions about the DOHGS measurements. Ambrose et al. (2013) also suggest that the observations of very high RGM concentrations indicate multiple forms of RGM and that the concentrations can be explained by oxidation of Hg(0), with O₃ and NO₃ being the likely nighttime oxidants. We have discussed these reactions in detail previously (Hynes et al., 2009) and concluded that they cannot play any role in homogeneous gas phase oxidation of Hg(0). Ambrose et al. (2013) cite recent work on this reaction by Rutter et al. (2012) stating that “On the basis of thermodynamic data for proposed reaction mechanisms, purely gas-phase Hg(0) oxidation by either O₃ or NO₃ is expected to be negligibly slow under atmospheric conditions; however, in the case of O₃-initiated Hg(0) oxidation, the results of laboratory kinetics studies unanimously suggest the existence of a gas-phase mechanism for which the kinetics can be treated as second-order.” We would suggest that a careful reading of the cited work by Rutter et al. (2013) demonstrates the opposite conclusion. We provide additional discussion of these issues in the SI and again conclude that O₃ and NO₃ can play no role in the homogeneous gas phase oxidation of Hg(0).

5.0 Future Mercury Intercomparisons:

The discrepancies that are discussed above suggest a need for a careful independent evaluation of mercury measurement techniques. The approaches used during the evaluation of instrumentation for the NASA Global Tropospheric Experiment (GTE) and the Gas-Phase Sulfur Intercomparison Experiment (GASIE) evaluation offer good models for such an evaluation. The Chemical Instrument and Testing Experiments (CITE 1-3) (Beck et al., 1987; Hoell et al., 1990; Hoell et al., 1993) were a major component of GTE establishing the validity of the airborne measurement techniques used in the campaign. The GASIE experiment (Luther and Stetcher, 1997; Stetcher et al., 1997) was a ground based intercomparison of SO₂ measurement techniques that might be particularly relevant to issues associated with mercury measurement. In particular, GASIE was a rigorously blind intercomparison that was overseen by an independent panel consisting of three atmospheric scientists none of whom were involved in SO₂ research. We would suggest that a future mercury intercomparison should be blind with independent oversight. Based on the RAMIX results it should consist of a period of direct ambient sampling and then manifold sampling in both reactive and unreactive configurations. For example an unreactive configuration would consist of Hg(0) and oxidized mercury in an N₂ diluent eliminating any possibility of manifold reactions and offering the possibility of obtaining a manifold blank response. Such a configuration would allow the use of both denuder and pyrolysis measurements since it is reasonable to conclude, based on the current body of experimental evidence, that denuder artifacts are associated with ambient sampling with water vapor and ozone as the most likely culprits. A reactive configuration would be similar to the RAMIX manifold configuration with atmospheric sampling into the manifold and periodic addition of Hg(0) and oxidized mercury over their ambient concentrations. The combination of the three sampling configurations should
enable instrumental artifacts to be distinguished from reactive chemistry in either the manifold itself or, for example, on the KCl denuder.

6.0 Conclusions

We deployed a 2P-LIF instrument for the measurement of Hg(0) and RGM during the RAMIX campaign. The Hg(0) measurements agreed reasonably well with instruments using gold amalgamation sampling coupled with CVAFS analysis of Hg(0). Measurements agreed to 10-25% on the short term variability in Hg(0) concentrations based on a 5 minute temporal resolution. Our results also suggest that the operation of the RAMIX manifold and spiking systems were not as well characterized as Finley et al. (2013) suggest. We find that the calculated concentration spikes consistently overestimated the amount of Hg(0) introduced into the RAMIX manifold by as much as 30%. This suggests a systematic error in concentration calculations rather than random uncertainties that should not produce a high or low bias.

We made measurements of TOM, and hence TOM by difference, by using pyrolysis to convert TOM to Hg(0) and switching between pyrolyzed and ambient samples. The short term variation in ambient Hg(0) concentrations is a significant limitation on detection sensitivity and suggests that a two channel detection system, monitoring both the pyrolyzed and ambient channels simultaneously is necessary for ambient TOM measurements. Our TOM measurements were normally consistent, within the large uncertainty, with KCl denuder measurements obtained with two Tekran Speciation Systems and with our own manual KCl denuder measurements. The ability of the pyrolysis system to measure higher RGM concentrations was demonstrated during one of the manifold HgBr₂ spikes but the results did not agree with those reported by the UW DOHGS system. We would suggest that it is not possible to reconcile the different measurement approaches to TOM. While there is other evidence that KCl denuders may experience artifacts in the presence of water vapor and ozone the reported discrepancies cannot explain the very large differences reported by the DOHGS and Tekran speciation systems.

Similarly, the differences between the DOHGS and 2P-LIF pyrolysis measurements suggest that one or both of the instruments were not making reliable, quantitative measurements of RGM. We suggest that both instrumental artifacts, an incomplete characterization of the sampling manifold, and limitations in the measurement protocols make significant contributions to the discrepancies between the different instruments and that it would be rash to draw significant implications for the atmospheric cycling of mercury based on the RAMIX results. This is particularly true of the RGM results. If one were to conclude that the discrepancies between the DOHGS and speciation systems sampling ambient oxidized mercury are accurate and reflect a bias that can be extrapolated to global measurements then it means that atmospheric RGM concentrations are much higher than previously thought and that we have little understanding of the atmospheric cycling of mercury. What is not in dispute is the urgent need to resolve the discrepancies between the various measurement techniques. The RAMIX campaign provided a valuable guide for the format of any future mercury intercomparison. It clearly demonstrated the need to deploy high accuracy calibration sources of Hg(0) and oxidized mercury, the need for multiple independent methods to measure...
elemental and oxidized mercury and to clearly characterize and understand the differences reported by
instruments that are currently being deployed for measurements.

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Table 1: RAMIX Manual KCl Denuder Sampling

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- * evidence from TDP’s for presence of PBM
- Measurements for UNR Speciation system made at similar times. The Spec 2 measurements are uncorrected values.

Figures.
Figure 1. Comparison of Hg(0) readings from the UM, UW and UNR Tekrans over the first 260 hours of UM measurements. The absolute concentration difference relative to the UNR instrument is shown in black for the UM Tekran and in red for the DOHGS (UW) Tekran.
Figure 2: a). A seven hour sequence of GEM measurements from September 5th that included two manifold spikes. Shown are the sequence of GEM measurements from the UNR, UW and UM Tekrans together with the 5 minute averages of the 2P-LIF signal. b) An expanded concentration scale focusing on ambient measurements.

Figure 3: Seven hour measurement period from September 5th. The % difference of the UNR (black line) and UW (red line) Tekrans and the UM 2P-LIF (blue line) measurements relative to the UM Tekran is shown.
Fig 4: 22 hour sampling period from September 1\textsuperscript{st} and 2\textsuperscript{nd}. Comparison of the UM (red line) and UNR (green line) Tekrans with the UM 2P-LIF (black line) concentrations. The concentrations for each instrument are scaled to force agreement during the second manifold spike at hour 33. This is the data from SI Fig. 3 with the concentration scale expanded to shown only ambient data.
Fig 5: A section of the 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 concentrations for each instrument are scaled to force agreement during the second manifold spike at hour 33. This is the data from SI Fig. 3 with the concentration scale expanded to shown only ambient data between hours 29 and 32.
Figure 6: September 7th an ozone interference test. A comparison of the UM, UW and UNR Tekrans and the UM-2P-LIF measurements. The “expected” concentration calculated from the ambient Hg(0) concentration prior to the spike plus the calculated spike concentration is also shown.
Fig 7: September 14 measurements hours 17-19 (5-7pm). The background subtracted 2P-LIF signals from the ambient (black) and pyrolyzed sampling lines (red) are shown. The gaps correspond to times when the laser was blocked to check power and background. The means and 1 standard deviation of each sample are shown. The absolute Hg(0) concentrations are obtained by scaling the ambient Hg(0) signal to the absolute Hg(0) concentration reported by the UNR Tekran during the Hg(0) manifold spike.
Fig. 8: September 14 measurements hour 17-19. 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.
Fig 9: TOM concentrations calculated from the difference between the pyrolyzed and ambient sample concentrations together with 2SE in the TOM concentrations. The reported HgBr$_2$ spike concentrations and DOHGS measurements are also shown.
Fig. 10: September 16th KCl manual denuder measurements. The raw data for the temporal decomposition profiles (TDP) for the denuder D1 is shown.
denuder measurements. The calibrated temporal decomposition profiles (TDP) for the tandem denuder pair, D1 and D3 are shown.
Fig. 12: The ozone concentration and absolute humidity for a 35 hour sampling period on September 13th and 14th that included two ozone spikes and only sampled ambient TOM.
Fig. 13. Expected denuder recovery based on the formula determined by McClure et al. which varies between a typical value of ~70% dropping to ~50% during the ozone spikes. The figure also shows the reported recoveries i.e. the ratio of RGM as measured by either the UNR speciation systems or the 2P-LIF system divided by the value reported by the DOHGS system.