Use of Multiple Lines of Evidence to Understand Reactive Mercury Concentrations and Chemistry in Hawai‘i, Nevada, Maryland, and Utah, USA

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ABSTRACT: To advance our understanding of the mercury (Hg) biogeochemical cycle, concentrations and chemistry of gaseous oxidized Hg (GOM), particulate-bound Hg (PBM), and reactive Hg (RM = GOM + PBM) need to be known. The UNR-RMAS 2.0 provides a solution that will advance knowledge. From 11/2017 to 02/2019, the RMAS 2.0 was deployed in Hawai‘i, Nevada, Maryland, and Utah to test system performance and develop an understanding of RM at locations impacted by different atmospheric oxidants. Mauna Loa Observatory, Hawai‘i, impacted by the free troposphere and the marine boundary layer, had primarily −Br/Cl RM compounds. The Nevada location, directly adjacent to a major interstate highway and experiences inputs from the free troposphere, exhibited −Br/Cl, −N, −S, and organic compounds. In Maryland, compounds observed were −N, −S, and organic-Hg. This site is downwind of coal-fired power plants and located in a forested area. The location in Utah is in a basin impacted by oil and natural gas extraction, multiday wintertime inversion episodes, and inputs from the free troposphere. Compounds were −Br/Cl or −O, −N, and −Br/Cl. The chemical forms of RM identified were consistent with the air source areas, predominant ion chemistry, criterion air pollutants, and meteorology.

INTRODUCTION

Atmospheric mercury (Hg) chemistry is a poorly understood component of the Hg biogeochemical cycle. Gaseous elemental Hg (GEM) in the atmosphere is oxidized to form gaseous oxidized Hg (GOM). Both can interact with aerosols to form particulate-bound Hg (PBM).1 GEM can be deposited to and re-emitted from surfaces (e.g., Miller and Gustin2). GOM can be deposited to surfaces and emitted back to air as GEM.3 Since oxidation reactions can occur quickly and deposited GEM and GOM can be re-emitted, the lifetime of gaseous Hg in the atmosphere may be as short as a few hours.3 While in the free troposphere/stratosphere, the lifetime is on the order of 4–12 months. Currently, the only commercially available instrument for measurement of the three forms of atmospheric Hg is the automated Tekran 2537/1130/1135. The instrument components collect GOM, GOM, and PBM, respectively. The KCI-coated denuder used to collect GOM has been shown to underestimate concentrations by up to 13 times.3−7 There are artifacts associated with the PBM measurement, and the 1135 unit has been shown to collect GOM.5,8 In addition, it has been demonstrated that GOM collected by the denuder may be reduced to GEM9 and there are likely other artifacts associated with the KCI denuder and the particulate filter that we are not aware of. Because of the limitations of this instrument, this research focused on using a membrane-based system for the collection of reactive Hg (RM = GOM + PBM) compounds to quantify total RM concentrations as well as identify RM compounds present in ambient air at field sites with different atmospheric chemistry. The University of Nevada, Reno- Reactive Mercury Active System (UNR-RMAS) has been a technology in the making for over 10 years and has been deployed at sites in Nevada,9 Florida, and10 Australia and Tasmania10 (see Graphical Abstract). The UNR-RMAS 2.011 was used in this study at locations in Hawai‘i, Nevada, Maryland, and Utah. In 2019, it was also deployed at Svalbard, Norway, and in Nanjing, China. The RMAS 2.0 uses cation exchange membranes (CEM) to quantify RM concentrations and nylon membranes to identify RM chemistry.5,11 CEM are digested and analyzed for RM
concentrations. Nylon membranes are thermally desorbed to determine the chemistry of the RM compounds on the membrane. For additional information on improvements to the RMAS system, see the Supporting Information and Luippold et al. (2020).11

The RMAS 2.0 has a few known limitations, including the need to be deployed for 3 days to 2 weeks. In addition, better understanding of limitations of the membranes as collection surfaces is needed though recent work has increased confidence in the CEM ability to quantify RM. For example, a paper recently submitted by Lyman et al.12 compared concentrations measured using a calibrated dual channel system that quantified GEM by scrubbing air of RM using a CEM and measured total gaseous Hg (TGM) by pulling air through a pyrolyzer, allowing for calculation of RM (RM = TGM – GEM). When the RMAS 2.0 collected data alongside the dual channel system, RMAS 2.0 CEM average values were the same as the dual channel system (34 ± 5 versus 36 ± 13 pg m−3, respectively). The dual channel system was calibrated using permeated HgCl2 and HgBr2.

In this research, the utility of the RMAS 2.0 as an RM measurement method was investigated. CEM and nylon membrane RM concentrations and chemistry for each deployment period were compared with meteorology and criterion air pollutant data from the field sites. Potential sources of RM compounds were determined using back trajectory analyses. Ion chromatography was used to determine anion chemistry on membranes. Given previous observations and the improved RMAS system, research hypotheses included (1) the RMAS 2.0 will collect more RM than the Tekran system, but this will vary depending on the chemistry of the RM compounds; (2) breakthrough observed could be impacted by artifacts or RM chemistry; (3) understanding breakthrough associated with the membranes will allow for developing a better understanding of what membranes capture; and (4) comparison of nylon and CEM concentrations will provide a framework for better understanding of RM chemistry, and differences can be explained based on the dominant RM forms present. To address these research hypotheses, a year of RM and ancillary data were collected for four field sites. Specific thermal desorption profiles from each site were compared with corresponding back trajectories, anion chemistry, and ancillary data. The percentage of different RM compounds present was also evaluated.

**METHODS**

**Field Sites.** The National Oceanic and Atmospheric Administration (NOAA) Mauna Loa Observatory (MLO) is on the island of Hawai‘i (19.5392, −155.5792; 3397 masl). This site is far from major sources of pollution and an optimal air sampling location in the free troposphere (www.esrl.noaa.gov/gmd/obop/mlo/). NOAA operates a Tekran 2537/1130/1135 speciation system as a part of the AMNet network (2010 to present) (see Graphical Abstract). Deployment began on February 7, 2018 and ended on March 6, 2019, with membranes collecting RM for 2 weeks (n = 27).

The location in Reno, NV (39.5375, −119.8047; 1371 masl) was at the College of Agriculture, Biotechnology, and Natural Resources Agricultural Experiment Station Valley Road Greenhouse Complex (GH). Membranes were deployed for 2 weeks from March 15, 2018 until July 17, 2018, and from then on for 1 week until March 12, 2019 (n = 42). The GH site is heavily influenced by road sources due to proximity to I-80, a major US interstate highway. This location hosted the UNR-RMAs past, and during this project a Tekran 2537/1130 system.

The Piney Reservoir site (MD) is north of Frostburg, Maryland (39.7053, −79.0122; 769 masl) located near the Maryland—Pennsylvania border and is downwind of coal-fired power plants in Ohio, Pennsylvania, and West Virginia. Deployments at this location were for 2 weeks and spanned from December 20, 2017 to November 12, 2018 (n = 18). Previous studies at this field site indicated that it has elevated GOM concentrations due to direct impacts by local power plant sources from the west.13–15 This site operated a Tekran 2537/1130/1135 speciation system under AMNet (2008 to present).

Horsepool Monitoring Station, Utah (UT) (40.1434, −109.4689; 1567 masl) is in the Uintah Basin south of Vernal and Dinosaur National Monument. This area has significant oil and natural gas activity. The basin sometimes has persistent, multiday winter inversion episodes and high ozone16 that may impact oxidation chemistry of the Hg collected on the membranes. Deployments here were for 2 weeks and spanned from November 21, 2017 until February 27, 2019 (n = 27).

Details on the membranes, deployment, percent breakthrough calculations, and thermal desorption are described in the Supporting Information. All dates presented hereafter represent the end date of the sampling period.

**Peak Deconvolution of Thermal Desorption Profiles.** A quantification method was developed to deconvolute and define individual GOM compound peaks from the thermal desorption profiles of field samples.11 Based on the pure standard compounds and calibration of exact desorption temperature,17 GOM compounds peak in the following ranges: 80–85 °C for [−O], 90–110 °C for [Br/Cl], 125−135 °C for [−N], 150−155 °C for [−S], and 180–190 °C for methylmercury (MeHg) or organic-bound compounds. Note that there are time periods when multiple overlapping compounds were found. The exact chemistry of the compounds is unknown; a method is needed to identify RM compounds other than the standard thermal desorption profiles already established. By comparing field sample desorption profiles to the profiles generated using pure compounds, the profile peaks from the field sample were assigned compound types. For peak deconvolution method details, see the Supporting Information. Fine particles will be collected on the filters; because of this, we are currently working on a system to separate GOM and PBM.

**Tekran Speciation System.** Tekran systems were maintained, and data was processed following AMNet protocols (http://nadp.slh.wisc.edu/lib/manuals/AMNet-2003_Operations_Manual_v_3-0.pdf). Data collected that did not meet AMNet A or B quality standards were omitted from this study. For details on the position of the inlets, see the Supporting Information.

**Ion Chromatography.** Anions on nylon membranes were determined using ion chromatography (IC), providing a framework for the assessment of whether identified RM compounds (i.e., by thermal desorption) were similar to major anions present in air. For more details on the ion chromatography method, see the Supporting Information.

**Back Trajectories.** The hybrid single-particle Lagrangian integrated trajectory model (HYSLIGHT), developed by NOAA Air Resources Laboratory,17,23 was used to calculate 24 h back trajectory simulations (GDAS 1° archive). Trajectories were
created throughout the membrane sampling periods and then analyzed using the gridded frequency distribution method. For details and selected modeling parameters, see the Supporting Information text and Table SI 2.

Ancillary Data and Statistical Analyses. Sources of ancillary data are presented in Table SI 3. Details of regression analyses are in the Supporting Information. Maps were made using ArcMap 10.6.1.

■ RESULTS AND DISCUSSION BY SITE

MLO. At MLO, RM concentrations on the CEM and nylon membrane were $133 \pm 5$ pg m$^{-3}$ (mean $\pm$ standard deviation) and $77 \pm 5$ pg m$^{-3}$, respectively, for the entire sampling campaign. During this time, the Tekran system collected $82 \pm 38$ pg m$^{-3}$ RM. RM concentrations were highest at MLO compared to the other sampling locations (Figure 1a). Mean breakthrough for CEM was $5.0 \pm 1.0\%$, and $1.2 \pm 0.3\%$ for nylon membranes. It is noteworthy that at MLO, there were two time periods when the CEM and Tekran RM concentrations were not statistically significantly different, and two in which the overlap was close. Looking at temperature, and relative and absolute humidity across the time period, there was nothing unique about this time period (data not shown).

There were two common thermal desorption profile shapes observed at MLO. Most (89%) had a single bell-shaped peak

![Figure 1. RM concentrations collected on CEM and nylon membranes at the four field sites and Tekran system RM concentrations (MLO 1130/1135/2537, GH 1130/2537, MD 1130/1135/2537). The gray box highlights the duration of the Kilauea volcanic eruption. Note differences of the scale on y-axes. Error bars (1σ) are shown where $n \geq 3$.](https://dx.doi.org/10.1021/acs.est.0c02283)
in the −Br/Cl range (Figure 2 MLO a). The profile shown in Figure 2a was dominated by −Br/Cl compounds based on peak deconvolution, with 74% of the RM desorbing the in the −Br/Cl temperature range of 90–110 °C. Ten day back trajectory analyses for this sample indicated that air moved to MLO over the Pacific Ocean, and >25% of this air resided within the marine boundary layer (Figure 3, brown grids). The percentage of the points within 100 km of the site was 3%. All profiles obtained at MLO for the entire sampling campaign can be found in Figure SI 1. Additional trajectories are shown in Figure SI 2.

The second thermal desorption profile shape at MLO was bimodal, with peaks in the −Br/Cl range and in the −N range. N peaks were only detected at MLO after the 2018 Kilauea eruption began. These samples (n = 3) were collected on 04/06/18, 18/06/18, and 30/07/18. The desorption profile for 18/06 (Figure 2 MLO b) included 78% −N, 12% −Br/Cl, and 10% −S. Back trajectories for this deployment showed that air came to MLO from over the Pacific Ocean then up from the south over the Kilauea eruption. Many trajectories resided beneath the marine boundary layer (Figure 3b, brown), indicating a mix of influences from the free troposphere and within the marine boundary layer from the direction of Kilauea, with contributed air within 100 km being 4%. For additional profiles with ion chemistry and back trajectories for MLO, see Figure SI 4. All ion chemistry is presented in Table SI 3.

Figure 2. Examples of the common desorption profile shapes from each field site. Black curves with black data points are the average thermal desorption profile from replicate (n = 3) nylon membranes. Curves without data points are the deconvoluted peaks of RM compounds, where yellow = −Br/Cl, brown = −Br/Cl/N, blue = −O, gray = −N, green = −S, and purple = organic-Hg. Percentages represent relative amounts of RM compounds in each sample compared to the total RM. The profiles from each site are labeled alphabetically to correspond with the back trajectories shown in Figures 3 and 4. Two curves were deconvoluted in the Br/Cl temperature range for UT-a.

The Kilauea eruption from 04/05 to 04/08/18 was the largest lower East Rift Zone eruption and caldera collapse in 200 years.19 GEM concentrations at MLO during this period were not significantly correlated with criterion air pollutants (p = 0.96; Table SI 4). Selected air pollutants, RM, and GEM measurements during the time period of the eruption are plotted in Figure SI 3. Sulfur dioxide (SO2) concentrations increased after the eruption as did RM concentrations; however, O3, relative humidity, and GEM showed no trends. RM chemistry and ancillary criterion air pollutant data suggest that the eruption may have influenced the chemistry of the compounds present; however, we do not have enough data to understand this.
RM, particularly −Br/Cl compounds, was likely formed from ocean and free tropospheric influences. Lin et al.\textsuperscript{20} found that increased O\textsubscript{3} precursor emissions (NO\textsubscript{x} and hydrocarbons) from Eurasia caused elevated autumn O\textsubscript{3} concentrations at MLO from 1996−2011 relative to the previous decade (1980−1995). Observations were coupled with models and CO tracers that indicated increased airflow from the west to MLO during the fall.\textsuperscript{20} Thermal desorption profiles from 08/18−11/18 had pronounced −Br/Cl peaks with no secondary or widened peaks in the −N range. O\textsubscript{3} for this time were <40 ppb and not elevated relative to the rest of the sampling campaign. MLO 10 day back trajectories (Figure SI 2) did not reside in the Eurasian source boxes for significant times. These trends indicate that the RM compounds on the membranes were not likely influenced by Eurasian oxidants. For discussion of correlations between membrane concentrations, Tekran data, and ancillary data for all sites, please see the Supporting Information.

Nylon membrane RM concentrations were always lower than CEM concentrations at MLO (Figure 1a). However, there was variation in the difference with the nylon membranes being 10 to 87\% lower than the CEM. The largest differences in concentrations occurred from August through October, when nylon membrane RM concentrations were 30 to 87\% lower than CEM. There were no significant correlations...
between percent difference values (Equation 1) and the ancillary data ($p > 0.05$).

MLO nylon membranes analyzed for anions by IC ($n = 9$) identified \( SO_4^{2-} \) that ranged from 11 to 100% of all anions detected. The next most prevalent anion measured was chloride (Cl$^-$) at 9–64% (Table SI 3). The presence of Cl$^-$ supports the profile peak deconvolution results of –Br/Cl compounds in corresponding thermal desorption profiles. Although \( SO_4^{2-} \) was measured on the membranes, Hg–S compounds were not present in the thermal desorption profiles. It is possible that RM compounds were not generated by oxidation associated with S compounds. HgSO$_4$ is typically generated from heterogeneous oxidation.$^{25,26}$ Both S and Cl would be emitted from the ocean.

Another high-elevation study by Maruszczak et al.$^{23}$ using a polyethersulfone cation exchange membrane and a Tekran system, was done at the Pic du Midi Observatory for 6 months. In this work, the authors found concentrations of \( \pm 58 \) pg m$^{-3}$ in dry free troposphere air and that CEM concentrations were 1.3 times higher. Given that the Hg compounds were likely Br-based and air was dry, the denuder was likely working fairly efficiently. During our study, RM concentrations measured by the Tekran at MLO reached these values when the volcanic eruption was occurring. Otherwise, values were lower, and thermal desorption profiles indicated that compounds were primarily –Br- and –Cl-based. Case studies demonstrated that a significant component of sampled air was derived from the marine boundary layer; this is supported in this study by RH being >35% most of the time (see excel file in the Supporting Information).

Marszczak et al.$^{23}$ also suggested that adding the flush data from the Tekran improved the agreement with their membranes. For MLO, the two flush concentrations were provided by NOAA and added to the RM concentrations. Linear regression analyses of the available data points for 2018 (<1/2 of all data) resulted in the following equation: RM + flush = 1.04(RM) + 1.8. $r^2 = 0.995$, $p < 0.001$ and $n = 1205$. Thus, an additional artifact associated with the Tekran is loss of RM with the initial flush cycles. The correlation between the Tekran and CEM data for this location was $r^2 = 0.23$ and $p = 0.01$ (Table SI 4). Thus, the correlation was poor.

**GH.** Mean RM concentration on the CEM at GH was 60 ± 40 pg m$^{-3}$ and on the nylon membranes was 44 ± 3 pg m$^{-3}$ (Figure 1b). The Tekran system was offline most of 2018. While operating, mean RM concentrations were 5 ± 4 pg m$^{-3}$, and concentrations may be underestimated for the 1135 unit deployment data shown in Figure 2 GH (a) was 0.7%, low relative to other nylon membranes from GH (see excel file in the Supporting Information). HYPLIT trajectories for profile 3a showed that the GH site was influenced by long-range transport over the Pacific Ocean. Of the calculated trajectories, 0.02% resided in the Northern Eurasian source box, 0.01% in the East Asian source box, 0.1% in the San Francisco, California source box, and 8.8% within 100 km. The grid frequency distribution showed that trajectories were concentrated over the Pacific Ocean (yellow, 5–25% of trajectories; Figure 3). These trajectory patterns for transport to GH were similar to those reported in Pierce et al.$^{25}$

For the deployment ending on 25/09/18 (Figure 3 GH b), the thermal desorption profile peaked in the –Br/Cl range and had a secondary peak in the –S range. The total residence time for trajectory points within the Northern Eurasian source box and East Asian source box was 0.2%. The frequency distributions indicated that air traveled over East Asia (Figure 3 GH b), then over the Pacific Ocean to the GH site in Nevada with 4.3% of the trajectory points within 100 km of the site. It is possible that the –S was derived from the marine boundary layer or from Asian sources as described in VanCuren and Gustin.$^{26}$

Profile Figure 2 GH c had very low concentrations and contained –O, –Br/Cl/N, –S, and organic-Hg compounds. The dominant peak was between the –Br/Cl and –N temperature ranges. Trajectories demonstrated a similar pattern to the previously discussed GH samples (Figure 3) with 6% of the trajectory points residing in the Northern Eurasian source box, 2% in the East Asian source box, 2.1% in the San Francisco source box, and 5% within 100 km of the site. These trajectories were similar to those reported in Pierce et al.$^{23}$ Trajectories in brown grids indicated 25% probability of residence in the marine boundary layer. High-elevation air parcels moved from Eurasia and dropped into the marine boundary layer before arriving at the GH site. Percent breakthrough of this sample (10%) was higher than for the samples GH a and b, and total RM on this nylon membrane was lower, indicating the possibility that the membrane did not collect RM chemistry associated with –N as effectively as the other cases where –Br/Cl was dominant. This supports the hypothesis that the nylon membranes collect compounds with varying efficiencies in ambient air. Additional trajectories with ion chemistry are in Figure SI 4, and GH profiles are in Figure SI 6.

At the GH, the first instance of nylon concentrations exceeding CEM concentrations was observed. Nylon membrane RM concentrations at the GH site ranged from 90% lower to 23% higher than the CEM (Figure 1). During July through September, 2018, nylon membrane concentrations exceeded CEM concentrations by 5–23%, the maximum measurements for temperature (26.9 °C) and O$_3$ (73 ppb) were observed, along with higher solar radiation than in other seasons (>240 W m$^{-2}$). It is thought that if O$_3$ directly formed HgO, this would make an aerosol that could be rapidly deposited; however, Hg oxide forms measured at the GH were measured as both GOM and PBM using the CEM and PTFE system, respectively.$^3$

Another high-elevation study reported by Maruszczak et al.$^{23}$ for the same location and measured by the Tekran. This is within the range of 1.7 to 13 was not used. When the Tekran was online, CEM RM and concentrations may be underestimated for the 1135 unit could be O$_2$–Hg–SO$_4$–Hg–NO$_3$ or other unidentified compounds.

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Ion chromatography results for the GH ($n = 15$) showed the presence of SO$_4^{2-}$ in all samples (Table SI 3). Cl$^-$ was present
in all samples analyzed except two, and the presence of Cl$^-$ supports the observations of Br/Cl in GH samples. For Figure 3 GH b, the anion concentrations were 93% SO$_4^{2-}$ and 7% Cl$^-$ ions that coincided with the compounds found on peak deconvolution results for this profile: 48% Br/Cl/N and 28% S. An additional profile with this pattern is shown in Figure SI 4 (sample 5). Anions detected in these samples match the compounds desorbed from the membrane.

**MD.** Mean RM concentration for samples collected at MD was $17 \pm 2$ pg m$^{-3}$ for CEM with a mean breakthrough of 14 ± 7%. For nylon membranes, the mean RM concentration was $6 \pm 2$ pg m$^{-3}$ with a breakthrough of 8 ± 3%. This site had the lowest total RM concentrations on membranes.

There were two common profiles at this location (Figure 2 and Figure SI 4 samples 6 and 7). All MD thermal desorption profiles had peaks in the −N range, but the second common profile shape (sample b) had low concentrations and a large component of organic-Hg compounds in addition to −N. Additional thermal desorption profiles are shown in Figure SI 7.

Sample a was characterized by a strong −N peak (69% peak) with −S (19%) and −Br/Cl (12%) compounds (Figure 2). Sample Figure 2 MD a, along with other profiles sharing this shape, had high percent breakthrough (12%). This suggests, similar to GH results, that RM compounds collected on the nylon membrane when N-based have a lower capture efficiency than the other GOM compounds on the membranes. Trajectory points for this sample did not reside in any defined source boxes for significant times. During this sample deployment, the MD site was influenced by air traveling across the continental United States with air masses originating from the west and southwest (Figure 4 MD a) and 1% within 100 km of the site. There was a high density of trajectories, >50% of total (red), over Pennsylvania, Ohio, and West Virginia, areas with documented coal burning activity that release SO$_2$, NO$_x$, chlorine, and Hg$^{13,14}$. The influence of coal burning near the area is reflected by the composition of the RM species on the membrane. The airflow pattern further supports that coal-fired power plant emissions influenced RM chemistry at MD.

The profile for Sample b was bimodal, with organic-Hg compounds (54%), −N (15%), −S (3%), and −Br/Cl (28%) compounds (Figure 2 MD b). The samples at this site with high organic-Hg were collected from May through July, 2018. Biogenic compounds, such as isoprene and nitrogen oxides, have been observed to be highest during the summer months.
in North America. Additionally, MD samples dominated by organic-Hg compounds had low breakthrough, 2% or less, for nylon membranes. In contrast, when −N compounds dominated, the breakthrough was high (see excel spreadsheet in the Supporting Information). When organic-Hg is present, regardless of the presence of −N on the sample, the breakthrough percentage was low, indicating that organic-Hg compounds may enhance the sorption efficiency of the nylon membranes for −N compounds. Back trajectories showed that air flowed predominantly from the southeast over the Atlantic Ocean. Many trajectories were within the calculated marine boundary layer and then followed a path over land (Figure 4 MB). Four percent of the trajectory points were within 100 km of the site. IMPROVE data collected at this site (PRAAMS, MD08) showed that organic matter (hence organic compounds) and sulfate are the dominant compounds on the aerosols (data not shown), supporting our observations.

The MD nylon membrane RM concentrations were 31 to 85% lower than CEM RM concentrations for the entire campaign. The low efficiency of the nylon membranes relative to CEM is thought to be attributed to the predominance of −N compounds at MD. No MD samples were analyzed by IC.

**UT.** Mean RM concentration of all CEM collected at UT was 38 ± 4 pg m⁻³. Nylon membranes had a mean RM concentration of 19 ± 2 pg m⁻³ (Figure 1d). Mean breakthrough percentages for CEM and nylon membranes were 19 ± 6 and 9 ± 4%, respectively.

UT sample 5a had a profile dominated by −Br/Cl compounds with a small peak at high temperature in the organic-Hg range (Figure 2 UT a). The airflow patterns were from the west. The total residence times for trajectory points during sample deployment was 4% for the Northern Eurasian source box, 1% for the East Asian source box, and 2% for the San Francisco source box. Greater than 25% of trajectory points traveled below the marine boundary layer over the East Asian source box (Figure 4a). In this case, 7% of the trajectory points were within 100 km of the site.

Trajectories for sample UT b showed that air passed over the Asian continent, Alaska, Canada, and the Pacific Ocean (Figure 4). Ten percent of total trajectory points resided in the Northern Eurasian source box, and 1% of points resided in the East Asian source box. Some trajectories resided beneath the boundary layer over East Asia, with a heightened density of trajectories (yellow, 5–25%), and 2% of the trajectory points were within 100 km of the site. For additional trajectories with ion chemistry, see Figure SI 9, and for all trajectories, Figure SI 10. Peak deconvolution proportions were 28% −O, 31% −Br/Cl, 39% −N, and 3% organic-Hg compounds. Ninety-two percent of ions detected by IC were SO₄²⁻, and 8% were NO₃⁻ (for additional ion concentrations, see Table SI 4). The presence of SO₄²⁻ on the membrane does not correspond with peak deconvolution results, indicating that (1) −S compounds were not participating in the oxidation of GEM, (2) the compounds degraded in air or on the membrane, or (3) membranes were sorbing other S gases/particles that do not match our standard profile. Based on evidence provided by peak deconvolution and back trajectories, RM chemistry was dominantly influenced by long-range transport from Asia. Long-range transport of Hg from Eurasia and Asia has been observed in other studies for the Western United States. In addition, elevated SO₄²⁻ was observed on IMPROVE network filters collected at Great Basin National Park associated with long-range transport from Eurasia and elevated O₃ events.

Thermal desorption profiles indicated that the primary compounds were −Br/Cl- and −N-based in winter 2017; −O, −Br/Cl, and −N compounds in spring 2018; −Br/Cl and −O in summer 2018; −O, −Br/Cl, −N, and −S in fall 2018; and −O, −Br/Cl, −N, and −S compounds in winter 2018 (see profiles in the Supporting Information). It is important to note that in winter 2017, there were no multiday inversion episodes. More data and discussion will be presented in a future publication (Lyman et al., in prep).

**MAJOR IMPLICATIONS**

UNR-RMAS 2.0 CEM RM concentrations were higher than Tekran RM concentrations in all cases, except for four 2-week periods given the error bars at MLO. Tekran data exhibited similar temporal trends as CEM measurements, and in general, were not well correlated (Table SI 4). RM compounds were not collected as efficiently by the Tekran, and this varied by location. This is consistent with past studies comparing CEM and Tekran RM concentrations. At the GH and MD sites that had more diverse thermal desorption profiles, the temporal trends between Tekran and membrane measurements did not follow similar patterns and concentrations were low (<6 pg m⁻³ on average for each deployment period for Tekran measurements). This reflects the lower efficiency of the KCl-coated denuder to collect −S, −N, and organic-Hg RM compounds. It is noteworthy that the HgCl₂ was the form that was used for the development of the KCI denuder and good recovery rates were achieved with this compound. Thus, during the good agreement periods at MLO, it is likely that this reflects input from the marine boundary layer. It is noteworthy that this air would be humid, and this indicates that the membranes are not specifically affected by RH. In addition, Gustin et al. found that absolute humidity had no influence on CEM concentrations.

All sites had lower RM concentrations on the nylon membranes than the CEM (Figure 1), except for GH during the summer of 2018 that were dominated by −Br/Cl-containing compounds. Thus, the CEM were more efficient at collecting total RM than nylon membranes, but the ratio of nylon membrane RM concentrations to CEM concentrations varied for different RM compounds due to RM chemistry.

In order from lowest to highest, breakthrough was MLO (CEM, 5 ± 1%; nylon, 1.2 ± 0.3%) < GH (CEM, 13 ± 1; nylon, 7 ± 2%) < MD (CEM, 15 ± 7%; nylon, 8 ± 3%) < UT (CEM, 19 ± 6%; nylon, 9 ± 4%). For the nylon membranes, this is attributed to the presence of −N compounds. This may also be true for the other locations for the CEM, or other conditions could be influencing these membranes. Further work is needed to better understand the limitations of the membranes under a variety of conditions. For example, whether reactions are occurring on the membranes needs to be better understood. However, because CEM concentrations have been found to be similar to a calibrated dual channel system (Lyman et al.), the CEM is more accurate than the Tekran system. In addition, because membranes did not significantly lose ambient RM and thermal desorption profiles remained the same after holding for 6 months (Dunham-Cheatham et al.), it suggests that membranes retain RM compounds. Limited work has demonstrated that the CEM retains HgCl₂ and HgBr₂ after being spiked and deployed.

Deconvolution results from nylon membranes provide a means of identifying the dominant RM compounds, as well as
mixtures. This is important information for future studies on developing methods that quantify RM chemistry.

IC results in some cases agreed with the compounds determined based on thermal desorption profiles. However, at some locations, SO$_4^{2-}$ was measured using IC and there were no S-based RM compounds detected in the thermal desorption profiles. This suggests that reactions were not occurring on the membranes forming S compounds. The least common ion in the samples was Br$^-$, which made up only 1% of the total ions when detected. It is possible that if there were bromine ions on the nylon membranes, they were not in the Br$^-$ form.

As noted, the exact compounds thermally desorbed from the nylon membranes are unknown. However, comparing thermal desorption behavior with that of pure standard compounds is a significant step forward in discerning RM chemistry.

This work demonstrated that the UNR-RMAS 2.0 provides a more quantitative measure of RM concentrations than the Tekran system, and a step toward understanding of RM chemistry. The UNR-RMAS 2.0 membrane data paired with peak deconvolution, ion chromatography, back trajectories, and site air chemistry relationships revealed important information for understanding RM concentrations and chemistry across space and time.

Deployment of the RMAS 2.0 at locations with a Tekran 2537/1130/1135 system would provide a framework for better understanding of the discrepancy between more accurate RM concentrations versus those measured by the Tekran. However, given that the chemistry of RM compounds depends on the oxidants in air and some RM compounds are more efficiently measured by the Tekran (halogenated), correction of past Tekran data will be a rough estimation.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.est.0c02283.

Details on the membranes, deployment, percent breakthrough calculations, and thermal desorption (PDF)

Data for all deployments (XLSX)

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Notes

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