The use of Pb, Sr, and Hg isotopes in Great Lakes precipitation as a tool for pollution source attribution

Laura S. Sherman^{a*}, Joel D. Blum^a, J. Timothy Dvonch^b, Lynne E. Gratz^c, and Matthew S.

Landis^d

a. University of Michigan, Department of Earth and Environmental Sciences, 1100 N. University

Ave., Ann Arbor, MI 48109, U.S.A.

b. University of Michigan, Air Quality Laboratory, 1415 Washington Heights, Ann Arbor, MI

48109, U.S.A.

c. University of Washington-Bothell, 18115 Campus Way NE, Bothell, WA 98011, U.S.A.

d. U.S. EPA, Office of Research and Development, Research Triangle Park, NC 27709, U.S.A.

*Corresponding author. Phone: 607-592-3026. Email: lsaylors@umich.edu

Abstract

The anthropogenic emission and subsequent deposition of heavy metals including mercury (Hg) and lead (Pb) presents human health and environmental concerns. Although it is known that local and regional sources of these metals contribute to deposition in the Great Lakes region, it is difficult to trace emissions from point sources to impacted sites. Recent studies suggest that metal isotope ratios may be useful for distinguishing between and tracing source emissions. We measured Pb, strontium (Sr), and Hg isotope ratios in daily precipitation samples that were collected at seven sites across the Great Lakes region between 2003 and 2007. Lead isotope ratios (${}^{207}Pb/{}^{206}Pb = 0.8062$ to 0.8554) suggest that Pb deposition was influenced by coal combustion and processing of Mississippi Valley-Type Pb ore deposits. Regional differences in Sr isotope ratios (87 Sr/ 86 Sr = 0.70859 to 0.71155) are likely related to coal fly ash and soil dust. Mercury isotope ratios (δ^{202} Hg = -1.13 to 0.13‰) also varied among the sites, likely due to regional differences in coal isotopic composition and fractionation occurring within industrial facilities and in the atmosphere. These data represent the first combined characterization of Pb, Sr, and Hg isotope ratios in precipitation collected across the Great Lakes region. We demonstrate the utility of multiple metal isotope ratios in parallel with traditional trace element multivariate statistical modeling to enable more complete pollution source attribution.

Keywords

heavy metals, stable isotopes, source attribution, Great Lakes, wet deposition

Abbreviations

Mississippi Valley-Type (MVT), National Institute of Standards and Technology (NIST), Standard Reference Material (SRM), mass-dependent fractionation (MDF), mass-independent fractionation (MIF), gaseous elemental mercury (GEM), gaseous oxidized mercury (GOM), particle bound mercury (PBM), University of Michigan Air Quality Laboratory (UMAQL), inductively coupled plasma high resolution mass spectrometry (ICP-HRMS), Positive Matrix Factorization (PMF), Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT), multi-collector inductively coupled plasma mass spectrometry (MC-ICP-MS)

1. Introduction

The emission and deposition of hazardous trace metals such as mercury (Hg) and lead (Pb) are of significant concern because they can affect human and environmental health (Clarkson et al., 2007; Evers et al., 2008; Laraque and Trasande, 2005). These metals are potent neurotoxins and are especially harmful to developing fetuses and young children (Clarkson et al., 2007; Laraque and Trasande, 2005). In fact, due to high fish methylmercury concentrations, all water bodies in the Great Lakes region currently have fish consumption advisories (U.S. EPA, 2009). Significant research has been devoted to understanding the sources, transport, deposition, and impacts of these metals in the region (Flegal et al., 1989; Hall et al., 2005; Keeler and Dvonch, 2005; Landis and Keeler, 2002; Landis et al., 2002; Risch et al., 2012).

Atmospheric deposition is a primary source of metals such as Hg to ecosystems in the Great Lakes region (Fitzgerald et al., 1991; Landis and Keeler, 2002). Research over the last two decades has shown that local and regional sources (especially in the Chicago, Illinois and Detroit, Michigan urban and industrial areas) contribute significantly to this deposition (Flegal et

al., 1989; Gratz et al., 2013a; Keeler and Dvonch, 2005; Landis et al., 2002). Mercury is emitted to the atmosphere in the U.S. primarily by coal combustion, metals processing, waste incineration, and chemical and cement production (Pacyna et al., 2006; Pirrone et al., 2010).
Similarly, major U.S. anthropogenic sources of Pb are coal and oil combustion, metals mining and processing, battery recycling, and oil refining (Graney and Landis, 2013; Komárek et al., 2008; Nriagu and Pacyna, 1998). Previous studies have estimated source contributions using a variety of methods including measurement of metals in precipitation (Glass and Sorensen, 1999; Gratz et al., 2009; Keeler et al., 2005; Keeler et al., 2006; White et al., 2009), meteorological analyses (Cohen et al., 2004; Cohen et al., 2007; Dvonch et al., 2005; Landis and Keeler, 2002), and multivariate statistical receptor modeling (Gratz and Keeler, 2011; Gratz et al., 2013b; Keeler et al., 2006). However, due to complex atmospheric chemistry, transport patterns, meteorological conditions, the mixing of emissions from multiple sources, and uncertainties associated with receptor modeling, it remains difficult to quantify the relative impact of emissions from different sources on metal deposition.

Radiogenic isotope ratios of metals such as Pb and strontium (Sr) are not measurably modified by industrial, atmospheric, or environmental processes and can be used to characterize emissions from different sources (Geagea et al., 2007; Graney and Landis, 2013; Grousset and Biscaye, 2005; Simonetti et al., 2000a; Simonetti et al., 2000b). For several decades, Pb isotope ratios measured in ice cores, snow, weekly-integrated precipitation, aerosols, soils, and lichens have been used to distinguish between deposition resulting from leaded gasoline combustion, metal smelters, coal-fired power plants, and waste incinerators (DesJardins et al., 2004; Erel et al., 1997; Geagea et al., 2007; Haack et al., 2002; Kaste et al., 2003; Rabinowitz and Wetherill, 1972; Simonetti et al., 2000b; Simonetti et al., 2000c; Sturges and Barrie, 1987; Sturges and Barrie, 1989; Sturges et al., 1993). However, source-receptor relationships can be determined most accurately by analyzing short-term daily samples (Dvonch et al., 1999; Graney and Landis, 2013; Gratz et al., 2010; Hoyer et al., 1995) or sub-daily event-based samples (White et al., 2009; White et al., 2013). A recent study in the Great Lakes region used Pb isotope ratios measured in daily precipitation samples in combination with air mass back trajectory analyses and trace element concentrations to distinguish between anthropogenic Pb emission sources (Graney and Landis, 2013). In some cases, Sr isotope ratios have been used in conjunction with Pb isotope ratios to further constrain metal sources (Geagea et al., 2007; Grousset and Biscaye, 2005; Simonetti et al., 2000a; Simonetti et al., 2000b). In addition, recent methodological developments now allow the precise measurement of Hg isotopes in precipitation samples and Hg isotope ratios may be used to distinguish between emissions sources (Chen et al., 2012; Gratz et al., 2010; Sherman et al., 2012; Sun et al., 2013).

The combination of multiple isotopic systems may ultimately improve source apportionment capabilities as well as allow characterization of the isotopic composition of atmospheric end-members for ecological studies of base cation (e.g., calcium, Ca) and metal (e.g., Hg and Pb) cycling in sensitive ecosystems (Bailey et al., 1996; Blum et al., 2002; Demers et al., 2013; Erel et al., 1997; Kaste et al., 2003; Miller et al., 1993). Very little data exists with which to evaluate the metal isotopic variability of either atmospheric sources or deposition in the Midwestern U.S. For example, although several previous studies have used Sr isotope ratios as a proxy tracer for Ca transport through forest ecosystems, these studies relied on a limited number of Sr isotope ratio measurements in precipitation (e.g., Bailey et al., 1996; Blum and Erel, 1997; Blum et al., 2002; Miller et al., 1993). We conducted this study both to characterize the values and variability of Pb, Sr, and Hg isotope ratios in precipitation in the Great Lakes region and to evaluate the potential application of multiple isotope systems to improve metal source attribution estimates.

1.1. Lead, strontium, and mercury isotopes

There are four stable isotopes of Pb with the following approximate abundances: ²⁰⁸Pb (52.4%), ²⁰⁷Pb (24.1%), ²⁰⁶Pb (22.1%), and ²⁰⁴Pb (1.4%). ²⁰⁴Pb is non-radiogenic and was not measured in this study because of its low relative abundance. ²⁰⁸Pb, ²⁰⁷Pb, and ²⁰⁶Pb are formed by the radioactive decay of 232 Th (half-life = 14 billion years), 235 U (half-life = 0.7 billion years), and 238 U (half-life = 4.5 billion years), respectively. The ratios of these isotopes vary in geologic materials due to differences in the relative concentrations of the parent and daughter isotopes and the age of the rock or ore body. In the U.S., the primary Pb producing ores are Mississippi Valley-Type (MVT) lead-zinc-copper ores, which have distinctly lower ²⁰⁷Pb/²⁰⁶Pb ratios than most other ores (MVT 207 Pb/ 206 Pb = 0.714 to 0.781; Doe and Delevaux, 1972; Goldhaber et al., 1995; Heyl et al., 1974; Komárek et al., 2008). These distinct Pb isotope ratios have been used to identify MVT ore-derived Pb in U.S. leaded gasoline and metallurgical emissions (Bollhöfer and Rosman, 2001; Graney et al., 1995; Graney and Landis, 2013; Komárek et al., 2008; Simonetti et al., 2000c; Sturges and Barrie, 1989). After the phase-out of leaded gasoline in the U.S. in the 1980s and 1990s and decreased primary Pb ore production (Herrick and Friedland, 1990; Shen and Boyle, 1987; Veron et al., 1993), the major U.S. sources of Pb to the atmosphere became more diverse and now include detectable emissions from facilities that recycle scrap metal from multiple sources (Rabinowitz, 2005; Sturges and Barrie, 1987).

Strontium is present in all Ca-bearing geologic materials including rocks, soils, and coals. There are four stable isotopes of Sr with the following approximate abundances: ⁸⁸Sr (82.6%), ⁸⁷Sr (7.0%), ⁸⁶Sr (9.9%), and ⁸⁴Sr (0.6%). Of these, only ⁸⁷Sr is radiogenic and forms by the radioactive decay of ⁸⁷Rb (half-life = 48.8 billion years). ⁸⁷Sr/⁸⁶Sr ratios are distinct in rocks of different ages and lithologies, as well as in soils, vegetation, and surface waters (Blum and Erel, 2003; Burke et al., 1982). Strontium is also present in coal and is concentrated in fly ash after combustion (Hurst and Davis, 1981; Hurst et al., 1991). Previous studies have used Sr isotopes to trace sources of airborne dust to soils and lake sediments, as well as fly ash leachates into ecosystems (Hurst et al., 1991; Hurst et al., 1993; Miller et al., 1993; Simonetti et al., 2000c; Spivak-Birndorf et al., 2012; Stancin et al., 2006; Straughan et al., 1981). However, only a few studies have measured Sr isotope ratios in precipitation samples (e.g., Bailey et al., 1996; Blum and Erel, 1997; Bullen et al., 1996; Miller et al., 1993; Simonetti et al., 2000b; Simonetti et al., 2000c).

In contrast to Pb and Sr, Hg isotopes are non-radiogenic and isotopic variability is caused by fractionation during biogeochemical processes. There are seven stable isotopes of Hg with the following approximate abundances: ²⁰⁴Hg (6.8%), ²⁰²Hg (29.7%), ²⁰¹Hg (13.2%), ²⁰⁰Hg (23.1%), ¹⁹⁹Hg (16.9%), ¹⁹⁸Hg (10.0%), and ¹⁹⁶Hg (0.2%). Mercury isotope ratios are reported using delta notation (Equation 1).

$$\delta^{\text{xxx}}\text{Hg}(\%) = ([(^{\text{xxx}}\text{Hg}/^{198}\text{Hg})_{\text{sample}}/(^{\text{xxx}}\text{Hg}/^{198}\text{Hg})_{\text{SRM 3133}}] - 1) * 1000$$
(1)

In Equation 1, ^{xxx}Hg is an isotope of Hg and SRM 3133 is the National Institute of Standards and Technology (NIST) Standard Reference Material (SRM) (Bergquist and Blum, 2007). Mercury isotopes can undergo mass-dependent fractionation (MDF) during a number of biotic and abiotic processes (Bergquist and Blum, 2007; Estrade et al., 2009; Kritee et al., 2009; Kritee et al., 2007; Rodríguez-González et al., 2009; Zheng et al., 2007) and mass-independent fractionation (MIF) primarily during photochemical reactions (Bergquist and Blum, 2007; Buchachenko, 2001; Zheng and Hintelmann, 2010). Mass-independent fractionation is reported as the deviation of a measured isotope ratio from the theoretically predicted ratio based on the MDF law (Equation 2).

$$\Delta^{\text{xxx}}\text{Hg} = \delta^{\text{xxx}}\text{Hg} - (\delta^{202}\text{Hg} * \beta)$$
⁽²⁾

In Equation 2, β for kinetic reactions is equal to 0.252 for ¹⁹⁹Hg, 0.502 for ²⁰⁰Hg, 0.752 for ²⁰¹Hg, and 1.493 for ²⁰⁴Hg (Bergquist and Blum, 2007).

It is especially difficult to quantify Hg source-receptor relationships because emitted Hg can be transported globally as a reduced elemental gas (GEM) or deposited near sources as gaseous oxidized Hg (GOM) and particle bound Hg (PBM) (Lin and Pehkonen, 1999; Schroeder and Munthe, 1998). Recent studies have begun to characterize Hg isotope ratios in source materials such as coals and other studies have begun to trace atmospheric Hg emissions using Hg isotopes (Biswas et al., 2008; Chen et al., 2012; Gratz et al., 2010; Lefticariu et al., 2011; Rolison et al., 2013; Sherman et al., 2012; Sun et al., 2013; Yin et al., 2014; Yin et al., 2013). Coals from North America, China, and Russia generally display negative δ^{202} Hg values (mean = -1.21%, 1SD = 0.64‰, n = 89) and slightly negative Δ^{199} Hg values (mean = -0.09%, 1SD = 0.15‰, n = 89) (Biswas et al., 2008; Lefticariu et al., 2011; Sherman et al., 2012; Sun et al., 2013; Yin et al., 2014). In addition to isotopic differences among combusted coals, fractionation of Hg isotopes occurs within coal-fired power plants. Sherman et al. (2012) observed extreme negative δ^{202} Hg values (<-3.0‰) in rainfall collected near a large coal-fired power plant in Florida. Subsequently, Sun et al. (2013; 2014) modeled Hg fractionation in two Chinese coalfired power plants and found that the light isotopes of Hg were preferentially oxidized and absorbed onto fly ash whereas the heavy isotopes of Hg were preferentially emitted as GEM. It is therefore possible that near-field deposition of the oxidized Hg species (i.e., PBM and GOM) resulted in the negative δ^{202} Hg values observed in precipitation near the Florida power plant

(Sherman et al., 2012). Although more research is needed to fully understand Hg isotope fractionation within industrial sources, these studies suggest that Hg isotope ratios may be used to identify emissions from sources such as coal-fired power plants.

2. Methods

2.1. Sample collection

Daily precipitation samples used in this study were collected at five sites in Michigan, one rural site in Vermont, and one site impacted by industrial emissions in Ohio between July 2003 and January 2007 (Figure 1). The precipitation samples analyzed during this study represent a subset of the total collected samples. As shown in Figure 1, these sites were located in proximity to a variety of local and regional emission sources including coal-fired power production, iron and steel production, non-ferrous metals processing, municipal waste and sewage incineration, and cement production. The precipitation collection sites in Michigan were operated by the University of Michigan Air Quality Laboratory (UMAQL) as part of a study to assess sources of Hg deposition (UMAQL, 2009). These sites included two southern Michigan urban locations: Grand Rapids (GRD; 42.9600N, 85.1491W) and southwest Detroit (FRT; 42.3030N, 83.1054W); one southern Michigan rural location: Dexter (DXT; 42.4167N, 83.9019W); and two northern Michigan rural locations: Eagle Harbor (EGL; 47.4596N, 88.1491W) and Pellston (PLN; 45.4487N, 84.6776W). The rural site at Underhill, Vermont (LCH; 44.5283N, 72.8684W) was located ~25 km east of Lake Champlain and was operated by UMAQL from 1992 to 2007 (Gratz et al., 2009; Keeler et al., 2005). The site impacted by industry at Steubenville, Ohio (STB; 40.3794N, 80.6195W) was operated through a cooperative agreement between the U.S. Environmental Protection Agency (U.S. EPA) and UMAQL from

2002 to 2007 (Keeler et al., 2006; White et al., 2009; White et al., 2013). This site was located in close proximity to numerous emission sources including five large coal-fired power plants within a 50 km radius, iron/steel and non-ferrous metal production facilities, slag processing, and incinerators (Keeler et al., 2006; Vedantham et al., 2014; White et al., 2009). Previous studies have identified elevated Hg deposition at this site (Keeler et al., 2006; White et al., 2009) as well as high concentrations of other hazardous air pollutants, including particulate matter (Dockery et al., 1993; Vedantham et al., 2014).

Daily wet-only precipitation samples were collected as described previously using modified wet-only automatic collectors (MIC-B; Landis and Keeler, 1997). Each collector had separate acid-cleaned glass/Teflon sampling trains for Hg samples and polypropylene sampling trains for trace element samples (Gratz and Keeler, 2011; Gratz et al., 2009; Keeler et al., 2006; Landis and Keeler, 2002). After collection, trace element samples were acidified with trace metal-grade HNO₃ to a concentration of 0.2% (v/v) and metal concentrations were measured by inductively coupled plasma high resolution mass spectrometry (ICP-HRMS; Thermo Scientific Element2) as reported previously (UMAQL, 2009; Gratz and Keeler, 2011; Keeler et al., 2006; White, 2009). Previous research has demonstrated that metals, including Pb, are stable in acidified precipitation samples stored in polypropylene bottles for at least ten years (Graney and Landis, 2013). After collection, Hg samples were oxidized with Hg-free BrCl to a concentration of 1% (v/v) as reported previously (UMAQL, 2009; Gratz and Keeler, 2011; Keeler et al., 2006; White et al., 2009). The original Teflon sampling bottles were tightly closed, sealed with Teflon tape, double bagged, and stored in a dark cold-room. We expect that oxidized precipitation samples stored under these conditions should be stable and fully retain Hg for many years (Parker and Bloom, 2005).

A subset of samples was chosen for isotopic analysis based on sample availability, adequate Hg mass, and collection date to provide maximal seasonal coverage. Because emissions from nearby sources may result in higher localized metal deposition (White et al., 2013), this data set may be biased toward the identification of local and regional emissions sources. In total, 32 precipitation samples were analyzed for Pb, Sr, and Hg isotope ratios (Table 1). Due to sample availability and/or low metal concentrations, 26 other samples were analyzed for only one or two of the three isotope systems investigated (Table 1).

2.2. Previous analyses

Positive Matrix Factorization (PMF) is an advanced mathematical receptor modeling tool implemented by the U.S. EPA that utilizes a matrix of trace element concentrations to retrospectively identify contributing atmospheric pollution sources by the presence of unique elemental tracers that have a significant influence on the modeled data set (U.S. EPA, 2014; Paatero, 1997). Mesoscale meteorological air mass back trajectories calculated using the NOAA Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model (Draxler and Rolph, 2010) can be paired with these quantitative source estimates. Using PMF, previous studies identified up to six sources that impacted metal deposition at the Great Lakes collection sites (UMAQL, 2009; Gratz and Keeler, 2011; Keeler et al., 2006; White et al., 2009). According to these analyses, Hg deposition at these sites is largely attributable to coal combustion (61–83% of total accounted for Hg deposition) whereas Pb deposition) and secondarily to coal combustion (<5%–40% of total accounted for Pb deposition) (UMAQL, 2009; Gratz and Keeler, 2011; Keeler et al., 2006; White et al., 2009; Gratz and Keeler, 2011; Keeler et al., 2006; White et al., 2009; Gratz and Keeler, 2011; Keeler et al., 2006; White et al., 2009; Gratz and Keeler, 2011; Keeler et al., 2006; White et al., 2009; Gratz and Keeler, 2011; Keeler et al., 2006; White et al., 2009; Gratz and Keeler, 2011; Keeler et al., 2006; White et al., 2009; Gratz and Keeler, 2011; Keeler et al., 2006; White et al., 2009; Gratz and Keeler, 2011; Keeler et al., 2006; White et al., 2009; Gratz and Keeler, 2011; Keeler et al., 2006; White et al., 2009; Gratz and Keeler, 2011; Keeler et al., 2006; White et al., 2009; Gratz and Keeler, 2011; Keeler et al., 2006; White et al., 2009; Gratz and Keeler, 2011; Keeler et al., 2006; White et al., 2009; Gratz and Keeler, 2011; Keeler et al., 2006; White et al., 2009; Gratz and Keeler, 2011; Keeler et al., 2006; White et al., 2009; Gratz and Keeler, 2011; Keeler et al., 2006; White et al.,

Underhill, Vermont, are impacted by regional Hg emissions from coal combustion and other sources that are transported from Midwestern urban and industrial centers such as Chicago, Detroit, and the Ohio River Valley (Gratz and Keeler, 2011; Liu et al., 2010). In contrast, Hg wet deposition at the Steubenville, Ohio site is dominantly impacted by local coal combustion and metals processing (Keeler et al., 2006; White et al., 2009; White et al., 2013).

2.3. Isotopic analyses: Lead

The selected precipitation samples were measured for Pb isotope ratios without preconcentration using ICP-HRMS following previously described methods (Graney and Landis, 2013). Lead isotopic standard NBS 981 was used to correct for internal mass bias using standard-sample bracketing with Pb concentrations matched to within 10%. All samples with Pb concentrations $>1 \mu g/L$ were diluted to $1 \mu g/L$. Two secondary standards, BCR-2 and TMRAIN-95, and 1% HNO₃ blanks were analyzed with each block of five samples. BCR-2 is a rock standard that was hot acid digested (Stancin et al., 2006) and diluted in 1% HNO₃ to concentrations ranging from 0.1 to 1 µg/L. TMRAIN-95 is a synthetic rainfall standard that was analyzed at 0.3 µg/L throughout the analytical session. Measured Pb isotope ratios for the BCR-2 standard across all sample concentrations were within error of reported values (this study: mean ${}^{207}Pb/{}^{206}Pb = 0.8333$, 2SD = 0.0042, n = 19, mean ${}^{208}Pb/{}^{206}Pb = 2.0658$, 2SD = 0.0065, n = 19; Weis et al., 2006: mean ${}^{207}Pb/{}^{206}Pb = 0.8332$, 2SD = 0.0009, n = 11, mean ${}^{208}Pb/{}^{206}Pb =$ 2.0649, 2SD = 0.0004, n = 11). Measured analytical uncertainties for the TMRAIN-95 standard were similar to those for BCR-2 (mean ${}^{207}Pb/{}^{206}Pb = 0.8138$, 2SD = 0.0040, n = 19; mean 208 Pb/ 206 Pb = 2.0032, 2SD = 0.0097, n = 19). Lead isotope ratios measured in sample replicates (n = 11) at all concentrations were the same within the reported uncertainties (see Appendix A).

Blank intensities remained constant throughout the analytical session and represented an average of $1.1 \pm 0.9\%$ (1SD, range = 0.2–6.7%) of the ²⁰⁸Pb sample signal intensity. Blank correction did not significantly change the Pb isotope ratios and the reported data were not blank corrected.

2.4. Isotopic analyses: Strontium

Splits of the selected precipitation samples (15–240 mL, 20–230 ng of Sr) were filtered to 0.45 μ m using acid-cleaned filters (n = 50, see Appendix B). Where possible, enough sample was filtered to obtain >50 ng of Sr. The samples were then evaporated to dryness on a hot plate at 80 °C and reconstituted in 0.5 mL of ultrapure 3 N HNO₃. Strontium was separated from the matrix by cation exchange chromatography using columns packed with Sr-specific resin (Eichrom Sr-Spec) as described previously (Stancin et al., 2006). After the samples were loaded on the columns, matrix elements were eluted in 2.9 mL of 3 N HNO₃ and Sr was eluted in 2 mL of 0.05 N HNO₃. The samples were then evaporated to dryness and loaded onto tungsten filaments in 1 μ L of 1 N HCl. Strontium isotope standards (NBS 987) and de-ionized water blanks were processed in the same manner. Procedural blanks contained <1% of the average sample Sr mass and were thus negligible (mean Sr blank = 0.38 ng, 1SD = 0.09 ng, n = 4).

Strontium isotope ratios were measured by thermal ionization mass spectrometry (Finnigan MAT 262) according to previously published methods (Stancin et al., 2006). The ⁸⁷Sr/⁸⁶Sr ratios were corrected for internal mass bias to an ⁸⁶Sr/⁸⁸Sr ratio of 0.1194. During the analysis period for this study (October 2013–January 2014), the measured mean ⁸⁷Sr/⁸⁶Sr ratio for NBS 987 was 0.71022 \pm 0.000024 (2SD, n = 26).

2.5. Isotopic analyses: Mercury

Prior to isotopic analyses, Hg concentrations were measured in the precipitation samples using atomic absorption spectrometry (see Appendix B). Mercury in the precipitation samples was then separated and concentrated according to previously described methods (Demers et al., 2013; Sherman et al., 2012). Briefly, samples (266–977 mL in volume) were diluted to a volume of 1 L with 1% BrCl (v/v) and neutralized with 0.3 mL of 30% NH₂OH HCl (w/w). Ten percent SnCl₂ (w/w) was then slowly added to each sample through a three-port cap. The resulting GEM was pulled into acidic 1% KMnO₄ (w/w) solutions using a vacuum pump. Procedural standards (n = 6) and blanks (n = 1) were processed in the same manner. Mercury in procedural standards consistently had high recovery in the final solutions (mean = 98%, 1SD = 6%, n = 6). With the exception of one sample, all of the Hg in the samples also had high recovery after separation and concentration (84–120%, mean = 101%, 1SD = 8%, n = 40; Appendix C). For unknown reasons, the Hg in one sample (STB 141) had lower recovery (77%) and was, therefore, excluded from the data set.

Mercury isotopic compositions were measured in the concentrated sample and standard solutions using continuous-flow cold vapor generation multi-collector inductively coupled plasma mass spectrometry (MC-ICP-MS; Nu Instruments Nu plasma) according to previously published methods (Blum and Bergquist, 2007). The UM-Almadén secondary standard was analyzed at least five times in each analytical session. Maximum sample analytical uncertainty is estimated as the greater of the 2SD variability of a given isotope ratio measured in the procedural standards or in the UM-Almadén secondary standard. Analytical uncertainty was observed to increase at concentrations $<2 \ \mu g/L$. Because many of the samples in this study were analyzed at concentrations $<2 \ \mu g/L$ (0.9–1.9 $\mu g/L$, n = 26), uncertainties were calculated for UM-

Almadén secondary standards analyzed both at <2 μ g/L (0.8–1.9 μ g/L) and ≥2 μ g/L (2.0–5.0 μ g/L) (Appendix C).

2.6. Meteorological analyses

In addition to Pb, Sr and Hg isotope ratios and previous PMF receptor modeling, we explored the meteorology associated with each precipitation event. We utilized HYSPLIT back trajectories (Draxler and Rolph, 2010) that were made publically available online by the University of Michigan Department of Atmospheric, Ocean, and Space Science (University of Michigan, 2012). These trajectories were calculated every six hours starting at 500 m using EDAS40 data from the National Weather Service (40 km, 3 h resolution). Surface meteorological maps and atmospheric sounding data were used to characterize frontal systems, surface wind directions, and vertical stability (National Weather Service; University of Wyoming).

2.7. Statistical analyses

Statistical significance was assessed using SAS 9.4 statistical analysis software (SAS Institute Inc., 2013). Wilcoxon nonparametric tests were conducted using the NPAR1WAY procedure utilizing the exact option to calculate *p*-values based on linear rank statistics. These tests were used to compare 207 Pb/ 206 Pb ratios between samples collected at rural and urban/impacted sites, 87 Sr/ 86 Sr ratios between samples collected in Steubenville and Grand Rapids/Detroit and δ^{202} Hg values between samples collected in Steubenville and Grand Rapids.

3. Results and Discussion

3.1. Lead isotope ratios

Abbreviated isotopic results are presented in Table 1 and complete data are presented in Appendix C. Measured Pb isotope ratios in the precipitation samples are shown in Figure 2A–B along with previous measurements of North American coals (Chow and Earl, 1972; Díaz-Somoano et al., 2009), MVT ores (Doe and Delevaux, 1972; Goldhaber et al., 1995; Heyl et al., 1974), and recent precipitation samples from the Great Lakes region (i.e., post 1990) (Graney and Landis, 2013; Simonetti et al., 2000c). The precipitation samples collected during this study displayed similar Pb isotope ratios to those of previously analyzed North American precipitation samples and North American coals (Figure 2A–B). There were significant differences between precipitation samples collected at the rural sites (Figure 2A) and those collected at the urban/industrial sites (Figure 2B) (Z = 3.4992, p = 0.0005). Precipitation samples collected at the rural sites (DXT, EGL, LCH, and PLN) displayed less Pb isotopic variability and significantly higher 207 Pb/ 206 Pb ratios (Figure 2A; 207 Pb/ 206 Pb: 0.8336 to 0.8543, mean = 0.8445, 1SD = 0.0063, n = 18) than samples collected at the urban/industrial sites (FRT, GRD, and STB; Figure 2B; 207 Pb/ 206 Pb: 0.8062 to 0.8554, mean = 0.8345, 1SD = 0.0098, n = 33). These differences may be due to the variable influence of local Pb sources at the urban/industrial sites in contrast with more consistent, regionally-sourced Pb deposition at the rural sites (Graney and Landis, 2013). For example, in some cases, the lower ²⁰⁷Pb/²⁰⁶Pb ratios measured in precipitation collected at the urban/industrial sites may be due to the influence of emissions from MVT ore processing (see section 3.4.1).

3.2. Strontium isotope ratios

Precipitation samples analyzed during this study were characterized by ⁸⁷Sr/⁸⁶Sr ratios between 0.70859 and 0.71155 (Table 1, Figure 3). These values are generally higher than those measured previously in precipitation collected in Montreal, Canada (Simonetti et al., 2000c), but similar to those measured in precipitation collected in the northeastern U.S. (Bailey et al., 1996; Blum et al., 2002; Miller et al., 1993). As shown in Figure 3, there is a significant division in Sr isotopic composition between samples collected in Steubenville, Ohio and those collected in Grand Rapids, Michigan and Detroit, Michigan (Z = 4.2267, p < 0.0001). The Steubenville samples display significantly higher ⁸⁷Sr/⁸⁶Sr ratios (mean = 0.71042, 1SD = 0.00054, n = 16) than those collected at the urban sites in Michigan (GRD: mean = 0.70945, 1SD = 0.00055, n = 14; FRT: mean = 0.70879, 1SD = 0.00016, n = 4). There are at least two possible contributing explanations for these differences: (i) suspended soil dust (and possibly sea-spray aerosol) impacting these sites displays regionally variable ⁸⁷Sr/⁸⁶Sr ratios or (ii) coal combustion in each region produces fly ash that has differing ⁸⁷Sr/⁸⁶Sr ratios.

During the last glacial maximum, all of the precipitation collection sites except for Steubenville were glaciated. As a result, soils in the Grand Rapids and Detroit areas are relatively young and derived from glacial sediments, which are a mixture of Canadian Shield and Michigan Basin materials (Dreimanis and Goldthwait, 1973). In contrast, Steubenville is located ~50 km south of the furthest extent of glaciation (Hedman et al., 2009). Soils in the Steubenville area are more deeply weathered and derived from the local underlying bedrock, which consists of Pennsylvanian through Permian shales, siltstones, sandstones, and limestones (Brockman, 1998). It is possible that the older, bedrock-derived soils in the Steubenville area display different ⁸⁷Sr/⁸⁶Sr ratios than the younger glacially-derived soils in the Grand Rapids and Detroit areas. However, if soil isotopic composition was the primary factor controlling the observed differences in precipitation ⁸⁷Sr/⁸⁶Sr ratios, we might expect to observe similar ⁸⁷Sr/⁸⁶Sr ratios across all of the glaciated sites in Michigan. Instead, precipitation collected in Pellston and Eagle Harbor displays ⁸⁷Sr/⁸⁶Sr ratios that are more similar to those observed in Steubenville, while precipitation collected in Dexter displays seasonally variable ⁸⁷Sr/⁸⁶Sr ratios (see Appendix B). Similarly, the lack of correlation between ⁸⁷Sr/⁸⁶Sr ratios in precipitation with collection site distance from the Atlantic Coast suggests that the abundance of sea-spray aerosol (modern ocean ⁸⁷Sr/⁸⁶Sr = 0.7092; DePaolo and Ingram, 1985; Hodell et al., 1990; McArthur et al., 2001; McArthur et al., 2006) is not a controlling factor of measured ⁸⁷Sr/⁸⁶Sr ratios.

We suggest that it is likely that fly ash emitted by coal-fired power plants and locallydeposited in Michigan and Ohio exhibits differing ⁸⁷Sr/⁸⁶Sr ratios. Strontium is present in coals in relatively low concentrations (Swaine, 1990) but is concentrated as a component in fineparticulate fly ash, reaching concentrations of up to 3,000 µg/g (Hurst and Davis, 1981; Hurst et al., 1991; Straughan et al., 1981). Most coal-fired power plants in the U.S., including those located near Grand Rapids and Steubenville, use electrostatic precipitators to remove particulates from the flue gas stream (Brown et al., 1999). Although electrostatic precipitators remove ~80% of the fine particulate matter in the flue gas, U.S. power plants emitted an estimated ~68,000 megagrams of fine particulates in 2005 (U.S. EPA, 2008; Clack, 2012). From 2004 to 2005, coal-fired power plants in Ohio purchased Appalachian Basin bituminous coal primarily from Ohio (45%), West Virginia (24%), and Kentucky (21%) (U.S. EIA, 2004-2005). In contrast, during the same time period, coal-fired power plants in Michigan purchased mainly subbituminous Powder River Basin (PRB) coal from Wyoming (47%) and Montana (28%) (U.S. EIA, 2004-2005). Carboniferous Appalachian Basin coals from Ohio, West Virginia, and Kentucky formed in coastal swamps and deltas (~300 Ma) (Heckel, 1995). In contrast, the much

younger Paleogene PRB coals from Wyoming and Montana formed in brackish inland seas and swamps (~60 Ma) (Flores and Hanley, 1984). Due to their different depositional environments, composition, rank, and ages, coals from these two regions likely have differing Sr isotope ratios. However, only a limited number of studies have measured Sr isotope ratios in coals and fly ash (Hurst et al., 1991; Hurst et al., 1993; Spivak-Birndorf et al., 2012). A recent study found that fly ash from U.S. sub-bituminous coals displayed lower 87 Sr/ 86 Sr ratios (mean = 0.71121, 1SD = 0.00088, n = 5) than fly ash from U.S. bituminous coals (mean = 0.71309, 1SD = 0.00075, n = 3) (Spivak-Birndorf et al., 2012). These data do not necessarily represent the full range of Sr isotope ratios displayed by U.S. coals or coals combusted in Michigan and Ohio. However, these data suggest that ⁸⁷Sr/⁸⁶Sr ratios vary systematically in coals depending on their characteristics. It is therefore possible that sub-bituminous PRB coals combusted in Michigan may produce fly ash with lower ⁸⁷Sr/⁸⁶Sr ratios than the bituminous Appalachian Basin coals combusted in Ohio. If these isotopic differences are preserved in precipitation, they would be most easily observed at urban/industrial collection sites located near coal-fired power plants (i.e., Grand Rapids, Detroit, and Steubenville). The observed precipitation ⁸⁷Sr/⁸⁶Sr ratios in Grand Rapids, Detroit, and Steubenville may therefore be influenced by the isotopic composition of locally-emitted fly ash.

3.3. Mercury isotope ratios

Great Lakes precipitation samples were generally characterized by negative δ^{202} Hg values (-1.13 to 0.13‰; Figure 4), positive Δ^{199} Hg values (-0.20 to 0.99‰; Figure 4), slight positive Δ^{200} Hg values (mean = 0.20‰, 1SD = 0.08‰; see Appendix B), and negative Δ^{204} Hg values (mean = -0.36‰, 1SD = 0.16‰; see Appendix B). With the exception of a few samples

collected in Steubenville, the precipitation δ^{202} Hg values were similar to those observed previously in North American precipitation collected at sites impacted by mixed Hg emissions sources (Chen et al., 2012; Demers et al., 2013; Donovan et al., 2013; Gratz et al., 2010; Sherman et al., 2012). Similarly, with the exception of one sample from Detroit that displayed an extreme negative Δ^{199} Hg value (-0.87‰; Table 1 and Figure 4; see section 3.4.2), the observed Δ^{199} Hg values were similar to those previously measured in precipitation (Chen et al., 2012; Demers et al., 2013; Donovan et al., 2013; Gratz et al., 2010; Sherman et al., 2012). It has been hypothesized that positive Δ^{199} Hg values in precipitation result from the preferential retention of the odd-mass-number isotopes of Hg in water droplets during photochemical reduction and loss of Hg (Chen et al., 2012; Gratz et al., 2010). It is likely that the MIF measured in the Great Lakes precipitation samples occurred due to similar photochemical processes (see Appendix B).

Previous PMF estimates indicate that the majority of Hg deposition at all of the precipitation collection sites is attributable to coal combustion (UMAQL, 2009; Gratz and Keeler, 2011; Keeler et al., 2006; White et al., 2009). It is possible, therefore, that the δ^{202} Hg values observed in these samples are related to the isotopic composition of combusted coal and fractionation occurring within coal-fired power plants. In a similar manner to Sr isotopes, the precipitation δ^{202} Hg values are significantly different between samples collected in Grand Rapids, Michigan and Steubenville, Ohio (Z = 3.6834, *p* < 0.0001). Samples collected in Grand Rapids display higher, less variable δ^{202} Hg values (mean = -0.04‰, 1SD = 0.09‰, n = 9) than those collected in Steubenville (mean δ^{202} Hg = -0.47‰, 1SD = 0.29‰, n = 16). This may be related to differences between the isotopic composition of PRB coal combusted in Michigan and Appalachian Basin coal combusted in Ohio. However, because only a limited number of PRB

coals have been measured for Hg isotope ratios (Biswas et al., 2008), it is difficult to fully assess this possibility. It is also possible that Hg fractionation and removal within power plants as well as proximity to power plants results in the deposition of Hg with distinct δ^{202} Hg values in these two areas. As described above, Sun et al. (2013; 2014) modeled Hg fractionation within coalfired power plants and found that emitted oxidized Hg species display more negative δ^{202} Hg values than emitted GEM. The Steubenville site was surrounded by coal-fired power plants, with the closest facility located ~15 km away and five facilities within a 50 km radius (Keeler et al., 2006; White et al., 2009). It is therefore possible that the more negative δ^{202} Hg values observed in samples collected at this site were caused by enhanced local deposition of PBM and GOM that displayed negative δ^{202} Hg values. Similarly, it is possible that the more positive δ^{202} Hg values observed in samples collected in Grand Rapids could be due the oxidation and deposition of Hg that was emitted as GEM by more disparate coal-fired power plants. Although there are no published experimental studies of Hg isotope fractionation during atmospheric oxidation of GEM, it is possible that this process may cause additional MDF of Hg isotopes such that the resulting oxidized species display higher δ^{202} Hg values. Because our current understanding of Hg fractionation within power plants and in the atmosphere is limited, we cannot conclusively evaluate these possibilities.

3.4. Precipitation case studies

3.4.1. Steubenville, Ohio: February 9 & February 14, 2005 (sample STB 180)

Although nearly all of the precipitation samples analyzed during this study represent an individual precipitation event, due to operator error a few of the samples included integrated deposition from more than one discrete precipitating event. As a result, these multi-event

samples may have been impacted by a wider variety of sources. This may dilute source-specific isotope ratios or trace element signatures, making source attribution more difficult.

A multi-event sample collected in Steubenville (sample STB 180) incorporated rain from two precipitation events: a short event on February 9, 2005 (16:00–24:00, UTC) and continuous rain on February 14, 2005 (2/14 10:00–2/15 4:00, UTC). The event on February 14 deposited substantially more rainfall (~22 mm) than the prior event (~6 mm) and, therefore, was likely responsible for most of the metal deposition in this sample (White et al., 2009). Precipitation on February 14 was associated with a low-pressure system that moved across the Great Lakes and an accompanying cold front that passed over Ohio from west to east. Air mass transport to Steubenville in advance of the cold front was southerly. For 24 hours prior to the event, southerly surface winds were light (<2 m/s) but these intensified during the event (5–6 m/s). As a result of these conditions, it is likely that local emissions from the coal-fired power plants and metallurgical processing facilities to the south of the Steubenville site (Vedantham et al., 2014; White et al., 2009) impacted deposition during this event.

Despite containing rainfall from more than one event, sample STB 180 exhibited distinct Pb and Hg isotope ratios (Figures 2, 3 and 4, Table 1). This sample displayed the lowest observed Pb isotope ratios (207 Pb/ 206 Pb = 0.8088, 208 Pb/ 206 Pb = 1.9591) and the most negative measured δ^{202} Hg value (-1.13 ± 0.13‰, 2SD). In addition, PMF source attribution estimates for this sample were distinct in comparison to the other Steubenville samples (Figure 5A–B) (White, 2009). As shown in Figure 5, Pb deposition in Steubenville was generally dominated by emissions from coal combustion and incinerators/smelters while Hg deposition was dominated by coal combustion. In contrast, Pb and Hg deposition in sample STB 180 were impacted by coal combustion to a lesser degree relative to the other samples. Instead, Pb and Hg deposition

in this sample were strongly influenced by scrap metal processing (Figure 5). Using PMF, emissions from scrap metal processing near the Steubenville site have been identified as distinct from emissions related to metal smelters based on high concentrations of vanadium, chromium, and iron along with low zinc concentrations (White, 2009). The low Pb isotope ratios measured in sample STB 180 may indicate that these local scrap metal facilities processed metal that was originally mined as MVT Pb ores.

It is possible that Hg associated with these local scrap metal processing facilities displays more negative δ^{202} Hg values. This may be related either to the isotopic composition of the processed materials or fractionation occurring within the facilities. To date, no studies have measured Hg isotope fractionation within metallurgical facilities. However, Hg isotopes have been used to track Hg deposited to soils near metal smelters (Gray et al., 2013; Ma et al., 2013; Sonke et al., 2010). These soils generally display negative δ^{202} Hg values (mean = -0.99‰, 1 SD = 0.82‰, n = 104), suggesting that emissions from metals processing facilities may also display negative δ^{202} Hg values (Gray et al., 2013; Ma et al., 2013; Sonke et al., 2010).

It is also possible that enhanced deposition of PBM due to cold cloud processes could have resulted in the more negative δ^{202} Hg value observed in sample STB 180. Although precipitation during this event was composed of rain and not snow, cold cloud nucleation processes were likely dominant during February in Steubenville. During cold cloud processes, ice crystals can nucleate preferentially through heterogeneous processes involving crustal mineral particles (Baustian et al., 2012). Subsequent winter time precipitation results in significantly lower Hg deposition due to decreased within-cloud Hg scavenging (Burke et al., 1995; Hoyer et al., 1995). However, previous studies in the Great Lakes region have observed that winter ambient PBM concentrations are higher than in the summer and that ambient PBM

concentrations are correlated with PBM concentrations in precipitation (Burke et al., 1995). As a result, winter samples collected in Steubenville likely contain more PBM emitted by local coalfired power plants, which display more negative δ^{202} Hg values. However, in general, there is no correlation among the Steubenville samples between δ^{202} Hg values and collection season.

3.4.2. Detroit, Michigan: July 27, 2003 (sample FRT 23) and August 2, 2003 (sample FRT 24)

The Detroit, Michigan precipitation collection site was located in southwest Detroit (Figure 1). This site was heavily influenced by emissions from nearby heavy-duty truck traffic and industrial sources including coal-fired power plants, oil refineries, coke ovens, iron and steel production facilities, and a large sewage sludge incinerator (Keeler and Dvonch, 2005). Many of these facilities were located within 4 km of the collection site to the south and southwest (Liu et al., 2007).

Precipitation on July 27, 2003 (21:00–22:00, UTC; sample FRT 23) was the result of a low-pressure system over Ontario, Canada. HYSPLIT back trajectories suggest relatively rapid air mass transport to the site from the southwest. Following this event, a high-pressure system moved into the area, resulting in local stagnation. Afternoon concentrations of pollutants including ground-level ozone (60–90 ppb), SO₂ (40–70 ppb), and fine particulate matter (18–40 μ g/m³) were highly elevated for several days (U.S. EPA, AirData). In the late afternoon on August 1, 2003, a low-pressure system moved over Lake Michigan from the west, resulting in precipitation in the early morning on August 2, 2003 (4:00–6:00, UTC; sample FRT 24). As described below, it is likely that stagnation prior to this event resulted in the enhanced deposition of locally emitted pollutants at the Detroit site (sample FRT 24).

Lead isotope ratios in these Detroit samples were similar to those of samples collected in Grand Rapids, Michigan (Figure 3). According to previous PMF estimates, Pb collected in sample FRT 23 was primarily associated with emissions from iron and steel production whereas Pb in sample FRT 24 was associated with emissions from both coal combustion and iron and steel production (UMAQL, 2009). Although sample FRT 23 did exhibit slightly lower Pb isotope ratios than the other Detroit samples (207 Pb/ 206 Pb = 0.8266, 208 Pb/ 206 Pb = 2.0344; Table 1), these ratios are not as low as those observed in some of the Steubenville samples or MVT Pb ores (Figure 2). It is likely that increased recycling of metals from multiple sources makes it more difficult to identify deposition resulting from iron and steel production using Pb isotope ratios.

Mercury isotope ratios measured in the Detroit precipitation samples were highly variable (Figure 4). As shown in Figure 6A, in addition to coal combustion, according to PMF estimates, Hg collected in sample FRT 23 was more heavily impacted by emissions from iron and steel production, likely from facilities located to the southwest of the site. This sample displayed a more negative δ^{202} Hg value (-0.50 ± 0.10‰, 2SD) than the other Detroit samples. These data support the hypothesis that Hg emissions from metals processing facilities display negative δ^{202} Hg values.

The variability in Δ^{199} Hg values between the Detroit samples is even more striking (Figure 6B). Sample FRT 24 displayed an extreme negative Δ^{199} Hg value (-0.87 ± 0.15‰, 2SD). Most analyzed rainfall samples have slightly positive Δ^{199} Hg values (Chen et al., 2012; Demers et al., 2013; Donovan et al., 2013; Gratz et al., 2010; Sherman et al., 2012) and no previous studies have measured similarly negative Δ^{199} Hg values in rainfall. As described above, rainfall during this event occurred after a period of several days of local stagnation

characterized by elevated ground-level ozone, SO₂, and fine particulate matter concentrations. It is likely that these pollutants were emitted by local sources and trapped in the area by the persistent high-pressure system and light surface winds. It is possible either that locally emitted Hg displayed extreme negative Δ^{199} Hg values or that negative MIF occurred in the atmosphere during this event. Negative Δ^{199} Hg values have been observed in Arctic snow (Sherman et al., 2010) and in vegetation (Blum et al., 2012; Blum et al., 2014; Carignan et al., 2009; Das et al., 2012; Demers et al., 2013; Estrade et al., 2010; Tsui et al., 2012) likely due to heterogeneous photochemical reduction of Hg from snow crystals and photochemical reduction of thiol-bound Hg, respectively (Sherman et al., 2010; Zheng and Hintelmann, 2010). It is unlikely that these summer, low-elevation air masses (max elevation of 24-hour HYSPLIT back trajectories = 250 m) contained significant snow crystals. Additionally, highly elevated Δ^{200} Hg values, which have been suggested to indicate heterogeneous upper atmospheric processes (Chen et al., 2012), were not observed in these samples (see Appendices B and C). However, we cannot rule of out the possibility that the locally stagnant conditions and elevated sulfur concentrations caused distinct atmospheric conditions resulting in the observed negative MIF. Whether the negative MIF in this sample was due to deposition of Hg emitted by an isotopically distinct Hg source or the presence of distinct atmospheric chemistry, these unique conditions were not discernable using trace metal concentrations and PMF receptor modeling alone.

4. Conclusions

This study represents the first combined measurement of Pb, Sr, and Hg isotope ratios in precipitation and was conducted both to explore the isotopic variability of precipitation in the Great Lakes region as well as the potential usefulness of multiple metal isotopes for

identification of pollution sources. These data will improve isotopic characterization of atmospheric end-members for ecosystem isotope studies. We also observed isotopic differences between precipitation collection sites that were attributable to local and regional differences in anthropogenic source emissions such as coal combustion and MVT Pb ore processing.

In several cases, the isotopic results support previous source apportionment results based on trace element concentrations and PMF receptor modeling. For example, Pb isotope ratios suggest that a significant proportion of the Pb deposition in the Great Lakes region results from coal combustion. Observed differences in Hg and Sr isotope ratios between sites in Michigan and Ohio suggest that coal combustion is a primary source of these metals and largely controls the measured isotope ratios. In addition, Pb and Hg isotope ratios indicate that industrial sites such as Steubenville, Ohio are variably influenced by other industrial sources such as MVT Pb ore processing.

These results corroborate previous PMF receptor modeling and may be used in future modeling efforts to constrain a model's numerical solution and reduce associated uncertainties. We have demonstrated that metal isotope ratios can provide independent corroboration of these estimates and add to the "weight of evidence" needed to identify and link pollution sources with metal deposition. However, these isotopic data often do not provide a simple tracer with which to definitively identify and distinguish between source emissions. Future source attribution using multiple metal isotope ratios will require improved characterization of the isotopic composition of both raw materials and point source emissions. In addition, a more complete understanding of Hg fractionation both within industrial facilities and in the atmosphere is needed. While more research is necessary to further develop these isotopic methods for source

attribution, our results suggest that the use of multiple metal isotope ratios as an additional tool for pollution source attribution holds great promise.

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Table Caption

Table 1. Abbreviated isotope results for precipitation samples. Collection site abbreviations are as follows: Detroit, Michigan (FRT), Dexter, Michigan (DXT), Eagle Harbor, Michigan (EGL), Grand Rapids, Michigan (GRD), Pellston, Michigan (PLN), Steubenville, Ohio (STB), Underhill, Vermont (LCH). Analytical uncertainties for the measured ⁸⁷Sr/⁸⁶Sr ratios (2SE) represent error in the last two or three presented decimal places. Ratios that were not measured due to low metal concentrations are listed as "n.d." Samples that were collected but not archived for future analyses and could not be analyzed are listed as "sample not retained."

Precipitation Date	Site ID	Sample ID	δ ²⁰² Hg (‰)	Δ ¹⁹⁹ Hg (‰)	²⁰⁷ Pb/ ²⁰⁶ Pb	²⁰⁸ Pb/ ²⁰⁶ Pb	⁸⁷ Sr/ ⁸⁶ Sr ±2SE
07/04/04	DXT	741	-0.35	0.36	0.8438	2.0620	0.710081±35
07/12/04	DXT	742	-0.51	0.29	0.8421	2.0541	0.709807±29
01/12/05	DXT	775	sample n	ot retained	0.8404	2.0581	0.708924±32
06/13/05	DXT	803	sample n	ot retained	0.8490	2.0674	0.710721±55
07/24/05	DXT	816	sample n	ot retained	0.8339	2.0449	0.710415±36
09/22/05	DXT	826	sample n	ot retained	0.8336	2.0435	0.709850±31
05/14/05	EGL	683	sample n	ot retained	0.8511	2.0728	n.d.
06/23/05	EGL	692	sample n	ot retained	0.8386	2.0482	0.709669 ± 24
06/28/05	EGL	693	sample n	ot retained	0.8497	2.0753	$0.710208{\pm}40$
07/27/03	FRT	23	-0.50	-0.20	0.8266	2.0344	0.709004±25
08/02/03	FRT	24	0.03	-0.87	0.8360	2.0460	0.708753±26
06/09/04	FRT	99	0.10	0.19	0.8310	2.0409	0.708610±28
05/13/05	FRT	206	sample n	ot retained	0.8294	2.0417	0.708773 ± 78
11/02/03	GRD	135	0.07	0.24	0.8413	2.0602	0.708891±62
05/21/04	GRD	182	-0.06	0.40	0.8423	2.0608	0.709303±26
05/30/04	GRD	183	0.02	0.51	0.8445	2.0663	0.709818±24
10/23/04	GRD	215	-0.17	0.14	0.8411	2.0569	0.708591 ± 30
01/13/05	GRD	234	sample n	ot retained	0.8322	2.0452	0.709155±30
03/30/05	GRD	244	sample n	ot retained	0.8062	1.9917	0.709286±19
05/13/05	GRD	251	-0.05	0.46	0.8430	2.0610	0.709087 ± 42
06/05/05	GRD	256	sample n	ot retained	0.8386	2.0497	0.709465 ± 25
06/30/05	GRD	262	sample n	ot retained	0.8331	2.0427	0.709659 ± 24
07/23/05	GRD	267	-0.12	0.52	0.8345	2.0486	0.709986 ± 44
09/22/05	GRD	278	-0.07	0.27	0.8336	2.0433	0.710371±33
11/05/05	GRD	288	0.09	0.33	0.8402	2.0566	0.709243±35
11/09/05	GRD	289	sample n	ot retained	0.8366	2.0452	0.708992 ± 23
10/02/06	GRD	352	-0.06	0.51	0.8331	2.0466	0.710463±27
8/04-06/03	LCH	163	-0.47	-0.13	S	sample not reta	ined

10/20-21/03	LCH	180	0.07	0.36	sample not retained		
07/01/04	LCH	264	-0.06	0.16	0.8472	2.0649	0.710322±41
8/12-13/04	LCH	280	-0.17	0.99	sample not retained		ained
07/05/05	LCH	368	-0.15	-0.09	0.8480	2.0788	0.709348±25
08/31/05	LCH	382	-0.20	0.42	0.8505	2.0562	0.710083±43
09/17/05	LCH	385	sample not retained		0.8493	2.0686	0.710282±34
04/19/04	PLN	428	0.13	0.66	0.8439	2.0687	0.710122±24
05/23/04	PLN	436	-0.07	0.35	0.8543	2.0937	0.710397±35
08/09/05	PLN	500	sample not retained		0.8338	2.0447	0.710107±32
08/19/05	PLN	502	-0.31	0.15	0.8480	2.0691	0.710112±47
05/11/06	PLN	534	-0.08	0.11	0.8438	2.0649	n.d.
06/14/04	STB	141	Recovery <80%		0.8554	2.0825	0.709959 ± 24
08/19/04	STB	157	-0.30	0.85	0.8299	2.0324	0.709702 ± 33
8/28-30/04	STB	159	-0.37	0.74	0.8425	2.0531	0.710496 ± 28
09/08/04	STB	160	-0.68	0.51	0.8252	2.0267	0.711469±43
01/11/05	STB	177	-0.68	0.05	0.8249	2.0140	0.709678 ± 50
02/09 &							
02/14/05	STB	180	-1.13	-0.14	0.8088	1.9591	0.71051±37
03/28/05	STB	186	-0.60	0.33	0.8390	2.0573	0.709951±58
04/02/05	STB	187	-0.63	0.14	sample not retained		
04/20/05	STB	188	-0.09	0.48	sample not retained		
05/13/05	STB	190	-0.37	0.41	0.8363	2.0510	0.710468 ± 31
06/28/05	STB	198	-0.50	-0.14	0.8349	2.0457	0.711553 ± 34
06/30/05	STB	200	-0.34	0.07	0.8332	2.0385	0.710435 ± 101
07/26/05	STB	212	sample n	ot retained	0.8440	2.0552	0.710781±21
10/23/05	STB	230	sample not retained		0.8376	2.0498	0.710633±97
11/14/05	STB	236	-0.80	0.02	0.8496	2.0689	0.710597±65
01/02/06	STB	245	-0.63	0.19	0.8393	2.0567	$0.710140{\pm}22$
04/21/06	STB	266	-0.15	0.28	sample not retained		
06/22/06	STB	275	-0.14	0.73	0.8438	2.0583	0.710287±36
01/05/07	STB	322	-0.07	0.09	n.d.	n.d.	0.710036±35

Figure Captions

Figure 1. Map of precipitation collection sites and sources in the Great Lakes region. Depicted sources are those that emitted >1 kg of Hg per year in 2005 as reported in the U.S. EPA National Emissions Inventory and Environment Canada National Pollutant Release Inventory (U.S. EPA, 2005; NPRI, 2008). Collection sites are marked with yellow stars and sources are depicted using symbols as indicated in the legend. Site name abbreviations are as follows: Detroit, Michigan (FRT), Dexter, Michigan (DXT), Eagle Harbor, Michigan (EGL), Grand Rapids, Michigan (GRD), Pellston, Michigan (PLN), Steubenville, Ohio (STB), Underhill, Vermont (LCH).

Figure 2. Pb isotope ratios in precipitation samples and sources. Previously published analyses of North American coals are shown as x's (Chow and Earl, 1972; Díaz-Somoano et al., 2009), North American MVT Pb ores are shown as black squares (Doe and Delevaux, 1972; Goldhaber et al., 1995; Heyl et al., 1974), and North American precipitation samples (collected after 1990) are shown as black triangles (Graney and Landis, 2013; Simonetti et al., 2000c). Precipitation samples collected at rural (Figure 2A: DXT, EGL, LCH, and PLN) and urban sites (Figure 2B: FRT, GRD, and STB) are shown as colored diamonds. Site name abbreviations are the same as in Figure 1. Case study sample STB 180 is labeled (see section 3.4). In both plots, analytical uncertainty (2SD) for precipitation samples based on measurement of the BCR-2 and TMRAIN-95 standards is less than the size of the symbols.

Figure 3. ⁸⁷Sr/⁸⁶Sr versus ²⁰⁸Pb/²⁰⁶Pb in precipitation samples. Precipitation samples are shown using colored symbols by collection site with the same site name abbreviations as in Figure 1. Case study samples FRT 24 and STB 180 are labeled (see sections 3.4). Analytical uncertainties

for ⁸⁷Sr/⁸⁶Sr ratios (2SD) are shown as error bars. Analytical uncertainty for ²⁰⁸Pb/²⁰⁶Pb ratios (2SD) based on measurement of the BCR-2 and TMRAIN-95 standards is less than the size of the symbols.

Figure 4. δ^{202} Hg (‰) versus Δ^{199} Hg (‰) in precipitation samples. Precipitation samples are shown using colored symbols by collection site with the same site name abbreviations as in Figure 1. North American precipitation samples (post 1990) impacted by mixed sources are shown as black triangles (Chen et al., 2012; Demers et al., 2013; Donovan et al., 2013; Gratz et al., 2010; Sherman et al., 2012), and North American precipitation samples directly impacted by a local coal-fired power plant are shown as black circles (Sherman et al., 2012). Case study samples FRT 24 and STB 180 are labeled (see sections 3.4). Representative 2SD analytical uncertainty based on analyses of procedural standards is shown.

Figure 5. Percentage of attributable metal deposition associated with sources based on PMF versus isotope ratios for Steubenville, Ohio precipitation samples. Each precipitation sample is represented by one isotope ratio (y-axis) and four or five source contributions (x-axis). Case study sample STB 180 is labeled (see section 3.4). A). Percentages of attributable Pb deposition previously associated with each source factor by PMF (White, 2009) versus measured 208 Pb/ 206 Pb ratio. B). Percentages of attributable Hg deposition previously associated with each source factor by PMF (White, 2009) versus measured source factor by PMF (White, 2009) versus measured 208 Pb/ 206 Pb ratio. B). Percentages of attributable Hg deposition previously associated with each source factor by PMF (White, 2009) versus measured 208 Pb/ 206 Pb ratio. B). Percentages of attributable Hg deposition previously associated with each

Figure 6. Percentage of attributable Hg deposition associated with sources based on PMF versus Hg isotope ratios for Detroit, Michigan precipitation samples. Each precipitation sample is

represented by one isotope ratio (y-axis) and four source contributions (x-axis). Case study samples FRT 23 and FRT 24 are labeled (see section 3.4). A). Percentages of attributable Hg deposition previously associated with each source factor by PMF (UMAQL, 2009) versus measured δ^{202} Hg value. B). Percentages of attributable Hg deposition previously associated with each source factor by PMF (UMAQL, 2009) versus measured Δ^{199} Hg value.

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