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Sedimentary Mercury Enrichments as a Tracer of Large Igneous Province Volcanism

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ABSTRACT

Volcanic activity associated with the emplacement of Large Igneous Provinces (LIPs) has been linked to most Phanerozoic extinctions/episodes of major environmental change. In recent years, mercury (Hg) enrichments and elevated mercury/total organic carbon (Hg/TOC) ratios have been increasingly utilized as a marker of volcanism in sedimentary records deposited distally from LIPs. The proxy is based on the premise that volcanism is a major natural source of the element to the atmosphere, and was especially important prior to anthropogenic emissions. To date, end-Permian and end-Triassic records illustrate the strongest use of Hg as a volcanic proxy; aided by supporting evidence (including Hg isotopes) for LIP eruptions and/or volatile emissions. Sedimentary records of several other events also document Hg enrichments in at least one region, suggesting regional or global Hg-cycle perturbations potentially linked to volcanism at those times. The Cenomanian-Turonian Oceanic Anoxic Event appears to be an exception, with Hg/TOC peaks documented in a small minority of studied records, suggesting minimal Hg-cycle disturbance at that time. Even for events that apparently featured a global-scale Hg-cycle perturbation, variable Hg enrichments across individual archives of that same crisis indicate that the complex biogeochemical cycling of the element can strongly influence local/regional aquatic, biological, or sedimentary processes to alter the precise signature of any worldwide disturbance. Recent studies are beginning to investigate these complexities, but further work is needed to fully explore the nuances of Hg in the geological record, and how it can be best employed as a proxy for LIP volcanism.

11.1. INTRODUCTION

Volcanic activity related to the emplacement of Large Igneous Provinces (LIPs) has long been thought to play a significant causal role in several episodes of environmental perturbation during the Phanerozoic Eon, often resulting in major mass extinctions (e.g., Rampino & Strothers, 1988; Wignall, 2001, 2005; Courtillot & Renne, 2003; Bond & Wignall, 2014; Bond & Sun, Chapter 3 this volume). This hypothesis of volcanically stimulated climatic/biospheric stress is based chiefly on a very strong correlation between the established dates of Phanerozoic events of environmental/biospheric change and the radiotopic ages of igneous rocks associated with various LIPs (Fig. 11.1; see also recent studies by Svensen et al., 2012; Blackburn et al., 2013; Sell et al., 2014; Burgess & Bowring, 2015; Renne et al., 2015; Schoene et al., 2015; Davies et al., 2017; and the review of Kasbohm et al., Chapter 2 this volume). However, correlating the ages of LIP volcanics and extinction/climate episodes in this way is hindered by uncertainties attendant in the determination and comparison of stratigraphic age models and radioisotopic dates, and a lack of precise ages for several geologic events and LIP eruptions. Moreover, the volcanic
activity of many LIPs began prior to or continued after the environmental/biospheric crises with which they are associated. This last point relates to the proposed causal mechanisms linking LIP volcanism to climate/extinction events revolving around the release of volcanic gases (e.g., CO₂ and SO₂) to the atmosphere, rather than the emplacement/eruption of igneous material (see Mather & Grasby, 2013b, 2016, 2017; Davies et al., 2017). It has been proposed that the most environmentally damaging gas emissions associated with LIPs were not necessarily magmatic in origin, but instead might have resulted from the metamorphism of volatile-rich sediments by the emplacement of intrusive sills (e.g., Svensen et al., 2004, 2009; McElwain et al., 2005; Ganino & Arndt, 2009; Burgess & Bowring, 2015; Davies et al., 2017). Recently, sedimentary mercury (Hg) enrichments (as characterized by elevated Hg concentrations and Hg/total organic carbon ratios) have been proposed as a potential proxy for volcanic activity and volatile release (Sanei et al., 2012). A major natural source of Hg to the atmosphere is emission of the element as a trace volcanic gas, which has the potential for global distribution through the atmosphere (see below and Fig. 11.2a). Consequently, increased Hg concentrations in sedimentary strata that record episodes of mass extinction/environmental perturbation are frequently interpreted as evidence for large-scale volcanism and/or volatile release at those times, which may have acted as the trigger for the biospheric/climatic crisis. Such peaks in Hg content have been reported from records of all five major Phanerozoic extinctions, as well as various other climatic perturbations such as the Oceanic Anoxic Events (OAEs) and the Paleocene-Eocene Thermal Maximum (e.g., Sanei et al., 2012; Percival et al., 2015; Font et al., 2016; Thibodeau et al., 2016, 2014; Fendley et al., 2019, in press) and the Paleocene-Eocene Thermal Maximum (PETM) (e.g., Fantasia et al., 2018, 2019). In this context, it is important to note that the proposed causal mechanisms linking LIP volcanism to climate/extinction events involve the emission of gases into the atmosphere rather than the emplacement/eruption of igneous material (see Mather & Grasby, 2013b, 2016, 2017).
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Figure 11.2 (a) The natural mercury cycle, a simplified illustration from Percival et al. (2018). Atmospheric and oceanic residence times of Hg are indicated (Slemr et al., 1985; Gill and Fitzgerald, 1988; Blum et al., 2014; Gustin et al., 2015). Processes affecting the source, sink, and transportation of mercury through the ocean-atmosphere system are indicated as follows: (1) Volcanic emission of mercury, chiefly as gaseous elemental mercury (Hg\textsubscript{0}) to the atmosphere; (2) Hydrothermal emission of Hg to the ocean at midocean ridges; (3) Presumed release of hydrothermal Hg to the ocean from submarine ocean plateau volcanism, similar to that at midocean ridges; (4) Possible emission of thermogenic Hg to the atmosphere following heating of organic-rich sediments by intruding sills; (5) Removal of atmospheric Hg to land, water, or forest canopy, either as direct uptake of gaseous Hg\textsubscript{0} by plants or deposition of particulate Hg (dry deposition) and soluble oxidized mercury (Hg\textsuperscript{2+}) following interactions between atmospheric Hg\textsubscript{0} and atmospheric oxidizing agents such as halogen, nitrile, ozone, and hydroxyl radicals (wet deposition); (6) Air-water interchange of Hg\textsubscript{0}; (7) Conversion of mercury species between Hg\textsubscript{0}, Hg\textsuperscript{2+}, and MeHg (monomethylmercury and dimethylmercury) through multiple biotic and abiotic reactions in aquatic environments (for example, Fitzgerald et al., 2007; Selin, 2009; Blum et al., 2014; Bowman et al., 2015); (8) Reduction of soil Hg to Hg\textsubscript{0}, which is subsequently reemitted to the atmosphere; (9) Interchange of mercury between soil and forest canopy through emission of soil Hg and decay of leaves that have taken up Hg; (10) High abundance of sulfate- and/or iron-reducing bacteria in reduced wetland environments promoting methylation of Hg\textsuperscript{2+} to organophilic monomethylmercury (MMHg), which can adsorb onto organic matter; (11) Deposition of sediments in soil Hg-OM complexes to peats or coals; (12) Riverine runoff (and potentially deposition) into lacustrine or marine environments of detrital Hg bound to either organic matter or clay minerals; (13) Uptake of Hg\textsuperscript{2+} or MMHg by aquatic biota; (14) Bioaccumulation of organophilic MMHg up the food chain; (15) Deposition of Hg into sediments as Hg-OM complexes; (16) Potential remobilization and release of sedimentary Hg into the aquatic realm. (b) Mass independent fractionation (MIF) of mercury in a simplified illustration of the mercury cycle. Grey lettering indicates a source or inventory of Hg at the Earth’s surface, and changes in the speciation of mercury (Hg\textsubscript{0} = inert gaseous elemental mercury; Hg\textsuperscript{2+} = reactive oxidized mercury; MeHg = mono- or di-methylmercury; Hg-OM = organo-mercury complexes). Black lettering indicates inventories of Hg and their average $\Delta^{199}$Hg composition in the modern (values taken from data in the model of Sonke, 2011; and the review of Blum et al., 2014), and pathways of Hg where MIF may occur. Bold arrows also indicate pathways of Hg where MIF occurs (e.g., photochemical reduction in surface-marine waters or raindrops, and uptake of Hg by plants/lichens). Thinner dashed arrows indicate pathways of Hg where MIF is not known to occur.
and the possible conclusions regarding how mercury operated during times of extinction/environmental perturbation and the use of this element as a proxy for volcanism are discussed.

### 11.1.1. The Natural Mercury Cycle

Volcanic outgassing represents a major natural (nonanthropogenic) source of mercury to the Earth’s surface in the modern (e.g., Schroeder & Munthe, 1998; Pyle & Mather, 2003; Bagnato et al., 2007, 2011, 2014). Volcanic Hg is largely emitted to the atmosphere as a gaseous elemental species (Hg⁰), with a smaller fraction erupted as oxidized or particulate species that have shorter atmospheric lifetimes and are likely to be deposited locally (Bagnato et al., 2007). Hg⁰ has a typical atmospheric residence time on the order of about 1 year. Removal of the element from the atmosphere typically occurs via oxidation by various species (e.g., halogen radicals, nitrous oxide, ozone) to Hg²⁺, followed by either “dry” or “wet” deposition to the terrestrial and aquatic realms (see Fig. 11.2a; Schroeder & Munthe, 1998; Selin, 2009). Additionally, Hg⁰ in the atmosphere can be directly taken up by plant foliage (e.g., Ericksen et al., 2003; Demers, 2013; Enrico et al., 2016), and transferred into soils via leaf-litter deposition (Obrist, 2007; Wang et al., 2016). A small fraction of gaseous Hg⁰ may also be directly adsorbed into soils (Obrist et al., 2011). The majority of soil Hg is thought to be buried bound to organic compounds, but a small fraction may be reduced and returned to the atmosphere as Hg⁰ (e.g., Lindberg et al., 1998; Obrist, 2007; Wang et al., 2016; Obrist et al., 2018). In the aquatic realm, a range of biotic and abiotic reactions can transform mercury between elemental Hg⁰ (which may then be reemitted to the atmosphere), oxidized Hg²⁺, or toxic monomethyl-/dimethylmercury (see review by Fitzgerald et al., 2007).

The methylation of mercury typically occurs in poorly oxygenated aquatic settings (e.g., bottom waters and sediment pore waters) and is largely facilitated by microbes (e.g., Fitzgerald & Lamborg, 2014). Ultimately, aquatic mercury is either evaded to the atmosphere as Hg⁰ or deposited to sediments, typically bound to organic compounds. Consequently, concentrations of the element tend to correlate well with the total organic carbon (TOC) content in modern sediments (e.g., Sanei & Goodarzi, 2006; Outridge et al., 2007; Liu et al., 2012; Ruiz & Tomyisya, 2015), resulting in a Hg/TOC ratio that should remain relatively constant under stable conditions in a particular setting. Thus, studies of sedimentary mercury normalize the Hg concentration to the TOC content (see Sanei et al., 2012). An increased Hg content in waters with no change in TOC will lead to higher Hg/TOC ratios that might indicate an increased external influx of Hg to that environment (see, e.g., Jin & Liebezeit, 2013; Sanei et al., 2014; Daga et al., 2016). Sulfides and clay minerals can also scavenge mercury from the water column, and in certain environments deposition of mercury with those species might override the usual sedimentary Hg-organic matter relationship (see Sanei et al., 2012; Charbonnier & Föllmi, 2017; and Percival et al., 2018). However, more work is needed to determine in what sedimentary contexts it is most prudent to normalize Hg against which species, and normalizing against all of them may be of use for some records (see Grasby et al., 2019). Southern Ocean diatomite sediments of Holocene age have also been noted as featuring large Hg concentrations, suggesting that biogenic silica might also be a potentially important sink of oceanic Hg in this context (Zaferani et al., 2018), although it is currently unknown whether this relationship also applies in other oceanographic settings or if biogenic silica would preserve changes to the global Hg cycle in the deep-time geological record. Submarine volcanism also emits Hg to the aquatic realm (e.g., Lamborg et al., 2006; Bagnato et al., 2014); however, the influence of subaqueous volcanic Hg emissions appears to be limited to only the regional marine environment immediately proximal to that volcanic system (Bowman et al., 2015). Consequently, it is not clear to what extent submarine volcanism can perturb the global Hg cycle.

### 11.1.2. Mercury Isotopes

In addition to sedimentary Hg concentrations and Hg/TOC ratios, variations in the natural stable isotopes of Hg are also being applied to understanding past Hg perturbations in the geological record. Mercury has seven stable isotopes: ^{198}\text{Hg} (0.16%), ^{199}\text{Hg} (10.04%), ^{200}\text{Hg} (16.94%), ^{201}\text{Hg} (23.14%), ^{202}\text{Hg} (13.17%), ^{203}\text{Hg} (29.73%), and ^{204}\text{Hg} (6.83%), which can undergo both mass-dependent (MDF) and mass-independent (MIF) fractionation in the natural environment. MDF, reported as variations in the ^{202}\text{Hg}/^{198}\text{Hg} ratio in delta notation (Δ^{202}\text{Hg}), occurs during many transformations including redox, biological, and phase changes. In contrast, large MIF for Hg only occurs during photochemical reactions with odd isotopes displaying MIF during aqueous photochemical reactions (reported as Δ^{199}\text{Hg}), and even MIF occurring during gas phase atmospheric photochemical reactions (reported as Δ^{200}\text{Hg}) (see Blum et al., 2014). This review focuses on the application of sedimentary mercury isotope compositions to understanding past volcanic events, focusing on odd-isotope MIF (Δ^{199}\text{Hg}); a more comprehensive review of mercury-isotope systematics in natural processes is given by Blum et al. (2014). Constraints on the isotopic composition of mercury emissions from modern volcanism are limited to only one study (Zambardi et al., 2009), which reported mercury with a ^{8/20}\text{Hg} composition between 0‰ and -2‰, and no measurable MIF (i.e.,...
Δ^{199}Hg = 0 ‰). This work represented only one volcanic system and focused on an arc-setting not analogous with LIPs, for which the isotopic signature of volcanically emitted Hg^0 might have been rather different. However, the results are consistent with other investigations of geogenic Hg-isotope compositions (in settings where the Hg has not undergone any atmospheric or aqueous cycling) that also reported similar δ^{202}Hg compositions and a lack of MIF (see Blum et al., 2014). It should be noted though that MDF can be caused by biological, physical, and chemical processes at the Earth’s surface, and potentially post-depositionally (see Blum et al., 2014; Thibodeau et al., 2016). Thus, several studies recommend caution in interpreting sedimentary δ^{202}Hg values between 0 ‰ and -2 ‰ as evidence of volcanic influx, as they may not be representative of the original source composition (e.g., Thibodeau et al., 2016; Grasby et al., 2017; Thibodeau & Bergquist, 2017). Nonetheless, trends in MDF may still reflect important changes in the Hg cycle and sources to it (Grasby et al., 2017), and may become more useful in the future with more understanding of Hg MDF preservation in sediments.

Consequently, trends in MIF of mercury (specifically Δ^{199}Hg) are currently more helpful for investigating records of environmental change potentially driven by volcanism, as volcanic (and nonphotochemically cycled) Hg likely has an isotopic signature distinct from inventories of that element that has undergone significant cycling through surface reservoirs. Mostly, odd-isotope MIF of mercury is thought to be dominated by two main processes at the Earth’s surface. First, the photochemical reduction of aqueous Hg^{2+} to Hg^0 in surface waters and raindrops (Bergquist & Blum, 2007; Demers et al., 2013), which produces negative Δ^{199}Hg values in the resultant Hg^0 gas (and consequently a slightly negative Δ^{199}Hg composition in the atmospheric inventory) and a residual pool of aqueous Hg^{2+} with a more positive Δ^{199}Hg signature (Fig. 11.2b). Second, during the uptake of Hg^0 by foliage (e.g., Carignan et al., 2009; Demers et al., 2013), there is additional MIF that enriches the plant matter in Hg with a Δ^{199}Hg composition even more negative than that of atmospheric mercury (Fig. 11.2b). Consequently, soils typically have negative Δ^{199}Hg compositions because the pathway of Hg to them is dominated by leaf litter fall (e.g., Demers et al., 2013). In general, marine sediments receive Hg from two main sources: (1) incorporation of aqueous Hg^{2+} deposition with positive Δ^{199}Hg and (2) runoff from terrestrial soils with negative Δ^{199}Hg (Blum et al., 2014). Thus, Hg in sediments typically features either significant positive or negative MIF, depending on the relative proportion of the two major pathways of Hg entering the ocean or body of water (Thibodeau & Bergquist, 2017). Positive sedimentary Δ^{199}Hg values reflect a predominantly atmospheric Hg influx most commonly observed in strata that were deposited in a marine environment far from land. In contrast, nearshore-marine and terrestrial sediments where Hg deposition is dominated by runoff from land tend to have negative Δ^{199}Hg compositions, consistent with an influx of Hg from soils. Because volcanic Hg is thought to be devoid of MIF at the point of emission, it has been proposed that a sedimentary Hg enrichment that lacks a MIF signature could reflect an increased output of Hg by volcanoes, which has either overwhelmed or bypassed the MIF inducing processes detailed above (e.g., Thibodeau et al., 2016). However, it should be noted that considerable work is still needed to fully understand all processes that can result in isotopic variations in Hg reservoirs (Blum et al., 2014). For example, it has recently been proposed that negative MIF could also arise in sediments deposited under strongly euxinic marine conditions (Zheng et al., 2018).

### 11.2. DISCUSSION OF GEOLOGICAL MERCURY RECORDS

#### 11.2.1. Do Stratigraphic Records of All Major Events Document Hg-Cycle Perturbations?

Of key significance in interpreting geological records of the past Hg cycle is the necessity to investigate multiple stratigraphic archives from different locations. The importance of adopting this multisite approach is demonstrated by the mercury records of the Cenomanian-Turonian OAE 2 (94 Ma). Increased Hg concentrations and Hg/TOC ratios were initially reported from OAE 2 strata in the southern part of the Western Interior Seaway (Texas, USA) and Demerara Rise in the central Atlantic, from which a Hg-cycle perturbation was inferred (Scaife et al., 2017; Percival et al., 2018). However, further studies of boreal, Tethyan, and Indian Ocean records have shown that the majority of OAE 2 records do not feature clear evidence of increased Hg contents or Hg/TOC ratios; thus, it appears that any Hg-cycle perturbation during OAE 2 was at most only regional in extent (Percival et al., 2018). Similarly, sedimentary strata recording the Smithian-Spathian extinction (250 Ma) feature elevated Hg/TOC in the Arctic (Grasby et al., 2013b, 2016; Hammer et al., 2019) but more variable trends across South China, Indian Tethyan, and Panthalassic records (Wang et al., 2019; Shen et al., 2019a). Therefore, it should not be assumed that Hg enrichments observed from one area are manifestly indicative of a global Hg-cycle perturbation.

At present, widespread records of Hg enrichments spanning multiple continents exist for only a few events. The end-Ordovician extinction (444 Ma), end-Permian
extinction (252 Ma), end-Triassic extinction (201 Ma), Toarcian OAE (183 Ma), and end-Cretaceous extinction (66 Ma), all feature Hg enrichments that can be correlated on the basis of biostratigraphic or carbon-isotope information across numerous sedimentary sequences spanning multiple regions/continents and both hemispheres (Sanei et al., 2012; Grasby et al., 2013b, 2016, 2017; Sial et al., 2013, 2014, 2016, 2020; Percival et al., 2015, 2017, 2018; Font et al., 2016; Thibodeau et al., 2016; Gong et al., 2017; Jones et al., 2017; Fantasia et al., 2018, 2019; Keller et al., 2018; Wang et al., 2018; Burger et al., 2019; Fendley et al., 2019; Lindström et al., 2019; Meyer et al., 2019; Shen et al., 2019b, 2019c; Smolarek-Lach et al., 2019; Them et al., 2019). There are also widespread stratigraphic records of the Frasnian-Famennian (372 Ma), Devonian-Carboniferous (359 Ma), and Capitanian (end-Guadalupian, 260 Ma) extinctions, the Valanginian “Weissert” Event (135 Ma), and Paleocene-Eocene Thermal Maximum (PETM, 54 Ma) that feature Hg enrichments (Grasby et al., 2016; Charbonnier et al., 2017, in press; Keller et al., 2018; Kwon et al., 2018; Racki et al., 2018a, 2018b; Jones et al., 2019; Kalvoda et al., 2019; Liu et al., 2019; Paschall et al., 2019). However, it should be noted that the Toarcian, end-Cretaceous, and PETM Hg records show significant variation across the studied sites, indicating that local sedimentary and/or environmental factors strongly influenced Hg deposition in some locales at those times, in addition to any postulated global perturbation by volcanism (Percival et al., 2015, 2018; Jones et al., 2019). There is also no unambiguous candidate LIP for the end-Ordovician extinction currently, and it has been proposed that Hg enrichments in uppermost Ordovician-lowermost Silurian strata are redox controlled rather than volcanically sourced (Shen et al., 2019b).

Mercury enrichments have been also noted from Tethyan records of the early Aptian OAE 1a (121 Ma), and Aptian-Albian OAE 1b (113 Ma), as well as in a number of other Tethyan black shale horizons of Hauterivian-Barremian (Early Cretaceous, 133–125 Ma) age (Charbonnier & Föllmi, 2017; Charbonnier et al., 2018; Sabatino et al., 2018). However, because these Hg-cycle perturbations are all observed in Tethyan records, they do not give a global overview of the Hg cycle at those times. Similarly, although stratigraphic archives of two Cambrian events show Hg enrichments (Faggetter et al., 2019; Pruss et al. 2019), those records are currently restricted to single areas and might only document a regional Hg influx and/or local-process controls on mercury deposition.

It is less clear to what extent smaller non-LIP eruptions (e.g., arc volcanism) can perturb the local or global mercury cycle, with the quantities of Hg gas emitted during these more minor events assumed to be lower. Marine sediments studied for the Toarcian and Cenomanian OAEs that were deposited proximally to arc systems did not show any Hg enrichments except within the event strata, suggesting that any local eruptions either did not perturb the local mercury cycle, or that any enrichments were not preserved in the geological record (Percival et al., 2015; Scaife et al., 2017; Fantasia et al., 2018). However, Racki et al. (2018a) did indicate possible correlations between Hg enrichments in marine sedimentary rocks and large arc eruptions. Local eruptions might be more easily preserved in the terrestrial realm. Andean eruptions (together with local wildfires) have been proposed as resulting in Hg enrichments of modern-day wetland and lake sediments (e.g., Ribeiro Guevara et al., 2010; Daga et al., 2016). Percival et al. (2018) speculated that high Hg values in uppermost Cretaceous strata from Montana might be related to local arc volcanism (which is also evidenced by abundant tuff beds), although Fendley et al. (2019) concluded that local volcanism was unlikely to have been the cause of these peaks, and the Deccan Traps were also volcanically active when those sediments were deposited so a LIP source for the mercury could not be excluded.

### 11.2.2. Nonvolcanic Influences on Sedimentary Hg/TOC Variations

In addition to the widespread documentation of Hg-cycle perturbations, a further step in demonstrating that mercury enrichments can act as a tracer for volcanic activity is the confirmation that peaks in Hg/TOC ratios do not appear in sediments deposited at times when large-scale volcanism was not occurring. Figure 11.3 shows two such records studied by Grasby et al. (2013b) and Percival et al. (2015). The Kimmeridge Clay core records more than 1 Myr of Late Jurassic time, not thought to coincide with LIP emplacement. Very low Hg/TOC ratios, featuring minimal variations, are consistent with a lack of large-scale Hg emissions to the atmosphere at that time (Percival et al., 2015). Smith’s Creek records elevations in Hg/TOC at the termination of the end-Permian extinction event and the Smithian-Spathian boundary, both of which are thought to coincide with pulses of Siberian Trap volcanism (Grasby et al., 2013b). However, the remainder of the sequence, which documents some 5 Myr of Early Triassic time, features relatively low and constant Hg/TOC values. Both archives document local redox changes evidenced by variability in TOC, pyrite abundance, and/or trace metal content (Morgans-Bell et al., 2001; Grasby et al., 2013a), but neither shows a correlation between those redox changes and Hg/TOC ratios, suggesting that localized variations in marine oxygenation do not necessarily influence Hg deposition. In contrast, other studies (of Late Cambrian and Ordovician-Silurian boundary records) have concluded that local changes in redox...
could indeed have influenced mercury deposition, based on a correlation of Hg and Hg/TOC with glauconite and pyrite content (Shen et al., 2019b; Pruss et al., 2019). However, it should be noted that no oceanic crust is preserved from the time of interest for either study, so a LIP source cannot be totally excluded in either case. Also, pyrite sequestration has been proposed as playing an important role in the draw-down of Hg from LIP volcanism in some settings, especially where organic-matter burial is limited (Sanei et al., 2012).

However, lithological variations arising from redox changes certainly can impact Hg/TOC ratios. The Toarcian and Aptian OAEs in particular demonstrate this issue. In 10 out of 13 Toarcian OAE archives studied for mercury, Hg content increases (Percival et al., 2015; Fantasia et al., 2018; Them et al., 2019). However, 4 of those records also document marked elevations in TOC, potentially overprinting any volcanic signal at those locations (see Percival et al., 2015). A similar phenomenon has been proposed for Aptian OAE records, where high TOC contents are thought to mute the magnitude of observed Hg/TOC peaks (Charbonnier & Föllmi, 2017). A younger Pleistocene record of Mediterranean sapropels might also illustrate this point, with Hg/TOC ratios noticeably lower in more organic-rich sapropel strata compared with background sediments, despite constant MIF values indicating that there was no change in the source of Hg to the immediate area (Gehrke et al., 2009). In fact, mercury concentrations increase within the sapropels, but are likely overprinted by the greater increase in TOC content. Very low TOC contents can also hinder interpretation of Hg/TOC ratios, as the low TOC values result in a very high Hg/TOC ratio that might vary dramatically depending on tiny changes in the TOC content. At present, Grasby et al. (2016) recommend not employing the Hg/TOC ratio for TOC contents under 0.2 wt%, in order to avoid generation of false Hg/TOC peaks that are unrelated to any increased Hg content (see Grasby et al., 2019, for a more in-depth discussion). Future increases in the precision of TOC measurements may mean that this recommendation can be revised. There is little published information on how sediment heating/oxidation may affect Hg/TOC ratios, although one study of Cenomanian-Turonian OAE 2 records suggests that TOC loss during organic-matter maturation may exaggerate Hg/TOC values and generate “false” volcanic peaks, in which case preservation and/or type of organic matter should be scrutinized when interpreting Hg/TOC.

Figure 11.3 Hg trends during nonvolcanic times. Published Hg/TOC trends from the Early Triassic and Late Jurassic long-term records of Smith Creek (Canada) and the Kimmeridge clay (England, UK), respectively. Temporal information and Hg/TOC data for the Kimmeridge clay (including ammonite zones) is from Percival et al. (2015) and for Smith Creek from Grasby et al. (2013b). The positioning of times of LIP volcanism associated with the end-Permian (PT) and Smithian-Spathian intervals at Smith Creek is postulated based on Grasby et al. (2013b).
trends (Charbonnier et al., 2020). Regardless, the key point in interpreting Hg/TOC trends is that elevated Hg/TOC ratios should not be interpreted as a volcanic signal unless they are correlated with an increase in the Hg concentration (i.e., there has been an increased influx/burial of Hg above and beyond the normal drawdown by organic material). Both increased Hg/TOC values derived from a fall in TOC content, and elevated Hg concentrations unaccompanied by a rise in Hg/TOC, cannot be unambiguously stated as being derived from an increase in the overall input of Hg (from volcanism or otherwise) because an alternative driver such as lithological changes could also have caused those variations (see also Grasby et al., 2019).

Finally, in addition to volcanic Hg emissions, there may be alternative sources of mercury that can perturb ocean-atmosphere reservoirs. Heating of organic-rich sedimentary rocks by intrusive sills may have supplemented magmatic Hg with thermogenic emissions of the element (see, e.g., Svensen et al., 2004; Percival et al., 2015; Jones et al., 2019), although such outgassing would likely still be related to LIP emplacement. Another possibility is a major release of Hg from soils and forests, since they are the largest surface reservoir of the element (Fitzgerald & Lamborg, 2014). A large runoff of terrigenous material during a time of enhanced continental weathering could cause a sedimentary enrichment in Hg, potentially also recycling any volcanically emitted Hg (Them et al., 2019; Hammer et al., 2019). Burning of coal/biomass via magmatic intrusions or wildfires may also release Hg as Hg\(^0\) gas or bound to coal fly ash or charcoal particles (e.g., Sanei et al., 2012; Grasby et al., 2017). Thermogenic, run-off, and wildfire processes are thought to have acted during a number of Phanerozoic environmental perturbations, and may act as supplementary, recycled, or alternative sources of Hg to sediments (Grasby et al., 2019). However, an influx of Hg to marine sediments from wildfire or terrigenous sources can be tested on the basis of whether a Hg enrichment correlates with other evidence such as charcoal/fly ash, Hg-isotopic excursions, or other markers of terrigenous material. Thus, attributing global enrichments in Hg to volcanism is strongest when also correlated with other possible markers of volcanism.

11.2.3. Correlating Hg enrichments with other evidence of LIP volcanism

For two events associated with LIP volcanism, the end-Triassic extinction and Toarcian OAE, atmospheric contents of other gases potentially derived from volcanism or metamorphism of volatile-rich country rocks by LIP magmas (e.g., carbon dioxide, CO\(_2\); sulfur dioxide, SO\(_2\)) have been reconstructed on the basis of plant fossils from the same stratigraphic records investigated for mercury (McElwain et al., 2005; Steinthorsdottir et al., 2011, 2018; Bacon et al., 2013). A close (though not perfect) correlation is observed between trends in CO\(_2\), SO\(_2\) (end-Triassic only), and Hg and Hg/TOC (Fig. 11.4). The existence of these correlations within single sedimentary records may strengthen the case for production of all three gases by volcanic and/or thermogenic emissions, and volcanogenically triggered Hg-cycle perturbations during those two events.

However, for records of some geological events, there is clear geochronological/geochemical evidence of volcanism from other markers that do not match sedimentary Hg enrichments (although they are not proxies of volcanic gases). As already discussed, it is unclear whether there was a global perturbation in the Hg cycle during OAE 2, but there is clear evidence of volcanic activity at that time from a globally observed shift in sedimentary osmium-isotope ratios to unradiogenic (mantle) values (Turgeon & Creaser, 2008; Du Vivier et al., 2014, 2015; also see Dickson et al., Chapter 10 this volume, for an overview of osmium-isotope stratigraphy associated with LIP volcanism). The unclear evidence for sedimentary Hg enrichments suggests that this volcanism did not greatly influence the Hg cycle, possibly because a large part of the LIP volcanism at that time was likely submarine and any Hg emissions may have been less efficiently dispersed, generating the major spatial variability in mercury records associated with that event (see Percival et al., 2018). Similarly, while peaks in Hg and Hg/TOC have been reported from numerous end-Cretaceous sequences, the LIP volcanism associated with Cretaceous-Paleogene times (the Deccan Traps in India) commenced ~300 kyr earlier (e.g., Renne et al., 2015; Schoene et al., 2015), and is well documented in the sedimentary record by osmium-isotope evidence for major weathering of mantle-derived basalts (Ravizza & Peucker-Ehrenbrink, 2003; Robinson et al., 2009). Currently, there is little evidence for a perturbed Hg cycle during those 300 kyr (see Percival et al., 2018), suggesting that perhaps the initial Deccan volcanism had less of an impact on the global Hg inventory up until just prior to the extinction. There is also geochronological evidence that the Permian Siberian Traps (Russia), Toarcian Karoo-Ferrar (South Africa and Antarctica), and Valanginian Paraná-Etendeka (Namibia and Brazil) LIPs were volcanically active at times outside of those marked by sedimentary Hg enrichments (see Percival et al., 2018; Xu et al., 2018). Consequently, it is possible that not all volcanism leads to major perturbations in the atmospheric and surface reservoirs of Hg, which may instead be related to the amount, emplacement, and type of volcanism. Factors such as submarine versus subaerial eruptions, explosive versus effusive events, and intrusion of Hg- (organic-) rich country lithologies by intrusive LIP magmas may be particularly important in determining
Figure 11.4 Evidence of atmospheric Hg versus other volcanogenic gases. Correlations between Hg/TOC records and reconstructed changes in atmospheric CO$_2$ and SO$_2$ based on plant fossil evidence for the Triassic-Jurassic record of Astartekløft (Greenland), and the Toarcian OAE (T-OAE) record of Bornholm (Denmark). For both the Triassic and Toarcian records, the Hg/TOC, CO$_2$, and SO$_2$ (Triassic only) data are all from samples taken from these sections, permitting direct stratigraphic correlation. For Astartekløft, lithological and biostratigraphic information and carbon-isotope data are from Hesselbo et al. (2002); positioning of the end-Triassic extinction horizon (ETE) is based on Mander et al. (2013); Hg/TOC data are from Percival et al. (2017); CO$_2$ data are from Steinthorsdottir et al. (2011); SO$_2$ data are from Steinthorsdottir et al. (2018). For Bornholm, lithological and biostratigraphic information and carbon-isotope data are from Hesselbo et al. (2000); Hg/TOC data are from Percival et al. (2015); CO$_2$ data are from McElwain et al. (2005). For the CO$_2$ data at both locations, light green trends indicate modern standardization; dark green trends indicate Carboniferous standardization (see McElwain et al., 2005; and Steinthorsdottir et al., 2011, for details).
when LIP volcanism results in major Hg-cycle perturbations (see discussion in Percival et al., 2018). Thus, while Hg/TOC peaks likely do not record all volcanic events associated with LIP volcanism, they may potentially indicate the times of major volcanogenic volatile release.

11.2.4. Can Hg Isotopes Indicate That Sedimentary Hg Enrichments Are Volcanically Derived?

In addition to correlations of Hg/TOC peaks with other volcanic markers, Hg-isotope studies are being increasingly employed as a tool to identify a volcanic fingerprint of mercury versus other potential sources of Hg influx. Thibodeau et al. (2016) reported $\Delta^{199}$Hg within error of zero that correlated with elevated Hg contents and Hg/TOC ratios across Triassic-Jurassic interval strata, hypothesized to result from a huge influx of volcanic mercury (i.e., devoid of MIF, see Introduction and Fig. 11.2) that had overwhelmed the normal pathways taken by the element to reach sediments.

It is interesting that, in contrast to the aforementioned end-Triassic study, Hg-isotope records of the end-Ordovician, end-Permian, Toarcian, and end-Cretaceous events do not show zero MIF correlate with sedimentary Hg enrichments (Sial et al., 2016; Gong et al., 2017; Grasby et al., 2017; Them et al., 2019; Wang et al., 2018). For records of these events, stratigraphic archives deposited relatively proximally to land tend to display shifts toward more negative MIF values that have been interpreted as a large influx of terrestrial Hg from a nearby land mass, which are sometimes correlative with elevations in Hg/TOC ratios (e.g., Grasby et al., 2017) (Fig. 11.5), although not always perfectly so (e.g., Them et al., 2019). In contrast, many of the studied marine records document slightly positive $\Delta^{199}$Hg values throughout the studied stratigraphic sequences, which as described above are likely indicative of photochemical reduction during/after atmospheric deposition of Hg to the marine realm. The continuation of such photochemical reduction despite the increased influx of Hg to these records (suggested by correlative Hg/TOC peaks) highlights the durability of this MIF pathway. Thus, a Hg/TOC peak correlative with positive MIF could be indicative of a substantial rise in atmospheric Hg content, possibly resulting from volcanogenic outgassing. In uppermost Permian sections, there is a clear pattern of the records closest to land documenting negative MIF, while more distally deposited records feature positive MIF, and Hg enrichments were observed at all locations (Grasby et al., 2017; Wang et al., 2018). Taken together, these records support an increase
in the global Hg inventory during the end-Permian mass extinction, but suggest that the normal pathways present in biogeochemical cycling and MIF of mercury were largely maintained, although potentially intensified by the elevated Hg abundance at the Earth’s surface during that time.

While there are still relatively few published Hg-isotope data sets from geological records where Hg/TOC peaks have been used to infer volcanic activity, the majority of those Hg-isotope trends do not appear to directly record the original volcanic source, assuming that LIPs emitted Hg devoid of MIF as proposed on the basis of the single study of a modern arc volcano (Zambardi et al., 2009). Rather, Hg isotopes appear to be sensitive to the different pathways taken by Hg to reach sediments deposited proximally or distally to the paleoshoreline (Grasby et al., 2017; Thibodeau & Bergquist, 2017; Bergquist, 2017). However, the use of Hg-isotopes to reconstruct these pathways could still be useful for interpreting how the overall Hg cycle was perturbed. In addition, with a better understanding of MDF, the combined interpretation of MIF and MDF shifts may also help to indicate changes in sources to the global Hg pool.

11.3. SUMMARY

Mercury enrichments have been documented in sedimentary records of all five major mass extinctions and several other episodes of environmental perturbation that occurred during the Phanerozoic. These studies indicate that the mercury cycle was perturbed, at least on a regional scale, during all of those events. As volcanism is a major natural source of mercury to the Earth’s surface, and sediments the ultimate (and geologically rapid) sink, it appears that mercury may show significant promise as a sedimentary proxy for large-scale volcanic activity. The correlation between mercury enrichments and reconstructed increases in other volcanic gases during at least two major events associated with volcanism supports this possibility. Furthermore, the absence of such elevated Hg concentrations in strata that record times not associated with major volcanism lends further credence to the influence of large-scale volcanic eruptions on the global mercury cycle. At the present time, mercury isotopes appear to be a good indicator of pathways taken by mercury (of whatever origin) to reach the sediment, but may not always record an original volcanic source. However, knowledge of these pathways may nonetheless allow more accurate reconstruction of a mercury cycle perturbation and whether any increases in the global inventory were caused directly by volcanogenic processes or by output from other reservoirs such as biomass, soils, and coal.

However, it is also apparent that there are a number of complexities associated with the use of the mercury proxy that remain to be fully understood. Even if an increase to the global mercury inventory is indicated by a broadly correlative enrichment in sedimentary mercury levels from numerous stratigraphic archives across multiple continents, spatial variability in the global signature will inevitably result from sedimentological and/or depositional processes specific to individual environments, as well as the nature of the volcanism itself. And in some cases, those same sedimentological/volcanological controls will mean that a Hg-cycle perturbation might not be recorded at all in some stratigraphic archives, potentially meaning that any Hg enrichment cannot be globally correlated. The clear lack of correlation between mercury enrichments and other geochronological and/or geochemical evidence for Large Igneous Province volcanism is suggestive that not all eruptive events will manifestly result in a major output of mercury to the environment, certainly on a global scale, and possibly even locally to the Province. Finally, there is also the potential for other Earth surface processes, such as increased runoff of soil/plant material, to elevate the flux of Hg to sediments on a local or possibly global scale. Although such an input may also simply reflect intense recycling of originally volcanic mercury, and global-scale increases in weathering rates are typically associated with times of large-scale volcanism. In short, it is essential to study multiple sites covering different facies in order to gain a clearer overview of the global mercury cycle, which will not necessarily be represented within every single sedimentary record or even an individual large geographical region. Future work should focus on exploring the many nuances of sedimentary mercury in order to better understand the element as a proxy for volcanism, investigating how mercury isotopes can be best employed in understanding past mercury perturbations, and confirming whether those disturbances were global or regional in extent during different Phanerozoic events by expanding the geographical distribution of stratigraphic records studied for mercury.

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