Mercury evidence for pulsed volcanism during the end-Triassic mass extinction

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ABSTRACT

The Central Atlantic Magmatic Province (CAMP) has long been proposed as having a causal relationship with the end-Triassic extinction event (~201.5 Ma). In North America and northern Africa, CAMP is preserved as multiple basaltic units interbedded with uppermost Triassic to lowermost Jurassic sediments. However, it has been unclear whether this apparent pulsing was a local feature, or if pulses in the intensity of CAMP volcanism characterized the emplacement of the province as a whole. Here, six geographically widespread Triassic–Jurassic records, representing varied paleoenvironments, are analyzed for mercury concentrations and mercury/total organic carbon (Hg/TOC) ratios. Volcanism is a major source of mercury to the modern environment. Clear increases in Hg and Hg/TOC are observed at the end-Triassic extinction horizon, confirming that a volcanically induced global mercury-cycle perturbation occurred at that time. The established correlation between the extinction horizon and lowest CAMP basalts allows this sedimentary mercury excursion to be stratigraphically tied to a specific flood basalt unit for the first time, strengthening the case for volcanic
mercury as the driver of sedimentary Hg/TOC spikes. Additional Hg/TOC peaks are also documented between the extinction horizon and the Triassic–Jurassic boundary (separated by ~200 kyr), supporting pulsatory intensity of CAMP volcanism across the entire province and providing the first direct evidence for episodic volatile release during the initial stages of CAMP emplacement. Pulsatory volcanism, and associated perturbations in the ocean–atmosphere system, likely had profound implications for the rate and magnitude of the end-Triassic mass extinction and subsequent biotic recovery.

Significance Statement

The end of the Triassic Period (~201.5 million years ago) witnessed one of the largest mass extinctions of animal life known from Earth history. This extinction is thought to have coincided with and been caused by one of the largest known episodes of volcanic activity in Earth’s history. This study examines mercury concentrations of sediments from around the world that record this extinction. Mercury is emitted in gaseous form during volcanism, and subsequently deposited in sediments. We find numerous pulsed elevations of mercury concentrations in end-Triassic sediments. These peaks show that the mass extinction coincided with large-scale, episodic, volcanism. Such episodic volcanism likely perturbed the global environment over a long period of time and strongly delayed ecological recovery.

1. Introduction

The end of the Triassic Period was marked by a major mass extinction event (~201.5 Ma; e.g. 1, 2), one of the five largest environmental perturbations of the Phanerzoic Eon. Significantly increased extinction rates of marine fauna, and major turnovers in terrestrial vegetation and vertebrate groups, have been well documented (e.g. 3–8). The end-Triassic mass extinction predated the onset of the Jurassic by ~100–200 kyr, as defined by the first occurrence of the Jurassic ammonite species Psiloceras spelae (9). The sedimentary record of the extinction correlates with a large (up to 6%) negative excursion in organic-carbon isotopes ($\delta^{13}$C_TOC: Figure 1A), indicative of a severe carbon-cycle perturbation coincident with the biotic crisis (e.g. 10–13). Moroccan strata that record this global carbon-cycle perturbation are transected by the lowest documented flows of the Central Atlantic Magmatic Province (CAMP). Consequently, the end-Triassic
extinction has been postulated as precisely coincident with the onset of known CAMP volcanism (e.g. 2, 12, 14, 15).

CAMP represents the most aerially expansive continental Large Igneous Province (LIP) known on Earth, consisting of volumetrically large-scale flood-basalt sequences covering at least \(7 \times 10^6\) km\(^2\) across four continents and both hemispheres (Figure 2A: ref 16). In North America and Morocco, CAMP basalts are interbedded with continental sediments that have precise temporal constraints and are stratigraphically well correlated with marine sedimentary records (Figure 1A: refs 12, 14). The apparent episodic emplacement of CAMP basalts over several 100 kyr, at least in Morocco and North America, is a key feature of this LIP.

The oldest known CAMP basalts are the lower Moroccan unit (termed the Lower Formation in the High Atlas and the Tasguint Formation in the Argana basin: 17, 18). This unit is overlain by two further Moroccan basalt units: the Middle and Upper Formations in the High Atlas (the former named the Alemzi Formation in Argana: 17, 18). For clarity, the High Atlas names are used henceforth in this study. These three basalt groups are interbedded with sedimentary deposits. A fourth extrusive unit, the Recurrent Formation, is locally preserved higher in the Moroccan sequence, with much thicker sediments between it and the Upper Formation (Figure 2B; 14, 17, 19). These four basalt units are defined and correlated on the basis of distinct igneous geochemistry. Based on geochemical correlation with North American CAMP units, which are temporally constrained by astrochronological and radioisotopic geochronology, the Moroccan Lower–Upper Formations are thought to have erupted in quick succession, with several 100 kyr then passing before the eruption of the Recurrent Formation (2, 14).

At least three major CAMP units are documented in North America: the Orange Mountain, Preakness, and Hook Mountain Basalts in the Newark Basin (20), with time-equivalent basalts known from other North American Basins. The Orange Mountain Basalt overlies thin, usually lacustrine, sediments deposited above the extinction horizon, and is thought to have been extruded 14–20 kyr after that event (2, 14, 21). Radioisotopic dating has demonstrated that the Preakness Basalt and Hook Mountain Basalt were emplaced 270 kyr and 620 kyr later, respectively (2). The Newark Basin Orange Mountain and Hook Mountain basalts have been geochemically established as equivalent to the Moroccan Middle and Recurrent Formations, respectively (14); the Preakness Basalts have no known Moroccan counterparts (2, 14). Consequently, North American and Moroccan records suggest that CAMP was emplaced in at least three major pulses of basalt extrusion over ~700 kyr, with the products of the first major pulse further divisible into at least three or four geochemically and stratigraphically distinct units. Thus, a total of at least six CAMP units are documented in...
Morocco/North America (Figure 2B), which were relatively close to one another at the end of the Triassic Period. However, more geographically dispersed CAMP basalts are also known from southern Europe, Brazil, and elsewhere in western Africa, with potentially different temporal relationships with the end-Triassic extinction and the dated CAMP flows (16). Therefore, it is not clear whether the intensity of CAMP volcanism was pulsatory across the entire province, or whether the apparent pulsing recorded in North America and Morocco was a local feature of the much larger scale LIP.

Proxy records of volcanic volatiles can aid in reconstructing the history of CAMP volcanism and its impacts. Analyses of pedogenic carbonates suggest increases in atmospheric $p$CO$_2$ following emplacement of each of the Newark CAMP basalts (CAMP pulses 3, 5, and 6 in Figure 2B), supporting a pattern of globally incremental emplacement (22). However, the effect of local processes, such as diagenesis, on this record cannot be ruled out. Reconstructions of $p$CO$_2$ based on stomatal indices (albeit at low temporal resolution) show no such pulsing during the Triassic–Jurassic transition (6, 23). Nor is there yet evidence of episodic CO$_2$ increases associated with the early Moroccan CAMP pulses (CAMP pulses 1, 3, and 4 in Figure 2B) that were extruded coincident with, and in the immediate aftermath of, the extinction event.

Here, the volatile emissions and ocean-atmosphere impact of CAMP volcanism is investigated by analysis of sedimentary mercury (Hg) concentrations across multiple Triassic–Jurassic sedimentary archives. Volcanism is known to be a major natural source of mercury, emitting it as a trace volcanic gas (24). Gaseous elemental Hg has a typical atmospheric residence time of 0.5–2 year (25), allowing the element to be globally distributed before being drawn down and eventually deposited in sediments. Several Phanerozoic events have previously been linked to approximately coeval LIPs through documented increases in sedimentary mercury concentrations, including the end-Permian and end-Cretaceous extinctions and Toarcian Oceanic Anoxic Event (e.g. 26–29). Importantly, sedimentary drawdown of Hg is typically achieved via organic matter (30, 31), although sulphides and clays may also play a role (32–34). Consequently, sedimentary Hg concentrations are typically normalized against total organic carbon (TOC) to account for the effect on Hg drawdown by changes in organic matter deposition rates when looking for evidence of an elevated supply of Hg to the environment (26).

In addition to interrogating the pulsatory history of CAMP volatile emissions, mercury analysis of uppermost Triassic sediments also provides a unique opportunity to test the assumption that Hg-enriched sediments were deposited precisely coincident with the eruption of LIP basalts. There are excellent age constraints on numerous end-Triassic records (including those containing CAMP basalts), and the precise
correlation between the end-Triassic extinction horizon and lowest Moroccan CAMP basalt (the Lower 119 Formation) is well established. Such temporal constraints may allow some Hg/TOC peaks in uppermost 120 Triassic sediments to be directly correlated with specific CAMP basalt units. This direct association between 121 Hg/TOC excursions and specific basalt flows has not been possible for other events due to the poor 122 preservation, or limited stratigraphic control relative to the sedimentary record, of many LIPs.

A recent study on the Triassic–Jurassic boundary section at New York Canyon (Nevada, USA) 123 showed an abrupt increase in Hg concentrations and Hg/TOC ratios correlated with the negative excursion in 124 $\delta^{13}C_{\text{org}}$ that marks the end-Triassic extinction horizon (35). These Hg excursions were attributed to volcanic 125 processes operating during the emplacement of CAMP. Here, the New York Canyon results are greatly 126 expanded by analyzing six further sedimentary records from around the world, to test whether the end-Triassic 127 mercury perturbation was a global phenomenon. The possibility of multiple episodic peaks in sedimentary 128 mercury is also investigated, to examine whether the documented pulsatory nature of CAMP emplacement 129 occurred province-wide, or was limited to specific areas of the LIP. The synchrony of any Hg excursions with 130 respect to the earliest CAMP flows is also assessed using the established stratigraphic correlation between the 131 end-Triassic extinction horizon and the stratigraphically lowest known CAMP basalts.

2. Study areas

End-Triassic records of both marine and terrestrial environments are known from a number of 135 locations around the world (Figure 1 in ref 36). In this study, the mercury records from six geographically 136 widespread sections are presented, representing a variety of marine and terrestrial paleoenvironments (Figure 137 2A): St Audries Bay (UK: restricted shallow-marine), Kuhjoch (Austria: open shallow-marine), Arroyo Malo 138 (Argentina: back-arc shallow marine), Astartekløft (Greenland: fluvio-deltaic), Partridge Island (Canada: 139 lacustrine), and Igounane (Morocco: evaporitic-lacustrine). See Supplementary Information S1 for details on 140 all the studied sections, and Supplementary Figure S2 for a full correlation among all the above sections and 141 other end-Triassic records.

3. Results and Discussion

3.1. Mercury as a recorder of CAMP volcanism

Clear excursions in Hg/TOC ratios and/or Hg concentrations are observed at five of the six studied 147 locations (St Audries Bay, Kuhjoch, Arroyo Malo, Astartekløft, and Partridge Island). The onsets of these
excursions are stratigraphically coincident with the globally observed $\delta^{13}$C negative excursion that marks the extinction horizon (Figure 3). Four sections also record additional peaks in Hg/TOC above that level. Crucially, at all sites where Hg has been normalized to TOC, the Hg/TOC peaks result from elevated Hg concentrations rather than decreased TOC content (see Supplementary Figure S4). For Partridge Island and Igounane sediments, Hg/TOC ratios were deemed unreliable due to the very low TOC content (typically below analytical uncertainty) in sedimentary samples. Consequently, Hg signals at these two locations are presented without normalization to TOC. The mercury trends generated in this study are also compared to the existing New York Canyon record (35) in Figure 3, which appears to have a subtly different trend in sedimentary Hg/TOC increase, potentially resulting from atmospheric or local sedimentological processes.

The observed peaks in sedimentary Hg and Hg/TOC strongly suggest that a perturbation to the global mercury cycle took place during the Triassic–Jurassic transition, beginning coincidentally with the end-Triassic extinction. The absence of a recorded mercury perturbation in sediments at Igounane is interpreted to result from their deposition being below the oldest known CAMP flows (thus at a time preceding the onset of CAMP basalt extrusion). A lack of change in terrestrial spores from Argana sediments below the CAMP basalts further suggest that these sediments were deposited prior to the end-Triassic extinction (14), and thus before the onset of CAMP volcanism.

The correlation between Hg excursions and the extinction horizon in the other five studied records is strongly suggestive of a perturbation to the global mercury cycle at that time. Variations in marine redox during the extinction may have influenced the marine Hg cycle, but the records at both Kuhjoch and Arroyo Malo do not show a consistent stratigraphic correlation between the observed Hg/TOC peaks and lithological or geochemical evidence for redox changes (37, this study). Additionally, the mercury excursions preserved in the terrestrial records from Astartekløft and Partridge Island could not have been caused by changes in the oceanic mercury inventory. Consequently, an atmospheric Hg perturbation is the most plausible cause.

The Hg perturbation also coincided with the established onset of an increase in atmospheric $\rho$CO$_2$, based on Hg/TOC and stomatal density records from Astartekløft (23: Supplementary Figures S2 and S5). This correlation suggests a geologically simultaneous increase in atmospheric Hg and CO$_2$, plausibly originating from magmatic degassing during CAMP emplacement. Emissions of both gases could also result
from thermogenic gas release from kerogen in subsurface organic-rich sediments intruded by (CAMP-associated) sills. Thermogenic emissions have been previously suggested as a key contributor to LIP atmospheric perturbations (e.g. 38, 39). Thermogenic volatiles also explain the observed negative excursion in \( \delta^{13}C \) at the extinction horizon more satisfactorily than magmatic carbon emissions (40). However, peaks in Hg/TOC stratigraphically above the extinction horizon are not marked by distinct negative excursions in \( \delta^{13}C \). Consequently, there is less evidence for these later mercury perturbations resulting from thermogenic emissions, and magmatic Hg emissions are a more probable cause.

Additional evidence for a volcanic origin of the perturbation to the global Hg cycle during the end-Triassic extinction comes from the established correlation between the lowest known CAMP flow (the Lower Formation in the High Atlas) and the extinction horizon. This correlation allows the Hg/TOC increase at the extinction horizon to be precisely stratigraphically matched with that particular unit of CAMP (Figure 1B: see also Supplementary Figure S2 and refs 14, 15). Consequently, it is highly likely that volcanic Hg associated with this lowest CAMP flow contributed to the global mercury perturbation during the end-Triassic extinction. The ability to correlate stratigraphically a Hg excursion directly with an individual CAMP basalt flow greatly strengthens the use of this element as a proxy for volcanism.

### 3.2. The pulsatory release of magmatic volatiles

In addition to the mercury excursion at the extinction horizon, four sections record distinct peaks in sedimentary Hg/TOC higher up in the stratigraphy. These higher peaks are most clearly distinct at Kuhjoch and Arroyo Malo, but may also be recorded at New York Canyon and St Audries Bay (Figure 3). Pulsatory CAMP volcanism has been inferred from the stratigraphic record of CAMP basalts in North America and Morocco (17–20). However, these lithological records only prove an apparent pulsing of CAMP basalts in specific locations of what is a much larger scale province. Episodic volcanism across the entire extent of CAMP has been inferred from pedogenic carbonate \( p\text{CO}_2 \) reconstructions (22), but these concretionary carbonates can be impacted by local (diagenetic) processes. The observed pulsatory signal of Hg perturbations in multiple, globally distributed, sedimentary records across the Triassic–Jurassic transition provides independent evidence that the intensity of CAMP volcanism, and likely CAMP emplacement, occurred in a non-continuous manner.
Although the mercury records shown in Figure 3 do show excursions in Hg/TOC in Jurassic strata and support the continuation of CAMP volcanism into the Jurassic, most of the Hg/TOC peaks are documented between the end-Triassic extinction horizon and the Triassic–Jurassic boundary. The data (Figure 3) suggest that at least two, and possibly three, major volcanic episodes occurred between the extinction and the beginning of the Jurassic (as defined by ammonite biostratigraphy). This interval is estimated as lasting only 100–200 kyr based on U–Pb geochronology and astrochronology (e.g. 1, 2, 41, 42). Ar–Ar and U–Pb geochronology suggests that the oldest three Moroccan basalt units (the Lower, Middle and Upper Formations: units 1, 3, and 4 in Figure 2B) and the first of the major North American CAMP pulses (units 2–4 in Figure 2B) were likely all emplaced during this interval (2, 14, 17). It is possible that the observed three pulses in Hg/TOC observed in strata between the extinction horizon and the Triassic–Jurassic boundary may directly relate to atmospheric volatile release from the first three CAMP basalt units in the Moroccan Argana Basin, and their North American equivalents. However, further correlative work is needed to confirm such a hypothesis. The later North American CAMP basalts, and the Moroccan Recurrent Formation (units 5 and 6, Figure 2B), were emplaced 300–600 kyr after the extinction (2, 14, 21 and references therein). Thus, the Hg/TOC peaks between the extinction horizon and the Triassic–Jurassic boundary cannot be related to these later flows.

The observed record of multiple pulses in Hg/TOC is direct evidence that volatile release associated with CAMP volcanism was also pulsatory, likely including episodic emissions of carbon, sulfur, and mercury. Moreover, pulsatory emissions are shown to have occurred throughout the first 100–200 kyr immediately following the end-Triassic extinction, in addition to the documented later pulses of volcanic emissions (22). Consequently, the large increase in atmospheric \( \rho \text{CO}_2 \) during the extinction event may have arisen as a series of episodic carbon-cycle perturbations to the atmosphere (and subsequently the ocean). Pulsatory perturbations of the ocean–atmosphere system caused by episodic volcanic events (and release of volatiles) associated with CAMP may explain the documented prolonged period of ecosystem deterioration and delayed recovery of benthic fauna during the emplacement of CAMP (35, 43–45).

4. Conclusions
Investigation of the global history of CAMP volatile emissions is important for understanding the
development of this igneous province and its potential environmental impact. This study demonstrates that the
mercury cycle was perturbed on a global scale during the Triassic–Jurassic transition. Mercury excursions are
recorded in five of the six sections studied; the one section with no record of Hg enrichment was likely
deposited prior to the onset of CAMP volcanism. The onset of Hg enrichment occurred synchronously across
the globe, coincident with the end-Triassic extinction and associated global carbon-cycle perturbation. The
presence of Hg/TOC excursions in sedimentary records of terrestrial and marine paleoenvironments, across
both hemispheres, indicates that atmospheric mercury concentrations likely increased substantially. This
atmospheric perturbation probably resulted from the emplacement of CAMP and the associated large-scale
emission of magmatic volatiles, and potentially thermogenic volatiles from intruded country rock (including
mercury). The appearance of a global Hg excursion at the end-Triassic extinction horizon and multiple
Hg/TOC peaks between it and the Triassic–Jurassic boundary is further evidence that pulses in the intensity of
CAMP volcanism (and associated volatile release) were not limited to North America and Morocco, but
representative of the entire province. The direct correlation between the oldest preserved flows of CAMP and
the end-Triassic extinction horizon allows the earliest pulse of elevated mercury in the sedimentary record of
this time to be linked directly with these basalts. This correlation supports a volcanic origin for the increased
mercury abundances, and represents the first time that a globally observed mercury excursion has been tied to
a specific basalt unit from a Large Igneous Province. The recording of multiple Hg/TOC excursions between
the extinction horizon and the Triassic–Jurassic boundary highlights that the initial stages of CAMP
emplacement were marked by multiple episodes of volcanic volatile release. Repeated volcanically driven
perturbations of the ocean-atmosphere system in the 100–200 kyr during and immediately following the end-
Triassic extinction may have had important implications for the biospheric impact of CAMP.

**Methods**

Mercury analysis was undertaken on the RA-915 Portable Mercury Analyzer with PYRO-915
Pyrolyzer, Lumex, at the University of Oxford (26). Where previous TOC determinations were not available,
new data were measured using either a Strohlein Coulomat 702 (46) or Rock-Eval VI (47) at the University of
Oxford. δ^{13}C_{org} analyses were performed on decarbonated Arroyo Malo samples (prepared at the University of
Oxford) with a Thermo Scientific Flash 2000 HT Elemental Analyzer (EA) coupled to a Thermo Scientific
MAT253 isotope-ratio mass spectrometer via a Confl o IV open-split interface at the Stable Isotope Laboratory at the Open University (Milton Keynes, UK). For full method details, see Supplementary Information S1.

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**Figure Captions:**

**Figure 1:** A: Stratigraphic correlation of the end-Triassic extinction with the Moroccan Lower Formation CAMP basalt. Argana lithology, carbon-isotope, and paleomagnetic data are from 14. Newark lithology and $p$CO$_2$ data are from 22, paleomagnetic and astrochronological data are from 48; carbon-
isotope data from 12; trilete spore data from 49. St Audries Bay lithology and astrochronology are from 42 and 50, biostratigraphy and carbon-isotope data from 10 and 50, trilete spore data from 51, paleomagnetic data from 52 and 53. Stratigraphic correlation of CAMP units between Argana, Newark, and St Audries Bay is based on 12 and 42. Kuhjoch biostratigraphy, lithology, and carbon-isotope data are from 11. The end-Triassic extinction horizon (marked as ETE) and Triassic–Jurassic boundary (marked as TJB) are also shown. B: Example of new Hg/TOC data from this study (Kuhjoch, Figure 3) are shown to stratigraphically correlate with the lowest CAMP basalt unit that intersects the end-Triassic extinction at Argana. See Supplementary Figure S2 for a full stratigraphic correlation of end-Triassic records.

Figure 2: A: Paleogeographic reconstruction of the end-Triassic world, with the modern continents overlain. The locations of the six studied sections are indicated (A: St Audries Bay, UK; B: Kuhjoch, Austria; C: Arroyo Malo, Argentina; D: Astartekløft, Greenland; E: Partridge Island, Canada; F: Igounane, Morocco). The New York Canyon section in Nevada, USA (G: note different color) studied by 35, and the Central Atlantic Magmatic Province (CAMP) are also shown (based on Figure 1 from ref 2). B: Summarized composite stratigraphy of Moroccan and North American CAMP basalts, following the stratigraphic relationships and ages (in Myrs) from 2 and 14. The Hickory Grove basalt is included with the Preakness due to their geochemical similarity. The age of the end-Triassic extinction (marked as ETE) is also indicated.

Figure 3: Comparison of Hg/TOC data from St Audries Bay, Kuhjoch, Arroyo Malo, Astartekløft, and New York Canyon (35), and Hg data from Partridge Island and Igouane (where TOC contents were below error). Carbon-isotope data, biostratigraphy (ammonite First Appearance), lithology, and magnetostratigraphy are also shown to allow stratigraphic correlation of the end-Triassic extinction horizon (marked as ETE) and Triassic–Jurassic boundary (marked as TJB). Lithological data is from St Audries Bay (10), Kuhjoch (11), Arroyo Malo (54), Astartekløft (10), Partridge Island (this study; shown in figure), Argana (14), Igouane (this study; shown in figure) and New York Canyon (35). Carbon-isotope and biostratigraphic data is from St Audries Bay (10 and 50), Kuhjoch (11), Arroyo Malo (54 and this study), Astartekløft (10), Partridge Island (this study), Argana (14) and New York Canyon (35). Line M indicates the magnetostratigraphic correlation, below the extinction horizon, between St Audries Bay, Partridge Island, Argana, and Igouane. The gray shading illustrates the
stratigraphic gap between line M and line ETE. Magnetostratigraphic data is from St Audries Bay (52 and 53), Partridge Island (55) and Argana (14). Note for Astartekløft the expanded horizontal scale, and the gaps in data due to sand beds with negligible TOC. Full $\delta^{13}$C, Hg, TOC and Hg/TOC data for each individual section are reported in the Supplementary Tables S3 and Supplementary Figure S4. (For interpretation of the references to color in this figure, the reader is referred to the web version of this article).
CAMP BASALT STRATIGRAPHY

End-Triassic World

- Recurrent Formation/
  Hook Mountain Basalt
- Preakness Basalt/
  Hickory Grove Basalt (5A)
- Upper Formation/
  York Haven Intrusive
- Middle Formation (Alemzi Fm.)/
  Orange Mountain Basalt
- North Mountain Basalt
- Lower Formation (Tasguint Fm.)

Sediments deposited between basalt units
Lithology*: CAMP basalt Mudstones (color indicates the color of the mud in outcrop)

*Applicable only at Partridge Island, Argana, and Igounane. Other lithological data sourced from previously published works (see figure caption).

From Thibodeau et al. (2016)