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Shallow ocean oxygen decline during the end-Triassic mass extinction

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ABSTRACT

The end-Triassic mass extinction (ETME) was associated with intensified deep-water anoxia in epicontinental seas and mid-depth waters, yet the absolute oxygenation state in the shallow ocean is uncharacterized. Here we report carbonate-associated iodine data from the peritidal Mount Sparagio section (Southern Italy) that documents the ETME ($\sim 200~\text{Ma}$) in the western Tethys. We find a sharp drop in carbonate I/(Ca + Mg) ratios across the extinction horizon and persisting into the Early Jurassic. This records local dissolved oxygen and iodate decline in the near-surface ocean of low-latitude Tethys due to the development of depleted oxygen concentrations. Consequently, during the ETME even shallow-water animals, such as the megalodonts seen at Mount Sparagio, were likely the victims of oxygen-poor conditions. The shallow ocean deoxygenation coincides with the synchronous spread of deeper anoxic waters and widespread anoxic deposition on continental shelves and slopes. An upwards expansion of the mid-water oxygen minimum zone in the latest Triassic shoaled the oxycline and triggered a major marine crisis.

1. Introduction

Deoxygenation of the upper ocean is a threat to modern marine ecosystems due to global warming and has been observed as a consequence of past warming events at numerous episodes in Earth history (Jenkyns, 2010; Breitburg et al., 2018; Lu et al., 2018; Oschlies, 2021; Song et al., 2021). The end-Triassic mass extinction (ETME; ~200 Ma) is cotemporaneous with a prominent expansion of marine anoxia that is closely linked to the hothouse climate associated with Central Atlantic Magmatic Province (CAMP) volcanism (Ruhl et al., 2011; He et al., 2020). Existing data demonstrate that strengthened anoxic conditions during the ETME were prevalent across many semi-enclosed basins of Europe (Luo et al., 2018; Beith et al., 2021; Fox et al., 2022) and the middepth waters of open ocean (Jost et al., 2017; He et al., 2020), whilst the pelagic deeper ocean remained fully-ventilated (Wignall et al., 2010; Fujisaki et al., 2020). Nevertheless, it is largely unknown with respect to

the absolute redox state in the upper water column of shallow marine locations across the Triassic–Jurassic (T–J) transition, as the shallow marine ecosystem accounts for the vast majority of aerobic marine organisms and their habitats. Hence, filling this knowledge gap is crucial for tracking the anoxic/hypoxic-extinction link across the ETME.

Carbonate-associated iodine is widely used as a redox proxy to constrain oxic to hypoxic conditions (dissolved oxygen content $[O_2] = \sim 10$ to $\sim 100~\mu mol/kg$) in the upper water column (Lu et al., 2010; Hoogakker et al., 2018; Lu et al., 2018; Pohl et al., 2021). The modern ocean has a relatively uniform concentration of iodine due to its long residence time of $\sim 300~kyr$. However, the speciation of iodine in the local water column is dependent upon a redox-sensitive pathway between iodate (IO₃⁻) and iodide (I⁻) (Luther and Campbell, 1991; Rue et al., 1997). Under low oxygen conditions iodate is readily reduced to iodide (Luther and Campbell, 1991; Rue et al., 1997), and will convert back to its oxidized form in the presence of abundant dissolved oxygen.

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The kinetics of iodide oxidation are slow relative to those of reduction (Chance et al., 2014; Hardisty et al., 2020, 2021). Only iodate is readily incorporated into the calcite lattice, substituting for the ${\rm CO_3}^{2-}$ ion (Podder et al., 2017; Feng and Redfern, 2018), which allows the concentrations of this structurally substituted iodine to directly reflect water column redox state during the deposition of carbonate (Lu et al., 2010). Thus, the utilization of I/(Ca + Mg) in ancient carbonates affords an opportunity to trace in situ [O₂] variations through time in shallow marine environments. Because of the slow oxidation kinetics of iodide, the I/(Ca + Mg) proxy can also incorporate a contribution from regional mixing between adjacent water masses (Lu et al., 2020; Hardisty et al., 2021). Further, carbonate I/(Ca + Mg) can also qualitatively track depth of the oxycline in the water column where the [O2] decreases more sharply. Carbonate deposited within the shallow ocean realm reflect surface or near-surface seawater dissolved iodate, and is considered to be imparted by the expansion or contraction of a proximal oxygen minimum zone (OMZ) or by fluctuations in the depth of the oxycline (Zhou et al., 2016; Lu et al., 2018).

Here we present a high-resolution I/(Ca + Mg) record from an upper Rhaetian–lower Hettangian peritidal carbonate succession (Mount Sparagio section, Southern Italy) that was located in the western Tethys (see geochemical data in Table S1). We show the first evidence of a prominent decline in dissolved oxygen levels in the shallow ocean across

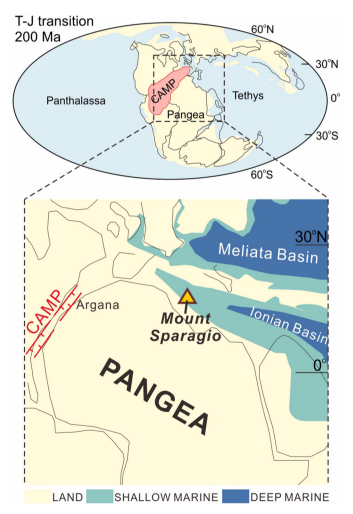


Fig. 1. Paleogeographic map for the shallow marine Mount Sparagio section of western Tethys during the Triassic–Jurassic (T–J) transition. This map is based on Todaro et al. (2018). Yellow triangle indicates the location of the studied Mount Sparagio section. CAMP: Central Atlantic Magmatic Province. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

the ETME. Low oxygen conditions appear to have persisted into the early Hettangian shallow ocean.

2. Palaeogeography and stratigraphy

The Mount Sparagio (MS) section from western Sicily (Southern Italy) was located at low-latitude of ~15°N in a shallow carbonate platform of the western Tethys (Fig. 1) (Todaro et al., 2017). The studied succession records upper Rhaetian to lower Hettangian peritidal carbonates (Todaro et al., 2017; He et al., 2020). The subtidal facies of Upper Triassic strata are characterized by the occurrence of megalodontids, calcareous algae and a benthic foraminifera association with Triasina hantkeni, Aulotortus sp., Auloconus permodiscoides indicative of a Rhaetian age (Todaro et al., 2017). The extinction horizon is marked by the last occurrence of megalodontids and the characteristic Rhaetian benthic foraminifer Triasina hantkeni and occurs immediately below a thin oolitic limestone that is unique to this level at ~200 m height (Fig. 2). The bloom of the typical Jurassic species Thaumatoporella parvovesiculifera associated with only rare Aeolisaccus sp., at a short distance above this oolitic horizon is a further evidence of the extinction interval, and is followed by a slow recovery of the Jurassic benthic community in the earliest Hettangian (Todaro et al., 2018; He et al., 2020).

3. Material and methods

A total of 49 well-preserved micritic limestone samples were measured for carbonate-associated iodine concentration. Bulk carbonate rocks were cut into small rock cubes to trim the weathered crust. This was followed by grinding to a fine powder using a TEMA laboratory agate disc mill. Around 20 mg of sample powder was first rinsed by ultrapure water three times and dried. The protocol of carbonateassociated iodine extraction is a modification after the work of Lu et al. (2010). For each sample approximately 5 mg of cleaned dry powder was then weighed and treated with 3% (ν/ν) nitric acid using an ultrasonic bath at room temperature. This carbonate leaching step was completed within 15 min to minimize the potential for iodine escape at low pH conditions. The samples were centrifuged and the supernatant containing the leachate was mixed with a 0.5% (v/v) HNO₃, 0.5% (v/v) ammonium hydroxide, 3% (v/v) methanol solution. The ammonium hydroxide is required to stabilize the iodine in solution and minimize sample washout times during inductively coupled plasma mass spectrometer (ICP-MS) analysis.

Methanol is added to the solution as a source of carbon to promote a signal enhancement for iodine measurements. Carbon has been known to enhance the signal for iodine on ICP-MS for some time and several studies have systematically investigated this effect (Grindlay et al., 2013; Ariga et al., 2019). The effect is not fully understood but is attributed to charge transfer reactions occurring in the plasma of the ICP-MS which allow difficult to ionize elements such as iodine to be ionized and thus detected by the mass spectrometer. Studies have shown that the source of carbon is not important to the effect (Grindlay et al., 2013). Methanol was chosen for this work due to its ready availability and simplicity to work with. Enhancement of the iodine signal with the addition of carbon has been found up to 2.5× that without (Ariga et al., 2019). No increase in background signal due to the addition of carbon for iodine was observed in this work.

Analysis of these solutions must be complete within 24 h of dissolution. An aliquot was measured for concentrations of Ca and Mg using a ThermoFisher iCAP 7400 radial inductively coupled plasma optical emission spectrometer (ICP-OES). Samples and calibration standards were internally standardized using 1 mg L $^{-1}$ Y and Lu. A further aliquot was analysed for iodine using a ThermoFisher iCAP Qc ICP-MS in the Aqueous Analytical Facility, University of Leeds. Samples and calibration standards were internally standardized using 5 mg L $^{-1}$ Te and the standards matrixed matched to the samples by addition of 50 mg L $^{-1}$ Ca. The instrumental precisions for Ca, Mg and I are better than 1%. The

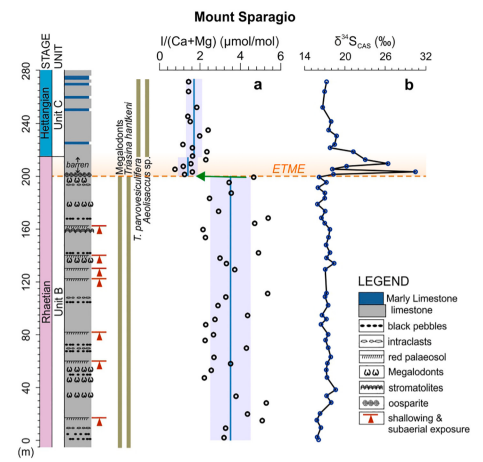


Fig. 2. I/(Ca + Mg) and $\delta^{34}S_{CAS}$ profiles of Mount Sparagio section from Latest Triassic to Early Jurassic. Stratigraphic depth (m) and the lithological log are presented alongside the stages, with stratigraphic units of Todaro et al. (2018). Carbonate-associated sulfate $\delta^{34}S_{CAS}$ data are from He et al. (2020). The orange horizontal dash line and shadowed field indicate the end-Triassic mass extinction (ETME) at this location. The green arrow indicates the sharp drop in I/(Ca + Mg) across the ETME. The occurrence of red palaeosols in Unit B indicate subaerial exposure of the site. The vertical blue lines and light purple bands in a show average I/(Ca + Mg) values with uncertainty windows of 1σ standard deviations pre-, during and post-ETME. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

ICP-MS was tuned for highest sensitivity to iodine. Repeated measurements of the reference material JCp-1 (coral, *Porites* sp.) yielded a I/Ca of 4.52 \pm 0.14 µmol/mol, which can also be expressed as a carbonate-associated iodine concentration of 5.48 \pm 0.17 µg/g, n=14 (see Table S2 for JCp-1 measurement data), comparable to the published acid-leachable iodine concentration of 5.43 \pm 0.07 µg/g, n=8 (Lu et al., 2010) and the certified total measure of iodine concentration of 5.5 \pm 0.2 µg/g, n=5 (Chai and Muramatsu, 2007).

4. Results and discussions

We first rule out potential lithological and diagenetic controls on the I/(Ca+Mg) dataset (Table S1). Samples are dominantly micritic limestone with uniform $CaCO_3$ contents mostly above 90% and lean in organic matter (He et al., 2020), suggesting minimal lithological control on the variations of I/(Ca+Mg) or contamination by non-carbonate phase and organic-bounded iodine (Glock et al., 2019). Mn/Sr ratios in carbonates have been suggested as a measure of the degree of post-

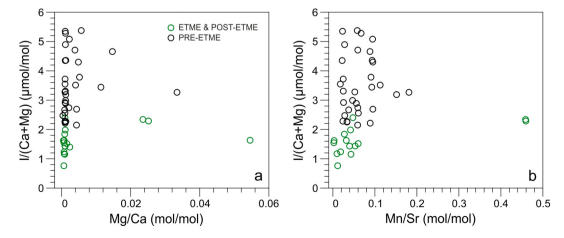


Fig. 3. Correlation between I/(Ca + Mg) molar ratios and elemental molar ratios of carbonates from Mount Sparagio section. a Cross-plot of I/[Ca + Mg] and Mg/Ca shows no correlation ($R^2 < 0.001$), suggesting no alteration of redox-proxy values from minor dolomitization (Mg/Ca < 0.06 mol/mol). b Cross-plot of I/[Ca + Mg] and Mn/Sr displays no correlation ($R^2 = 0.002$), suggesting minimal diagenetic imprint. Mn and Sr data are from He et al. (2020). ETME: end-Triassic mass extinction.

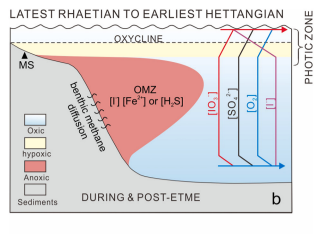
depositional alteration (Brand and Veizer, 1980) while post-depositional dolomitization can be assessed by Mg/Ca ratios. No correlation is observed with the Mg/Ca or Mn/Sr (Fig. 3), which indicates a minor influence of dolomitization or diagenetic imprint on the structurally incorporated iodine. Hence, the carbonate I/(Ca + Mg) changes at the MS section likely indicate primary signals of water column redox conditions (Lu et al., 2010).

Our I/(Ca + Mg) profile from MS section documents higher baseline values with an average of 3.5 \pm 1.0 μ mol/mol (N = 32) throughout the pre-extinction period in the late Rhaetian (Fig. 2a). These new I/(Ca + Mg) data add to the existing low-resolution global data compilation for the T–J transition (~200 Ma) (Lu et al., 2018) when seawater iodine concentration depicts a substantial rise from the low plateau phase (~1 μ mol/mol) in the Permian–Triassic to high levels (~3 μ mol/mol) in the Early Jurassic. Hence, our I/(Ca + Mg) data from MS section validate the previous finding of a long-term increase of oceanic I/Ca ratios from the Triassic to the Jurassic. This change was attributed to a net reduction of oxygen consumption in the upper ocean due to post-ETME radiation of modern-type eukaryotic phytoplankton (Lu et al., 2018).

Nevertheless, the absolute I/(Ca + Mg) values at MS section fluctuate between 2 μ mol/mol and 6 μ mol/mol throughout the pre-ETME late Rhaetian (Fig. 2a). This fluctuating I/(Ca + Mg) record may have resulted from a periodic shallowing of the depositional site as indicated by frequent occurrence of red palaeosols (Fig. 2), which is consistent with facies stacking evidence of shallowing-upward cycles (subtidal-intertidal-supratidal) in these peritidal sediments (Todaro et al., 2017). Alternatively, fluctuation in seawater iodine abundance may have been driven by frequent shallowing of water column oxycline that overlies a proximal OMZ, where dissolved iodate were completely reduced to iodide (Zhou et al., 2016; Lu et al., 2018).

In stark contrast to the pre-extinction interval, I/(Ca + Mg) data record a sharp decline from $\sim 5~\mu mol/mol$ to as low as 0.8 $\mu mol/mol$ (average of 1.4 \pm 0.5 μ mol/mol (N = 6)) in the latest Rhaetian (Fig. 2a), which coincides precisely with the mass extinction horizon. The extinction is also characterized by the sudden disappearance of megalodont bivalves and the foraminifer Triasina hantkeni, and a synchronous positive S-isotope excursion in carbonate-associated sulfate (δ^{34} S_{CAS}; Fig. 2b). Hyperthermal conditions around the T-J transition are thought to have initiated the spread of marine anoxia via increased eutrophication, oxygen consumption and reduction in oxygen solubility in warmer surface waters (He et al., 2020). The large decrease in carbonate I/(Ca + Mg) ratios across the ETME (MS) indicates a depletion of the dissolved iodate pool due to decreased [O₂]. The concurrent positive δ³⁴S_{CAS} shift suggests extensive anoxia and burial of pyrite on continental shelves and slopes at this time (He et al., 2020). The spread of anoxic waters would have been enhanced by oceanic sulfate paucity, which would have suppressed the anaerobic oxidation of methane in the sediments, leading to increased benthic methane flux and net oxygen consumption on the seafloor (He et al., 2020). Anoxic conditions likely expanded from the mid-depth OMZ into shallower waters (Fig. 4b), causing hypoxia and iodate depletion in the surface waters. It is also noteworthy that such correlative redox changes through the shallow to mid-depth waters, as demonstrated by coupled I/(Ca + Mg) and $\delta^{34}S_{CAS}$ trends, are commonly seen in many other prominent anoxic events in the geologic past (Lu et al., 2010; Gill et al., 2011; Owens et al., 2017; Edwards et al., 2018). Thus, the shallowing of the oxycline and upwards invasion of anoxic waters across the ETME is likely to explain the sharp decline in carbonate I/(Ca + Mg) ratios. Indeed, uranium isotope evidence for contemporary deeper water anoxia was found in the adjacent Lombardy Basin (Jost et al., 2017), which likely suggests an upwards expansion of OMZ across the ETME in the area.

The I/(Ca + Mg) record stays at low level (1.7 \pm 0.4 μ mol/mol (N = 11) on average) across the T–J boundary and into the earliest Hettangian (Fig. 2a), indicating continued iodate depletion due to hypoxic conditions in the shallow ocean. However, these post-ETME carbonates exhibit slightly higher iodine concentration compared to those at the



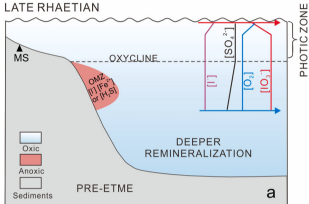


Fig. 4. Schematic diagram of global oceanic redox change through the Triassic–Jurassic transition. Iodine speciation systematics is adapted from Lu et al. (2018). Relative concentrations of redox-sensitive elements: iodate ($\mathrm{IO_3^-}$), iodide ($\mathrm{I^-}$) and sulfate ($\mathrm{SO_4^{2-}}$) are indicated. OMZ: Oxygen Minimum Zone. Anoxic-ferruginous and euxinic water column conditions are demarcated with [Fe²⁺] and [H₂S] respectively. ETME: end-Triassic mass extinction. a Pre-ETME 'Mesozoic-type' water column redox state with well-oxygenated upper ocean and restricted OMZ in mid-depth waters. b Redox state during and post-ETME showing an expanded OMZ and shallowing of the oxycline, leading to hypoxic conditions and iodate depletion in the shallow ocean. Low sulfate concentration in the ocean likely increased the benthic methane flux to the bottom-water and the oxygen demand (He et al., 2020).

major phase of ETME, associated with the recovery of sulfate δ^{34} S (Fig. 2b), which may indicate a lesser degree or persistence of anoxia/ hypoxia. Further, the uppermost Rhaetian-lower Hettangian sedimentology of the MS section records a minor facies transition to a relatively deeper peritidal setting, evident by the thin oolitic limestone bed at a flooding surface immediately above the extinction horizon and higher occurrence of marly limestone in the lower Hettangian (Fig. 2). Hence, it is possible that MS section saw a transition to slightly deeper facies preceding the ETME, resulting in an improved connection to the open ocean. Thus, the shallow ocean hypoxic condition revealed from the post-extinction MS section likely indicates an open marine signal of dysoxia. These commonly hypoxic conditions likely prevailed in the wider shallow ocean in the post-extinction early Hettangian, possibly with an oscillating redox state as seen in the European epicontinental sea (Beith et al., 2021; Fox et al., 2022). The shallow ocean hypoxia during the T-J transition may have also varied spatially although uranium isotope evidence suggests deeper water column anoxia was widespread until the middle Hettangian Stage (Jost et al., 2017). Despite low oxygen levels in the post-ETME shallow ocean, it did not prolong the extinction or inhibit the local recovery of marine communities, although some post-ETME species found at the MS section (alga Thaumatoporella

parvovesiculifera and benthic foraminifers Siphovalvulina sp.) (Todaro et al., 2017, 2018) could be tolerant to oxygen-poor conditions. Furthermore, hypoxic conditions in the earliest Hettangian may have had a limited geographic spread and likely frequently alternated with periods of oxic water column conditions (He et al., 2022), as evidenced by the presence of a diverse benthic fauna in some shallow-water areas during the earliest Jurassic (Atkinson and Wignall, 2020; Wignall and Atkinson, 2020). Some other environmental stressors for the ETME, such as ocean acidification (Fox et al., 2020, 2022), may have receded immediately after the extinction, allowing the shallow ocean ecosystem to recover in the early Hettangian.

In summary, our new I/(Ca + Mg) record from the MS section in Southern Italy, combined with published $\delta^{34}S_{CAS}$ data provide evidence of oxygen decline in the shallowest realms of western Tethys during the ETME. We attribute this major redox shift to a combined consequence of local [O₂] decrease and possible upwards expansion of oxygen depletion from a mid-depth OMZ to the shallow ocean. The onset of the shallow ocean deoxygenation event was clearly synchronous with the loss of megalodont bivalves and foraminifer Triasina hantkeni (Fig. 2), suggesting a cause-and-effect relationship between [O2] scarcity and ecological stress even in exceptionally shallow-water Tethyan areas. This adds to the growing evidence that indicates a global anoxia/ hypoxia-extinction link in a variety of marine settings during the ETME (Jaraula et al., 2013; Jost et al., 2017; Luo et al., 2018; He et al., 2020; Beith et al., 2021; Fox et al., 2022; Kipp and Tissot, 2022). Many similar events through the Mesozoic and Paleogene (e.g., Toarcian oceanic anoxic event, Paleocene-Eocene Thermal Maximum) were accompanied by hyperthermal events and upper ocean hypoxia (Jenkyns, 2010; Lu et al., 2010; Zhou et al., 2014, 2016; Song et al., 2021), which together serve as potential analogues to explore the possible outcomes of ongoing anthropogenic warming.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Acknowledgements

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.gloplacha.2022.103770.

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