Isotopic constraints on ocean redox at the end of the Eocene

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A multi-million-year decrease in global temperatures during the Eocene was accompanied by large reorganisations to ocean circulation, ocean chemistry and biological productivity. These changes culminated in the rapid growth of grounded ice on Antarctica during the Eocene–Oligocene climate transition (EOT), ∼34 million years ago. However, while it is likely that environmental perturbations of this magnitude altered the oceanic oxygen inventory, the sign and magnitude of the response is poorly constrained. We show that euxinic, hydrographically restricted conditions developed in the Austrian Molasse Basin during the EOT. The isotopic compositions of molybdenum and uranium captured by sediments accumulating in the Molasse Basin at this time reveal that the global extent of sulfidic conditions during the EOT was not appreciably different to that of the Early Eocene greenhouse world. Our results suggest that the early Cenozoic oceans were buffered against extreme long-term changes in oxygenation.

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1. Introduction

The spread of deoxygenated zones in the modern ocean threatens the stability of marine ecosystems and nutrient cycling. Predicting the pattern of future deoxygenation trends is difficult because the key drivers – water temperature, organic matter production, and water mass ventilation – may interact with each other in non-linear ways over short and long timescales (Oschlies et al., 2018). In the early part of the Cenozoic Era, the Earth moved from a greenhouse world with temperatures ∼10 °C warmer than pre-industrial during the early Eocene Climatic Optimum (EECO, ∼50–53 Ma) (Cramwinkel et al., 2018; Zachos et al., 2008) to a glaciated world in the early Oligocene, with the inception of an extensive Antarctic ice sheet during the Eocene–Oligocene Transition (EOT, ∼33.7 Ma) (Lear et al., 2008; Katz et al., 2008). This ∼20-million-year climatic transition presents an opportunity to study how the movement of seawater oxygen concentration [O2] interacted over multi-million year timescales.

Published data provide contradictory information about the direction and magnitude of early Cenozoic changes in seawater [O2].

A long-term decline in deep-ocean temperatures from 53–34 Ma should have increased the [O2] of global deepwaters due to the higher solubility of oxygen in colder water. The magnitude of this increase can be roughly estimated from the temperature declines inferred from TEX86, Mg/Ca, δ18O and pollen/spore assemblage proxy data at high northern (Eldrett et al., 2009) and southern (Bijl et al., 2009; Hollis et al., 2009; Liu et al., 2009) latitudes and from gas solubility coefficients, to ∼0.5–3 ml/l (Weiss, 1970). High-latitude surface temperature changes are disproportionately important for deep-ocean [O2] due to the formation of deepwaters that occupy a large volume of the ocean interior. Before ∼50 Ma, deepwater production was concentrated in the Southern Ocean (Hobbein et al., 2012; Nunes and Norris, 2006) where early Cenozoic surface temperature decreases were particularly large (Bijl et al., 2009; Hollis et al., 2009; Liu et al., 2009).

The effect of declining temperatures on deepwater [O2] would have been partly balanced by the remineralisation of exported organic matter into the ocean interior. However, organic matter export was spatially and temporally variable during the Eocene: during the EECO, high ocean temperatures and microbial metabolic rates enhanced the efficiency by which exported organic matter was recycled (Moore et al., 2008; Olivarez-Lyle and Lyle, 2006), possibly leading to lower deep ocean [O2] than at present (Sexton et al., 2011). A second peak in organic matter export after ∼37 Ma
has been attributed to a greater oceanic nutrient inventory and/or rapid ocean overturning (Diekmann et al., 2004; Diester-Haas and Zahn, 1996). The situation is further complicated by patterns of organic matter export in different ocean basins (Moore et al., 2008), temporal changes in the rate of ocean mixing after the EECO (Sexton et al., 2006) and changes in deep ocean ventilation linked to the onset of Northern Component Deepwater formation after ~49 Ma (Hohbein et al., 2012; Sexton et al., 2006). It is thus unclear how organic matter export would have balanced lower temperatures at a global scale to control net \( [O_2] \).

Here we use Mo and U isotope compositions of shales deposited in the euxinic and partially restricted Molasse Basin (MB) to constrain global ocean oxygen distributions at the end of the long-term Eocene cooling trend. The MB was a tectonically controlled narrow seaway that formed north of the Alpine thrust belt as the European Plate was subducted beneath the Adriatic Plate during the Paleogene (Fig. 1). The regional tectonic context is described in detail by Sissingh (1997), Wagner (1998) and references therein. The sedimentary infill is Late-Eocene to Miocene in age and consists of a series of sandstones, limestones, marls and shales (Schulz et al., 2002; Sissingh, 1997). The basin was connected to the Tethys Ocean via a shallow strait to the Valence Basin in the southwest, and to the Para-Tethys Sea via shallow seaways to the north and east (Sissingh, 1997). We focus our study on the Oberschauersberg-1 (OSCH-1) core, in which organic-rich shales of the Eocene–Oligocene age Schöneck Fm are underlain by the Ampfing Sandstone and overlain by the Dynew Marl (Fig. 2). The OSCH-1 sedimentary succession is representative of the MB (Schulz et al., 2002, 2005) (Fig. S1).

2. Methods

2.1. Molybdenum and uranium isotopes

Molybdenum and uranium isotope measurements were performed at The Open University. Powdered mudrocks were accurately weighed and mixed with an aliquot of either a \(^{97}\)Mo–\(^{100}\)Mo double spike or IRMM 3636 U double spike, aiming for a \( \text{Mo}_{\text{sample}}/\text{Mo}_{\text{sample}} \) of \( \sim 0.6 \) or a \( U_{\text{spike}}/U_{\text{sample}} \) of \( \sim 0.04 \). Samples were digested with \( 3:1 \) concentrated HNO\(_3\) and HCl at 150°C to dissolve the non-detrital fraction. Mo was separated from matrix elements with AG1-X8 anion exchange resin using protocols developed by Pearce et al. (2010), as modified by Dickson et al. (2016). Mo isotope ratios were measured in low resolution on a Thermo-Finnigan Neptune MC-ICP-MS, using a CETAC Aridus II desolvolizing nebulizer system for sample introduction, and fitted with nickel sample and H skinner cones. Molybdenum isotope ratios are reported as the per mil deviation from a standard solution and calculated using the following equation:

\[
\delta^{98}\text{Mo} = \left( \frac{^{98}\text{Mo}_{\text{sample}}}{^{95}\text{Mo}_{\text{sample}}} \right) / \left( \frac{^{98}\text{Mo}_{\text{std}}}{^{95}\text{Mo}_{\text{std}}} \right) - 1 \times 1000
\]

All Mo isotope data were determined relative to an in-house Mo solution standard (Fisher Chemicals, ICP Mo standard solution, lot 9920914-150). They have subsequently been normalised to NIST 3134, firstly using a -0.37‰ offset between the two standards (Goldberg et al., 2013), which was confirmed by repeated measurements of NIST 3134 relative to the OU standard over the measurement period (0.37±0.06‰, 2 S.D., n=70), and secondly by adding 0.25‰ following the recommendation of Nägler et al. (2015). External reproducibility of the measurements was assessed by repeatedly processing the USGS SDO-1 shale standard through the full chemical and analytical procedure, which yielded a \( \delta^{98}\text{Mo} \) of 1.07±0.04‰ (2 S.D., n=16), similar to published values (Goldberg et al., 2013). Total procedural blanks measured by isotope dilution were negligible, between 1–4 ng.

U was purified from the sample matrix with established U chromatography protocols using UTEVA resin. \(^{238}\text{U}/^{235}\text{U} \) ratios were measured in low resolution on a Thermo-Finnigan Neptune MC-ICP-MS, set up with nickel jet sampler and X skinner cones, and equipped with a CETAC Aridus II desolvolizing nebulizer system for sample introduction. Uranium isotope ratios are reported as the per mil deviation from CRM-112A standard:

\[
\delta^{238}\text{U} = \left( \frac{^{238}\text{U}_{\text{sample}}}{^{235}\text{U}_{\text{sample}}} \right) / \left( \frac{^{238}\text{U}_{\text{CRM112A}}}{^{235}\text{U}_{\text{CRM112A}}} \right) - 1 \times 1000
\]

External reproducibility was assessed by measuring several standards repeatedly over the analytical period. The solution standard SRM 950A gives a \( \delta^{238}\text{U} \) of 0.03±0.06‰ (n=82, 2 S.D.), similar to previously published values (Andersen et al., 2017). Measurements of seawater yielded a \( \delta^{238}\text{U} \) value of -0.39±0.05‰ (n=8, 2 S.D.), similar to published values (-0.39±0.01‰ (Andersen et al., 2017)); USGS SDO-1 shale yielded a \( \delta^{238}\text{U} \) of -0.09±0.06‰ (n=20, 2 S.D.), similar to published values (Tissot and Dauphas, 2015). Total procedural blanks measured by isotope dilution were 6–12 pg.

2.2. Osmium isotopes

Osmium isotopes were measured at Royal Holloway University of London. Sample powders were digested with a \(^{180}\text{Os}^{185}\text{Re}\) spike solution and 8 ml 3 : 1 HNO\(_3\) and HCl in flame-sealed Carrus tubes at 220°C for 48 hours. After digestion, Os was purified by solvent extraction using CHCl\(_3\) and concentrated hydrobromic acid (Cohen and Waters, 1996), followed by microdistillation in 2M H\(_2\)SO\(_4\)/CrO\(_3\)(VI) (Birck et al., 1997). Re was purified from the digested inverse aqua regia by solvent extraction into iso-amyl alcohol and back extraction into high-purity water (Birck et al., 1997).

Os solutions were loaded onto pre-cleaned and de-gassed high-purity Pt filaments and measured as osmium oxide species using peak-hopping on the axial SEM of an Isotopx Phoenix TIMS. Oxygen-corrected \( ^{187}\text{Os}^{188}\text{Os} \) ratios were mass corrected to a \( ^{192}\text{Os}^{188}\text{Os} \) ratio of 3.083 (Brandon et al., 2006) and blank-corrected. Accuracy and precision were monitored by measuring the DROsS standard (Luguet et al., 2008). \( ^{187}\text{Os}^{188}\text{Os} \) ratios aver-
aged 0.1601 compared to a reference value of 0.16092 (Nanne et al., 2017). Precision was ~2.3% over the course of this study (n=6). Os concentrations were calculated by isotope dilution using mass-bias corrected $^{180}$Os/$^{188}$Os ratios. Procedural blanks were $\sim$0.06 ± 0.07 pg (n=3, 1 S.D.) with an average $^{183}$Os/$^{188}$Os ratio of 0.77.

Re solutions were measured as ~0.5 ppb solutions on a Neptune Plus MC-ICP-MS using a standard introduction system (wet plasma, SIS). To correct for instrumental mass fractionation, sample aliquots were doped with 10 ppb NIST 3636 to obtain a W/Re ratio of ~20. $^{185}$Re, $^{187}$Re, $^{184}$W, $^{186}$W and $^{189}$Os were measured simultaneously using Faraday detectors equipped with $^{101}$W and $^{103}$W resistors (Re), $^{185}$Re/$^{187}$Re ratios were corrected to a $^{186}$W/$^{184}$W ratio of 0.92767 using an exponential mass bias law. Re concentrations were calculated by isotope dilution using corrected $^{185}$Re/$^{187}$Re ratios. A small correction for $^{187}$Os inference on $^{181}$Re was made by monitoring $^{188}$Os. For all samples, this correction was insignificant. External reproducibility for Re concentrations was estimated with separate digestions of well-characterised mudrock standard 00N118 (Dickson et al., 2015), yielding a concentration of 57.05 ppb ± 0.2% (2 S.D., n=5). Procedural blanks were insignificant (4.9 ± 0.9 pg, 2 S.D., n=3). The relative uncertainty on age-corrected $^{187}$Os/$^{188}$Os$_{DP}$ ratios based on analyses of three 00N118 standards in the sample batches analysed for this study is ~1.9%.

### 2.3. Fe-speciation

Highly reactive non-sulfidized Fe pools were determined using well-established sequential leaching procedures at the University of Leeds (Poulton and Canfield, 2005), followed by concentration measurements using Atomic Adsorption Spectroscopy. Pyrite Fe was determined by chromous chloride distillation (Canfield et al., 1986). Replicate analyses gave a RSD of <5% for all extraction steps. Total Fe concentrations used to calculate Fe$_{HR}$/Fe$_T$ ratios were determined by XRF.

### 2.4. Major elements

Major element oxides were measured by X-Ray Fluorescence (XRF) analysis at the University of Edinburgh School of GeoSciences using a Philips PW2404 wavelength dispersive XRF spectrometer. Elemental intensities were corrected for background and known peak overlap interferences, and medium-term instrumental drift was taken into account using a drift normalisation monitor. Calibration lines were produced from a large number of reference materials (ca. 20 for major elements) encompassing a wide range of silicate compositions. Accuracy and precision for most elements in the analysed rock standards are better than ±3% and 1% RSD respectively.

### 2.5. Minor elements

For trace element analysis by ICP-MS, sample and standard rock powders were prepared in a clean room environment using Teflon-distilled reagents and ultra-pure water (18.2 MΩ). Rock powders were digested in 4:1 concentrated HF/HNO$_3$, dried and refluxed in 5 M HNO$_3$; these solutions were rinsed into a 125 ml polypropylene bottle and made up to 100 g with ultra-pure water. The
solutions were analysed at the Open University using a standard Agilent 7500 Series ICP-MS. The analyses were carried out in a single batch, including one rock-standard (USGS SDO-1), a procedural duplicate and a blank which were repeatedly analysed for blank and drift correction purposes. In addition, a minimum of six out of eight rock standards (including WS-E, W-2, DNC-1, JB-2, JB-3, BCR-2 BHVO-1 and BIR-1) were analysed to calibrate the instrument. Accuracy and precision, estimated from the repeat analyses of SDO-1, were better than 5% (2 RSD) for all elements.

2.6. Bulk organic geochemistry

%TOC was measured using a Rock Eval 6 at the University of Oxford following standard analytical protocols (Behar et al., 2001). Relative uncertainty on %TOC measurements was ~0.2%, assessed by repeated analyses of an in-house shale standard (SAB-1).

3. Results and discussion

3.1. Severity and timing of local redox conditions during the EOT

Mo and U are both redox-sensitive metals, whose burial increases in marine sediments as local conditions become anoxic and sulfidic and/or as their concentrations in seawater increase. Mo burial occurs due to water-column precipitation of Fe-Mo-S particulates as sulfate reduction leads to the accumulation of aqueous H₂S (Helz et al., 2011; Vorlicek et al., 2019), or by sorption to exported organic particulates (Dahl et al., 2017; Dickson et al., 2019; Tessin et al., 2019). U burial occurs by the organic or inorganic reduction of U(VI) to U(IV) within the sediment coupled to diffusion of U across the sediment-water interface (Andersen et al., 2017).

Our new data from the OSCH-1 core reveal an interval of major trace-metal burial that was triggered by the development of water-column euxinia in the MB. This interval clearly stands out when the trace metal stratigraphy is divided into three zones. In the lowest zone (a, Fig. 2) concentrations of the redox-sensitive metals Mo and U average ~14 μg/g and 8 μg/g respectively; in zone b they are much higher (61 μg/g and 14 μg/g), and in zone c they average 9 μg/g and 6 μg/g. Sedimentation rates, approximately estimated from nanofossil zone boundary ages transferred from ODP Site 1218 (Coxall and Wilson, 2011) (Figs. 2 and 3), vary by a factor of ~3, while La/Yb ratios indicate subtle changes in detrital sediment provenance that are probably linked to the tectonically driven evolution of the MB across the study period (Fig. S3). Importantly, however, detrital sediments have little to no effect on Mo and U trends: enrichment factors (EF) of Mo and U, calculated by normalising Mo/Al and U/Al ratios to Upper Continental Crust averages of 0.000135 and 0.000027 respectively (Rudnick and Gao, 2003), follow the major trends in the raw concentrations (Fig. 2). In zone b, ~93–99% of the accumulated Mo and ~64–92% of the accumulated U is non-detrital. The low proportions of detrital Mo and U also means that the isotopic compositions of these elements discussed in section 3.2, particularly in zone b, are dominated by the non-detrital fraction.

Determining the cause of the large changes in trace metal burial throughout zones a–c requires us to independently constrain local redox conditions and/or local seawater concentrations. To constrain local redox conditions, we use the Fe-speciation proxy that quantifies the amount of highly reactive (Fe₅⁶₃₄) and pyritised (Fe₅₇₈₉) iron in sediments as a fraction of the total Fe (Fe₅) inventory (Raiswell et al., 2018) (Fig. 2). Sediments in zones a–c have consistently high Fe₅₇₈₉/Fe₅ ratios >0.38, signifying an anoxic depositional environment. Fe₅₇₈₉/Fe₅₃₄ ratios are mostly >0.7, suggestive of euxinic conditions with H₂S present in the water column. In zone a, however, low Mo concentrations, pyrite frambooids >4.5 μm with wide size distributions, and the minor presence of glauconite grains (Schulz et al., 2002) suggest that the high Fe₅₇₈₉/Fe₅₃₄ ratios probably represent extensive diagenetic pyritization, with sulfidic conditions dominantly limited to sediment porewaters. In zone c, Fe₅₇₈₉/Fe₅₃₄ ratios frequently decrease below ~0.7, signifying ferruginous (sulfur-limited) conditions. When viewed in context with low stearane/hopane ratios and high C₂₃₇₉/S ratios (Schulz et al., 2002, 2005), our new Fe-speciation data indicate that deposition in zone c probably occurred in a low-salinity setting due to a combination of relative sea-level fall and influx of shallow seaways to the north (Rögl, 1999). We conclude that from zones a–b the increase in trace-metal concentrations in the OSCH-1 core is related to the development of basal euxinia, while the diminished trace-metal concentrations passing from zone b to c is linked to a declining local seawater inventory as the MB became isolated from the open ocean. The presence of aryl-isoprenoids from the base of zone b (Schulz et al., 2002, 2005) (Fig. 2) demonstrates that the local chemocline rose into the photic zone from this time, probably with higher aqueous H₂S concentrations that facilitated conversion of Mo₅₇₈₉ to particle-reactive thiomolybdates Mo₅₅₄₉(SX)₅⁻ (Helz et al., 2011; Vorlicek et al., 2019). Mo removal may have been accelerated by better preservation of sinking organic matter under locally stagnant conditions.

We can tie the increase in Mo and U concentrations observed in the MB during zone b to the EOT, when the mass of grounded ice on Antarctica increased over a period of ~600 krysts (Coxall and Wilson, 2011) (Fig. 3). In ODP Site 1218 in the Central Pacific Ocean, the nanofossil NP20/21 zonal boundary occurs below the initial increase in benthic foraminifera δ¹⁸O that defines the first pulse of the EOT, and the NP21/22 zonal boundary occurs above the peak of the second increase in δ¹⁸O that marks ends of the EOT (Coxall and Wilson, 2011). Zone b in the OSCH-1 core is contained entirely within zone NP21 (Schulz et al., 2002). The biostatigraphic correlation is supported by our new ¹⁸⁷Os/¹⁸⁸Os measurements from zone b in the OSCH-1 core that range from 0.26–0.48 (Table S1). The Late Eocene features a unique global minimum in seawater ¹⁸⁷Os/¹⁸⁸Os of 0.22–0.28 that can be thus employed as a chemostratigraphic marker to tie the OSCH-1 core stratigraphy to the period of time immediately prior to the EOT (Ravizza and Paquay, 2008; Ravizza and Peucat-Ehrenbrink, 2003) (Fig. 3). The position of the Eocene/Oligocene boundary in the OSCH-1 core is not exact, and is approximately placed at the lithostratigraphic boundary between zones a and b.

Local tectonics preconditioned the MB to the development of local euxinia over timescales of millions of years (Schulz et al., 2002; Sissingh, 1997; Wagner, 1998). However, the correspondence in timing between the start of zone b and the EOT suggests that the ~70 m of eustatic sea-level fall associated with polar glaciation (Lear et al., 2008; Katz et al., 2008) may have been the final trigger for the short, intense period of euxinia that developed in the MB. Sea-level fall would have significantly reduced the transfer of water across shallow sills (Schulz et al., 2005) into a basin that was probably only ~400–800 m deep in the late Eocene (Schulz et al., 2002; Sissingh, 1997). Importantly, by triggering euxinia, sea-level fall may have initiated a feedback of regionally enhanced organic matter burial that contributed to pCO₂ decline across the EOT (Cramwinckel et al., 2018; Pearson et al., 2009). Age-equivalent organic-rich shales occur throughout the inter-connected basins of the Para-Tethys (Veto, 1957), suggesting that the depositional conditions documented in the OSCH-1 core were regionally extensive.

It has been shown that several species of planktonic foraminifera and coccolithophores adapted to oligotrophic conditions went extinct during the EOT, while species adapted to more eutrophic conditions became established (Dunkley-Jones et al., 2008; Pearson et al., 2008; Wade and Pearson, 2008). It is possible that nutri-
3.2. Global redox conditions during the EOT

The development of locally euxinic conditions during the EOT in the MB in zone b allows us to use the isotopic compositions of Mo ($\delta^{58}$Mo) and U ($\delta^{238}$U) to constrain the global signature of anoxia and euxinia in the world’s oceans at the end of the long-term Eocene cooling interval. The sedimentary compositions of Mo and U in settings such as these can be related directly to the isotopic compositions of these metals in the open ocean, acting as ‘yardsticks’ of global seawater chemistry (Andersen et al., 2017; Dickson, 2017).

Molybdenum and uranium are long-lived elements in the oceans, with residence times on the order of ~400,000–500,000 years (Dunk et al., 2002; Miller et al., 2011). A large fraction of the global burial flux of Mo occurs into oxic sediments containing Mn and Fe oxyhydroxides, with isotopic compositions that are generally lower than weathering input fluxes of ~0.6‰ (Kendall et al., 2017). In sulfidic settings, Mo burial occurs as thiomolybdate species or by adsorption to organic matter (Helz et al., 2011; Dahl et al., 2017; Dickson et al., 2019) with isotopic compositions higher than weathering inputs. The global burial flux of isotopically light Mo thus leverages the global seawater composition to a high value of ∼2.3‰ today (Nakagawa et al., 2012; Goldberg et al., 2013). In situations where the oxic burial flux contracts at the expense of the burial of Mo into sulfidic sediments – such as during times when oceanic deoxygenation becomes more extensive – the seawater $\delta^{58}$Mo would become lower. Sediments deposited in highly sulfidic water columns, with restricted water mass ventilation, have $\delta^{58}$Mo compositions close to open-ocean seawater, due to its near-quantitative removal from the water column (Brüské et al., 2019; Neubert et al., 2008; Noordmann et al., 2015). For U isotopes, the seawater composition (today ∼−0.4‰ (Tissot and Dauphas, 2015)) is largely controlled by the burial of U(VI) following biotic or abiotic reduction of U(VI) at the sediment-water interface (Andersen et al., 2017). This U(IV) burial flux has an effective isotopic composition that is ∼0.6‰ higher than seawater, but the exact sediment-seawater difference depends on the diffusion and reduction rates and the overlying seawater composition (Andersen et al., 2014). Thus, at times of expanded oceanic deoxygenation and increased burial of U(IV), the isotopic composition of U in seawater becomes lower (Andersen et al., 2014, 2017; Tissot and Dauphas, 2015).

In zone b, where Fe-speciation data and the presence of aryl-isoprenoids indicate persistently euxinic conditions, $\delta^{58}$Mo values are highest (1.6–1.8‰), and thus closest to the global seawater value (Fig. 2, Fig. S2). In zone a, $\delta^{58}$Mo values of ∼1.0‰ are probably fractionated from basin seawater under weakly sulfidic conditions. In zone c, low and variable $\delta^{58}$Mo values (∼−0.4–1.0‰) were probably sensitized to local weathering inputs and sequestration pathways, in view of evidence for low salinity conditions and very low Mo concentrations.

We argue that the $\delta^{58}$Mo fluctuations in zone b are related to changes in the fractional removal of Mo from MB seawater. In this zone, euxinic conditions, as recorded by Fe-speciation, persist throughout intervals of both low (∼1.0‰) and high (∼1.7‰) $\delta^{58}$Mo values, thus limiting the effect of redox-driven changes to Mo cycling; $\delta^{58}$Mo increases in tandem with [Mo], which suggests rapid conversion and burial of Mo as MoS$_2$ at higher [H$_2$S]. Maxima in $\delta^{58}$Mo occur higher up-section in concert with increasing concentrations of aryl-isoprenoids, a trend that is suggestive of a shoaling chemocline and euxinic conditions spanning a larger fraction of the water column. Particular shuttling cannot fully explain the fluctuations of $\delta^{58}$Mo in zone b, because it would tend to lower $\delta^{58}$Mo at times of higher [Mo] by delivering oxyhydroxide-adsorbed Mo to the seafloor, which is opposite to what we observe.

An important line of evidence for the drawdown of dissolved Mo in the MB during zone b comes from the inverse correlation between $\delta^{58}$Mo and $\delta^{238}$U (Fig. 4). This relationship emerges in restricted marine basins where the burial of isotopically heavy U(IV) outpaces U resupply by seawater exchange, causing the local basin water $\delta^{238}$U to become lighter (Andersen et al., 2018; Brüské et al., 2019; Bura-Nakic et al., 2018). As the drawdown of Mo and U progresses, the isotopic compositions of the sediments evolve towards the seawater endmember, i.e. $\delta^{238}$U becomes lower and $\delta^{58}$Mo becomes higher. $\delta^{58}$Mo and $\delta^{238}$U are only inversely corre-
lated in zone b, suggesting that in zone a, isotopic compositions are controlled by redox processes and in zone c by mixing with local weathering fluxes. Thus the Mo-U covariation trend supports the idea that the highest $\delta^{98}\text{Mo}$ compositions in zone b ($\sim$1.75‰) must have been close to the global seawater composition during the EOT due to basin drawdown. The slope of the correlation between $\delta^{98}\text{Mo}$ and $\delta^{238}\text{U}$ is close to that seen for the modern Cariaco Basin (Fig. 4). Given that the highest sediment $\delta^{98}\text{Mo}$ values in the Cariaco Basin are fractionated from global seawater by $\sim$0.5‰ (Brüseke et al., 2019), seawater during the EOT could have been as high as $\sim$2.2‰ (Fig. 5). The highest $\delta^{238}\text{U}$ values in zone b sediments ($\sim$0.2‰) constrain the seawater $\delta^{238}\text{U}$ during the EOT to $\sim$0.4‰, assuming an effective fractionation from seawater of $\sim$0.6‰ (Andersen et al., 2014, 2017; Brüseke et al., 2019; Bura-Nakic et al., 2018). This value is close to the modern seawater value and is indistinguishable from estimates obtained from EOT-age Fe-Mn crusts (Wang et al., 2016) and from carbonates deposited in the Early Eocene (Clarkson et al., 2021).

The global seawater $\delta^{98}\text{Mo}$ composition for the Early Eocene ($\sim$56–54 Ma) has been estimated as 1.94±0.24‰ from sediments deposited in the Arctic Ocean (Dickson et al., 2012; Dickson and Cohen, 2012). This value is thought to capture the prevailing redox state of the Early Eocene global ocean as conditions for the quantitative removal of Mo became briefly established in the Arctic Ocean during the Paleocene–Eocene Thermal Maximum ($\sim$56 Ma) and Eocene Thermal Maximum 2 ($\sim$54 Ma) (Dickson and Cohen, 2012). This value characterizes the global redox state of the oceans slightly before the peak of Early Eocene warmth, and equates to a slightly larger fraction of the global seafloor being anoxic and sulfidic than today (Dickson, 2017). Our inferred range of global seawater $\delta^{98}\text{Mo}$ for the EOT ($\sim$1.7–2.2‰) overlaps this value (Fig. 5).

$\delta^{238}\text{U}$ values likewise vary little from the early Eocene to the EOT, as recorded by pelagic carbonates (Clarkson et al., 2021), Fe-Mn crusts (Wang et al., 2016) and shales (this study). The correspondence of all three sedimentary archives (carbonates, crusts, shales) is a powerful indication of the robustness of the observations (Fig. 5). Taken at face value, our observations indicate that the extent of anoxic and sulfidic conditions on the global seafloor was insensitive to the long-term changes in global climate and ocean circulation that occurred during the middle–late Eocene. We note however that the area of seafloor anoxia may fluctuate $\sim$3–10 fold even within the uncertainties of the $\delta^{238}\text{U}$ method (Wang et al., 2016; Clarkson et al., 2021) and a similar magnitude of change might also be expected for the area of sulfidic deposition within the uncertainties of $\delta^{98}\text{Mo}$. These uncertainties leave room for smaller long-term changes in whole-ocean redox that would be unsolvable by these methods.

3.3. Response of global ocean redox to long-term cooling

Wang et al. (2016) concluded that the invariance of low-resolution Fe-Mn crust $\delta^{238}\text{U}$ across Early Cenozoic cooling interval probably reflects an increase in oxygen consumption in the open ocean that balanced an expected increase in $\text{O}_2$ solubility at lower temperatures. While increases in biogenic silica export rates in the Southern Ocean at $\sim$36–38 Ma are consistent with this suggestion (Dickmann et al., 2004; Diester-Haas and Zahn, 1996), biogenic silica export rates decreased sharply at approximately the same time in the equatorial Pacific Ocean (Moore et al., 2008). It is therefore not straightforward to invoke a whole-ocean change in organic matter export to explain the near-constancy of the $\delta^{238}\text{U}$ or $\delta^{98}\text{Mo}$ signatures across this interval. We suggest that the expansion of anoxic depositional conditions documented in the MB during the EOT, and perhaps in the wider Para-Tethys Sea (Veto, 1987), may have partly balanced any increase in U and Mo burial into more oxygenated sediments in the open ocean.

The response of global oxygenation to short-term environmental triggers has been studied for brief hyperthermal events such as the PETM (Dickson et al., 2012; Clarkson et al., 2021) and Cretan-
ceous Oceanic Anoxic Events such as OAE 2 (Jenkyns, 2010; Poul
ton et al., 2015). The response of global ocean oxygenation to long-
term environmental gradients is considerably less well-known. The
apparent stability in seafloor redox conditions across the multi-
million-year Early Cenozoic cooling interval, constrained by both
$\delta^{38}$Mo (this study) and $\delta^{238}$U (this study, Wang et al., 2016), sug-
gests that the oceans may be insensitive to extreme long-term changes in oxygenation. This insensitivity probably stems from the
non-linear interaction of different Earth System feedbacks and pro-
cesses (tectonics, ocean circulation, biogenic productivity, temper-
ature) operating over timescales of $10^{3-4}$ years.

4. Conclusions

We present a detailed, multi-proxy study of the MB sedimentary
succession that spans the EOT. New Os isotope data refine
existing biostratigraphic constraints on the age of the deposits and
demonstrate that an interval of basin euxinia developed during the
EOT. We do not question that large-scale tectonics associated with
the Alpine thrust belt pre-conditioned the basin to these de-
positional conditions but suggest that sea-level falls associated with
polar glaciation may have tipped the balance in favour of hydro-
graphic restriction and the accumulation of HS$_2$ in the MB water
column. These conditions are suitable for the application of redox-
sensitive isotope systems to constrain global-scale redox conditions
in the oceans. Mo and U isotope compositions of the sedimentary
deposits in the OSHC-1 core negatively co-vary in the interval of
euxinic deposition, supporting a model of metal drawdown from the
basin seawater. The sediments can thus be used to place con-
straints on the seawater isotope compositions of these elements at
$\sim$34 Ma, which are indistinguishable from similar estimates for the
Early Eocene ($\sim$54–56 Ma), prior to the commencement of long-
term global cooling. Our data imply that the long-term Eocene
cooling trend did not result in large-scale changes in the distri-
bution of anoxic and sulfidic conditions on the global seafloor. We
note, however, that the uncertainties associated with estimating seawater $\delta^{38}$Mo signatures in particular are still significant enough to
courage future refinement to these observations.

CRediT authorship contribution statement

Alexander J. Dickson: Conceptualization, Funding acquisition,
Investigation, Methodology, Project administration, Supervision, Vi-
sualization, Writing – original draft. Marie-Laure Bagard: Investi-
gation, Methodology, Project administration, Supervision, Valida-
tion, Writing – review & editing. Joachim A.R. Katchinoff: Investi-
gation, Writing – review & editing. Marc Davies: Investigation,
Project administration, Supervision, Writing – review & editing. Anthony S. Cohen: Conceptualization, Funding acquisition, Project administration, Supervision, Writing – review & editing.

Declaration of competing interest

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

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gaea database (www.pangaea.de).

Appendix A. Supplementary material

Supplementary material related to this article can be found on

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