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Iron limitation in the Western Interior Seaway during the Late Cretaceous OAE 3 and its role in phosphorus recycling and enhancing organic matter preservation

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Abstract

The sedimentary record of the Coniacian-Santonian Oceanic Anoxic Event 3 (OAE 3) in the North American Western Interior Seaway is characterized by a prolonged period of enhanced organic carbon (OC) burial. This study investigates the role of Fe in enhancing organic matter preservation and maintaining elevated primary productivity to sustain black shale deposition within the Coniacian-Santonian-aged Niobrara Formation in the USGS #1 Portland core. Iron speciation results indicate the development of a reactive Fe limitation coeval with reduced bioturbation and increased organic matter preservation, suggesting that decreased sulfide buffering by reactive Fe may have promoted enhanced organic matter preservation at the onset of OAE 3. An Fe limitation would also provide a feedback mechanism to sustain elevated primary productivity through enhanced phosphorus recycling. Additionally our results demonstrate inconsistencies between Fe-based and trace metal redox reconstructions. Iron indices from the Portland core indicate a single stepwise change, whereas the trace metal redox proxies indicate fluctuating redox conditions during and after OAE 3. Using Fe speciation to reconstruct past
redox conditions may be complicated by a number of factors, including Fe sequestration in
diagenetic carbonate phases and efficient sedimentary pyrite formation in a system with limited
Fe supply and high levels of export production.

Introduction

Iron availability in marine systems can significantly influence organic carbon (OC)
accumulation in sediments by affecting rates of both primary productivity and organic matter
preservation. In the modern ocean, increased Fe delivery or “fertilization” is known to enhance
primary productivity in high nutrient, low chlorophyll regions (e.g. Boyd et al., 2000). In
sediments, the availability and early diagenesis of Fe are known to play complex roles in OC
accumulation by inducing changes in C, P, and S cycling (Canfield, 1989; Raiswell and Canfield,
1996). For example, Meyers (2007) outlined the role of Fe in the “sulfide buffer and phosphorus
trap hypothesis,” where reactive Fe minerals (especially Fe (oxyhydr)oxides) act to buffer sulfide
buildup within porewaters through formation of Fe-sulfide mineral phases (FeS, FeS$_2$). Once all
available reactive Fe is sulfidized, free sulfide accumulates in porewaters, which promotes P
recycling in the absence of Fe (oxyhydr)oxides and reduces bioturbation and organic matter
oxygen exposure time (Canfield, 1989; Raiswell and Canfield, 1996; Meyers, 2007; Tribovillard
et al., 2015).

As some of the most dramatic examples of OC burial in the geologic record, Mesozoic
Oceanic Anoxic Events (OAEs) provide unique opportunities for studying OC burial processes
as each event varied in duration and geographic extent. The Cenomanian-Turonian OAE 2, for
example, is characterized by widespread black shale deposition (Schlanger et al., 1987; Tsikos et
al., 2002), and a distinctive and relatively short-lived (~600 ka: Sageman et al., 2006) positive
carbon isotope excursion (Arthur et al., 1987; Jenkyns, 2010). In contrast, OC-rich sedimentation
during the Coniacian-Santonian OAE 3 was restricted to the equatorial Atlantic and adjacent
continental shelves and seaways (e.g., März et al., 2008; Locklair et al., 2011; Wagreich, 2012),
including the North American Western Interior Seaway (WIS). Due to the long duration (>3
Myrs) and higher rates of OC burial in some locations as compared to OAE 2, OAE 3 represents
an important perturbation to the global carbon cycle. Irrespective of its exact geographic extent,
OC burial during OAE 3 was elevated, widespread, and prolonged, and must have required
internal oceanic feedback mechanisms.

This study investigates Fe speciation before, during and after OAE 3 in the WIS to
understand its role in prolonged OC burial. A comparison of these results with S concentrations
and previously published records of bioturbation (Savrda, 1998) and organic matter quality
(Tessin et al., 2015) is used to assess how fluctuations in Fe availability can affect “sulfide
buffering” and OC preservation, particularly at the onset of enhanced OC burial. Evolution of Fe
speciation is also compared to TOC/P and P/Al ratios to evaluate how changes in Fe chemistry
affected “P trapping” before, during, and after OAE 3. Finally, comparison of Fe speciation with
redox sensitive trace metals provides an opportunity to test whether Fe-based redox
reconstructions are supported by other redox proxies in highly productive sedimentary systems
where pyrite formation may be Fe limited.

Background

Western Interior Seaway

During maximum transgression, the WIS extended from the Gulf of Mexico to the Arctic
Ocean and from central Utah to central Iowa (Fig. 1). The sedimentary record of the WIS
includes episodic deposition of OC-rich shales and chalks, including those recorded during the two maximum transgressions, the Greenhorn and the Niobrara Transgressions. The Niobrara Formation was deposited during the latter transgression from the Late Turonian to the Early Campanian (Scott and Cobban, 1964; Locklair et al., 2011) and is formally divided into two members: the basal Fort Hays Limestone and the overlying Smoky Hill Chalk (Fig. 1). Previous research has identified OAE 3 in the Smoky Hill Chalk based on a positive $\delta^{13}C$ excursion and elevated OC concentrations (e.g. Locklair et al., 2011; Tessin et al., 2015).

Paleontological and sedimentary evidence indicates that in western Kansas and central Colorado, the Fort Hays was initially deposited in water depths of 15–50 m and that the water column progressively deepened to water depths of 150–300 m during Smoky Hill deposition (Hattin, 1982). The focus of this study, the USGS Portland core, was located in the deepest portion of the WIS during the Late Cretaceous (Fig 1; Sageman and Arthur, 1994). During maximum transgression, the influx of Tethyan water from the south may have delivered nutrients, including reactive Fe into the WIS (e.g. Meyers et al., 2005). Other sources of Fe and nutrients to the seaway include continental runoff from the Sevier Highlands (e.g. Flögel et al., 2009) and volcanic ash falls, which are recorded as numerous bentonite layers.

Trace metal and iron redox proxies

Trace metal concentrations in ancient sediments and rocks allow for the reconstruction of marine redox conditions (Tribovillard et al., 2006). Cadmium and zinc accumulate in sediments under sulfidic conditions. Cadmium exhibits a nutrient-like behavior and is delivered to sediments with OM (Piper and Perkins, 2004). Zinc is delivered to the sediments via organic acid complexes or through adsorption on Fe-Mn oxyhydroxides (Fernex et al., 1992; Algeo and
Maynard, 2004). During early diagenesis, Cd and Zn are released to porewaters from organic matter and are authigenically enriched in sediments as CdS and ZnS in the presence of free sulfide (Huerta-Diaz and Morse, 1992). Sedimentary Mo accumulation is thought to require a threshold concentration of H$_2$S for molybdate to be transformed to the particle reactive thiomolybdate (i.e. the thiomolybdate switch) and subsequently scavenged by OM and Fe–S mineral phases (Erickson and Helz, 2000; Chappaz et al., 2014). Thus, more sulfidic conditions are thought to be required for Mo accumulation than for Cd and Zn accumulation. Rhenium also accumulates in reducing sediments but unlike Mo, Zn, and Cd, sedimentary enrichment is not thought to require the presence of free H$_2$S (Crusius et al., 1996).

Iron speciation has also been used as a redox proxy to distinguish oxic, ferruginous or euxinic conditions in ancient black shales. This approach is based on the observation that Fe, particularly highly reactive Fe (Fe$_{HR}$), is enriched in sediments deposited beneath an anoxic water column (e.g. Raiswell and Anderson, 2005; Lyons and Severmann, 2006). Therefore, Fe$_T$/Al ratios of >0.5 and Fe$_{HR}$/Fe$_T$ of >0.38 are indicative of an anoxic water column (e.g. Poulton and Canfield, 2005; Lyons and Severmann, 2006). Studies of modern euxinic basins (e.g. Black Sea, Cariaco Basin) have identified the source of Fe enrichment in anoxic basins as reactive Fe shuttled from suboxic shallow shelf sediments and subsequently precipitated within the water column in the deeper euxinic basin (Lyons and Severmann, 2006; Severmann et al., 2010). The ratio of Fe$_{py}$/Fe$_{HR}$ can be further used to distinguish between euxinic and non-euxinic anoxic systems because significant sulfidization of the Fe$_{HR}$ pool (>0.7–0.8) has been used to indicate the presence of dissolved H$_2$S in the water column (Raiswell and Canfield, 1998; März et al., 2008; Poulton and Canfield, 2011; Poulton et al., 2015).
Certain factors, including basin geometry, high sedimentation rates, and low Fe export efficiency from the shelves, are known to complicate the interpretation of Fe-speciation results by decreasing \( \text{Fe}_{\text{HR}} / \text{Fe}_T \) and \( \text{Fe}_{\text{py}} / \text{Fe}_{\text{HR}} \) values (Raiswell and Canfield 1998; Anderson and Raiswell 2004; Raiswell and Anderson 2005; Lyons and Severmann, 2006). These complications are generally thought to obscure the signal of anoxic conditions rather than erroneously indicate anoxic/euxinic conditions. Basin geometry, in particular the shelf to basin ratio, is of interest in the current study because “shelf” settings are defined as <200 m water depth, which means that most, if not all, of the WIS is considered to be a shelf and not deep basin setting.

Methods and materials

The USGS #1 Portland core was drilled and continuously cored near Cañon City, CO (Dean and Arthur, 1998). The 75-m thick Late Cretaceous Niobrara Formation section of the Portland core was sampled at 0.5 m resolution at the USGS Core Research Center in Denver, CO (Fig. 1). Chemostratigraphy for the core is based on carbon isotope and total organic carbon records presented in Tessin et al. (2015), which was used to identify pre-OAE, OAE, and post-OAE intervals. In Fig. 1, the three defined intervals from Tessin et al. (2015) are further divided. The pre-OAE Interval (Interval 1) has been partitioned into Interval 1a, which includes the Fort Hays Limestone and Interval 1b, which includes the lower shale limestone member of the Smoky Hill Chalk (Scott and Cobban, 1964). The OAE Interval (Interval 2) is divided into 2a, 2b, and 2c based on previously published total organic carbon (TOC) values (Tessin et al., 2015; Fig. 1). Intervals 2a and 2c are defined by TOC values >3%. The post-OAE Interval (Interval 3) was not subdivided.

Samples analyzed for bulk elemental concentrations (Al, Fe, S, P, Mo, Re, Cd, and Zn)
were ground to <75 μm and homogenized in an alumina shatterbox to minimize trace element contamination. Analyses were completed at ALS Laboratories in Vancouver, BC. Whole rock samples were digested with perchloric, hydrofluoric, nitric, and hydrochloric acids. Concentrations were determined by inductively coupled plasma-atomic emission spectrometry (ICP-AES) and inductively coupled plasma-mass spectrometry (ICP-MS). GBM908-10, GBM908-5, OREAS 90 and MRGeo08 standards were used to verify elemental concentrations. Accuracy and precision are reported in Supplemental Table 1.

Sequential iron extractions were completed following Poulton and Canfield (2005), and pyrite Fe extractions following Canfield et al. (1986). The pyrite fraction is stoichiometrically determined following precipitation of chromium reducible sulfide as ZnS. Iron phases measured are outlined in Table 1. Carbonate-associated Fe (Fe\textsubscript{carb}), magnetite Fe (Fe\textsubscript{mag}), and Fe\textsubscript{oxyhydr}oxides (Fe\textsubscript{ox}) are considered highly reactive (HR) because these phases react with sulfide on timescales of months to years (Poulton and Canfield, 2005). The Fe\textsubscript{HR} pool also includes pyrite Fe (Fe\textsubscript{py}), which represents Fe that has reacted with sulfide in the water column or during early diagenesis in the sediments (Raiswell and Canfield, 1998; Poulton and Raiswell, 2002). The Fe\textsubscript{HR} pool is thus defined as Fe\textsubscript{carb} + Fe\textsubscript{ox} + Fe\textsubscript{mag} + Fe\textsubscript{py} and poorly and non-reactive Fe (Fe\textsubscript{NR/PR}) is calculated as Fe\textsubscript{T} – Fe\textsubscript{HR}. Concentrations of Fe\textsubscript{carb}, Fe\textsubscript{ox}, and Fe\textsubscript{mag} were measured on ICP-MS (Thermo iCAP Q) within the STARLAB at Central Michigan University. Analytical precision and accuracy, determined from replicate analyses (n = 21) of a certified standard (SCP Science) were better than 5%. Calculated pyrite Fe concentrations generally replicated with <5% precision. All geochemical data discussed in this paper are archived in Pangaea (www.pangaea.de).
Results

Iron

The different Fe phases identified by sequential extractions and pyrite Fe measurement are plotted in Fig. 2. Carbonate-associated Fe ($\text{Fe}_{\text{carb}}$) is generally elevated during Interval 1, with concentrations up to 0.2 wt. %, whereas during Intervals 2 and 3, $\text{Fe}_{\text{carb}}$ concentrations are consistently < 0.05 wt. %. Iron (oxyhydr)oxides ($\text{Fe}_{\text{ox}}$) consistently accounts for between 0.01–0.1 wt. % throughout the Portland record. Magnetite-associated Fe ($\text{Fe}_{\text{mag}}$) is generally low but varies throughout the three intervals. During Interval 1, $\text{Fe}_{\text{mag}}$ concentrations range from 0.01–0.07 wt. %. Concentrations of $\text{Fe}_{\text{mag}}$ are consistently < 0.01 wt. % during Interval 2 and range between 0.02–0.04 wt.% during Interval 3. Pyrite-associated Fe ($\text{Fe}_{\text{py}}$) increases notably leading into Interval 2 to concentrations of between 0.2–1.1 wt.% During Intervals 2 and 3, an average of 88% of $\text{Fe}_{\text{HR}}$ is in the form of $\text{Fe}_{\text{py}}$.

The $\text{Fe}_T$/Al values are the most variable and elevated during Interval 1, with values ranging from 0.07 to 1.60 (Fig. 2). During Intervals 2 and 3, $\text{Fe}_T$/Al values average 0.51 and 0.46, respectively. Values of $\text{Fe}_T$/Al are elevated during Intervals 2a and slightly elevated during Interval 2c. Values of $\text{Fe}_{\text{HR}}$/Fe$_T$ range between 0.36 and 0.88 with an average value of 0.60 throughout the record. The lowest values (0.12 and 0.33) of $\text{Fe}_{\text{py}}$/Fe$_{\text{HR}}$ occur during Interval 1a. Values of $\text{Fe}_{\text{py}}$/Fe$_{\text{HR}}$ increase abruptly in Interval 1b with values ranging between 0.71 and 0.93 for the remainder of the record.

Trace metals

Trace metal/Al ratios are plotted as ppm (Mo, Cd, Zn) or ppb (Re) to weight % ratios and compared to previously published Mo and TOC concentrations from Tessin et al. (2015) in Fig.
3. The records of Mo concentrations and Mo/Al exhibit similar patterns throughout the Portland core, and generally follow the TOC record (Fig. 3). Interval 1 is characterized by low Mo concentrations (an average of 0.3 ppm) and Mo/Al ratios (an average of 0.2). Interval 2 exhibits the most variable Mo and Mo/Al values, with two peak periods of elevated Mo concentrations during Intervals 2a and 2c separated by low concentrations during Interval 2b. During Interval 3, Mo concentrations and Mo/Al values become more stable, ranging from 8.5–36 ppm and 1.9–10.8, respectively. The record of Mo/TOC exhibits similar trends to the Mo and Mo/Al record, with the lowest values recorded during Interval 1 (1.0). Within Interval 2, Mo/TOC concentrations are elevated during Intervals 2a and 2c, with low values recorded during Interval 2b. However, unlike the Mo and Mo/Al records, Mo/TOC values within Interval 3 are elevated and variable, ranging from 2.8–40.9.

Rhenium, Cd, and Zn behave similarly to Mo in the Portland core (Fig 3). Throughout Interval 1, Re/Al, Cd/Al, and Zn/Al values are generally low (an average of 3.8, 0.08, and 27, respectively). The highest Re/Al, Cd/Al, and Zn/Al values occur during Intervals 2a and 2c, whereas Interval 2b exhibits values similar to Interval 1. During Interval 3, values remain elevated and range between 7–85, 0.4–3.1, and 7–91, respectively.

Sulfur and Phosphorus

Sulfur results are compared to previously published TOC, bioturbation, and Rockeval results from Tessin et al. (2015) and Savrda (1998) in Fig. 4. Bioturbation derived Interpreted Oxygenation Curves (IOC) from Savrda (1998) are based on the vertical distribution of laminites and oxygen-related ichnocoenoses. The IOC record has been smoothed by integrating data to 0.5 m resolution to match the resolution of the geochemical data. IOC values generally decrease
throughout Interval 1, with the lowest values recorded within Interval 2, indicating reduced bioturbation during the OAE (Savrda, 1998). Hydrogen and oxygen index values, produced through RockEval pyrolysis, are proxies for the hydrogen- and oxygen-richness of organic matter. Hydrogen index values are markedly elevated in Interval 2, as compared to Interval 1, highlighting an increase in the hydrogen-richness of organic matter. Conversely, oxygen index values are elevated in Interval 1 as compared to Interval 2, indicating a decrease in the oxygen-richness of organic matter (Tessin et al., 2015).

Weight % ratios of TS/Al indicate increasing S accumulation within the sediments throughout the record (Fig 4). During Interval 1, TS/Al averages 0.13 compared to an average of 0.51 and 0.38 during Intervals 2 and 3. Excess S/TS records (calculated as (TS - CRS)/TS) illustrate that between 13 and 44% of TS is non-CRS (Fig 5). Due to low concentrations of S and CRS in samples between 58 and 78 m in the core, errors on $S_{\text{excess}}/TS$ increase but values are near 0%.

Phosphorus values are plotted as P/Al and TOC/P ratios in Figure 5. Trends in P geochemistry are based on total P and rather than organic or reactive P because it has been observed that P speciation records can be significantly altered by sample handling and diagenesis and that $P_T/Al$ and TOC/P are the most reliable records of changes in P geochemistry in black shales (Algeo and Ingalls, 2007; Kraal et al., 2010). Calculations of P/Al ratios are shown as weight % ratios, while TOC/P values are calculated as molar ratios. During Interval 1, P/Al are highly variable, ranging between 0.009 and 0.170, with an average value of 0.04. During Intervals 2 and 3, P/Al values become less variable and average 0.025 and 0.018, respectively. TOC/P values are low throughout Interval 1, with an average molar ratio of 15. Molar ratios of
TOC/P range between 26 and 306 during Interval 2. During Interval 3, TOC/P molar ratio values stabilize with an average value of 99.

Discussion

Fe speciation and limitation

Comparison of Fe speciation and trace metal concentrations throughout the Portland record suggests that changes in Fe speciation and abundance are decoupled from redox conditions. During Interval 1a, the Fe/T/Al record exhibits significant variability (Fig. 2). Elevated Fe/T/Al (>1) and Fe/hr/Fe/T ratios (>0.38) both indicate sedimentary Fe enrichments. Samples enriched in Fe/hr are characterized by low Al/T (0.29–0.37 wt. %) and Fe/T (0.43–0.5 wt. %) concentrations, as well as high carbonate concentrations (94.8–96.7 wt. %; SI Table 2). While elevated Fe/T/Al and Fe/hr/Fe/T ratios could be interpreted as an indicator of periodic deposition under an anoxic water column, this interpretation is inconsistent with trace metal results. No enrichments in trace metals (Mo, Re, Cd, or Zn) are observed during Interval 1 (Fig. 3). This interval is also characterized by significant bioturbation and the presence of benthic macrofossils, both of which support well-oxygenated conditions (Savrda, 1998).

It has been shown in modern, carbonate-rich sediments that anomalously high Fe/T/Al values can occur when Fe/T is <0.5 wt. % due to incorporation of Fe^{2+} from anoxic porewaters into early diagenetic carbonate cements (Clarkson et al., 2014). Samples in the Portland core with <0.5 wt. % Fe, generally exhibit elevated Fe/T/Al ratios (>1) and have large contributions of carbonate-associated Fe (>40% of total Fe), whereas carbonate-associated Fe comprises less than 8% of total Fe in samples with >1 wt. % Fe (Fig. 2). The significant increase in Fe_{carb} suggests early diagenetic transformations of Fe within the sediments and pyrite formation limited by...
sulfide supply (Raiswell et al., 2011). The latter is also supported by the S excess/TS data, showing that all available sulfide has been sequestered into pyrite. Our results agree with the modern compilation of Clarkson et al. (2014) that suggests that Fe speciation from carbonate-rich samples <0.5 wt. % Fe may not accurately record redox conditions.

During Interval 1b, \( \text{Fe}_{\text{py}}/\text{Fe}_{\text{HR}} \) values increase abruptly and remain elevated throughout the remainder of the record, suggesting efficient sulfidization of reactive Fe preceding, during and after the OAE (Fig. 2). This transition marks a shift from sulfide limited to reactive Fe limited pyrite formation in WIS sediments. Ratios of \( \text{Fe}_{\text{HR}}/\text{Fe}_T \) consistently falling above the anoxic threshold value of 0.38 and elevated \( \text{Fe}_{\text{py}}/\text{Fe}_{\text{HR}} \) values suggest water column precipitation of \( \text{Fe}^{2+} \) via pyrite formation under euxinic conditions preceding, during, and after the OAE (Fig. 2). However, persistent euxinia is inconsistent with trace metal distributions and biotic evidence, which indicate fluctuating redox conditions (Savrda, 1998). Redox sensitive trace metals (Mo, Re, Cd, and Zn) all exhibit consistent patterns throughout the Portland record, wherein trace metal accumulations track TOC concentrations. In particular, during Intervals 1b and 2b, low trace metal concentrations indicate relatively well-oxygenated conditions, despite consistently elevated \( \text{Fe}_{\text{py}}/\text{Fe}_{\text{HR}} \). Additionally, Intervals 1b and 2b are associated with increased bioturbation, which is inconsistent with persistent euxinia (Savrda, 1998). Conversely, during Intervals 2a and 2c, elevated Mo, Cd, Re, and Zn concentrations and dominantly laminated (not bioturbated) sediments (Savrda, 1998) support the presence of euxinic conditions indicated by elevated \( \text{Fe}_{\text{py}}/\text{Fe}_{\text{HR}} \) and \( \text{Fe}_{\text{HR}}/\text{Fe}_T \) ratios.

The differences between \( \text{Fe}_{\text{py}}/\text{Fe}_{\text{HR}} \) and Mo, two proxies for euxinia, are highlighted in Fig. 6. Based on modern observations, Scott and Lyons (2012) proposed that Mo enrichment levels can be correlated with specific redox conditions: oxic (up to 2 ppm), anoxic (chemocline
located within sediment, 2–25 ppm), intermittently euxinic conditions (chemocline located
within water column, 25–100 ppm), and permanently euxinic (>100 ppm). However, Mo
concentrations >25 ppm are only recorded within Intervals 2a and 2c, despite Fe$_{py}$/Fe$_{HR}$ values
>0.7 throughout Intervals 1b, 2, and 3.

Changes in the trace metal inventory of the ocean, particularly within a partially restricted
basin, could affect interpretation of trace metal redox proxies. Low Mo/TOC values in the
Niobrara Formation relative to modern marine sediments indicate significant basin restriction
and/or a reduced global Mo seawater budget (Tessin et al., 2015). However, variability in the
Mo/TOC record, specifically higher Mo/TOC values coeval with elevated Mo and TOC
concentrations, suggests that Mo is not being forced by changes in the Mo budget of the ocean
(Fig. 3). Additionally, the consistency of trends across all trace metals, paired with biological
evidence, demonstrates that the trace metal distributions are likely primarily controlled by redox
conditions. Therefore, the lack of variability in Fe$_{py}$/Fe$_{HR}$ values suggests that this proxy may
have been insensitive to redox fluctuations displayed by trace metal enrichments during and after
OAE 3 in the WIS (Fig. 6).

The disagreement between the trace metal distributions and the Fe speciation could be
explained by either significant porewater sulfidization of reactive Fe or downward moving
sulfidization fronts. Elevated TOC concentrations during Intervals 2 and 3 indicate a significant
increase in organic matter flux to the sediments, which would result in elevated sedimentary
sulfate reduction, increased porewater H$_2$S concentrations, and enhanced porewater sulfidization
of reactive Fe. While it is generally assumed that water column sulfide is required to scavenge
Fe$^{2+}$ and produce elevated Fe$_{py}$/Fe$_{HR}$ and Fe$_{HR}$/Fe$_{T}$ ratios, a recent study from the Peel-Harvey
Estuary in Australia illustrated that significant Fe sulfidization in sediments can occur despite
deposition under a <2m deep, oxic water column (Kraal et al., 2013). The Peel-Harvey estuary sediments similarly exhibit large Fe\textsubscript{HR} enrichments despite the absence of an anoxic water column. Efficient Fe sulfidization has also been recorded within the surface sediments of the highly productive Achterwasser lagoon in the SW Baltic Sea, despite oxic conditions at the sediment-water interface (Neumann et al., 2005). These observations of significant early diagenetic sulfidization of reactive Fe within the surface sediments of modern sites support the possibility of pore water, rather than water column, sulfidization of reactive Fe within the WIS. Downward moving sulfidization fronts associated with peak export productivity during Intervals 2a and 2c could also have pyritized reactive Fe deposited in the underlying Intervals 1b and 2b. Trace metal and TOC results indicate that sediments in Intervals 1b and 2b were deposited under conditions characterized by enhanced oxygenation and/or reduced export productivity. Enhanced levels of sulfate reduction under elevated export productivity during Intervals 2a and 2c could have led to porewater H\textsubscript{2}S accumulation (for example, around a sulfate-methane transition zone), which could "burn down" into underlying organic- and Fe-poor sediments, sulfurizing any available Fe\textsubscript{HR}. Diagenetic sulfidization of reactive Fe in sediments underlying high TOC sediments has been recorded in a number of modern settings including the Black Sea, Kau Bay, and the Arabian Sea OMZ (Neretin et al., 2004; Middelburg, 1991; Schenau et al., 2002) and also occurred below Mediterranean sapropels (Passier et al., 1996). These sulfidization processes occur far below the sediment-water interface, and are not associated with any OC or trace metal enrichments, which could explain the discrepancy between trace metal, TOC, and Fe results within Intervals 1b and 2b. Reactive Fe consumption within sediments, whether through early diagenetic sulfidization near the surface sediment or through post-depositional sulfide burn down, could...
occur more readily under low Fe availability. Despite elevated Fe$_{HR}$/Fe$_T$ ratios, the Fe$_T$/Al record rarely shows Fe enrichments after Intervals 1a (Fig. 2). A marked increase in Fe$_T$/Al during Interval 2a (up to 0.88) and a more modest increase during Interval 2c (up to 0.58) paired with elevated Fe$_{HR}$/Fe$_T$ and Fe$_{py}$/Fe$_{HR}$ values (>0.38; >0.7) are consistent with pyrite formation in a euxinic water column. In general, however, the majority of Fe$_T$/Al ratios from Intervals 1b to 3 are below the average shale value of 0.55, strongly supporting that Fe could be the limiting pyrite formation in the WIS.

Although Fe$_T$/Al values are relatively low, a significant portion of this Fe is Fe$_{HR}$.

Potential sources of reactive Fe to the deep basin of the WIS include Fe shuttled from the shallower to the deeper parts of the basin, the influx of nutrient-rich Tethyan water to the WIS during the Niobrara transgression, and Fe associated with volcanic ash. Regardless of the relative influence of these Fe sources, Fe$_T$/Al ratios generally suggest that, especially during Intervals 1b, 2b, and 3, Fe delivery to the sediments is low, which could promote porewater sulfide accumulation and reactive Fe consumption. While Fe speciation has not been measured elsewhere in the WIS during OAE 3, in depth analysis of Fe-S-C relationships in the Berthoud State #4 core indicated that, similar to the Portland core, pyrite formation was reactive Fe limited during deposition of the Niobrara Formation (Dean and Arthur, 1989; Fig. 1). Furthermore, average Fe$_T$/Al ratios within the Berthoud and the Aristocrat Angus cores are 0.39 and 0.38, respectively, within the Niobrara subunits that are equivalent to our OAE and post-OAE intervals (Dean and Arthur, 1998; Locklair et al., 2011; SI Table 2; Fig. 1). Ratios of Fe$_T$/Al below crustal values (0.55) at all three sites support the conclusion that low Fe delivery was a persistent phenomenon throughout the deep basin in central Colorado during and after OAE 3. Efficient Fe retention in shallower, nearshore sediments due to expansion of sedimentary sulfidic conditions...
could limit Fe shuttling, reducing the supply of reactive Fe to the deepest regions of the WIS.

Recent work by Scholz et al. (2014) demonstrates that a relatively narrow window of redox conditions that promote Fe release exists, such that slightly more reducing conditions on the "shelf" could limit Fe supply to the deepest portions of the basin.

Our results suggest that Fe limitation, paired with either elevated early diagenetic sulfdization of reactive Fe in surface sediments or downward moving sulfdization fronts, can lead to significant porewater pyrite formation, suggesting that particular attention should be taken when applying the Fe$_{py}$/Fe$_{HR}$ proxy to reconstruct water column redox at sites characterized by low Fe availability and elevated export productivity.

Implications of low Fe availability

The ability of Fe to buffer porewater sulfide has been proposed as a mechanism to decrease organic matter preservation in sediments (e.g. Meyers et al., 2005; Meyers, 2007; Tribovillard et al., 2015). When reactive Fe is readily available, it can buffer porewater H$_2$S accumulation via Fe-sulfide formation. Conversely, when reactive Fe is limited, H$_2$S can accumulate in porewaters more rapidly (Meyers et al., 2005). Resulting high porewater H$_2$S concentrations create inhospitable conditions for infauna and, subsequently, can reduce the degree of bioturbation (Meyers et al., 2005; Meyers 2007). A reduction in bioturbation and bioirrigation, in turn, influences the composition of the porewater and solid constituents in sediments because infaunal organisms ventilate the substrate with oxygen-rich waters, stimulating microbial activity and reducing sedimentary organic matter preservation (Aller, 1978; Kristensen, 2000; Zonneveld et al., 2010).
During the Coniacian-Santonian, a strong relationship is found between S concentrations, bioturbation-derived IOC, and TOC in the Portland core (Fig. 4). Interval 1a is highly bioturbated and characterized by low TOC, indicating that burrowing organisms and the associated oxidative processes resulted in near complete OC loss from the sediments. When oxygen limitation reduces macrofaunal activity, beginning at the end of Interval 1, the reduction in bioturbation reduced sedimentary ventilation and oxygen exposure time. The increase in Fe$_\text{py}$/Fe$_\text{HR}$, S$_\text{excess}$/TS, and TS/Al ratios near the end of Interval 1 (Figure 2) suggests that a drawdown of reactive Fe by H$_2$S consumption is coeval with the observed onset of reduced benthic activity. Coeval changes in Hydrogen Index and Oxygen Index (HI and OI) values indicate enhanced preservation of hydrogen-rich, oxygen-poor organic matter in laminated or microbioturbated sediments. These combined results support that a reactive Fe limitation led to increased porewater sulfide concentrations, reduced bioturbation and bioirrigation, and increased organic matter preservation. Changes in organic matter composition (HI and OI) occur alongside significant increases in measured TOC concentrations, highlighting the importance of changes in organic matter preservation during periods of enhanced OC burial.

Another consequence of a reactive Fe limitation is the possible sulfurization of organic matter (Zaback et al., 1993; Meyers, 2007; Tribovillard et al., 2015). Organic matter sulfurization occurs when excess sulfide (not reacted with Fe) modifies organic matter functional groups, such as lipids and carbohydrates. During and after OAE 3, 20–45% of S is in excess of pyrite S and S$_\text{excess}$ generally tracks TOC, indicating that excess S may be associated with organic matter accumulation (Fig. 4). Sulfurization would make organic matter more resistant to degradation, further promoting organic matter preservation throughout Intervals 2 and 3.
By reducing sedimentary P sequestration (the so-called “Phosphorus Trap”), an Fe limitation could also directly impact nutrient recycling within sediment porewaters, helping to sustain elevated levels of primary production throughout the prolonged interval of black shale deposition. Iron speciation is known to control benthic P recycling and can therefore impact the return of nutrients buried with organic matter to the photic zone. Generally, as organic matter degrades in surface sediments, phosphate (PO$_4^{3-}$) is released to porewaters where it can be adsorbed on Fe oxyhydroxides, providing an efficient sedimentary sink for P. However, under O$_2$ depleted conditions, Fe oxyhydroxides are dissolved and thus, P adsorption is limited (Rozan et al., 2002). In the presence of free H$_2$S, Fe will precipitate as Fe-sulfides (e.g. FeS and FeS$_2$), which have a low affinity for PO$_4^{3-}$ (Krom and Berner, 1980). Enhanced P recycling in the presence of sulfide has been suggested as a means to sustain primary productivity (Van Cappellen and Ingall, 1994), indicating that a change in how nutrients are recycled from the sediments could be as important as changes in external nutrient input for nutrient dynamics and primary productivity in the WIS.

Elevated P/Al and low TOC/P values during Interval 1 indicate P sequestration in sediments. Conversely, during Intervals 2 and 3, P/Al values decrease and TOC/P values increase suggesting that P is no longer efficiently sequestered in the sediments (Fig. 5). Changes in P burial are coeval with increased pyrite abundance and low siderite, magnetite, and Fe (oxyhydr)oxides abundances (Figure 2), indicating that increased sedimentary P release is caused by Fe sulfidization. Our results support enhanced P recycling during the OAE and post-OAE intervals because (1) TOC/P records are consistently elevated during Intervals 2 and 3, as compared to Interval 1, and (2) P/Al ratios are very low compared to average shale values,
despite significant increases in organic matter deposition during Intervals 2 and 3, which would deliver P to the sediments.

Implications for the development of OAEs

Elevated Fe delivery was proposed to be a primary control on low OC accumulation within the WIS during OAE 2 by reducing organic matter preservation and enhancing sedimentary phosphate burial (Meyers, 2007). Our results indicate that, during OAE 3, Fe cycling had the opposite effect and promoted prolonged black shale deposition both during and after the OAE. Efficient organic matter preservation and P recycling during the Coniacian-Santonian could provide a feedback mechanism that explains the prolonged duration of black shale deposition during OAE 3 within the WIS as compared to OAE 2.

The role of P release from sediments during Cretaceous OAEs has been the focus of recent work in other ocean basins. Geochemical evidence of enhanced sedimentary P recycling during OAE 2 has been found from the Tethys, WIS, and proto-Atlantic Ocean (Mort et al., 2007; Tsandev and Slomp, 2009; Kraal et al., 2010). Furthermore, the termination of OAE 2 is associated with an increase in sedimentary P sequestration, indicating that P burial was a possible control on the duration of the event (Kraal et al., 2010). Conversely, sustained sedimentary P recycling in the WIS could have prolonged OC burial after the OAE 3 interval if remineralized P was returned to the photic zone.

Phosphorus recycling in the WIS appears to be distinct from the Proto-Atlantic during OAE 3. Previous work on OAE 3 from the Demerara Rise indicated that fluctuating ferruginous and euxinic conditions caused periods of massive P deposition that probably decreased levels of primary productivity (März et al., 2008). While the samples in our study were collected at a
lower resolution than the Demerara Rise study, our results do not provide similar evidence of P burial during OAE 3, even during better oxygenated periods such as Interval 2b. Specifically, the new results indicate that reactive Fe, not sulfate, was consistently limiting within the WIS throughout Intervals 1b, 2, and 3, thereby, inhibiting P burial throughout OAE 3 and afterwards. The ferruginous/euxinic fluctuations observed at the Demerara Rise, but not the WIS, could be attributed to either periodic influence of oxic South Atlantic water masses at the Demerara Rise or sulfate drawdown in the restricted Proto-Atlantic (März et al., 2008). Similar ferruginous/euxinic redox fluctuations associated with variable continental Fe delivery were observed in OAE 2 deposits from Tarfaya, Morocco; however, efficient P recycling was also observed throughout OAE 2 (Poulton et al., 2015). Further detailed and high-resolution geochemical work is, therefore, necessary to look for relationships between Fe speciation and enhanced P recycling at other Cretaceous sites that experienced prolonged periods of black shale deposition. Additionally, the relationship between changes in organic matter composition, bioturbation and Fe speciation should be determined during OAEs to evaluate whether an Fe limitation was important to OAE development in other regions, or only within the WIS.

Conclusions

Iron speciation results from OAE 3 suggest that a reactive Fe limitation may have stimulated and maintained enhanced OC burial after the initial development of anoxia within the WIS by increasing organic matter preservation in sediments and enhancing nutrient recycling. Preceding the onset of increased OC burial, Fe speciation results record a shift from predominantly non-sulfide Fe phases (carbonate, (oxyhydr)oxides, and magnetite) to
predominantly sulfide Fe phases. Intervals 2 and 3 are characterized by stable, elevated pyrite formation despite other proxies recording significant redox variability, suggesting efficient Fe sulfidization, either within surface sediments or due to downward migration of sulfidization fronts. Significant sulfidization of Fe, paired with low Fe\textsubscript{T}/Al ratios, indicates that reactive Fe may be limiting sedimentary pyrite formation.

Our results highlight two complications for using Fe as a redox proxy on geologic timescales: (1) similar to modern observations, reactive Fe is sequestered in early diagenetic carbonate phases during Interval 1a producing anomalously elevated Fe\textsubscript{T}/Al values independent of redox conditions, and (2) efficient sulfidization of reactive Fe within the Fe-limited WIS, obscured redox variability before, during and after the OAE.

A reactive Fe limitation could promote organic matter preservation in sediments because the amount of reactive Fe was insufficient to buffer porewater H\textsubscript{2}S concentrations through Fe-sulfide formation. Coeval changes in Fe, S, bioturbation and organic matter composition support that changes in Fe availability reduced the sedimentary “sulfide buffer,” reducing bioturbation and producing conditions more conducive to organic matter preservation. During Interval 1a, the availability of non-sulfide Fe phases promoted P sequestration. However, as porewater H\textsubscript{2}S accumulation increased, sulfidization of reactive Fe phases promoted P release from the sediments. This change in phosphate cycling could have prolonged enhanced OC burial during and after OAE 3 by stimulating enhanced primary productivity in the WIS, helping to extend the duration of OAE 3 as compared to OAE 2.

Distinct C-S-P cycling in the WIS during the Coniacian-Santonian, as compared to the Proto-Atlantic, during supposedly stable and widespread anoxic periods in the Late Cretaceous oceans, indicates that there was a significant amount of spatial variability between open shelf
settings, continental margins, and epicontinental seas such as the WIS. Detailed studies of stratigraphically equivalent sediment intervals in high resolution, using a range of sedimentological and geochemical tools will enable a more complete understanding of the biogeochemical cycles of the Cretaceous greenhouse ocean.

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Figure captions

Figure 1: (a) Map of the Cretaceous Western Interior Seaway (WIS) with Turonian paleobathymetry estimates (adapted from Sageman and Arthur, 1995). Darker colors represent relatively deeper depths. The USGS #1 Portland core location is marked with a star. Circles denote the Aristocrat Angus and Berthoud State cores. (b) Stratigraphy and total organic carbon (TOC) for the USGS Portland core with adapted intervals from Tessin et al. (2015). Gray bars denote Intervals 1b, 2b, and 3.

Figure 2: Iron concentrations and speciation results from the Portland core (a) Fe phases measured by sequential extractions ($\text{Fe}_{\text{carb}}$, $\text{Fe}_{\text{ox}}$, and $\text{Fe}_{\text{mag}}$), $\text{Fe}_{\text{py}}$, and $\text{Fe}_{\text{NR/PR}}$. The thick line indicates $\text{Fe}_T$ (wt. %). (b) Weight % ratios of $\text{Fe}_T$/Al. Dashed line indicates the average $\text{Fe}_T$/Al shale value of 0.55. (c) $\text{Fe}_{\text{HR}}/\text{Fe}_T$, with the anoxic threshold of 0.38 indicated by the dashed line. (d) $\text{Fe}_{\text{py}}/\text{Fe}_{\text{HR}}$, with the euxinic $\text{Fe}_{\text{py}}/\text{Fe}_{\text{HR}}$ threshold of 0.7 designated by a dashed line. Gray bars denote Intervals 1b, 2b, and 3.

Figure 3: Total organic carbon (TOC) and trace metal records ($\text{Mo}/\text{Al}$ (ppm/wt. %), $\text{Mo}$ (ppm), $\text{Mo}/\text{TOC}$ (ppm/wt. %), $\text{Re}/\text{Al}$ (ppb/wt. %), $\text{Cd}/\text{Al}$ (ppm/wt. %), and $\text{Zn}/\text{Al}$ (ppm/wt. %)) from the USGS Portland core. Gray bars denote Intervals 1b, 2b, and 3. TOC and Mo (ppm) records are from Tessin et al. (2015).

Figure 4: Total organic carbon (TOC), Total Sulfur (TS)/Al (wt. % ratios), $\text{S}_{\text{excess}}$/TS, ichnofacies-derived interpreted oxygenation curve (IOC; smoothed to 0.5 m resolution), and
Oxygen Index and Hydrogen Index values. Gray bars denote Intervals 1b, 2b, and 3. TOC and RockEval results are from Tessin et al. (2015). IOC record is adapted from Savrda (1998).

Figure 5: P/Al (weight % ratios) and TOC/P (molar ratios) records from the USGS Portland core. Dashed lines indicate the average shale P/Al ratio of 0.08 and the Redfield C:P ratio of 106:1. Gray bars denote Intervals 1b, 2b, and 3.

Figure 6: Crossplot of Fe$_{py}$/Fe$_{HR}$ and Mo (ppm; Tessin et al. 2015). Gray circles indicate samples from Interval 1; blue squares indicate samples from Interval 2 and green triangles indicate samples from Interval 3. Dashed lines indicate interpreted euxinic conditions for Mo concentrations (Scott and Lyons, 2012) and Fe$_{py}$/Fe$_{HR}$ (März et al., 2008; Poulton and Canfield, 2011).
Figure 1
Figure 2

Fe (%)

Depth (m)

Fe\_HR/Fe\_T

Fe\_py/Fe\_HR

Fe\_ox

Fe\_mag

Fe\_carb

Fe\_HR\_phases

A. B. C. D.

Interval 1

Interval 2

Interval 3

"OAE 3" Interval
Figure 3
Figure 4

The figure shows a graph with several axes and data points. The axes include:

- Depth (m)
- Total Organic Carbon (%)
- $S_{excess}/TS$
- Hydrogen Index
- Oxygen Index
- Interpreted Oxygenation Curve
- TS/Al

The graph is divided into three intervals:

- Interval 1
- Interval 2
- Interval 3

Each interval is marked with different symbols and colors for data representation.
Figure 5.
Figure 6.
Table 1: Description of iron phases and their corresponding extraction procedure adapted from März et al., 2008

<table>
<thead>
<tr>
<th>Iron Phase</th>
<th>Extraction Procedure</th>
</tr>
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<tbody>
<tr>
<td>Fe&lt;sub&gt;carb&lt;/sub&gt;: Fe bound to carbonate, including siderite and ankerite</td>
<td>10 mL 1 M Na-acetate (pH 4.5, shake for 48 hours)</td>
</tr>
<tr>
<td>Fe&lt;sub&gt;ox&lt;/sub&gt;: Fe bound as oxyhydr(oxides) including goethite and hematite</td>
<td>10 mL citrate-buffered Na-dithionite (pH 4.8; shake for 2 hours)</td>
</tr>
<tr>
<td>Fe&lt;sub&gt;mag&lt;/sub&gt;: iron bound as magnetite</td>
<td>10 mL Ammonium oxalate (pH 3.2; shake for 6 hours)</td>
</tr>
<tr>
<td>Fe&lt;sub&gt;tot&lt;/sub&gt;: total Fe including silicates</td>
<td>ALS procedure outlined above</td>
</tr>
<tr>
<td>Fe&lt;sub&gt;py&lt;/sub&gt;: Fe bound to chromium (II)-reducible sulfur</td>
<td>15 mL CrCl&lt;sub&gt;2&lt;/sub&gt; (boiling for 2 hours; precipitated in ZnS)</td>
</tr>
</tbody>
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