Redox element record shows that environmental perturbations associated with the T-OAE were of longer duration than the carbon isotope record suggests – the Aubach section, SW Germany

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With 7 figures and 2 tables

Abstract. The Early Jurassic Toarcian Oceanic Anoxic Event (T-OAE) with its associated carbon-isotope excursion (CIE) was possibly one of the most pronounced periods of widespread oxygen deficiency in the Mesozoic ocean. The event has been extensively studied in order to understand the processes triggering the environmental perturbations and the extreme oxygen depletion in many marine basins. However, comparatively little focus has been placed on the end of the positive CIE and the stratigraphic coherent end of anoxic-euxinic conditions. In the present study, we constrain the stratigraphic extent of anoxic-euxinic conditions and define the termination of the positive CIE in the Swabo-Franconian Basin covering the Lower Toarcian strata using carbon-isotope ratios, organic matter pyrolysis and redox-sensitive element concentrations of outcrop samples from the Aubach section. Bulk organic carbon-isotope values, corrected for changes in type of organic matter using the Hydrogen Index (HI), suggest that the amplitude of the negative CIE in organic matter is as little as 3.3–3.5 ‰, in contrast to 4.5 ‰ change in δ13Ccarb in the same section. Enrichment in redox-sensitive proxies (V/Al and DOP-T) and %TOC suggest that environmental perturbations associated with the T-OAE continued until the upper falciferum Zone in the Aubach section. This indicates that anoxic–euxinic conditions terminated in the same stratigraphic interval in which δ13C values return to steady, light values at ~–28 ‰ (termination of positive CIE). This synchronism in the return to normal marine conditions is also observed in the southern Paris Basin, but not in the Cleveland Basin.

Key words. Toarcian Oceanic Anoxic Event, δ13C, carbon isotope stratigraphy, redox proxies, Jurassic environment

1. Introduction

The Toarcian Oceanic Anoxic Event (T-OAE; ~182 Myr ago) represents one of the most pronounced episodes of widespread ocean oxygen deficiency in the Phanerozoic. The T-OAE resulted in enhanced organic matter (OM) burial, numerous geochemical anomalies (Frimmel et al. 2004, Jenkyns 2010, Thi...
bault et al. 2018), and global climate warming (Bailey et al. 2003, Jenkyns 2010, Korte et al. 2015, Ullmann et al. 2020). One of the most prominent geochemical features associated with the T-OAE is the fluctuation in the δ13C record. A long-lasting positive carbon isotope excursion (pCIE) (Hesselbo et al. 2000, Hermoso et al. 2009a, Hermoso et al. 2009b, Hermoso et al. 2012) is marked by a large-scale negative CIE (nCIE) (Fig. 1A) (Jenkyns et al. 2002, McArthur et al. 2008, Korte et al. 2015). Previous work has shown that the Early Toarcian nCIE commenced in the upper tenuicostatum/polymorphum Zone (ammonite biozonation) and culminated in the lower falciferum/serpentinitum Zone as shown by the δ13C reference curve for the Early Toarcian based on published sections (Ruebsam and Al-Husseini 2020). The nCIE has been detected in marine and terrestrial OM and marine carbonate from successions across Europe (Jenkyns 1988, Röhl et al. 2001, Kemp et al. 2005, Suan et al. 2008, Hermoso et al. 2009a, Hermoso et al. 2009b), N-Africa (Krencker et al. 2020), S-America (Fantasia et al. 2018a, Al-Suwaidi et al. 2010), N-America (Them et al. 2017a), Japan (Gröcke et al. 2011, Kemp et al. 2019), and Arctic Russia (Suan et al. 2011), indicating that the global carbon cycle was greatly influenced.

Ocean anoxic events have traditionally been defined on the basis of widespread deposition of laminated black shale (Jenkyns 1988, Jenkyns 2010), yet the %TOC in a succession may be heavily influenced by

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**Fig. 1.** A) The δ13C reference curve by Ruebsam and Al-Husseine (2020) modified to include the Pliensbachian-Toarcian nCIE, termination of the T-OAE and end of the Toarcian pCIE. B) Paleogeographic map of the Laurasian Seaway in the Early Jurassic with the studied locality, the Aubach section (star) situated in the Swabo-Franconian Basin, Cleveland Basin: C and Paris Basin: P (modified from e.g. Ruebsam et al. 2018) Emerged landmasses (green), organic-rich sediment (grey, darker grey represent a higher %TOC).
local effects, and thus the understanding of the T-OAE as a global climate perturbation contrasts to a definition applicable only to a limited set of palaeoenvironments. Such a definition is therefore not optimal for global considerations (Wignall et al. 2005) as global climate change likely transcended such restricted definitions. Now, generally, the T-OAE is defined by patterns in the coeval C-isotope curve, e.g., the onset limb of the positive the δ13C perturbation caused by the widespread burial of organic matter (cf. Jenkyns 2010) (Fig. 1A). This definition sets the beginning of the OAE where δ13C values start to change and the end at the climax of the positive δ13C excursion (pCIE) values decrease. The Toarcian pCIE is interrupted by the nCIE (e.g., Hermoso et al. 2009), which can prove it challenging to recognize its beginning.

When the end of the T-OAE is based on the stratigraphic datum where the most positive δ13C values are reached, it terminates in the early serpentinum/falciferum Zone (Hesselbo 2008, Hermoso et al. 2013). Yet, previous studies show that black shale formation was not restricted to the lower falciferum Zone in the Swabo-Franconian Basin (Fantasia et al. 2018a, Röhl and Schmid-Röhl 2005, Van de Schootbrugge et al. 2013). Furthermore, other geochemical proxies indicate that environmental and climatic conditions were still perturbed after the nCIE, far from having returned to pre-excursion levels in many sections. Organic matter characteristics, trace element and biomarker data from the nearby Dotternhausen section (Schouten et al. 2000, Schwark and Frimmel 2004, Suan et al. 2015, Röhl et al. 2001, Them et al. 2018, Dickson et al. 2017) and Swiss successions (Fantasia et al. 2018b, Montero-Serrano et al. 2015) show that intense oxygen-depletion continued in the Swabo-Franconian Basin and that redox conditions became more oxidized on the continental margin after the carbon isotope anomaly. In the Paris Basin (France) and the Cleveland Basin (UK) enrichment in redox-sensitive elements indicate that anoxia also continued beyond the nCIE and lasted into the upper falciferum Zone (Hermoso et al. 2013, Thibault et al. 2018). This all states well that the environmental perturbation of the T-OAE played out differently from basin to basin. However, comparatively little focus has been placed on how the continuation of anoxia and high %TOC correlate with the return to steady light δ13C values at the end of the Toarcian pCIE.

In the present study, we investigate the evolution of the seawater chemistry through the Early Toarcian in the Aubach section (Swabo-Franconian Basin, SW Germany) in relation to the stratigraphic extent of the Toarcian nCIE and the termination of the Toarcian pCIE by combining δ13C and TOC(wt%) with redox-sensitive element concentrations using new high-resolution geochemical data. The evaluation of the nCIE and termination at the peak pCIE is based on δ13Corg and δ13C_carb as well as on changes in OM sources. The degree of pyritization based on total Fe (DOP-T) and V/Al, among various geochemical proxies for anoxic/euxinic conditions, have been selected to assess the stratigraphic extent of anoxia in Aubach.

2. Geological setting

The Aubach section is located in the Wutach river area (Baden-Württemberg, SW Germany) between the Swabian Alps and the Black Forest. The outcrop succession consists of Lower Jurassic sediments deposited in the shallow-marine Swabo-Franconian Basin that was part of the Laurasian Seaway, which connected the Tethys to the Boreal Ocean (Fig. 1B) (Bjerrum et al. 2001). In the Early Jurassic, this seaway bordered the northwestern part of the Tethyan epicontinental shelf in the South and the Boreal Sea in North (Fig. 1; Bailey et al. 2003).

The section consists of marls with limestone horizons in the Pliensbachian spinatum Zone (Fig. 2). The Lower Toarcian interval of the section comprises bioturbated light-grey marl constituting the Blaugraue Mergel and Aschgraue Mergel interbedded by two laminated shale beds in the semicelatum Zone. The lower and middle falciferum/serpentinitum Zone consist of dark laminated shales characterized by distinct wavy to lenticular microlamination with two thick limestone horizons. Benthic fauna is absent (Riegraf et al. 1984). A predominance of dark claystone, black shale and condensation/discontinuity surfaces demonstrate a general transgressive trend in these Lower Jurassic strata (Pienkowski et al. 2008).

The Lower Jurassic strata of the Aubach section became exposed during the development of the west European rift system in the Oligocene resulting in the uplift of the Black Forest massif (Ziegler 1992, Demoulin et al. 1998). Rivers subsequently cut deep and narrow incisions during the Quaternary, and recent land-slides have served to produce excellent exposures of the Lower Jurassic succession in the investigated region (Demoulin et al. 1998).
3. Material and Methods

3.1. Sampling

Biostratigraphic subdivision for the Aubach outcrop section is taken from Riegraf et al. (1984), which is the most recent zonation available for this section. A total of 123 bulk-rock samples were collected from uppermost Pliensbachian to Lower Toarcian sediments with a resolution of ~20 cm in the uppermost Pliensbachian, 5–10 cm in the lower Toarcian and 20–100 cm in the higher *elegans* subzone. During sampling, the outermost weathered surface sediment was removed, and only fresh material was collected.

3.2. Stable isotopes

Stable isotope analysis was performed on bulk rock samples using an IsoPrime triple collector Isotope Ratio Mass spectrometer at the Department of Geosciences and Natural Resource Management at the University of Copenhagen (IGN). A total of 72 and 160 samples were analysed to determine δ¹³Ccarb/δ¹⁸Ocarb and δ¹³Corg, respectively.

Crushed sample material, equivalent to 0.6 mg carbonate, was placed into 3.5 ml glass Labco vials and sealed with plastic lids for δ¹³Ccarb and δ¹⁸Ocarb analysis. Vials were flushed with helium to remove atmospheric gasses and subsequently reacted with nominally anhydrous phosphoric acid at 70 °C for 90 min. The measured ¹³C/¹²C and ¹⁸O/¹⁶O ratios were normalized against a CO₂ reference gas. The reproducibility of the analyses was 0.12 ‰ for δ¹³C (2σ) and 0.16 ‰ for δ¹⁸O (2σ) based on 15 standards to 45 samples in each batch of analyses. Isotope ratios were corrected with the in-house reference material LEO (δ¹³C = +1.96 ‰; δ¹⁸O = –1.93 ‰; see Ullmann et al. 2013).

For organic δ¹³C analysis, powdered samples were decarbonated by placing the sample in a 50 ml polypropylene centrifuge tube and treating with 10 % HCl for 1 hour at 60 °C. The samples were then rinsed with demineralized water, centrifuged, and rinsed again until neutrality was reached (following the method of Gröcke et al. 1999). These decarbonated samples were then freeze-dried to remove potential liquids. Decarbonated bulk rock material with a weight equivalent to 60 µg organic carbon was packed in tin capsules for δ¹³Corg analysis. Samples were fluxed in an oven heated to 1100 °C, where oxygen was added to react the OM to CO₂. The measured ¹³C/¹₂C ratios were corrected against the in-house reference AKsill_9 (δ¹³C = −25.30 ‰; 18 standards to 57 samples). Carbon-isotope ratio reproducibility was found to be 0.07 ‰ (2σ of repeat measurements of the lab standard).

The in-house reference material LEO (δ¹³C 1.96 ‰ and δ¹⁸O −1.93 ‰) and AKsill_9 (−25.30 ‰) are calibrated to the international reference materials NBS-18 and NBS-19 and USGS-24 respectively, ensuring accurate determination of isotope ratios.

3.3. Organic geochemistry

Bulk-rock samples were crushed using a mill and sieved to grain sizes smaller than ~250 μm. Organic matter pyrolysis was carried out on 100 mg size at the Department of Reservoir Geology, Geological Survey of Denmark and Greenland (GEUS) using a HAWK instrument (Wildcat Technologies, USA). This instrument was calibrated using the IFP 16000 standard reference material to yield results similar and comparable to those obtained by the Rock-Eval 6 instrument. Total Organic Carbon contents (TOC wt%), Hydrogen Index (HI) and Oxygen Index (OI) were determined.

The calcium carbonate concentrations of 123 samples were determined by calculating the loss of weight after decarbonisation. A subset of 93 of these decarbonated samples were analysed for TOC and Total Sulphur (TS) using an Eltra CS 500 Analyzer at IGN.

3.4. X-ray fluorescence (XRF) analysis

XRF analysis was undertaken to determine the concentrations of redox-sensitive elements. XRF measurements were performed on 120 powdered samples stored in plastic containers covered with 0.016 millimetre thick polypropylene film following Lenniger et al. (2014). Element concentrations of the samples were determined using a hand-held Niton TmX13tGoldd+ XRF device (hh–XRF) equipped with an Ag anode (measurements at 6–50 kV and 0–200 μA max), and inductively coupled plasma mass spectrometry (ICP-MS) at GEUS. The geochemical composition of 120 samples was determined based on three replicates measured for 100 seconds using the ‘test all geo filter’ program on the hh–XRF that allows determination of 32 elements. Correction-factors for hh–XRF data against ICP-MS measurements and sulphur concentration to %TS data, are all tightly constrained (Table 1).
4. Results

4.1. Stable isotopes

Organic δ\(^{13}\)C data vary between −24.5 ‰ and −32.4 ‰ (Fig. 2). Two short-term negative excursions of ~2.3 ‰ are observed at the Pliensbachian–Toarcian boundary. Marked negative excursions occur coeval with the black shales in the *semicelatum* Zone (~4 ‰ in the Tafelfleins and ~2 ‰ in the two Seegrasschiefer shales). The pronounced and well-known Early Toarcian nCIE commences in the uppermost *semicelatum* Zone and is here characterized by four steps (Fig. 2). This CIE culminates in the *exaratum* subzone reaching a total decrease of 6.3 ‰ (lowest value −32.4 ‰) after which δ\(^{13}\)C\(_{\text{org}}\) values steadily increase. δ\(^{13}\)C\(_{\text{org}}\) values reach the plateau of the Toarcian pCIE of −26.6 ‰ to −25.6 ‰ in the upper part of the *elegans* subzone and decrease to ~−28 ‰ in the lower part of the *falciferum* subzone, marking the end of the Toarcian pCIE (Fig. 2).

The δ\(^{13}\)C\(_{\text{carb}}\) values range from −6.5 ‰ to +1.7 ‰. The δ\(^{13}\)C\(_{\text{carb}}\) profile fluctuates around 0 ‰ in the upper *spinatum* Zone and records minimum values of −2.5 ‰ in CaCO\(_3\)-rich beds occurring immediately below the Pliensbachian-Toarcian transition (Fig. 2). In the lower *semicelatum* Zone, δ\(^{13}\)C\(_{\text{carb}}\) values increase to ~0.6 ‰ and further to ~1.3 ‰, where a significant drop of 4.5 ‰ interrupts the positive trend. The δ\(^{13}\)C\(_{\text{carb}}\) minimum values of ~−3.2 ‰ and ~−2 ‰ in the *exaratum* subzone occur just below and above the Unterer Stein, respectively (Fig. 2). Very light (~−6.5 ‰) δ\(^{13}\)C\(_{\text{carb}}\) values occur in the middle of the Unterer Stein limestone bed and record an isotopic fluctuation of 7.8 ‰ in amplitude (Fig. 1S). These extremely light δ\(^{13}\)C\(_{\text{carb}}\) values of the limestone are regarded as anomalies and have been excluded from Fig. 2.

The δ\(^{18}\)O\(_{\text{carb}}\) values decrease generally from ~−5.3 ‰ to ~−6.8 ‰ across the full section and display high variability in the Unterer Stein ranging from −8.0 ‰ to −3.8 ‰ (Fig. 1S).

4.2. Organic matter

All samples show T\(_{\text{max}}\) of 420–435 °C (Table S2) corresponding to a level of thermal maturity just below the oil window (435–470 °C) (Hunt 1996). Organic matter types have been classified using the Van Krevelen-type diagram after Hunt (1996); HI as a function of OI (mg HC/g TOC; Fig. 3). Marl and carbonate layers of the *spinatum* Zone and *semicelatum* Zone include kerogen type III with low HI (<150 mg HC/g TOC) and high OI values (25–185 mg CO\(_2\)/g) (Figs. 2 and 3). The OM of the two Seegrasschiefer shales and the Tafelfleins shale (*semicelatum* Zone) mainly contain Type II kerogen (300–600 mg HC/g TOC), and shales of the lower part of the *falciferum* Zone contain a mix of kerogen Type I and II with high HI (300–830 mg HC/g TOC) and low OI values (<80 mg CO\(_2\)/g TOC). Previous studies have established the marine origin of the Lower Toarcian strata in SW Germany (Küspert 1982, Moldowan et al. 1986, Schouten et al. 2000, Röhl et al. 2001, Frimmel et al. 2004) where kerogen I (HI >600) therefore indicates marine algae. Kerogen Type II is derived from marine plankton and kerogen Type III from oxygenated marine plants. Type III kerogen samples all display low TOC (0.5–1 wt%), whereas Type I–II kerogen samples range from 5–14 wt% TOC with the highest TOC in the Tafelfleins shale (13 wt%) and during the Early Toarcian nCIE (up to ~14 wt%). Above the nCIE, TOC decreases to 5–7 wt% and further to 4–5 wt%, coinciding with δ\(^{13}\)C\(_{\text{org}}\) value stabilizing at ~−28 ‰ in the upper *falciferum* Zone (Fig. 2).

To test whether the changing source or maturity of OM affected the δ\(^{13}\)C\(_{\text{org}}\) data, Spearman’s rank correlation was used to assess the data (δ\(^{13}\)C\(_{\text{org}}\) vs T\(_{\text{max}}\): ρ=0.41, p<0.01, n=70) (Table 2). The data was assessed as a whole, and also subdivided into pre-exursion (pre-CIE samples originating from the *spinatum* to the upper *semicelatum* Zone) and non-exursion (all samples outside the *falciferum* zone), as the

<table>
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<th>Element</th>
<th>Slope (a)</th>
<th>Intercept (b)</th>
<th>(r^2)</th>
<th>p-values &lt;0.01</th>
<th>N</th>
<th>r</th>
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<td>Fe (ppm)</td>
<td>0.9718</td>
<td>3616</td>
<td>0.9836</td>
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<td>0.99</td>
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<tr>
<td>V (ppm)</td>
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<td>−17.97</td>
<td>0.8959</td>
<td>yes</td>
<td>9</td>
<td>0.95</td>
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<tr>
<td>Al (ppm)</td>
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<td>−7802.4</td>
<td>0.9927</td>
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<td>S (ppm)</td>
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<td>3401</td>
<td>0.8435</td>
<td>yes</td>
<td>62</td>
<td>0.92</td>
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Table 1. Correction-factors for hh-XRF data against ICP-MS measurements and sulphur concentration to %TS data.
Fig. 2. Lithology, stratigraphy, δ\(^{13}\)C\(_{\text{carb}}\) and δ\(^{13}\)C\(_{\text{org}}\) values and HI-corrected δ\(^{13}\)C\(_{\text{org}}\) trends of the Aubach succession. δ\(^{13}\)C\(_{\text{carb}}\) and HI-corrected δ\(^{13}\)C\(_{\text{org}}\) show a comparable amplitude of the nCIE of −4.5 ‰ and 3.3–3.5 ‰, respectively. Note that HI-corrected δ\(^{13}\)C\(_{\text{org}}\) as well as the δ\(^{13}\)C\(_{\text{org}}\) reach steady light values of −28 ‰ in the upper falciferum Zone, concurrent with TOC concentrations of more than 5 wt%. The source of organic matter continues to be of kerogen Type II from the base of the falciferum Zone to top of the Early Toarcian. The shaded area marks the uncertainty of the HI-corrected based on the two corrections used. δ\(^{13}\)C\(_{\text{carb}}\) values marked by empty circles may be diagenetic.
The effect of OM source on $\delta^{13}C$ may be different across the excursion, as demonstrated by Lenninger et al. (2014) and Suan et al. (2015). As a whole, the HI values (an analogue for OM source) showed a moderate but significant relationship to $\delta^{13}C$ ($\rho=-0.64$, $p<0.01$, $r^2=0.42$, $n=70$). However, excluding the CIE data, the pre-CIE data and data outside the falciferum Zone show a significant, moderate to strong correlation between HI and $\delta^{13}C$ ($\rho=-0.57$, $p<0.01$, $r^2=0.68$, $n=31$ and $\rho=-0.70$, $p<0.01$, $r^2=0.68$, $n=37$, respectively) (Table 2).

The significant moderate/strong correlation between HI and $\delta^{13}C$ in the data (excluding the CIE interval) indicates that the $\delta^{13}C_{\text{org}}$ signal is influenced by OM source (e.g. Tyson 1995). Lenninger et al. (2014), and Suan et al. (2015) developed a correction that compensates for fluctuations in $\delta^{13}C_{\text{org}}$ caused by changes in OM type through establishing the background of HI and $\delta^{13}C_{\text{org}}$ in the basin, via the non-excursion HI-$\delta^{13}C$ linear relationship (Fig. 4, Table 2).

We estimated the relationship for the Aubach section

<table>
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<td>$n$</td>
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<td>70</td>
<td>37</td>
</tr>
<tr>
<td>$\rho&lt;0.01$</td>
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<td>yes</td>
<td>yes</td>
</tr>
<tr>
<td>$r^2$</td>
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<td>0.42</td>
<td>0.68</td>
</tr>
<tr>
<td>$a$</td>
<td>0.005</td>
<td>0.004</td>
<td>0.004</td>
</tr>
<tr>
<td>CIE amplitude</td>
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<td>$-5.58,^\circ\text{C}$</td>
<td>$-3.5,^\circ\text{C}$</td>
</tr>
</tbody>
</table>
and applied the resulting correction to our $\delta^{13}C$ data. The outcome is a reduction in the magnitude of the nCIE from 6.3 ‰ resulting in a negative $\delta^{13}C$ excursion of only 3.3–3.5 ‰ (Fig. 2). After correction the pCIE reach a maximum $\delta^{13}C_{org}$ values of $-25.6$ ‰ corresponds to an increase of HI-corrected $\delta^{13}C_{org}$ to $+3.5$ ‰.

4.3. Element data

4.3.1. Vanadium

The vanadium concentration varies over the investigated section from below the detection limit (<3 ppm) to 270 µg/g (Fig. 5). The V-enrichment is determined relative to average shale composition (V = 130 ppm and V/Al = 14.8 µg/g/wt%) (Wedepohl 1971). Vanadium content is near or below the detection limit in the limestone beds (upper spinatum bed, Unterer Stein, Oberer Stein). V-enrichment occurs in the Tafelfleins and Seegrasschiefer shales (semicelatum Zone) and from the first step of the negative CIE in the upper semicelatum Zone. In the falciferum Zone, the V/Al ratio increases to ~36 and gradually declines to ~10 above the Oberer Stein, coinciding with the disappearance of laminated black shale. A significant V-enrichment of V/Al ratios up to ~37 is found in the elegans subzone.

4.3.2. Sulphur concentration and DOP-T (S/Fe)

The sulphur concentration varies in the succession from ~1 wt% to 4.5 wt% (Fig. 5). Sulphur concentrations of >2 wt% only occur in the black shales in the semicelatum Zone, and coincide with high %TOC. S/Fe is referred to as degree of pyritization based on total Fe (DOP-T), which has been proposed as a simple, good indicator for redox conditions during deposition (Algeo and Maynard 2008, Berner et al. 2013). DOP-T 0–0.25 indicates oxic, 0.25–0.50 indicates suboxic–dysoxic conditions, and DOP-T >0.50 corresponds to anoxic–euxinic conditions (Algeo and Maynard 2004). DOP-T values have been excluded in the dataset where TOC <0.5 wt% (Algeo and Maynard 2008). DOP-T of 0.25–0.50 is thought to indicate dysoxic conditions in the spinatum and semicelatum zones, while DOP-T of 0.60–0.70 reflects anoxic–euxinic conditions in the semicelatum Zone black shales. DOP-T increases significantly with the initiation of the Early Toarcian nCIE and remains between

Fig. 4. The Hydrogen Index (HI)-correction of $\delta^{13}C_{org}$ accounts for changes in organic matter in the Early Toarcian. The linear relation between HI and $\delta^{13}C_{org}$ of pre-CIE samples (red), samples outside falciferum Zone (green) and all samples including the CIE. Our HI-correction is based on $\delta^{13}C_{org}$ as a function of HI, based on samples pre-CIE and samples outside the falciferum Zone.
0.50 and 1 until the *elegans-falciferum* subzone boundary, and fluctuates around 0.50 up to the top of the investigated interval, indicating slightly improved oxygen availability.

5. Discussion

5.1. The Toarcian $\delta^{13}$C record in the Swabo-Franconian Basin

The trends in $\delta^{13}$C$_{\text{carb}}$ and $\delta^{13}$C$_{\text{org}}$ co-vary throughout the succession indicating that an environmental imprint on the carbon isotope signal is dominant in both carbonate and organic matter. The Toarcian $\delta^{13}$C record of the Aubach section has been correlated to the $\delta^{13}$C reference curve by Ruebsam and Al-Husseini (2020) (Fig. 6), in order to define the stratigraphic extent of the Toarcian nCIE and pCIE in Aubach based on a compilation of $\delta^{13}$C$_{\text{org}}$, $\delta^{13}$C$_{\text{carb}}$ and HI-corrected $\delta^{13}$C (Fig. 6). The correlation shows the onset of the nCIE in the uppermost part of the *semicelatum* Zone, where $\delta^{13}$C values start to decrease in four to five abrupt steps, culminating in the lower *exaratum* subzone (HI-correction of $\delta^{13}$C$_{\text{org}}$ removes a step). The nCIE terminates below the Oberer Stein in the lower *elegans* subzone, where $\delta^{13}$C values reach a plateau.

5.1.1. The positive CIE

The Toarcian pCIE is interrupted by the nCIE (e.g., Hermoso et al. 2009), which can make it challenging to recognize its beginning. In Aubach, the $\delta^{13}$C$_{\text{carb}}$ record increases to ~0.6 ‰ with the first occurrence of black shale in the *semicelatum* Zone and further to +1.3 ‰ by the initiation of long-term black-shale deposition in the upper *semicelatum* Zone (Fig. 2). One of these
increases in $\delta^{13}C_{\text{carb}}$ values may mark the beginning of the Toarcian pCIE. Hermoso et al. (2013) argued that the Toarcian pCIE initiated at the Pliensbachian–Toarcian boundary, based on increasing $\delta^{13}C_{\text{carb}}$ values in the Paris Basin (Fig. 7). Thallium isotopes from Alberta (Canada) support this, showing evidence for global marine deoxygenation of ocean water beginning at the Pliensbachian–Toarcian boundary (Them et al. 2018). Alternatively, the beginning of the Toarcian pCIE may occur already in the late Pliensbachian period as suggested by previous work (e.g. Ruebsam and Al-Husseini 2020).

The Toarcian pCIE culminates in a plateau at $\delta^{13}C_{\text{org}}$ values of $\sim$26.5 to $\sim$25.6‰, followed by the falling limb just above the Oberer Stein. The $\delta^{15}C_{\text{org}}$ record stabilizes at $\sim$28‰ in the upper part of the falciferum Zone (Fig. 6). The Toarcian pCIE reaches HI-corrected $\delta^{13}C_{\text{carb}}$ values of +3.5‰, somewhat similar to $\delta^{13}C_{\text{carb}}$ negative excursion recorded in Spanish and Portuguese brachiopods +5.1‰ and +5.8‰, respectively (Ullmann et al. 2020). The stabilization of the $\delta^{13}C_{\text{org}}$ record at around $\sim$28‰ marks the return to normal marine $\delta^{13}C_{\text{org}}$ conditions in the Aubach sediments (Fig. 6). The return to near-steady light $\delta^{13}C$ values recorded in the $\delta^{13}C$ reference curve span from the uppermost falciferum Zone to the lower part of the bifrons Zone (Ruebsam and Al-Husseini 2020). Thus, carbon-isotope values stayed relatively heavy for a substantial period of time after the culmination of the Toarcian pCIE. The elevated $\delta^{13}C$ values may be a result of continued widespread organic matter burial, as high TOC values in levels post-dating the nCIE have been documented in several basins in Europe (this study, Röhl et al. 2001, Hermoso et al. 2013), Canada (Them et al. 2017a) and in Northern Siberia (Suan et al. 2011). The interval of high TOC post nCIE has been

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**Fig. 6.** The $\delta^{13}C$ profile of Aubach correlated to the $\delta^{13}C$ reference curve by Ruebsam and Al-Husseini (2020) to determine the stratigraphic extent of the Toarcian nCIE and pCIE. Note that the termination of the Toarcian pCIE has been set in the upper falciferum Zone $\delta^{13}C_{\text{org}}$, stabilizing at $\sim$28‰. The beginning of the Toarcian pCIE has not been determined.
estimated to have lasted 600–1800 kyr in Northern Siberia (Suan et al. 2011).

5.2. The magnitude of Toarcian negative CIE

The exact magnitude of the nCIE is difficult to determine as differences in amplitude occur in different carbon reservoirs. This has made it challenging to estimate the amount of $^{13}$C-depleted carbon needed to create such a significant negative excursion as the Toarcian CIE (Kemp et al. 2005, Suan et al. 2015).

Our new $\delta^{13}$C$_{org}$ profile for the Aubach section displays a nCIE magnitude of $-6.3$ ‰. Other European sections record (uncorrected for OM source) similar amplitudes of $-6$ ‰ to $-7$ ‰ in the $\delta^{13}$C$_{org}$ of bulk sedimentary organic matter (Jenkyns and Clayton 1997, Hesselbo et al. 2000, Röhl et al. 2001, Kemp et al. 2005, Hesselbo and Pienkowski 2011, Xu et al. 2018). Some researchers examined only the wood record in order to avoid bias introduced by OM source mixing, yet amplitudes may be as high as $-8$ ‰ (Hesselbo et al. 2007, Pieńkowski et al. 2016, Hesselbo et al. 2000). However, degradation of fossil wood can alter $\delta^{13}$C depending on preservation stage (Ruebsam et al. 2020, Hesselbo et al. 2007). Changes in the hydrological cycle may also play a role and water availability has shown to affect the leaf gas-exchange and thereby cause up to 6 ‰ bias in $\delta^{13}$C of terrestrial plants during the PETM (Diefendorf et al. 2010). Increasing atmospheric CO$_2$ ($p$CO$_2$) may also affect terrestrial plant $\delta^{13}$C, resulting in higher isotopic fractionation if $p$CO$_2$ is more than 500 ppm (Schubert and Jahren 2013). Ruebsam et al. (2020) estimated $p$CO$_2$ during the nCIE to increase to as much as 1700 ppm. Magnitude variations of the nCIE between terrestrial and marine plants are seen in compound specific isotope ratios: $\delta^{13}$C of short-chain n-alkanes (marine indicators) give $\sim$2–3 ‰ whereas the $\delta^{13}$C of long-chain n-alkanes (terrestrial plants) is $\sim$4–5 ‰ in Yorkshire, UK (French et al. 2014).

Interestingly, the magnitude of the nCIE in the $\delta^{13}$C$_{carb}$ record varies between individual sections. The magnitude of the nCIE is $\sim$4.5 ‰ in our record from Aubach, up to 6 ‰ in successions of the Paris Basin (Sancerre-Couy) and Mochras drillcore, Wales (Jenkyns et al. 2002), where as it is only $\sim$3.5 ‰ at Peniche (Hermoso et al. 2009b). Differences in magnitude of the $\delta^{13}$C$_{carb}$ may be attributed to the sensitivity to diagenesis or to local effects. The latter can be significant especially in deposits of shallow marine epicontinental seas, such as the Laurasian Seaway, in which the dissolved inorganic carbon $\delta^{13}$C is more sensitive to changes in the water column compared to pelagic sediments of the deep ocean (Immenhauser et al. 2008). The carbonate carbon-isotope ratios from the Unterer Stein limestone bed are very low, and the $\delta^{18}$O$_{carb}$ signals are highly variable, both suggesting that these samples are diagenetically altered. These samples have therefore been excluded from the interpretation (Fig. S1).

The drivers for the Toarcian nCIE are the subject of controversial discussion. Several causes have been proposed over the years. One of the first, suggested by Küspert (1982), was the recycling of isotopically light carbon in the restricted water column. Volcanic CO$_2$ sourced from the Karoo Large Igneous Province may have injected $^{13}$C-depleted into the atmosphere causing the nCIE (Hesselbo et al. 2007, Hermoso et al. 2009b). However Gröcke et al. (2009) reject magmatism as the cause for the nCIE as $\delta^{13}$C values from volatile matter and vitrinite reflectance from two coal transects associated with dykes from the Karoo Basin, S Africa show no significant change in $\delta^{13}$C (which would be expected by thermogenic CH$_4$ generation). Data indicate that the dyke intrusions occurred when the coal still had a high moisture content prohibiting CH$_4$ generation (Gröcke et al. 2009). Alternatively, an injection of $^{13}$C-depleted carbon from methane release from various reservoirs (gas hydrates, wetlands, permafrost or from fossil wood degradation) may be responsible for the Toarcian nCIE, possibly secondarily paced by astronomical-forcing (Hesselbo et al. 2000, Them et al. 2017a, Ruebsam et al. 2019, Kemp et al. 2005, Pieńkowski et al. 2016).

5.2.1. HI-correction of $\delta^{13}$C$_{org}$

Schouten et al. (2000) established that the difference in magnitude of the nCIE between carbon reservoirs was not a result of changes in a photosynthetic fractionation factor caused by increasing $p$CO$_2$ during the T-OAE, because marine molecular-compound specific $\delta^{13}$C of the OM displayed similar amplitudes as $\delta^{13}$C$_{carb}$. However, fluctuations in OM source affect $\delta^{13}$C$_{org}$ (Van de Schootbrugge et al. 2013) and may account for the difference between marine $\delta^{13}$C$_{org}$ and $\delta^{13}$C$_{carb}$; this can be corrected for by applying the HI-correction as in Lenniger et al. (2014) and Suan et al. (2015; see results).

The matrix effect of high carbonate concentrations can lower the HI value and thereby affect the accuracy
of the HI-correction (Suan et al. 2015). Yet this is not the case in the Aubach section as strong HI similarities occur in coeval sediments of other sections in the vicinity of the Wutach area (e.g., Denkingen and Dotternhausen, SW Germany; Suan et al. 2015). By applying this HI-correction on δ13Corg, the amplitude of the nCIE is reduced to 3.3–3.5 ‰ in Aubach (Fig. 2). Thus, the excursion magnitude becomes comparable to that observed in δ13Ccarb (4.5 ‰), the brachiopod record of Spain (4.2 ‰; Ullmann et al. 2020) and marine and lacustrine compound specific isotopes (2–4 ‰; French et al. 2014, Schouten et al. 2000, Xu et al. 2017). Three other German sections, Denkingen and Dotternhausen (Suan et al. 2015) and Rietheim (Montero-Serrano et al. 2015) report nCIE of 3–4 ‰ after HI-correction, and is in concert with the δ13Ccarb fluctuation in all the sections. Thus, correction for changing in OM seems of great importance in understanding the severe environmental perturbation. For comparison with other successions, we will discuss the uncorrected δ13Corg curve, as the uncorrected curve relates directly to local environmental changes such as redox.

5.3. Environmental changes in the Early Toarcian in Aubach

A selection of redox sensitive elements (V/Al and DOP-T) were used to determine the redox conditions in the sediments of the Aubach section (Fig. 5). Vanadium is fixed in sedimentary organic matter in anoxic waters, when V(V) is reduced to V(IV), which is adsorbed onto settling organic compounds (Breit and Wanty 1991). V is remobilized back into the water column when OM is oxidized for dysoxic waters above and below modern anoxic zones in the oceans (Wang et al. 2017). DOP-T (S/total Fe) is a simple, good indicator for redox conditions during deposition based on the degree of pyritization (Algeo and Maynard 2008, Berner et al. 2013). The formation of pyrite is limited by the abundance of reactive Fe and H2S in local environment. Framboidal pyrite precipitating in the euxinic water column/pore water forms via a metastable phase, Fe-sulfide (FeS), and then transforms into pyrite (FeS2) if the sulfur species are available. Sulphate-reducing bacteria control the production of sulfur and sulfides available for pyrite formation (e.g., Berner et al. 2013).

In general suboxic to dysoxic conditions characterized the semicelatum Zone as indicated by DOP-T of 0.25–0.5. Simultaneously, low V/Al ratios combined with low %TOC, Kerogen Type III OM and the marly bioturbated lithology support the more oxygenated conditions in the Swabo-Franconian Basin prior to the nCIE. Most likely the V was remobilized post-depositionally during OM oxidation explaining the relationship between low TOC and low V concentrations (Fig. 5). The discrete black shale horizons within the semicelatum Zone coincide with high V/Al ratios and DOP-T >0.50, supporting the interpretation of short intervals of anoxia (Röhl et al. 2001). The HI-correction removes the light δ13Corg values occurring in the shales, suggesting a strong lithological control on the organic matter (kerogen type). These short-term intervals of oxygen deficiency are also recognized in some basins: Yorkshire, UK (McArthur et al. 2008, Thibault et al. 2018) and Dotternhausen (Röhl et al. 2001), which may suggest a connection between these basins or concurrent local outbreaks of anoxia.

5.3.1. Expression of the T-OAE in the Aubach section

We adopt the definition of an OAE by the positive δ13C perturbation, setting the beginning of the OAE where δ13C values start to increase and the end at the climax of the rising δ13C limb of the positive excursion (Jenkyns 2010). This definition sets the end of the T-OAE in the lower falciferum Zone near the Oberer Stein (Fig. 6). Long-term V/Al enrichment and DOP-T >0.5 indicates that anoxia–euxinia initiated in the uppermost semicelatum Zone, synchronous with the first negative step in δ13Corg (Fig. 5). Thus, long-lasting anoxia and enhanced OM burial in the Aubach section is delayed compared to the beginning of the T-OAE, and may be directly related to the environmental changes linked to the nCIE. Global warming in Early Toarcian led to enhanced continental weathering, especially during the CIE (Them et al. 2017b, Cohen et al. 2007, Percival et al. 2016, Xu et al. 2018). The increased weathering boosted the supply of phosphorous and other nutrients to the ocean, which enhanced the primary productivity in the photic zone (Röhl et al. 2001). Oxidation of this OM-lean to oxygen deficiency deeper in the water column, and even photic zone euxinia (Ruebsam et al. 2018).

Vanadium usually accumulates in the sediment fixed in organic compounds. This may explain the low V/Al ratios in the two organic matter-lean limestone beds in the anoxic–euxinic interval (Fig. 5). Aryl isoprenoid and isorenieratane biomarkers occur in the Un terer Stein and Oberer Stein in the nearby Dotternhausen
Fig. 7. Correlation of the Aubach section (Swabo-Franconian Basin), Sancerre-Couy core (Paris Basin; Hermoso et al. 2013) and Yorkshire section (Cleveland Basin; Littler et al. 2010, Thibault et al. 2018) based on the correlation in Ruebsam and Al-Husseini (2020). The correlation to the Rietheim section (Swiss Jura; Montero-Sancarro et al. 2015) is based on their definition of the nCIE and the return to steady light $\delta^{13}$C$_{org}$. The coherence between termination of the Toarcian pCIE and end of TOC $>5\%$ and anoxic–euxinic conditions in the upper *falciferum-serpentinum* Zone occurs in the Aubach section and in the Sancerre-Couy core. However, in the Yorkshire section DOP-T of 0.5–1 continues after $\delta^{13}$C$_{org}$ has stabilized, and in the nearby Rietheim section V-enrichment and TOC $>5\%$ only occurs during the nCIE.
section, indicating that the limestone beds formed during photic zone anoxia–euxinia (Schouten et al. 2000, Schwark and Frimmel 2004). V/Al enrichment and DOP-T of 0.5–1 suggest that anoxic–euxinic conditions prevailed until the late *falciferum* Zone in the Aubach section, accompanied by TOC content >5% (Fig. 5). Thus, anoxia and deposition of organic-rich shales continued after the defined end of the OAE in the lower *falciferum* Zone (Fig. 6). The continuation of anoxia after the maximum of the positive excursion has also been observed in nearby Dotternhausen section and in Sancere Couy, Paris Basin (Röhl et al. 2001, Hermoso et al. 2013). Thallium isotopes from Dotternhausen remain high throughout the Early Toarcian suggesting that anoxia continued after enhanced organic matter burial ended (Them et al. 2018).

Interestingly, the end of anoxic–euxinic conditions (with abatement of V/Al enrichment and elevated DOP-T 0.5–1) and the termination of TOC >5% occur synchronously with the return to steady light $^{13}$C$_{org}$ values in the Aubach section (Fig. 5 and 7). This also appears to be the case in the southern Paris Basin (Sancere-Couy Core) where [V/Al] and TOC content decrease just after the return to steady light $^{13}$C values (Fig. 7). The %TOC stabilizes synchronously with $^{13}$C$_{org}$ in Yorkshire as well, suggesting a coupling between the proxies. However, a coherence between redox sensitive proxies (DOP-T) and $^{13}$C is not present (Fig. 7). V/Al ratios and DOP-T are not documented for the nearby Dotternhausen section, so a direct comparison is not possible. However, redox sensitive biomarkers (Pr/Ph and Aryl isoprenoid ratios) decrease where $^{13}$C$_{org}$ return to steady light values in lower *bifrons* Zone (Fig. 3 in Schwark and Frimmel 2004), suggesting the same mirroring of $^{13}$C and redox sensitive proxies here, whereas TOC of more than 5% occur throughout the *bifrons* Zone (Röhl et al. 2001).

In contrast, the Iberian Range, Spain (Reolid 2014), Morocco, N-Africa (Krencker et al. 2020), and Peniche section, Lusitanian Basin (Fantasia et al. 2019, Hermoso et al. 2009b) were more oxygenated, and long-term anoxia never developed during the T-OAE. The transect between anoxic and oxic locations is well illustrated across the southern epi-continental margin (S Germanic and Swiss locations) displaying the transition from the anoxic Swabo-Franconian Basin (this study, Röhl et al. 2001, Dickson et al. 2017) to the dysoxic–anoxic northern shelf (Fig. 7) (Montero-Serrano et al. 2015) to the oxic Lombardian Basin on the continental margin (Fantasia et al. 2018). The T-OAE as an environmental perturbation is thus not necessarily synchronous in different basins, yet the nCIE of the T-OAE still serves well as a detailed stratigraphic correlation (Ruebsam and Al-Husseini 2020). The non-identical redox conditions during and after the T-OAE in European sections illustrate well that the drivers of anoxia was, at least partly, regionally influenced. Redox sensitive proxies (V/Al and DOP-T) indicate that anoxic-euxinic conditions prevailed in the inner-epicontinental Sea until the upper *falciferum* Zone, concurrent with the return to stable light $^{13}$C values.

### 5.4. Decoupling between $^{13}$C and organic matter burial

Deposition of black shale (TOC >5%) in the Aubach section preceded the stratigraphic extent of the positive limb of $^{13}$C excursion (Fig. 2) and by definition the end of the T-OAE (cf. Jenkyns 2010; Fig. 6). This suggests that local scale black shale formation did not solely drive $^{13}$C$_{org}$. If $^{13}$C$_{org}$ were exclusively governed by local processes, primary productivity and organic carbon burial (e. g., Scholle and Arthur 1980), the TOC content and $^{13}$C$_{org}$ would be expected to stratigraphically mirror each other, if we assume that type of organic matter and growth rate were constant. An increase in these local processes will thereby lead to an increase in $^{13}$C of the local dissolved inorganic carbon (DIC) and finally in the $^{13}$C$_{org}$ and $^{13}$C$_{carb}$ (Scholle and Arthur 1980). The fact that black shale deposition (%TOC >5%) precedes the rising $^{13}$C$_{org}$ values document a partial decoupling of the pCIE and organic carbon burial in the Aubach section (Fig. 2). This suggests that the local pCIE were only partly influenced by regional seawater circumstances, and thus other factors must have impacted the local $^{13}$C$_{DIC}$. Changes in atmospheric CO$_2$ ($p$CO$_2$) levels would affect the local $^{13}$C$_{DIC}$ as atmosphere and the upper mixed layer are in equilibrium (Revelle and Suess 1957). A global decline in $p$CO$_2$ is expected after the nCIE, as a result of the widespread organic carbon burial (Ruebsam et al. 2018) and intensified chemical weathering indicated by elevated Osmium isotope values (Them et al. 2017b). The local pCIE may be a result of 1) a global decline in $p$CO$_2$ after the nCIE, and 2) local changes in bioproductivity and organic carbon burial. However, this call for further investigation.

We observe a stratigraphic relationship between termination of black shale formation (TOC >5%) and the end of the Toarcian pCIE in the Aubach,
Sancerre Couy and Yorkshire locations (Fig. 7). Previous studies have also shown that black shale formation continued after the maximum of the Toarcian pCIE in basins worldwide (Them et al. 2017a, Röhrl et al. 2001, Hermoso et al. 2013, Suan et al. 2011) by comparison to what is expected from theoretical models (Ullmann et al. 2020) as the maximum of the Toarcian pCIE is commonly used to define the termination of the T-OAE. This mismatch between proxies, warning us that prolonged anoxia locally occur long after the peak in the carbon cycle perturbation call for a revision of the definition of the T-OAE. Furthermore, the T-OAE may serve as an analogue for the anthropogenic carbon perturbation. This suggests that the environment recovery will be substantially delayed in many places, after we have the release of $^{12}C$ to the atmosphere under control.

6. Conclusions

This study examines carbon isotope, redox element and Rock Eval records across the Early Toarcian interval in the Aubach section, SW Germany, which are compared to published data from other European sections. We find that the negative carbon isotope event (nCIE) record in the Aubach section is reduced from $\delta^{13}C_{org}$ 6.3 ‰ to 3.5–3.3 ‰ after applying a HIC-correction which is comparable to the carbonate $\delta^{13}C$ record (4.5 ‰). This shows that it may be worth accounting for variable OM sources to understand the magnitude of the nCIE better.

The onset of long-term anoxia indicated by V/Al enrichment and DOP-T of 0.5–1 and TOC of more than 5 % initiate in the upper semicelatum Zone. This occurs concurrently with the first negative shift in $\delta^{13}C_{org}$. The redox element record indicates that anoxic–euxinic conditions prevailed until the late falciferum Zone, concurrent with the return to stable $\delta^{13}C_{org}$ values at $\sim$28 ‰. We define the termination of the Toarcian positive CIE based on the return to steady light $\delta^{13}C_{org}$ values in the upper falciferum Zone, which we believe marks the return to normal marine conditions in the Aubach section. This coherence between return to stable $\delta^{13}C_{org}$ values and the end of V/Al enrichment and TOC >5% also occurs in the Sancerre-Couy (southern Paris Basin). In Yorkshire (Cleveland Basin), there is only synchronism between the $\delta^{13}C_{org}$ and %TOC trends but not in redox conditions. The return to normal marine conditions seems to be synchronous in the some European basins but not all, highlighting the need for further investigation.

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References


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Figure 1S: Lithology and carbon and oxygen isotopic trends in the uppermost Pliensbachian and lower Toarcian sediment of the Aubach section. δ13C carb values from the Unterer Stein limestone bed (empty circles) have been excluded from the interpretation because of diagenetic overprint (see main text). Note, that the negative CIE of 6.3 ‰ develops over four – five steps in δ13C org, marked by the red arrows, and of only 4.5 ‰ in δ13C carb.

Table 1S: Labels, δ13C org and δ13C carb values and stratigraphic height (m) of the investigated samples.

Table 2S: Organic matter pyrolysis data with stratigraphic height (m). HI and OI are presented in mg HC/g TOC and mg CO2/g, respectively.

Table 3S: Element data, total S, total Fe, TOC wt%, %CaCO3. DOP-T (degree of pyritization based on total Fe) have been excluded from the results were TOC <0.5 wt% and in the limestone bed (2.90–3.08 m). TOC data in italic are measured using organic matter pyrolysis and the others with the Eltra CS 500 Analyzer.

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