Organic matter content and type variation in the sequence stratigraphic context of the Upper Devonian New Albany Shale, Illinois Basin

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Organic matter quantity and type are important parameters in conventional source rock evaluation and unconventional shale oil/gas reservoir characterization. Understanding the stratigraphic distribution of organic matter content and type in black shale successions is critical for identifying potentially productive intervals, because organic matter can adsorb large amounts of oil and gas. Detailed examination of total organic carbon content, organic petrographic composition, and high-resolution (8 cm spacing) geochemical proxies in a sequence stratigraphic framework were conducted on an early mature (0.55% R0) New Albany Shale core of the Illinois Basin to study the influence of relative sea-level fluctuations on paleoproductivity, clastic supply, bottom-water redox conditions, and their combined control on total organic carbon distribution pattern and organic maceral variation. Marine organic matter including amorphous organic matter and alginite (mainly derived from Tasmanites cysts) is the dominant organic matter in the New Albany Shale. Terrestrial organic matter accounts for <10% of total organic matter. Within a sequence stratigraphic context, total organic carbon content increases in transgressive systems tracts, reaches a maximum before the maximum flooding surface, and shows relatively low values in highstand systems tracts. Comparatively, low total organic carbon contents at maximum flooding surfaces reflect a combination of low sedimentation rates, elevated bottom-water oxygenation, and high biogenic silica dilution. Stratigraphically, amorphous organic matter content increases in transgressive systems tracts, reaches a maximum near the maximum flooding surface, and decreases in highstand systems tracts. Enrichment of broken Tasmanites cysts and their detrital infills are an indicator of high-energy environments and could possibly indicate lowstand systems tracts. The stratigraphic distribution of organic matter content and type may result in cyclic stratigraphic variations of hydrocarbon generation potential and oil saturation, and influence the development of secondary organic pores when the New Albany Shale is within the oil and gas windows.

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1. Introduction

Organic matter (OM) richness, type, and thermal maturity are key parameters used in the evaluation of conventional and unconventional petroleum systems (Tissot and Welte, 1984; Peters and Cassa, 1994; Jarvie, 2012a, 2012b; Hackley and Cardott, 2016). The total organic carbon (TOC) content in source rocks determines the quantity of generated petroleum and the type of kerogen controls the type of generated hydrocarbons (Tissot and Welte, 1984). OM-hosted pores play a significant role in oil and gas storage and migration in unconventional petroleum systems because of their large specific surface area and organic affinity (Passey et al., 2010; Schieber, 2010; Loucks et al., 2012; Liu et al., 2017). However, most organic pores are <1000 nm (Wang et al., 2009; Schieber, 2010; Loucks et al., 2012) and contribute differently in shale oil vs shale gas systems because of the difference in fluid properties of oil vs gas (Jarvie, 2012a, 2012b; Bohacs et al., 2013). OM content in source rocks is not uniformly distributed stratigraphically or spatially (Creaney and Passey, 1993; Slatt and Rodriguez, 2012; Dong et al., 2017), and OM is not homogeneous in composition, but is composed of a mixture of organic macerals (e.g., Robl et al., 1992; Stasiuk and Fowler, 2004; Mastalerz et al., 2012, 2013; Hackley and Cardott, 2016; Liu et al., 2017, 2019; Abarghani et al., 2018; Harris et al., 2018; Ponsaing et al., 2018). Therefore, documenting TOC distribution and organic maceral variation in a sequence stratigraphic context is of great significance for the evaluation of conventional source rocks and unconventional shale oil/gas resources.

Primary controls on OM accumulation in marine sediments include paleoproductivity, preservation conditions (bottom-water redox conditions), clastic supply, as well as sedimentation rate (Sageman et al., 2003; Rimmer et al., 2004; Bohacs et al., 2005, 2013). Sedimentation rate can affect preservation conditions of OM and tie accumulation of organic-rich mudstones to a sequence stratigraphic framework.
OM in petroleum source rocks can be classified into four types of kerogens on the basis of H/C versus O/C ratios (Tissot et al., 1974; Peters and Cassa, 1994). However, examining them with a reflected-light microscope reveals added complexity through the recognition of a variety of organic macerals which have different sources and formation processes (Peters and Cassa, 1994; Taylor et al., 1998). Stratigraphic variation of OM type has been previously studied in black shale successions (Pasley et al., 1991; Gregory and Hart, 1992; Robison et al., 1996; Bialkowski et al., 2000; Schieber, 2001; Harris et al., 2018). For example, the stratigraphic variability of the distribution of organic macerals is illustrated by the observation that in the New Albany Shale and Chattanooga Shale, alginite derived from Tasmanites cysts is concentrated at maximum flooding surfaces (MFSs) (Schieber, 2001; Lazar, 2007).

The Upper Devonian New Albany Shale of the Illinois Basin is an organic-rich formation with TOC contents ranging from ~1% to ~20% (Chou et al., 1991), and its thermal maturity ranges from vitrinite reflectance (R$_t$) 0.5% near the basin margin to 1.5% at Hicks Dome, Illinois (Spatroš et al., 2010; Mastalerz et al., 2013). Although geochemically OM in the New Albany Shale is classified as type II kerogen on the basis of Rock-Eval pyrolysis (Chou et al., 1991; Akar et al., 2015), it is nonetheless composed of varying organic macerals including amorphous organic matter (AOM), alginite, liptodetrinite, vitrinite, inertinite, and solid bitumen (SB) when examined with a reflected-light microscope (Mastalerz et al., 2012, 2013; Wei et al., 2016; Liu et al., 2017, 2019). Commercial oil and gas production from the New Albany Shale has been reported where it has oil-window maturity (Hamilton-Smith et al., 1991; Nuttall et al., 2015).

Lazar (2007) studied the controls of paleo-redox conditions, paleopedoactivity, and clastic dilution on organic matter accumulation in each unit of the New Albany Shale. Because Lazar (2007) only had TOC contents available, he operated under the assumption of uniform marine OM, although he noted that petrographic examination of these rocks shows considerable variability in the relative abundances of organic macerals (also pointed out by Mastalerz et al. (2012, 2013) and Liu et al. (2017, 2019)) and that quantitative information on their abundances could affect the conclusions reached in his study. This study builds on the earlier work of Lazar (2007) and adds to the data set with more detailed information on the stratigraphic variation of TOC content, and augments it with newly acquired data on the types and proportions of organic macerals. The goal of this investigation is to understand the controls on TOC content from a perspective of organic maceral accumulation in the sequence stratigraphic context of the Upper Devonian New Albany Shale of the Illinois Basin. Specific objectives are to: (1) document TOC content distribution in greater detail; (2) reevaluate the influence of relative sea-level fluctuations on paleo-productivity, clastic supply, bottom-water redox conditions, and their combined control on OM accumulation; and (3) examine the stratigraphic variability of organic macerals (AOM, alginite, vitrinite, and inertinite) within the New Albany Shale. Results of this study should allow for more sophisticated assessment of OM accumulation in time-equivalent Upper Devonian black shales deposited in adjacent epicontinental basins such as the Ohio Shale of the central Appalachian Basin, the Antrim Shale of the Michigan Basin, the Chattanooga Shale of the southern Appalachian Basin, and the Bakken Shale of the Williston Basin (Schieber, 1998; Schieber and Lazar, 2004).

2. Geological background

The Upper Devonian New Albany Shale of the Illinois Basin was deposited in an epicontinental sea during the late Devonian (Lineback, 1964, 1968; Beier and Hayes, 1989). It unconformably overlies the Middle Devonian North Vernon Limestone and underlies the Falling Run Bed, a phosphatic lag deposit (Campbell, 1946). Where the Falling Run Bed is absent, it is unconformably overlain by the early Mississippian Underwood, Henryville, and Jacobs Chapel Beds, or directly overlain by the Mississippian Rockford Limestone or the Mississippian New Providence Shale if these beds are absent (Campbell, 1946; Lineback, 1964, 1968). The New Albany Shale is composed of the Blocher, Selmer, Morgan Trail, Camp Run, and Clegg Creek Member in ascending order (Lineback, 1964, 1968). Lithologically, the New Albany Shale is composed of black laminated to banded shales and greenish-gray, bioturbated shales (Lineback, 1964, 1968; Lazar, 2007).

A complete New Albany Shale core of 39.43 m thickness that was described by Lazar (2007) for a sequence stratigraphic study, has been further examined and analyzed in order to study the distribution of TOC and organic macerals within a sequence stratigraphic context. The core is from Daviess County, Indiana (Fig. 1) with a coordinate of X = 494,611, Y = 4,276,705 (North American Datum 83; Lazar, 2007). The core (1–3 Kavanaugh) was drilled by Deka Exploration Inc. in 1994. OM in this core indicates a thermal maturity of 0.55% R$_t$. Being marginally mature, OM in this core has not undergone significant thermal transformation, although early diagenesis could to some degree alter OM through chemical and microbial degradation (Curtis, 1980; Tissot and Welte, 1984).

Using the conceptual framework of Schieber (1998) and Schieber and Lazar (2004), the studied New Albany Shale core can be divided into four basin-wide depositional sequences based on integration of sedimentological, paleontological, geophysical, and geochemical data (Fig. 2) (Lazar, 2007). Within those sequences, we identified six major lithofacies based on sedimentological (color, lamination, and banding) and ichnological (bioturbation intensity) characteristics: (1) black laminated to banded carbonate-bearing shale, (2) black laminated to banded shale, (3) black to brownish laminated shale with abundant lag deposits, (4) black-gray cyclic shale, (5) black massive to banded shale with interbedded gray bioturbated shale, and (6) black massive to banded shale (Fig. 3).

The black laminated to banded carbonate-bearing shale lithofacies (Figs. 2, 3A) is limited to unit 1TST (TST of sequence 1) with varying content of carbonate minerals (calcite and dolomite) and a bioturbation index (BI) of 0–1. The laminae are mainly composed of silt-sized quartz and recycled dolomite (Fig. 4) and formed through intermittent reworking of bottom sediments by storms and waves which winnowed the seabed and removed the fine-grained fraction (e.g., clay minerals and clay-sized particles). The black laminated to banded shale lithofacies (Figs. 2, 3B) is most common in units 1HST (BI = 0–2) and 3TST (BI = 0–1). The black to brownish laminated shale with abundant lag deposits (Figs. 2, 3C) is the major lithofacies in units 2LST (lowstand systems tract, BI = 0–1) and 4LST (BI = 0–2). Laminae and lag deposits, which can be cm-thick, formed via intensive reworking of bottom sediments by strong storms and waves (Schieber, 1998). The brownish color is due to an abundance of pyrite and Tasmanites microfossils, unicellular green algae that belong to the class Prasinophyceae (Tappan, 1980). The black-gray cyclic shale lithofacies (Figs. 2, 3D) occurs in the upper half of unit 2TST (BI = 2–4) and the entire 2HST (BI = 3–4). The formation of these black-gray cyclic shales has been interpreted to reflect alternation of anoxic and oxic environments resulting from fluctuations of relative sea level (Calvert et al., 1996). As such the black-gray cycles most likely represent distal parasequences (Schieber,
that reflect a high-frequency interplay of sediment buildup (shallowing) and intermittent small increases of sea level (deepening) as detailed by Spencer (2013). The black massive to banded shale with interbedded gray bioturbated shale lithofacies (Figs. 2, 3E–F) is restricted to unit 3HST (BI = 0–2). Gray bioturbated beds are organic-poor beds. The black massive to banded shale lithofacies (Figs. 2, 3G) is common in units 4TST (BI = 0–1) and 4HST (BI = 0–1). Generally, shales deposited in HSTs are more intensively bioturbated than those in TSTs within a given depositional sequence.

3. Analytical methods

Portable x-ray fluorescence (pXRF) instrumentation can provide rapid, non-destructive, and quantitative geochemical analysis of mudstone samples (Rowe et al., 2012). Major and trace elements in 482 samples were measured on the core at a spacing of approximately 8 cm with a pXRF analyzer (Thermo Niton XL3t GOLDD+) at the Indiana Geological and Water Survey. The measurements were carried out in situ on the cleaned surfaces of core slabs and conducted on a round area with a diameter of 11 mm. The pXRF analyzer was run in “Test All Geo” mode for 75 s. Three USGS certified reference materials (one carbonatite and two shale: COQ-1, SDO-1, and SBC-1) were analyzed at the beginning and end of each analytical session to examine the accuracy and precision of the analyses (McLaughlin et al., 2016). In this study, the analysis errors for major elements silicon (Si), aluminum (Al), titanium (Ti), phosphorus (P) are 0.78%, 2.88%, 3.27%, and 8.49%, respectively. Errors for trace elements molybdenum (Mo), nickel (Ni), vanadium (V), U, thorium (Th), and cobalt (Co) are 13.24%, 21.24%, 23.31%, 34.71%, 38.43%, and 43.17%, respectively, which are much higher than those of major elements.

Because portable XRF spectrometers have a lower detection accuracy than high-energy XRF spectrometers, contents of elements Si, Al, Ti, P, U, Mo, V, Ni, Th, and Co obtained with pXRF analyzer were calibrated with 44 previously analyzed New Albany Shale samples from Lazar (2007) (Supplementary Fig. 1). Major elements Si, Al, and Ti and trace elements U, V, and Mo exhibit high regression coefficients of determination ($R^2 > 0.7$), whereas coefficients of determination for P and trace elements Ni, Th, and Co are relatively low (Supplementary Fig. 1). Although accuracy is low, trace element ratio profiles can still indicate stratigraphic trends of paleo-redox conditions. Because U is commonly associated with OM in marine shales (Swanson, 1960; Leventhal, 1981; Fertl and Chilingar, 1988; Bohacs and Schwalbach, 1994; Lüning and Kolonic, 2003), the TOC content of the 482 samples was calculated based on the relationship between TOC and U content (Eq. (1)) established with 43 previously analyzed New Albany Shale samples (Lazar, 2007). This relationship is very similar to the results of Fertl and Chilingar (1988) (Fig. 5).

\[
\text{TOC (wt\%)} = 4.18 \ln(U \text{ ppm}) - 6.26
\]  

(1)

The TOC content of seventeen samples was measured using a LECO elemental analyzer (SC832DR). The seventeen samples were selected to
supplement the forty-four measurements of Lazar (2007) from the same core and to cover the unmeasured intervals. During the TOC measurements, 0.25 g powdered samples were acidified to remove carbonate minerals and then combusted in the LECO elemental analyzer. The amounts of generated CO2 were converted to organic C content in samples. A duplicate sample was measured to ensure the accuracy of measurements. Of the 61 samples (17 in this study and 44 from Lazar (2007)) with TOC contents, thirty-three samples were taken from ten systems tracts in four depositional sequences for organic petrographic study. Vitrinite reflectance of shale samples was measured on polished whole-rock pellets using a Zeiss RS-III microscope with 50 measurements.

Semi-quantitative organic petrographic compositions (AOM, alginite, liptodetrinite, vitrinite inertinite, and SB) of shale samples were determined via point-counting (500 points counted only on OM). The maceral groups vitrinite and inertinite were not classified into macerals because of their scarcity. The liptinite group includes AOM, alginite, and liptodetrinite because they are dominant oil-prone macerals in the New Albany Shale (Liu et al., 2017, 2019) and marine black shales in general (Mastalerz et al., 2018). Organic macerals were identified and photographed using a reflected-light microscope (Leica DFC310 FX) in reflected white light (for occurrence, color, and morphology) and blue light (for fluorescence properties) with oil immersion. A field emission scanning electron microscope (FEI Quanta 400 FEG) was used to examine shale composition and AOM using a correlative microscopy approach (reflected-light and SEM microscopy). The methods of correlative microscopy have been presented in Liu et al. (2017).

4. Results

4.1. Stratigraphic variation of TOC content

The TOC content of the New Albany Shale measured in this study ranges from 0.18% to 16.29%, with an average content of 6.53% (Fig. 6). Samples with the same lithofacies and next to each other show high variations of TOC content (e.g., at 630.75 m; Fig. 6), suggesting high heterogeneity of organic richness in black shales. The Clegg Creek Member (sequence 4) exhibits the highest TOC content (average TOC content 8.22%) in the New Albany Shale. Within a given depositional sequence, TOC content increases in the TST, reaches a maximum before getting to the MFS, and maintains relatively low values during the HST (Fig. 6). Stratigraphically, the TOC profile displays a very similar trend to the gamma-ray profile except in the Blocher Member (Fig. 6).

Calculated TOC content with Eq. (1) ranges from 0 to 12.53%, with an average content of 6.45%, which is very close to the measured average TOC content of 6.53%. The calculated TOC profile is very similar to the measured TOC profile and closely matches the gamma-ray profile (Fig. 6), suggesting that U is the main contributor to gamma radiation in the New Albany Shale. As with the trend of measured TOC in the sequence stratigraphic context, calculated TOC content increases in the TST, reaches a maximum before getting to the MFS, and maintains relatively low values during the HST with the exception of the Blocher Member (Fig. 6). Similar to the gamma-ray profile, the calculated TOC content is relatively constant in the Blocher Member (Fig. 6).

4.2. Paleoproductivity

Even though the New Albany Shale is characterized by high clay content (Mastalerz et al., 2013), there is a significant contribution of biogenic silica (Schieber, 1996; Schieber et al., 2000). Biogenic silica in the New Albany Shale is primarily sourced from radiolaria, planktonic organisms with an opaline test (Schieber, 1996; Schieber et al., 2000). High biogenic silica content (Sibio) typically indicates high paleoproductivity (Ross and Bustin, 2009), and for this study Sibio content was calculated using the following formula (Eq. (2); Ross and Bustin, 2009):

\[ \text{Sibio} = \frac{\text{Sisample}}{\text{Al}^{\text{background}}} \times \text{Al}^{\text{sample}} \]  \hfill (2)

A Si:Al ratio of 2.55 is used as the background value, which is the lowest Si:Al ratio (at 623.90 m) in the New Albany Shale. This interval is gray bioturbated shale and is mainly composed of detrital components, such as clay minerals (mostly illite), silt-sized quartz, dolomite, albite, potassium (K)-feldspar and minor pyrite, muscovite, and biotite (Fig. 7). Biogenic silica is very rare to absent in this interval. A background Si:Al ratio of 3.11 (Wedepohl, 1971) has been used in many previous studies (Ross and Bustin, 2009; Dong et al., 2017; Zhao et al., 2017; Harris et al., 2018), but at that ratio, a large number of samples would
have negative $S_{\text{bio}}$ content. Because that is implausible, a background Si:Al ratio of 3.11 is considered too high for the New Albany Shale. Likewise, Harris et al. (2018) reported that a background Si:Al ratio of 3.11 would result in negative values for calculated biogenic silica in the Upper Devonian Duvernay Formation (Western Canada Sedimentary Basin). Given that the background Si:Al ratio of 3.11 is based on an average composition that includes data from a large number of shales across the globe and without differentiation of biogenic vs detrital sources of SiO$_2$ (Wedepohl, 1971), universal application of a Si:Al background ratio of 3.11 should be discouraged. As shown in this study, a plausible minimum Si:Al ratio that is intrinsic to a given stratigraphic interval can be derived and should help a more realistic assessment of biogenic silica inputs.

In the studied core, total Si content ranges from 4.07 to 38.90% (average 25.52%), out of which 0 to 35.14% (average 6.01%) is calculated as of biogenic origin, suggesting that 24% of total Si is biogenic. Sequence 4 has the highest biogenic Si content (average $S_{\text{bio}}$ 9.51%) in the New Albany Shale (Fig. 6). The highest $S_{\text{bio}}$ (35.14%) content occurs near the MFS of sequence 4, accounting for 90% of total Si. Within a given depositional sequence, $S_{\text{bio}}$ content increases in the TST, reaches a maximum at the MFS, and decreases in the HST (Fig. 6).

Phosphorus is an essential nutrient for marine phytoplankton and all organisms in general (Schoepfer et al., 2015), and its content in marine sediments can be related to surface ocean productivity (Schenau et al., 2005). P content in the New Albany Shale core is relatively constant (close to 0.04%). P content in the Clegg Creek Member (average content
0.06%), however, is higher than in the rest of the core (average content 0.03%). The high P content at the bottom of unit 4LST results from the enrichment of reworked conodont fragments. The P profile shows a similar distribution pattern to the Sibio profile, but the P content maximum at the MFS is not as clear as in the case of Sibio (Fig. 6).

4.3. Clastic supply

Al is abundant in aluminosilicates such as clay minerals and feldspars, and Ti is contained in heavy minerals such as rutile (Brumsack, 2006). Al and Ti are chemically conservative during diagenesis (Brumsack, 2006) and have been used to serve as proxies for clastic supply (Sageman et al., 2003; Rimmer et al., 2004; Dong et al., 2017). Al and Ti contents range from 0 to 10.20% (average 7.65%) and 0 to 0.56% (average 0.41%), respectively. Stratigraphically, Al and Ti contents display the same trend, and show an opposite distribution pattern compared to the Sibio profile, with Al and Ti content at a minimum (but Sibio maximum) at the MFS (Fig. 6). Within a given depositional sequence, Al and Ti contents decrease in the TST, reach a minimum at the MFS, and increase in the HST, with the exception of the Blocher Member (Fig. 6).

4.4. Bottom-water redox conditions

There is a variety of trace element-based geochemical proxies that have been suggested as indicators of relative oxygenation of bottom waters (e.g., Adams and Weaver, 1958; Hatch and Leventhal, 1992; Jones and Manning, 1994; Rimmer et al., 2004). These proxies, however, may provide contradictory results, and should not be applied rigidly (Rimmer, 2004; Rimmer et al., 2004; Lazar, 2007; Ocubalidet et al., 2018). Instead, they are best used as relative indicators of levels and trends of bottom-water oxygenation (Rimmer, 2004; Rimmer et al., 2004).

\[ \mathrm{U(UO_2(CO_3)_3}^{4-}) \] is soluble in seawater, but it will be converted to the less soluble \( \mathrm{U(O_2, U_2O_7, or U_3O_8)} \) in reducing environments (Langmuir, 1978; Anderson et al., 1989; Klinkhammer and Palmer, 1991; Algeo and Tribovillard, 2009; Cumberland et al., 2016). \( \mathrm{U(IV, +4 oxidation state)]} \) occurs in marine OM as either U dioxide (\( \mathrm{UO_2)} \) in an absorbed state or as U-OM compounds (Swanson, 1960; Cumberland et al., 2016). Th is typically associated with heavy minerals and is relatively immobile during diagenesis (Adams and Weaver, 1958). Therefore, high U/Th ratios of marine sediments are generally thought to indicate oxygen-deficient conditions (Adams and Weaver, 1958; Jones and Manning, 1994).

In this study, the U/Th ratio profile closely matches the calculated TOC profile (Fig. 6). Stratigraphically, the U/Th ratio increases in the TST, reaches a maximum before the MFS, and maintains relatively low values during the HST. Most of the U/Th ratio values are higher than 0.75, which may indicate dysoxic to anoxic environments (Tyson and Pearson, 1991; Jones and Manning, 1994) (Fig. 6).

Enrichment factors (EF) have been used to evaluate the enrichment of trace metals in organic-rich shales relative to average shales (Brumsack, 2006; Tribovillard et al., 2006, 2012; Algeo and Tribovillard, 2009). EF are calculated as (Eq. (3)):

\[
X_{\text{EF}} = \left( \frac{X_{\text{sample}}}{X_{\text{average shale}}} \right)_{\text{Al}}
\]

where X and Al represent the concentrations of elements X and Al, respectively. Samples were normalized using the post-Archean average shale (PAAS) composition (Taylor and McLennan, 1985). 

\( U_{\text{EF}} \) ranges from 0 to 50.08, with an average value of 11.26, suggesting a common anoxic environment (Algeo and Tribovillard, 2009; Tribovillard et al., 2012) during the deposition of the New Albany Shale. \( U_{\text{EF}} \) with values of 0 probably results from the relatively high detection limit of pXRF analyzer (samples with low U content were not detected). Stratigraphically, the \( U_{\text{EF}} \) profile displays the same distribution pattern as the U/Th profile (Fig. 6).

V and Ni are preferentially concentrated in tetrapyrrole compounds under anoxic conditions (Lewan and Maynard, 1982), and \( V/(V + Ni) \) ratios from whole-rock analyses have been used to indicate
**Fig. 6.** Stratigraphic distribution of lithofacies and profiles of gamma-ray log, total organic carbon (TOC) content, and geochemical proxies of paleoproductivity, clastic supply, and bottom-water redox conditions of the New Albany Shale in the studied core. Proposed threshold values for the degree of bottom-water oxygenation are from Tyson and Pearson (1991); Hatch and Leventhal (1992); Jones and Manning (1994); Algeo and Tribovillard, 2009. LST = lowstand systems tract; TST = transgressive systems tract; HST = highstand systems tract; MFS = maximum flooding surface; O = oxic; D = dysoxic; A = anoxic. *Sequence stratigraphic context of the New Albany Shale is modified from Lazar (2007). †Measured TOC content of 17 samples in this study and 44 samples from Lazar (2007). ‡TOC content calculated from U content. pXRF data can be found in Supplementary data sets.
environmental conditions at the time of deposition (Hatch and Leventhal, 1992). For the studied interval, the V/(V + Ni) ratio does not show significant variations stratigraphically, and is generally higher than 0.46, suggesting dysoxic to anoxic conditions (Tyson and Pearson, 1991; Hatch and Leventhal, 1992) (Fig. 6).

Co is strongly determined by the abundance of clastic materials (Tribovillard et al., 2006), and the Ni/Co ratio of mudstones has been used to indicate oxygenation conditions (Jones and Manning, 1994). Similar to the V/(V + Ni) ratio, the Ni/Co ratio does not show significant variations stratigraphically (Fig. 6). Across the studied core, the Ni/Co ratio is typically lower than 7.00, which would suggest dysoxic to oxic conditions (Tyson and Pearson, 1991; Jones and Manning, 1994) (Fig. 6).

Mo tends to be enriched in marine sediments under anoxic conditions (Calvert and Pedersen, 1993; Algeo and Lyons, 2006; Tribovillard et al., 2006). Mo content is generally normalized to Al content to compensate for the dilution of biogenic minerals (common diluents are calcium carbonate and biogenic silica; Tribovillard et al., 2006), and this normalized Mo content has in the past been proposed as an indicator of paleo-redox conditions of marine black shales (Rimmer et al., 2004). In this study, the Mo/Al ratio and the U/Th ratio display similar distribution patterns stratigraphically except in sequence 4, where the Mo/Al ratio reaches a maximum at the MFS (Fig. 6). Covariation of U and Mo has been reported in modern and paleo-marine systems (Algeo and Tribovillard, 2009; Tribovillard et al., 2012). No threshold values have been proposed to define paleo-redox conditions based on the Mo/Al ratio. Mo$_{eq}$ ranges from 0 to 283.17, with an average value of 69.85. Mo/Al ratio and Mo$_{eq}$ show exactly the same distribution pattern (Fig. 6) because Mo$_{eq}$ is normalized with Al content.

4.5. Organic maceral variation

Organic macerals are microscopically recognizable individual constituents of OM (Stach et al., 1982; Taylor et al., 1998). Organic macerals observed in this study include AOM, alginate, liptodetrinite, vitrinite, inertinite, and SB (Fig. 8). In marine organic-rich shales, AOM and alginate are of marine origin (Kus et al., 2017), whereas vitrinite and inertinite are derived from terrestrial woody materials (Stach et al., 1982; Taylor et al., 1998). AOM, derived from microbially degraded phytoplankton, zooplankton, and bacterial biomass, refers to structureless OM in coals and organic-rich shales (Teichmüller, 1989; Taylor et al., 1998; Pacton et al., 2011; Kus et al., 2017). AOM commonly has small mineral inclusions, such as clay minerals and clay-sized particles (e.g., quartz, feldspar, and pyrite), suggesting its degradation origin (Fig. 9). Alginate in the New Albany Shale is mainly in the form of Tasmanites cysts (Fig. 8). Liptodetrinite occurs as small fluorescenting OM particles (Fig. 8E), and is likely of algal origin. SB is secondary OM generated from primary oil-prone macerals (Jacob, 1989; Mastalerz et al., 2018), such as AOM and alginate (Liu et al., 2019). A detailed petrographic study of OM in the New Albany Shale was presented in Liu et al. (2017, 2019).

OM in the New Albany Shale is dominated by either AOM or alginate (Table 1). The terrestrial OM (vitrinite + inertinite) content is <10% (Table 1). Vitrinite content is typically lower than inertinite content (Table 1) because inertinite is chemically inert and vitrinite is possibly partially oxidized during transport to the place of deposition (Wei et al., 2016; Liu et al., 2019). Because this core is at the early mature stage (0.55% Ro), relatively low but varying contents of SB were observed (Table 1). This SB likely results from thermal degradation of labile OM (Liu et al., 2019).

The organic petrographic composition of the New Albany Shale varies stratigraphically (Fig. 10). AOM dominates except in the unit 1HST, the Selmer Member (sequence 2), the unit 3TST, and the unit 4LST (Table 1; Fig. 10). Within sequences 1, 2, and 4, AOM content increases in the TST, reaches a maximum near the MFS, and decreases in the HST (Fig. 10). There is a slight decrease of AOM content at the MFS (Fig. 10). Because OM is dominated by alginate in the Selmer Member, the stratigraphic distribution pattern of AOM content is not as obvious as in sequence 1 and 4 (Fig. 10). Compared to other sequences, sequence 3 does not show an AOM content maximum near the MFS, but within the middle of the HST (Fig. 10). In contrast to AOM, alginate displays an opposite pattern stratigraphically (Fig. 10). Alginate can be enriched in certain intervals of the studied core. For example, alginate accounts for 83.1% of total OM in sample NAS–15, whereas samples above and below it are dominated by AOM (Table 1; Figs. 10, 11). Terrestrial OM (vitrinite + inertinite) content is relatively constant (~2.5% of total OM) throughout most of the studied core except at the top where terrestrial OM accounts for 6–10% of total OM (Table 1; Figs. 10, 12). Ripley et al. (1990) also reported a high abundance of terrestrial OM (6–14%) within the upper part of the Clegg Creek Member. The increase of terrestrial OM at the top of the New Albany Shale is accompanied by increasing Al and Ti contents (Figs. 6, 10).
Fig. 8. Photomicrographs of organic macerals in reflected white light and oil immersion (A–C, F–H) and in fluorescence mode (D–E) in the New Albany Shale in the studied core. (A–B) Amorphous organic matter. AOM occurs as organic streaks. (C) Alginite in the form of Tasmanites cysts. The algal cysts are compacted. (D) The same field of panel C in fluorescence mode. Alginite shows greenish-yellow fluorescence at this maturity. (E) Liptodetrinite and alginite in fluorescence mode. (F–G) Vitrinite and inertinite occurring as dispersed particles in the matrix. Inertinite in panel G shows cellular structure (red arrows). (H) Solid bitumen. Note the embayments against mineral particles. AOM = amorphous organic matter; AL = alginite; LPD = liptodetrinite; V = vitrinite; I = inertinite; SB = solid bitumen. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
5. Discussion

5.1. Paleoproductivity

OM preserved in marine sediments comes mainly from primary production of phytoplankton in the ocean (e.g., Müller and Suess, 1979; Schoepfer et al., 2015), together with a component of terrestrial OM supplied from adjacent land (e.g., Goñi et al., 1997; Hedges et al., 1997; Opsahl and Benner, 1997; Ponsaing et al., 2018). High paleoproductivity is favorable for enhanced OM burial (Pedersen and Calvert, 1990; Schoepfer et al., 2015). Biogenic silica content is considered a useful proxy for paleoproductivity (Ross and Bustin, 2009). Zhao et al. (2017) reported a positive correlation between TOC and excess (biogenic) silica content in the Wufeng and Longmaxi Formations of the Sichuan Basin (China). In this study, the Clegg Creek Member has the highest average Si_{bio} content (9.51%), P content (0.06%), and TOC content (8.22%) in the New Albany Shale (Fig. 6). High paleoproductivity could be one reason for high TOC content in the Clegg Creek Member, an assumption that is supported by the common presence of early diagenetic silica, derived from the dissolution of radiolaria (Schieber, 1996; Schieber et al., 2000), in Tasmanites cysts and microcrystalline quartz in the matrix in sequence 4 (Fig. 13). However, when abundant biogenic silica contributes to the shale composition, it can lower OM content in shales (Figs. 6, 13). Bohacs et al. (2005) and Passey et al. (2010) also suggested that primary productivity might have a dual impact on OM accumulation in organic-rich sediments, with organic-carbon-rich organisms enhancing OM content, but siliceous and calcareous organisms diluting OM content in sediments.

The stratigraphic distribution pattern of Si_{bio} content (Fig. 6) suggests increasing paleoproductivity in the TST and decreasing paleoproductivity in the HST. Lash and Blood (2011) and Dong et al. (2017) reported elevated diagenetic silica during the deposition of TSTs in the Devonian gas shales of the Appalachian Basin (USA) and the Middle and Upper Devonian Horn River Group of the Horn River Basin (Canada), respectively. The stratigraphic distribution pattern of Si_{bio} content, however, can alternatively be explained by decreasing clastic supply in the TST and increasing clastic supply in the HST (Fig. 6) with paleoproductivity held stable throughout a given sequence. Stratigraphically, the Si_{bio} profile displays an opposite distribution pattern relative to Al and Ti, with the Si_{bio} content maximum coinciding with the Al and Ti content minimum at MFSs (Fig. 6). Therefore, Si_{bio} variation in the New Albany Shale may not so much indicate paleoproductivity, but rather reflect changes of dilution by clastic materials (Fig. 6).

Mass flux of organic carbon per unit area (in g cm\(^{-2}\) kyr\(^{-1}\)) is a better way to represent the surface ocean productivity at the time of deposition of organic-rich sediments if a high-resolution chronostatigraphic framework is available (Algeo et al., 2011, 2013; Schoepfer et al., 2015). However, OM consumption during settling, early diagenesis, and thermal

\[ \text{mass flux} = \text{primary productivity} - \text{OM consumption} \]

\[ \text{OM consumption} = \text{settling} + \text{early diagenesis} + \text{thermal} \]
and Ti products in marine organic-rich sediments. What is worth noting is that Al and Ti paleoproductivity, clastic supply has a dual impact on OM accumulation which in turn can reduce paleoproductivity (Killops and Killops, 2005). This can imply limited supply of nutrients (e.g., N, P, Si, and Fe) to the basin, for OM accumulation due to dilution. However, a too low clastic supply can still be preserved if paleoproductivity is high enough (Pedersen et al., 2018). The stratigraphic distribution pattern of primary productivity of OM exists in the New Albany Shale (Fig. 6) proposed by Lazar (2007), and high Al and Ti contents typically indicate high sedimentation rate and possibly high detrital flux.

Al and Ti contents in the Blocher Member display a different pattern from the other three sequences (Fig. 6), which show decreasing Al and Ti contents in the TST and increasing Al and Ti contents in the HST with the minimum at the MFS. The reason for this difference could be that when sea level started to rise during the deposition of the Blocher Member, clastic materials were not yet effectively delivered to the basin (Brett et al., 2004). The Blocher Member contains calcareous fossil fragments and recycled (clastic) dolomite, as shown by laminae composed of Tentaculites and recycled dolomite (Fig. 14), which are features uncommon in the other three sequences.

5.2. Clastic supply

Devonian clastics in the Illinois Basin were supplied from the east, the Acadian orogeny (Dickinson et al., 1983; Ettensohn, 2008; Schieber, 2016). The sedimentation rate of the studied New Albany Shale interval is extremely low (~ 0.24 cm/ka; Lazar, 2007). Relative sea-level fluctuations played a significant role in transporting detrital materials to the Illinois Basin. Generally speaking, detrital materials are restricted landward during transgressions and transported to the basin during regressions. The stratigraphic distribution patterns of Al and Ti contents (Fig. 6) suggest decreasing clastic supply in the TST and increasing clastic supply in the HST. The TOC profile exhibits an opposite trend compared to the Al and Ti profiles (Fig. 6), which indicates that clastic supply is not favorable for OM accumulation due to dilution. However, a too low clastic supply can imply limited supply of nutrients (e.g., N, P, Si, and Fe) to the basin, which in turn can reduce paleoproductivity (Killops and Killops, 2005). In addition, low clastic supply limits the burial of OM by clastic materials and thus keeps OM within the surficial zone of active microbial degradation for longer time intervals (Bohacs et al., 2005). Therefore, similar to paleoproductivity, clastic supply has a dual impact on OM accumulation in marine organic-rich sediments. What is worth noting is that Al and Ti contents (in %) may not represent the real flux of siliciclastic detritus because of the dilution of biogenic minerals (Tribovillard et al., 2006). However, Al and Ti contents correlate well with the sequence stratigraphic framework of the New Albany Shale (Fig. 6) proposed by Lazar (2007), and high Al and Ti contents typically indicate high sedimentation rate and possibly high detrital flux.

Al and Ti contents in the Blocher Member display a different pattern from the other three sequences (Fig. 6), which show decreasing Al and Ti contents in the TST and increasing Al and Ti contents in the HST with the minimum at the MFS. The reason for this difference could be that when sea level started to rise during the deposition of the Blocher Member, clastic materials were not yet effectively delivered to the basin (Brett et al., 2004). The Blocher Member contains calcareous fossil fragments and recycled (clastic) dolomite, as shown by laminae composed of Tentaculites and recycled dolomite (Fig. 14), which are features uncommon in the other three sequences.

5.3. Bottom-water redox conditions

The preservation potential of OM decreases in the presence of oxygen due to microbial degradation (Wignall, 1991). Nonetheless, OM can still be preserved if paleoproductivity is high enough (Pedersen and Calvert, 1990). The stratigraphic distribution of redox-sensitive trace element ratios suggests that paleo-redox conditions varied during the deposition of the New Albany Shale (Beier and Hayes, 1989; Lazar, 2007; Ocubalidet et al., 2018) (Fig. 6). The U/Th and V/(V + Ni) ratios suggest common dysoxic to anoxic conditions, whereas the Ni/Co ratio suggests common dysoxic to oxic conditions (Fig. 6). Previous studies have proposed that application of trace element ratios to indicate paleo-redox conditions should be in a relative sense, rather than strictly applying proposed thresholds (Rimmer, 2004; Rimmer et al., 2004;
Fig. 10. Stratigraphic distribution of lithofacies, gamma-ray intensity, total organic carbon (TOC) content, and organic petrographic composition of the New Albany Shale in the studied core. LST = lowstand systems tract; TST = transgressive systems tract; HST = highstand systems tract; MFS = maximum flooding surface; OM = organic matter; AOM = amorphous organic matter; V = vitrinite; I = inertinite. aSequence stratigraphic context of the New Albany Shale is modified from Lazar (2007). bMeasured TOC content of 17 samples in this study and 44 samples from the same core (Lazar, 2007). cTOC content calculated from U content.
For example, Ocubalidet et al. (2018) reported that the Ni/Co ratio indicates higher oxygenated conditions than the \( V/(V + Ni) \) ratio in the New Albany Shale. Based on geochemical and petrographic paleo-redox proxies and sedimentologic observations from cores and thin sections, Lazar (2007) concluded that bottom water was mainly dysoxic with intermittent anoxia and oxic conditions during the deposition of the New Albany Shale. Schieber (2009) also reported a common presence of agglutinated benthic foraminifera in the New Albany Shale, suggestive of bottom-water oxygenation.

\[ \frac{V}{V + Ni} \] and Ni/Co ratios are relatively constant stratigraphically (Fig. 6) and thus of limited use for detecting variable paleo-redox conditions of bottom waters in this study. Because \( U_{37} \) and MoEF profiles show the same distribution patterns as \( U/Th \) and Mo/Al profiles, respectively (Fig. 6), they will not be discussed separately. The stratigraphic distribution of \( U/Th \) and Mo/Al ratios indicates that oxygen content in bottom water decreases towards the MFS with the rise of sea level and increases towards the sequence boundary with the drop of sea level (Fig. 6). Interestingly, according to the \( U/Th \) and Mo/Al ratio profiles, the most oxygen-deficient condition (as indicated by metal ratios) does not occur at the MFS where relative sea level is presumed at a maximum (Fig. 6). A potential explanation for this could be that at the MFS strong storms that reworked the substrate and generated lag deposits at MFSs (Fig. 15) were able to supply oxygen to bottom waters via water column mixing. Because redox-sensitive metals would have been mobilized very close to the sediment-water interface under these conditions, they could have diffused out of the sediments into the overlying water (Shaw et al., 1994; Morford and Emerson, 1999). The mismatch between the \( U/Th \) and Mo/Al ratios at the MFS of sequence 4 (Fig. 6) is difficult to explain, probably due to the difference between the degree of enrichment of authigenic U and Mo (Algeo and Tribovillard, 2009; Tribovillard et al., 2012). The lag deposits at the MFS are mainly composed of pyrite, recrystallized radiolaria, Tasmanites cysts filled with authigenic quartz and pyrite, and microcrystalline quartz in the matrix (Figs. 13, 15). The Tasmanites cysts with authigenic infills suggest extremely low sedimentation rate (Schieber, 1996), otherwise the algal cysts would be compacted (Figs. 8C−D, 11). The \( U/Th \) and Mo/Al ratio profiles display a similar trend to the TOC profile (Fig. 6), suggesting that oxygen-deficient conditions are favorable for OM accumulation.

The \( U/Th \) and Mo/Al ratios indicate lower bottom-water oxygenation in the Clegg Creek Member compared to the other members, which could be another reason for high TOC content in the Clegg Creek Member besides high paleoproductivity (Fig. 6). A productivity-anoxia feedback (Ingall et al., 1993) may have contributed to OM accumulation during the deposition of the Clegg Creek Member (Ocubalidet et al., 2018). In the Blocher Member, the \( U/Th \) and Mo/Al ratios are relatively constant and close to the values in the HSTs of sequences 2 and 3 (Fig. 6). A potential explanation could be that relative sea level was still low during the deposition of the Blocher Member and bottom waters were still oxygenated to some extent because storms and waves, as indicated by silty laminae and lenses (Figs. 14, 16), would mix oxygen-deficient bottom waters and oxygenated surface waters. U and Mo do not tend to be enriched under such conditions (Langmuir, 1978; Anderson et al., 1989; Klinkhammer and Palmer, 1991; Calvert and Pedersen, 1993; Tribovillard et al., 2006; Algeo and Tribovillard, 2009); a possible explanation of low gamma-ray intensities observed in the Blocher Member (Fig. 6). Ocubalidet et al. (2018) also suggested that the paleoenvironment was dysoxic to oxic during the deposition
of the Blocher Member based on C-S-Fe relationships and redox-sensitive trace element ratios.

5.4. Organic maceral variation

Organic macerals follow specific evolutionary pathways during thermal maturation (Peters and Cassa, 1994; Liu et al., 2019). Alginite derived from Tasmnites cysts has higher hydrocarbon generation potential (Revill et al., 1994; Vigran et al., 2008), but is more resistant to thermal degradation compared to AOM (Vigran et al., 2008; Liu et al., 2019). The stratigraphic distribution of organic macerals can result in stratigraphic variability of the hydrocarbon generation potential of source rocks (Bohacs, 1993; Chandra et al., 1993; Slatt et al., 2012; Ade and Trindade, 2017). For example, Slatt et al. (2012) reported that cyclic variations of hydrocarbon generation potential in the Barnett Shale resulted from fluctuations of relative sea level at the parasequence scale. Ade and Trindade (2017) reported higher hydrogen index in the TST than in the HST in the southeastern Paraná Basin (Brazil).

The stratigraphic distribution of organic macerals within the New Albany Shale indicates that sea-level fluctuations control selective preservation of organic macerals by influencing bottom-water redox conditions and clastic supply. In the LST, organic macerals are dominated by alginite (Fig. 10). Relative sea level is low and bottom water is oxygenated in the LST, and under such conditions OM undergoes severe
microbial breakdown. Labile OM, which will turn into AOM later in diagenesis, would be largely destroyed and relatively resistant algal cysts which will turn into alginite during diagenesis will be enriched. Schieber (2001) also reported that alginite was more abundant than bituminite (a synonym of AOM; Kus et al., 2017) at the base of black shale packages within a given depositional sequence. At the onset of transgression, sea level rises and oxygen content in the bottom water decreases. Labile OM will be better preserved because of reduced microbial degradation. There is a slight decrease of AOM content at the MFS (Fig. 10), probably because sedimentation rates were so low that labile OM stayed on the sea floor long enough to be partially destroyed. The U/Th and Mo/Al ratio profiles also suggest that the most oxygen-deficient conditions did not occur at MFSs (Fig. 6). Schieber (2001) also found that alginite derived from Tasmanites cysts was enriched at MFSs in the Chattanooga and New Albany Shale of the eastern USA. As sea level drops, oxygen content in bottom water increases and microbial breakdown of labile OM intensifies. Better oxygenation of bottom waters is indicated by gray bioturbated beds in HSTs (Figs. 2, 3). In the Selmier Member, bioturbation is common and intense (Figs. 2, 3) and OM is dominated by alginite (Fig. 10), suggesting that relatively oxic conditions are not favorable for AOM preservation. In sequence 3, however, the AOM content reaches a maximum in the middle of the HST (Fig. 10). A potential explanation for this could be that increasing clastic supply in the HST improved preservation of AOM because timely cover by clastic materials prevented AOM from microbial breakdown. This is further supported by the coincidence of AOM and Al content maximum at about 625.0 m (Figs. 6, 10). Organic petrographic study of more closely spaced samples is needed to understand the stratigraphic variation of different types of OM and infer the control of depositional conditions on the differential preservation of organic macerals.

Authigenic precipitates of quartz and pyrite in Tasmanites cysts have been reported from Devonian black shales of the eastern USA (Schieber, 1996; Schieber et al., 2000; Schieber and Baird, 2001) as a consequence of very low sedimentation rates. These cyst fills are common in the TST and HST, and most prominent at the MFS where sedimentation rates were minimal. In the LST, filled Tasmanites cysts occur as well, but the fill in this case consists of detrital materials, such as grains of quartz, K-feldspar, recycled dolomite, mica, pyrite, and clay minerals (Fig. 17). Schieber and Lazar (2004) and Lazar (2007) pointed out that these detrital infills as well as an abundance of broken alga cysts (Fig. 17) are best explained as a consequence of sediments reworking by storm-induced bottom currents during lowstands.

5.5. Influence of relative sea-level fluctuations on OM accumulation

In this study, the stratigraphic distribution of TOC and geochemical proxies indicates that bottom-water redox conditions are one of the controlling factors for OM accumulation within a sequence stratigraphic context (Fig. 6). Clastic supply and paleoproductivity exert additional influences via dilution or preservation.

Relative sea-level fluctuations influence paleoproductivity, clastic supply, bottom-water redox conditions, and their combined control on OM accumulation in epicontinental seas (Arthur and Sageman, 2005). OM content has been linked to sequence stratigraphic context in a number of previous studies (Bohacs, 1993; Chandra et al., 1993; Creaney and Passey, 1993; Slatt and Rodriguez, 2012; Ade and Trindade, 2017; Dong et al., 2017; Byun et al., 2018; Harris et al., 2018), and in general terms, OM preferentially accumulates in TSTs because of lower clastic dilution and bottom-water oxygenation. OM is not as easily preserved in HSTs because of increasing clastic dilution and bottom-water oxygenation due to
shallowing, and is thought to reach a maximum at the MFS (Creaney and Passey, 1993). As pointed out above, in this study TOC maxima do not coincide with MFSs, presumably because OM residence times were such that substantial amounts of initial OM were remineralized prior to burial. Byun et al. (2018) reported similar TOC distribution patterns in the Middle Ordovician Jigunsan Formation of Korea, but attributed it to the reduction of productivity of benthic biota towards the MFS. The TOC distribution pattern seen in relation to sequence stratigraphic variability in the New Albany Shale may also apply to other black shale successions, which may have different depositional settings and histories. It is noteworthy that the stratigraphic variation of TOC content in black shale successions with high thermal maturity may not represent the variation of OM content when shales were deposited because of petroleum generation and expulsion (Tissot and Welte, 1984) and migration of solid bitumen within black shales (Hackley and Cardott, 2016; Liu et al., 2019).

Fig. 15. (A) Scanned core slab showing lag deposits at the MFS of sequence 3 at 632.64 m. (B) Scanned core slab showing lag deposits at the MFS of sequence 4 at 613.85 m. The carbonate nodules suggest low sedimentation rate. (C) Scanned polished thin section of the lower two lags in panel B. (D) SEM image (backscattered electron image) of the rectangular framed area in panel C. Note the Tasmanites cysts filled with authigenic quartz. T = Tasmanites cyst; Q = quartz; Py = pyrite.

Fig. 16. (A) Scanned core slab showing silty laminae in the Blocher Member. Note the combined-flow ripple in the lower right corner. The laminae are mainly composed of recycled dolomite, quartz, and pyrite. Also note silt-filled burrows and laminae disruptions, an indication of benthic life at the seafloor. (B) Scanned polished thin section of the sample in panel A. Laminae of several grains thick are very common, suggesting frequent storm and wave reworking of bottom sediments.
6. Conclusions

Detailed examination of TOC content, organic petrographic composition, and high-resolution (8 cm spacing) geochemical proxies of the New Albany Shale allow us to evaluate the influence of relative sea-level fluctuations on paleoproductivity, clastic supply, bottom-water redox conditions, and their complex interactions on OM enrichment in marine organic-rich sediments. The key contribution of this study is understanding stratigraphic variation of OM content and type within a sequence stratigraphic context. Specific conclusions can be drawn:

1. OM content and type in the New Albany Shale vary stratigraphically. The stratigraphic distribution of TOC content and geochemical proxies indicates that bottom-water redox conditions are one of the controlling factors for OM accumulation in the New Albany Shale. Paleoproductivity and clastic supply have dual impacts on OM accumulation; they can enrich or dilute OM.

2. Relative sea-level fluctuations influence paleoproductivity, clastic supply, bottom-water redox conditions, and their combined control on OM accumulation in the New Albany Shale. Within a sequence stratigraphic context, TOC content increases in TSTs, reaches a maximum near the MFS, and decreases in HSTs. Increasing clastic supply in HSTs can enhance the preservation of AOM and shift the AOM content maximum to the middle of HSTs. Enrichment of broken and detrital filled Tasmanites cysts indicates high-energy environments and LST conditions.

3. OM in the New Albany Shale is dominated by either AOM or alginit (mainly derived from Tasmanites cysts). Terrestrial OM (vitrinite + inertinite) content is <10%. AOM content increases in TSTs, reaches a maximum near the MFS, and decreases in HSTs. Increasing clastic supply in HSTs can enhance the preservation of AOM and shift the AOM content maximum to the middle of HSTs. Enrichment of broken and detrital filled Tasmanites cysts indicates high-energy environments and LST conditions.

4. Successful development of oil and gas from tight shale reservoirs requires a good understanding of stratigraphic distribution of TOC and hydrocarbon generation potential because OM plays different roles in shale oil vs shale gas plays (Jarvie, 2012a, 2012b; Bohacs et al., 2013). The results of this study can help better evaluate unconventional petroleum systems.

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