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THE ORGANIC GEOCHEMICAL VARIABILITY IN DEPOSITIONAL ENVIRONMENTAL PROCESSES WITHIN A STRATIGRAPHIC SEQUENCE OF THE WOODFORD SHALE IN THE ARDMORE BASIN, OKLAHOMA

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THE ORGANIC GEOCHEMICAL VARIABILITY IN DEPOSITIONAL ENVIRONMENTAL PROCESSES WITHIN A STRATIGRAPHIC SEQUENCE OF THE WOODFORD SHALE IN THE ARDMORE BASIN, OKLAHOMA

A THESIS APPROVED FOR THE SCHOOL OF GEOCIENCES

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To my parents and my sisters for their undying support, and bountiful of love... To my beloved friends who have supported me and helped me throughout my academic journey...

Thank you all for being the light in my life!

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ABSTRACT

The Devonian-Early Mississippian Woodford Shale underlying the subsurface of Carter and Murray Counties in the Ardmore Basin, Oklahoma, USA, has attracted growing interest as an unconventional reservoir. Heterogeneity has always been assumed within samples collected from different locations, while uniformity is often assumed in a stratigraphic sequence of the shale both in chemical and physical properties. However, the Woodford Shale is composed of characteristically different layers, a soft bed comprising of higher TOC, higher ductility and clay, as well as a hard layer that is lower in TOC, higher in brittleness and in silica content which greatly contributes to the variation in biomarker distributions. The bulk of the interval shows pristane/n- C_{17} vs. phytane/n- C_{18} ratios, C_{27}/C_{29} sterane ratios, diasterane/sterane ratios, and Ts/Tm ratios indicating a reduced, marine depositional environment, low in maturity.

The variations between the soft and hard beds are responsible for the variability of the biomarkers rather than solely depth. These variations can be seen within the C_{30} hopanes that show higher values within the harder beds relative to the higher TOC softer beds. Samples have been taken at 6" intervals over a 40ft. outcrop of the Woodford shale to investigate variability in organic facies, source input, and depositional environment conditions.

Shale samples have been characterized by Rock-Eval to determine TOC and hydrocarbon potential to identify variations of biomarkers in TOC rich intervals and TOC lean intervals. Additionally, shale extracts were processed by gas chromatography (GC), gas chromatographymass spectrometry (GC-MS) and GC-MS-MS to acquire biomarker traces that aid in the construction of paleoenvironmental conditions. Furthermore, the Frasnian-Famennian mass extinction which occurred at the Late Devonian is reflected to have occurred within this interval based on significant changes in the geochemical data. Variations in the biomarker distributions signify the heterogeneity of the shale over this 40ft interval, which in turn will yield fundamental information related to fine differences that affect reservoir quality in exploration and production of unconventional reservoirs.

CHAPTER I

1. Introduction

1.1 The Woodford Shale of Oklahoma

Fossil fuels have been central to energy resources, relying on hydrocarbon properties to fulfill energy needed to power everyday life. Fossil fuels have become the most sought after and effective source to power energy for life on Earth. It is the vital component to natural gas, oil, and coal for which researchers and industries are endlessly exploring today. The fame of oil and gas rose to fruition when the first oil well was drilled on August 27, 1859 near Titusville in northwestern Pennsylvania (Binet et al., 2002). Today, oil and natural gas have become the most important sources of energy and the engine to modern society, energizing the industry, heating homes, providing fuel for vehicles and all modes of mobile transportation. The increasing necessity for the utilization of hydrocarbons has allowed for the progression and advancement in technology for efficiency and improvement to enhance refineries and transportation of materials. According to Parra et al. (2013) development of shale plays has risen exponentially, and nearly 1,000,000 total wells have already been drilled in the United States according to U.S. Energy Information Administration (2019). Hydrocarbons are the product of continuous generation and accumulation following thermal degradation of biologically derived organic matter deposited in a sedimentary environment (Mulik and Erdman, 1963). The quality and quantity of the organic matter may vary laterally and vertically depending on the process of sedimentation, environmental conditions upon deposition, and the processes that occur post-deposition (diagenesis/thermal alteration). The organic rich-sediments in oil-forming basins are referred to as source rocks and reflect the process of sedimentation over geologic time (Moldowan et al., 1985). According to Peters and Cassa (1994), there are three criterias to consider sediments as "generative source rocks" and these

depend on quantity (amount of organic matter), quality (type of organic matter), and thermal maturity (extent of burial and heating).

The Woodford Shale has risen to fame as the most important and productive hydrocarbon source rock in Oklahoma as well as being the source to the majority of the state's hydrocarbon reserves (Slatt and Rodriguez, 2012; Philp and deGarmo, 2020). Galvis et al. (2018) noted the Woodford Shale, Late Devonian-Early Mississippian in age, has excellent source/reservoir rock properties, which enabled optimal production of oil and gas. Moreover, the Woodford Shale has unconventional reservoir properties that many researchers are continuing to explore and uncover. Unlike the typical conventional reservoirs, unconventional shale reservoirs require hydraulic stimulations to generate commercial rates of hydrocarbons (Temraz et al., 2016). According to Hammes et al. (2013) and Soliman et al. (2015) unconventional shale reservoirs, which include tight-gas sandstone, as well as gas and oil shales, are reservoirs that essentially require additional special recovery operations. Therefore, the inherently complex process of expulsion in unconventional shale reservoirs gives way to the accumulation of significant volumes of oil and gas (Comer, 2008). The major questions for scientists are related to the process of organic material incorporation and preservation into the sediments. In addition, it is appropriate to understand the depositional conditions and settings conducive for the organic matter to be converted into the desired petroleum components (Dembicki Jr., 2017). The mystery of its unconventional properties and the exploration of the Woodford Shale has led to a series of questions that inspired this research.

The approach of this thesis research is to undertake a very high resolution organic geochemical analysis of mudrocks (shales) within a small interval of interbedded siliceous, clay-shale, and chert of the Woodford group. The geochemical data would be integrated with the

geological facets of sequence stratigraphy to uncover further knowledge of the depositional environmental processes within the Ardmore Basin that created the Woodford Shale. The goal is to determine and reconstruct processes that occurred upon deposition in the Ardmore Basin that yield the fine variabilities found in biomarkers. This is to correlate relationships between the current stratigraphic model, and biomarker distributions. The outcome of this research is the provision of fine-scale depositional processes and environmental condition changes that impact large-scale changes seen in the shales today.

1.2 Biomarkers

This research explores and investigates the realm of biomarkers in organic geochemistry. Biomarkers relay the story of the environment in which the sediments were deposited and the organisms that thrived during the Late Devonian – Early Mississippian. Biomarkers are compounds derived from precursors present in living organisms that once thrived (Connan, 1981). Biological markers help scientists determine the environmental conditions during the deposition of sediments, the burial conditions in diagenesis, the thermal maturation the sediments experienced during catagenesis, the degree of biodegradation, and the nature of the source materials of the sediments (Eganhouse, 1997; Philp and deGarmo, 2020). The resulting information that biomarkers provide are helpful tools to correlate with sequence stratigraphy to build on the story of sediment deposition and processes occurring to the organic matter during and after deposition. Exploring biomarkers has been one of the leading geologic methods in exploration, development, and production for oil and gas industries as well as environmental industries (Dahl et al., 1994). Biomarkers mostly retain the molecular structure that they inherit from their biological source (Pratt et al., 1992). Although, the biomarkers do experience alteration and biodegradation during early diagenesis and catagenesis, this does not eradicate the source signature. The resilience of the carbon-based molecule structure within biomarkers provide geochemists with evidence of the contribution from either bacterial, algal, vascular plant, or animal sources that had accumulated in the sediments. Therefore, the varying geochemical output properties reflect the source input variations and the thermal alteration processes that occur post deposition (Moldowan et al., 1985). Organic matter can be found in two forms which are insoluble kerogen, and soluble bitumen where most of the biomarkers have been found to reside.

For biomarkers to become the effective and reliable indicators that they are today, they require optimal preservation conditions. Optimal preservation of organic matter upon and after deposition depend upon an environment that is anoxic (Volkman, 1988). This will commonly lead to organic-rich, oil-prone petroleum source rocks (Peters et al., 2005). According to Pratt et al. (1992), there are 3 stages for desired organic matter preservation: the first stage of preservation is restriction from oxidation and remineralization, followed by the prevention of destruction caused by both aerobic and anaerobic microbial process (Pratt et al., 1992; Romero and Philp, 2012) and lastly, the organic matter must retain its structure throughout the burial process (diagenesis) (Pratt et al., 1992). Therefore, the controls for optimal organic matter preservation are oxygen content within the water column, the biological productivity or the organisms thriving in the environment, water circulation/stratification, as well as sedimentation rate (Demaison and Moore, 1980). Furthermore, Kirkland et al. (1992) considered anoxic bottom-water conditions and limited water circulation to be the factors that inhibited aerobic microbial activity, which contributed to the enhanced preservation of the Woodford Shale. Different reducing conditions with varying Eh levels affect the biomarker precursor and their product pathways, which need to be taken into consideration (Moldowan et al., 1985).

1.3 Regional Geology

Devonian-Early Mississippian black shales have a complex depositional history reflecting a broad range of fundamental earth processes, such as paleoclimate, paleogeography, eustasy, and tectonism (Ettensohn, 1992). The Earth was in a greenhouse state during the Devonian and the warm climate provided a platform for high productivity. Tectonic activities have played a significant role in the evolution of black shale deposition as it provides the framework for the formation of restricted basins, favorable for organic-rich mud deposition (Klemme and Ulmishek, 1991; Ettensohn, 1997). With an overwhelming rate of subsidence over sedimentation, the sea floor became restricted, allowing for the water column to become stratified and dysoxic to anoxic (Ettensohn, 1992). Oklahoma is home to three major provinces that are the products of Early Paleozoic tectonic activities, namely the Oklahoma Basin, the Southern Oklahoma Aulacogen, and the Ouachita trough (Philp and deGarmo, 2020). The area of interest in this study is the Woodford Shale outcrop located within the Ardmore Basin of Oklahoma (Figure 1; Ham et al., 1964).



Figure 1. Major geologic provinces of Oklahoma encasing the Southern Oklahoma Aulacogen that comprises the Anadarko, Marietta, and Ardmore Basin (Ham et al., 1964).

The Ardmore Basin is a fault-bounded northwest depression of Pennsylvanian age in South Central Oklahoma (Granath, 1989). The Southern Oklahoma Aulacogen (SOA) is a major depocenter for the accumulation of the Woodford shale, due to an extensional failure in the form of a three-arm rift; SOA arose from the failed arm of the continental lithosphere extension which formed a NW-SE trending trough (Figure 2; Hoffman et al., 1974; Allen 2000; Brueske et al., 2016). The SOA is comprised of the Anadarko, Marietta, and Ardmore Basin (Figure 1; Ham et al., 1964). From the Cambrian to Early Devonian, shales, limestones, and sandstones were deposited within the mega-graben, that was a product of the failed arm of the SOA (Carlucci et al., 2014). After this period of deposition, and approaching the end of the Devonian, the Acadian Orogeny occurred, and eroded sediments that led to the deposition of the Woodford, Sycamore, Caney, and Springer formations well into the Early Pennsylvanian (Allen, 2000). Following this event, the Wichita Orogeny caused a major pulse of deformation, uplifting the Criner Hills 16,000ft above sea level. Then erosion of these structures provided sediments to be deposited into the Ardmore and Marietta Basin throughout the Late Mississippian (Figure 3; Allen, 2000). Further structural deformation occurred as the Arbuckle Orogeny caused the folding of the Caddo anticline and the Arbuckle anticlines (Figure 4; Northcutt and Campbell, 1995; Allen, 2000). The Ardmore Basin was mostly affected by tectonic activities in the Late Mississippian – Pennsylvanian. However, processes of sedimentation began starting from the Cambrian to Late Devonian. The phases of these events are illustrated in Figure 3 (Hoffman et al., 1974). The Late Devonian to Early Mississippian Woodford Shale was deposited over a span of 33 million years in the Ardmore Basin within Oklahoma, Central USA, and reflects a multiple history of deformation and deposition (Figure 5; Ham et al., 1973; Paxton et al., 2006; Galvis et al., 2018).



Figure 2. Map of Oklahoma and the tectonic activities during the early Paleozoic showing the Southern Oklahoma Aulacogen (Allen, 2000; Brueske et al., 2016). Two of the arms of the threearm rift joined to form the Early Paleozoic continental margin, while the third arm formed the SOA. CO: Colorado; OK: Oklahoma; MO: Missouri; AR; Arkansas; LA: Louisiana; TX: Texas; NM: New Mexico; WM: Wichita Mountains, AM; Arbuckle Mountains; WVF: Washita Valley Fault; MVF: Mountain View Fault.



Figure 3. The tectonic events that occurred from the Cambrian until the Permian to illustrate the basinal history of the Ardmore Basin in Oklahoma (Hoffman et al., 1974).



Figure 4. Tectonic map illustrating the location of the study within the Ardmore Basin at the southern flank of the Arbuckle Mountains (Northcutt and Campbell, 1995).



Figure 5. Cross-section across the Southern Oklahoma Aulacogen including the Woodford Shale in the Ardmore Basin. (Ham et al., 1973).

1.4 Paleogeography and climate

Understanding the variability within the sediment input also assists in the understanding of the global framework that sets the stage for erosional and depositional processes that have occurred. During the early Cambrian, marine limestones, sandstones, and shales were deposited into the SOA, over a broad epeiric sea, the Oklahoma Basin (Carlucci et al., 2014). Then, sea level experienced a eustatic drop that exposed the Devonian carbonate platform and created extensive incised valleys and karst topography on the North American Craton. The first stage of eustatic sea level fall provided the platform for the Woodford Shale to be deposited in a restricted marine stratified and dysoxic to anoxic environment throughout the Devonian (Ettensohn, 1992).

Marine transgressions overwhelmed the North American midcontinent craton during the Late Devonian times as shown in Figure 6. The rising sea level filled the NW-SE trending trough (that is deeper from the SE and shallows towards the N), which allowed for favorable accumulation of organic matter (Galvis et al., 2018). Therefore, the Woodford shale was deposited on a highly time-transgressive 2nd order sea level cycle as the Oklahoma shelf subsided and the Kaskaskia Sea lapped onto the Laurussian Craton (Figure 6; Kvale and Bynum, 2014; Philp and deGarmo, 2020). The Middle Woodford was deposited as a Transgressive System Tract (TST) which graded upwards into the Upper Woodford Highstand System Tract (HST; Romero and Philp, 2008; Philp and deGarmo, 2020). Though the Woodford Shale was deposited during a 2nd order sea level cycle, a superimposed 3rd order sequence stratigraphic parasequence developed as part of the Devonian Sea level curve (Figure 7a ; Slatt and Rodriguez, 2012; Philp and deGarmo, 2020). While this occurred, the maximum flooding surface and underlying Woodford allowed for the deposition of organic rich fine-grained sediments. Interestingly, the Late Devonian was also marked by the expansion of vascular land plants that may have increased chemical weathering and caused run-off of land-plant nutrients into the open marine setting (Algeo and Scheckler, 1998). This is a significant event that affected the organic matter input introduced into the basin. It will be discussed in detail in the following chapters.



Figure 6. Paleogeography of the mid-continent North America within the Late Devonian and Early Mississippian illustrating the marine transgression that overwhelmed the Oklahoma region (Robinson and Kirschbaum, 1995; modified from Comer, 2008; Blakey, 2011; and illustrated by Galvis, 2017).

Percival et al. (2019) determined that the Late Devonian (383-359Ma) was a time marked by numerous environmental and biotic crises, which included one of the Big Five mass extinction events, namely the Frasnian-Famennian extinction. There was widespread development of marine anoxia recorded by the appearance of organic-rich laminated shales in the stratigraphic record (Buggisch and Joachimski, 2006). There were two anoxic episodes during the Late Frasnian, which are widely known as the Lower Kellwasser and the Upper Kellwasser. There were many factors that affected the climate during the Late Devonian, including extra-terrestrial impacts, large scale volcanic activity potentially linked to the Viluy Traps in Siberia, orogenic uplifts and erosions, as well as the expansion of vascular-rooted terrigenous flora (Wang, 1992; Algeo et al., 1995; Algeo and Scheckler, 1998).

1.5 Sequence Stratigraphy and the Woodford Shale

The Woodford Shale in the Ardmore Basin is among the few complete sections in which all its members (Upper Woodford, Middle Woodford, and Lower Woodford) are exposed and preserved in outcrop at the Speake Ranch location (Galvis et al., 2018). The sequence stratigraphic model is highly correlative to other existing North American Devonian black shales that exhibit these unconventional features, such as the Antrim Shale (Michigan Basin), Ohio Shale (Appalachian basin), New Albany Shale (Illinois Basin), Bakken Shale (Williston Basin), and Exshaw Formation (Western Canada Basin) (Comer, 2008). The Woodford shale has widely been discussed as a formation that is generally deposited in an open marine environment (Slatt and Rodriguez, 2012; Galvis et al., 2018). Moreover, the deposition of the Woodford Shale is highly dependent and controlled by factors such as eustatic sea level fluctuations among many other factors. The cyclic fluctuations in eustatic sea level has caused variability in topographic relief, karsting, periodic subaerial exposure, and subterranean collapse which produced significant local depressions as show in Figure 7b. As sea level periodically drops at Lowstand System Tracts (left falling curve on the Sea Level Curve on Figure 7a), the carbonate platform experienced a periodic subaerial exposure that carved the depressions/minibasins (Slatt and Rodriguez, 2012). The combination of minibasins and sea level fluctuation provides the foundation of variability in sediment influx and organic matter accumulations both vertically as well as laterally, therefore resulting in stratigraphic variability. During a Transgressive System Tract, the rising of the seas

allowed for circulation, deposition of organic rich sediments, and oxygenation of these depressions. Then, in the incoming Highstand System Tract as sea level regressed, the oxygenated waters oxidized the land or marine derived organic matter and allowed for the deposition of fine sediments poor in organic matter. The Lower Woodford and Middle Woodford were primarily deposited during a Transgressive System Tract, with a massive organic rich condensed section and a maximum flooding surface. The ideal preservation of organic matter, where total organic carbon is seen at a maximum, is at a time of Lowstand System Tract, when oxygenation circulation was poor, and the basin was restricted (Figure 8). Biomarkers are important tools in reconstructing the paleoenvironment as they retain conditions of the environment that include levels of oxygenation and eutrophication, in which determine the resulting water depth fluctuation (Philp, and Degarmo, 2020). Specifically, considering the Eh and pH conditions of anoxic environments are important to consider, as they change with varying oxygenation levels. This is caused by their dependency of bacterially mediated sulfate reduction resulting from varying levels of anoxia that may have occurred (Moldowan et al., 1985). Therefore, the key stratigraphic feature of the Woodford shale is the variation of high TOC values that are found in mini-basin topographic lows. Ensuing tectonic events from the Cambrian to the Permian shaped the topography of the N. American mid-continent and created the platform for accumulation of organic rich sediments in restricted basins. Therefore, topography is the control for the variability we see in the lithofacies of the Woodford formation.



b.)

a.)



Figure 7(a). Sea Level curve displaying relative positions during a Highstand System Tract, Transgressive System Tract, and Lowstand System Tract; and (b). model of the sea level fluctuation stages relative to the depositional and erosional process (Slatt et al., 2016; Slatt and Rodriguez, 2012).

The Woodford Shale is comprised of 13 identified lithofacies and divided into three members which are the Upper Woodford (UW), Middle Woodford (MW), and Lower Woodford

(LW) (Galvis et al., 2018). For the purpose of this research, the division of the members has been classified by their lithologic attributes (Slatt et al., 2012). The geologic provinces in Oklahoma feature a range of geologic formations, ranging from shallow water carbonate platform to clastic abyssal plains, that contain variability within each sequence stratigraphy and reflect different depositional environments (Figure 8; Comer, 2008). There is a consistent alternation of brittle (hard) and ductile (soft) layers throughout the Middle and Upper Woodford. The hard layers are siliciclastic rich sourced from biogenic radiolaria, while the soft layers are clay rich and are the most organic rich intervals (Galvis et al., 2018). The application of this hard-soft couplet to the industry is very helpful as the beds are stratigraphically targeted for drilling. The ductile layer target is the oil zone and the source rock generating the desired hydrocarbons, whereas the brittle zone is the reservoir (Slatt, 2013). This trend has been recognized to be the hard-soft couplet (Slatt et al., 2012).



Figure 8. Model of water stratification, and anoxic environment within the minibasins for the Woodford Shale in the Ardmore Basin, Oklahoma.

1.6 *Objectives*

There is rising interest in the Late Devonian Woodford Shale to many researchers and the oil and gas industry as a prolific unconventional reservoir and a highly generative source rock. Multiple processes that occurred during deposition, such as cycles of sea level changes, the expansion of rooted vascular plants, climate changes and atmospheric composition changes, directly affected the influx of nutrients to the sediments deposited in the Late Devonian Woodford Shale and contributed to the variabilities seen in the formation (Philp and DeGarmo, 2020). Many studies have been accomplished over sedimentological analysis, stratigraphic description, structure analysis, and seismic analysis locally, but a high-resolution geochemical study over the section, which specifically encases one of the 'Big Five' global extinction events (Frasnian-Famennian time), is uncommon. The high-resolution analysis is hypothesized to yield significant variability that can help determine reservoir quality and production.

Galvis et al. (2018) presented a detailed rock-based and stratigraphic documentation of the entire Woodford Shale outcrop from the Speake Ranch, Oklahoma, for his M.S. thesis. He identified the Woodford shale to be a transgressive section composed of three different claystones, black argillaceous shale, two types of siliceous shales, siliceous dolomitic shale, two types of limestones and mudstones, as well as two distinct types of cherts (Galvis, 2017). The seven main lithofacies of the 13 identified have high frequency of vertical interbedding of hard (chert) and soft (siliceous, clay-shale) beds within the Middle and Upper members, due in part to the sea level changes (Galvis et al., 2018).
1.7 Purpose of study

- I. The purpose of the study is to conduct a high resolution organic geochemical investigation understanding the effect of depositional processes, lithofacies, and stratigraphy on the distribution of biomarkers of the Late Devonian Early Mississippian Woodford Shale.
- II. Integrate geochemical analysis into the sequence stratigraphic framework of the Woodford Shale established by Galvis (Portilla, 2018).
- III. Compare and determine minor variabilities (if any) of the samples throughout the 40ft section of the Woodford Shale from the Speake Ranch. Recognize to what degree the complex depositional architecture would control the distribution of various organic compounds.
- IV. Identify the Frasnian-Famennian extinction boundary within the section based upon organic geochemistry.

CHAPTER II

2. METHODOLOGY

2.1 Study Area and Sample Location

All the samples analyzed for this research were obtained from Dr. Roger Slatt's core laboratory in Sarkey's Energy Center at the University of Oklahoma campus. These samples were collected from an exposed outcrop at the Speake Ranch within the Ardmore Basin, located between the Carter and Murray County of Oklahoma as shown in Figure 9 (Cardott, 2012; Galvis, 2017). This specific location lies on the southwestern end of the Arbuckle mountains (Galvis et al., 2018). This private quarry on the Speake Ranch, with the geographic coordinates 34°22'40.36" N and 97°20'17.26" W, has the complete, but exposed, sections of the Woodford Formation; making it one of the few preserved outcrops to exist in South Central Oklahoma. The exposed sections span to 335ft of true stratigraphic thickness that comprise the Upper Woodford, Middle Woodford, and Lower Woodford as well as the basal and upper contact with Hunton Group and Sycamore formation respectively. Due to the unconformity boundary of the Hunton, the Woodford Shale across Oklahoma can vary in thickness from 350 to 400ft, displaying the inverse relationship between the Woodford and the Hunton that highlights the effect of paleotopography on the distribution of thicknesses and consequently lithofacies (Ham, 1973; Fay, 1989; Galvis et al., 2018).







Figure 9. Top figure is location of Ardmore Basin in Oklahoma between the Carter and Murray county which is included in the SCOOP play. Middle figure is the location of Speake Ranch Quarry just west off of I-35 in Southern Oklahoma. Bottom figure is the exposed Speake Ranch Outcrop displaying the complete section of the Woodford Formation including the underlying Hunton Formation, and the overlying Sycamore Formation (Cardott, 2012; Galvis, 2017).

2.2 Lithofacies Classification and discussion

The Woodford Shale is comprised of 13 identified lithofacies and divided into three members, which are the Upper Woodford (UW), Middle Woodford (MW), and Lower Woodford (LW) (Galvis et al., 2018). The Woodford members were divided and classified based on its lithologic attributes (Slatt and Rodriguez, 2012). The Lower Woodford is predominantly a black-gray fissile, siliceous/carbonaceous shale, with silicified logs (Molinares, 2013). The Middle Woodford is a dominant pyritic, black shale characterized by the highest Gamma Ray output in response to high TOC (Molinares, 2013). The Upper Woodford is primarily quartzose, phosphatic, gray-black shale dominated by phosphate lenses and pyrite nodules (Molinares, 2013). Within each member lies more variability in the lithofacies and lithologic characteristics as shown in Figure 10. At a finer scale, the Upper and Middle Woodford inhibit intervals of interbedded chert and fissile shale, causing frequent high-low cyclicity within the Gamma Ray log as shown in Figure 10 (Slatt and Rodriguez, 2012; Galvis, 2017). The oscillation of the Gamma Ray throughout the Woodford shale has been recognized as the soft and hard couplet correlating to its lithofacies characteristics (Figure 10; Galvis, 2017).



Figure 10. Stratigraphic column and outcrop Gamma Ray depth log of the Woodford Shale from Speake Ranch. The colors of the lithofacies correlate to those in Figure 11. The Upper Woodford is overwhelmed with phosphate concretions within the chert lithofacies indicated by the red circles (Galvis, 2017).

The soft layer is higher in ductility, as it is dominated by abundant clay content, which contains both detrital and authigenic quartz. The soft layers are the intervals with the highest TOC values (Slatt and Rodriguez, 2012). Detrital quartz is often found near continents, along shelf (especially the northwestern shelf of the Anadarko Basin), and basin depocenters where bottom water turbidity flows converge (Comer, 2008). Meanwhile, the hard layers are higher in brittleness

caused by the diagenetically-recrystalized radiolarians and have relatively lower TOC values (Slatt and Rodriguez, 2012). There is a clear trend that shows the ratio of hard to soft beds, which increases upward in the Upper Woodford beginning from the Middle Woodford as more siliclastic facies are introduced into the section. The 13 lithofacies identified can be divided into two major lithologic features which can be further subdivided into various shales and claystones. Brown claystone, green claystone, green silty claystone, black argillaceous shale, black siliceous shale, brown siliceous shale, and siliceous dolomitic shale fall under the soft bed intervals that are spread throughout the Woodford Formation (Figure 11; Galvis, 2017; Galvis et al., 2018). The second major features that fall under the indurated, blocky bed intervals are comprised of crystalline limestone, micritic limestone, dolomitic limestone, siliceous mudstone, black chert, and nonorganic chert (Figure 11; Galvis, 2017; Galvis et al., 2018). The 13 identified lithofacies are divided into seven major lithofacies of claystones, shales, limestones, mudstones, and chert as described below (Figure 11).



Figure 11. The 13 identified lithofacies within the Woodford Shale of South Central Oklahoma distinguished between the soft and hard layers (Galvis, 2017).

2.2.1 Argillaceous Shales

The argillaceous shale is a member of the soft bed containing the most clay content and is one of the most fissile lithofacies (Galvis, 2017). These layers are found to be predominantly black with high TOC values compared to other lithofacies (Galvis, 2017).

2.2.2 Siliceous Shales

The siliceous shales are among the soft lithofacies that are the most quartz-rich (Galvis, 2017). These lithofacies are predominantly black and gray, but brown siliceous shales are often found interbedded within the Upper Woodford. These intervals are typically slightly indurated and are composed of microcrystalline quartz, clay minerals, and *Tasmanites* (Galvis, 2017). The gray/black siliceous shales exhibit thicker parting layering among all the other softer lithofacies throughout all the members of the Woodford Shale.

2.2.3 Brown Siliceous Shales

The brown siliceous shale is a member of the soft fissile-laminated layer. This lithofacies dominates the Upper and Middle Woodford interlayering between the brittle black chert. The total organic carbon within this layer ranges from 3-12wt% (Galvis, 2017).

2.2.4 Siliceous-Dolomitic Shales

The siliceous dolomitic shale is one of the less frequent of the soft lithofacies with moderate quartz and clay content (Galvis, 2017). This interval has TOC values ranging from 8-14wt% and is often found in the Lower and Middle Woodford.

2.2.5 Siliceous Mudstones

The siliceous mudstone is among the few lithofacies that contain well-preserved and silicified radiolarians and *Tasmanites*. This lithofacies of the ductile layer member dominates throughout all three members of the Woodford (UW, MW, and LW). Its total organic carbon ranges from 3-8% and falls under the relatively lower organic content abundance (Galvis, 2017).

2.2.6 Cherts

The black chert layer is among the most dominant and abundant silica-rich lithofacies of the hard interval members with quartz richness reaching from 87 to 100% (Galvis, 2017). Its brittle characteristic is owed due to the replacement of radiolaria by chalcedony quartz and pyrite (Galvis, 2017). These intervals are found in increasingly thick partitions approaching the Upper Woodford from the Lower Woodford. Its increase in abundance is related to the opening of the ocean and is often found along continental margins or distal areas of cratonic basins such as the Ardmore Basin, Marietta Basin, and Anadarko Basin (Comer, 2008).

2.2.7 Dolomitic Mudstone

Dolomitic mudstone is a member of the hard lithofacies that is less frequent among the others in this lithofacies group. It has very high calcareous content with low TOC values less than 4 wt% (Galvis, 2017).

2.3 Experimental

2.3.1 Rock-Eval Pyrolysis and Leco – TOC

Five hundred thirty-six outcrop samples from the Woodford outcrop at the Speake Ranch within the Ardmore Basin located between the Carter and Murray counties of Oklahoma were collected at approximately 6" intervals. As far as possible, the samples were collected below the weathering zone, although with extensive network of microfractures that had developed over time, this was not always possible. Aliquots (3-4g) of each sample were subjected to Rock-Eval analysis to obtain TOC, HI, OI, S₁, S₂, S₃ values at GeoMark. A TOC value of >1 wt% is indicative of a good petroleum potential rock and the Woodford Shale ranges from 0.05 to 33.9wt% (Peters and Cassa, 1994). The purpose of this research is to consider the relatively high and low TOC values into the correlation of geochemical data variations.

2.3.2 Laboratory Experimental Procedures

2.3.2.1 Preliminary Samples Preparation

A total of 40 samples of Speake Ranch outcrop were collected from Dr. Roger Slatt's core lab in Sarkey's Energy Center at the University of Oklahoma. In this study, the samples included both the soft and hard beds of the Upper Woodford and Middle Woodford at 190ft-230ft (Fig. 4). This interval is of interest since it is thought to straddle the Frasnian-Famennian (F-F) boundary. An aim of this study is to determine whether this F-F boundary could be recognized by a highresolution biomarker investigation. Each sample was crushed to powder with a porcelain mortar and pestle and sieved through a US Standard mesh No. 25. The resulting powder material was then stored in glass jars to be used for extraction and fractionation procedures.

2.3.2.2 Extraction of Organic Matter

Approximately 50g of the rock powder were placed in a Soxhlet extraction system, which consisted of the Soxhlet device, thimble, and glass wool to extract the organic constituents of the powdered shale using 350-400ml mixture of methanol (CH₃OH) and dichloromethane (CH₂Cl₂) (1:1). The mixture of methanol and dichloromethane provided an efficient extraction method since it utilized solvents of different polarity. The bitumen is diluted in a less polar solvent (DCM) and a more polar solvent (MeOH) to enhance the expansion of clays and allow for the access to the organic compounds residing within the rock sample (Brassell et al., 1992). Within the Soxhlet apparatus, extraction solvent vaporizes from the 500ml round-bottom flask up through the side arm of the apparatus, condenses at the top, and drips down into the thimble containing the powdered sample, before it finally returns to the round-bottom flask through a siphon. This cycle of mildly hot solvent running through the samples was repeated over a span of 24 hours, to extract the maximum amount of organic matter from the crushed rock samples. The resulting solvent containing the extract was concentrated with a rotary evaporator, prior to transfer to a 4ml vial. This process workflow is illustrated in Figure 12 (a).

(a).



(b).



Figure 12(a). Process workflow of sample processing of the Woodford Shale for organic biomarker analysis; (b). Enhanced process workflow of sample processing of the Woodford Shale for organic biomarker analysis displaying the detailed fractionation process.

2.3.2.3 De-Asphaltening Process – Bitumen Fractionation

A known weight of bitumen (approximately 100mg) was transferred into a 50ml centrifuge tube and excess n-pentane (C_5H_{12}) was added to precipitate the asphaltenes. The bitumen solution was placed in a freezer for approximately 15 hours to ensure complete precipitation of the asphaltenes. Next, the mixture was centrifuged and the maltene solution in the pentane was decanted into a 250ml round bottom flask as described below. The asphaltene precipitate was dissolved in DCM and transferred to a 4ml vial, dried, and quantified.

2.3.2.4 Maltene Fractionation

The pentane containing the dissolved maltene fraction was transferred from the 50ml centrifuge tube into a 250ml round-bottom flask and the excess solvent removed using the rotary evaporator. The maltenes were fractionated into saturates, aromatics, and polar NSO constituents using a 50ml glass column packed with a small amount of glass wool and 8.5g of activated alumina (Al₂O₃), which had been rinsed using n-hexane (C₆H₁₄). The extended workflow of the extraction and fractionation procedure can be found in Figure 12(b). The saturate fraction was eluted with 16ml of n-hexane (C₆H₁₄). The aromatic fraction was eluted using 25ml of a (7:3) mixture of n-hexane (C₆H₁₄) and dichloromethane (CH₂Cl₂). Finally, the polar resins were eluted using 25ml (1:1) of a mixture of dichloromethane (CH₂Cl₂) and methanol (CH₃OH). The saturate and aromatic fractions were screened by GC, and selected sample fractions were then analyzed by GC-MS for biomarker distributions.. Only 37 of the samples were able to be fractionated and screened due to low extract.

2.3.2.5 *Gas Chromatography*

All 37 samples were diluted in a 4ml vial using n-hexane to prepare for the GC and GC-MS analyses. Gas chromatography has widely been used as a separation tool of complex mixtures, by volatizing organic compounds at high temperatures. An Agilent Technologies 6890 GC was used to screen all the samples. The GC was equipped with a splitless capillary injection system and a 60m J& W Scientific DB-1 122-0162 fused silica column (0.250mm i.d. and 0.2 µm liquid film coating. The GC was programmed at an initial temperature of 40°C to be held until 1.5-minute post sample injection, where it is followed by a gradual temperature rise of 4°C/min. The temperature is programmed to reach 310°C and be held constant for 24 minutes until method completion. The diluted sample $(1\mu L)$ is introduced into the GC inlet, volatized, and transferred to the GC column by a Helium gas carrier at a flow rate of 1.4ml/min. The GC column then separates the compounds based on factors such as polarity, shape, stereochemistry, volatility and size. The purpose of the temperature ramp is to further enhance volatilization and separation for the larger molecules. Though, extreme high temperature was avoided as it would cause premature degradation of the samples and affect the resulting data outcome. As the constituents elute from the column, they pass through a flame ionization detector (FID) where they are combusted and ionized, and a signal is recorded relative to their abundance. The more volatile components elute first, and are the earliest peaks in the chromatograms, as shown in Figure 13 (Brassell et al., 1992; Philp, 2000).



Figure 13. Gas chromatograms displaying the distribution and separation of the n-alkanes. Upper right chromatogram shows the saturate and aromatic fractions (Philp, 2000).

2.3.2.6 Gas Chromatography – Mass Spectrometry

Thirty-seven of the saturate and aromatic fractions were diluted in dichloromethane to a concentration of 4mg/ml and injected with C_{27} -deuterated cholestane (107ppm concentration) used as a standard. An Agilent Technologies 7890A GC was used to screen all the samples with a splitless capillary injection system and a 60m J & W Scientific DB-5MS 122-5562 fused silica column coupled with an Agilent Technologies 5975 XL Mass Selective Detector. Just like the GC, the GC-MS is programmed at an initial temperature of 40°C for 1.5min post sample injection, then it is gradually increased to a temperature of 315°C where it is held constant for 50 minutes. Single-ion monitoring or multiple ion detection was utilized to assess the distribution and concentration

of the hydrocarbons residing within the samples. The coupling of a gas chromatograph to a mass spectrometer helps to determine and identify more complex molecules such as biomarkers that tend to show at relatively low concentrations (Philp, 2000). Unlike the GC, which is specifically characterized by retention times and abundances, the GC-MS aids in the identification of molecules and is most helpful for biomarker analysis (Brassell et al., 1992).

CHAPTER III

3. Results and Discussion

3.1 Geochemical Screening

Rock-Eval pyrolysis coupled with TOC measurements have widely been used for source rock characterization (Dahl et al., 2004). Rock samples are introduced to an atmosphere of inert nitrogen and heated to produce organic products. A flame ionization detector is incorporated to detect the compounds emitted from the rock to be utilized as parameters that characterize the source rock. Essentially, Rock-Eval pyrolysis is the volatization of hydrocarbons residing in the potential source rock (Peters, 1986). The amount of hydrocarbons, CO and CO_2 released are measured to determine quantity, type, and thermal maturity of potential source rocks. Rock-Eval pyrolysis alone is insufficient in examining source rock characterization, as it only quantifies the TOC, petroleum potential, and dominant kerogen type (Demaison and Moore, 1980; Cooper and Barnard, 1984; Isaksen and Ledje, 2001). Throughout the heating process, the temperature programmed pyrolysis detects 5 peaks with increasing temperature that records the degradation of the organic matter in the rocks (Figure 14; Pepper and Corvie, 1995; McCarthy et al., 2011). S₁ is the first peak detected and represents the free hydrocarbons. The free hydrocarbons represent the bitumen in the sample generated in the subsurface that are released during the first stage of heating (Peters, 1986). S₂ represents the hydrocarbons generated by thermal breakdown of kerogen in the source rock at second stage of heating, which reflects the hydrocarbon potential (Peters, 1986). T_{max} is the peak pyrolysis temperature of S_2 at which maximum hydrocarbons are generated. Knowing the T_{max} value reached during S_2 is important to reveal the history of the rock and the thermal alteration the rock has experienced. S_3 is derived from the evolution of CO_2 during thermal cracking of kerogen measured by a thermal conductivity detection. This is followed by S₄,

representing the oxidation of residual organic carbon that is separated into CO and CO₂ components (McCarthy et al., 2011). The very last peak of pyrolysis is the S_5 , referring to the carbon dioxide resulted from the decomposition of carbonate material (McCarthy et al., 2011). Table 1 displays the geochemical screening parameters used in this research as defined by Peters and Cassa (1994) and McCarthy et al. (2011).



Figure 14. Programmed temperature of Rock-Eval 6 pyrolysis displaying the resulting products and heat signatures (McCarthy et al., 2011).

The petroleum-generating potential of the source rock is directly dependent on volume, organic richness and thermal maturity. Combining the pyrolysis measurements provides information on the potential of petroleum generation, kerogen type, and maturity, which are important for source rock characterization. It is also important to note that with increasing temperature during pyrolysis, abundant expulsion of hydrocarbons results, and the reactive kerogen is consequently consumed, which will decrease the TOC wt% and hydrogen, therefore resulting in a decrease of S_2 . In addition, a decrease in S_2 and hydrogen content is also caused by a process of hydrocarbon adsorption onto the clay minerals of the samples, that are converted to non-volatile char and light materials (Katz, 1983; Peters, 1986; Dahl et al., 2004). The mineral matrix plays a significant role on S_2 pyrolysis yield, and associated T_{max} value. The T_{max} represents the temperature of maximum hydrocarbon generation (S_2). Therefore, abrupt changes in lithology within the sequence, such as the hard and soft lithofacies of Speake Ranch samples, will cause changes in the T_{max} values (Dahl et al., 2004).

Parameter	Definition	Units
TOC	Total Organic Carbon	Wt%
S1	Free Hydrocarbon released at 300°C	mg HC/ g rock
S2	Residual Hydrocarbon	mg HC/ g rock
\$3	Evolution of CO_2 during thermal cracking	$Mg CO_2/g rock$
Tmax	Hydrocarbons generated during S2	°C
HI	Hydrogen Index, quantified hydrocarbons from S2 (HI = S2*100/TOC)	mg HC/g TOC
OI	Oxygen Index, quantified Carbon Dioxide from S3	mg HC/g TOC

Table 1. Geochemical screening parameters defined by Peters and Cassa (1994) and Tissot and Welte (1984) for Rock-Eval pyrolysis and Leco-TOC.

3.2 Organic Richness

Evaluation of organic richness through Rock-Eval pyrolysis includes a quantifiable organic carbon assessment. According to Peters (1986) what constitutes an optimal source rock generative potential is a TOC wt% greater than 2% coupled with S₁ values greater than 2 mg HC/g rock and S₂ values greater than 10 mg HC/g rock. All the members of the Woodford Shale formation from Speake Ranch are abundant in TOC with values ranging from 0.05 to 33.9 wt% (Galvis, 2017). The Lower Woodford displays the highest organic rich interval, owing to its clay rich composition (Figure 15; Galvis, 2017). Kennedy et al. (2014) found that there is a direct relationship between the high surface area of detrital clay minerals with organic carbon, which indicates that abundance of clay provides for an extended preservation capability of organic matter. This property aligns with the characteristics of the Woodford Shale. The Middle Woodford and Upper Woodford from Speake Ranch have TOC values ranging from 0.62% to 15.90%. This classifies the Woodford Shale from Speake Ranch to be an excellent potential source rock.



Figure 15. TOC abundance distribution across all the Woodford members from Speake Ranch studied by Galvis (2017). The Lower Woodford displays the most organic rich intervals within the soft lithofacies intervals. The Middle Woodford shows nearly equivalent ratio of hard to soft lithofacies. The Upper Woodford has the lowest average TOC values and a higher hard to soft lithofacies ratio. The TOC and Gamma Ray plots illustrate that high TOC is not always correlative to high Gamma Ray response. The higher Gamma Ray plots may be an effect of the radioactive component, phosphatic particles (Galvis, 2017).

There is an apparent variability throughout the Speake Ranch interval seen in Figure 16 correlating with Gamma Ray, which has been described as the soft and hard couplet by Slatt and Rodriguez (2012). It is clearly illustrated that the hardness depth-log oscillation is correlative to the variations of the TOC, defining the hard-soft bed couplet (Figure 16). The average TOC wt% shows a clear and common trend of decreasing values towards the Upper Woodford from the Lower Woodford. However, a resulting high TOC wt% (5-35%) in the Upper Woodford may be a result of possible migrated hydrocarbons (Torres, 2020). The frequency of the TOC fluctuations within the 40ft sequence may be a product of geochemical and sedimentological effects of water column anoxia (Kennedy and Wagner, 2011). There is a covariance between TOC wt% and

lithofacies earlier discovered by Galvis (2017), where the quartz-rich brittle layers have a negative correlation to TOC wt% richness whist the clay-rich ductile beds have a positive relationship with organic matter abundance (Figure 17). Significant fluctuations of TOC wt% can be found at very small scales, especially in the studied 40ft interval, where TOC values are observed to have jumped from 5.19 (hard bed) to 12.9 (soft bed) wt% within just 1ft. It is hypothesized that regardless of the location of the sample within the 40ft interval, similar rock lithofacies from the middle or upper of the section will have similar TOC wt% abundances (discussed in Chapter IV). Clay minerals are initially formed in soils prior to erosion and transportation to continental margins, then climate and continental processes alter their mineralogy and affect their preservation capacity (Kennedy and Wagner, 2011). In contrast, the lower TOC contents are commonly found within beds containing cherts resulting in dilution of organics caused by syn-deposition of biogenic silica (Roberts and Mitterer, 1992; Galvis, 2017). Throughout this research the defining term to identify TOC abundance will be dependent on the brittle-ductile (hard-soft bed) couplet; where the soft bed intervals that occur at every foot have relatively higher TOC wt%. In each foot of the Speake Ranch intervals, the average bed thicknesses per rock type were measured in order to estimate the soft-to-hard ratio, relating the cumulative thickness of soft beds over the cumulative thickness of hard beds per each foot (Galvis, 2017). One foot of Speake Ranch interval may be made of 50/50 of hard and soft, however there are still several combinations where the alternation of soft and hard beds stack to accommodate the 50/50 within that foot.







Figure 17. Soft vs. hard relationship with TOC. The softer bed intervals have positive correlations with increasing TOC, while the hard lithofacies intervals have negative correlations with TOC (Galvis, 2017). Hard bed lithofacies are enriched in quartz content and depleted in clay content, while the soft bed lithofacies are enriched in clay content. The outlier symbols of the hard bed and soft beds with significantly low quartz content, may be an indication of dolomites.

These relatively lower TOC values which are found within the brittle beds, may be a product of the predominant regressive 3rd order hemicycles (Molinares, 2013). As sea level regressed the water column shallows and introduces oxygen to bottom waters through vertical mixing. This produces turbulence in sediments that caused disturbances to the anoxia of bottom water sediments ideal for organic matter preservation (Figure 18). In support of Molinares (2013), Philp and DeGarmo (2020) interpreted that lower TOC values are the results of paleoweathering from subaerial exposure as a consequence of sea level regression during climate transition from Late Devonian to Early Mississippian time. Kennedy and Wagner (2011) have suggested that bioturbated intervals, yield lower organic carbon values due to remineralization and removal of organic carbon on sediment surface by oxygenation, which could also apply to the Speake Ranch samples. Considering the above-mentioned possibilities, it is likely that the lower TOC intervals of the Woodford Shale from Speake Ranch resulted from loss of conditions necessary for organic

matter preservation. These conditions include subaerial exposure, turbulence, and disruption of the anoxic water columns. The sequence stratigraphy mentioned in earlier chapters explained that the ratio of the hard to soft lithology increases towards the Upper Woodford from the Middle Woodford, leading to a higher frequency of lower TOC values toward the Upper Woodford. Furthermore, the Si/Al record provides additional support for the association of clay minerals with organic matter, indicating that there is an increase of clay minerals, relative to quartz, in each interval with relatively higher TOC values (Figure 19). The presence of clay-rich composition not only provides favorable conditions for carbon preservation, but it also reflects climatic conditions and sea level fluctuations of the environment relating to marine carbon deposition and burial, which is explored in this research (Kennedy and Wagner, 2011). The bulk geochemical parameters earlier defined such as TOC, HI, OI, Si/Al, maturity, and GR are shown in Figure 19.



Figure 18. Model of depositional environment and water column s for both higher water levels and lower water levels. Higher water levels cause vertical mixing of the water column and introduces oxygen to the system. Lower water levels cause a restricted water column accompanied with stratification (Tulipani et al., 2013).





3.3 Organic Matter Type

Kerogen maceral composition, which include liptinite, amorphinite, vitrinite, and inertinite, have been utilized for source rock evaluation. Liptinite macerals such as alginite, sporinite, cutinite, and resinite, are dominant within Type I kerogens and are oil producing (Dahl et al., 2004). Amorphinite macerals are insoluble macerals common in Type II Kerogen and are the most oil prolific. Vitrinite macerals originate from woody plant tissue and are predominantly found in Type III kerogens (Dahl et al., 2004). Finally, inertinite macerals are dominant within Type IV kerogen profile which generate little or no petroleum (McCarthy et al., 2011). The plot of the Hydrogen Index vs. T_{max} identified the Woodford Shale to be primarily a Type II kerogen type characterization will be explained in the next section.



Figure 20. Kerogen classification and maturity assessment utilizing T_{max} vs. Hydrogen Index (HI) parameter plot useful in aiding primary interpretation of source rock analysis. The Speake Ranch samples are primarily a Type II kerogen with several samples plotting as Type III.

3.4 Kerogen Type

Kerogen is the high molecular weight insoluble organic residue in which can generate hydrocarbons during pyrolysis or maturation. Kerogen can be classified into four types, depending on Hydrogen Index (HI) and Oxygen Index (OI) values. The pseudo van Krevelen plot has widely been used to define the 4 types of kerogen to determine its compositional variances and organic matter types (Figure 21; Tissot et al., 1974). The HI is quantified by normalizing S₂ peak to TOC multiplied by 100 which will result in the quantified hydrocarbons from S₂ relative to TOC measured in mg HC/g TOC (McCarthy et al., 2011; Dahl et al., 2014). The OI is quantified from S_3 relative to TOC which measures the quantified carbon dioxide measured in mg CO₂/g TOC (McCarthy et al., 2011). The following sections will discuss the use of biomarkers to identify the influx of organic material and processes that occurred during deposition. Both influx of organic material and processes that occur upon deposition affect the molecular characteristics of the organic material within the sample, thus controlling the generation of petroleum (Hsu et al., 2003). Source rock quality and richness (HI and S_2) have been observed to generally decrease with depth and follow the trends of the Gamma Ray log intensity (Dahl et al., 2004). However, this trend is not clearly visible in the 40ft section of the Woodford Shale from Speake Ranch characterized in this study (Figure 19). Additionally, TOC alone is not an accurate measurement of hydrocarbon generation within the sediment (Dembicki, 2009). Knowing the amount of carbons and hydrogens present, as well as the maturity of the source rock, is required to determine a more accurate quantification of hydrocarbon generation.

Type I kerogen is highly oil prone and possibly gas prone as well, yielding a high HI value and a low OI value (Peters, 1986). Type II kerogen will generate both oil and gas and is characterized by decreasing OI values while initial HI remains nearly constant (McCarthy et al., 2011). Type III kerogen has low HI values and high OI values and is very prone to generate gas and maybe some oil (Peters, 1986). Many of the Upper and Middle Woodford Shale samples plot as Type I and Type II mixed kerogen, reflecting samples that are in the early oil generative window and sourced from a marine environment (Figure 21). Furthermore, the HI values for the Speake Ranch samples range from 25 to 577 mg HC/g TOC along with OI values that range from 3 to 194 mg CO_2 /g TOC.

The HI values of Speake Ranch have a majority of its samples varying between 400–600 mg HC/g TOC. This interval is indicative of oil generative quality from a marine source based on the parameters define by Peters and Cassa (1994). Nearly all the samples of the Upper Woodford and Middle Woodford from the Speake Ranch plot within the 400-600 mg HC/g TOC interval. However, there are outliers present within the section, displaying values below 200 mg HC/g TOC which are often associated with dilution of the TOC by sediment input from a terrigenous source (Figure 19 and Figure 21). The majority of these outliers can be found distributed within the Middle Woodford of this study. Several of Speake Ranch samples in Figure 21 also display a Type III and IV kerogen, which may be a direct effect of organic matter source input and/or thermal maturation. These distributions are primarily found within the Middle Woodford, and at the boundary of the Upper Woodford and Middle Woodford (Figure 19).



Figure 21. Pseudo van Krevelen type kerogen type plot plotting the HI vs. the OI to assess the kerogen type of the Woodford samples from Speake Ranch. 40 of the studied samples indicate a Type I and Type II kerogen. Outlier sample points within the circles are most likely to a result from organic matter source input.

3.5 Thermal Maturity from Rock-Eval

Thermal maturity can be determined from a combination of vitrinite reflectance, T_{max} , S_1 , and S_2 ratios. There are 3 stages of maturation which include diagenesis, catagenesis, and metagenesis. Diagenesis is the thermally immature phase, where the physical, chemical, and biological transformations occur prior to burial or with increasing thermal gradient while still retaining the molecular structure of the compounds (Peters and Cassa, 1994). Catagenesis is the mature and post mature stage comprising the oil window which is the interval for oil and gas production. It also consists of the wet gas zone containing less than 98% of methane, owing to the light hydrocarbons resulting from cracking (Peters and Cassa, 1994). Finally, metagenesis is the post mature stage/phase which comprises the dry gas zone with more than 98% of methane (Peters

and Cassa, 1994). Maturation pathways help decipher the level of thermal stress the samples in this study have experienced.

Based on the T_{max} values, the Woodford Shale from Speake Ranch is predominantly immature with values ranging from 419 to 434° C, averaging at 426° C. Based on the values proposed by Peters (1986) this constitutes an overall immature source rock characteristic (<435° C). Furthermore, the maximum T_{max} value (434° C) indicates that the Speake Ranch samples are within the early oil window. In a 40 ft interval we would not expect any variations in thermal maturity as earlier defined. However, Figure 22 shows slight variations possibly attributed to the mineral composition within the interval. The variations in the T_{max} do not coincide with oscillations in the TOC values, suggesting that any variabilities within the T_{max} value are not only a reflection of organic matter, but also an effect of weathering. It is insufficient to rely on T_{max} for the evaluation of maturity as errors are likely to be associated due to the changing mineralogical effects within the 40ft sequence. The sulfur-carbon bonds in comparison to the carbon-carbon bonds require a lower bond-breaking temperature, which results in lowering of oil generation by $\sim 30^{\circ}$ C (McCarthy et al., 2011). Thus, presence of sulfur within the lithofacies may lower the T_{max} . The characterization of maturity supported by vitrinite reflectance is necessary. The vitrinite reflectance can be calculated using T_{max} derived from Rock-Eval data [%Rc=(0.0180*T_{max}-7.16] (Jarvie, 1991; Wust et al., 2013). These samples result in values ranging from 0.27 to 0.65% Rc with a mean at 0.53% Rc indicating a relatively low maturity. The average value closely coincides with the mean from direct petrographic measurements of vitrinite reflectance (0.60%Ro) provided by Cardott (2017; written communication to Galvis, 2017) in Figure 23. This was taken over 40 measurements at random within a single depth, varying from 0.49-0.72% Ro. However, Galvis (2017) calculated an average over the entirety of Woodford Shale to be at 0.79% Rc making it higher than the average calculated for the interval in this study. The discrepancy may be attributed to his measurements over the 536 samples from Speake Ranch vs. the 37 (Upper Woodford and Middle Woodford) samples analyzed in this research. Additionally, lower thermal maturity values discovered in Speake Ranch may be attributed to tectonic uplift of the Arbuckle Mountains during the Pennsylvanian (Ham et al., 1973; Galvis, 2017). The following sections will utilize biomarker parameters to further determine the conclusion that there are slight thermal maturity variations within Speake Ranch samples.



Figure 22. Depth plot of T_{max} showing variations within the 40ft sequence. There is little to no variation in the distribution of the maturity parameter, which justifies suggesting that thermal alteration is not a factor contributing to observed biomarker variations. Gray intervals indicate soft bed lithofacies, while white intervals indicate hard beds.



Figure 23. Distribution of vitrinite reflectance in South Central Oklahoma. Petrographic vitrinite reflectance of Speake Ranch ranges from immature to early oil window, with a mean of 0.60% Ro (Cardott, 2012; Galvis, 2017).

3.6 Gas Chromatography

As proposed in previous research, biomarker analysis is useful in determining fluctuating levels of anoxia upon deposition and organic matter source (Connock, 2015; Jones, 2017; Galvis, 2017; and Torres, 2020). This research explores the biomarker families used to identify changing organic matter sources and processes occurring within the paleoenvironment. These biomarkers include the n-alkanes, steranes, terpanes, diasteranes, tetracyclic polyprenoids, aryl isoprenoids, alkylnaphthalenes, and alkylphenanthrenes screened on the GC-FID and GC-MS. The analysis of these biomarker families will take into account their relationships with depth, maturity, biodegradation and lithology.

3.6.1 n-Alkane distribution

The n-alkanes are found within the saturate fractions of source rock extracts and crude oils screened by the GC. They are specifically useful as a tool for correlation and organic matter evaluation (Peters et al., 2005). The n-alkanes derive from lipids within living organisms, such as higher-order plants (odd-carbon-numbered n-alkanes, specifically n-C₂₇, n-C₂₉, and n-C₃₀ or algae which are illustrated in the lower carbon number homologs (Eglinton and Hamilton, 1967; Smith, 1968). The Woodford Shale samples from Speake Ranch have experienced one of the earliest levels of biodegradation, which is the preferential removal of the n-alkanes, and isoprenoids (Volkman et al., 1984). Based on several chromatograms, there is a slight odd-even predominance between the $n-C_{15} - n-C_{17}$ (I; See Appendix II for compound structure) which are typically attributed to algal/cyanobacteria origin as shown in Figure 24 (Han et al., 1967; Grice et al., 1996). Higher carbon number alkanes that maximize at n-C₂₉ with an odd-even predominance are known to be indicative of higher plant input (Eglinton et al., 1962; Grice et al., 1996). However, the preferential removal of the n-alkanes in the Speake Ranch samples make it hard to discern any nalkane trend which may lead to misinterpretations. Thus, additional supporting biomarker analysis is essential to further specify the organic matter source.



Figure 24. Gas chromatogram of the saturate fraction from the extract of the sample at a depth of 229.0 ft from the Woodford Shale at Speake Ranch.

3.6.2 Pristane and Phytane

The acyclic isoprenoid hydrocarbons pristane and phytane are determined by GC. In immature sediments, the pristane (Pr; **II** see Appendix II for compound structure) and phytane (Ph; **III**) reflect the oxidation fluctuations of the phytyl side chain of chlorophyll a and b in phototrophic organisms (Brooks et al., 1969; Powell and McKirdy, 1973; Goosens et al., 1984; Grice et al., 1996). Accordingly, differences in pristane/phytane (Pr/Ph) ratio reflect the level of anoxia within

the water column. Connock (2015) classified that the Upper Woodford was deposited in a dysoxicsuboxic environment whilst the Middle Woodford was deposited in a predominantly anoxic environment as indicated by the Pr/Ph values (Table 2). Similarly, the Woodford Shale from Speake Ranch shares the same characteristics as those classified by Connock (2015). Using the pristane/n-C₁₇ vs. the phytane/n-C₁₈ plot, Speake Ranch samples display values below 1 for both parameters indicating low maturity samples deposited under reducing conditions, with an input of mixed organic matter (Figure 25). These data agree with the identified depositional environment framework of the Woodford Shale present in Late Devonian-Early Mississippian as previously reported by Romero and Philp (2012). The outliers of the hard bed intervals with a Pr/Ph values >1 in Figure 25 are possibly a product of biodegradation effects rather than depositional environmental conditions. However, apparent variations in the Pr/Ph within the depth log shown in Figure 26, follow variations in lithology (tracking the relative TOC wt % abundance of soft and hard beds). This suggests that although the Woodford Shale was deposited predominantly in an anoxic environment, there were periodic oxic to sub-oxic disturbances (Figure 26). It is insufficient to determine organic matter source and nature of depositional environment from Pr/Ph alone. Lower Pr/Ph ratios do not necessarily indicate reducing environments but may also be an effect of lithology and maturity (Marynowski et al., 2000).



Figure 25. $Pr/n-C_{17}$ vs. $Ph/n-C_{18}$ illustrating the redox conditions for the Woodford Shale from the Speake Ranch. All the studied source rock samples illustrate a Type II and Type II/III kerogen of marine and marine-terrigenous mixed organic matter deposited within a reducing environment. The red symbols indicate soft bed lithofacies, while green symbols indicate hard bed lithofacies.

It is expected that lower Pr/Ph values would coincide with high TOC wt% and the data from the Speake Ranch samples confirm that a positive correlation is present (Figure 26). Significant elevated shifts of Pr/Ph coincide with intervals of lower TOC wt%, owing to an oxic environment that caused poor preservation of organic matter. For more accurate evaluation of the relationship between organic carbon preservation and anoxia, more redox evaluator parameters were also utilized in this research.
Pr/Ph	0.76	0.75	0.66	1.05	0.64	0.80	0.94	0.92	1.10	0.78	0.71	0.63		'	0.73	0.78	1.55	0.84	1.15	0.85	0.76	1.65	0.69	0.93	0.77	0.75	,	1.28	0.75	1.33	2.56	0.86	0.77	2.13	0.77	0.79	0.66	0.72
S1/TOC	15.50	15.09	15.58	11.16	17.11	14.95	20.36	19.37	18.80	16.85	21.52	6.19	13.53	10.99	14.27	13.30	18.18	11.79	11.40	13.31	14.97	8.72	9.32	9.52	5.37	9.71	10.92	12.87	5.06	11.7.1	14.56	12.27	11.77	11.71	11.88	13.57	9.63	12.92
IO	4.92	5.59	3.72	25.25	3.36	4.07	3.72	5.44	4.30	74.72	10.09	5.69	205.50	93.92	3.75	9.68	71.51	7.35	6.50	6.36	2.93	12.20	3.11	4.58	3.37	3.63	183.19	18.67	4.26	10.83	171.39	3.06	22.72	24.34	4.29	3.36	4.11	1036
IH	430.67	580.73	496.28	445.05	564.64	532.30	530.01	616.00	596.58	90.45	538.43	507.98	14.56	136.14	489.04	484.59	208.72	577.94	546.75	514.38	541.87	499.52	543.02	590.53	472.08	526.37	40.71	520.25	527.41	532.76	83.46	569.64	483.59	487.17	528.57	577.49	591.58	490.72
Tmax (°C)	419	426	419	429	432	428	430	432	431	425	428	422	413	426	422	425	426	425	429	424	423	426	422	427	421	429	434	433	432	429	429	429	428	426	428	434	430	428
S3 (mg CO2/g)	0.59	0.29	0.48	0.51	0.31	0.46	0.34	0.68	0.49	1.33	1.09	0.62	1.27	5.25	0.39	0.27	1.23	1.14	0.42	0.77	0.36	0.51	0.33	0.6	0.34	0.37	2.07	0.59	0.46	0.38	1.15	0.34	2.34	0.74	0.48	0.33	0.4	115
S2 (mg HC/g)	51.68	30.14	64.02	8.99	52.06	60.15	48.39	77	68.01	1.61	58.15	55.37	0.09	7.61	50.86	13.52	3.59	89.58	35.32	62.24	66.65	20.88	57.56	77.36	47.68	53.69	0.46	16.44	56.96	18.7	0.56	63.23	49.81	14.81	59.2	56.71	57.62	54 47
S1 (mg HC/g)	1.55	0.5	1.75	0.24	1.08	1.33	1.12	1.82	1.95	0.09	1.39	1.19	0.06	0.3	0.99	0.26	0.15	2.32	0.86	1.38	1.45	0.76	1.41	1.87	11.11	1.38	0.07	0.68	1.82	0.66	0.13	2.26	1.54	0.52	1.25	1.53	1.47	1 72
TOC (wt%)	12	5.19	12.9	2.02	9.22	11.3	9.13	12.5	11.4	1.78	10.8	10.9	0.618	5.59	10.4	2.79	1.72	15.5	6.46	12.1	12.3	4.18	10.6	13.1	10.1	10.2	1.13	3.16	10.8	3.51	0.671	1.11	10.3	3.04	11.2	9.82	9.74	
Lithology	Soft	Hard	Soft	Hard	Soft	Soft	Soft	Soft	Soft	Hard	Soft	Soft	Hard	Soft	Soft	Hard	Hard	Soft	Hard	Soft	Soft	Hard	Soft	Soft	Soft	Soft	Hard	Hard	Soft	Hard	Hard	Soft	Soft	Hard	Soft	Soft	Hard	Soft
Depth (ft.)	190.5	191	191.5	192	193.5	194.5	195.5	196.5	197.5	198	199.5	200.5	202	203.5	204.5	205	207	208.5	209	210.5	211.5	212	213.5	214.5	215.5	217.5	218	219	221.5	222	223	224.5	225.5	226	227.5	228.5	229	230.5
Sample	SR-190b	SR-191a	SR-191b	SR-192a	SR-193b	SR-194b	SR-195b	SR-196b	SR-197b	SR-198a	SR-199b	SR-200b	SR-202a	SR-203b	SR-204b	SR-205a	SR-207a	SR-208b	SR-209a	SR-210b	SR-211b	SR-212a	SR-213b	SR-214b	SR-215b	SR-217b	SR-218a	SR-219a	SR-221b	SR-222a	SR-223a	SR-224b	SR-225b	SR-226a	SR-227b	SR-228b	SR-229a	SR-230h

Table 2. Bulk geochemical data of the Woodford Shale from Speake Ranch, including S1, S2, S3, HI, OI, Tmax, S1/TOC, and Pr/Ph Pr/Ph



Figure 26. Integration of TOC depth plot and Pr/Ph depth plot generated in this study. Pr/Ph parallels the oscillation of TOC which coincides with the preservation of abundant organic matter within anoxic intervals. Gray intervals indicate soft bed layers, while the white intervals indicate hard bed layers. The red line is the Upper Woodford and Middle Woodford boundary.

More distinctions can be determined between the soft vs. hard of the Pr/Ph values approaching the Upper Woodford, where oxic periods are found to be more frequent (Figure 26). The Pr/Ph values have shown to be consistently anoxic until 205ft, and depths beyond this interval show that the sequence experienced episodes of oxic to sub-oxic disturbances (Figure 26). As TOC-rich intervals have high clay content, it would be expected that these intervals would experience consistently high levels of anoxia (low Pr/Ph). However, the correlation of Pr/Ph vs. C₂₇ diasterane/steranes (formulas can be found in Appendix III) show that although clay content stay at nearly consistent values, Pr/Ph values are seen to vary (Figure 27). An elaborated discussion regarding the relationship between clay content, anoxia, and organic richness will follow in the

next sections. Peters et al. (2005) advised that the Pr/Ph in rock and oil samples within the oilgenerative window should be used with caution as a correlative tool for depositional redox conditions. As samples of Speake Ranch values fall between 0.8-3, although still reflecting depositional redox conditions, may be influenced by variable source inputs, differing rates of early generation, and variations of thermal maturity (Peters et al., 2005). It is hypothesized in this research that optimal preservation of organic matter results from multiple factors other than anoxia. The highly oscillating TOC must be a result of combined depositional effects, further elaborated in the following sections.



Figure 27. Pr/Ph vs. C₂₇ diasteranes/steranes (Appendix III; modified from Moldowan et al. 1994). The arrow shows the expected direction of thermal maturity effect, which is not considered for these specific samples since they are all of relatively low-maturity rocks. The diagram shows that Pr/Ph varies as clay content stays nearly consistent.

3.6.3 Steranes

Steranes are derived from the precursor steroid molecules (sterols) which contain 27, 28, 29, and 30 carbon atoms and are present in both higher plants and algae (Waples and Machihara, 1990). Steranes can have various conformations, in which the stereochemistry can change according to the level of thermal stress experienced. During diagenesis, these precursor molecules

are converted into a more stable saturate hydrocarbon biomarkers (Grantham, and Wakefield, 1987; Waples and Machihara, 1990). The thermal stability of steranes is lower than that of hopanes in both free and bound fractions due to their differing chemical structures and incorporation into the kerogen (Wu and Geng, 2016). The sterane family has proved to be reliable in the definition of organic matter source, lithology, maturation, biodegradation, and depositional environment conditions used in this research. Sterane distributions are monitored by the GC-MS using the characteristic ion at m/z 217(Figure 28; Table 3).

Peak No.	Compound	Peak No.	Compound
1	Diapregnane	16	5α14β17β 20R Ergostane + 24-Ethyl- 13α17β-Diacholestane 20R (C ₂₉)
2	Homopregnane	17	5α14β17β Ergostane 20S
3	$13\beta17\alpha$ Diacholestane 20S (C ₂₇)	18	5α14α17α Ergostane 20R
4	13β17α Diacholestane 20R (C ₂₇)	19	5α14α17α Stigmastane 20S
5	$13\alpha 17\beta$ Diacholestane 20S (C ₂₇)	20	5α14β17β Stigmastane 20R
6	$13\alpha 17\beta$ Diacholestane 20R (C ₂₇)	21	5α14β17β Stigmastane 20S
7	24-Methyl-13 β 17 α Diacholestane 20S (C ₂₈)	22	5α14α17α Stigmastane 20R
8	24-Methyl-13 β 17 α Diacholestane 20R (C ₂₈)	23	C ₃₀ 5α14α17α 24-n-Propylcholestane 20S
9	$5\alpha 14\alpha 17\alpha$ Cholestane 20S + 24-Methyl- 13 α ,17 β -Diacholestane 20S (C ₂₈)	24	C_{30} 5a14 β 17 β 24-n-Propylcholestane 20R
10	5α4β17β 20R Cholestane +24-Ethyl- 13β,17α-Diacholestane 20S (C ₂₉)	25	C_{30} 5 α 14 β 17 β 24-n-Propylcholestane 20S
11	$\begin{array}{c} 5\alpha 14\beta 17\beta \ 20S \ Cholestane + 24-Methyl-\\ 13a, 17\beta \ Diacholestane \ 20R \ (C_{28}) \end{array}$	26	C_{30} 5 α 14 α 17 α 24-n-Propylcholestane 20R
12	5α14α17α Cholestane 20R		
13	24-Ethyl-13β,17α-Diacholestane 20R (C ₂₉)		
14	24-Ethyl-13α,17β-Diacholestane 20S (C ₂₉)		
15	5a14a17a Ergostane 20S		

Table 3. Peak identification of the steranes of sample SR-223a at depth 223ft within the Upper Woodford of Speake Ranch.



Figure 28. Distribution of the steranes and diasteranes monitored at m/z 217. Molecular structure of sterane from Pratt et al. (1992; **IV**, **V**, **VI**; **Appendix II**).

3.6.3.1 Organic Matter Source

Regular steranes are useful tools to evaluate and identify the photosynthetic biota (organisms/plants) that could be derived from precursor sources present in marine, terrigenous, or lacustrine environments (Waples and Machihara, 1990). The most common regular steranes used to determine organic matter input include the C_{27} cholestanes (IV) that are abundant within plankton and marine invertebrates, C_{28} ergostanes (V) which are derived from organisms present in lacustrine environments, and the C_{29} sitostanes (VI) derived from land plants. The resulting C_{27} , C₂₈, and C₂₉ steranes acquired from the source rock extracts of Speake Ranch were used to construct a sterane ternary diagram [i.e. % $C_{27} = C_{27} \alpha \alpha \alpha 20R/(C_{27} \alpha \alpha \alpha 20R + C_{28} \alpha \alpha \alpha 20R + C_{29})$ ααα 20R); Figure 30]. The steranes from the Speake Ranch cluster very closely to each other and suggest that the samples were deposited in an open marine environment (abundance of C₂₇ relative to C₂₉) with a major source contribution from phytoplankton (Figure 29(a.); modified from Huang and Meinschein, 1979). The modified plot by Moldowan et al. (1985) in Figure 29(b.) illustrate Speake Ranch values plotting below the defined marine shale parameter. As C₂₈ steranes are shown to be present in low abundances within these samples, weathering may have further reduced these abundances. Overall, there are no significant variations found for the distribution of the steranes within the sterane ternary plot, suggesting consistent and invariant influx of organic matter input. However, with higher resolution of biomarker analysis, variations of source input may uncover a principal marine origin with an incorporation of terrigenous input into the system, which is why utilizing the $C_{27} \alpha \alpha \alpha 20 R/C_{29} \alpha \alpha \alpha 20 R$ steranes for the ratio will reveal the relative proportion of marine vs. terrigenous/algal input (Figure 30).



Figure 29(a). Ternary plot of the C_{27} - C_{28} - C_{29} steranes to indicate organic matter source for the Woodford Shale. Majority of the samples are seen clustering in an abundance of C_{27} steranes indicating an open marine environment (sterane ternary plot recreated from Huang and Meinschein, 1979); (b). distribution determined by Moldowan et al. (1985) for source rocks of varying ages.





C ₂₇ diasterane/sterane	0.32	0.30	0.30	0.30	0.31	0.30	0.31		0.32	0.28	0.31	0.30			0.30	0.29	0.31	0.31	0.29	0.32	0.30	0.29	0.33	0.31	0.31	0.33		0.30	0.34	0.32	0.29	0.33	0.34	0.29	0.34	0.35	0.37	0.34
C ₂₈ BNH/Norhopane	2.18	1.97	1.93	1.71	2.09	2.05	2.07		2.08	1.65	2.07	2.07			2.03	1.72	1.34	1.94	1.67	2.02	1.93	1.62	1.94	2.02	2.18	1.92		1.59	2.17	1.54	1.47	2.28	2.25	1.47	2.13	2.11	2.21	2.23
C ₂₈ BNH/C ₃₀ Hopane	0.50	0.53	0.52	0.29	0.39	0.34	0.42		0.36	0.25	0.38	0.42			0.37	0.27	0.23	0.38	0.26	0.35	0.38	0.27	0.34	0.35	0.39	0.39		0.24	0.35	0.26	0.23	0.39	0.39	0.24	0.38	0.36	0.37	0.37
C ₁₉ /C ₂₃ TT	0.09	0.10	0.10	0.08	0.09	0.08	0.09		0.08	0.06	0.10	0.10			0.09	0.07	0.06	0.09	0.06	0.08	0.09	0.06	0.08	0.08	0.07	0.09		0.04	0.06	0.05	0.05	0.08	0.07	0.05	0.08	0.07	0.07	0.09
C _{23 TT} /C ₂₄ Tet	2.95	2.91	2.95	3.64	3.12	3.13	2.69	,	3.07	2.91	3.13	2.98			3.12	2.88	3.58	3.06	2.82	2.80	3.11	2.55	2.90	2.77	3.04	3.03	,	2.63	2.40	2.27	2.91	3.00	2.80	3.03	3.47	2.84	2.97	3.25
C ₂₈ /C ₂₉ Sterane	0.39	0.39	0.39	0.40	0.40	0.41	0.39	,	0.40	0.38	0.39	0.42			0.40	0.35	0.39	0.38	0.37	0.39	0.38	0.37	0.37	0.33	0.38	0.41		1.27	0.42	0.40	0.36	0.42	0.42	0.36	0.44	0.43	0.42	0.42
C ₂₇ /C ₂₉ Sterane	1.14	1.22	1.22	1.00	1.03	1.04	1.06	,	1.05	0.91	1.08	1.12			1.11	0.90	1.01	1.09	0.95	1.06	1.08	0.92	1.06	1.03	1.11	1.12	,	1.17	1.00	0.93	0.93	1.16	1.13	0.94	1.19	1.08	1.05	1.20
C ₃₀ sterane index	2.48		3.26	2.48	1.89	2.45	2.30	,	2.49	2.02	1.98	2.20			2.17	2.34	1.97	2.04	2.34	2.26	2.51	2.01	2.16	2.43	3.24	2.88		1.74	2.53	3.04	2.41	2.42	2.76	2.50	2.49	2.56	2.52	2.49
Pr/Ph	0.76	0.75	0.66	1.05	0.64	0.80	0.94	0.92	1.10	0.78	0.71	0.63	,		0.73	0.78	1.55	0.84	1.15	0.85	0.76	1.65	0.69	0.93	0.77	0.75	,	1.28	0.75	1.33	2.56	0.86	0.77	2.13	0.77	0.79	0.66	0.72
TOC (wt%)	12.00	5.19	12.90	2.02	9.22	11.30	9.13	12.50	11.40	1.78	10.80	10.90	0.62	5.59	10.40	2.79	1.72	15.50	6.46	12.10	12.30	4.18	10.60	13.10	10.10	10.20	1.13	3.16	10.80	3.51	0.67	11.10	10.30	3.04	11.20	9.82	9.74	11.10
Lithology	Soft	Hard	Soft	Hard	Soft	Soft	Soft	Soft	Soft	Hard	Soft	Soft	Hard	Soft	Soft	Hard	Hard	Soft	Hard	Soft	Soft	Hard	Soft	Soft	Soft	Soft	Hard	Hard	Soft	Hard	Hard	Soft	Soft	Hard	Soft	Soft	Hard	Soft
Depth (ft.)	190.50	191.00	191.50	192.00	193.50	194.50	195.50	196.50	197.50	198.00	199.50	200.50	202.00	203.50	204.50	205.00	207.00	208.50	209.00	210.50	211.50	212.00	213.50	214.50	215.50	217.50	218.00	219.00	221.50	222.00	223.00	224.50	225.50	226.00	227.50	228.50	229.00	230.50
Sample	SR-190b	SR-191a	SR-191b	SR-192a	SR-193b	SR-194b	SR-195b	SR-196b	SR-197b	SR-198a	SR-199b	SR-200b	SR-202a	SR-203b	SR-204b	SR-205a	SR-207a	SR-208b	SR-209a	SR-210b	SR-211b	SR-212a	SR-213b	SR-214b	SR-215b	SR-217b	SR-218a	SR-219a	SR-221b	SR-222a	SR-223a	SR-224b	SR-225b	SR-226a	SR-227b	SR-228b	SR-229a	SR-230b

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There is a clear distinction as shown in C_{27}/C_{29} sterane ratio $[C_{27}/C_{29}=C_{27} \alpha \alpha \alpha 20 R/C_{29} \alpha \alpha \alpha$ 20R sterane] depth plot of a trend between the soft (gray interval) and the hard (white interval) beds (Figure 30). The C_{27}/C_{29} sterane ratio is often used as a marine organic matter indication when values exceed one (Grantham and Wakefield, 1988). Apart from association with abundant organic matter with the soft lithofacies, the softer beds have also shown to be deposited in a marine environment with an abundance of C₂₇ steranes (Figure 31). Conversely, the harder beds generally have C_{27}/C_{29} sterane at values less than 1 (C_{29} steranes abundant) throughout the studied interval, illustrating a relatively higher terrigenous input and a more oxic depositional environment. The ratio that is observed to illustrate values less than 1 are due to a relative increase in the C_{29} sterane, and subsequently a decrease in C₂₇ sterane from the measurements of their relative concentrations (Figure 32; C₂₇/total steranes to C₂₉/total steranes). It is suggested that as the Late Devonian was a time of expanding terrigenous vegetation and higher nutrient input, the resulting fluctuation reflects weathering, widespread anoxia, and eutrophication. According to the sequence stratigraphy more terrigenous input should be naturally observed within the Upper Woodford due to its higher proportion of silica to clay content (Torres, 2020). Therefore, the decrease in clay content in the Upper Woodford signifies an increase of terrigenous source input that dilutes the marine clays, thus reducing the availability of acid sites. The significance of acid sites is that it provides a platform for catalyzing rearrangement of the sterol backbone (van Kaam-Peters et al., 1998). Another possible explanation for the frequent fluctuations between marine and terrigenous input is that as the shoreline retreated, the depositional environment may have become more restricted, which causes CaCO₃ to supersaturate the water column. This resulted in the stimulation of algal blooms from marine ocean water anoxia caused by the Devonian expansion of terrigenous land plants and extensive emission of CO₂ (Philp and DeGarmo, 2020). Furthermore, when the

 C_{27}/C_{29} sterane ratio is combined with Pr/Ph ratio, it would appear to suggest that the marine dominated soft beds with high TOC values were deposited in a primarily anoxic condition (Figure 31). Alternatively, the lower TOC intervals of the hard beds were deposited under a more oxidized environment, with a higher input of the C₂₉ steranes (Figure 31 highlighted intervals). Using the C_{27}/C_{29} ratio depth plot against Pr/Ph defined from Hossain et al. (2009) can help determine types of organic matter input and the depositional environmental condition as shown in Figure 33. Observed trends in Figure 31, sterane index along with relative abundances in Figure 32, and plot of C₂₇/C₂₉ sterane against Pr/Ph (Figure 33) all point toward the same conclusion of the marine/algal source input in an anoxic depositional environment of the Woodford Shale. It is apparent that the relatively lower TOC values are associated with elevated C₂₉ sterane abundances within the Upper Woodford and Middle Woodford. However, caution is needed for sediments indicating a terrigenous source environment (showing an abundance of C₂₉) as it is not to confuse such an input from cyanobacteria (green-blue algae) (Boon et al., 1983; Brassell et al., 1988; Telnaes and Cooper, 1991) or chlorophyta (green algae) (Fowler and Douglass, 1987; Waples and Machihara, 1990) which have also been proposed to contribute C₂₉ steranes into the sediments. Grantham and Wakefield (1988) have also suggested that C₂₉ steranes could result from a contribution of brown algae. Table 4 displays all the values of biomarker parameters evaluating source organic matter input. Intervals without values are due to data limitations from the lack of maltene yield from extraction.



Figure 31. Geochemical logs correlating TOC fluctuations, with the associated marine vs. terrigenous source input (C_{27}/C_{29} sterane), and the role of anoxia (Pr/Ph) in its distribution. Highlighted zones indicate that low preservations of organic matter are associated with high C_{29} input and oxidizing environmental deposition conditions. Gray zones are the soft bed intervals, and the white zones are the hard bed intervals.



Figure 32. The plot on the right display the relative abundances of C_{27} and C_{29} [C_{27} 14 α ,17 α -+14 β ,17 β -Steranes(20S+20R)/total steranes; C_{29} 14 α ,17 α -+14 β ,17 β -Steranes (20S+20R)/total steranes; Circles: C_{29} /total steranes]. Decrease in C_{27}/C_{29} steranes coincide with a relative increase in C_{29} steranes.



Figure 33. C_{27}/C_{29} steranes vs. Pr/Ph plot characterizing the Woodford Shale from Speake Ranch as algal rich organic matter deposited in an anoxic environment. (modified from Hossain et al., 2009). Samples plotting below 0.55 indicate terrigenous input into the anoxic environment.

The abundance of the C_{30} steranes (VII) in sediments is a strong indication of marine organic matter input (Peters and Moldowan, 1991). The C_{30} sterane index (Sterane Index: $C_{30}/(C_{27} C_{30}$) steranes) has widely been used to discern the amount of marine biotic input (Peters et al., 1986 and Peters et al., 2005). This ratio is suggested to decrease with regression of sea level, as there are less input of marine-sourced organic matter and more of terrigenous-sourced organic matter. The Speake Ranch samples have values ranging from 2.3 - 5.6% averaging at 4.1% (Figure 30 and 32; Table 4). Moldowan et al. (1985), Peters et al. (2005) and Hays et al. (2012) determined that C_{30} sterane index values >4% are indicative of marine pelagophyte algae, and thus indicating a marine depositional environment. This places majority of the values within the 40ft interval of the Speake Ranch well above that parameter which supports the characterization of marine environment origin. Values plotting below this 4% parameter are most likely due to the elevated C_{29} sterane (terrigenous input) relative to the C_{27} sterane values found within the harder lithofacies (Figure 32). Although, as mentioned in the earlier section, the C₂₉ sterane could also be synthesized by green algae (Schwark and Empt, 2006). The C₃₀ sterane index values illustrate an overall marine source input as shown in Figure 32. The C_{30} sterane index is seen to increase approaching the Upper Woodford, although ironically it has been proposed that the terrigenous input increases prior to the Frasnian-Famennian extinction boundary suggested by Torres (2020) shown in Figure 32. The Frasnian-Famennian (F-F) extinction boundary marks the loss of a wide marine biota, which Torres (2020) has defined to be at 194 to 210 ft. This interpretation was defined by abrupt changes in biomarker parameters. However, a specified interval was unable to be determined by Galvis (2017) and Torres (2020) due to a lack of high resolution biomarker analysis.

A narrow interval in which the F-F boundary is proposed to have occurred will be discussed in Chapter V. It is expected that terrigenous input should overwhelm the Upper Woodford as the sediments within this endmember were deposited during a Highstand System Tract. As shoreline retreated, erosional events increase with progradation, which introduces terrigenous material into the basin and disturbed the marine productivity, which may contribute to the elevated C_{29} steranes input (Figure 18).

3.6.3.2 Thermal maturity indicators

The T_{max} and R_c measurements indicated that the Woodford Shale from the Speake Ranch is largely immature throughout the 40ft section. To validate and support the accuracy of the Tmax and R_c values, several biomarker maturity parameters were also utilized. As discussed earlier, the sterane and hopanes structures will be converted into more stable conformations with increasing levels of thermal stress (Grantham, and Wakefield, 1987; Waples and Machihara, 1990). Reliable assessment of maturity can be determined by utilizing the epimerization of the H atom at C-20 in the S configuration of C_{29} 5 α , 14 α , 17 α -sterane (VI), specifically for immature to early mature samples (Mackenzie et al., 1980). The 20R configuration in steroid precursors within living organisms, are converted into a mixture of the 20R and 20S configurations with burial and maturation (Requejo et al., 1994). With increasing maturity, the proportion of the 20S increases at the expense of the 20R epimer. Thus, changes in the 20S/(20S+20R) ratio for any of the C₂₇-C₂₈-C₂₉ steranes can be used to evaluate relative maturity levels. The C₂₉ sterane isomers and epimers are generally used for this purpose since all four isomers and epimers are resolved chromatographically (Waples and Machihara, 1990). Seifert and Moldowan (1986) proposed that the isomerization at C-14 and C-17 in the 20S and 20R of the C₂₉ steranes provide for a reliable maturity parameter, $\beta\beta/(\beta\beta+\alpha\alpha)$, which is independent of organic matter source input. An increase

in the $\alpha\beta\beta$ steranes relative to the less stable $\alpha\alpha\alpha$ steranes reaches equilibrium slower than the 20S and 20R, thus making it effective at higher thermal maturity. $C_{29} \beta\beta/(\beta\beta+\alpha\alpha)$ [$C_{29}(14\beta,17\beta-\alpha)$] Stigmastane $(20S+20R)/[14\beta,17\beta-+14\alpha,17\alpha$ -Stigmastane (20S+20R))] is a useful and reliable maturity indicator to be incorporated with C₂₉ 20S/(20S+20R) [C₂₉ (14 α , 17 α -Stigmastane (20S)]/[14 α , 17 α -Stigmastane (20S+20R))]. However, it is also important to note that elevated values of $\beta\beta$ steranes may be a product of reaction to sulfur (ten Haven et al., 1986). Schmid (1986) indicated that heating the $\alpha\alpha\alpha$ cholestane with sulfur gave little isomerization at C-20, however, a substantial amount formed of the $\alpha\beta\beta$ 20R and 20S. The plot of C₂₉ 20S/(20S+20R) vs. C₂₉ $\beta\beta/(\beta\beta+\alpha\alpha)$ in Figure 34 defined by Peters (1999) have been used as an indicator of organic maturity. The Woodford Shale samples from the Speake Ranch show higher maturity values than indicated for immature characteristics defined by Seifert and Moldowan (1980) in Figure 34. This could possibly reflect migrated hydrocarbons (Torres, 2020) and/or the presence of sulfur upon deposition. The outlier may be due to partial sterane degradation that resulted in increase of the 20S, by selective removal of the 20R epimer by bacteria. Nevertheless, the clustering of the samples in Figure 34 confirms the similar maturity values for all the samples in this study and indicate that maturity is not playing any significant role in variations of any of the biomarker distributions.



Figure 34. Maturity plot of the 20S/(20S+20R) vs. $C_{29}\beta\beta/(\beta\beta+\alpha\alpha)$ isomers. This plot illustrates clustering of the Woodford Shale samples from Speake Ranch due to their similar levels of maturity. Upper Woodford and Middle Woodford may indicate samples of migrated HC (modified from Peters et al., 1999). Blue circle symbols indicate Middle Woodford and, red triangle symbols indicate Upper Woodford samples. Outlier may be due to partial sterane biodegradation that resulted in increase of the 20S, by selective removal of the 20R epimer by bacteria.

3.6.3.3 Lithology

It was earlier noted that organic matter rich intervals coincide with clay-rich intervals. Therefore, it is expected for those clay-rich intervals to be deposited in a more anoxic environment. The diasteranes have been found to indicate acidic (clay or water chemistry) and oxic conditions during diagenesis (Rubinstein et al. 1975; Sieskind et al., 1979; Moldowan et al., 1985; van Kaam-Peters et al., 1998; Brincat and Abbott, 2001). Acidic sites on clays catalyze sterane conversion to diasteranes during diagenesis (Moldowan et al., 1985; Grantham and Wakefield, 1988).Therefore, diasteranes serve as an indicator of the lithology and oxicity. Moldowan et al. (1985) found that samples from Toarcian shale from southwest Germany, have Pr/Ph values that increase with C₂₇ diasteranes/(diasteranes+steranes) (**VIII**; C₂₇diasterane/sterane). In Speake Ranch, Pr/Ph values significantly vary as C₂₇ diasterane/sterane values stay nearly consistent shown in Figure 27 (formula can be found in Appendix III). However, the Speake Ranch samples in Figure 35 reveal

that approaching the Upper Woodford higher values of clay content determined by C_{27} diasterane/sterane coincide with relatively higher TOC and anoxic environmental condition. Additionally, lower relative clay content indicated by lower C_{27} diasterane/sterane coincides with more oxic conditions starting from 204.5ft within the section. This shows an improved resolution of the relationship between clay abundance and anoxic deposition than the parameter shown in Figure 27.

Mello et al. (1988) noted that the diasterane/sterane ratio using the C_{27} diasteranes and steranes [(C_{27} (20S+20R $\beta\alpha$ - and $\alpha\beta$ -Diacholestane)/(20S+20R 5 α ,14 α ,17 α -Cholestane and $20S+20R 5\alpha, 14\beta, 17\beta$ -Cholestane))] is another correlative tool to distinguish carbonates from siliclastic rocks, whereby a lower diasterane/sterane ratio indicates clay-poor carbonate source rocks (Figure 36). Furthermore, high diasterane concentrations relative to regular steranes are thought to be typical of clay rich source rocks, but they have also been observed in extracts from organic-lean carbonates (Moldowan et al., 1991). van Kaam-Peters et al. (1998) found that the diasterane/sterane ratio does not always correlate directly with clay content as it may be affected by level of oxidation as well. The Speake Ranch samples have low values of diasteranes/steranes ratios below 0.5, ranging between 0.3-0.5, which indicates anoxic carbonate source rocks. Carbonate deposition seems to predominate in the Upper Woodford, illustrating that as the shoreline retreated the depositional environment became increasingly restricted causing supersaturation of the water column by CaCO₃. Plots of Pr/Ph vs. diasterane/sterane and C₂₇/C₂₉ sterane ratio vs. C_{27}/C_{29} diasterane ([C_{27} 13 β ,17 α -Diacholestane 20R+13 α ,17 β -Diacholestane 20S]/[24-Ethyl-13 β ,17 α -Diacholestane20R+-13 α ,17 β -Diacholestane 20S]) ratio are shown in Figure 36 and both agree on a similar conclusion. The Pr/Ph (0.2-2.5) vs. diasterane/sterane (0.3-0.5) have values indicating anoxic cabonates and C_{27}/C_{29} sterane ratio (0.8-1.3) vs. C_{27}/C_{29}

diasterane ratio (1.61-3.63) indicate mixed marine source input. The plots correspond to a predominantly anoxic carbonate deposition with a mixed input of marine and terrigenous organic matter. Chapter IV discusses the effects of varying levels of anoxia on organic carbon preservation.



Figure 35. C_{27} diasterane to non-arranged sterane ratio $[C_{27} 13\beta, 17\alpha$ -Diacholestane $(20S+20R)]/[C_{27} 14\alpha, 17\alpha + 14\beta, 17\beta$ -Cholestane (20S+20R)] as a correlative tool to indicate lithology and level of anoxia. High diasterane values over the regular steranes indicate clay rich intervals, which coincide with the soft beds of the Speake Ranch samples deposited in an anoxic environment, determined by the Pr/Ph ratio.



Figure 36 (a). Pr/Ph ratio vs. diasterane/sterane ratio is used to determine the lithology and the extent of anoxia the sediments were deposited (modified from Moldowan et al., 1994) (b). C_{27}/C_{29} sterane ratio vs. C_{27}/C_{29} diasterane ratio to further support organic matter source influx of predominantly marine with mixed marine (modified from Ghori, 2002). Ratio description are included in the Appendix.

3.6.4 Terpanes

Terpanes are biomarkers derived from bacterial (prokaryotic) membrane lipids (Ourisson et al., 1982; Peters et al., 2005). The dominant bacterial family of terpanes are the hopanoids. The two families that the terpanes are subdivided into are based on the number of rings, where those with 27 to 35 carbons belong to the pentacyclics, often known as the "hopanoid" and "nonhopanoids" (hopanes; IX), and the tricyclics and tetracyclic terpanes have 21 to 35 carbon atoms (Waples and Machihara, 1990). The dominant molecular structure of the triterpanes contains 5-rings, where the E-ring contains 5 carbon atoms. Similarly, gammacerane contains the same number of rings but has 6 carbon atoms within the E-ring. Tricylic terpanes, hopanes and gammacerane are useful for indicating certain depositional environments. A typical terpane distribution is monitored on the GC-MS at m/z 191 as shown in Figure 37 with peak identifications in Table 5.



Figure 37. GC-MS chromatogram displaying the distribution of the tricyclic terpanes and hopanes screened at m/z 191. Peaks nomenclature are displayed in Table 5.

Peak	Compound	Peak	Compound
No.		No.	
1	C ₂₀ Tricyclic terpane	19	C ₃₀ Hopane
2	C ₂₁ Tricyclic terpane	20	C ₃₀ Tricyclic terpane
3	C ₂₂ Tricyclic terpane	21	C ₃₁ Tricyclic terpane
4	C ₂₃ Tricyclic terpane	22	Moretane
5	C ₂₄ Tricyclic terpane	23	C ₃₁ Hopane 22S
6	C ₂₅ Tricyclic terpane	24	C ₃₁ Hopane 22R
7	C ₂₄ Tetracyclic terpane	25	Gammacerane
8	C ₂₆ Tricyclic terpane 22S+22R	26	C ₃₂ Hopane 22S
9	C ₂₈ Tricyclic terpane 22S	27	C ₃₂ Hopane 22R
10	C ₂₈ Tricyclic terpane 22R	28	C ₃₃ Hopane 22S
11	C ₂₉ Tricyclic terpane 22S	29	C ₃₃ Hopane 22R
12	C ₂₉ Tricyclic terpane 22R	30	C ₃₄ Hopane 22S
13	Ts	31	C ₃₄ Hopane 22R
14	Tm	32	C ₃₅ Hopane 22S
15	C ₂₈ Bisnorhopane	33	C ₃₅ Hopane 22R
16	C ₂₉ Ts	34	
17	C ₃₀ Diahopane (C ₃₀ D)		
18	C_{29} 17 β 21 α Moretane		
	(Normoretane)		

Table 5. Peak identification of the terpanes at m/z 191 fragmentogram.

3.6.4.1 Organic Matter Source

Tricyclic terpanes (X) have been found to derive from precursors produced by algae and have been known to occur in small abundances (or absent) in oils and extracts from terrigenous sources (Seifert and Moldowan, 1980; Philp et al., 1992; Peters et al., 2005). Moreover, an abundance of tricyclic terpanes can often be found in immature sediments where the short-chain member of the tricyclic terpanes are believed to be related to bacterial input (Aquino Neto et al., 1986; Chicarelli et al., 1983). Aquino Neto et al. (1992), Simoneit et al. (1993), and Araujo et al. (2018) believe that the source of tricyclic terpanes is associated with samples rich in *Tasmanites* algae, which is highly associated with the Woodford shale from Speake Ranch. Tasmanites play a significant role in the production of TOC and biogenic silica, as they contribute to the mechanics of "frackable" intervals for the unconventional play of the Woodford Shale (Slatt, 2013). This further justifies the algal source for the Speake Ranch Woodford samples rather than higher vascular plant input that increased the relative C₂₉ sterane abundance. Araujo et al. (2018) suggested that C_{23} tricyclic terpanes (C_{23} TT) are often dominant in marine sourced oils while C_{24} tetracyclic (C_{24} Tet) and C_{20} tricyclic (C_{20} TT) terpane compounds are found to be more abundant in oils of terrigenous origin. The values of C_{23} TT/ C_{24} Tet from the Woodford samples are consistently above one, justifying the predominance of marine input. It is also observed that lowering of the C₂₃ TT/C₂₄ Tet coincides with the oxic-suboxic interval indicated by Pr/Ph and C₂₇ diasterane/sterane ratios (Figure 38). However, approaching the Upper Woodford, values can be seen to slightly decrease, most likely as a consequence of the Devonian terrigenous land plant bloom and sea level fluctuation marking the Frasnian-Famennian boundary (Figure 38). Philp et al. (1992) have suggested that tricyclic terpanes were found in oils from a saline lacustrine environment in China, which reflects the function of tricyclic terpanes as salinity parameter. In addition to a source input indicator, C_{24} tetracyclic terpanes have been found to occur in highly saline depositional environments within evaporite-carbonate sequences (Connan et al., 1987). The preservation of organic matter in highly saline evaporitic environments is probably due to oxygen depleted waters accompanied by increased salinity and water stratification (Sammy, 1985). However, this does not align to the mudstone lithofacies nature of the Woodford shale, and it is unlikely that the abundance of C_{24} tetracyclic terpane is associated with carbonate evaporitic environments.



Figure 38. Depth logs of C_{23} TT/ C_{24} Tet, C_{35}/C_{34} homohopane, C_{27} diasterane/sterane and Pr/Ph to correlate origin of organic source matter using terpane parameters and its correlation to anoxia. Highlighted interval show increase in C_{23} TT/ C_{24} Tet of marine indication along with low values of C_{35}/C_{34} Homohopane, C_{27} diasterane/sterane and Pr/Ph parameters reflecting anoxic marine sediments.

3.6.4.2 Depositional Environments

The C₃₅ homohopane index [HHI = $C_{35}/\Sigma(C_{31}-C_{35})$ (22S +22R)] is a parameter to evaluate redox conditions especially in marine sediments during diagenesis (Moldowan et al., 1991). High HHI values which indicate relative dominance of C₃₅HH is referred to marine source deposited under anoxic conditions. Low HHI is often associated to prevailing sub-oxic to oxic environment, where preservation of organic matter is low. The Woodford Shale samples from the Speake Ranch have HHI values varying from 5-9% (Table 6). However, the utilization of C_{35}/C_{34} HH [C_{35}/C_{34} $\alpha\beta(22S+22R)$ HH], in addition to HHI, can further discern and confirm anoxic conditions. The C_{35}/C_{34} HH has been widely used as a redox indicator as well, due to its sensitivity to oxic conditions, as the C₃₅ HH values are elevated in reducing conditions (Peters et al., 2005). The side chain of the C₃₅ HH rapidly degrades when oxidized and therefore will make C₃₄ HH relatively higher in abundance (Tulipani et al., 2013). Higher C₃₅ homohopane values relative to C₃₁-C₃₅ homohopanes are often associated to marine carbonates or evaporites (Connan et al., 1987; ten Haven et al., 1988; Mello et al., 1988; Clark and Philp, 1989). Additionally, high C₃₅ HH is also a good indicator of highly reducing (low Eh) marine conditions (Peters and Moldowan, 1991). Therefore, the sensitivity of C_{35}/C_{34} HH ratio to depositional environment and rock lithology makes it a helpful parameter used in this research.

The C₃₅/C₃₄ HH of Speake Ranch samples show values ranging from 0.58-0.91, and according to Peters and Moldowan (1993) and Bishop and Farrimond (1995) the values exceeding 0.8 is indicative of carbonate source rocks (Table 6; Figure 38). The C₃₅/C₃₄ HH can be used in tandem with C₂₉ Ts/C₃₀ diahopane to further identify source facies of Speake Ranch. The correlation between C₃₅/C₃₄ HH against C₂₉ Ts/C₃₀ diahopane is a reliable indicator of lithology, source facies, and level of oxicity. The rearranged hopanes 17α -diahopane (C₃₀ diahopane: C₃₀D)

and 18α -30-norneohopane (C₂₉Ts) are sensitive to redox conditions (Peters et al., 2005). The oxygen content in the depositional environment can also be evaluated using C₂₉Ts/C₃₀D ratio. High values (>1) of C₂₉Ts/C₃₀D ratio show shale sediment deposition under oxidizing conditions (Peters and Moldowan, 1993; Peters et al., 2005). The Speake Ranch samples have very low values of the C₂₉Ts/C₃₀D ratio (0.07-0.1) in Figure 39. These values are observed to be marine shale deposited in anoxic environmental conditions.



Figure 39. C_{35}/C_{34} Homohopane vs. C_{29} Ts/ C_{30} D plot to determine lithology and oxicity level of depositional environment. The Speake Ranch samples were largely deposited in anoxic environment with oxic to sub-oxic disturbances (Zumberge, 2000).

3.6.4.2.1 Ts and Tm

Ts (18α -22,29,30-trisnorneohopane) and Tm (17α -22,29,30-trisnorhopane) are a pair of C₂₇ hopanes that are commonly used to identify maturity and anoxia levels of the depositional environment (Waples and Machihara, 1990; El-Sabagh et al., 2018). This is a more reliable

parameter in relation to the sterane biomarkers, as the 20S/(20S+20R) sterane ratio remains in equilibrium after the peak oil generation window, while the Ts and Tm continue to change with increasing thermal maturity. Clear illustration of the consistency in maturity parameter C₂₉ 20S/(20S+20R) in Figure 40, shows fluctuating values in Ts/Tm. Values of Ts/Tm in the Speake Ranch samples range from 0.2 to 0.38 in reflection to the immature sediments that are deposited in saline conditions. This agrees with the fact that the organic rich layers are mainly deposited under anoxic environmental conditions with intervals of oxic to sub-oxic that are frequent within such small intervals.



Figure 40. Geochemical logs to determine anoxia proxies (Ts/Tm) of Speake Ranch deposition, and if (at all) output values have been affected by thermal maturation determined using C_{29} sterane 20S/(20S+20R).

Gammacerane (XI: Appendix III) has been used to assess water column stratification and an aid to identify photic zone anoxia during sediment deposition. Gammacerane (XI) is proposed to be derived from tetrahymanol present in bacterivorous ciliates that thrive at the boundary of water salinity differences in the water column (Grantham et al., 1983). Torres (2020) suggested that higher proportions of gammacerane have been found at the base of the Lower Woodford, Middle Woodford, and Upper Woodford of Speake Ranch. This suggests that stratification and water column salinity played a significant role within the sequence transitions. The use of gammacerane index (GI=G/G+H) parameter has been helpful to determine water column stratification (Sinninghe Damste et al., 1995). The GI of the observed samples plot at values below 0.2 (marker for water salinity and stratification), where the Middle Woodford experienced a gentle decline and moved to a gradual increase within the Upper Woodford (Figure 41). This trend agrees with the discovery made by Torres (2020). The values from GI illustrates a range from 0.12-0.17, showing subtle changes in salinity levels. From the abundance of the gammacerane it appears that saline conditions and density driven water column stratification weakly persisted throughout deposition and periodically euxinia existed as indicated by the presence of aryl isoprenoids discussed below. In the Middle Woodford the depositional conditions were more restricted, possibly resembling a pseudo-lacustrine depositional environment later supported by tetracylic polyprenoids to be discussed in the following sections. This relationship can be seen in some intervals within Figure 41. Increased water salinity attributes to density stratification and causes a reduction in oxygen content in bottom waters (lower Eh), consequently causing lower Pr/Ph values. A modified plot from Mello et al. (1988) shows that water column stratification does not necessarily covary with varying levels of anoxia (Figure 42). Connock (2015) found similar GI trends in the Woodford Shale from Wyche quarry, and suggested trend of GI is highly correlative

to the C_{40} aromatic carotenoids (paleorenieratane and isorenieratane) where competition ensued between the potential source organism (bacterial ciliates) and the sulfur-oxidizing chemoautotrophic bacteria.



Figure 41. Depth plots for indication of water salinity, water column stratification, and anoxia. GI define the water column stratification, TPP/C₂₇Diasterane and Hopane/C₂₇ $\alpha\alpha\alpha$ S+R sterane define the organic matter source, and Pr/Ph defines the water column anoxia. No correlation of the GI to Pr/Ph and organic matter source parameters can be determined.



Figure 42. Plot of the relationship between Gammacerane Index and Pr/Ph of the Woodford Shale. There is no direct relationship between Pr/Ph and Gammacerane index, where heightened gammacerane values would yield lower Pr/Ph values (modified from Mello et al., 1988).

The 17α , 18α , 21β -28, 30-bisnorhopane (**XII**) is one of the three C₂₈ hopane stereoisomers first reported in the Monterey Shale by Seifert and Moldowan (1978). Banks et al. (1967) and Grantham et al. (1979) suggested that small concentrations of the 28, 30-bisnorhopane (**XII**) in the extract of a pre-Devonian source rock were related to ferns. Therefore, its presence could provide an indication of vascular land plant input within the Speake Ranch samples. However, it was later discovered that 28, 30-bisnorhopane (**XII**) may have also been derived from anaerobic bacteria in anoxic depositional environments (Katz, 1983). The latter study is more suitable for the application of this research. It is interpreted that a change in environment from restricted to marine conditions may strongly decrease sulfur oxidizing chemoautotrophic bacterial activity, which controls the abundance of bisnorhopane (Tornabene et al., 1979; Curiale and Odermatt, 1989). Therefore, the water column. Moreover, according to Hughes et al. (1995) the abundance of bisnorhopanes are direct results of sulfur rich environments, making it a possible tool to identify sulfur content for this 40ft interval of the Woodford Shale from Speake Ranch. Mackenzie et al. (1984) measured the bisnorhopane (XII) content relative to the 30-nor- $17\alpha 21\beta$ -hopane (XIII) defined by C₂₈ BNH/C_{29} norhopane ratio to identify the anoxic levels of the depositional environment in which was used in the Draupne formation in North Sea by Dahl (2004). Additionally, Curiale and Odermatt (1989) also suggested that utilizing the bisnorhopane index (C_{28} BNH/ C_{30} hopane) would be helpful to determine anoxic conditions. The transition from low or absent bisnorhopane content to higher C_{28} BNH/ C_{29} norhopane ratios may be a reflection of a change from archipelago setting to a basin deepening setting of widespread marine conditions, just as found at the Draupne Formation in West Greenland (Mackenzie et al., 1984; Miller 1990; Rattey and Hayward, 1993; Dahl, 2004). It is found within Speake Ranch that the C₂₈ BNH/C₂₉ norhopane ratio varies considerably over the 40ft interval and additionally, bisnorhopane abundance covaries with the organic richness. As Curiale et al. (1985) indicated that high bisnorhopane abundance correlates with Pr/Ph. Similarly, the C28 BNH/C29 norhopane ratio of Speake Ranch does coincide with the trend of anoxia determined by the Pr/Ph from its negative correlation shown at 226ft (Upper Woodford), 223ft (Upper Woodford), and 207ft (Middle Woodford) (Figure 43). The majority of relatively lower C₂₈ BNH/C₂₉ norhopane values are found within the softer bed intervals (Table 6). Tornabene et al. (1979) determined from a study that the C₂₈ BNH/C₂₉ norhopane ratio does not co-vary with lithology nor the organic matter quality, but instead is associated with anoxia. A decreasing C₂₈ BNH/C₂₉ norhopane trend reflects the change from restricted to more general marine conditions causing termination or strong reduction of the sulfur oxidizing chemoautotrophic bacterial activity in the water column responsible for controlling the synthesis

of bisnorhopane (Tornabene et al., 1979). A later section within this chapter will discuss the correlation of anoxia as determined by bisnorhopane and Pr/Ph ratios to the amount of sulfur present in the water column determined by dibenzothiophenes.



Figure 43. Speake Ranch anoxia proxies utilizing C_{28} BNH/ C_{30} hopane, C_{28} BNH/ C_{29} norhopane, and Pr/Ph to observe frequency of anoxia.

DBT/PHEN	0.82	0.69	0.87	0.22	0.68	0.59	0.74	0.57	0.56	0.25	0.68	0.70			0.65	0.49		0.79	0.49	0.85	0.79	0.86	0.95	0.87	0.88	0.95	,	0.39	0.78	0.55	0.32	0.74	0.70	0.38	0.80	0.67	0.57	0.54
C ₃₀ D/C ₂₉ Ts	0.08	0.09	0.09	0.07	0.09	0.09	0.08		0.08	0.08	0.08	0.08	,	,	0.10	0.08	0.08	0.07	0.07	0.09	0.09	0.08	0.09	0.09	0.08	0.08	,	0.07	0.07	0.08	0.07	0.08	0.08	0.08	0.08	0.08	0.09	0.09
Ts/Tm	0.31	0.33	0.33	0.31	0.28	0.30	0.26		0.28	0.29	0.26	0.28			0.29	0.30	0.34	0.30	0.29	0.29	0.28	0.27	0.30	0.29	0.29	0.27		0.35	0.33	0.34	0.38	0.35	0.33	0.35	0.33	0.34	0.35	0.38
IHH	8.16	5.80	6.96	5.46	5.34	7.02	6.91		8.53	7.59	5.86	6.39	,		6.78	6.76	6.80	5.32	7.28	6.37	6.47	9.82	6.15	6.10	6.94	7.02	,	8.40	6.74	8.57	7.22	6.28	6.32	7.12	6.27	6.76	8.19	7.50
C ₃₅ /C ₃₄ HH	06.0	0.80	0.74	0.61	0.64	0.81	0.77	,	0.84	0.69	0.72	0.74		,	0.81	0.65	0.68	0.64	0.67	0.70	0.73	0.91	0.65	0.64	0.82	0.93	,	0.77	0.68	0.82	0.68	0.71	0.67	0.67	0.58	0.73	0.83	0.69
C ₂₈ BNH/Norhopane	2.18	1.97	1.93	1.71	2.09	2.05	2.07	,	2.08	1.65	2.07	2.07	,	,	2.03	1.72	1.34	1.94	1.67	2.02	1.93	1.62	1.94	2.02	2.18	1.92	,	1.59	2.17	1.54	1.47	2.28	2.25	1.47	2.13	2.11	2.21	2.23
C ₂₈ BNH/C ₃₀ Hopane	0.50	0.53	0.52	0.29	0.39	0.34	0.42	,	0.36	0.25	0.38	0.42	,	,	0.37	0.27	0.23	0.38	0.26	0.35	0.38	0.27	0.34	0.35	0.39	0.39	,	0.24	0.35	0.26	0.23	0.39	0.39	0.24	0.38	0.36	0.37	0.37
Pr/Ph	0.76	0.75	0.66	1.05	0.64	0.80	0.94	0.92	1.10	0.78	0.71	0.63		,	0.73	0.78	1.55	0.84	1.15	0.85	0.76	1.65	0.69	0.93	0.77	0.75	,	1.28	0.75	1.33	2.56	0.86	0.77	2.13	0.77	0.79	0.66	0.72
TOC (wt%)	12.00	5.19	12.90	2.02	9.22	11.30	9.13	12.50	11.40	1.78	10.80	10.90	0.62	5.59	10.40	2.79	1.72	15.50	6.46	12.10	12.30	4.18	10.60	13.10	10.10	10.20	1.13	3.16	10.80	3.51	0.67	11.10	10.30	3.04	11.20	9.82	9.74	11.10
Lithology	Soft	Hard	Soft	Hard	Soft	Soft	Soft	Soft	Soft	Hard	Soft	Soft	Hard	Soft	Soft	Hard	Hard	Soft	Hard	Soft	Soft	Hard	Soft	Soft	Soft	Soft	Hard	Hard	Soft	Hard	Hard	Soft	Soft	Hard	Soft	Soft	Hard	Soft
Depth (ft.)	190.50	191.00	191.50	192.00	193.50	194.50	195.50	196.50	197.50	198.00	199.50	200.50	202.00	203.50	204.50	205.00	207.00	208.50	209.00	210.50	211.50	212.00	213.50	214.50	215.50	217.50	218.00	219.00	221.50	222.00	223.00	224.50	225.50	226.00	227.50	228.50	229.00	230.50
Sample	SR-190b	SR-191a	SR-191b	SR-192a	SR-193b	SR-194b	SR-195b	SR-196b	SR-197b	SR-198a	SR-199b	SR-200b	SR-202a	SR-203b	SR-204b	SR-205a	SR-207a	SR-208b	SR-209a	SR-210b	SR-211b	SR-212a	SR-213b	SR-214b	SR-215b	SR-217b	SR-218a	SR-219a	SR-221b	SR-222a	SR-223a	SR-224b	SR-225b	SR-226a	SR-227b	SR-228b	SR-229a	SR-230b

Woodford Shale from Speake Ranch. Emp	
. Geochemical parameters defined to determine redox conditions of the	s are due to the lack of maltene from extraction.
Table 6.	intervals

3.6.5 C₄₀ Aromatic carotenoids and aryl isoprenoid

Anoxic conditions in marine environments are a product of a depleted supply of oxygen and a lack of water circulation at marine bottom waters due to temperature and density stratification (Koopmans et al., 1996). The zone of oxygen depletion is commonly termed as photic zone euxinia (PZE) which can extend from marine bottom waters into the photic zone as indicated by the presence of the green sulfur bacteria, Chlorobiaceae. The green sulfur bacteria is an anoxygenic photosynthetic bacteria that requires the photic zone to be in H_2S waters (Koopmans et al., 1996). According to Koopmans et al. (1996), Chlorobiaceae utilizes reverse tricarboxylic acid cycle to fix CO₂. The presence of PZE can be determined by the abundance of C₄₀ diaromatic carotenoids (Grice et al., 1996). The paleorenieratane (XIV), isorenieratane (XV), and renieratane (XVI) are derivatives of the C_{40} diaromatic carotenoids detected at the mass fragmentogram m/z 133-134 (Figure 44; Table 7). Isorenieratane (XIX), a diaromatic carotenoid diagenetic and catagenetic product, correlates with a distinct enrichment in δ^{13} C. This provides for the initial evidence of the applicability of isorenieratane and its derivatives, such as aryl isoprenoids, for the implication of photic zone euxinia, which will be utilized in this study. Percival et al. (2019) found that there is a positive correlation between TOC and δ^{13} C. Furthermore, samples from Dongcun and Yangdi in China have shown that the δ^{13} C closely tracks with oxygenation levels, where the values are at a maximum at the lowering of oxygenation (Joachimski et al., 2002). The paleorenieratane (XIV) and the isorenieratane (XV) are present at higher concentrations in the Woodford Samples from Speake Ranch (Figure 44). A consistent proportion of higher paleorenieratane to the isorenieratane are consistent throughout the Speake Ranch section despite the lithofacies variations. The proportions of the C₄₀ carotenoid paleorenieratane/isorenierane were plotted against Pr/Ph (Figure 45). This ratio serves as evidence that the carotenoids are affected with varying levels of anoxia. All of the soft bed lithofacies plot in the reducing area, while the

hard beds (yellow symbol) vary within areas defining oxidizing and reducing conditions. Due to data limitation only 13 Speake Ranch samples were able to be observed.



Figure 44. Identification of the C_{40} carotenoids screened at summed m/z 133+134 at depth 204.5ft within the Middle Woodford. Top chromatogram is at depth 205ft from the soft bed, and bottom chromatogram is at depth 204.5ft from the hard bed.



Figure 45. Paleorenieratane/Isorenieratane ratio vs. Pr/Ph ratio illustrating a mainly reducing condition of the Speake Ranch samples defined by the concentration and distribution of carotenoids. However, hard beds (yellow symbols) are seen to vary within areas of oxidation and anoxia.

Aryl isoprenoids are the diagenetic products of aromatic carotenoids, whose formation result from the cleavage of the isorenieratane chain. Koopmans et al. (1996) has stated that there is a broad range of diagenetic and catagenetic derivatives of the diaromatic carotenoid isorenieratane, a pigment of *Chlorobiaceae* found in sedimentary rocks. These aryl isoprenoids are comprised of C₄₀, C₃₃, C₃₂, diaryl isoprenoids, and short-chain aryl isoprenoids. The compounds of interest in addition to paleorenieratanes and isorenieratanes, are the aryl isoprenoids, which are extremely useful in evaluating variability in the persistence of PZE. Distribution of the aryl isoprenoids are found in the mass fragmentogram m/133-134 (Figure 46; Table 7). The assessment of PZE is done through the calculation of the short-chain C_{13} - C_{17} of the 2,3,6-trimethylated aryl isoprenoids (2,3,6-TMB) (XVII) substitution proportion in relation to the intermediate chain aryl isoprenoid, C₁₈-C₂₂ 2,3,6-trimethylated aryl isoprenoids (3,4,5-TMB) (XVII), to obtain the aryl isoprenoid ratio (AIR). The AIR is helpful to determine the relative amounts of aryl isoprenoids from Chlorobiaceae indicated by lower values, whereas higher values will indicate cyanobacteria input (Schwark and Frimmel, 2004). It is known that if there is more of an abundance of the short chain components it is an indication of a more intense aerobic degradation of aryl isoprenoids (Schwark and Frimmel, 2004).

Development of PZE which is the enhancement of anoxia and sulfide within the photic zone of the water column, is additional support to indicate restriction within the basin that may have been caused by early erosional events of the pre-Woodford. Schwark and Frimmel (2004) suggested that the development of a low AIR coupled with low Pr/Ph values within the samples is suggestive of more persistent PZE conditions, while high values of AIR (>3) and Pr/Ph (>1) values indicate short termed episodic PZE. Schwark and Frimmel (2004) concluded that AIRs are never below 0.5 indicating that PZE has always been episodic. The Woodford Shale has shown values

ranging from 0.34-1.18, indicating variable episodic PZE (Figure 47). Additionally, not much variation can be seen within the boundary separating the Middle Woodford and Upper Woodford at Speake Ranch. A distinctive fluctuation is found between 210-220ft which may indicate unstable PZE and may be a time marker of F-F coinciding with the Pr/Ph depth plot.



Figure 46. GC-MS chromatogram showing the distribution of aryl isoprenoids in the aromatic fraction at the summed m/z 133+134. 2,3,6-TMB aryl isoprenoids (**XXII**) are indicated by the letter b, and the 3,4,5-TMB (**XXIII**) are indicated by the letter a.
Peak	Compound	Peak	Compound
No.		No.	
1	C ₁₃ TMB	10a	C ₂₂ 3,4,5 TMB
2	C ₁₄ TMB	10b	C ₂₂ 2,3,6 TMB
3a	C ₁₅ 3,4,5 TMB	11a	C ₂₄ 3,4,5 TMB
3b	C ₁₅ 2,3,6 TMB	11b	C ₂₄ 2,3,6 TMB
4a	C ₁₆ 3,4,5 TMB	12a	C ₂₅ 3,4,5 TMB
4b	C ₁₆ 2,3,6 TMB	12b	C ₂₅ 2,3,6 TMB
5a	C ₁₇ 3,4,5 TMB	13a	C ₂₆ 3,4,5 TMB
5b	C ₁₇ 2,3,6 TMB	13b	C ₂₆ 2,3,6 TMB
6а	C ₁₈ 3,4,5 TMB	14a	C ₂₇ 3,4,5 TMB
6b	C ₁₈ 2,3,6 TMB	14b	C ₂₇ 2,3,6 TMB
7a	C ₁₉ 3,4,5 TMB	15a	C ₂₉ 3,4,5 TMB
7b	C19 2,3,6 TMB	15b	C ₂₉ 2,3,6 TMB
8a	C ₂₀ 3,4,5 TMB	16a	C ₃₀ 3,4,5 TMB
8b	C ₂₀ 2,3,6 TMB	16b	C ₃₀ 2,3,6 TMB
9a	C ₂₁ 3,4,5 TMB	17a	C ₃₁ 3,4,5 TMB
9b	C ₂₁ 2,3,6 TMB	17b	C ₃₁ 2,3,6 TMB

Table 7. Peak Identification for the 2,3,6- and 3,4,5 trimethyl substitution for the aryl isoprenoid chromatogram at m/z 133-134 in Figure 48.

The distribution and variability of aryl isoprenoid composition is mainly affected by the primary environmental conditions during sediment deposition, burial, as well as diagenesis (Schwark and Frimmel, 2004). This in turn can also reflect the productivity of *Chlorobiaceae*, which could be explained by the extent of anoxia that intruded through the photic zone or to the availability of nutrients (Schwark and Frimmel, 2004). The concentration of *Chlorobiaceae* biomass found in sediment is mainly controlled by the persistence of PZE over a time interval, which is controlled by the "stability of water-column" driven by climate (Schwark and Frimmel, 2004). They also concluded that the high surface productivity and the decomposition of organisms via sulfate-reducing bacteria within the sediments as well as anoxia bottom waters during deposition would lead to an expansion of the euxinic zone.

The C_{18} aryl isoprenoid ratio was calculated from the proportions of the C_{18} 2,3,6-trimethyl aryl isoprenoid (**XVII**) to the C_{18} 3,4,5-trimethyl substitution (**XVIII**) to determine redox

conditions of the depositional environment. Relatively low values of C_{18} AIR plotting below 10 (Figure 47) coincide with low Pr/Ph ratios indicating anoxic waters and presence of H₂S, later confirmed by tetracyclic polyprenoids as well. The instability of C_{18} aryl isoprenoid ratio parameters is also monitored in the same interval where this behavior is found in the Pr/Ph and AIR (2,3,6/3,4,5-trimethyl-substituted aryl isoprenoid; Formulas can be found in Appendix III). In summary, the Upper and Middle Woodford members from Speake Ranch show evidence for bottom water anoxia including periods of oxic to sub-oxic as indicated by Pr/Ph values, and periodic photic zone euxinia as indicated by the aryl isoprenoid ratio.

Sample	Depth (ft.)	Lithology	TOC (wt%)	Pr/Ph	AIR	C ₁₈ -Aryl Isoprenoid	Gammacerane Index	C35/C34 HH	HHI
SR-190b	190.50	Soft	12.00	0.76	0.84	4.04	0.15	0.90	0.08
SR-191a	191.00	Hard	5.19	0.75	0.75	5.17	0.16	0.80	0.45
SR-191b	191.50	Soft	12.90	0.66	1.12	10.96	0.17	0.74	0.07
SR-192a	192.00	Hard	2.02	1.05	1.15	0.72	0.15	0.61	0.38
SR-193b	193.50	Soft	9.22	0.64	0.83	6.94	0.16	0.64	0.05
SR-194b	194.50	Soft	11.30	0.80	0.60	5.16	0.13	0.81	0.07
SR-195b	195.50	Soft	9.13	0.94	0.62	4.24	0.16	0.77	0.07
SR-196b	196.50	Soft	12.50	0.92	0.56	6.49	-	-	-
SR-197b	197.50	Soft	11.40	1.10	0.57	3.67	0.14	0.84	0.09
SR-198a	198.00	Hard	1.78	0.78	0.52	8.70	0.15	0.69	0.08
SR-199b	199.50	Soft	10.80	0.71	0.72	4.34	0.16	0.72	0.06
SR-200b	200.50	Soft	10.90	0.63	0.78	8.27	0.15	0.74	0.06
SR-204b	204.50	Soft	10.40	0.73	0.68	4.41	0.15	0.81	0.07
SR-205a	205.00	Hard	2.79	0.78	0.59	5.39	0.14	0.65	0.07
SR-207a	207.00	Hard	1.72	1.55	0.34	4.23	0.14	0.68	0.07
SR-208b	208.50	Soft	15.50	0.84	0.83	5.47	0.14	0.64	0.05
SR-209a	209.00	Hard	6.46	1.15	0.85	9.05	0.14	0.67	0.07
SR-210b	210.50	Soft	12.10	0.85	0.95	6.20	0.14	0.70	0.06
SR-211b	211.50	Soft	12.30	0.76	1.55	6.90	0.14	0.73	0.06
SR-212a	212.00	Hard	4.18	1.65	0.34	7.62	0.13	0.91	0.10
SR-213b	213.50	Soft	10.60	0.69	1.06	6.73	0.13	0.65	0.06
SR-214b	214.50	Soft	13.10	0.93	0.56	4.73	0.14	0.64	0.06
SR-215b	215.50	Soft	10.10	0.77	0.78	7.98	0.14	0.82	0.07
SR-217b	217.50	Soft	10.20	0.75	1.26	9.79	0.12	0.93	0.07
SR-219a	219.00	Hard	3.16	1.28	0.17	6.88	0.15	0.77	0.08
SR-221b	221.50	Soft	10.80	0.75	0.62	8.41	0.14	0.68	0.07
SR-222a	222.00	Hard	3.51	1.33	0.46	12.73	0.14	0.82	0.09
SR-223a	223.00	Hard	0.67	2.56	0.40	4.76	0.13	0.68	0.07
SR-224b	224.50	Soft	11.10	0.86	-	-	0.13	0.71	0.06
SR-225b	225.50	Soft	10.30	0.77	0.75	4.91	0.15	0.67	0.06
SR-226a	226.00	Hard	3.04	2.13	0.67	11.54	0.14	0.67	0.07
SR-227b	227.50	Soft	11.20	0.77	0.72	9.27	0.15	0.58	0.06
SR-228b	228.50	Soft	9.82	0.79	0.79	10.75	0.15	0.73	0.07
SR-229a	229.00	Hard	9.74	0.66	0.62	6.93	0.17	0.83	0.08
SR-230b	230.50	Soft	11.10	0.72	1.18	8.41	0.16	0.69	0.08

Table 8. Biomarker proxies for water structure and chemistry in the Woodford Shale at Speake Ranch. Pr/Ph, HHI, and C_{35}/C_{34} HH indicating the redox state of the Woodford paleoenvironment, AIR and C_{18} Aryl Isoprenoid signify the H₂S presence within the photic zone as well as the stability of the PZA, and the C_{27}/C_{29} steranes used as a correlation to the effects of the organic matter input source.





3.6.6 Tetracyclic Polyprenoids

The C_{30} tetracyclic polyprenoids (XIX) which are detected by GC-MS using the ions at m/z 191 and m/z 259 are highly specific indicators of brackish lacustrine environment and freshwater algae (Holba et al., 2003). They are represented by two peaks that signify the 21S and 21R isomers, suggested to be derived from green algae (*Chlorophyta*) (Figure 48; Li et al., 1996; Poinsot et al., 1997; Holba et al., 2003; Araujo et al., 2018). Not only do their abundances indicate lacustrine environment dominance over marine environment, but they also indicate a predominance of fresh-water algal precursor or preferential preservation of saturate hydrocarbons within non-marine conditions (Holba et al., 2000; Araujo et al., 2018). When tetracylic polyprenoids (TTP) (XIX) are correlated with tricyclic terpane ratios $[C_{26}/C_{25} TT = C_{26} 13\beta, 14\alpha$ tricyclic terpane)/(C_{25} 13 β ,14 α -tricyclic terpane)] the distinction between marine or lacustrine source rock organic matter is enhanced (Holba et al., 2000). Furthermore, abundance of the pentacyclic terpanes, C_{30} 17 α , 21 β -hopane (C_{30} H) identify the significant presence of microbially reworked organisms (Mello et al., 1988; Araujo et al., 2018). The combination of C₂₆/C₂₅ tricyclic terpane (C₂₆/C₂₅ TT), C₃₀ tetracyclic polyprenoid/C₂₇ diasterane (TPP/C₂₇ diasteranes), and, Hopane/ $C_{27} \alpha \alpha \alpha (S+R)$ sterane ratios provide a stronger correlation platform to identify lacustrine vs. marine paleoenvironments. If values are high enough to indicate lacustrine environment, then these parameters may indicate that the marine environment had become so restricted it became pseudo lacustrine. TPP/C₂₇ diasteranes values are well below 0.4, illustrating a marine depositional environment (Casilli et al., 2014; Araujo and Azevedo, 2016). Additionally, based on Figure 49, harder beds tend to illustrate higher TPP C₃₀ values relative to the C₂₇ diasteranes. According to Zumberge et al. (1987), Mello et al. (1988), Casilli et al. (2014), and Araujo et al. (2018) values of high Hopane/ $C_{27} \alpha \alpha \alpha$ (S+R) sterane (>5), high C_{26}/C_{25} TT (>1) and high TPP/ C_{27} diasterane (>1) are associated with a lacustrine environment, which were found in crude oils from Espirito Santo

Basin, Brazil. While, low Hopane/C₂₇ $\alpha\alpha\alpha(S+R)$ sterane (0.58-3.70), low C₂₆ TT/C₂₅ TT (0.78-0.85) and low TPP/C₂₇ diasterane (0.07-0.24) reflect a marine environment. The values of Speake Ranch samples illustrated in Figure 49 and Table 9 clearly depict that these observed samples have low Hopane/C₂₇ $\alpha\alpha\alpha(S+R)$ sterane with some intervals plotting above 10. TPP/C₂₇ diasteranes ratios range from 0.1 to 3, showing low, but largely varying values. Additionally, the C₂₆/C₂₅ TT have values that are commonly invariant around 1. The highlighted intervals in Figure 49 show that relatively higher Hopane/C₂₇ $\alpha\alpha\alpha(S+R)$ sterane and TPP/C₂₇ diasteranes illustrate the pseudo lacustrine environment. Red highlighted zones are those increased parameters within the Middle Woodford, and the yellow highlighted interval indicates elevated parameter values within the Upper Woodford. In summary, data show that lowest TOC values coincide with all the combined effects of oxicity and no specificity of water stratification, while elevated TOC values can be found to associate with the highest indication of anoxia and elevated gammacerane values.



Figure 48. Fragmentogram of C_{30} tetracyclic polyprenoid (**XII**) on mass spectra 259. SR-192a is a hard bed at depth 192 ft, and SR-193b is a soft bed at 193.5 ft.



depositional environment. Highlighted intervals indicate intervals when the environment of deposition for the Woodford Shale Figure 49. Variation in the TPP screened at m/z 259 along with its correlation to gammacerane Index, C₃₀Hopane/C₂₇ aaaS+R sterane, C₂₆ TT/C₂₅, Ts/Tm to determine intervals that illustrate the level of restriction in the water column of the Woodford at Speake ranch was highly restricted.

DBT/PHEN	0.82	0.69	0.87	0.22	0.68	0.59	0.74	0.57	0.56	0.25	0.68	0.70	I	ı	0.65	0.49	i	0.79	0.49	0.85	0.79	0.86	0.95	0.87	0.88	0.95	i	0.39	0.78	0.55	0.32	0.74	0.70	0.38	0.80	0.67	0.57	0.54
GI	0.15	0.16	0.17	0.15	0.16	0.13	0.16		0.14	0.15	0.16	0.15			0.15	0.14	0.14	0.14	0.14	0.14	0.14	0.13	0.13	0.14	0.14	0.12		0.15	0.14	0.14	0.13	0.13	0.15	0.14	0.15	0.15	0.17	0.16
C26/C25 TT	1.03	0.94	0.96	0.85	0.95	0.94	0.93		0.97	0.89	0.92	0.98	,		0.83	0.96	0.94	0.92	0.96	0.92	0.97	0.97	0.97	1.00	0.87	0.96		0.87	0.93	0.94	0.95	0.93	0.84	0.94	0.78	0.88	0.81	0.84
$C_{30} \; H/ \; C7 \alpha \alpha \alpha S + R$	0.71	0.47	0.48	19.48	0.91	1.72	1.11		1.34	8.60	0.75	0.61	,	,	09.0	3.04	28.45	0.92	7.50	2.06	1.03	7.99	4.45	1.73	1.16	0.87		5.64	4.21	6.99	10.20	1.61	2.14	6.04	3.46	3.05	2.88	1.07
TPP/C ₂₇ Diasterane	0.12		0.10	0.23	0.14	0.13	0.13		0.12	0.27	0.12	0.08		,	0.14	0.23	0.22	0.12	0.20	0.12	0.11	0.20	0.12	0.13	0.12	0.11		0.31	0.19	0.28	0.30	0.19	0.21	0.28	0.22	0.24	0.23	0.22
Pr/Ph	0.76	0.75	0.66	1.05	0.64	0.80	0.94	0.92	1.10	0.78	0.71	0.63	ı	ı	0.73	0.78	1.55	0.84	1.15	0.85	0.76	1.65	0.69	0.93	0.77	0.75	ı	1.28	0.75	1.33	2.56	0.86	0.77	2.13	0.77	0.79	0.66	0.72
TOC (wt%)	12.00	5.19	12.90	2.02	9.22	11.30	9.13	12.50	11.40	1.78	10.80	10.90	0.62	5.59	10.40	2.79	1.72	15.50	6.46	12.10	12.30	4.18	10.60	13.10	10.10	10.20	1.13	3.16	10.80	3.51	0.67	11.10	10.30	3.04	11.20	9.82	9.74	11.10
Lithology	Soft	Hard	Soft	Hard	Soft	Soft	Soft	Soft	Soft	Hard	Soft	Soft	Hard	Soft	Soft	Hard	Hard	Soft	Hard	Soft	Soft	Hard	Soft	Soft	Soft	Soft	Hard	Hard	Soft	Hard	Hard	Soft	Soft	Hard	Soft	Soft	Hard	Soft
Depth (ft.)	190.50	191.00	191.50	192.00	193.50	194.50	195.50	196.50	197.50	198.00	199.50	200.50	202.00	203.50	204.50	205.00	207.00	208.50	209.00	210.50	211.50	212.00	213.50	214.50	215.50	217.50	218.00	219.00	221.50	222.00	223.00	224.50	225.50	226.00	227.50	228.50	229.00	230.50
Sample	SR-190b	SR-191a	SR-191b	SR-192a	SR-193b	SR-194b	SR-195b	SR-196b	SR-197b	SR-198a	SR-199b	SR-200b	SR-202a	SR-203b	SR-204b	SR-205a	SR-207a	SR-208b	SR-209a	SR-210b	SR-211b	SR-212a	SR-213b	SR-214b	SR-215b	SR-217b	SR-218a	SR-219a	SR-221b	SR-222a	SR-223a	SR-224b	SR-225b	SR-226a	SR-227b	SR-228b	SR-229a	SR-230b

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3.6.7 Dibenzothiophene and phenanthrene

Dibenzothiophenes (**XX**) are important sulfur-containing aromatic compounds with two benzene rings connected to a central thiophene ring (Figure 50; Li et al., 2012). This compound is thought to be derived from a thermal reaction between sulfur and the organic matter contained within the sediment producing dibenzothiophene (**XX**). The dibenzothiophene/phenanthrene (DBT/Phen) ratio is often used to assess the presence and abundance of reduced sulfur within the organic matter, primarily hydrogen sulfide (H₂S) and polysulfides (H₂Sn) (Hughes et al., 1995). This is a helpful parameter to indicate anoxia within the depositional environment as sediments must be anoxic for the accumulation of reactive sulfur (Hughes et al., 1995). Dibenzothiophenes are monitored on the GC-MS of the aromatic fraction at m/z 184, 198, and 212 (Figure 50), while the phenanthrenes (**XXI**) are observed at m/z 178, 192, 206 as shown in Figure 51.



Retention time

Figure 50. GC-MS chromatogram monitored for the variation in the dibenzothiophenes (**XVI**) at m/z 184+198+212. The m/z 184 is used to identify the dibenzothiophene, m/z 198 is used to identify the methyldibenzothiophene, and m/z 212 is used to identify the dimethyldibenzothiophene.



Retention time

Figure 51. GC-MS chromatogram displaying the distribution of the phenanthrenes (**XV**) series observed at summed m/z 178+192+206. The m/z 178 is used to identify the phenanthrene, m/z 192 is used to identify the methylphenanthrenes, and m/z 206 is used to identify the dimethylphenanthrenes.

According to Hughes et al. (1995) all oils from marine, predominantly carbonate source rocks have (DBT/Phen) ratios>1 and Pr/Ph ratios <1. Marine siliclastic and fluvial/deltaic source rocks have DBT/Phen ratios<0.5 and Pr/Ph ratios >1 as defined by Hughes et al. (1995). The Woodford has Pr/Ph ratios <3 whereas those generated from nonmarine carbonaceous shales and coals have Pr/Ph ratios >3 as defined in Table 9. Thus, the DBT/Phen ratio alone is a good discriminator of siliclastic vs. nonsiliclastic source rock lithologies. Sediments with the highest DBT/Phen ratios indicate elevated sulfur concentrations. Based on the DBT/Phen vs. Pr/Ph plot, the source rocks can be divided into 4 zones that determine its depositional environment and lithology (Table 10). According to Figure 52, the Speake Ranch samples are mostly of Zone 3 representing sulfide-rich sediments of lacustrine derived environment. However, several of the hard samples plot in the Zone 2, representing thiophene-rich sediments originating from a marine and lacustrine environment. Concentrations of DBT correlated to the organic carbon and clay content of the sediments, suggesting that the clay and organic carbon contents are two prime factors

that are controlling dibenzothiophene levels (Yang, 1998). Changing extents of anoxia (accompanied by changes in the variability of ferric iron) could have made different amounts of elemental sulfur available to the depositional systems corresponding to the two different lithofacies. The combined parameters Pr/Ph, bisnorhopanes, tetracylic polyprenoids, and gammacerane, with dibenzothiophene to phenanthrene, show evidence that sulfur rich intervals coincide with anoxic intervals as well as water column stratification, that provide a suitable environment for an abundant preservation of organic matter as shown in Figure 53. The highest values of TOC wt% are found within the intervals that show high values of all the mentioned environmental condition.



Figure 52. The Woodford samples from Speake Ranch show that the major lithology of the source rock is lacustrine/ marine shale of low Pr/Ph and DBT/Phen values. There are several of the hard bed intervals that indicate sulfate poor lacustrine environment (modified from Hughes et al., 1995).

ZONE	DBT/PHEN	PR/PH	DEPOSITIONAL ENVIRONMENT	LITHOLOGY
1 A	> 3	< 1	Marine	Carbonate
1B	1-3	< 1	Marine and Lacustrine (sulfate-rich)	Carbonate and Mixed*
2	< 1	< 1	Lacustrine (sulfate-poor)	Variable
3	< 1	1-3	Marine and Lacustrine	Shale
4	< 1	> 3	Fluvio/Deltaic	Carbonaceous shale and Coal

* Mixed includes marls, argillaceous carbonates and silaceous/phosphatic rocks

Table 10. Defining parameter for DBT/Phen with Pr/Ph outputs classified into 4 zones with 4 distinct environments and its associated lithology.



Figure 53. Correlation of redox conditions (Pr/Ph, C_{28} BNH/ C_{30} hopane, and C_{28} BNH/norhopane), presence of sulfur in water column (DBT/PHEN), and water column stratification (gammacerane), to the abundance of organic matter (TOC wt%) of the Woodford Shale from Speake Ranch. Gray intervals indicate the soft beds, and white intervals indicate the hard beds.

CHAPTER IV

4. Discussion

4.1 High Resolution Analysis of Biomarker Distributions

As discussed earlier, the Woodford Shale has frequent variations in the organic matter abundance within the Upper and Middle Woodford members. This positive correlation of TOC to the hard-soft bed couplet are controlled by changing lithofacies. The purpose of this research is to determine whether the variations found within the biomarkers are an effect of a compilation of multiple factors that include sea level changes, onset of terrigenous influx, tectonic disturbances, varying levels of anoxia, and marine biota productivity. In prior chapters, the effects of maturity have been eliminated because maturity changes are not expected or observed over a 40ft interval.

For the high-resolution analysis purposes discussed in this section, Speake Ranch samples with similar TOC values were taken at different depths of the 40ft interval. The aim of this chapter is to determine if intervals of similar TOC wt% would yield similar distributions of biomarkers. The chosen samples were 5 samples with relatively higher TOC wt% (11%) and 3 samples with relatively lower TOC wt% (~3%) found at varying depths within the Upper Woodford and Middle Woodford of Speake Ranch (Figure 54). All bulk geochemical source rock evaluation parameters, such as S₁, S₂, S₃, and T_{max}, were consistent throughout the chosen depths (Table 11). Significant variations were seen at depth 230.5 ft (SR-230b, Upper Woodford) for the soft bed group which shows the lowest HI value and highest OI value, possibly signifying a different kerogen type from the other samples, or one that is heavily weathered. Anoxia (based on the Pr/Ph) is consistent within intervals of high TOC. However, the soft bed at 197.5 ft has a higher Pr/Ph value, indicating a more oxic condition as shown in Table 11. In addition to that, SR-205a, of the hard bed group,

shows a Pr/Ph value indicating anoxic water column upon deposition. These two outlier values contradict the hypothesis that organic rich layers are always found in highly reducing environments. However, these values may be affected by weathering, as the Speake Ranch samples were collected from an exposed outcrop.

Sample	Depth (ft.)	Lithology	TOC (wt%)	S1 (mg HC/g)	S2 (mg HC/g)	S3 (mg CO2/g)	T _{max} (°C)	HI	OI	Pr/Ph
SR-230b	230.5	Soft	11.1	1.72	54.47	1.15	428	490.72	10.36	0.72
SR-227b	227.5	Soft	11.2	1.25	59.2	0.48	428	528.57	4.29	0.77
SR-224b	224.5	Soft	11.1	2.26	63.23	0.34	429	569.64	3.06	0.86
SR-197b	197.5	Soft	11.4	1.95	68.01	0.49	431	596.58	4.30	1.10
SR-194b	194.5	Soft	11.3	1.33	60.15	0.46	428	532.30	4.07	0.80
SR-226a	226	Hard	3.04	0.52	14.81	0.74	426	487.17	24.34	2.13
SR-219a	219	Hard	3.16	0.68	16.44	0.59	433	520.25	18.67	1.28
SR-205a	205	Hard	2.79	0.26	13.52	0.27	425	484.59	9.68	0.78

Table 11. High resolution source rock analysis of similar high TOC values distributed throughout the 40ft interval. Lighter green tables indicating Middle Woodford data.

4.1.1 Organic Matter Source

The C₃₀ sterane index is a direct reflection of marine source input (Waples and Machihara, 1990; Fildani et al., 2005). The soft bed group shown in Table 12 has consistent C_{27}/C_{29} sterane and C_{28}/C_{29} sterane values, indicating a predominantly marine source input along with a consistent C_{30} sterane index. The hard bed group shows consistent but contrasting values, except for SR-219a, indicating a higher marine input from C_{27}/C_{29} steranes and C_{28}/C_{29} steranes, but a lower C_{30} sterane index. It is possible that these values were also affected by weathering, as C_{30} steranes are already found in low concentrations in proportion to the $C_{27}-C_{28}-C_{29}$ steranes. The TST occur when sea level is rising (representing deposition of Middle Woodford), and the basin becomes flooded with less influx from terrigenous and reworked (inert) organic material. During the HST, more

terrigenous derived and reworked organic material (with lower HI values) comes into the basin (representing deposition of Upper Woodford). However, this can only be assumed for Speake Ranch samples as the 40ft interval only shows a dominating marine source input throughout the Upper and Middle Woodford in most of its soft bed lithofacies. As mentioned in an earlier chapter particulate organic matter in certain marine environments were found to contain C₂₉ steranes which were reasoned to be derived from green algae (chlorophyta), Prymnesiophycean algae, as well as cyanobacteria or *Tasmanites algae*, and not from land-plants (Grantham and Wakefield, 1987). From the depth logs of organic matter source input, high values of C₃₀ sterane index plotting beyond 4% are found periodically and correlates with the marine indicating ratios of the C_{27}/C_{29} steranes in Figure 54. Since all the organic matter source input parameters used in this study point toward a marine source input, it is interesting that we would see differences in the proportion of C_{29} and C_{27} steranes. The correlation of low C_{30} sterane index values with lower C_{27}/C_{29} sterane, C₂₃ TT/C₂₄ Tet, and C₁₉/C₂₃ TT ratios, may attest to the theory that bioturbated intervals yield lower organic carbon loading onto the sediment surfaces. This process caused remineralization and removal of organic carbon from sediment surface by oxygenation as previously suggested by Kennedy and Wagner (2011). The XRF-data of the Mo/Al ratio help determine whether water column mixing, and oxygenation co-vary with the low marine input intervals. The main elements analyzed by XRF are defined in Table 13. Interestingly, Mo/Al negatively correlates to all the organic matter source input parameters shown in Figure 54. This justifies that a lower marine source input indication and heightened "terrigenous" markers may just be a product of water column oxygenation. As sea level regressed, oxygenation is introduced by water column mixing and weathering of terrigenous sediments by erosion are introduced into the basin.

Sample	Depth (ft.)	Lithology	TOC (wt%)	Pr/Ph	Sterane index	C ₂₇ /C ₂₉ sterane	C ₂₈ /C ₂₉ sterane	C ₂₃ TT/C ₂₄ Tet	C ₁₉ /C ₂₃ TT
SR-230b	230.5	Soft	11.1	0.72	2.49	1.20	0.42	3.25	0.09
SR-227b	227.5	Soft	11.2	0.77	2.49	1.19	0.44	3.47	0.08
SR-224b	224.5	Soft	11.1	0.86	2.42	1.16	0.42	3.00	0.08
SR-197b	197.5	Soft	11.4	1.10	2.49	1.05	0.40	3.07	0.08
SR-194b	194.5	Soft	11.3	0.80	2.45	1.04	0.41	3.13	0.08
SR-226a	226	Hard	3.04	2.13	2.50	0.94	0.36	3.03	0.05
SR-219a	219	Hard	3.16	1.28	1.74	1.17	1.27	2.63	0.04
SR-205a	205	Hard	2.79	0.78	2.34	0.90	0.35	2.88	0.07

Table 12. Organic matter source high resolution analysis of similar high TOC values distributed throughout the 40ft interval to distinguish variability. Lighter green tables indicating Middle Woodford data.

Element	Proxy - Significance
Titanium (Ti)	Continental source and dust input
Zirconium (Zr)	Continental source
Silicon/Aluminum (Si/Al)	Quartz origin (biogenic or detrital)
Aluminum (Al)	Clay contents and feldspar
Potassium (K)	Clay contents and feldspar
Thorium (Th)	Clay contents and feldspar
Calcium (Ca)	Carbonate source and phosphates
Strontium (Sr)	Carbonate source and phosphates
Magnesium (Mg)	Carbonates, dolomitization
Manganese (Mn)	Carbonates, dolomitization
Uranium (U)	Organic matter richness, ?bitumen
Vanadium (V)	Bottom water anoxia, redox sensitive
Molybdenum (Mo)	Bottom water euxinia, redox sensitive
Sulfur (S)	Pyrite, reducing conditions, euxinia
Phosphorous (P)	Phosphate accumulation

Table 13. Main elements analyzed through XRF and their significance in chemostratigraphic interpretations. (Galvis, 2017; compiled in Turner, 2016)



Figure 54. Depth plots of the parameters defining the source of organic matter and its correlation to the Gamma Ray. Yellow highlighted intervals are the chosen samples for high resolution analysis with TOC values ~11%, and red highlighted intervals are chosen samples for low TOC ~3%.

4.1.2 Depositional Environment

The Pr/Ph values analyzed in the prior section suggested that a majority of the hard bed group 3% TOC (wt%) coincide with oxic depositional environment (high Pr/Ph), while high TOCs coincide with anoxic depositional environment (low Pr/Ph and high C_{35}/C_{34} HH). The anoxic intervals within the soft bed group 11% TOC (wt%) are mostly consistent, except for specific disturbances where intervals may have been more restricted in certain periods. Within these

selected intervals in Table 14, the C₂₈ BNH/C₃₀ hopane and the C₂₈ BNH/C₂₉ norhopane ratios show consistency in their values. The values of C₂₈ BNH/C₃₀ hopane and the C₂₈ BNH/C₂₉ norhopane are higher for intervals indicating anoxic water column (Table 14). Changing extents of anoxia (accompanied by changes in the variability of ferric iron) could have made different amounts of elemental sulfur available to the depositional systems corresponding to the two different lithofacies. Thus, the soft bed lithofacies which have been shown to be deposited in a more anoxic environment are enriched with the presence of sulfur by DBT/Phen (Table 15). While, hard bed lithofacies have lower sulfur content.

Sample	Dept h (ft.)	Lithology	TOC (wt%)	Pr/Ph	C ₃₅ /C ₃₄ HH	Homohopane Index	Ts/Tm	C30D/C29 Ts	C ₂₈ BNH/C ₃₀ hopane	C ₂₈ BNH/C ₂₉ norhopane
SR-230b	230.5	Soft	11.1	0.72	0.69	0.08	0.38	0.09	0.37	2.23
SR-227b	227.5	Soft	11.2	0.77	0.58	0.06	0.33	0.08	0.38	2.13
SR-224b	224.5	Soft	11.1	0.86	0.71	0.06	0.35	0.08	0.39	2.28
SR-197b	197.5	Soft	11.4	1.10	0.84	0.09	0.28	0.08	0.36	2.08
SR-194b	194.5	Soft	11.3	0.80	0.81	0.07	0.30	0.09	0.34	2.05
SR-226a	226	Hard	3.04	2.13	0.67	0.07	0.35	0.08	0.24	1.47
SR-219a	219	Hard	3.16	1.28	0.77	0.08	0.35	0.07	0.24	1.59
SR-205a	205	Hard	2.79	0.78	0.65	0.07	0.30	0.08	0.27	1.72

Table 14. Redox condition high resolution analysis of similar high TOC values distributed throughout the 40ft interval to distinguish variability. Lighter green tables indicating Middle Woodford data.

4.1.3 Marine vs. Lacustrine

Anoxic intervals could largely be restricted and create a marine environment to become pseudo lacustrine and therefore may contribute to the abundance of TOC. Heightened Hopane/C₂₇ $\alpha\alpha\alpha$ S+R values have indicated a lacustrine freshwater organic matter input (Mello et al., 1988; Araujo et al., 2018). The low Hopane/C₂₇ $\alpha\alpha\alpha$ S+R ratio value intervals are found within the soft bed group of Speake Ranch (~1-2). In accordance to the trend we have been observing regarding organic matter source input, it is expected that soft bed lithofacies would have relatively lower Hopane/ C_{27} $\alpha\alpha\alpha$ S+R ratio values. As throughout this study, soft bed lithofacies are found to be enriched in marine source input. A relatively higher Hopane/ C_{27} $\alpha\alpha\alpha$ S+R (3.46) can be found signifying more of a lacustrine freshwater input at SR-227b ft (227.5 ft) shown in Table 15. This outlier does not correlate to the variation of organic matter source input in Table 13 nor the variations in the redox parameters in Table 14. However, this outlier does correlate with an elevated DBT/Phen value and suggests that this value may be affected by the presence of sulfur in the water column. It is clear that the GI is consistent throughout both the hard bed group and soft bed group just as the DBT/Phen values shown in Table 15. From these 8 chosen samples, only one interval can be identified as pseudo lacustrine (SR-205a) in Table 15. Otherwise, there aren't any further and significant value distributions conclusive of lacustrine vs. marine implications within these 8 samples.

Sample	Depth (ft.)	TOC (wt%)	Lithology	Pr/Ph	TPP/C27 diasterane	Hopane/ C7αααS+R	C26/C25 TT	GI	DBT/Phen
SR-230b	230.5	11.1	Soft	0.72	0.22	1.07	0.84	0.16	0.54
SR-227b	227.5	11.2	Soft	0.77	0.22	3.46	0.78	0.15	0.80
SR-224b	224.5	11.1	Soft	0.86	0.19	1.61	0.93	0.13	0.74
SR-197b	197.5	11.4	Soft	1.10	0.12	1.34	0.97	0.14	0.56
SR-194b	194.5	11.3	Soft	0.80	0.13	1.72	0.94	0.13	0.59
SR-226a	226	3.04	Hard	2.13	0.28	6.04	0.94	0.14	0.38
SR-219a	219	3.16	Hard	1.28	0.31	5.64	0.87	0.15	0.39
SR-205a	205	2.79	Hard	0.78	0.23	3.04	0.96	0.14	0.49

Table 15. Lacustrine vs. Marine source high resolution analysis of similar high TOC values distributed throughout the 40ft interval to distinguish variability. Lighter green intervals indicate Middle Woodford data, and white intervals indicate Upper Woodford.

4.2 Evidence of Frasnian – Famennian Extinction Boundary

The Frasnian-Famennian extinction boundary is marked by the loss of a wide and diverse range of marine biotic organisms, such as brachiopods, rugose corals, coral stromatoporoids, pelagic conodonts, and benthic faunas within the shallow marine settings, similar to most extreme crises (Tulipani et al., 2015; Ma et al., 2016). This extreme loss of biodiversity within the Devonian time interval came to be known as a "reef gap", reflecting a transitional cause to the instigation of climate change from Devonian Greenhouse to the Carboniferous Icehouse (Zhang et al. 2019). Percival et al. (2016) suspected that the underlying cause for the loss of marine biodiversity was the widespread of marine anoxia and the consequential high rates of continental weathering, shown through the occurrence and appearance of black organic rich shales within the stratigraphic record. Ma et al. (2016) also believed that it was the combined forces of marine anoxia, continental weathering and the association of multiple volcanic/hydrothermal activities that caused the fluctuating and erratic climate cooling and warming which affected sea level dynamics. According to Wang et al. (2018), the Frasnian-Famennian boundary was marked by a preceding two-phase event called the "Kellwasser Event" and the "Annulata event", which signified the horizonal pulses leading up to the extinction. The Kellwasser Event is characterized by tectonic processes that include the large-scale volcanic activity associated with the Viluy Traps in Siberia, orogenic uplift, erosion, and the expansion of vascular rooted terrigenous flora (Wang et al., 2018). Alternatively, the Annulata horizon marked a great anoxic event that corresponded with a major marine transgression and global volcanic activity. The Late Devonian was not only recognized as a time marked by severe extinction, but it was also a time of evolution of the terrigenous forests and seed plants, which may have affected the dynamics of the ecosystem (Zhang et al. 2019). All of these processes that have been suspected to cause the wide extinction of marine biodiversity could possibly be traced by organic geochemical biomarker analysis.

The massive loss of marine biota did not occur abruptly, but rather in a transitional progress. Therefore, to identify the Frasnian-Famennian boundary, extreme climatic and environmental shifts need to be identified. This can be done by organic geochemical analysis

through biomarkers to track large shifts in organic matter input. It is possible that the influx of weathered terrigenous organic matter from a Transgression System Tract to a regression of sea level were introduced into the ocean and disturbed the marine organic productivity. Thus, stimulated algal blooms resulted in anoxic water columns and provided the mechanism that led to the end-Frasnian extinction (Philp and DeGarmo, 2020). It is also hypothesized that with such a high-resolution study over the Woodford interval from Speake Ranch, an accurate identification of the F-F boundary can be established. Moldowan et al. (1985) suggested that low values of Ts/Tm are an indication of different organic material which are often related to bacterial and/or terrigenous input. Not much can be deduced for the Ts/Tm of the Speake Ranch, as the ratio values are nearly linear and consistent low as previously discussed in Chapter 3. No significant shifts in organic matter source input can be dictated within the 40ft interval of the Woodford Shale. Biomarkers utilized in this section have been targeted to define signs of euxinia, distinct fluxes of terrigenous input along with algal blooms, and a shift in anoxia.

Previous studies from Nowaczewski (2011), Connock (2015), deGarmo (2015), Jones (2017), and Torres (2020) have made efforts to locate the Frasnian-Famennian (F-F) extinction boundary within the Woodford Shale using geochemical evidence. Nowaczewski (2011) indicated that there are biomarker anomalies which include elevated C₃₀ hopanes at the expense of C₂₉ hopanes, high abundance of gammacerane, and isorenieratane as markers to determine the F-F boundary. His findings demonstrated that the F-F boundary was a time marked by an increase in marine eukaryotic preservation within a stratified, anoxic marine water column. Connock (2015) and deGarmo (2015) indicated that the F-F boundary is marked by elevated H₂S presence and persistent PZE prior to marine biota loss. The increased nutrients from the PZE conditions and terrigenous blooms cause eutrophication and algal blooms. Moreover, Connock (2015) determined

that the F-F is located within the Middle Woodford at two occurrences of euxinic conditions of the Woodford. Both Over (1992) and Turner (2016) suggested that the F-F boundary to be just below the Middle Woodford and Upper Woodford contact boundary using lithology, chemostratigraphy, and conodont stratigraphy. Jones (2017) also indicated that the F-F boundary to be slightly below the Middle Woodford and Upper Woodford contact boundary as well. More specifically, Torres (2020) indicated that the F-F event to be located between 202ft to 220ft of the Speake Ranch section. The purpose of this research is to mark the exact interval of where this boundary could be located.

Schwark and Frimmel (2004) suggested a crossplot of the Pr/Ph ratio against aryl isoprenoid ratio to better assess the extent and persistence of photic zone anoxia during deposition of Jurassic Possidonian Shale. The distribution indicates the anoxic event coincides with facies changes as a reflection of rapid transgression. Grice et al. (2005) suggested that sulfide toxicity was a driving factor in the extinction and in the protracted recovery. Furthermore, indication of euxinia can be further assessed by the abundance of dibenzothiophene (DBT), where high contents of this compound in parallel to high values of aryl isoprenoids are suggestive of rapid input of terrigenous matter that consumed the oxygen within the water column and caused for further reduction optimal for DBT production (Mizukami et al., 2013). Combining all the geochemical parameters used to assess the paleoenvironment condition of the Ardmore Basin in Oklahoma, it seems that the F-F boundary is not a single bed but a transitional event, where the environment shifts with changing conditions. In agreement to Zhang et al. (2019), the F-F boundary is followed by preceding erratic environmental conditions. These are dictated by 3 intervals shown in Figure 55. The initial drop of GI, C₂₇/C₂₉ sterane, C₂₃/C₂₄ Tet, DBT/Phen, HHI, as well as enhanced Pr/Ph and TPP/C₂₇ diasterane could be a possible mark for the Maximum Flooding Surface (MFS) in

preparation for the F-F at 191-192ft. The MFS has been determined by prior studies by Galvis (2017) and Torres (2020) from their correlation of the Gamma Ray to the Speake Ranch section. However, due to the lack of data prior to MFS, the interval we have discovered as MFS may just be a preceding pulse of environmental instability leading up to the F-F boundary. The following highlighted event upsection is manifested based on the elevated and erratic fluctuation of values in Pr/Ph, HHI, AIR, TPP/C₂₇ diasterane, DBT/Phen, C₂₃TT/C₂₄Tet, and C₂₇/C₂₉ Sterane at 205.5ft to 215ft. Finally, the interval at 218.5ft to 227.5ft has shown significant instability and a transition to elevated and erratic values of Pr/Ph, homohopane index, TPP/C₂₇, GI, and AIR in correlation to a shift to decreasing C₃₅/C₃₄ HH and C₂₃ TT/C₂₄ Tet that serve as a marker of the Frasnian-Famennian boundary (Figure 55).



Figure 55. Organic geochemical parameters to define the anoxia, eutrophication, and organic matter input to determine the Frasnian – Famennian extinction boundary in the Woodford Shale of the Ardmore Basin in Oklahoma. Highlighted intervals signify the pulses leading up to the F-F.

Chapter V

5. Conclusion

Previous studies (Galvis, 2017; Torres, 2020) set lithostratigraphy and sequence stratigraphy frameworks for the Woodford Shale in southern Oklahoma at the Speake Ranch. The challenge posed by those two studies was the characterization of the Woodford shale at a smaller scale to observe the geochemical changes within the hard-to-soft cycles. This study is the first to explore a high resolution organic geochemical analysis to understand the effects of depositional processes, lithofacies, and stratigraphy over a 40ft interval.

Throughout the Upper and Middle Woodford, oscillations of soft and hard lithofacies are found at every 6 inches. Soft lithofacies, which are the clay abundant intervals, provide for a better petroleum source rock (high TOC) than the siliceous hard bed lithofacies. Maturity parameters $(T_{max}, C_{29} 20S/(20S+20R))$, and $C_{29} \beta\beta/(\beta\beta+\alpha\alpha)$ steranes) explored in this study show that the Speake Ranch samples are consistently immature. Therefore, variations in the biomarker distributions are independent of thermal alteration. There exist covariances of the lithofacies with biomarker distribution variations, which provide insight concerning the environmental deposition of the Woodford Shale from Speake Ranch. The organic matter source determined by C_{27}/C_{29} sterane, C₃₀ sterane index, and C₂₃ TT/C₂₄ Tet, illustrate a predominantly marine source input. Several intervals within the 40ft sequence have shown lower marine source input within the hard bed lithofacies and were thought to be derived from elevated terrigenous input. However, these elevated terrigenous markers identified by C₂₉ steranes, and C₂₄ tetracyclic terpanes may have been attributed to the input of marine algae (Tasmanites) instead. The retreating shoreline causes restriction in the water column causing CaCO₃ to supersaturate, thus stimulating algal blooms (Philp and DeGarmo, 2020). Torres (2020) also suggested that upwellings leading to chert

deposition provide nutrient-rich waters, which caused for algal blooms. The result of this study also documented the high-resolution analysis of biomarkers defining redox and euxinia of the depositional environment of the Woodford shale. It is anticipated that high sulfur content in the water column and high levels of anoxia are the optimal environmental conditions for organic matter preservation. Multiple biomarker parameters that are used to evaluate redox conditions, such as Ts/Tm, Pr/Ph, relative concentrations of bisnorhopanes, and homohopanes, illustrate that anoxia prevailed throughout soft bed lithofacies deposition. The combined parameters Pr/Ph, gammacerane, bisnorhopanes, tetracyclic polyprenoids, with and the dibenzothiophene/phenanthrene ratio, show evidence that sulfur rich intervals coincide with anoxic intervals along with providing a suitable environment for abundant preservation of organic matter. The analysis of elevated tetracylic polyprenoid concentrations combined with tricyclic terpanes, hopanes, steranes, and diasteranes reveal that there may have been three intervals within the Speake Ranch samples that became pseudo lacustrine depositional environments affecting total organic carbon yield.

A closer look into the biomarker distributions does attest to a major input of marine source. However, the analysis from XRF (Mo/Al) has also revealed that the fluctuations found within the organic matter source parameter indicate that bioturbated intervals yield lower organic carbon loading onto the sediment surface. This is due to remineralization and removal of organic carbon by oxygenation. In addition to that theory, mineral composition of the rock, whereby an abundance of clay is present, provides for optimal preservation properties. Therefore, variations in the abundance of organic matter preservation are controlled by varying levels of anoxia, water stratification, and mineral composition. A combined study of indicators for euxinia, distinct fluxes of terrigenous input along with algal blooms, helped define the extinction of the marine biota during the Late Devonian. This study explored the idea that the F-F extinction boundary was a transitional and gradual event that was preceded by two pulses of extreme conditions that preceded the extinction. No significant and definite shifts could be determined at the high-resolution sampling in this study. However, the events were signified by erratic distribution of the biomarkers within these intervals, indicating an instability of fluxes of organic matter source, anoxia, and euxinia.

In summary, the complex features of the unconventional Woodford Shale from Speake Ranch can be discerned using a high-resolution analysis that help integrate small scale environmental conditions to the sequence stratigraphy. However, the characteristics of the Middle and Upper Woodford members of this study may not reflect the Woodford Shale across Oklahoma, as this section may vary depending on the location. This could possibly be attributed to the enclosed mini-basin fill deposits nature of the section and may only be accurately compared to sections of other areas identified with the same nomenclature and interval thicknesses.

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Appendix I: Naphthalenes and Phenanthrenes

Naphthalenes and phenanthrenes are among the most common aromatic hydrocarbons in crude oils and can be used to determine thermal maturity of oils and extracts. When the relative abundance of 1,2,7-trimethylnaphthalene (14) predominates over the 1,2,5-trimethylnaphthalene (TMN; 17), this is an indication of early oil window or immature extract (Strachan et al., 1988; Akinlua et al., 2007; Philp and DeGarmo, 2020). The Speake Ranch samples illustrate that the 1,2,7-TMN predominates over the 1,2,5-TMN throughout the Middle Woodford. However, in the Upper Woodford abundance of 1,2,7-TMN is observed to significantly decrease, while 1,2,5-TMN predominates. This clearly suggests that the Middle Woodford is immature, while the Upper Woodford experiences higher thermal maturity. The 1,2,5-TMN and the 1,2,7-TMN are suggested to originate from constituents of angiosperms, such as the β -amyrin (Strachan et al., 1988; Ogala and Akegbobi, 2014). Furthermore, these compounds can form as diagenetic products of oleananetype triterpenoids (Chaffee and Johns, 1982; Chaffee et al., 1984; Ogala and Akegbobi, 2014). Golovko et al. (2014) indicated that the alkylated naphthalenes are less resistant to thermal alteration than the phenanthrenes and their alkylated derivatives. The thermal alterations of the methylnaphthalenes (MN) and dimethylnaphthalenes (DMN) occur upon removal of the n-alkanes (Fisher et al., 1998; Peters et al., 2005). Additionally, the trimethylphenanthrenes are altered during removal of isoprenoids, while tetramethylphenanthrenes remain unaltered until steranes are significantly depleted (Fisher et al., 1998; Peters et al., 2005). Naphthalenes and the alkylated naphthalenes were screened on the GC-MS at m/z 128, 142, 156, 170, 184, 198, 212, 226 (Figure 1; Table 1). Alkylnaphthalenes are not only a product of different genesis and different deposition ages, but they are also the products of alteration from varying organic matter sources. The effect of sedimentation environment and influence of catagenesis cause a secondary conversion of the alkylnaphthalenes into more stable molecules with substituents in the β -positions (Ogala and Akaegbobi, 2014; Golovko et al., 2014). Therefore, the relative changes in β - and α -isomers of the alkylnaphthalenes have been used to indicate thermal maturity as well. The stable 2-ethyl,3-methylnaphthalene ($\beta\beta$) relative to the less stable 1-ethyl,2-methylnaphthalene ($\alpha\beta$) ($\beta\beta$ -/ $\alpha\beta$ -ethyl-methylnaphthalene) is largely enhanced throughout the samples of Speake Ranch, and is also seen to frequently oscillate (Figure 2). This indicates that secondary conversions of the alkylnaphthalenes to the more stable β -positions had resulted, and therefore illustrates an increase in the degree of rock transformation.



Retention time

Figure 1. Distribution of the naphthalenes and the alkylated naphthalenes screened on the GC-MS at m/z m/z 128, 142, 156, 170, 184, 198, 212, 226.

Peak No	Compound	Peak No	Compound
110.		110.	
1	1-Ethylnaphthalene	16	1,2,4-Trimethylnaphthalene
2	2,6-Dimethylnaphthalene	17	1,2,5-Trimethylnaphthalene
3	2,7-Dimethylnaphthalene	18	1,3,5,7-Tetramethylnaphthalene
4	1,3+1,7-Dimethylnaphthalene	19	1,3,6,7-Tetramethylnaphthalene
5	1,6-Dimethylnaphthalene	20	1,2,4,7-Tetramethylnaphthalene
6	1,5-Dimethylnaphthalene	21	1,2,5,7-Tetramethylnaphthalene
7	1,2-Dimethylnaphthalene	22	2,3,6,7-Tetramethylnaphthalene
8	2-Ethyl,3-methylnaphthalene ($\beta\beta$)	23	1,2,6,7-Tetramethylnaphthalene
9	1-Ethyl,2-methylnaphthalene ($\alpha\beta$)	24	1,2,3,6-Tetramethylnaphthalene
10	1,3,7-Trimethylnaphthalene	25	1,2,5,6-Tetramethylnaphthalene
11	1,3,6-Trimethylnaphthalene	26	Dibenzothiophene
12	1,4,6-+1,3,5-Trimethylnaphthalene	27	4-Methyldibenzothiophene
13	2,3,6-Trimethylnaphthalene	28	3-+2-Methyldibenzothiophene
14	1,2,7-Trimethylnaphthalene	29	1-Methyldibenzothiophene
15	1,4,7+1,2,7-Trimethylnaphthalene	30	Dimethyldibenzothiophene

Table 1. Peak identifications for m/z 128, 1492, 156, 170, 184, 198, 212, 226 of naphthalenes and alkylnaphthalenes.

The naphthalene and its alkylated derivatives have also been used as a proxy for biodegradation. The level of biodegradation experienced can be seen using DBR [DBR=1,6-dimethylnaphthalene (5)/1,5-dimethylnaphthalene (6)], TBR [TBR= 1,3,6-trimethylnaphthalene (11)/1,2,4-trimethylnaphthalene (16)], and TeBR [TeBR= 1,3,6,7-tetramethylnaphthalene (19)/1,3,5,7-tetramethylnaphthalene (18)] ratios developed by Fisher et al. (1998). The reduction of the ratio values should indicate rising levels of biodegradation according to Fisher et al. (1998). There is an apparent oscillation from 190-194ft across DBR, TBR, and TeBR depth plots, whereby lower values are found within the harder beds, and relatively higher values are observed at the softer beds of Speake Ranch (Figure 2). The reduction of the DBR, TBR and TeBR values within the harder beds (190-194ft) suggests that these intervals experienced relatively higher levels of biodegradation than their overlying and underlying softer beds. This confirms the earlier finding

that utilized the XRF data of Mo/Al to determine weathering, where hard bed lithofacies experienced higher level of weathering. The DBR, TBR, and TeBR show an interesting trend whereby at the contact of the Upper and Middle Woodford, a significant shift is found. Values are seen to shift to lower values in DBR and TBR within the Upper Woodford, while values in TeBR experienced an elevated shift. The decreasing shift in values for DBR and TBR indicate that significant level of biodegradation occurred in the Upper Woodford.



Figure 2. Depth plots of the biodegradation parameters utilizing the alkylated naphthalenes. DBR, TBR, TeBR, MNR, $\beta\beta$ - $/\alpha\beta$ -ethyl-methylnaphthalene ratio and 1,2,7/1,2,5-TMN.

It was earlier discussed that relative concentrations of the alkylnaphthalenes generally reflect the presence of gymnosperms and terrigenous input into the system. Resins of higher plants, bryophytes, containing cadalene carbon skeleton convert to DMN and TMN with catagenetic effects, thus substitutions of the alkylnaphthalenes also make a reliable indicator of terrigenous input (de Leeuw et al., 1991). Golovko et al. (2014) suggested that marine organic matter derived oils are dominated by DMN over the MN, TMN, and naphthalenes. Additionally, he also suggested that a predominance of TMN over DMN, MN, and naphthalenes is found within continental oils, suggesting terrigenous source matter input (Golovko et al., 2014). The ternary plot of DMN-TMN-TeMN, reveal that the Speake Ranch is depleted with the DMN concentration, and has relatively higher values of TMN and TeMN (Figure 3). This suggests a predominant terrigenous source input. However, this does not align to the organic matter source finding that we utilized using sterane and terpane biomarkers. These values had possibly experienced alteration from weathering as well at the removal of n-alkanes as previously discussed.



Figure 3. Ternary plot of DMN, TMN, and TeMN to determine relative concentrations of the alkylated derivatives of the naphthalenes.

Phenanthrenes behave in a similar manner to the naphthalenes with increasing alkylation. Though, they are more resistant to biodegradation than alkylnaphthalenes (Philp and deGarmo, 2020). The abundance of phenanthrenes over the alkyl derivatives, methylphenanthrenes, is suggested to be attributed to pyrolysis and combustion derived PAH illustrating a source from natural fires (Youngblood and Blumer, 1975). The distribution of the phenanthrenes and their alkylated derivatives are screened on the GC-MS at m/z 178, 192, 206, 220, 234 (Figure 4). Therefore, phenanthrene/methylphenanthrene (phenanthrene/3-+2-+9-+1-methylphenanthrene) ratio has widely been used as a parameter to evaluate level of weathering (Radke et al., 1987). It is suggested that with increasing oxidation within the system, the phenanthrenes are thought to be more stable than their methylated derivatives (Clayton and King, 1987 Gieskes et al., 1990; Speczik et al., 1995; Marynowski and Wyszomirski, 2008). Thus, an increase in the phenanthrene/methylphenanthrene ratio is an indication of weathering, possibly a modern surface weathering from sea level regression or paleoweathering. An overall increase towards Upper Woodford at Speake Ranch does suggest that weathering was enhanced approaching the Upper Woodford (Figure 2). However, the phenanthrene/methylphenanthrene illustrates a shift to depletion in values in the Upper Woodford and this is possibly due to the predominating methylphenanthrene reflecting paleoweathering. An important factor to note is that the effect of paleoweathering by enhanced concentration of methylphenanthrene may be accurate if the mechanism of its generation is controlled by sedimentary methylation (Alexander et al., 1995; Jiang et al., 1998).



Figure 4. Phenanthrene and methylphenanthrene distribution screened on the GC-MS at m/z 178, 192, 206, 220, 234.

Appendix II: Compound Structures



I: C₁₇ n-Alkane









III: Phythane

IV: C₂₇ Sterane

V: C₂₈ Sterane



VI: C₂₉ Sterane



VII: C₃₀ Sterane



VIII: C₂₇ Diasterane



IX: Hopane



X: Tricyclic Terpane



XI: Gammacerane



XII: C₂₈ Bisnorhopane



XIII: C₂₉ Norhopane



XIV: Paleorenieratane



XV: Isorenieratane



XVI: Renieratane



XVII: 2,3,6-TMB



XVIII: 3,4,5-TMB

XIX: Tetracyclic Polyprenoid (C₃₀)



XX: Dibenzothiophene



XXI: Phenanthrene

XIII: Tetracyclic Polyprenoid (C₃₀)

Appendix III: Abbreviations and formulas for calculation of geochemical biomarkers

Maturity Parameters	Formula
$C_{29} 20S/(20S + 20R)$	C ₂₉ [14 α ,17 α -Stigmastane (20S)]/[14 α ,17 α -Stigmastane (20S+20R)]
$C_{29} \beta\beta/(\beta\beta+\alpha\alpha)$	C ₂₉ [14 β ,17 β -Stigmastane (20S+20R)]/[14 β ,17 β -+14 α ,17 α -Stigmastane (20S+20R)]

Organic Matter Source	Formula	
$C_{27}\%, C_{28}\%, C_{29}\%$	C_{27}, C_{28}, C_{29} [$C_{27} = C_{27} \alpha \alpha \alpha 20R / (C_{27} \alpha \alpha \alpha 20R + C_{28} \alpha \alpha$	
	C ₂₉ ααα 20R Steranes]	
C27/C29 Steranes	$(C_{27} \alpha \alpha \alpha (S+R) + C_{27} \alpha \beta \beta (S+R))/(C_{29} \alpha \alpha \alpha (S+R) + C_{29})$	
	$\alpha\beta\beta$ (S+R))	
Sterane Index	$C_{30}/(C_{27}-C_{30}) = [C_{30} \ 14\alpha, 17\alpha + 14\beta, 17\beta \ \text{Steranes} \ (20\text{S}+20\text{R})]/\Sigma C_{27},$	
	C_{28} , C_{29} , C_{30} [14 α ,17 α - + 14 β ,17 β Steranes (20S+20R)]	
C ₂₃ TT/C ₂₄ Tet	C ₂₃ Tricyclic terpane/C ₂₄ Tetracyclic terpane	
C ₂₆ /C ₂₅ TT	C_{26}/C_{25} Tricyclic terpanes	
C ₃₀ Hopane/C ₂₇ Diasterane	C ₃₀ Hopane/ C ₂₇ 13β,17α-Diacholestane	
C ₃₀ TPP/C ₂₇ Diasterane	C_{30} Tetracyclic Polyprenoid/ C_{27} 13 β ,17 α -Diacholestane	
C ₂₇ /total steranes	$(C_{27} \alpha \alpha \alpha (S+R) + C_{27} \alpha \beta \beta (S+R)) / (C_{27}-C_{30} \text{ Steranes})$	

Environmental Indicators	Formula
	C ₃₅ 17α,21β-Pentakishomohopane (22S+22R)/C ₃₄
C35/C34 HH	17α , 21β -Tetrakishomohopane (22S+22R)
	$C_{35}/(C_{31} - C_{35}) = C_{35} \ 17\alpha, 21\beta$ -Pentakishomohopane
Homohopane Index (%)	$(22S+22R)/[C_{31}+C_{32}+C_{33}+C_{34}+C_{35} 17\alpha,21\beta$
_	Homohopanes (22S+22R)]
AID	$(C_{13}-C_{17})/(C_{18}-C_{22})$ 2,3,6-Trimethyl substituted aryl
AIR	isoprenoids
Gammacerane Index	Gammacerane/ C ₃₀ Hopane
Pr/Ph	C ₁₉ Pristane/C ₂₀ Phytane
To/Tm	18α-22,29,30-Trisnorneohopane / 17α-22,29,30-
1 \$/ 1111	Trisnorhopane
C ₃₀ Diahopane/C ₂₉ Ts	C_{30} 17 α Diahopane/ C_{29} Ts
C ₂₈ BNH/C ₃₀ Hopane	C ₂₈ Bisnorhopane/C ₃₀ Hopane
C ₂₈ BNH/C ₂₉ Norhopane	C ₂₈ Bisnorhopane/C ₂₉ Hopane
C Amliconnerid	C_{18} 2,3,6-Trimethyl substituted aryl isoprenoids/ C_{18}
C ₁₈ -Aryi isopienolu	3,4,5-Trimethyl substituted aryl isoprenoids

Lithology Proxies	Formula	
C ₂₇ Dia/Sterane	$[C_{27} 13\beta, 17\alpha$ -Diacholestane (20S+20R)]/ $[C_{27} 14\alpha, 17\alpha$ - + 14 β ,17 β -Cholestane (20S+20R)]	
Diasterane/Steranes	$[C_{27} 13\beta, 17\alpha + 13\alpha, 17\beta - Diacholestane (20S+20R)]/ [C_{27} 14\alpha, 17\alpha + 14\beta, 17\beta - Cholestane (20S+20R)]$	
C ₂₇ /C ₂₉ diasteranes	$[C_{27} 13\beta, 17\alpha$ -Diacholestane 20R + 13α, 17β- Diacholestane 20S]/[24-Ethyl-13β, 17α- Diacholestane 20R + -13α, 17β-Diacholestane 20S]	

APPENDIX IV

GC-Chromatograms

TAP759



TAP751























TAP765



TAP767













TAP 772













TAP724



SATURATES M/Z 191

SR-190b



SR-191b







SR-193b



SR-194b



SR-195b


SR-197b



SR-198a







SR-200b









SR-208b







SR-210b



SR-211b







SR-213







SR-221b







SR-225b





SR-227b





SR-229a





WDFD SATURATES M/Z 217

SR-190b





SR-192a









SR-199b







SR-207a















































SR-194b











SR-198a





SR-200b




SR-205a





SR-209a





SR-211b





SR-213b







SR-219a





SR-222a





SR-225b





SR-227b











WDFD M/Z 178.3, 192.3, 206.3

SR-190b





SR-191b











SR-199b





SR-204b





SR-207a





SR-209a







SR-213b







SR-219a





SR-222a









