

OPEN

Received: 17 May 2018 Accepted: 7 November 2018

Published online: 30 November 2018

Improved Kerogen Models for Determining Thermal Maturity and Hydrocarbon Potential of Shale

Vikas Agrawal (1) & Shikha Sharma

Kerogen is the insoluble component of organic-rich shales that controls the type and amount of hydrocarbons generated in conventional and unconventional reservoirs. Significant progress has recently been made in developing structural models of kerogen. However, there is still a large gap in understanding the evolution of the molecular components of kerogen with thermal maturation and their hydrocarbon (HC) generative potential. Here, we determine the variations in different molecular fragments of kerogen from a Marcellus Shale maturity series (with VRo ranging from 0.8 to 3) using quantitative 13C MultiCP/MAS NMR and MultiCP NMR/DD (dipolar dephasing). These molecular variations provide insight into the (1) evolution of the molecular structure of kerogen with increasing thermal maturity and, (2) the primary molecular contributors to HC generation. Our results also indicate that old model equations based on structural parameters of kerogen underestimate the thermal maturity and overestimate the HC generation potential of Marcellus Shale samples. This could primarily be due to the fact that the kerogen samples used to reconstruct old models were mostly derived from immature shales (VRo <1) acquired from different basins with varying depositional environments. We utilized the kerogen molecular parameters determined from the Marcellus maturity series samples to develop improved models for determining thermal maturity and HC potential of Marcellus Shale. The models generated in this study could also potentially be applied to other shales of similar maturity range and paleo-depositional environments.

The advent of unconventional shale gas drilling has necessitated the need to develop a better understanding of the spatiotemporal variations in type and quality of organic matter in shale source rocks. Kerogen is a high molecular weight organic matter (OM) that serves as source and reservoir of all the hydrocarbons in these shales. Kerogen is formed by the degradation, condensation, and polymerization of biomolecules contributed by different sources of OM^{1-3} . Kerogen formed in the diagenetic stage of burial later cracks to form oil and gas in catagenetic and metagenetic stages of burial. The type and amount of HC generated, its sorption/retention and release on hydraulic fracturing operations is controlled by the molecular structure of kerogen³⁻⁶. Therefore, it is critical to understand the chemical structure of kerogen at the molecular level. In addition, molecular parameters of kerogen can serve as a more robust proxy for determining thermal maturity and hydrocarbon potential in mature shales (with VRo >1) where traditional techniques such as vitrinite reflectance, SRA, and biomarker analysis fails.

Significant progress has been made in understanding the molecular structure of kerogen using destructive (pyrolytic) and non-destructive (spectroscopic) methods⁷⁻¹⁰. However, the results generated using pyrolytic experiments can be biased due to the interaction of products generated from the labile fraction of kerogen¹⁰ or because reactions taking place in laboratory conditions might not be representative of sedimentary basin conditions¹¹⁻¹⁴. Due to these limitations, non-destructive methods such as Fourier transform infrared (FT-IR), Raman spectroscopy (RS), X-ray photoelectron spectroscopy (XPS), X-ray absorption near edge structure (XANES), and ¹³C solid-state nuclear magnetic resonance (¹³C NMR) have been employed for the qualitative, semi-quantitative, and quantitative measurements of kerogen^{7,15-21}.

The most reliable and robust tool for determination of fractions of molecular components of kerogen is ¹³C solid state NMR²². Numerous studies have been conducted using ¹³C solid-state NMR for kerogen characterization^{15–17,19,23–29}. A few recent attempts have been made to develop realistic structural models of kerogen^{30,31}. However, the evolution of different molecular components of kerogen on thermal maturation and the primary contributors to HC generation are still not well understood, especially for mature shales. A recent study by

Department of Geology and Geography, West Virginia University, Morgantown, WV, United States. Correspondence and requests for materials should be addressed to S.S. (email: shikha.sharma@mail.wvu.edu)

Sample ID	TOC	S2	VRo	Sample ID	TOC	S2	VRo
BG-1 UM ⁱ	4.68	5.11	1	WV-6 LM ⁱⁱ	9.10	0.47	2.5
BG-1 LM ⁱ	15.40	15.84	0.81	BL-3H UM	5.31	0.06	2.93
WV-7 UM ⁱⁱ	3.13	1.41	1.4	BL-3H LM	9.24	0.15	2.96
WV-7 LM ⁱⁱ	12.91	10.67	1.4	MIP-3H UM	4.14	0.12	2.94
MW-1 UM	8.63	1.84	1.49	MIP-3H LM	8.86	0.42	2.98
MW-1 LM	7.45	1.38	1.61	MIP 3H MT	3.14	0.08	2.92
WV-6 UM ⁱⁱ	3.52	0.02	2.5	MIP 3H MM	6.64	0.30	2.96
				MIP 3H MO	5.35	0.26	2.97

Table 1. The TOC, S2 and VRo values of samples selected for from 6 different Marcellus Shale wells in the Appalachian basin. ⁱValues from *Agrawal and Sharma*, 2018¹⁹. ⁱⁱValues from *Agrawal and Sharma*, 2018³³.

Agrawal and Sharma, 2018¹⁹ indicated that kerogen structural parameters used in previous models for determining HC generative potential²⁸ and thermal maturity^{26,32} could over or under estimate these values.

In this study, we determine the stability/reactivity of different molecular fragments of kerogen with thermal maturation and predict the primary contributors of hydrocarbons in a Marcellus shale maturity series (VRo ranging from 0.8 to 3). The variations in molecular parameters of kerogen were determined using quantitative MultiCP/MAS NMR and MultiCP NMR/DD (dipolar dephasing). Most of the previous investigations used CP/MAS NMR technique, a semi-quantitative technique that could not differentiate between non-protonated carbons (from protonated carbons) and mobile groups (from immobile groups). However, in this study, by using MultiCP/MAS NMR with and without dipolar dephasing, it was possible to determine the fraction of non-protonated carbons, protonated carbons, mobile groups and immobile groups (Supplementary Table 1). The correlation of structural components of kerogen with maturation and HC potential parameter (S2), was used to develop regression models (linear and multiple) for accurate estimation of thermal maturity and hydrocarbon generated. The structural parameters of 15 kerogen samples were determined using ¹³C multiple CP/MAS and multiple CP/MAS plus dipolar dephasing technique detailed in Agrawal and Sharma, 2018¹⁹. A total of 15 samples were obtained from six Marcellus shale wells (BG-1, WV-7, WV-6, MIP-3H, MW-1 and BL-3H) across a thermal maturity gradient in Appalachian basin (Table 1). TOC, S2 and VRo values of six samples were obtained from Agrawal and Sharma^{19,33} as indicated in Table 1.

Results and Discussion

Different aliphatic and aromatic carbon chains have characteristic chemical shifts in an NMR spectra^{34–37} as shown in Table 1 of Supplementary Information. Dipolar dephasing method with multiple CP method allows the quantification of mobile aliphatic and protonated (and non-protonated) aromatic carbon chains along with other aliphatic and aromatic chains^{8,22,38–40} as shown in Table 1 of Supplementary Information. The aliphatic and the aromatic fraction in the NMR spectra lies in 0-90 ppm and 90-165 ppm chemical shift range respectively. The fractions of different aliphatic and aromatic chains of kerogen were calculated using the peak area of the respective chemical shifts in an NMR spectra. The fractions of aliphatic carbon, alkyl (without heteroatoms), methoxy and amine, O and O₂ substituted alkyl carbons (ether and dioxy alkyl), total aromatic carbon, alkyl substituted aromatic, O-substituted aromatic (phenol), carboxyl and amide, aldehyde and ketone were determined using the multiple CP method (without dipolar dephasing). However, the fractions of mobile (freely rotating) and immobile (restricted rotation) methyl, mobile and quaternary alkyl (without heteroatoms), methoxy, protonated aromatic, non-protonated aromatic bridgehead carbon (fa^B) were determined using dipolar dephasing method with multiple CP (Table 1 in the Supplementary Information). Using these aliphatic and aromatic structural parameters, several lattice structural parameters such as average aliphatic carbon chain length (Cn'), mole fraction of bridgehead aromatic carbon(Xb), and SP²/SP³ hybridized carbon ratio were determined^{17,34,41}. The fractions of different structural parameters of 15 kerogen sample used in this study are shown in Table 1 of the Supplementary Information. Data of structural parameters of six samples are taken from Agrawal and Sharma¹⁹ (as indicated in Table 1 of the Supplementary Information). Multiple linear regression plots were made to determine the correlation of different structural parameters with VRo and S2. (refer to Figs 1-4 in Supplementary Information).

Kerogen models for thermal maturity. Thermal maturity is one of the most important parameter required for accurate determination of the hydrocarbon generated by source rocks. The traditional methods used for determining thermal maturity are the Tmax measurement using Source Rock Analyzer (SRA) and vitrinite reflectance measurement. However, Tmax may not be reliable in mature and over-mature samples⁴², and vitrinite is present as a major maceral only in Type III kerogen. Errors are involved in measuring reflectance on macerals other than vitrinite⁴³ or measuring reflectance of vitrinite whose thermal maturation is different from that of the bulk OM³. Biomarker ratios have also been used to determine the thermal maturity of source rocks⁴². However, due to thermal degradation and alteration of biomarkers on maturation^{33,44} and low extraction efficiency in high maturity samples, the results can be biased. Recent advancement in ¹³C solid-state NMR spectroscopic analysis, has led to an accurate quantification of different aliphatic, aromatic and lattice parameters of kerogen, even in over-mature shales¹⁹. Understanding the changes in these parameters on maturation can provide an important tool for determining thermal maturity in a broader maturity range.

In the Marcellus Shale the evolution of most of the structural parameters of kerogen such as immobile alkyl without heteroatoms (with restricted rotation), mobile (freely rotating), and alkyl-substituted aromatic carbons¹⁹

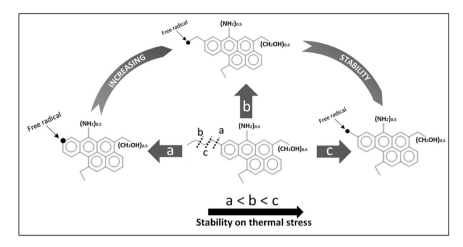


Figure 1. Stability of free radicals formed on thermal degradation. Structure of unit kerogen taken from Agrawal and Sharma, 2018¹⁹.

are primarily controlled by thermal maturity. Similar observations have also been made for shales from other basins basin^{17,23,28}. However, the cracking mechanism of different structural parameters of kerogen can vary because of the difference in their thermal stability. To determine the sensitivity of different kerogen structural parameters with thermal maturation, linear correlation plots were made using the structural parameters of kerogen and calculated vitrinite reflectance (Figs 1 and 2 of Supplementary Information). It has been previously shown that maturation leads to a decrease in aliphatic and an increase in aromatic carbon chains 17,23,26,28. However, the sensitivity and reactivity of carbon chains within the different aliphatic and aromatic carbon fractions are still not understood. Our results show that among different aliphatic chains, immobile alkyl chains are most prone to thermal degradation ($R^2 = 0.94$), followed by mobile and quaternary alkyl carbon chains ($R^2 = 0.61$) and mobile methyl carbon chains (R² = 0.54, Fig. 1 of Supplementary Information). In the alkyl carbon chains (without heteroatoms) the immobile methyl group is most resistant to breakdown during maturation ($R^2 = 0.20$). Immobile alkyl chains are mostly attached to the aromatic rings, restricting their rotation. The resistant nature of the immobile methyl group is possibly due to the instability of free radicals formed on thermal degradation (breaking of bond "a" in Fig. 1). The instability of the free radical is due to the disruption of resonance present the aromatic chains. However, for immobile alkyl groups, free radicals formed by thermal degradation (breaking of bond "c" in Fig. 1) is stabilized by the resonance of conjugated double bonds of the aromatic ring, making it the most stable amongst all the other aliphatic carbon chains.

In contrast, to the aliphatic structural parameters, the aromatic structural parameters showed relatively poor correlation to thermal maturation. Amongst the aromatic structural parameters, the highest correlation is observed between bridgehead aromatic carbons ($R^2 = 0.48$) followed by protonated aromatic carbon ($R^2 = 0.06$), O- substituted aromatic carbon ($R^2 = 0.06$), and alkyl substituted aromatic carbon chains ($R^2 = 0.02$). However, it is important to note that although individual aromatic parameters showed poor correlation with the vitrinite reflectance (Fig. 2 of Supplementary Information) the correlation of total aromatic with VRo is still significant ($R^2 = 0.90$). This indicates that good correlation of total aromatic carbon with increasing maturity as determined by several previous studies is actually due to increase in the relative percentage of aromatic carbon due to the higher breakdown of aliphatic carbon chains rather than formation of new aromatic carbons. The similar mole fraction of bridgehead aromatic carbon (Xb) in all mature kerogen samples (VRo >1, Fig. 2 in Supplementary Information) also supports that the total amount aromatic clusters do not increase significantly with maturity. This observation is contrary to the previous studies done on lower maturity samples 26,32 .

Regression models have been developed to determine thermal maturity using aromatic carbon percentage, the mole fraction of bridgehead carbon (Xb), SP² to SP³ carbon ratio 26,32,45 . In this study, we observed that the correlation coefficient of immobile alkyl chains (CnH₂n groups, with n > 1) was highest with R² = 0.94 (95% confidence interval or CI of slope = ± 2.99) followed by alkyl chains (immobile and mobile CnH₂n groups, with n > 1) with R² = 0.91 (95% CI of slope = ± 2.82), then by total aromatic carbon (R² = 0.90, 95% CI of slope = ± 1.89) and by total alkyl (without heteroatoms) with R² = 0.87 (95% CI of slope = ± 2.54) as shown in Fig. 2. Based on these observations, we propose four new regression models for estimating thermal maturity in shales with VRo between 0.8 to 3.0 (all equations have R² > 0.85). The correlation coefficents and 95% confidence interval (CI) of the slope of the equations proposed for determining VRo were determined using software XLSTAT. The equations are as follows:

$$VRo = 3.39 - 19.32 * IA$$
 (1)

where IA is fraction of immobile alkyl chains (CnH₂n groups) with n >1.

$$VRo = 3.41 - 14.74 * TA_2$$
 (2)

where TA_2 is fraction of total alkyl chains (CnH₂n groups) with n >1.

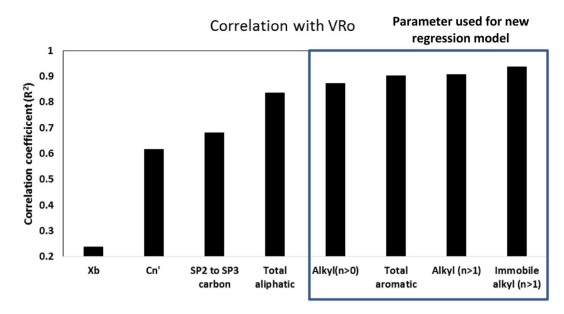


Figure 2. Correlation of different kerogen structural parameters with calculated vitrinite reflectance.

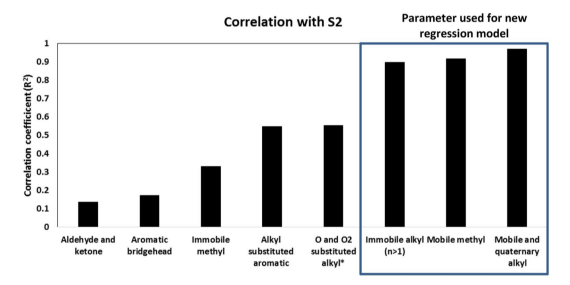


Figure 3. Correlation of different structural parameters of kerogen with true HC potential (S2).

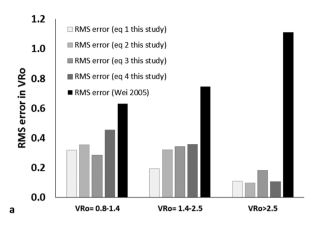
$$VRo = -5.84 + 9.69 * F_{ar}$$
 (3)

where F_{ar} is fraction of total aromatic chains.

$$VRo = 3.55 - 11.05 * TA$$
 (4)

where TA are total alkyl chains (CnH_2n groups) with n > 0.

Kerogen models for hydrocarbon potential. A recent study done on kerogen from Marcellus Shale by Agrawal and Sharma, 2018^{19} indicates that previously build kerogen models used for determining HC generative potential overestimate the S2 values at least by a factor of two. This can lead to underestimation of the total amount of HC generated in the reservoir. We determined primary contributors to HC generation by evaluating the correlation of different structural parameters of kerogen with S2. We observed that the carbon chains that had highest correlation coefficient with S2 were mobile and quaternary alkyl group ($R^2 = 0.97, 95\%$ CI = ± 0.32), mobile methyl group ($R^2 = 0.92, 95\%$ CI = ± 0.55) and immobile alkyl chains ($R^2 = 0.90, 95\%$ CI = ± 0.17) (Fig. 3 in the Supplementary Information). Additionally, the relatively smaller correlation coefficient of immobile methyl, aromatic bridgehead carbon, aldehyde and ketone groups, and O and O₂ substituted alkyl chains with S2 (Fig. 3), indicates there was little or no contribution from these structural components to the total hydrocarbons generation potential, contradicting the observations of Longbottom *et al.*²⁸. Therefore, we propose a new equation



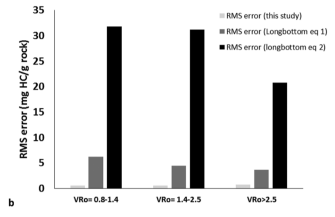


Figure 4. Comparison of RMS error of (a) thermal maturity and (b) HC generation, determined using models proposed in this study with previous kerogen models.

for determining the hydrocarbon potential that is based on a multiple regression model using the fractions of structural parameters: mobile and quaternary alkyl group, mobile methyl group and immobile alkyl chains:

$$S2 = 0.60 \times MM + 2.02 \times MA + 0.12 \times IA - 1.85 \tag{5}$$

where MM is mobile methyl chains (-CH₃ groups), MA is mobile alkyl chains (CnH₂n groups, n > 1) and IA is immobile alkyl chains (CnH₃n groups, n > 1).

The coefficient of determination (R^2) between the true S2 and modeled S2 was 0.98, and the RMS (root mean square) error of prediction was \pm 0.70 mg hydrocarbon per gram of rock. It was also, observed that the by adding additional structural parameters in the multiple regression model, the coefficient of determination (R^2) between true vs. modeled S2 did not improve. This observation further suggests that the structural parameters used in the model are the primary contributors of hydrocarbon potential.

Model validation. To validate the model proposed for determining thermal maturity, we determined the VRo values from the four equations proposed in this study and compared it with the true VRo values and values determined using previous regression model proposed by Wei *et al.*²⁶ (shown in Table 2 of the Supplementary Information). The VRo values predicted using the newly build regression model were comparable to the true values with RMS error in VRo for all the samples were 0.21, 0.26, 0.26, 0.31 using equations 1–4 respectively. However, the VRo value predicted using equations proposed by Wei *et al.*²⁶ underestimated the true values with a total RMS error 0.91 in VRo values (Fig. 4a, Table 2 of the Supplementary Information). The RMS error for predicting VRo in different maturity ranges using the proposed equations were also significantly less than the previously proposed model (Fig. 4a, Table 2 of the Supplementary Information).

To validate the model proposed for determination of hydrocarbon potential, we determined the total amount of HC generated using the modeled values of S2 and compared it with the true amount of HC generated. The true amount of total HC generated (in mg) per gram of rock is determined by equation $S2_{net} = S2_{org} - S2_{pd}$ (org stands for original and pd for the present day). $S2_{org}$ is calculated by the dividing the TOC_{org} by HI_{org} . TOC_{org} is calculated using equation given by Peters *et al.*⁴²:

$${\rm TOC_{org}} = 83.33~({\rm HI_{pd}})({\rm TOC_{pd}})/{\rm HI_{org}}~(1-f)(83.33-{\rm TOC_{pd}})~+~({\rm HI_{pd}})({\rm TOC_{pd}})$$

where $f=1-[\mathrm{HI_{pd}}\,(1200-(\mathrm{HI_{org}/1-PI_{org}})/\mathrm{HI_{org}}\,(1200-(\mathrm{HI_{pd}/I-PI_{pd}}]\,\mathrm{HI_{org}}\,\mathrm{obtained}$ for immature Marcellus shale is approximately 250 mg HC/g rock⁴⁶. Similar $\mathrm{HI_{org}}$ were obtained from HI vs. VRo plot (Y-intercept of the

curve Fig. 5 in Supplementary Information). PI $_{\rm org}$ is assumed to be 0.02 for the most immature source rocks⁴². The S2 values modeled in this study are the S2 $_{\rm present \, day}$. The total amount of HC generated calculated using the true S2 values and modeled values are shown in Table 3 in the Supplementary Information. We compared these values with true amount of HC generated and HC generated using previous regression models (Table 3 in Supplementary Information). The amount of HC generated predicted using the newly build regression model were comparable to the true values with RMS error for all the samples 0.70 mg HC/g rock. However, the amount of HC generated predicted using the two equations proposed by Longbottom *et al.*²⁸ underestimated the true values with RMS error 3.93 mg HC/g rock and 27.05 mg HC/g rock (Fig. 4b, Table 3 in Supplementary Information). The RMS error for predicting HC generated in different maturity ranges using the proposed model was also significantly less than the previously proposed model (Fig. 4b, Table 3 in Supplementary Information). The higher RMS error of older models could be attributed to the fact that they (1) utilized kerogen derived from shales that were all below VRo <1 and, (2) used shale samples acquired from different basins where the structure of kerogen might vary significantly due to variations in sources of OM and depositional conditions.

The major strength of new models proposed in this study is that they were generated using kerogen extracted from samples across the entire maturity range of hydrocarbon generation, and therefore more accurately represent the source rock in mature shale plays like Marcellus as compared to older models. Future work will focus on acquiring samples from other shales to test the efficacy of our models. It is also plausible that for more accurate estimation of HC generative potential and maturity, similar kind of models need to be developed for individual plays instead of utilizing generalized models.

Methods

13C solid-state NMR analysis. Solid-state NMR experiments were performed on a Bruker Advance III 400 spectrometer operating at 400-MHz 1H and 100-MHz 13 C frequencies at Environmental NMR Service at Old Dominion University in Norfolk, Virginia, USA. The 13 C chemical shifts were referenced to tetramethyl silane, using the COO resonance of glycine in the α-modification at 176.46 ppm as a secondary reference. Quantitative 13 C NMR spectra for all the kerogen samples were acquired using the high-spinning speed multi-ramped amplitude cross polarization/magic angle spinning technique developed by Johnson and Schmidt-Rohr⁴⁷. This multiple-cross polarization (multiCP) technique is a simple, robust way to obtain quantitative solid-state 13 C NMR spectra of kerogen, with good signal-to-noise ratio. The spectra were measured at a spinning speed of 14 kHz, where spinning sidebands are fairly small (<3%) and have little overlap with center bands. 13 C multiCP/MAS with dipolar dephasing was performed under the same conditions as for 13 C multiCP/MAS but combined with a dipolar dephasing time of 68 μs⁴⁰ to differentiate nonprotonated C from total C and to determine fractions of mobile groups (with no restricted rotation). The relative proportion of different carbon chains as shown in Supplementary Table 1 were obtained from 13 C NMR spectrum using an NMR peak fitting program *TopSpin* (area of spectra from 0–240 ppm was considered to be 100% as detailed in Agrawal and Sharma 19).

SRA analysis. Approximately 80 mg of powdered (200 mesh) shale sample was weighed into a SRA crucible and placed in the autosampler and held isothermally at 300 °C for 3 minutes. The free hydrocarbons are volatilized during this isothermal heating which is quantitatively detected by the FID detector and reported as milligrams (mg) of S1 per gram of rock. The free CO_2 is simultaneously liberated which is detected by the IR cell and reported as milligrams (mg) of S3 per gram of rock. The temperature is increased after the isothermal period at the rate of 25 °C/minute until 600 °C. Pyrolytic degradation of the kerogen takes place between 300 °C and 600 °C generating HCs. These hydrocarbons are also detected by the FID are labeled as S2, reported as mg of S2 per gram of rock. The temperature at the peak of S2 generation is known as Tmax. It is used to estimate the thermal maturity of shale samples. Vitrinite reflectance is calculated using equation (VRo) = $0.018 \times \text{Tmax} - 7.16^{48}$. Residual carbon is also measured by SRA and is reported as S4. TOC of the sample is calculated using the equation $0.1 \times [0.082 \times (\text{S1} + \text{S2}) + \text{S4}]$, in wt %. After every five sample WFT Source Rock Standard 533 (P/N 810-141) was run. SRA analysis was performed at the National Energy Technology Laboratory and at IsoBioGEM lab in Morgantown.

References

- Huc, A. & Durand, B. Etude des acides humiques et de l'humine de sédiments récents considérés comme précurseurs des kérogènes. in 53-72 (1974)
- 2. Tissot, B. P. & Welte, D. H. Petroleum Formation and Occurrence. (Springer-Verlag, 1984).
- 3. Vandenbroucke, M. & Largeau, C. Kerogen origin, evolution and structure. Org. Geochem. 38, 719–833 (2007).
- 4. Behar, F. & Vandenbroucke, M. Chemical modelling of kerogens. Org. Geochem. 11, 15-24 (1987).
- 5. Vandenbroucke, M. Kerogen: from Types to Models of Chemical Structure. Oil Gas Sci. Technol. 58, 243-269 (2003).
- 6. Ho, T. A., Criscenti, L. J. & Wang, Y. Nanostructural control of methane release in kerogen and its implications to wellbore production decline. Sci. Rep. 6, 28053 (2016).
- 7. Tong, J., Han, X., Wang, S. & Jiang, X. Evaluation of Structural Characteristics of Huadian Oil Shale Kerogen Using Direct Techniques (Solid-State 13C NMR, XPS, FT-IR, and XRD). Energy Fuels 25, 4006–4013 (2011).
- 8. Cao, X., Yang, J. & Mao, J. Characterization of kerogen using solid-state nuclear magnetic resonance spectroscopy: A review. *Int. J. Coal Geol.* **108**, 83–90 (2013).
- Cao, X. et al. Chemical structure changes in kerogen from bituminous coal in response to dike intrusions as investigated by advanced solid-state 13C NMR spectroscopy. Int. J. Coal Geol. 108, 53–64 (2013).
- Horsfield, B. Practical criteria for classifying kerogens: Some observations from pyrolysis-gas chromatography. Geochim. Cosmochim. Acta 53, 891–901 (1989).
- 11. Behar, F., Vandenbroucke, M., Tang, Y., Marquis, F. & Espitalie, J. Thermal cracking of kerogen in open and closed systems: determination of kinetic parameters and stoichiometric coefficients for oil and gas generation. Org. Geochem. 26, 321–339 (1997).
- 12. Lewan, M. D. & Ruble, T. E. Comparison of petroleum generation kinetics by isothermal hydrous and nonisothermal open-system pyrolysis. *Org. Geochem.* 33, 1457–1475 (2002).

- 13. Freund, H. et al. Predicting oil and gas compositional yields via chemical structure–chemical yield modeling (CS-CYM): Part 1 Concepts and implementation. Org. Geochem. 38, 288–305 (2007).
- Walters, C. C., Freund, H., Kelemen, S. R., Peczak, P. & Curry, D. J. Predicting oil and gas compositional yields via chemical structure-chemical yield modeling (CS-CYM): Part 2 – Application under laboratory and geologic conditions. Org. Geochem. 38, 306–322 (2007).
- 15. Dennis, L. W., Maciel, G. E., Hatcher, P. G. & Simoneit, B. R. T. 13C Nuclear magnetic resonance studies of kerogen from Cretaceous black shales thermally altered by basaltic intrusions and laboratory simulations. *Geochim. Cosmochim. Acta* 46, 901–907 (1982).
- Witte, E. G., Schenk, H. J., Müller, P. J. & Schwochau, K. Structural modifications of kerogen during natural evolution as derived from 13C CP/MAS NMR, IR spectroscopy and Rock-Eval pyrolysis of Toarcian shales. Org. Geochem. 13, 1039–1044 (1988).
- 17. Kelemen, S. R. et al. Direct Characterization of Kerogen by X-ray and Solid-State 13C Nuclear Magnetic Resonance Methods. Energy Fuels 21, 1548–1561 (2007).
- 18. Petersen, H. I., Rosenberg, P. & Nytoft, H. P. Oxygen groups in coals and alginite-rich kerogen revisited. *Int. J. Coal Geol.* **74**, 93–113 (2008).
- Agrawal, V. & Sharma, S. Molecular Characterization of Kerogen and its Implications for Determining Hydrocarbon Potential, Organic matter sources and Thermal Maturity in Marcellus Shale. Fuel. https://doi.org/10.1016/j.fuel.2018.04.053 (2018).
- Miknis, F. P., Smith, J. W. & Maughan, E. K. & Maciel (5), G. E. Nuclear Magnetic Resonance: A Technique for Direct Nondestructive Evaluation of Source-Rock Potential. AAPG Bull. 66, 1396–1401 (1982).
- 21. Miknis, F. P., Netzel, D. A., Smith, J. W., Mast, M. A. & Maciel, G. E. 13C NMR measurements of the genetic potentials of oil shales. *Geochim. Cosmochim. Acta* 46, 977–984 (1982).
- 22. Mao, J. et al. Chemical and nanometer-scale structure of kerogen and its change during thermal maturation investigated by advanced solid-state 13C NMR spectroscopy. Geochim. Cosmochim. Acta 74, 2110–2127 (2010).
- 23. Patience, R. L., Mann, A. L. & Poplett, I. J. F. Determination of molecular structure of kerogens using 13C NMR spectroscopy: II. The effects of thermal maturation on kerogens from marine sediments. *Geochim. Cosmochim. Acta* 56, 2725–2742 (1992).
- 24. Mann, Al, Patience, Rl & Poplett, I. J. F. Determination of molecular structure of kerogens using 13C NMR spectroscopy: I. The effects of variation in kerogen type. *Geochim. Cosmochim. Acta* 55, 2259–2268 (1991).
- Lille, Ü., Heinmaa, I. & Pehk, T. Molecular model of Estonian kukersite kerogen evaluated by 13C MAS NMR spectra*. Fuel 82, 799–804 (2003).
- 26. Wei, Z., Gao, X., Zhang, D. & Da, J. Assessment of Thermal Evolution of Kerogen Geopolymers with Their Structural Parameters Measured by Solid-State 13C NMR Spectroscopy. *Energy Fuels* 19, 240–250 (2005).
- Werner-Zwanziger, U., Lis, G., Mastalerz, M. & Schimmelmann, A. Thermal maturity of type II kerogen from the New Albany Shale assessed by 13C CP/MAS NMR. Solid State Nucl. Magn. Reson. 27, 140–148 (2005).
- 28. Longbottom, T. L. *et al.* Organic structural properties of kerogen as predictors of source rock type and hydrocarbon potential. *Fuel* **184**, 792–798 (2016).
- Longbottom, T. L., Hockaday, W. C., Boling, K. S. & Dworkin, S. I. Effect of ocean oxidation on the chemical structure of marine kerogen. Org. Geochem. 106, 1–12 (2017).
- 30. Bousige, C. et al. Realistic molecular model of kerogen's nanostructure. Nat. Mater. 15, 576-582 (2016).
- 31. Ungerer, P., Collell, J. & Yiannourakou, M. Molecular Modeling of the Volumetric and Thermodynamic Properties of Kerogen: Influence of Organic Type and Maturity. *Energy Fuels* 29, 91–105 (2015).
- 32. Wang, Z. & Cheng, K. Study on the Thermal Evolution Degree of Source Rocks Developed in Early Paleozoic or Older Eras. In Organic Geochemistry: Development and Applications to Energy, Climate, Environment and Human History 478–480 (1995).
- 33. Agrawal, V. & Sharma, S. Testing Utility of Organogeochemical Proxies to Assess Sources of Organic Matter, Paleoredox Conditions, and Thermal Maturity in Mature Marcellus Shale. Front. Energy Res. 6 (2018).
- 34. Solum, M. S., Pugmire, R. J. & Grant, D. M. Carbon-13 solid-state NMR of Argonne-premium coals. *Energy Fuels* 3, 187–193 (1989).
- 35. Kuangzong, Q., Deyu, C. & Zhanguang, L. A new method to estimate the oil and gas potentials of coals and kerogens by solid state 13C NMR spectroscopy. *Org. Geochem.* 17, 865–872 (1991).
- 36. Baldock, J. A., Masiello, C. A., Gélinas, Y. & Hedges, J. I. Cycling and composition of organic matter in terrestrial and marine ecosystems. *Mar. Chem.* 92, 39–64 (2004).
- 37. Smernik, R. J. & Oades, J. M. Solid-state 13C-NMR dipolar dephasing experiments for quantifying protonated and non-protonated carbon in soil organic matter and model systems. *Eur. J. Soil Sci.* 52, 103–120 (2001).
- 38. Mao, J.-D. et al. Humic acids from particulate organic matter in the Saguenay Fjord and the St. Lawrence Estuary investigated by advanced solid-state NMR. Geochim. Cosmochim. Acta 71, 5483–5499 (2007).
- 39. Mao, J.-D. & Schmidt-Rohr, K. Recoupled long-range C–H dipolar dephasing in solid-state NMR, and its use for spectral selection of fused aromatic rings. *J. Magn. Reson.* **162**, 217–227 (2003).
- Mao, J.-D. & Schmidt-Rohr, K. Accurate Quantification of Aromaticity and Nonprotonated Aromatic Carbon Fraction in Natural Organic Matter by 13C Solid-State Nuclear Magnetic Resonance. Environ. Sci. Technol. 38, 2680–2684 (2004).
- 41. Hockaday, W. C. et al. Measurement of soil carbon oxidation state and oxidative ratio by 13C nuclear magnetic resonance. J. Geophys. Res. Biogeosciences 114, G02014 (2009).
- 42. Peters, K. E., Walters, C. C. & Moldowan, J. M. *The Biomarker Guide*. (Cambridge University Press, 2005).
- 43. Durand, B. Indices optiques, potentiel pétrolier et histoire thermique des sédiments. in Alpern, B. (Ed.), Pétrographie de la Matière Organique des Sédiments, Relations avec la Paléotempérature et le Potentiel Pétrolier Editions du CNRS (Centre National de la Recherche Scientifique) 205–215 (1975).
- Farrimond, P., Taylor, A. & Telnæs, N. Biomarker maturity parameters: the role of generation and thermal degradation. Org. Geochem. 29, 1181–1197 (1998).
- 45. Lupoi, J. S. et al. Assessment of Thermal Maturity Trends in Devonian–Mississippian Source Rocks Using Raman Spectroscopy: Limitations of Peak-Fitting Method. Front. Energy Res. 5 (2017).
- 46. Bruner, K. R. & Smosna, R. A Comparative Study of the Mississippian Barnett Shale, Fort Worth Basin, and Devonian Marcellus Shale, Appalachian Basin. 118 (2011).
- Johnson, R. L. & Schmidt-Rohr, K. Quantitative solid-state 13C NMR with signal enhancement by multiple cross polarization. J. Magn. Reson. 239, 44–49 (2014).
- 48. Jarvie, D. M. & Lundell, L. Hydrocarbon generation modelling of naturally and artificially matured Barnett Shale, Fort Worth Basin, Texas. Southwest Reg. *Geochem. Meet. Sept.* 8–9 1991 Woodl. Tex. (1991).

Acknowledgements

The research was funded by Department of Energy's National Energy Technology Laboratory (DE# FE0024297; DE# FE0004000) and National Science Foundation (NSF DEB-1342732) grants to S. Sharma. The authors thank Dr. Jingdong Mao and Wenying Chu from Old Dominion University, VA for ¹³C NMR analysis and Dr. Ajay Warrier from the WVU IsoBioGem Lab for help with sample preparation. West Virginia Geological and Economic Survey, Northeast Natural Energy and Southwestern Energy are acknowledged for providing samples for the study.

Author Contributions

V.A. and S.S. were involved in initial conception of idea of the research. V.A. conducted the analytical work needed for study and drafted the manuscript. V.A. and S.S. were both involved in data interpretation, revising and editing of the manuscript.

Additional Information

Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-018-35560-8.

Competing Interests: The authors declare no competing interests.

Publisher's note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit https://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2018