Full Length Article

Swelling of Kimmeridge kerogen by normal-alkanes, naphthenes and aromatics

Liang Huang, Atefeh Khoshnood, Abbas Firoozabadi

ABSTRACT

Despite recognition of kerogen swelling in liquid solvents, swelling trends in different classes of petroleum liquids remain unexplored. Modeling results based on the extended polymer theory have not been beyond curve-fitting of swelling measurements. This work centers on measurement of swelling of Kimmeridge kerogen in three classes of petroleum liquids: normal-alkanes, naphthenes and aromatics. The measured data show that swelling decreases as the molecular size increases for the three classes of solvents. Compared with naphthenes, normal-alkanes with flexible linear chains, and aromatics having π–π interaction with kerogen induce higher swelling. In the Kimmeridge kerogen, the swelling in both normal-alkanes and naphthenes monotonously decreases with the solvent solubility parameter and molar volume starting with hydrocarbon molecules with 5 carbon atoms. Compared with solvent solubility parameter, the effect of solvent molar volume dominates the swelling of Kimmeridge kerogen. The extended Flory-Rehner and regular solution theory framework does not describe the swelling behavior of Kimmeridge kerogen in petroleum liquids despite widespread applications of the framework. Results from the model show an increase in swelling in normal alkanes with molecular size which is opposite to our measurements.

1. Introduction

Petroleum fluids are generated from the organic matter of shale. The organic matter which often consists of kerogen has a cross-linked network with a complex molecular structure [1]. The maturation of kerogen includes both the bond-breaking and bond-making reactions. The bond-breaking reactions lead to the conversion of organic carbon in kerogen skeleton into volatile hydrocarbons, bitumen and solid residue [2]; the bond-making reactions contribute to the increase of the cross-linked degree of the organic network [3]. Kerogen has abundant nanopores and large specific surface areas, which facilitates the storage of the generated hydrocarbons [4,5]. Expulsion of petroleum hydrocarbons, also called primary migration, is the initial stage of petroleum migration from the source kerogen to the reservoir accumulation. The process remains poorly understood [6,7]. Kerogen is in dynamic equilibrium with the generated hydrocarbons during the expulsion process.

The producible hydrocarbon and non-hydrocarbon species are in three states in kerogen: free molecules in the pores, adsorbed molecules on the surfaces and dissolved molecules in the kerogen network [8]. The expulsion process is accompanied by chemical fractionation where saturated hydrocarbons and aromatics are preferentially expelled compared to polar compounds and polycyclic aromatic hydrocarbons such as asphaltenes and resins [6,7]. There have been many investigations on the chemical fractionation during expulsion [6,7,9–15]. These investigations have emphasized the role of either the pairwise interaction or the solubility difference between solvents and organic network. The mechanisms provide the process direction but not magnitude of the chemical fractionation [15]. Physical principles like the effect of fluid molecular size on the chemical fractionation remain less understood than chemical principles.

Kerogen is a soft nanoporous matter with flexible molecular structure [16–18]. The generated petroleum compounds when dissolved in kerogen matrix may induce the swelling of kerogen. Swelling of the kerogen network by different liquid solvents is well-known; it may be used to characterize its physical structure and chemical nature [3,6]. Swelling of kerogen immersed in various liquid solvents has been...
measured by various authors [2,3,6,10,19–28]. A swelling as high as 47% has been reported for the Kimmeridge kerogen without bitumen extraction from exposure to heptane [29]. In addition to the effect of bitumen, the high swelling in Ref. [29] may be due to the sample preparation; the kerogen was compressed into cylindrical pellets in the swelling experiments. To the best of our knowledge, there is no comprehensive data set on the swelling of the same type of kerogen sample in various normal-alkanes, napthenes and aromatics. In the past, liquid solvents in the same class with similar properties are selected in the swelling experiments. Measurements have also been performed on different classes of kerogen samples. No general trend of kerogen swelling of a given class of petroleum liquids has been reported to the best of our knowledge.

The regular solution theory framework has been widely adopted to describe the solubility behavior of liquid solvents in cross-linked organic materials such as polymers [30,31], coals [32,33], asphaltites [34,35] and kerogens [22,23]. The applicability of the theory has been found to be sensitive to the type of solvent interacting with the organic materials. The basic behaviors of solvents that have nonspecific interaction with the organic materials have been assumed to follow the regular solution theory [22,23]. In contrast, the theory may not describe the behaviors of solvents that interact with the organic materials via specific interaction [31–33]. A more recent work has concluded that the regular solution theory may not describe the behavior of asphaltites in organic solvents [35]. The nonspecific interactions are dominated by the dispersion forces. The specific interactions are exhibited when there is strong electron density or a positive charge (e.g. cation exchange) located on the bonded or nonbonded links of the interacting partners [36]. A non-classical example for a specific interaction is the hydrogen bond, which is prevalent in polymer–solvent [31] and coal–solvent systems [33].

The regular solution theory has been combined with the Flory-Rehner theory to describe swelling of kerogen immersed in solvents [3,6]. Apparent observations indicate that the swelling of kerogen in organic solvents may be described within the framework of the extended Flory-Rehner and regular solution theory framework [3,6]. These observations are based on limited solvent data. Only a small number of similar solvent groups are used in each class of petroleum hydrocarbons. Fig. 1 shows the predicted swelling of kerogen in normal-alkanes based on the theory with the fitting thermodynamic parameters of kerogen from ref [6]. The solubility parameter, cross-link density, and native swelling fraction used in the computations are 22.5 J^{1/2} cm^{−3/2}, 0.16 mol/cm³ and 0.76, respectively for Type II kerogen, and 23.3 J^{1/2} cm^{−3/2}, 0.25 mol/cm³ and 0.85, respectively for Type III kerogen [6]. The swelling reveals an increasing trend as the number of carbon atoms increases in normal-alkanes. One objective of this work is to provide systematic measurements on swelling of kerogen in normal alkane over a broad range to assess the applicability of the extended Flory-Rehner and regular solution theory framework. We will show that the results in Fig. 1 are not consistent with experimental data. Note that Fig. 1 is presented to examine the theoretical model using documented thermodynamic parameters. In this work, we will further assess the applicability of the theoretical model based on our experimental data in Kimmeridge kerogen, which can facilitate the improvement or development of new models in the future.

A key objective of this work is to present systematic experimental data on swelling of Kimmeridge kerogen immersed in normal-alkanes, napthenes and aromatics. The effect of solvent solubility parameter, solvent molar volume, solvent molecular size and molecular shape on kerogen swelling will be investigated. The systematic data are used to examine the validity of the extended Flory-Rehner and regular solution theory framework in describing the swelling of kerogen. Results of this work can facilitate the development of more comprehensive mechanisms for the chemical fractionation in kerogen during petroleum expulsion.

2. Methodology

2.1. Kerogen

The shale source rock is an outcrop sampled from the Blackstone band of the Kimmeridge Clay Formation in Dorset, UK. The kerogen sample used in the swelling experiment was extracted from the source rock by acid treatment and Soxhlet extraction process. The source rock was crushed into millimeter pieces by hammer and pincer and then processed in a wig-L-bug to reduce particle size to less than 200 mesh (74 µm). A 200-mesh sieve was used to separate particles of different sizes. The sample powder was then treated with the HCl solution (6 N) to remove carbonates. The sample was kept in solution for 48 hr while the container was held in a water bath at 55 °C and magnetic mixer constantly mixed the solution. Sonication was regularly used for complete mixing. The release of CO₂ at early stage of the process showed the effectiveness of HCl to remove carbonates. At the end of treatment with HCl, the solid residue was collected by filter paper and then dried in an oven overnight at a temperature less than 60 °C. Afterwards, the HF (24 wt%) and HCl (6 N) mixture solution (1:1) was used to remove silicates in the same process as in HCl treatment. The acid concentration, acid content and interaction time with sample were designed to be sufficient to remove the silicates including clay minerals, thus eliminating the effect of clay minerals on kerogen swelling. Subsequently, the extractable organic matter was removed by Soxhlet extraction with toluene from the previous solid residue. The sample was then dried in an oven. The temperature used for drying the sample at different steps of kerogen extraction was kept below 60 °C. This is the limit that changes the maturity of the kerogen. Starting with 10 g of shale source powder, the processing of the sample resulted in ~6.5 g of kerogen powder, which indicates a high kerogen content (greater than 60%). The total organic content (TOC) of Kimmeridge kerogen is reported to be about 62.5 wt% [37]. Elemental analysis [37] shows that Kimmeridge kerogen contains 62.52 wt% carbon, 6.49 wt% hydrogen, 5.58 wt % oxygen, 1.70 wt% nitrogen and 11.31 wt% sulfur. The kerogen type corresponds to immature type II according to the hydrogen/carbon (1.25) and oxygen/carbon (0.067) atomic ratio [37].

2.2. Liquid hydrocarbons

The swelling experiments were performed in three classes of organic solvents, including normal-alkanes, intermediate napthenes and aromatics. These liquid hydrocarbons represent the prevalent intermediate
components in petroleum liquids. A suite of liquid hydrocarbons with different number of carbon atoms were selected for each class of solvents. This selection was made to investigate the effect of solvent molecular size and solvent molecular shape on kerogen swelling. Our selection covers a broad range of chain length in normal-alkanes, aliphatic ring size in naphthenes and side methyl group of rings in aromatics. The difference between the three classes of hydrocarbons with the same number of carbon atoms represents the effect of solvent molecular shape. These liquid hydrocarbons cover a wide range of solvent solubility parameter and molar volume at 25 °C (Table 1). In addition to nonpolar solvents, tetrahydrofuran was also selected as a polar solvent to study kerogen swelling.

2.3. Test-tube swelling

Kerogen swelling is a measure of the change in volume of kerogen due to expansion of the organic network by a solvent. To measure swelling, roughly 0.2 g of the extracted kerogen powder was weighed into NMR tubes with outer diameter of 3 mm. NMR tubes are made of glass with uniform thickness. The 3-mm tubes were selected, instead of 5-mm tubes widely used in previous studies [3,6], to improve the accuracy of measurement. The key step in the swelling experiments is densely pack the kerogen in the NMR tube and then measure the initial kerogen height. Loose packing may result in kerogen shrinking in solvents if the kerogen swelling is low. Multiple tapping steps were adopted in the kerogen filling process to achieve the highest compaction. The sample tubes were then centrifuged twice at 2500 rpm for 5 min, and the initial height of dry sample was recorded when there was no change in height in hours, which guaranteed sufficient crosslink formation of kerogen network. The crosslinked and elastic properties of kerogen were assumed to be independent of solvent. The liquid hydrocarbons were subsequently added into each tube gradually while allowing sufficient time in crosslink formation of kerogen network.

In this section, we report the measured kerogen swelling by normal-alkanes, naphthenes and aromatics (Fig. 2, Table 1). For normal-alkanes, the swelling is measured in n-C₅ to n-C₁₀, n-C₁₂ and n-C₁₄. Normal-alkanes with carbon atoms more than 15 are solid at room temperature. The highest kerogen swelling ratio is measured in n-hexane and n-pentane with values of 1.14 ± 0.03 and 1.13 ± 0.01, respectively. As the number of carbon atoms increases in the chain of normal-alkanes, the swelling decreases in the kerogen sample. The averaged swelling in n-C₆ in three measurements is slightly higher than that in n-C₅, but the difference is within the standard deviations. The swelling ratio reaches a plateau for normal-alkanes with the number of carbon atoms higher than 10.

The swelling of kerogen in cycloalkanes is measured for C₅ to C₈, with values of 1.14 ± 0.04. The swelling of kerogen in cycloalkanes is modeled by the extended regular solution theory framework originally developed for swelling of polymer networks [3]. In this framework, the regular solution theory of Hildebrand [38] is extended by coupling with the Flory-Rehner theory of rubber elasticity [39]. In this treatment, the elastic contribution is integrated in the free energy of the kerogen-solvent system, which, in turn, is utilized to determine the swelling of kerogen network in solvent. The theoretical model has three assumptions [6]: (1) the dissolution of solvent in kerogen is in thermodynamic equilibrium; (2) kerogen network follows a rubber-like behavior; (3) the kerogen-solvent-solvent interactions are nonspecific. The assumption of rubber-like behavior implies that the amorphous kerogen can experience reversible deformation with exposure to solvent. To possess the rubber-like behavior, the kerogen network may not be over-crosslinked, and the interaction with solvent may not change the crosslinked and elastic properties of kerogen. In the theoretical model, the measured swelling of kerogen in various solvents were used to determine the three thermodynamic parameters of kerogen (solubility parameter, cross-link density, and native swelling fraction), which minimize the overall difference between experimental and theoretical swelling ratios. The solubility parameter represents the degree of interaction in kerogen, and is a measure of solubility [15]. The cross-link density is defined as the density of elastically effective number of chain segments bounded on both ends by crosslinks [6]. Native swelling fraction is equal to the volume fraction of kerogen network that minimizes its elastic strain energy [3]. It reflects the native deformation of kerogen network to reach the unstretched state. The general formulation of the theoretical model has been presented by Ertas et al. [3] for kerogen interaction in multicomponent solvents. In this work, we use the same model for a system in which kerogen interacts with a single excess solvent. Detailed formulation is supplemented in Appendix A.

3. Results

In this section, we report the measured kerogen swelling by normal-alkanes, naphthenes and aromatics (Fig. 2, Table 1). For normal-alkanes, the swelling is measured in n-C₅ to n-C₁₀, n-C₁₂ and n-C₁₄. Normal-alkanes with carbon atoms more than 15 are solid at room temperature. The highest kerogen swelling ratio is measured in n-hexane and n-pentane with values of 1.14 ± 0.03 and 1.13 ± 0.01, respectively. As the number of carbon atoms increases in the chain of normal-alkanes, the swelling decreases in the kerogen sample. The averaged swelling in n-C₆ in three measurements is slightly higher than that in n-C₅, but the difference is within the standard deviations. The swelling ratio reaches a plateau for normal-alkanes with the number of carbon atoms higher than 10.
and C10. The cycloalkanes are solid when the number of carbon atoms is higher than 10. The swelling decreases with the increase in the number of carbon atoms in the ring of naphthenes. The highest swelling ratio of kerogen is 1.14 ± 0.03 in cyclopentane, and the lowest swelling ratio is 1.02 ± 0.01 in cyclodecane. The swelling ratio of kerogen in aromatics is also presented in Fig. 2. A comparison among benzene, toluene and o-xylene in the kerogen sample shows that as the number of methyl groups attached to the base ring increases from zero to two, the swelling ratio decreases slightly from 1.12 ± 0.02 to 1.09 ± 0.01. The swelling ratio of tetrahydrofuran (1.14 ± 0.04) is slightly higher than the aromatic solvents (Table 1). However, we observed the change of color of tetrahydrofuran in the tube as it came in contact with the kerogen sample. This implies that some residual bitumen components dissolve in tetrahydrofuran, since kerogen is insoluble in organic solvent. The dissolution of organic components in solvent can affect the swelling of kerogen, indicating the measured result (1.14 ± 0.04) is most probably not accurate. We exclude tetrahydrofuran for further discussion.

4. Discussion

4.1. Effect of solvent molecular size and shape

Measurements of the three classes of liquid solvents in the Kimmeridge kerogen show a consistent trend; there is a decrease with the increase of solvent molecular size. Specifically, the swelling ratio descends as the chain length increases in normal-alkanes, the ring size expands in naphthenes, and the side methyl group rises in benzene ring of aromatics. This trend is correlated with the size exclusion effect which integrates the fluid molecular size with the pore accessibility in kerogen [41]. The increase in molecular size can reduce the fluid diffusivity in ultraconfining kerogen matrix, which results in less dissolution of fluid. The effect of fluid molecular size is consistent with our recent finding that [17] kerogen swelling induced by water molecules decreases with the increase of kerogen pore sizes. In addition to molecular size for each class of solvents, the comparison of kerogen swelling in the three classes of solvents with the same number of carbon atoms can help to investigate the effect of solvent molecular shape. The kerogen swelling is more pronounced in normal-alkanes with linear chains than that in naphthenes with cyclic rings. This is because the linear molecular chain with flexibility facilitates the fluid to diffuse in the porous channels. Compared with naphthenes, aromatics induce a larger swelling ratio of kerogen, which may be attributed to the π–π interaction between solvents and aromatic rings of kerogen macromolecules [4].

4.2. Effect of solvent solubility parameter and molar volume

Fig. 3 shows the plot of kerogen swelling and solvent solubility parameter. The swelling in normal-alkanes and naphthenes has the same trend; it decreases linearly with the increase of solvent solubility parameter. The two solvent classes have the same slope which is an indication of the type of interaction with the kerogen network. The decreasing linear trend has not been reported in the past to the best of our knowledge. The aromatics are very distinct from alkanes. Previous studies (based on modeling) have reported a bell-shaped [22–24] or increasing trend [6] in different classes of solvents. In this work, the swelling data in different classes of solvents are distinguished independent trends.

In addition to solvent solubility parameter, solvent molar volume is another key parameter that may correlate with the swelling of kerogen. The swelling of kerogen as a function of solvent molar volume is presented in Fig. 4. The solvent molar volume includes both the effect of solvent molecular size and molecular shape. A decreasing trend of kerogen swelling with increasing solvent molar volume is observed, which confirms the size exclusion effect in the ultraconfining kerogen matrix. The number of solvent molecules that can diffuse into the kerogen matrix diminishes as the solvent molar volume increases.
Similar to solvent solubility parameter, the dependence of kerogen swelling on solvent molar volume is distinct in normal-alkanes, naphthenes, and aromatics. A comparable exponential trend is observed in the two classes of alkanes, which demonstrates the interactions are similar.

In Figs. 3 and 4, the swelling data of aromatics show a different class and trend. Due to small range in solubility parameter and molar volume in the three aromatics, a definitive trend cannot be deduced.

In Fig. 2, the swelling of Kimmeridge kerogen reveals a decreasing trend as the molar volume of solvent increases and the solubility parameter of solvent gets closer to the general kerogen solubility parameter [3,6,22–24]. It is assumed that the swelling increases when the solubility parameter of solvent approaches that of kerogen. Compared with solvent solubility parameter, the molar volume of solvent dominates the swelling of Kimmeridge kerogen. Hruljova et al. [20] have also reported that the solubility parameter of solvent may not be a key parameter, while the molar volume of solvent could be a dominate parameter in determining the swelling of Kukersite kerogen. In their work, the kerogen swelling showed no clear relation with solvent solubility parameter and a linear trend was not observed with solvent molar volume.

4.3. Description by the extended regular solution theory framework

The extended regular solution theory framework has been used in the past to describe kerogen swelling by various authors [3,6]. Fig. 1 was prepared based on the three coefficients which were obtained from swelling measurements of alkanes and aromatics in ref [6]. As the figure shows the kerogen swelling increases with the increase in the number of carbon atoms in normal-alkanes. This theoretical trend is contrary to the experimental trend in Fig. 2. In this work, our goal is investigation of the applicability of the extended regular solution theory to the swelling of Kimmeridge kerogen in petroleum liquids. We will demonstrate that although the theoretical predictions can match well with the experimental results, the predictions are based on fitting of the three thermodynamic parameters of kerogen, and the fitting thermodynamics parameters especially the fitted solubility parameter of kerogen may not have a sound physical meaning.

The swelling ratio of kerogen in the three class of hydrocarbons from our measurements is modeled and presented in Fig. 5. We have used all of our data together (Fig. 5 a) and then all the data for n-alkanes and naphthenes (Fig. 5 b) and then separately (Fig. 5 c, d) to get the parameters of the model. The theoretical predictions for swelling ratio of kerogen in normal-alkanes and naphthenes match well with the experimental data (Fig. 5 b–d). In contrast, the theoretical results for aromatics are different from the experimental data (Fig. 5 a). This observation indicates that aromatics may have different interactions with the kerogen network compared with normal-alkanes and naphthenes. The difference can be due to the π–π interaction between aromatics and kerogen aromatic units [4].

Fig. 6 presents the theoretical predictions of kerogen swelling ratio as a function of solvent solubility parameter and molar volume based on experimental results of normal-alkanes and naphthenes. The swelling follows a bell-shaped trend with the solvent solubility parameter, which first increases to a maximum, and then decreases with further increase in solubility parameter. The bell-shaped trend is consistent with the reported experimental results for type I and III kerogen [23]. Note that the theoretical swelling shows a decreasing trend as the solvent solubility parameter varies from 14.4 to 18.7 J1/2cm−3/2, which is in qualitative agreement with the experimental data. The solvent solubility parameter is well defined. In contrast, the solubility parameter of kerogen cannot be measured directly. It is estimated from the swelling measurements. The kerogen solubility parameter has been reported to vary between 19 and 24.3 J1/2cm−3/2 [3,6,22–24], which is close to that of petroleum asphaltenes [35]. Table 2 presents the solubility parameter of kerogen based on description of the extended regular solution theory and our data. The solubility parameters are obtained from all of our data, as well as separating the data in different classes of solvents. Note that the kerogen solubility parameters in Table 2 are less than the solubility parameters of light liquid normal alkanes in Table 1; they are much less than the kerogen solubility parameters reported in the literature. As we have seen above there is inconsistency in the predictions from the literature solubility parameters with the trends of our normal alkane data. The other two parameters of the extended regular solution theory are in qualitative agreement with the values obtained in the past.

Going back to Fig. 5, we observe that the kerogen swelling from the extended regular solution theory matches the experimental results (except for aromatics). However, the thermodynamics parameters especially the solubility parameter may not have a sound physical meaning. The cross-link density is reported to increase with the poly-aromatic clusters in kerogen, which increases as the thermal maturity of kerogen from immature to overmature [42] or as the organic type of kerogen changes from type I, type II to type III [43]. The cross-link density (0.10–0.15 mol/cm3) for Kimmeridge kerogen (type II) from our work is lower than the reported results for type II kerogen (0.16 mol/cm3) and type IIIC kerogen (0.25 mol/cm3) [6]. The solubility parameter of kerogen in Table 2 (8.5–10.8 J1/2cm−3/2) is two times less than the values (19–24.3 J1/2cm−3/2) in the literature [3,6,22–24]. The large difference indicates that the extended Flory-Rehner and regular solution theory may not describe the swelling of kerogen.

The theoretical model can fit the measured swelling of Kimmeridge kerogen by adjusting the thermodynamic parameters of kerogen, but the optimized solubility parameter does not have a sound thermodynamic basis. Therefore, the theoretical model may not describe the swelling of Kimmeridge kerogen in petroleum liquids. Oja et al. [44] have reported a similar conclusion but from a different aspect. They concluded that the extended Flory-Rehner and regular solution theory is not suitable for Green River kerogen, because the average molecular weights between cross-links predicted by the theoretical model were clearly lower than the experimentally determined results. These observations indicate that the extended Flory-Rehner and regular solution theory may not be reliable for quantitative estimate, which can be attributed to the uncertainties in the assumptions and simplifications of the theoretical model. In addition to an enthalpic term, the kerogen-solvent interaction parameter can also contribute to an entropic term. The neglect of the entropic term in the theoretical model may increase the uncertainty. Furthermore, the theoretical model requires the kerogen to follow the rubber-like behavior, namely the interaction with solvent does not change the crosslinked and elastic properties of kerogen, which may be invalid for realistic kerogen samples with highly irregular structural segments and heterogeneous nature.

5. Conclusions

We draw two main conclusions from this investigation.

1. Kerogen swelling decreases as the chain length increases in normal-alkanes, the ring size expands in naphthenes, and the side methyl group increases in benzene ring of aromatics. Compared with naphthenes with cyclic rings, the swelling is higher in normal-alkanes with flexible linear chains and in aromatics having π–π interaction with kerogen. The swelling ratio of Kimmeridge kerogen in both normal-alkanes and naphthenes shows a linear decrease with the solvent solubility parameter and an exponential decrease with the solvent molar volume. The swelling in the two classes of kerogen has a similar decreasing slope, demonstrating similar interactions with the kerogen network. Compared with solvent solubility parameter, the effect of solvent molar volume dominates the swelling of kerogen in normal-alkanes and naphthenes.
The extended Flory-Rehner and regular solution theory framework may not describe the swelling of kerogen in petroleum liquids. The optimized thermodynamic parameters of kerogen do not have a sound thermodynamic basis. Alternative models may be pursued in the future.

**Table 2**

<table>
<thead>
<tr>
<th>Solvent</th>
<th>Solubility parameter $(J^{1/2} \text{cm}^{-3/2})$</th>
<th>Cross-link density (mol/cm³)</th>
<th>Native swelling fraction</th>
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</thead>
<tbody>
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<td>Alkanes + Aromatics</td>
<td>10.8</td>
<td>0.15</td>
<td>0.95</td>
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<td>Alkanes</td>
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<td>0.95</td>
</tr>
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<td>N-alkanes</td>
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<td>0.10</td>
<td>0.85</td>
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<td>Naphthenes</td>
<td>8.8</td>
<td>0.10</td>
<td>0.80</td>
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</table>

**CRediT authorship contribution statement**

Liang Huang: Data creation, Formal analysis, Validation, Visualization, Writing - original draft, Writing - review & editing.

Atefeh Khoshnood: Investigation, Methodology, Writing - original draft. Abbas Firoozabadi: Conceptualization, Funding acquisition, Project administration, Supervision, Writing - review & editing.

**Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

**Acknowledgments**

This work was supported by the member companies of the Reservoir Engineering Research Institute (RERI). We appreciate the support of the
National Natural Science Foundation of China (Grant No. 51774298) and the China Scholarship Council (CSC No. 201806440095). We would like to thank Mr. Peng Gao of Stanford University for his help on programming the extended Regular Solution Theory. We also thank Dr. Tianhao Wu and Dr. Ali Zidane of RERI for discussion.

Appendix A

One key relationship is the expression for the change in Gibbs free energy of the system which includes entropic and enthalpic terms. The changes in entropy originate from two sources: conformational variations and entropy of mixing. According to Flory-Rehner theory, an affine transformation in the network of a crosslinked polymer results in the following entropic changes [38],

$$\Delta S_n = -k_b \frac{N_e}{2} \left[ \alpha_x^2 + \alpha_y^2 + \alpha_z^2 - 3 - 2 \ln(\alpha_x, \alpha_y, \alpha_z) \right] \quad (1)$$

In Eq. (1), $\Delta S_n$ is the change in entropy due to conformational variations, $k_b$ is the Boltzmann constant, $\alpha_x$, $\alpha_y$, $\alpha_z$ are the fractional elongations along the $x$, $y$, and $z$ directions, respectively, $N_e$ is the elastically effective number of chain segments bounded on both ends by crosslinks. $N_e$ is defined as the total number of crosslinked units minus the number of terminal chains bounded at one end to a cross-linkage [38]. The terminal chains have no contribution to the elastic free energy.

By introducing native swelling fraction $\nu_{eq}$, kerogen volume fraction $\nu_0$, and assuming uniform elongation $\alpha_x = \alpha_y = \alpha_z = (\nu_{eq}/\nu_0)^{1/3}$, Eq. (1) can be rewritten as,

$$\Delta S_n = -k_b N_e \left[ 3 \frac{(\nu_{eq})^2}{\nu_0} - 3 - 2 \ln \left( \frac{\nu_{eq}}{\nu_0} \right) \right] \quad (2)$$

where $\nu_0$ is the kerogen volume fraction, $\nu_{eq}$ is the native swelling fraction. The native swelling fraction $\nu_{eq}$ is numerically equal to the volume fraction of kerogen network that minimizes its elastic strain energy [3]. The native swelling fraction $\nu_{eq}$ is an innate thermodynamic parameter of kerogen network, which is independent of solvent type and amount.

The entropy of mixing is given by [3],

$$\Delta S_m = -k_b n_1 \ln \nu_1 \quad (3)$$

where $\Delta S_m$ is the change in entropy due to introduction of solvent, $n_1$ and $\nu_1$ are the total number of solvent molecules and solvent volume fraction, respectively. Eq. (3) can be used to interpret the mixtures of kerogen and solvent when ignoring specific interactions. In this equation, the solute term is excluded since the equation is utilized to describe the contribution of solvent introduction to the associated change of conformational entropy. The contribution of solute to the change of conformational entropy is described in Eq. (2). The heat exchange due to mixing of solvent and kerogen network can be obtained by [3],

$$\Delta H_m = k_b T \chi_{X,0} n_1 \nu_0 \quad (4)$$

where $\Delta H_m$ is the change in enthalpy due to mixing of solvent and kerogen, $\chi_{X,0}$ is the interaction parameter between solvent (1) and kerogen network (0), and $T$ is temperature. An alternative way of writing Eq. (4) is by defining interaction energy density $b_{1,0} = \chi_{X,0}/\theta$, where $\theta$ is the molecular volume of the solvent. The molecular volume is converted from the molar volume of solvent [15,40], divided by Avogadro constant.

We can relate the enthalpy of mixing to solubility parameters through the regular solution theory [38],

$$b_{1,0} = \frac{(\delta_1 - \delta_0)^2}{k_b T} \quad (5)$$

where $b_{1,0}$ is interaction energy, $\delta_1$ and $\delta_0$ are solubility parameters of solvent and kerogen, respectively.

The change in Gibbs free energy from mixing of kerogen and solvent molecules is given by,

$$\Delta G = \Delta H_m - T (\Delta S_m + \Delta S_n) \quad (6)$$

where $\Delta G$ is the change in Gibbs free energy. By substituting Eqs. (2), (3), (4), (5) into Eq. (6), we arrive at,

$$\frac{\Delta G}{k_b T} = b_{1,0} \delta_0 n_1 \nu_0 + n_1 \ln \nu_1 + \frac{N_e}{2} \left[ 3 \left( \frac{\nu_{eq}}{\nu_0} \right)^2 - 3 - 2 \ln \left( \frac{\nu_{eq}}{\nu_0} \right) \right] \quad (7)$$

The expression for $\Delta G$ can be used to calculate the chemical potentials. The change in chemical potential of the solvent in contact with kerogen network with reference to pure liquid is,

$$\frac{\Delta \mu_1}{k_b T} = \frac{1}{k_b T} \frac{\partial \Delta G}{\partial n_1} \quad \text{(8)}$$

where $\Delta \mu_1$ is the change in chemical potential of the solvent. By substituting Eq. (7) into Eq. (8) and carrying out the derivative, we arrive at,

$$\frac{\Delta \mu_1}{k_b T} = b_{1,0} \delta_0 \omega^2 + \ln \nu_1 + \nu_1 + \frac{N_e \delta}{\omega} \left( \frac{1}{\omega^2 \nu_0} - \nu_0 \right) \quad (9)$$

where $\omega$ is the initial volume occupied by the kerogen network, and $\nu_1$ is the solvent volume fraction ($\nu_1 = 1 - \nu_0$). The kerogen volume fraction in the mixture is defined as $\nu_0 = \omega/(\omega + n_1 \theta)$. The volumetric swelling ratio is the inverse of the kerogen volume fraction, $Q_c = \nu_0^{-1}$. Now we write the equilibrium condition for the liquid absorbed in the kerogen network and the excess liquid. The swelling is modeled similar to the way the experiments are performed; there is always excess solvent after equilibrium has been reached. Therefore, $\Delta \mu_1^{\text{absorbed}} = \Delta \mu_1^{\text{excess}}$. Note that the theoretical model ignores the effect of osmotic pressure on solvent partitioning and kerogen swelling. We can write the working equation for kerogen swelling calculations,
Eq. (10) can be used to calculate kerogen swelling by excess solvent once we determine the three thermodynamic parameters of kerogen (solubility parameter δ0, native swelling fraction uw0 and crosslink density N/a0). Note that the three thermodynamic parameters reflect the innate properties of kerogen network, which are independent of solvent type and amount. In this treatment, the three thermodynamic parameters are first determined by optimizing the match of the swelling data and the theoretical model. For this purpose, a cost function ξ is defined to minimize the error of predictions and experiments [3,6],

\[
\xi = \frac{1}{N} \sum_{i=1}^{N} \left( \frac{Q_i^0}{Q_i} - 1 \right)^2 \]

(11)

In Eq. (11), ξ is the cost function, and N is the number of data points. Superscript p and e refer to predicted and experimental results, respectively. In this work, we adopt the marching method to find the combination of the three parameters that leads to the minimized cost function. The search region is 0–40 J/cm³·mol⁻¹/² for solubility parameter of kerogen, 0–1 mol/cm³ for crosslink density of kerogen and 0–1 for native swelling fraction. In this method, all the possible combinations of the three parameters are examined to ensure that the optimized combination is obtained.

References