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Molecular structure characterization of asphaltene in the presence of inhibitors with nanoemulsions†

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Molecular structure characteristics and morphological features of asphaltene can be significantly influenced by the addition of asphaltene inhibitors (AI). We have recently developed a novel concept of using nanoemulsions (NE) as carriers for controlled release of asphaltene inhibitors, which could prohibit the precipitation problem with reduced AI quantity. In this work, X-ray diffraction (XRD) was utilized to investigate the changes in the stacking behaviour of asphaltenes in the presence of three cases: (i) strong organic acids (dodecyl benzene sulfonic acid, DBSA), (ii) nanoemulsions (blank NEs), and (iii) nanoemulsion loaded DBSA (DBSA NEs). Based on the XRD and transmission electron microscopy (TEM) analyses, the stacking distance between aromatic rings of asphaltene was found to be increased by 22.2%, suggesting that the modification of the π system over the aromatic zone prevented the ultimate π - π interactions between asphaltene sheets. The evidence of multiple intermolecular interactions quantitatively obtained from Fourier-transform infrared spectroscopy (FTIR) supported our proposed mechanism for controlled release effect and long-term asphaltene stability, i.e., the decrease of the aromaticity and the reduction in the aliphatic side chains of asphaltene. The refractory nature of asphaltenes was examined by thermogravimetric analysis (TGA), which showed that the asphaltene structure was improved considerably and the coke yield was decreased by 62% due to the decrease of the cluster size and the increase of the stacking distance.

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1. Introduction

Asphaltene is a fraction that tends to cause most of the issues during crude oil transport and refining. The molecular structure of asphaltenes is the least understood among all petroleum fractions. The asphaltene aggregation and the formation of colloidal particles in crude oil can be attributed to a range of intermolecular interactions such as hydrogen-bonding, aromatic π - π stacking, a polar-polar interactions, and electrostatic attractions between asphaltene molecules.

Analytical techniques have been used to characterize the asphaltenes and to determine their molecular structures. Y-ray diffraction (XRD) is used to provide valuable information concerning the internal structure of asphaltenes and to study the crystallite parameters of the asphaltene clusters. O-13 A pile of aromatic sheets can be formed due to the stacking of asphaltene aromatic cores, which is possible to analyse by XRD because the asphaltene stacking may form the crystallites. In addition, further crystallite parameters in different crude oil fractions such as naphthene aromatics, polar aromatics, and saturate

fractions also can be determined by XRD.¹⁵ For example, AlHumaidan $et\ al.$, ¹⁶ studied the influence of thermal cracking on the molecular structure of asphaltene, and their results showed that the distance between two aromatic sheets $(d_{\rm m})$, distance between the two aliphatic chains or saturated rings (d_{γ}) , and the size of the aromatic sheets $(L_{\rm a})$ were not significantly affected by the thermal cracking intensity. The condensed sheets may also contain heteroatoms such as oxygen, sulfur, and nitrogen atoms which could act as free radicals for anchor points of bound metals such as Ni, V, Mo or Fe. Yin $et\ al.$, ¹⁷ investigated the interaction of V and Ni with condensed aromatic compounds. Their results showed that no interactions between metals and aromatic rings of asphaltene have been detected using UV-vis and fluorescence spectroscopy.

Chemical treatments are generally used to alter the macroscopic properties of asphaltenes and improve heavy oil properties by alteration of asphaltene structural characteristics and change of nanoaggregate size. Afraa *et al.* oused (anacardic acid, cardanol, and cardol (ACC)) as a solvent to investigate its effects on the aggregation behaviour of asphaltene by using XRD. The results revealed the modification of both the aromaticity and the layer distance between two aromatic sheets after the treatment with the solvent. Recently investigators have examined the effects of bio-oils on asphaltene structure. Properties of bio-oils on asphaltene structure of the substances such as waste cooking oil, and swine waste, does not

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only weaken the clustering of asphaltene but also improve the overall colloidal structure of asphaltene. However, the literature is struggling with how these compounds can affect the properties of asphaltene and what kind of molecular interactions are responsible for their activity.23-25

Even though asphaltene inhibitors are widely used in oilfields as common methods to prevent the aggregation and precipitation of asphaltene, 25,26 a major problem due to the heavy loss of inhibitors and hence high treatment cost makes asphaltene inhibition by inhibitors uneconomic. We have proposed a novel concept of using nanoemulsion as a carrier for AI and control its release inside asphaltene solution to increase asphaltene stability.27 The findings confirm that NEs could (i), significantly decrease the inhibitors usage (ii) improve the asphaltene stability, and (iii) extend the treatment time via the slow release of AIs with high efficiency. However, a complete molecular analysis of asphaltene and a systematic understanding of how asphaltene molecular structure changes with the addition of inhibitors have not yet been achieved.

In the present work, we attempted to provide novel insights into the stability and molecular modification structure of asphaltene in the presence of three cases: (i) pure DBSA, (ii) blank NEs, and (iii) DBSA NE. Based on the structural parameters obtained from combined XRD, transmission electron microscopy (TEM), and Fourier-transform infrared spectroscopy (FTIR), a stability mechanism was proposed to describe changes in asphaltenes and the effects of DBSA NEs thereon. The thermal stability and the refractory nature of asphaltene are also performed by a thermogravimetric analyser (TGA).

Experiment and methods 2.

2.1. Chemicals

Chemical materials were purchased from Sigma-Aldrich company including solvents such as xylene (>98.5%), nheptane (extra pure \geq 98%), toluene (extra pure \geq 98%), surfactants such as Tween 80 (sorbitan monooleate), sodium dodecyl sulfonate (SDS) and asphaltene inhibitor dodecylbenzene sulfonic acid (DBSA).

2.2. Methods and measurements

2.2.1. Synthesis of nanoemulsions with the presence and absence of inhibitor. The successful long-term stability of AI inside NEs needs different materials than currently used (surfactants) to provide resistant barriers that can prevent quick diffusion of AI from the nanodroplet core. As an example study, DBSA can provide steric and electrostatic repulsion together with hydrophobic and acid-base interactions between its molecules and ionic/non-ionic surfactants. Such a synergistic effect may overcome the hydrogen bond forces of water molecules and enhance the bounded molecules at the interface, resulting in more hydrophobicity of the oil phase. The procedure of synthesis of nanoemulsions with the presence and absence of AI was elaborated in the previous study,27 as depicted in the schematic diagram in Fig. S1 in the ESI.† Table 1 shows

the composition and concentrations of all three cases used in this work.

2.2.2. The extraction of asphaltene and sample preparation. The extraction of asphaltene was carried out based on ASTM D2007 by adding crude oil to *n*-heptane at a volume ratio of 1:40, which is led to asphaltene destabilization and increasing the asphaltene precipitation rate.28 The procedure of asphaltene extraction was reported previously.27 For preparation asphaltene solution, a (0.5 w/v %) of the asphaltene was dissolved in a mixture of two solvents: toluene/heptane (Heptol) volume ratio of 60:40. To observe the effect of DBSA, blank NEs, and DBSA NEs on asphaltene precipitation structure, samples were slowly added to the asphaltene solution under shear action in an (IKA T2S digital/Ultra-Turrax) homogenizer at a rotation speed of 7000 rpm until a complete combination of the NEs into the asphaltene solution. Then the asphaltene solution with and without DBSA, blank NEs, and DBSA NEs were kept for 3 days for totally evaporation at ambient temperature. The dried solid asphaltenes were degassed for 2 h at 70 °C in a vacuum oven prior to the analyses.

2.2.3. Molecular structure characterization of asphaltene. X-ray diffraction of the dried samples was obtained with a highpower (12 kW) X-ray diffractometer (D8 XRD) containing a graphite monochromator, rotating copper anode, and a scintillation counter. The diffraction profiles ($6^{\circ} \le 2\theta \le 90^{\circ}$) were measured at ambient temperature using Cu K α radiation (λ = 1.54055 Å) at 1 deg min⁻¹ scanning rate and a step size of 0.02 (2), the final procedure of calculating the crystalline parameters are presented in S2 (ESI†). The asphaltene particles morphology was inspected using a transmission electron microscope (FEI Titan Themis Cubed 300 TEM).

At spectra range from 500 to 4000 cm⁻¹, a resolution of 1 cm⁻¹, and 64 scans per spectra, the asphaltene samples were collected on a Bruker TENSOR-270 FTIR spectrometer. FTIR spectra were measured by the attenuated total reflectance (ATR) technique using a Bruker ATR A225 sample cell equipped with a diamond crystal of quadrate area of $2 \times 2 \text{ mm}^2$ and a spectral range of crystal from 10 000 to 10 cm^{-1} . The FTIR spectrometer was controlled using the OPUS software.

Studying the thermal behaviour of asphaltene in the presence of oxygen is necessary to have a better understanding of the thermal stability of asphaltenes. In a general thermogravimetric analysis (TGA) (a Perkin-Elmer analyzer model TGA-7HT), a known amount (9-15 mg) of a solid sample was put in an alumina or a platinum pan and placed in a small furnace. The weight of the sample was constantly measured at 10 °C min⁻¹ heating. For all cases, the temperature interval ranged from ambient temperature up to 800 °C.

Results and discussion

3.1. XRD studies of asphaltenes

3.1.1. Effect of DBSA on asphaltene structure. Fig. 1 shows that when the asphaltene is stabilized by the addition of 4 vol% DBSA, the crystalline parameters are changed. The distance between two aromatic sheets $d_{\rm m}$ of asphaltene is 3.26 Å, which increases to 3.70 Å for the asphaltene with DBSA. These results

Table 1 The composition and concentrations of all three cases

Sample	DBSA (vol%)	Xylene (vol%)	Tween 80 (vol%)	SDS (vol%)	Water (vol%)
DBSA	4.0	_	_	_	_
Blank NE	_	7	9.9	0.1	83
BDSA NE	1.0	7	9.9	0.1	82

are compatible with those reported in the literature. ²⁹⁻³¹ Correspondingly, the distance between the two aliphatic chains or saturated rings d_{γ} also increased from 4.70 to 5.25 Å. However, the height stack of the aromatic sheets vertical to the plane ($L_{\rm c}$) is decreased from 13.3 to 10.31 Å. The average number of aromatic sheets ($M_{\rm e}$) in asphaltene cluster is 4.65 Å and it decreases to 3.9 Å after the addition of DBSA molecules. The average diameter of the aromatic sheet $L_{\rm a}$ and the number of aromatic rings in the aromatic sheet ($R_{\rm a}$) are significantly changed with the addition of DBSA from an average of 7.1 Å and 2.66 for asphaltene to 6.2 Å and 3 for asphaltene with DBSA respectively. This can be attributed to the effectiveness of DBSA to interact with asphaltene molecules via acid-base interaction, which suggests that the asphaltene becomes looser.

3.1.2. Effect of blank NEs on asphaltene structure. The average interlayer distance d_{m} for asphaltene stacks with blank NEs are not significantly changed gradually from 3.26 Å for the asphaltene sample to 3.42 Å. Therefore, despite the fact that observed changes are considerably small, they give a good indication that the distance between two aromatic sheets of asphaltene cluster is slightly affected and loosened by the addition of blank NEs. The distance between the two aliphatic chains or saturated rings was around 5.01 Å for the asphaltene with blank NEs. The average diameter of the aromatic sheet L_a and the cluster diameter L_c for asphaltene have also decreased with the addition of blank NEs and ranges from an average of 7.1 and 13.3 Å to 6.7 and 11.3 Å respectively. However, the insignificant change in the average number of aromatic sheets $M_{\rm e}$, the average number of aromatic rings in each aromatic sheet R_a , and the number of carbons per aromatic structural C_{Au}

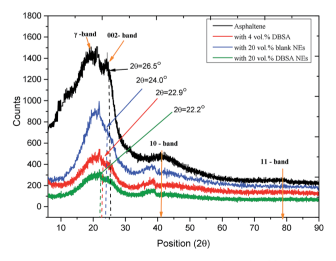


Fig. 1 $\,$ XRD pattern of asphaltenes with 4 vol% DBSA, 20 vol% blank NEs, and 20 vol% DBSA NEs.

compared with asphaltene sample were observed. Thus, these results suggest that blank NEs have a moderate effect on asphaltene structure comparing with using 4 vol% DBSA. This performance may be owing to the presence of different components in blank NEs such as surfactants, water, and xylene that provide intermolecular interactions with asphaltene molecules.

3.1.3. Effect of DBSA NEs on asphaltene structure. One interesting finding is the distance between the aromatic sheets (stacking space), which is significantly increased from 3.26 Å to 3.96 Å with the addition of DBSA NE to the asphaltene sample. The distance between the two aliphatic chains or saturated rings d_{γ} was also affected by the addition of DBSA NE and increased to 5.35 Å. In addition, the size of the aromatic sheets $L_{\rm a}$ has significantly altered with the addition of DBSA NEs from 7.1 Å to 6.1 Å. This result is consistent with a reduction of the number of aromatic rings in the aromatic sheet R_a from 3 to 2. The significant change in the size of the aromatic sheets is primarily due to the fact that, at the addition of DBSA NES, the aromatic C=C bonds are changed. Moreover, the M_e , R_a , and $C_{\rm Au}$ decreased to 3.9, 2.2, and 11.27 respectively. On the other hand, the cluster diameter L_c , decreased in size to 10.2 Å as the DBSA NEs is added. The decrease in the cluster diameter of asphaltene is owing to the loss of aliphatic carbon in the alkyl chains and naphthenes that is evidently reflected by the aromaticity f_a , which signifies the ratio of aromatic to total carbon per cluster. Because the aromaticity plays a key role in the mechanism of asphaltenes stacking, its modification due to the addition of DBSA NEs should be examined in order to provide more information about the parameters that would change the aromaticity. The determination of aromaticity has a significant advantage to give the aromaticity of aromatic carbon that has the ability to form stacks. It is shown from Table 2 that the aromaticity of the asphaltene sample was 0.35; however, after the addition of DBSA NEs this value reduced to 0.22. It is possible that DBSA NEs modify the structure of asphaltene cluster and make it less aromatic. The small value of the aromaticity can be due to either shortening of alkyl side chains or loss of aromatic rings. These findings also suggest that the increasing trend of $d_{\rm m}$ value for asphaltene with DBSA NEs is consistent with the aromaticity reducing trend. The reduction of aromaticity is indicative of the reduction in the molecular moieties that initiate the stacking and leads to a less ordered structure. It may be supposed that, in this case, the DBSA and other nanoemulsion's components were progressively released from NE to provide various interactions with asphaltene and involve mainly the breaking of alkyl groups that weakly bond the poly-aromatic basic units, which are saturated the H-bonding sites of asphaltenes and the fact that it

Table 2 The crystalline parameters obtained from XRD results for asphaltene without and with DBSA, blank NE, and DBSA NE

	Asphaltene	With DBSA	With blank NE	With DBSA NE	Equation ^a	Reference
d_{γ} , Å	4.70	5.25	5.01	5.35	$d_{\gamma} = \frac{5\lambda}{8\sin\theta_{(\gamma)}}$	22
$d_{ m m}$, Å	3.26	3.65	3.42	3.96	$d_{ m m}=\;rac{\lambda}{\sin heta_{(002)}}$	22
$L_{ m c}$, Å	13.3	10.3	11.3	10.2	$L_{\rm C} = \frac{0.45}{\rm FWHM_{002\text{-band}}}$	22
$M_{ m e}$	5	4	4	4	$M = \frac{L_{\rm C}}{d_{\rm m}} + 1$	23
L_{a} , Å	7.1	6.2	6.7	6.1	$L_{\rm a} = \frac{0.92}{1000000000000000000000000000000000000$	23
$R_{\rm a}$	3	2	3	2	$R_{\rm a} = \frac{L_{\rm a}}{2.667}$	23
C_{Au}	13	11	12	11	$C_{\rm Au} = \frac{L_{\rm a} + 1.23}{0.65}$	24
$f_{ m a}$	0.35	0.27	0.31	0.22	$f_{\rm a} = \ rac{A_{(002)}}{A_{(002)} + A_{(\gamma)}}$	24

^a Where $\lambda = 1.54055$ Å; θ_{γ} , θ_{002} , and θ_{10} are the diffraction angles of γ , 002, and 10 bands, respectively; FMHM is the full width at half maximum of diffraction peaks, all these values are presented in the Table S1 (ESI). d_{γ} : the distance between the two aliphatic chains or saturated rings, $d_{\rm m}$: the distance between two aromatic sheets, $L_{\rm a}$: the size of the aromatic sheets, $L_{\rm c}$: cluster diameter, $M_{\rm c}$: the average number of aromatic sheets, $R_{\rm a}$: the average number of aromatic rings in each aromatic sheet, $C_{\rm Au}$: the number of carbons per aromatic structural. $f_{\rm a}$: the aromaticity.

interacted with both the periphery and aromatic cores of asphaltene. This suggests that the asphaltene is prevented from interacting laterally between themselves and kept well-stable in the solution (i.e., increasing asphaltene stability). The significant cleavage in alkyl side chains also produced a decrease in the asphaltene cluster diameter $L_{\rm c}$ as well as the size of the aromatic sheets $L_{\rm a}$ and the average number of aromatic rings in each aromatic sheet $R_{\rm a}$ were affected.

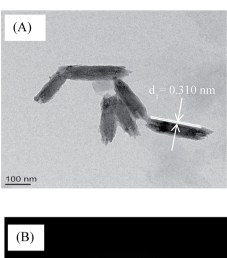
These findings for crystallite parameters of the asphaltene clusters and its aromaticity provide a further comparison between the three cases, which show that asphaltenes with 20 vol% DBSA NE has an approximately similar result as that of 4 vol% DBSA. However, the addition of 20 vol% blank NEs still have some effect but less strong compared with the other two cases. Asphaltene treatment with 20 vol% DBSA NEs (i.e., containing 0.04 mL DBSA) give closely the same values of crystallite parameters and aromaticity when treated with 4 vol% DBSA (i.e., about 0.8 mL). Thus, the reduction amount of asphaltene inhibitor (% R_{AI}) is reduced by 95%. The total chemicals usage (% R_{TC}) are again lower than 4 vol% DBSA by a factor of 10%. These results in very good agreement with the previous work²⁷ for reducing AI amount and the total chemicals used, which found that 4 vol% DBSA and 20 vol% of DBSA NEs gave the same values of the instability indexes and the sedimentation rates.

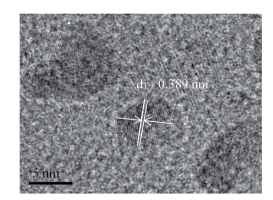
We further investigated the effects of DBSA NEs on asphaltene stacking with TEM in order to examine the evidence from the experimental work and support the theoretical predictions. TEM is particularly effective for providing a direct image of nanoaggregates and clusters in asphaltene samples. As shown in Fig. 2A, with the addition of DBSA NEs, the asphaltene cluster size is significantly decreased to about (5 nm), while the average interlayer spacing between aromatic rings is increased, indicating a gradual stabilization process. Fig. 2B demonstrations the selected area electron diffraction (SAED) images of

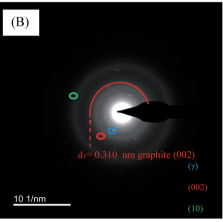
asphaltene with and without DBSA NEs, which further indicated the transformation of asphaltene sample from unstable to stable by reduction of the π - π stacking distance between asphaltene molecules. The interlayer distance between the periodic lattice fringes from the (002) plane of asphaltene with DBSA NEs was approximately 0.310 nm. This suggests that DBSA NEs allow significant change about 22.2% increase in the stacking distance between aromatic rings of asphaltene. These are in very good agreement with the findings obtained from our XRD that indicate the distance between aromatic sheets of asphaltene molecules is reduced in the presence of DBSA NEs.

3.2. The intermolecular interactions between asphaltene and DBSA NES

FTIR is one of the most important analytical techniques, which is used for the determination of functional properties of asphaltene. Despite its versatility, the study of the intermolecular interactions of asphaltene has received little attention by using this technique.32 In addition, the literature has been rarely cited about quantitative32-34 and qualitative35,36 analyses for the functional resolution of asphaltenes. The importance of FTIR studies not only to identify the existence and quantity of functional groups but also to provide further information such as the ratio of CH2 to CH3 and the relative amount of aliphatic and aromatic groups, which are essential criteria for crude oil. 37 Fig. 4 shows the absorption spectrum of extracted asphaltene obtained from FTIR, which showed similar functional groups as reported in other studies.38-40 The characteristic peaks of asphaltene molecular structure can be categorized into three types: aromaticity characteristic, aliphaticity characteristic, and polar functionality.39 Especially, for the aromaticity characteristic, the adsorption peaks at 750, 808, and 866 cm⁻¹ linked to the out-of-plane C-H bending in 1,2-disubstituted aromatic, 1,4-substituted aromatic, and 1,3-disubstituted aromatic,







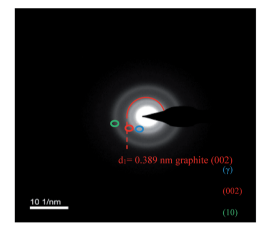


Fig. 2 (A) TEM image of asphaltene, and (B) HRTEM image of asphaltene in the presence of DBSA NEs. The insets are the selected area electron diffraction pattern (SAED), which show the stacking distance measured $d_{002} = 0.310$ and $d_{002} = 0.389$ for asphaltene without and with the presence of DBSA NEs respectively.

respectively, while the C=C stretching vibration in the aromatic structure was allocated at 1602 cm $^{-1}$. On the other hand, the aliphatic peaks at 2922 and 2852 cm $^{-1}$ were represented C–H stretching vibrations of CH $_2$ and CH $_3$, while the C–H bending vibration of CH $_2$ and CH $_3$ peaks were allocated at 1458 and 1375 cm $^{-1}$. For polar functionality, the absorbance peak at 3458 cm $^{-1}$ was represented stretching vibrations of –OH and/or –NH, while the absorbance peak at 1031 cm $^{-1}$ was signified stretching vibrations of S=O in sulfoxides. All functional groups present in are listed in Table S2 (ESI†).

Looking at Fig. 3, it is apparent that DBSA NEs spectra showed broad strong absorption peak assigned at 3365 cm⁻¹ and this can be attributed to the formation of hydrogen bonds of interlayer water molecules with an anion (DBSA) as well as with hydroxyl groups of non-ion of Tween 80 and SDBS. The colloidal structure of asphaltene can be modified by the addition of DBSA NEs, which provide multiple intermolecular interactions between the asphaltene molecules and the components of DBSA NEs. The ability of DBSA NEs in reducing π - π stacking and hence increase the stability of asphaltene can be mainly attributed to two mechanisms: acid-base interaction and hydrogen bond interaction. Further investigation for the hydrogen bond of asphaltene and -OH functional group of DBSA NE molecules was carried out by observing the -OH peak at the wave range of 3150 cm⁻¹ to 3650 cm⁻¹. First, the

absorbance of free and hydrogen bonded –OH of different concentrations of DBSA NE at carbon tetrachloride (CCl_4) was characterized. The hydrogen bonded –OH band started at 75 vol% of DBSA NE in CCl_4 that is the initial point of the formation of hydrogen bond of DBSA NE molecules. Then this solution (DBSA NE and CCl_4) was added to asphaltene solution (0.5 w/v % of asphaltene dissolved with 60/40 vol% Heptol) and FTIR spectrum was collected (Fig. 4).

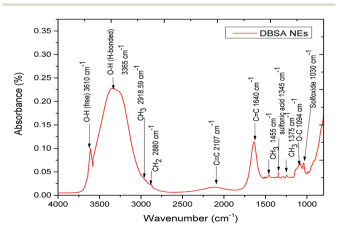


Fig. 3 FTIR of DBSA NEs with 25 vol% of carbon tetrachloride.

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asphaltene with DBSA NEs 2918.59 cm Amine 804 cm 0.30 0.25 2940 cm 2849.74 cm Absorbance (%) 0.20 0.10 3500 3000 2500 2000 1500 1000 4000 Wavenumber (cm⁻¹)

Fig. 4 FTIR of asphaltene with and without DBSA, showing the characteristic bands

Fig. 4 also presented that, after the addition of asphaltene, the intensity of free –OH bond at 3606 cm⁻¹ disappeared, while the peak of hydrogen-bonded –OH at 3365 cm⁻¹ improved. These results indicate that the addition of DBSA NEs to asphaltenes can provide strong hydrogen bonding of free –OH groups in the DBSA NEs with asphaltene molecules. In addition, the integrated areas under the FTIR spectrums of asphaltene with the presence and absence of DBSA NEs were measured after curve deconvolution and normalizing to quantitatively evaluate any modification. The results show that the integral area of asphaltene with DBSA NEs spectra increases from 2.58 to 5.35, indicating that asphaltene forms hydrogen bond with DBSA NEs.

Furthermore, the addition of DBSA NEs to asphaltene largely exhibited several additional peaks. For instance, the peak around 1300-1000 cm⁻¹ from the O=S=O stretching vibration shifts of the sulfonic group stretching vibration of asphaltene with DBSA NEs. This should be due to the interactions between the surface heteroatoms of asphaltene and the functional groups of DBSA. Thus, it can be suggested that the release of DBSA molecules can interact with asphaltene and absorb onto the surfaces, via hydrogen bond of O=S=O with O-H or C-H. Besides, the peaks between 660 and 900 cm⁻¹ were higher than for asphaltene without DBSA NEs, which were caused by the intensity vibration of NH3 and N-H wagging (shifts on Hbonding). The peak intensity at 1740 cm⁻¹ represented C=O (saturated aldehyde) or cis-RCH=CHR. The changes in out-ofplane O-H bonding in the aromatic rings of asphaltene appeared at 675 cm⁻¹, while the peak intensity at 1510 cm⁻¹ caused by NH2 scissoring.

As discussed in the previous study,²⁷ this system can overcome asphaltene intramolecular forces, creating new spaces, and involves multiple interactions between asphaltene and DBSA NE components including acid-base interactions, hydrogen bonding, polar- π interactions, cation- π interactions, and π - π interactions. These forces can provide additional electrostatic interactions with other ion-pair. For instance, hydrogen bond can be formed between asphaltene and DBSA NE due to the presence of the hydroxyl group in the chain length

with water and surfactants, while acid-base interaction can be appeared due to the presence of the nitrogen in the aromatic core of asphaltene. Thus, the use of a small concentration of DBSA with other components of DBSA NEs is adequate to provide complete coverage of the asphaltene nanoaggregates.

The quantitative comparison of major functional groups in asphaltenes can be investigated by measuring the integrated area of peaks and the intensities of a specific wavelength. To calculate the number of carbon atoms per alkyl side chain, the molar ratio of CH₂/CH₃ is required which can be attributed to the ratio of absorbance intensities at wavelengths of 2927 and 2957 cm⁻¹. Also, this ratio can be calculated from the length and the degree of the aliphatic chain by the eqn (1):⁴⁰⁻⁴³

$$N = \frac{n(\text{CH}_2)}{n(\text{CH}_3)} = \frac{A_{2927}}{A_{2957}} \times K \tag{1}$$

where K is a constant and specified to be 1.243 from correlating of 20 chemicals.⁴⁰ In addition, with the absorbance intensities of 1450 and 1600 cm⁻¹, the molar ratio of aromatic rings to aliphatic side chains can be correlated by eqn (2):

$$R = \frac{n(\text{aromatic C=C})}{n(\text{aliphatic C-H})} = \frac{A_{1600}}{A_{1450}}$$
 (2)

In addition, in order to know how many aliphatic and aromatic groups in asphaltene, the ratios of the integrated areas I_1 (2500–3000 cm⁻¹/1600 cm⁻¹) and I_2 (2500–3000 cm⁻¹/630–930 cm⁻¹) should be calculated.⁴⁴

Table 3 shows the values of N and R for asphaltene with and without DBSA NEs samples. It can be seen from these results that asphaltene with DBSA NEs has lower the higher number of carbon atoms per alkyl side chain and it also has lower aromaticity compared with pure asphaltene. It means that the pure asphaltene is the most aromatic sample, while the asphaltene with DBSA NEs is the least aromatic sample. On the other hand, asphaltene with DBSA NEs shows a higher ratio of I_1 and I_2 than asphaltene without DBSA NEs, indicating the shorter side chain of aliphatic substituents and more amounts of aliphatic and aromatic groups.

The tendency of asphaltene toward the formation of hydrogen bonding can be evaluated by calculating the ratio of the N-H and O-H intensities (3435 cm⁻¹) over the intensity of aromatic fraction (3100 cm⁻¹). These values in Table 3 imply that the pure asphaltene has the highest propensity toward aggregation by forming hydrogen bonding. The length of alkyl side chain can be attributed to the ratio of the intensities at a wavelength of 1455 and 1376 cm⁻¹. On the other world, the higher ratio means longer side chain. These results are in very good agreement with the aromaticity results obtained from XRD experiments.

Table 3 Asphaltene structure parameters with and without DBSA NEs from FTIR

Sample	N	R	I_1	I_2	I_{3435}/I_{3100}
Asphaltene	2.2	0.5	1.29	0.85	0.23
With DBSA NE	1.8	0.42	1.52	1.05	0.188

3.3. Mechanism on the effect of DBSA NEs on asphaltene

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structure

According to the experimental results, a possible mechanism was proposed and discussed as followed. The alteration of the asphaltene structure with the addition of DBSA NEs can be attributed to the interactions between the asphaltene molecules and the components of DBSA NEs. Therefore, the ability of DBSA NEs to stabilise asphaltene can be ascribed to several mechanisms such as (i) the slow release of DBSA from NEs can provide the electrophilic addition reaction between DBSA and the π electron of asphaltenes by initial breaking its SO₃-H bond and then bonding its dodecylbenzene sulfonate group with asphaltene molecules. It is worth noting that the DBSA NEs exhibit effective release kinetics due to the use mixtures of anionic and non-ionic surfactants, which can pack closely at the O/W interface and thereby make DBSA diffusion from NE slower. Moreover, the slower release rate can be attributed to the small size of DBSA NE (i.e., 21 nm), which can provide a high surface area of the system. In fact, higher surface area means a higher surface from which the inhibitor can be released; (ii) the presence of the relatively large lipophilic group (i.e., long-chain alkyl) of DBSA molecules formed adsorbed DBSA like a wedge with a thick tail in the periphery of the new structure.45 Similarly, the molecular structures of Tween 80 and SDS can shape themselves similar to lipophilic group of DBSA molecules that can prevent the π - π interactions between asphaltene molecules by providing steric interferences in those interactions, which therefore prevent asphaltene aggregation. The effectiveness of asphaltenes inhibitors is depended on the length and shape of aliphatic side chains. With increasing in the aliphatic chain length, the inhibitors become able to surround asphaltene molecules strongly; resulting in higher efficiency of the inhibitors;46,47 (iii) aromatic compounds of DBSA NEs (i.e., Tween 80, SDS, and xylene) with a certain rigidity of the DBSA head group are able to interact with asphaltene nanoaggregates by π - π and hydrogen-bond interactions; (iv) H2O can also provide an electrophilic addition to the asphaltene's π -electron by the catalysis of acids (e.g., DBSA) and cause the absorption band in the range of 2000-3000 cm⁻¹ as well as H₂O molecules can act as a bridge between asphaltene molecules by hydrogen bonding between H2O molecules and heteroatoms in asphaltenes. Fig. 5 shows asphaltene molecules with two different nano-colloidal species are achieved after asphaltene molecules stacking. The smaller species is a nanoaggregate of small aggregation number with an aromatic core and an alkane shell. Another species is a cluster with a small aggregation number of nanoaggregates. Whereas the overall length scales for these nano-species are obtained from TEM and XRD results share many agreements with the Yen-Mullins model.48 However, our proposed model involves the molecular structure of asphaltene after the addition of AI or NEs loaded with AI and this must be explicitly accounted for the energetics of aromatic systems, as discussed.

3.4. Thermogravimetric analysis (TGA)

The refractory nature of asphaltenes can be confirmed with the wide difference in weight loss, which is relatively insensitive to

the temperature. The combustion of pure asphaltenes and asphaltene with DBSA NEs in the presence of air causes weight loss, which has been measured by thermogravimetric analysis and it is presented in Fig. 6A. The presence of air is important to start the oxidation reaction; thus, at a higher temperature, the decomposition rate is very fast. This figure demonstrates that the behaviour of asphaltenes towards coke formation is totally burned off, showing a maximum weight loss at 600 °C. Similar findings were found by Savel'ev et al.,49 and Maity et al.,50 using air in the degradation of asphaltene and functionalized asphaltene. However, in the presence of air, about 97 wt% of another sample (asphaltene with DBSA NEs) is burned at around 600 °C.

The derivative curve clearly shows that two types of asphaltene particles are present due to two main weight losses at 460 $^{\circ}$ C and at 585 $^{\circ}$ C (Fig. 6B). The first one is named as soft asphaltene and is burned at a lower temperature. While the second one is named as hard asphaltene because it is burned at a higher temperature. These findings are similar to those reported in the literature which suggested the presence of two types of coke in asphaltene sample. 50-52 Asphaltene particles are subjected to very severe conditions above 450 °C, leading to form coke as final residue.53 Total volatilization is observed at 550 °C; at this temperature and with the presence of oxygen in the air, asphaltene was fully burned and the maximum weight loss is achieved. In addition, at this temperature, intermolecular interactions and weaker chemical bondings were destroyed such as sulfur bridges. However, asphaltene particles are not only changed to coke but also lighter compounds such as gases, oils and resins are expected to be formed. The final residue from asphaltene decomposition was 12.57% (2.33 mg) of the initial mass, which may consist of typical of polycyclic condensed aromatic hydrocarbon compounds with large aggregates as well as different elements such as Ni, Fe, and V that are stable at higher temperatures.

Comparing the thermal profile of asphaltene with the asphaltene with DBSA NEs, the derivative curve clearly develops three consecutive weight loss processes at around 300 °C, 460 °C, and at 585 °C, donating that three types of asphaltene particles are present. TGA curves show that asphaltene with DBSA presented an oscillation peak from 50 to 200 °C, this may be due to the loss of light organic compounds and the residual water (dehydration) obtained from the addition DBSA NEs to the asphaltene. The water molecules directly interact with asphaltene molecules via hydrogen bonding. Hence, it is noteworthy that the loss of water molecules from asphaltene is associated with minor structural changes; however, the framework is kept as the overall shape of the pattern changes little between ambient temperature and 220 °C. The monohydrate phase up to 200 $^{\circ}\mathrm{C}$ is stable.

After 200 °C asphaltenes with DBSA NEs suffer continuous decomposition up to around 600 °C. This can be attributed to releasing CO, CO2 and some hydrocarbons from asphaltene as well as to the loss of long alkyl chains of DBSA NEs components which linked with asphaltene molecules. However, the 2nd peak of the asphaltene sample appeared at 550 °C and it was Paper

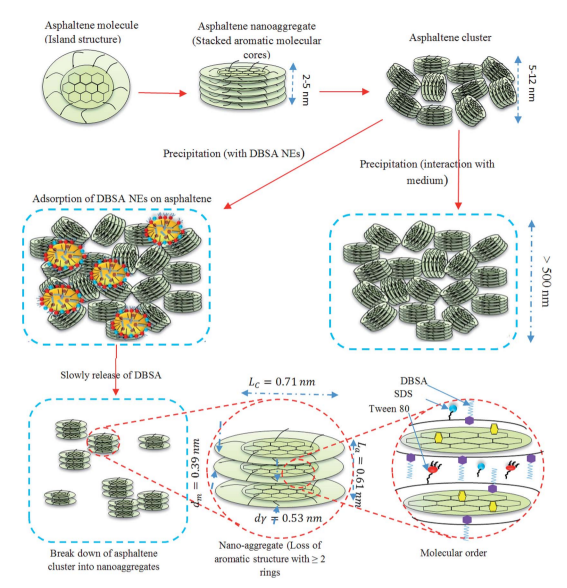


Fig. 5 Schematic representation of the progression of asphaltene molecule and formation of nanoaggregates, with most aging being seen in larger asphaltene clusters (or primary particles). The mechanisms of the adsorption of DBSA NEs on asphaltene and the slow release of DBSA from nanoemulsion affect the stacking distance and the cluster size.

shifted to a higher temperature around 585 °C for the sample asphaltene with DBSA NEs. The residue at the highest temperature for asphaltene with DBSA is very low about 4.71%, suggesting that formation of multiple intermolecular interactions like hydrogen bond, acid–base interaction, and π – π interaction between the adjacent molecules of DBSA NEs with asphaltene molecules, which give rise to the high temperature.⁵⁴

The formation of coke from asphaltene decreased when DBSA NEs were added to asphaltene, indicating that DBSA NEs tended to stabilize a proportion of asphaltene. The DBSA NEs can promote the loss of labile points of long alkyl chains in DBSA NEs components which linked with asphaltene molecules, releasing more volatile compounds for example gases. In the initial stages, many of lighter compounds present in asphaltene with DBSA NEs have been volatilized; however, only a few amounts of asphaltene remains during the reaction

to finally residue towards coke formation in a small amount compared with pure asphaltene. This can explain why asphaltene with DBSA NEs continuously change in a wide range of temperatures (200-585 °C). This behaviour may indicate that the yield of coke is affected by the presence of DBSA NEs components, reducing it around 62%. Also, the similarities in the overall shape of the weight loss curves not only confirm that DBSA NEs did not change the main degradation pathways of asphaltene but also indicate that no new materials are formed due to the interaction of DBSA NEs with asphaltene. However, the nature and the structure of hard asphaltene become softer when this is treated with the DBSA NEs as well as the coke yield is decreased. These findings of TGA are in very good agreement with the other results of XRD, TEM, and IFIR for both asphaltenes with and without DBSA NEs presented in this work.

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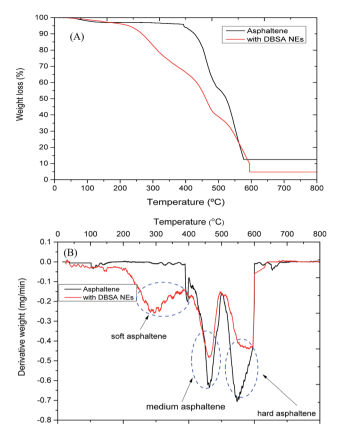


Fig. 6 (A) TGA, and (B) the derivative weight of asphaltene and asphaltene with DBSA NEs in air.

4. Conclusions

This work examined the detailed molecular structure of asphaltene in the presence of NEs, and proposed a mechanism of modification structure and stabilizing asphaltene. The experiments by using XRD and TEM confirmed the validity of the mechanism of stabilizing asphaltene by the release of DBSA and other components from NE, which were able to increase both the stacking distance between aromatic rings (about 22.2%) and aliphatic chains, and reduce the aromatic sheet diameter and the cluster size. These alterations in molecular structure lead to a decrease of the aromaticity of asphaltene, which in turn increase the solubility of aromatic compounds in the oil and improve the liberation of asphaltene aggregates. In addition, the π system over the aromatic zone is changed due to the disruption of the aromaticity that will, in turn, prevent eventual π - π stacking between asphaltene molecules, which is the main mechanism responsible for the formation of nanoaggregates and clusters.

The FTIR spectroscopic study verified that DBSA NE involves multiple interactions with asphaltene, building up a multicentric electron density in this region, which leads to deeper isosurfaces of van der Waals forces in the interacting zone. In addition, the quantitative study indicated that asphaltenes with DBSA NEs have a lower number of carbon atoms per alkyl side chain, lower aromaticity, shorter side chain of aliphatic

substituents, and more amount of aliphatic and aromatic groups compared with pure asphaltene.

The TGA results further demonstrated the alteration of asphaltene structure. It was found that during the chemical reaction between asphaltene and DBSA NEs, the structure of asphaltene is improved considerably and the coke yield is decreased around 62% due to the decrease of the cluster sizes and the increase of the stacking distance between aromatic sheets.

The findings of this study can contribute to the developments in deep in the enhanced oil recovery process (EOR), finishing processes, and the refining industry in general, as asphaltenes are the heaviest and most problematic hydrocarbon components in crude oil. Revealing the impact of NE to deliver and controlled release of AI on the molecular structure of asphaltene should help our understanding to suggest the best upgrading scheme for heavy crude oils, which is expected to be as a prospective method for preventing oilfield asphaltene.

Conflicts of interest

There are no conflicts to declare.

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