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Published in: Energy and Fuels

Link to article, DOI: 10.1021/acs.energyfuels.6b02767

Publication date: 2017

Document Version Peer reviewed version

Link back to DTU Orbit

Citation (APA):

Arya, A., Liang, X., von Solms, N., & Kontogeorgis, G. (2017). Modeling of Asphaltene Precipitation from Crude Oil with the Cubic Plus Association Equation of State. Energy and Fuels, 31(2), 2063-2075. DOI: 10.1021/acs.energyfuels.6b02767

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## Modeling of Asphaltene Precipitation from Crude Oil with the Cubic Plus Association Equation of State

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In this study, different modeling approaches using the Cubic Plus Association (CPA) equation of state (EoS) are developed to calculate the asphaltene precipitation onset condition and asphaltene yield from degassed crude oil during the addition of *n*-paraffin. A single model parameter is fitted to calculate the asphaltene onset condition during the addition of different nparaffin precipitants (n-pentane to n-hexadecane). Three parameters per precipitant are fitted to calculate the asphaltene yield during the addition of the precipitant. The results obtained from the model are compared with the experimental data for eight different crude oils. Results were also obtained for seven crudes using the Perturbed Chain Statistical Association Fluid Theory (PC-SAFT) EoS based approach described in the literature. The CPA EoS based approaches treat the asphaltene fraction as an associating component whereas the PC-SAFT EoS based approach does not. A comparison between the approaches shows that the CPA EoS based approaches, developed in this work, give more reliable results. The predictions from the PC-SAFT EoS based approach result in behavior that is unphysical: the mole fraction of nparaffin (at the precipitation onset) and the asphaltene yield do not decrease with the carbon number of *n*-paraffin. Furthermore, it is shown that the approach, developed in this work, can predict the onset conditions of asphaltene precipitation resulting from a blend of two or more crudes.

## 1. Introduction

A crude oil at room temperature can be divided into maltene (deasphalted fraction) and asphaltene fractions. The maltene fraction can be further divided into saturates, aromatics and resins fractions. The SARA (saturates, aromatics, resins, asphaltenes) analysis is used to measure the amount (on weight basis) of these four solubility fractions. All of these fractions are polydisperse in the oil and contain a large number of components. Asphaltenes are defined as the heaviest fraction of the oil, which is not soluble in *n*-pentane/*n*-heptane but soluble in aromatic solvents such as toluene [1-3]. Resins are the fraction of the oil, which is soluble in ethyl acetate at room temperature [2,4]. It is believed that resins are cross-associating (form hydrogen bonds) with asphaltenes to

keep asphaltenes soluble in the oil [5]. The asphaltene fraction may precipitate out of the crude oil during the addition of a paraffinic component. As the carbon number of *n*-paraffin increases, the solubility of the asphaltene fraction increases and the amount of precipitated asphaltenes decreases. For the higher carbon number (>11) *n*-paraffins, the amount of precipitated asphaltenes remains almost constant [5]. The asphaltenes containing crude is generally blended with one or more crudes to maintain the properties of feedstock (blended crude) to the refinery. In certain cases, the blending of incompatible crudes results in the asphaltene precipitation. The asphaltene precipitation can result in fouling and coking of preheat trains, heat exchangers and pipestill furnace tubes in the refinery [6]. Therefore, it is important to understand the asphaltene precipitation in order to avoid complications during the processing of crude oil.

There are several studies on the modeling of the asphaltene precipitation from the crude oil. Some of them consider the asphaltene fraction as polydisperse while others consider it as monodisperse. These studies can also be classified according to whether the asphaltene fraction is treated as a non-associating or an associating molecule. When the asphaltene molecule is considered not to associate with other asphaltene molecules, it is assumed that the asphaltene component already exists in a pre-aggregated form and only van der Waals forces are included in the asphaltene precipitation model. On the other hand, when the asphaltene molecule is allowed to associate with other asphaltene molecules, it is assumed that the asphaltene molecule exists as a monomer. In this case, the association between asphaltene molecules is an important criteria for precipitation, along with the van der Waals forces. This association is a way to account for strong polar forces such as hydrogen bonding. The framework of both PC-SAFT and CPA EoS allows the modelling of both van der Waals and association forces. However, we point out that the association term is not considered in this work for the PC-SAFT based model, where we follow the approach used previously in the literature [7].

## Asphaltene as a Non-associating and Pre-aggregated Component

The asphaltene precipitation from the crude oil has been modeled by several researchers. Hirschberg et al [2] used the Flory-Huggins theory in order to study the effect of miscible gas flooding on the asphaltene precipitation. They used the Soave-Redlich-Kwong (SRK) EoS for vapor-liquid equilibrium considering no asphaltene precipitation. They then used the Flory-Huggins model to calculate the asphaltene precipitation from the liquid phase, calculated from the SRK EoS. The asphaltene fraction was considered as a monodisperse component. Alboudwarej et al [8] used the regular Flory-Huggins model dividing the asphaltene fraction into multiple components of different molar mass, volume and solubility parameters. They assumed that the precipitated phase is ideal and consists of only the asphaltene components.

Akbarzadeh et al [9] used the same approach developed by Alboudwarej et al [8] and showed the temperature and pressure effect on the model predictions. They used a temperature dependent gamma distribution function to describe the temperature dependent molar mass distribution of the asphaltene fraction. Pazuki and Nikookar [10] modified the Flory-Huggins model and compared the asphaltene precipitation results with the original Flory-Huggins model and experimental data. They concluded that the modified model is better than the original one. Wiehe et al [6] developed the oil compatibility model in order to check whether the blending of two or more crudes is compatible with respect to the asphaltene precipitation. Wiehe et al [11] compared the oil compatibility model with the regular Flory-Huggins theory based model from Yarranton and co-workers [8, 9]. They concluded that both models can calculate the asphaltene onset condition as well as the maximum in volume of *n*-paraffin, as a function of the carbon number of *n*-paraffins at the onset of asphaltene precipitation. They also mentioned that the basic assumption of the oil compatibility model, that the solubility parameter of mixture at the onset is constant, contradicts the physical behavior. Tharanivasan et al [12] used the regular Flory-Huggins theory based model, developed by Yarranton and co-workers [8, 9], to calculate the asphaltene yield (ratio of mass of precipitated asphaltenes to mass of oil before addition of precipitant) as well as the onset condition of the blend of different crudes. They considered the asphaltene fraction as polydisperse components. Sabbagh et al [13] used the Peng-Robinson (PR) EoS. They divided the asphaltene fraction into multiple components of different molar masses based on the gamma distribution function. The precipitated phase was assumed to contain only the asphaltene components. They concluded that the PR model is not a universal predictor and not better than the previously developed regular solution model. Panuganti et al [14] and Tavakkoli et al [7] used a model, based on the PC-SAFT EoS, considering the asphaltene fraction as polydisperse components. They studied both light and heavy crudes. They used interaction parameters for the asphaltene-n-paraffin binary pairs in order to correlate both the onset conditions and asphaltene yield from the light crude. They kept the binary interaction parameters for asphaltene-n-paraffins binary pairs to a zero value in order to predict the asphaltene yield from the heavy crude, however, they used non-zero values for the binary interaction parameter of the asphaltene-saturates binary pair in order to correlate the onset conditions. Their model predictions contradict the general experimental observation [2,15,4], where the amount of precipitated asphaltenes decreases or remains constant with the carbon number of *n*-paraffin precipitant. This contradiction is discussed in detail in the "Results and Discussion" section of this work.

## Asphaltene as an Associating and Monomeric Component

Wu et al [16,17] used the SAFT EoS in the framework of McMillan-Mayer theory. In their model, asphaltenes and resins are considered as monodisperse pure pseudo-components and the interactions between them are screened through the continuous medium of solution containing

other components. They applied their model to calculate the asphaltene yield from the crude oil and the onset conditions from the reservoir oil. However, they did not study the onset conditions for the crudes during the addition of *n*-paraffin. Buenrostro-Gonzalez et al [15] modified the modeling approach of Wu et al [16,17] by using the SAFT-VR instead of the SAFT EoS. They applied their model to calculate both the onset conditions and asphaltene yields from the crude. However, they could not match both types of experimental data with the single set of parameters. Li and Firoozabadi [18] developed a modeling approach based on the CPA EoS considering the asphaltene fraction as a monodisperse component. They studied the asphaltene precipitation from the heavy oil. They did not study their modeling approach with respect to the asphaltene onset condition from the crude oil.

To our knowledge, nobody has so far studied the CPA equation of state for the asphaltene precipitation from the crude considering the asphaltene fraction as polydisperse. There is also no study on the blending of crudes with the CPA and PC-SAFT equations of state. In this study, different CPA approaches are developed to study the asphaltene yield and onset conditions. The PC-SAFT approach is referred from Tavakkoli et al [7]. Both CPA and PC-SAFT approaches are studied and compared with the experimental data. A total of eight crudes are studied. Crude-1 is studied with respect to both the asphaltene yield and onset conditions considering the asphaltene fraction as a polydisperse fraction. Crude-2 to 4 are studied with respect to asphaltene yield considering the asphaltene fraction as a polydisperse fraction, while Crude-5 to 8 are studied with respect to onset conditions considering the asphaltene fraction. The brief introduction about the CPA EoS is given here in order to familiarize the reader about model parameters. The reader can refer to the supplementary information for more details on the PC-SAFT EoS.

## 2. CPA EoS

The CPA EoS, proposed by Kontogeorgis et al [19], can be expressed for mixtures in terms of pressure P, as shown in equation (1). Please consult the "List of Symbol" and "Greek Letters" sections for the meaning of each symbols used in the following equations.

$$P = \frac{RT}{V_m - b} - \frac{a(T)}{V_m(V_m + b)} - \frac{1}{2} \frac{RT}{V_m} \left(1 + \rho \frac{\partial \ln g}{\partial \rho}\right) \sum_i x_i \sum_{A_i} (1 - X_{A_i})$$
(1)

$$a(T) = \sum_{i} \sum_{j} x_i x_j a_{ij}(T)$$
(2)

$$a_{ij}(T) = \sqrt{a_i(T)a_j(T)} (1 - k_{ij})$$
(3)

When the CPA EoS is used for mixtures, the conventional mixing rules are employed in the physical term (SRK) for the energy and co-volume parameters. The geometric mean rule is used for the energy parameter  $a_{ij}(T)$ . The interaction parameter  $k_{ij}$  is the only binary adjustable parameter of CPA in the applications for self-associating mixtures, e.g. glycol with n-alkanes.

The energy parameter of the EoS is given by a Soave-type temperature dependency, while *b* is temperature independent:

$$a_{i}(T) = a_{0,i} \left[ 1 + c_{1,i} \left( 1 - \sqrt{\frac{T}{T_{c,i}}} \right) \right]^{2}$$
(4)

where  $T_{c,i}$  is the critical temperature of the *i*<sup>th</sup> component.

The key element of the association term is  $X_{A_i}$ , which represents the mole fraction of site-A in molecule of component *i* not bonded to other sites, while  $x_i$  is the mole fraction of component *i*.  $X_{A_i}$  is related to the association strength  $\Delta^{A_iB_j}$  between two sites belonging to two different molecules, e.g. site A on molecule *i* and site B on molecule *j*, determined from:

$$X_{A_i} = \frac{1}{1 + \rho \sum_j x_j \sum_{B_j} X_{B_j} \Delta^{A_i B_j}}$$
(5)

where the association strength  $\Delta^{A_iB_j}$  in CPA is expressed as:

$$\Delta^{A_i B_j} = g(\rho) \left[ exp\left(\frac{\varepsilon^{A_i B_j}}{RT}\right) - 1 \right] b_{ij} \beta^{A_i B_j}$$
(6)

where:

$$b_{ij} = \frac{b_i + b_j}{2} \tag{7}$$

with the radial distribution function:

$$g(\rho) = \frac{1}{1 - 1.9n}, n = \frac{1}{4}b\rho$$
(8)

In the expression for the association strength  $\Delta^{A_iB_j}$ , the parameters  $\varepsilon^{A_iB_j}$  and  $\beta^{A_iB_j}$  are called the association energy and the association volume, respectively. These two parameters are used only for associating components, and along with the three additional parameters of the SRK term ( $a_0$ , b,  $c_1$ ), they are the five pure-compound parameters of the model. They are obtained by fitting vapor pressure and liquid density data. For inert components such as hydrocarbons, only the three parameters of the SRK term are required, which can either be obtained from vapor pressures and liquid densities or be calculated in the conventional manner (from critical data, acentric factor).

Following are the combining rules (CR), which are required for cross-associating systems. CR-1 and Elliott CR are generally used. Modified CR-1 and Customized CR-1 are used for solvating systems only.

(**-**)

(i) CR-1:

$$\varepsilon^{A_i B_j} = \frac{\varepsilon^{A_i B_i} + \varepsilon^{A_j B_j}}{2}, \quad \beta^{A_i B_j} = \sqrt{\beta^{A_i B_i} \beta^{A_j B_j}}$$
(ii) Elliott CR:
(9)

 $\Delta^{A_i B_j} = \sqrt{\Delta^{A_i B_i} \Delta^{A_j B_j}} \tag{10}$ 

(iii) Modified CR-1:

Cross association volume  $\beta^{A_iB_j}$  is fitted to experimental data and CR-1 rule is used for cross association energy  $\epsilon^{A_iB_j}$ .

## (iv) Customized CR-1:

Cross association volume  $\beta^{A_iB_j}$  and cross association energy  $\epsilon^{A_iB_j}$  are fitted to experimental data.

## 3. Modeling Approaches

#### Approach-1

Approach-1 is based on the CPA EoS and is the same approach as was proposed by Arya et al [20-22]. As shown in Table 1, the modeling approach-1 has a single component for the maltene fraction and six components for the asphaltene fraction. The asphaltene components are considered self-associating with two positive and two negative sites on each asphaltene molecules. The maltene component's molecule has one association site and can crossassociate with both positive and negative sites of the asphaltene components. The maltene component is not self-associating. The critical temperature and pressure  $(T_c, P_c)$  and acentric factor ( $\omega$ ) of maltene component are calculated from the Kesler-Lee correlations [23] based on the information of normal boiling point, molecular weight (MW) and specific gravity (SG). The MW and SG of maltene component are calculated from the experimental information of MW and SG of the crude and asphaltenes MW (750 Da) and SG (1.2 g/cc). The normal boiling point is calculated from the Pedersen correlation [24]. The critical temperature and pressure  $(T_c, P_c)$ , acentric factor ( $\omega$ ) and self-association volume are constant for all six asphaltene components and are referred from Arya et al [20-22]. The self-association energies of asphaltene components (Asp-4 to Asp6) are also kept constant to reduce the number of model parameters. In addition the self-association energies of asphaltene components (Asp1 to Asp3) are

assumed to be same. The cross-association volume between the asphaltenes and maltene is also fixed to the value of 0.05.

Mitchell and Speight [4] mentioned that the asphaltene yield decreases with the carbon number of *n*-paraffin. However, approach-1 cannot predict this physical behavior and precipitate all asphaltene components (Asp1 to 6) irrespective of the carbon number of nparaffin. To overcome this issue, one more model parameter is incorporated, which is the crossassociation energy between the *n*-paraffin and asphaltene components (except Asp6). It means that Asp1 to Asp5 have the same cross-association energy with a given *n*-paraffin. By doing this, the solubility of Asp1 in *n*-paraffin is higher than those of other asphaltene components (Asp2 to Asp5) in *n*-paraffin since the self-association energy of Asp1 is lower than those of other asphaltene components. If the cross-association energy for a given *n*-paraffin increases, the solubility of all asphaltene components (with fixed self-association energies) in a given nparaffin increases. Therefore, we can assign the higher value of the cross-association energy for *n*-paraffin precipitant as the carbon number increases and can correlate the asphaltene yield data. Li and Firoozabadi [18] also studied the CPA EoS for the calculations of asphaltene yield from the crude and they suggested that the cross-association energy between (A+R) and asphaltenes is different for different n-paraffins. They have not considered any crossassociation between n-paraffin and asphaltenes. However, in our approach, the crossassociation energy between aromatics+resins (A+R)/maltene and asphaltenes is the same for different *n*-paraffins and the cross-association energy between *n*-paraffin and asphaltenes is different for different *n*-paraffins. Speight [25] mentioned that the asphaltene yield is almost constant for *n*-paraffin of higher carbon number (around >11). Therefore, Asp6 component is treated as an *n*-paraffin undissolved asphaltene fraction and does not cross-associate with the n-paraffin precipitant. Thus, approach-1 needs two additional model parameters (the crossassociation energy between the precipitant and asphaltene components and amount of Asp6). It should be noted that the binary interaction parameter  $(k_{ij})$  between the *n*-paraffin and asphaltenes pair could also be used instead of the cross-association energy between them. However, it needs some modification of the modeling approach and is not shown in this work. In addition, one could also fix the amount of Asp6 for all crudes, for example 30% of asphaltenes amount from SARA analysis, which would result in minor deviations from the experimental data. Table 2 shows that there are total three model adjustable parameters (MAP1, MAP2, and MAP3). The model parameter MAP3 is different for different *n*-paraffins.

Comp	Amount	MW	T <sub>c</sub>	P <sub>c</sub>	ω	Self- association energy $(\varepsilon^{AA}/R)$	Self- association volume $(\beta^{AA})$	No. Of Sites
	wt%	g/mol	K	bar	-	(K)	-	-
Maltene	Xa	Calc <sup>d</sup>	Calc <sup>d</sup>	Calc <sup>d</sup>	Calc <sup>d</sup>	0	0	1 <sup>f</sup>
Asp1	Yb	750	1040.1	15.44	1.535	MAP1 <sup>e</sup>	0.05	2+,2-
Asp2	Yb	750	1040.1	15.44	1.535	MAP1 <sup>e</sup>	0.05	2+,2-
Asp3	Yb	750	1040.1	15.44	1.535	MAP1 <sup>e</sup>	0.05	2+,2-
Asp4	Yb	750	1040.1	15.44	1.535	6500	0.05	2+,2-
Asp5	Yb	750	1040.1	15.44	1.535	6600	0.05	2+,2-
Asp6	Z ⁰ (MAP) ⁰	750	1040.1	15.44	1.535	7000	0.05	2+,2-

# Table 1 Approach-1: Crude oil characterization and components parameters.

<sup>a</sup> X is the sum of wt% of saturates, aromatics and resins from SARA analysis.

<sup>b</sup> Y = (Asp wt% from SARA - Z)/5.

<sup>c</sup> Z is the amount of asphaltene sub-fraction (Asp6), which is not dissolved in *n*-paraffin.

<sup>d</sup> Calculated (Calc) from the Kesler-Lee correlations as mentioned in the text.

<sup>e</sup> Model Adjustable Parameter (MAP) determined from the experimental data.

<sup>f</sup> Maltene molecule has one site, which can cross-associate with both positive and negative sites of asphaltene molecule.

#### Table 2

Approach-1: Self-association and Cross-association energy parameter in Kelvin.<sup>b</sup>

	Asp1	Asp2	Asp3	Asp4	Asp5	Asp6	Maltene	<i>n</i> -paraffin
Asp1	MAP1 <sup>a</sup>	Elliott	Elliott	Elliott	Elliott	Elliott	MAP2 <sup>a</sup>	MAP3 <sup>a</sup>
Asp2	Elliott	MAP1 <sup>a</sup>	Elliott	Elliott	Elliott	Elliott	MAP2 <sup>a</sup>	MAP3 <sup>a</sup>
Asp3	Elliott	Elliott	MAP1 <sup>a</sup>	Elliott	Elliott	Elliott	MAP2 <sup>a</sup>	MAP3 <sup>a</sup>
Asp4	Elliott	Elliott	Elliott	6500	Elliott	Elliott	MAP2 <sup>a</sup>	MAP3 <sup>a</sup>
Asp5	Elliott	Elliott	Elliott	Elliott	6600	Elliott	MAP2 <sup>a</sup>	MAP3 <sup>a</sup>
Asp6	Elliott	Elliott	Elliott	Elliott	Elliott	7000	MAP2 <sup>a</sup>	0
Maltene	MAP2 <sup>a</sup>	0	0					
<i>n</i> -paraffin	MAP3 <sup>a</sup>	0	0	0				

<sup>a</sup> Model Adjustable Parameters (MAP1, MAP2 and MAP3) determined from the experimental data of asphaltene yield and onset conditions.

<sup>b</sup> Cross-association volume is 0.05 for all pairs where Elliott combining rule is not used.

## Approach-2

Approach-2 is based on the CPA EoS and similar to approach-1, except for a few differences, and is shown in Table 3. In approach-2, the maltene fraction is divided into saturates and (A+R) components. The critical temperature and pressure  $(T_c, P_c)$ , acentric factor ( $\omega$ ) and MW are fixed for all components and are taken from Li and Firoozabadi [18]. For asphaltenes MW, the value of 750 Da is used in this study, whereas Li and Firoozabadi [18] used the value of 1800 Da. Sabbagh et al [13] showed the relationship between MW and critical temperature and pressure  $(T_c, P_c)$  and acentric factor ( $\omega$ ), which were then used by Li and Firoozabadi [18]. There are two

reasons for the development of approach-2 over approach-1. The first reason is to show that the maltene fraction can further be divided into two components (and possibly more) and model can still be fitted to the experimental data. The second reason is to use the different set of component parameters, derived based on the information of experimental MW from the vapor pressure osmometry technique, for the heavy oil from the literature. Table 4 shows that there are total three model adjustable parameters (MAP1, MAP2, and MAP3). The model parameter MAP3 is different for different *n*-paraffins.

## Table 3 Approach-2: Crude oil characterization and components parameters.

Comp	Amount	MW	T <sub>c</sub>	P <sub>c</sub>	ω	Self- association energy $(\varepsilon^{AA}/R)$	Self-association volume ( $\beta^{AA}$ )	No. Of Sites
	wt%	g/mol	K	bar	-	(K)	-	-
Saturates	X1 a	460	930	13.4	0.9	0	0	0
A+R	X2 <sup>b</sup>	660	1074	10.85	1.5	0	0.05	1 <sup>f</sup>
Asp1	Yс	750	1474	7.07	2	MAP1 <sup>e</sup>	0.05	2+,2-
Asp2	Yс	750	1474	7.07	2	MAP1 <sup>e</sup>	0.05	2+,2-
Asp3	Yс	750	1474	7.07	2	MAP1 <sup>e</sup>	0.05	2+,2-
Asp4	Yс	750	1474	7.07	2	6500	0.05	2+,2-
Asp5	Yс	750	1474	7.07	2	6600	0.05	2+,2-
Asp6	Z <sup>d</sup> (MAP) <sup>e</sup>	750	1474	7.07	2	7000	0.05	2+,2-

<sup>a</sup> X1 is the wt% of saturates from SARA analysis.

<sup>b</sup> X2 is the sum of wt% of aromatics and resins from SARA analysis.

<sup>c</sup> Y = (Asp wt% from SARA - Z)/5.

<sup>d</sup> Z is the amount of asphaltene sub-fraction (Asp6), which is not dissolved in *n*-paraffin.

<sup>e</sup> Model Adjustable Parameter (MAP1) determined from the experimental data.

<sup>f</sup> (A+R) molecule has one site, which can cross-associate with both positive and negative sites of asphaltene molecule.

#### Table 4

Approach-2: Self-association and Cross-association energy parameters in Kelvin.b

	Asp1	Asp2	Asp3	Asp4	Asp5	Asp6	saturates	A+R	n-paraffin
Asp1	MAP1 <sup>a</sup>	Elliott	Elliott	Elliott	Elliott	Elliott	0	MAP2 <sup>a</sup>	MAP3 <sup>a</sup>
Asp2	Elliott	MAP1 <sup>a</sup>	Elliott	Elliott	Elliott	Elliott	0	MAP2 <sup>a</sup>	MAP3 <sup>a</sup>
Asp3	Elliott	Elliott	MAP1 <sup>a</sup>	Elliott	Elliott	Elliott	0	MAP2 <sup>a</sup>	MAP3 <sup>a</sup>
Asp4	Elliott	Elliott	Elliott	6500	Elliott	Elliott	0	MAP2 <sup>a</sup>	MAP3 <sup>a</sup>
Asp5	Elliott	Elliott	Elliott	Elliott	6600	Elliott	0	MAP2 <sup>a</sup>	MAP3 <sup>a</sup>
Asp6	Elliott	Elliott	Elliott	Elliott	Elliott	7000	0	MAP2 <sup>a</sup>	0
saturates	0	0	0	0	0	0	0	0	0
A+R	MAP2 <sup>a</sup>	0	0	0					
<i>n</i> -paraffin	MAP3 <sup>a</sup>	MAP3	MAP3 <sup>a</sup>	MAP3 <sup>a</sup>	MAP3 <sup>a</sup>	0	0	0	0

<sup>a</sup> Model Adjustable Parameters (MAP1, MAP2 and MAP3) determined from the experimental data of asphaltene yield and onset conditions.

<sup>b</sup> Cross-association volume is 0.05 for all pairs where Elliott combining rule is not used.

#### Approach-3

If we are only interested in calculating the precipitation onset condition, we can reduce the number of components and therefore the number of model adjustable parameters. Approach-3 is based on the CPA EoS and is similar to approach-2. Approach-3 has only one asphaltene component as shown in Table 5. There is no cross-association between the precipitant and asphaltene component but the cross-association between (A+R) and asphaltene components is present. Moreover, the self-association energy of asphaltene is fixed to 3000K (instead of 7000K) to show that the relative balance between the asphaltene and solvent is important rather than the absolute values of asphaltene self-association energy and cross-association energy of solvent. In other words, there can be multiple pairs of values of the asphaltene self-association energy (between (A+R) and asphaltene self-association energy and cross-association energy (between (A+R) and asphaltene self-association energy and cross-association energy (between (A+R) and asphaltene self-association energy (between (A+R) and asphaltene self-association energy (between (A+R) and asphaltene components), which can correlate the experimental data. The experimental values of MW are used, if they are available, without changing the critical temperature and pressure ( $T_c$ , $P_c$ ), acentric factor ( $\omega$ ) for saturates and (A+R) components. Table 6 shows that there is only one model adjustable parameter (MAP1).

#### Table 5

Approach-3: Crude oil characterization and components parameters.

Comp	Amount	MW	$T_c$	$P_c$	ω	Self- association energy ( $\varepsilon^{AA}/R$ )	Self-association volume ( $\beta^{AA}$ )	No. Of Sites
	wt%	g/mol	K	bar	-	(K)	-	-
Saturates	X1 a	460	930	13.4	0.9	0	0	0
A+R	X2 b	660	1074	10.85	1.5	0	0.05	1 <sup>d</sup>
Asp	X3 °	750	1474	7.07	2	3000	0.05	2+,2-

<sup>a</sup> X1 is the wt% of saturates from SARA analysis.

<sup>b</sup> X2 is the sum of wt% of aromatics and resins from SARA analysis.

<sup>c</sup> X3 is the wt% of asphaltene from SARA analysis.

<sup>d</sup> (A+R) molecule has one site, which can cross-associate with both positive and negative sites of asphaltene molecule.

#### Table 6

Approach-3: Self-association and Cross-association energy parameters in Kelvin.<sup>b</sup>

	Asp	saturates	A+R	n-paraffin
Asp	3000	0	MAP1 <sup>a</sup>	0
saturates	0	0	0	0
A+R	MAP1 <sup>a</sup>	0	0	0
<i>n</i> -paraffin	0	0	0	0

<sup>a</sup> Model Adjustable Parameter (MAP1) determined from the experimental data of asphaltene onset condition.

<sup>b</sup> Cross-association volume is 0.05 for all pairs.

#### Approach-4

Approach-4 is based on the PC-SAFT EoS and no association term is used since there is no associating component. Approach-4 is the same approach as was developed by Tavakkoli et al [7]. In approach-4, the crude is divided into saturates, A+R and multiple components of the asphaltene fraction. The number of asphaltene components depends upon the experimental data of different *n*-paraffins undissolved asphaltene amounts. For example, if the experimental data of *nC*5, *nC*7 and *nC*10 undissolved asphaltene amounts are available, three asphaltene components (nC5 insoluble but nC7 soluble asphaltene fraction, nC7 insoluble but nC10 soluble asphaltene fraction and nC10 insoluble asphaltene fraction) are used. The PC-SAFT parameters of saturates components are calculated from the MW correlations developed from the homologous series of *n*-paraffin. The standard *n*-paraffin PC-SAFT parameters from Gross and Sadowski [26] are used for this approach. The PC-SAFT parameters of the (A+R) and asphaltene components are also calculated from the MW and aromaticity ( $\gamma$ ) correlations developed from the polynuclear aromatic components by Gonzalez et al [27]. The zero value of aromaticity ( $\gamma$ ) corresponds to benzene derivatives and value of unity corresponds to polynuclear aromatics. Both correlations are mentioned in Table 7. The aromaticity value of (A+R) is tuned to match density of the crude. The aromaticity ( $\gamma$ ) values of all asphaltene components are assumed to be the same considering they have the same physical nature. The single value of aromaticity ( $\gamma$ ) and respective value of MW of all asphaltene components are tuned to match the precipitation data. In addition, binary interaction parameters  $(k_{ij}s)$  are also used to match the precipitation data. Tavakkoli et al [7] also tuned the segment diameter  $\sigma$  of asphaltenes to find a good match with the experimental data. However, the segment diameter  $\sigma$  is not used as an adjustable parameter in approach-4 in order to reduce the number of fitting parameters. When we are only interested in onset conditions, one asphaltene component, like approach-3, is used for approach-4 rather than dividing it into multiple components. Also, when the experimental MW of aromatics and resins fractions are available, (A+R) component can be divided into aromatics and resins components with the same value of aromaticity ( $\gamma$ ).

#### Table 7

Approach-4: Empirical correlations for the estimation of the PC-SAFT parameters for saturates and mixture of Polynuclear Aromatics (PNA) and benzene derivative components from Panuganti et al [14].

Correlation for Saturates	( $\gamma$ ) PNA correlation + (1 – $\gamma$ ) benzene derivatives correlation					
$m = 0.0257 \times MW + 0.8444$	$m = (1 - \gamma)[0.0223 \times MW + 0.751] + \gamma[0.0101 \times MW + 1.7296]$					
$\sigma(\text{\AA}) = 4.047 - \frac{4.8013 \times \ln(MW)}{MW}$	$\sigma(\text{\AA}) = (1 - \gamma) \left[ 4.1377 - \frac{38.1483}{MW} \right] + \gamma \left[ 4.6169 - \frac{93.98}{MW} \right]$					
$\ln(\epsilon/k)$ in $K = 5.5769 - \frac{9.523}{MW}$	$(\epsilon/k)$ in $K = (1 - \gamma)[0.00436 \times MW + 283.93] + \gamma \left[508 - \frac{234100}{(MW)^{1.5}}\right]$					

## 4. Results and Discussion

In this work, a total of eight different crudes are studied. The composition and properties of the crudes are mentioned in Table 8. Crudes-1, 2 and 8 are light crudes whereas Crudes-3 to 7 are relatively heavy crudes. The model parameters for the CPA EoS based approaches-1 to 3, calculated from the experimental data, and are shown in Tables 9 and 10. The model parameters and characterization of the crudes for approach-4 are shown in Tables 11 and 12. The binary interaction parameters  $(k_{ij}s)$  used for approach-4 are shown in Table 13. The experimental and calculated density using approach-4, after tuning the aromaticity ( $\gamma$ ) of A+R component, are compared in Table 14. For approaches-1 to 3, Peneloux volume corrections are used for the *n*-paraffin unless otherwise mentioned. It should be noted that no  $k_{ii}$  parameter is used for approaches-1 to 3. For all the crudes (except Crude-2), one of the CPA EoS based approaches-1 to 3 and the PC-SAFT EoS based approach-4 are compared with experimental data. For Crude-2, only approach-1 is studied. For Crudes-1, 3, and 4, the PC-SAFT parameters and oil characterization for approach-4 are referred from Tavakkoli et al [7]. For Crudes-5 to 8, the PC-SAFT parameters for approach-4 are from this work. Table 15 shows the comparison of the number of model adjustable parameters between the approaches for all the crudes. Approaches-1 and 2 are used for Crudes-1 and 2 and Crudes-3 and 4 respectively. One can also use either approach-1 or approach-2 for Crudes-1 to 4 but results are not presented here. Since we are only interested in onset conditions, approach-3 (instead of approaches-1 and 2) is used for Crudes-5 to 8.

Crude		1	2	3	4	5	6	7	8
Composition	Volatile	-	-	-	-	-	-	12.6	25.7
	Saturates	_ a	54.67	25	20.5	19.5	16.4	25.3	34.2
	Aromatics	<b>_</b> a	28.89	31.1	38	38.2	40.1	40.9	33.5
Composition	Resins	<b>_</b> a	12.66	37.1	19.6	26.8	28.7	4.9	3.2
	nC5 Asphaltenes	3.9	3.8	6.8	21.8	15.5	14.8	16.3	3.5
	solids	-	-	-	0.1	-	-	-	-
	Volatile	-	-	-	-	-	-	86	86
	Saturates	-	-	361	400	508	524	460	460
MW	Aromatics	-	-	450	508	522	550	522	522
(Da)	Resins	-	-	1108	1090	930	976	1040	1040
	Asphaltenes	-	-	7065	7662	2850	2910	-	-
	Crude	221.5	238.1	-	-	-	-	-	-
	Volatile	-	-	-	-	-	-	0.657	0.657
	Saturates	-	-	0.853	0.882	0.882	0.885	0.880	0.880
Density	Aromatics	-	-	0.972	0.997	0.995	1.003	0.990	0.990
(g/cc)	Resins	-	-	1.066	1.052	1.037	1.040	1.044	1.044
	nC5 Asphaltenes	-	-	1.192	1.193	1.203	1.203	-	-
	Crude	0.873	0.857	0.982	1.016	0.991	0.995	0.924	0.870

Experimental data of composition, MW and density of all crudes.

<sup>a</sup> Molar composition of saturates (50 mol%), aromatics (25 mol%) and hetero-compounds (25 mol%) are given in reference [2].

#### Table 9

The model adjustable parameters for Crudes-1 to 4 for the CPA EoS based approaches-1 and 2.

Crude	Self-association Energy $(\varepsilon^{AA}/R \text{ in K})$				Asp-6							
	Asp-1	Asp-2	Asp-3	Maltene	A+R	nC5	nC7	nC9	nC10	<i>nC</i> 12	nC16	(\vvv /0)
1	6450	6450	6450	3670	-	2400	2970	-	3135	3135	3135	1.2
2	6000	6000	6000	3480	-	2000	2610	2720	-	2920	-	0.5
3	6100	6100	6100	-	3440	1950	2380	-	-	-	-	0.5
4	6430	6430	6430	-	3580	2300	2550	-	-	-	-	2

The model adjustable parameters for Crudes-5 to 8 for the CPA EoS based approach-3.

Crude	Self-association Energy (K)	Cross-association Energy (K)				
	Asphaltenes	A+R	Toluene			
5	Default	1810	-			
6	Default	1845	-			
7	Default	1860	1410			
8	2600	1690	1260			

#### Table 11

Characterization of Crudes-1, 3 and 4 for the PC-SAFT EoS based approach-4 referred from Tavakkoli et al [7].

Componente	C	rude-1		Cr	Crude-3			Crude-4		
Components	MW (Da)	γ	wt%	MW (Da)	γ	wt%	MW (Da)	γ	wt%	
Saturates	207.43	0	46.8	361	0	25	400	0	20.5	
Aromatics	-	-	-	450	0.23	31.1	508	0.3	38	
Resins	-	-	-	1108	0.23	37.1	1090	0.3	19.6	
Aromatic+Resins	219.44	0.38	49.3	-	-	-	-	-	-	
<i>nC</i> 5 – <i>nC</i> 7 Asphaltenes	2900	0.2	2	4700	0.2	2.2	2500	0.4	4.3	
<i>nC</i> 7 − <i>nC</i> 10 Asphaltenes	2950	0.2	0.4	-	-	-	-	-	-	
<i>nC</i> 7 Asphaltenes	-	-	-	5600	0.2	4.6	2600	0.4	17.5	
<i>nC</i> 10 Asphaltenes	4307	0.2	1.5	-	-	-	-	-	-	

#### Table 12

The parameters for Crudes-5 to 8 for the PC-SAFT EoS based approach-4 from this work.

<b>.</b>		MW (Da	)	<u>.</u>	γ	
Crude	Saturates	A+R	Asp		A+R	Asp
5	508	637.3	2850		0.32	0.37
6	524	672.4	2910		0.32	0.37
7	460	800	6350		0.38	0.18
8	460	800	5200		0.30	0.25

The binary inte	eraction (k <sub>ij</sub> )	) parameters	s used for the	PC-SAFT	EoS based	approach-4 for	different	crudes. F	For
Crudes 1 and 4	1, parameter	rs are from T	avakkoli et al	[7].					
	-							-	

Crude	Component	nC5	nC7	nC10	nC12	nC16	Saturates
	Aromatic+ Resins	0.007	0.0065	0.006	0.006	0.005	0.007
1	<i>nC</i> 5 – <i>nC</i> 7 Asphaltenes	0.01 ª	0.007ª	0.006	0.005	0.005	0.003
I	<i>nC</i> 7 − <i>nC</i> 10 Asphaltenes	0.01 ª	0.007 a	0.006	0.005	0.005	0.003
	<i>nC</i> 10 Asphaltenes	0.01 ª	0.0075ª	0.0075ª	0.006 ª	0.006 ª	0.003
Λ	<i>nC</i> 5 − <i>nC</i> 7 Asphaltenes	0	0	0	0	0	-0.009
4	<i>nC</i> 7 Asphaltenes	0	0	0	0	0	-0.009
5,6,7,8	Asphaltenes	0	0	0	0	0	-0.02

<sup>a</sup> Modified in this work to decrease the deviations from the experimental data.

#### Table 14

The experimental density and calculated density using the PC-SAFT based approach-4, after adjusting aromaticity ( $\gamma$ ) of A+R, for Crudes-5 to 8.

Crude	Experimental Crude density (g/cc)	Calculated Crude density (g/cc)
1	0.873	0.872
3	0.982	0.951
4	1.016	0.997
5	1.001	1.002
6	0.995	1.011
7	0.928	0.927
8	0.871	0.841

The number of Model Adjustable (MAP) Parameters for all the approaches calculated from experimental data of asphaltene precipitation.

Number of Model Adjustable (MAP) Parameters calculated from experimental asphaltene precipitation data								
MAP	Crude-1	Crude-2	Crude-3	Crude-4	Crude-5	Crude-6	Blend of Crude- 7 and 8	
CPA EoS based approaches								
	Approach-1	Approach-1	Approach-2	Approach-2	Approach-3	Approach-3	Approach-3	
Self-association energy of asphaltene	1ª	1 <sup>a</sup>	1 <sup>a</sup>	1 <sup>a</sup>	0	0	1 (Crude-8)	
Cross-association energy with maltene/(A+R)	1	1	1	1	1	1	2 (Crude-7 and 8)	
Cross-association energy with precipitant	5 (5 precipitants)	4 (4 precipitants)	2 (2 precipitants)	2 (2 precipitants)	0	0	2 (Crude-7 and 8 with Toluene)	
Amount of Asp6	1	1	1	1	0	0	0	
Total	8	7	5	5	1	1	5	
PC-SAFT EoS based approach								
		PC-S	SAFT EoS based a	approach				
	Approach-4	PC-S Approach-4	SAFT EoS based a	approach Approach-4	Approach-4	Approach-4	Approach-4	
Asphaltene component MW	Approach-4 3 (3 components)	PC-5 Approach-4 Not studied	Approach-4 2 (2 components)	Approach Approach-4 (2 components)	Approach-4 0	Approach-4 0	Approach-4 2 (Crude-7 and 8)	
Asphaltene component MW Asphaltene components aromaticity	Approach-4 3 (3 components) 1 <sup>b</sup>	PC-5 Approach-4 Not studied Not studied	AFT EoS based a Approach-4 (2 components) 1 <sup>b</sup>	Approach Approach-4 2 (2 components) 1 <sup>b</sup>	Approach-4 0 1	Approach-4 0 1	Approach-4 2 (Crude-7 and 8) 2 (Crude-7 and 8)	
Asphaltene component MW Asphaltene components aromaticity $k_{ij}$ (asphaltene- saturates)	Approach-4 3 (3 components) 1 <sup>b</sup> 3 (3 components)	PC-5 Approach-4 Not studied Not studied	Approach-4 2 (2 components) 1 <sup>b</sup>	Approach Approach-4 (2 components) 1 <sup>b</sup>	Approach-4 0 1	Approach-4 0 1	Approach-4 2 (Crude-7 and 8) 2 (Crude-7 and 8) 1	
Asphaltene component MW Asphaltene components aromaticity $k_{ij}$ (asphaltene- saturates) $k_{ij}$ (asphaltene- precipitants)	Approach-4 3 (3 components) 1 <sup>b</sup> (3 components) 15 (3 components) (5 precipitants)	PC-5 Approach-4 Not studied Not studied Not studied	Approach-4 2 (2 components) 1 <sup>b</sup> 1	Approach Approach-4 (2 components) 1 <sup>b</sup> 1	Approach-4 0 1 1 0	Approach-4 0 1 1 0	Approach-4 (Crude-7 and 8) (Crude-7 and 8) 1 0	
Asphaltene component MW Asphaltene components aromaticity $k_{ij}$ (asphaltene- saturates) $k_{ij}$ (asphaltene- precipitants) $k_{ij}$ (saturates-(A+R))	Approach-4 3 (3 components) 1 <sup>b</sup> (3 components) 15 (3 components) (5 precipitants) 1	PC-5 Approach-4 Not studied Not studied Not studied Not studied	AFT EoS based a Approach-4 (2 components) 1 <sup>b</sup> 1 0 0	Approach Approach-4 (2 components) 1 <sup>b</sup> 1 0 0	Approach-4 0 1 1 0 0	Approach-4 0 1 1 0 0	Approach-4 (Crude-7 and 8) (Crude-7 and 8) 1 0 0	

<sup>a</sup> Self-association energy of Asp1, Asp2 and Asp3 components are assumed same.

<sup>b</sup> Aromaticity of all asphaltene components of the crude are assumed same.

## Asphaltene Precipitation from Light Crudes

Crude-1 is an Iranian crude whose information including the experimental data are referred from Hirschberg et al [2]. It is characterized using the CPA EoS based approach-1 and the PC-SAFT EoS based approach-4. The characterization of the crude and model parameters for approach-4 are referred from the work of Tavakkoli et al [7]. Fig. 1a shows the experimental asphaltene yield for nC5, nC7, and nC10 precipitants while Fig. 1b shows the experimental concentration of precipitant (nC7, nC10, nC12 and nC16) at the onset of asphaltene precipitation condition. Fig. 1a and 1b show that both approaches can correlate the data but the accuracy of approach-1 is higher than approach-4. It should be noted that approach-4, with all  $k_{ii}$ =0, cannot predict the asphaltene precipitation during the addition of *n*-paraffin. However, as shown in Fig. 1a, when  $k_{ii}$  is used, as shown in Table 13, approach-4 can correlate the asphaltene precipitation data. The higher value of  $k_{ii}$  (e.g.  $k_{ij}$  with nC5 is greater than that with nC7) results in more asphaltene yield. From Fig. 1b, one can analyze that the difference between the approaches-1 and 4 fitted results for the *nC*5 concentration at the onset point is large since the higher value of  $k_{ii}$  lowers the precipitant concentration at onset. From the approach-4 results, it can be concluded that it cannot predict both types of the experimental data (the asphaltene yield and onset condition for different *n*-paraffins) gualitatively. The approach-1 results of onset conditions are not very sensitive to the cross-association energy of *n*-paraffin since the crossassociation between n-paraffin undissolved asphaltene (Asp6) component and n-paraffin is not considered. Therefore, approach-1 can predict both types of experimental data qualitatively. For approach-1, the difference between the values of cross-association energies for nC5asphaltenes and nC7-asphaltenes pairs is higher than the difference between the values of cross-association energies for the nC7-asphaltenes and nC10-asphaltenes pairs. The two widely different values of the model parameters (cross-association energy in approach-1 and  $k_{ij}$  value in approach-4) related to nC5 and nC7 precipitants are due to the widely different values of respective asphaltene yield. From the fitted results using both approaches, as shown in Fig. 1a, it is observed that the asphaltene yield decreases at higher dilution of nC7 and nC10. Hirschberg et al [2] did not mention the error margin on the experimental data. Wang and Buckley [28] concluded from their experimental studies that the asphaltene yield increases with the amount of *n*-paraffin, reaches a maximum value, then decreases at higher amount of *n*paraffin. They observed the maximum in the asphaltene yield at around 30:1 to 40:1 nparaffin: oil volume ratio when nC6 or nC7 used. For nC5, they observed the maximum at around 80:1 *n*-paraffin:oil volume ratio. They mentioned that the overall error on asphaltenes amount measurement was around  $\pm 0.02$  wt%.



Fig. 1. Crude-1: (a) Asphaltene yield during the addition of nC5, nC7 and nC10 paraffins. (b) Amount of different *n*-paraffins (nC5, nC7, nC10, nC12, nC16) required at the onset of asphaltene precipitation. Experimental data are from Hirschberg et al [2]. All experimental data (Fig. 1a and 1b) are used for the parameters estimation.

Crude-2 is a Mexican crude whose information including the experimental data are referred from Buenrostro-Gonzalez et al [15]. The experimental data of the asphaltene yield for nC5, nC7, nC9 and nC12 precipitants are available. For this crude, only the CPA EoS based approach-1 is studied. Fig. 2 shows that approach-1 is able to correlate the data after calculating the model adjustable parameters. The PC-SAFT EoS based approach-4 could also predict this data but we could not fit the model due to the difficulty of estimation of a large number of model adjustable parameters. Buenrostro-Gonzalez et al [15] used the SAFT-VR EoS in McMillan-Mayer framework developed by Wu et al [16-17] to predict this experimental data (with seven adjustable model parameters), however, their model results are not as accurate as the results of this work. The reader is referred to their work for more detail.



Fig. 2. Crude-2: Asphaltene yield during the addition of *nC*5, *nC*7, *nC*9 and *nC*12 paraffins. Lines represent the correlations using approach-1. Symbols represent the experimental data from Buenrostro-Gonzalez [15].

## Asphaltene Precipitation from Heavy Crudes

Crude-3 and 4 are Russian and Venezuelan heavy oils respectively whose information including the experimental data are referred from Sabbagh et al [13]. They mentioned that the asphaltene yield data were repeatable to  $\pm 0.015$  (wt/wt). For approach-4, the oil characterization and model parameters are referred from Tavakkoli et al [7]. For these crudes, the experimental data of asphaltene yield for *nC*5 and *nC*7 precipitants are available. As seen from Fig. 3a and 3b, approach-2 can correlate the data better than approach-4. For approach-4, only one  $k_{ij}$ , of negative value, between saturates and asphaltenes is used. The  $k_{ij}$  value is used to correlate the onset condition.



Fig. 3. (a) Crude-3: Asphaltene yield during the addition of nC5 and nC7 paraffins. (b) Crude-4: Asphaltene yield during the addition of nC5 and nC7 paraffins. Experimental data are from Sabbagh et al [13] for both Crude-3 and 4.

Fig. 4 shows the predictions of asphaltene yield for Crude-3 for nC7, nC9, nC10 and nC11 precipitants using approach-4. One can see that for the nC10 precipitant, the asphaltene yield is zero and the asphaltene yield for nC11 is higher than that for nC9, when precipitant concentration is 40 cm<sup>3</sup>/g of oil. However, it contradicts the general experimental behavior from the literature, where the asphaltene yield always decreases or remains almost constant as the carbon number of paraffinic solvent is increased. Therefore, we hypothesize that approach-4 needs  $k_{ii}$  between asphaltenes and precipitant (like Crude-1) before calculating the PC-SAFT parameters of polydisperse asphaltene fraction. In other words, the asphaltene yield of the crude for different *n*-paraffins precipitants cannot be predicted using approach-4 but can be correlated considering  $k_{ij}$  between asphaltenes and precipitant. It should be noted that approaches-1 to 3 can also not predict the asphaltene yield for different *n*-paraffins and that is why the approach-2 predictions, like Fig. 4 for the approach-4 predictions, are not shown. The same behavior of the approach-4 results, like Fig. 4 for Crude-3, has been observed for Crude-4 but results are not shown here. Therefore, one must check physical behaviors (mole fraction of *n*-paraffin at the precipitation onset, discussed later, and the asphaltene yield should decrease with the carbon number of *n*-paraffin) while calculating the model parameters.



Fig. 4. Crude-3: Predictions of the asphaltene yield during the addition of different *n*-paraffins using approach-4.

## Asphaltene Precipitation Onsets for Heavy Crudes

When the asphaltene precipitation onset conditions are modeled for the reservoir fluid (at high T and P), the asphaltene fraction is usually considered as a monodisperse fraction/component since we are generally interested in calculating the onset conditions. Therefore, it is important to check whether the modeling approaches can predict the onset conditions when the asphaltene fraction is treated as a single component. Crude-5 and 6 are Cold Lake Bitumen and Athabasca heavy oils respectively whose information including the experimental data are referred from Wiehe et al [11]. The experimental data of asphaltene onset conditions for different *n*-paraffins are available for these crudes. Wiehe et al [11] mentioned that the precision of the onset data is within 0.1 vol%. Fig. 5a and 5b show the experimental onset data and model fitted results on molar and volume basis respectively for Crude-5. Fig. 5a shows that the experimental g-moles of *n*-paraffin at the onset condition decreases with the *n*-paraffin carbon number. Fig. 5b shows that the experimental data has a maximum at the *n*-paraffin carbon number of 9. Fig. 5a and 5b show that approach-3 correlates the experimental data qualitatively while approach-4 has major deviations. Approach-4 is studied with respect to scenarios-1 and 2. In scenario-1, the standard *n*-paraffin PC-SAFT parameters from Gross and Sadowski [26] are used while in scenario-2, the n-paraffin PC-SAFT parameters are calculated from the generalized MW correlations, as shown in Table 7. For all other crudes, approach-4 is studied with scenario-1 only. Approach-4 with scenario-1 introduces an inconsistent effect of carbon number that does not seem to exist in the experimental data as shown in Fig. 5. On the other

hand, approach-4 with scenario-2 gives linear trend as shown in Fig. 5. Therefore, the nonlinearity with scenario-1 is due to the inconsistent values of the PC-SAFT parameters for the *n*-paraffin (especially for *nC*7 to *nC*10). The other model parameters like the PC-SAFT parameters of other components and  $k_{ii}$  values are kept the same for scenarios-1 and 2. The approach-4 with scenario-2 model fitted results on volume basis, as shown in Fig. 5b, have considerable deviations due to the poor predictions of the fitted results on mole basis shown in Fig. 5a. It should be noted that approach-4 with scenario-2 still gives accurate prediction of the molar volume. Therefore, it can be concluded that the asphaltene onset precipitation is very sensitive to the PC-SAFT parameters and detailed study about it could give us more insight. For approach-4 with scenarios-1 and 2, using  $k_{ii}$  value for asphaltenes-*n*-paraffin pair could correlate the experimental onset data. The same conclusion, like Crude-5, can be drawn for Crude-6 from Fig. 6. From Table 8, one can observe that Crudes-5 and 6 have almost similar properties and SARA fractions. For approach-4, a negative  $k_{ii}$  value (-0.02) between saturates and asphaltenes is used for both Crudes-5 and 6 to decrease the deviations from the experimental data. Since there are multiple sets of MW and aromaticity values, which give the same modeling results (for approach-4), the experimental MW for the asphaltene component is used to reduce the number of adjustable parameters and the aromaticity value is calculated from the experimental data. The results of mole fraction of *n*-paraffin at the asphaltene onset condition for crudes-1 to 6 are mentioned in the supplementary information.



Fig. 5. Crude-5: (a) Amounts of *n*-paraffin on molar basis at the onset of asphaltene precipitation. (b) Amounts of *n*-paraffin on volume basis at the onset of asphaltene precipitation. Experimental data are from Wiehe et al [11]. Solid lines are from approach-3, dashed lines are from approach-4 with scenario-1, dotted lines are from approach-4 with scenario-2.



Fig. 6. Crude-6: Amounts of *n*-paraffin on volume basis at the onset of asphaltene precipitation. Experimental data are from Wiehe et al [11].

#### Asphaltene Precipitation Onsets from Blending of Crudes

In this section, the modeling approach to calculate the asphaltene precipitation onsets from the blends of Crudes-7 and 8 is discussed. Crudes-7 and 8 are from Gulf of Mexico and Middle East respectively whose information including the experimental data are referred from Tharanivasan et al [12]. They mentioned that the repeatability (in terms of relative standard deviation) of the measured yields was within  $\pm 14\%$  and  $\pm 5\%$  of the reported yield data at low (*n*-heptane mass fraction <0.6) and high (*n*-heptane mass fraction>0.6) dilution ratios respectively of crude oil or blend. The repeatability for the onset measurements was within  $\pm 4\%$  of the reported amount of *n*-heptane at the onset. The experimental data of *nC*7 precipitant mass fraction at the onset of asphaltene precipitation from the blends of Crude-7 or 8 with three different toluene concentrations are available as shown in Fig. 7. Also, the experimental data of *nC*16 mass fraction at the onset of asphaltene precipitation from the blend of Crude-7 with three different concentrations of Crude-8 are available as shown in Fig. 8. The approaches-3 and 4 parameters, as shown in Table 10 and 12 respectively, are calculated from the experimental data of Figs. 7 and 8 and the model fitted results are also shown in the same figures. As shown in Fig. 7, both approaches can correlate the experimental data. Fig. 8 shows

that only approach-3 can correlate the non-linear behavior whereas approach-4 shows almost a linear trend between two extremes, which represent pure Crude-7 result (left) and pure Crude-8 result (right). For approach-3, this non-linearity increases with the difference between the selfassociation energies for the Crude-7 and Crude-8 asphaltenes. In other words, a large difference between the solubility parameters of asphaltene component of Crudes-7 and 8 is required. However, for approach-4, the appropriate parameters could not be found, which can show the non-linearity. The minimum in this non-linear behavior is very important to decide the correct order of crudes to prepare the blend. For example, in this case, the minimum is close to the extreme of pure Crude-8 as shown in Fig. 8. Therefore, one should always prepare the blend by adding Crude-8 into Curde-7 and not the other way. The experimental data for the blend of Crude-7 and Crude-8 with three different toluene concentrations are also available as shown in Fig. 9. This figure shows that the approach-3 predictions have minor deviations while the approach-4 predictions have moderate deviations from the experimental data. One can also analyze that the trend between two extremes becomes more linear as the toluene concentration is increased. It would also be interesting to study the blend of three or more asphaltenic crudes, however, we could not find such experimental data from the literature.



Fig. 7. Crude-7and 8: Amount of *n*-Heptane at the onset of asphaltene precipitation for the blend of different concentrations of crude oil and toluene. Experimental data are from Tharanivasan et al [12].



Fig. 8. Amount of *n*-Hexadecane at the onset of asphaltene precipitation for the blend of different concentrations of Crude-7 and 8. Experimental data are from Tharanivasan et al [12].



Fig. 9. Amount of *n*-Heptane at the onset of asphaltene precipitation for the blends of Crude-7, 8 and toluene at three different concentrations of toluene. Symbols represent the experimental data from Tharanivasan et al [12].

## 5. Conclusions

Based on the results found in this work we conclude that the solubility of asphaltene fraction in solvent at precipitation onset condition can be calculated from experimental data of onset composition for different *n*-paraffins precipitants. The solubility of asphaltene fraction is not dependent upon the number of solvent (maltene) components. The asphaltene fraction can be treated as monodisperse (single component) rather than polydisperse (multiple components) to calculate onset composition of the precipitant. The results for Crudes-5 and 6 show that the CPA EoS based approach-3 (with one adjustable parameter) can accurately correlate the onset compositions of *n*-paraffins while the PC-SAFT EoS based approach-4 without association term (with two adjustable parameters) has moderate deviations. Approach-4 needs binary interaction parameters to correlate the onset composition. With the modeling approaches used in this work, the asphaltene yield during the addition of a precipitant can only be correlated with given experimental data of asphaltene yield with the same precipitant. The approaches consider the asphaltene fraction as multiple components having different solubility parameters to correlate asphaltene yield. For the CPA EoS based approaches-1 and 2, different selfassociation energies are used to define multiple asphaltene components. For the PC-SAFT EoS based approach-4, different pure component parameters are used to define multiple asphaltene components. In addition, one adjustable parameter linked to the specific precipitant is also used. For approaches-1 to 3, it is the cross-association energy between precipitant and asphaltene. For approach-4, it is the  $k_{ij}$  value between precipitant and asphaltenes. It is also shown from the results of Crudes-3 and 4 that when  $k_{ii}$  values (for *n*-paraffins- asphaltene pairs) for approach-4 are not used, the model predictions of asphaltene yield for different nparaffins precipitants are not in agreement with the observed physical behavior of asphaltene precipitation: (a) In the first case it has been observed that the asphaltene yield decreases or remains constant with the carbon number of *n*-paraffin (b) In the second case the mole fraction of *n*-paraffin precipitant at onset condition is known to decrease with the carbon number of *n*paraffin. A detailed study of the PC-SAFT parameters of *n*-paraffins (especially for *n*-heptane to n-decane) is required to establish monotonic trend of the mole fraction of n-paraffin precipitants at the onset conditions versus the carbon number of *n*-paraffin. It is also observed from the results of Crudes-7 and 8 that approach-3 correlates and predicts the non-linear trend of onset condition for the blend (Fig. 8 and 9) while approach-4 cannot. It should be noted that a similar approach based on the CPA EoS has already been studied with respect to the asphaltene precipitation onset conditions from the reservoir fluid in our previous studies [20-22].

## Acknowledgement

The authors wish to thank BP International limited (UK), Statoil (Norway) and Petrobras (Brazil) for supporting the work as part of the CHIGP (Chemical in Gas Processing) projects. The authors are grateful to Dr. Nikolaos Diamantonis and Dr. Leslie Bolton from BP International Limited (UK) for reviewing this article and providing constructive comments.

#### List of Symbols

=	Parameter in the energy term for component i
=	Angstrom (unit of length)
=	Co-volume parameter for component i
=	Parameter in the energy term for component <i>i</i> , dimensionless
=	Radial Distribution Function
=	Binary interaction parameter
=	Boltzmann constant
=	Kelvin
=	Number of segments/spheres per chain/molecule
=	<i>n</i> -Paraffin of carbon number <i>N</i>
=	Pressure
=	Critical Pressure
=	Gas constant

- T = Temperature
- $T_c$  = Critical Temperature
- $V_m$  = Molar volume
- $X_{A_i}$  = Mole fraction of site-A in molecule *i* not bonded to any other site/s
- $x_i$  = Mole fraction of component *i*

## Greek Letter

$\beta^{A_i B_j} = \beta$	Association volume between site A of component $i$ and site B of component $j$
$\Delta^{A_i B_j} = \mu$	Association strength between site A of component $i$ and site B of component $j$
$\varepsilon^{A_i B_j} = \mu$	Association energy between site A of component $i$ and site B of component $j$
$\epsilon_i$ = F	Potential energy between spherical segments of component i
γ = μ	Aromaticity (1 means 0% benzene derivative and 100% PNA)
$\rho$ = M	Molar density
$\omega = \mu$	Acentric factor

 $\sigma$  = Temperature independent diameter of segment (*m*)

## List of Abbreviations

Asp	=	Asphaltene
A+R	=	Aromatics+Resins
Calc	=	Calculated
CERE	=	Center for Energy Resources and Engineering
CPA	=	Cubic Plus Association
Da	=	Dalton
DTU	=	Technical University of Denmark
EoS	=	Equation of State
Exp	=	Experimental
MW	=	Molecular Weight
PC-SAFT	=	Perturbed Chain Statistical Association Fluid Theory
PR	=	Peng-Robinson
SAFT	=	Statistical Associating Fluid Theory
SAFT-VR	=	Statistical Associating Fluid Theory with Variable Range Potential
SARA	=	Saturates Aromatics Resins Asphaltene
SG	=	Specific Gravity
SRK	=	Soave Redlich Kwong

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