NEW TECHNIQUES AND METHODS FOR THE STUDY OF AGGREGATION, ADSORPTION, AND SOLUBILITY OF ASPHALTENES. IMPACT OF THESE PROPERTIES ON COLLOIDAL STRUCTURE AND FLOCCULATION

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NEW TECHNIQUES AND METHODS FOR THE STUDY OF AGGREGATION, ADSORPTION, AND SOLUBILITY OF ASPHALTENES. IMPACT OF THESE PROPERTIES ON COLLOIDAL STRUCTURE AND FLOCCULATION

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ABSTRACT

The solubility of Furrial asphaltene in toluene was 57 g L\(^{-1}\). However, using a new technique, based on the precipitation of this sample by the phenol PNP, we found that a fraction (2), comprising 47% of the asphaltene, is of low solubility. This suggested that this material constitutes the colloidal phase, and the rest acts as the dispersing fraction. This technique allowed the fractionation of asphaltenes in fractions A\(_1\), A\(_2\), and A\(_3\) according to solubility, going from practically insoluble (A\(_1\)) to low (A\(_2\), 1 g L\(^{-1}\)) to high (A\(_3\), around 57 g L\(^{-1}\)). The adsorption isotherms of asphaltenes on glass and silica in toluene consist of a sequence of steps or step-wise adsorption. The first layer or first step is formed by the adsorption

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of free asphaltene molecules and by small aggregates (aggregation number between 3 and 6) which saturate the glass or silica surface in the usual manner (L-type or H-type isotherms). However, we suggest that the second, third, and other asphaltene layers adsorb sequentially according to the above differences in solubility. The very slow changes with time and the negligible desorption from the surface measured for the above isotherms were interpreted as the effect of packing or the building up of a well packed layer. This would be achieved by the slow formation and rupture of bonds between neighboring molecules at the surface. Thus, molecules with difficulties to pack, adsorbed by a kinetically controlled process, are either rejected or relocated in a thermodynamic controlled process. The above results and ideas were used to improve the models for asphaltene and petroleum colloids and to underscore the importance of surfaces and colloid dispersants in asphaltene precipitation during the production of crude oils. For instance, the results described below suggest that colloids are constituted by a well packed and insoluble asphaltene core, impervious to the solvent, and by a loose packed periphery which, by allowing solvent penetration, keep the colloid in solution. According to this model, desorption of compounds in the above loosely packed periphery, such as the one promoted by a surface, would be the main cause of asphaltene precipitation from crude oils. In this case, solubility reductions caused by pressure drops during oil production would have a minor effect. Also, preliminary number average molecular weights $M_n$ for four asphaltenes, obtained using a new procedure, are presented here. The $M_n$ values obtained ranged from 780 to 1150 g/mol.

Key Words: Asphaltene-colloids; Asphaltene-aggregates; Asphaltene-adsorption-solubility

GLOSSARY OF SYMBOLS

**English Symbols**

$A$ Asphaltene

$A_1$ Asphaltene fraction insoluble in toluene isolated from $A_2$

$A_2$ Asphaltene fraction of low solubility in toluene, isolated by L-Type: precipitation with PNP, after PNP extraction

$A_3$ Asphaltene fraction soluble in toluene, isolated from Furrial asphaltenes after PNP extraction
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ALT-17  Crude sample from Maracaibo, Venezuela
Ao, Ai  Absorbance values
Am, Ai, and Aa  Asphaltene sample in the media, interface, and surface, respectively
C  Solution concentration
CMC  Critical micelle concentration
C-148  Crude sample from Maracaibo, Venezuela
D  Thermal diffusivity constant
DM-148  Crude sample from Maracaibo, Venezuela
DM-22  Crude sample from Maracaibo, Venezuela
DM-153  Crude sample from Maracaibo, Venezuela
H-Type  High energy type isotherm
k  First-order rate adsorption of asphaltene
k1, k−1  Diffusion rates
k2  Adsorption rate
K  Equilibrium constant
M  Average molecular weight of a free or non-aggregated asphaltene
M  Molecular weight
Ma  Average molecular weight
Mn  Number average molecular weight
MA  Methylated asphaltenes
N  Number of Avogadro
n  number of n-octyl groups added to the asphaltene molecule
NPE  Nonylphenol ethoxylated
OA  Octylated asphaltene
PNP  para-nitrophenol
PNT  para-nitrotoluene
p  Amount of asphaltene precipitated in the PNP experiments
PSD  Photothermal surface deformation method or technique
S  Surface area, surface or site on surface
THF  Solvent tetrahydrofurane
VLA-711  Crude sample from Maracaibo, Venezuela
VPO  Vapor pressure osmometry method for molecular weight determination

Greek Symbols

δ  Width of an asphaltene molecule
ν  Amount of material adsorbed on surfaces
ρ  Density
σ  Molecular area
1. INTRODUCTION

The idea regarding the presence of asphaltenes in crude oil as colloids dispersed by resins and other compounds has been around since the days of Pfeiffer and Saal (1). So far many experimental evidence has been gathered in support of this theory (2). Measurements of the size and shape of the colloids have been reported, and some theoretical studies regarding phase behavior have used the colloidal approach (3–5). Undoubtedly, this interest is due to the negative impact of asphaltene precipitation during production and other industrial operations, and hence research in this area is very active.

Despite all the evidence collected so far, many points remain obscure. This is expected in view of the very high complexity of asphaltenes discussed many times in the literature. A point, regarding phase behavior, is whether a molecular or colloidal approach should be used to represent solutions of asphaltenes. The literature on the subject is not clear in this regard, and models using molecular (6,7) and colloidal (see above) theories have been reported recently. The results shown below support the colloidal model very strongly, and we suggest that the molecular model should be abandoned in view of the absence of any experimental evidence supporting it.

Another point is related to the so-called asphaltene micelles forming at a particular critical micelle concentration (CMC). By definition micelles are surfactant aggregates forming sharply at the CMC in water with a sharp aggregation number. This very particular behavior is due to the hydrophobic effect, whereby the surfactant forms a closed volume with the lowest energy, to enclose the maximum number of hydrophobic tails. The undesired contacts between water and the hydrocarbon tails are thus kept to a minimum. Thus, a necessary condition for a CMC is the presence of water, and even in the case of ordinary surfactants in oily media, such CMC have been strongly questioned (8). A step-wise aggregation process for micelle formation in the oily case is now widely accepted (9).

For asphaltenes the above CMC process is more unlikely due to polydispersity. This would inhibit any packing procedure at any particular concentration, and the difficulties to pack in the aggregate could only be overcome by increasing the trend of the solute to be adsorbed; that is, by increasing the concentration in solution. In other words, the aggregate would grow in a step-wise form.

Other important aspects regarding structure, such as the nature of the material in the core and periphery of the colloid are also obscure, and are matters of much speculation.

Clearly, new techniques and methods should be developed in order to shed light on the above and similar matters. This paper condenses the description of new techniques and methods developed in our laboratories to study the solubility, aggregation, and adsorption properties of asphaltenes and the impact of these properties on colloid solubility and asphaltene flocculation. Part of the research with some of
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these techniques (PSD and thermal lens) has been published in recent times (10,11). The treatment of asphaltene solutions with PNP, and the treatment of crude oils with silica, devised to obtain information on phase solubility and colloidal structure, are described here for the first time. Also, preliminary published results on kinetic and asphaltene isotherm (12) are discussed in terms of new ideas regarding asphaltene tendency to adsorb on asphaltene surfaces, and packing ability.

We hope that the results and discussion given below will clarify some of the obscure points mentioned above.

2. EXPERIMENTAL

2.1. Materials and Area Determination

Some properties of the Venezuelan crude oils and their asphaltenes used in this work are shown in Table 1. Samples ALT-17, C-148, DM-22DM-153 and VLA-711 from Maracaibo Lake (in the western part of the country) are stable crude oils without precipitation problems, whereas Furrial, from Monagas state (in the eastern part of the country), has severe flocculation problems. The compounds para-nitrophenol (PNP) and p-nitrotoluene (PNT) were commercial samples used as received.

Asphaltenes, thoroughly Soxhlet extracted with n-heptane to remove resins, were obtained from Furrial crude oil by the addition of 40 volumes of n-heptane as described elsewhere (13). Commercial samples of silica gel were used as adsorbent. Using the dye method (14) and PNP as the dye, an area of 13.7 m²/g was found for silica in toluene when the molecular area $\sigma = 52.5 \text{Å}^2$, reported for the PNP (14) was used (see Eq. 1 below).

Table 1. Properties of Crude Oils and Asphaltenes

<table>
<thead>
<tr>
<th>Sample</th>
<th>Asphaltene</th>
<th>API</th>
<th>C</th>
<th>H</th>
<th>N</th>
<th>S</th>
<th>H/C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Furrial</td>
<td>10</td>
<td>21</td>
<td>85.5</td>
<td>6.9</td>
<td>1.6</td>
<td>3.4</td>
<td>0.97</td>
</tr>
<tr>
<td>ALT-17</td>
<td>0.25</td>
<td>30</td>
<td>86.8</td>
<td>7.1</td>
<td>1.2</td>
<td>2.7</td>
<td>0.98</td>
</tr>
<tr>
<td>C-148</td>
<td>0.23</td>
<td>35</td>
<td>83.9</td>
<td>7.1</td>
<td>0.90</td>
<td>–</td>
<td>1.00</td>
</tr>
<tr>
<td>DM-22</td>
<td>3.2</td>
<td>24</td>
<td>84</td>
<td>7.5</td>
<td>0.96</td>
<td>3.3</td>
<td>1.10</td>
</tr>
<tr>
<td>DM-153</td>
<td>10</td>
<td>14</td>
<td>81.1</td>
<td>8.2</td>
<td>1.64</td>
<td>6.6</td>
<td>1.2</td>
</tr>
<tr>
<td>VLA-711</td>
<td>2.2</td>
<td>31</td>
<td>83.9</td>
<td>7.5</td>
<td>1.2</td>
<td>4.6</td>
<td>1.06</td>
</tr>
<tr>
<td>A1</td>
<td>–</td>
<td>–</td>
<td>83.3</td>
<td>6.6</td>
<td>0.96</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*a*API gravities of crude oils. *b*Corresponding to asphaltenes. *c*Sample of low solubility in toluene isolated by treatment with PNP.
By a similar procedure a surface area of 91 m$^2$/g was found when Furrial asphaltene was employed as the adsorbing surface in n-heptane. Since the solubility of PNP in this solvent is very low, we used PNT as the dye and the $\sigma$ value for PNP. Surface areas $S$, in m$^2$/g, were calculated using Equation 1 (14).

$$S = \nu_m \sigma \times N_A 10^{-20}$$

In this equation $\nu_m$ is the amount of dye adsorbed in mg g$^{-1}$, $\sigma$ is the dye area in Å$^2$/molecule, and $N_A$ is the number of Avogadro.

## 2.2. Methods

### 2.2.1. Solubility and Precipitation Methods

Toluene solutions of PNP and asphaltene were prepared in separated flasks and then mixed in 100 ml volumetric flasks to make final concentrations of PNP in the range of 500 to 8000 mg L$^{-1}$ and a final constant concentration of 8 g L$^{-1}$ in asphaltene. The solutions were allowed to stand at room temperature for one week, and after this time the contents were filtrated, the filtrated was set aside and the precipitate solid was washed with toluene, dried, weighted, and dissolved in chloroform. This solution was washed once with aqueous sodium hydroxide (5%) to extract the PNP into the aqueous solution, which was then determined as phenolate by UV. The precipitated solids obtained before and after the extraction of PNP are hereafter called $p$ and $A_2$, respectively. The separation scheme used is shown in Figure 1.

The precipitated asphaltene $p$ is reported here in percentage of total asphaltene in solution. The amount of adsorbed PNP ($\nu$), coprecipitated with the asphaltenes, is in mg g$^{-1}$. The soluble fraction $A_3$ was isolated after evaporation of the above filtrate, dissolution of the residue in chloroform, and extraction of PNP, as described above. A solution of $A_2$ in toluene was prepared after mixing with $A_3$ in proportions (1:3 and 2:3). Sample $A_1$ was the residue in the thimble, obtained after a thorough Soxhlet extraction of $A_2$ with toluene.

The solubility of samples at room temperature was determined as usual, by preparing saturated solutions in toluene using the following procedure. A weighted amount of sample was added in small portions to toluene and contained in the test tube placed in an ultrasonic bath. Using a capillary a drop of solution was withdrawn and placed on filter paper. Undersaturated solutions give a continuous stain on the paper, whereas saturated solutions give a stain with solid at the application point. The amount of sample added just before solid appearance was equal to the solubility. These solubility measurements afforded the following results in g L$^{-1}$: $A_1$, 0.00; $A_2$, 1.14; and the solubility of $A_3$ was similar to the one for Furrial asphaltenes: 57 g L$^{-1}$.
Flocculation experiments were carried out using 25 ml of crude oil in the presence of 5 g of silica gel. When heating was required, the mixture was placed in a flask attached to a condenser and heated for 12 h, then allowed to cool and stand overnight. For the experiments below room temperature, the mixture was shacked for one day. After this time (in all cases) the mixture was allowed to reach room temperature, the solid (silica plus precipitate) was filtered, washed with n-heptane until a clear filtrate was obtained, dried, and weighted. The amount of precipitate asphaltene was obtained by weight difference. When required, the asphaltenes were extracted from the silica in a Soxhlet with a boiling mixture of chloroform–methanol 3:1.

**Figure 1.** Flow diagram showing the separation of $A_1$, $A_2$, and $A_3$ after the treatment of toluene solutions of Furrial asphaltenes with paraf-nitrophenol (PNP).
2.2.2. Adsorption Methods

Asphaltene-toluene solutions were prepared by toluene dilution of a stock solution (500 mg L\(^{-1}\)). The stock solution was prepared by weighting the appropriate amount of dried asphaltene, which was dissolved in toluene using ultrasonic bath. The solution was allowed to stand for two days before it was used.

For kinetic runs, small silica plates (2.5 × 1 cm\(^2\), 24 mg of silica) were introduced in small glass vials containing 8 mL of asphaltene-toluene solution. Changes in absorbance with time were continuously monitored by passing a He-Ne laser beam (632.8 nm) through the vials, and the intensity was registered by using a photodiode. In one experiment (asphaltene concentration equal to 200 mg L\(^{-1}\)) the solution was stirred by using a small magnetic stirrer. Since no difference in rate was observed with and without stirring, all experiments were carried out without stirring.

From these kinetic plots, the amount of adsorbed solute at any time in each case \(\nu_t\) (mg g\(^{-1}\)), as well as the corresponding solution concentration \(C_t\) (mg L\(^{-1}\)), could be obtained using Equations 2 and 3 below:

\[
\nu_t = \left(\frac{(A_0 - A_t) a}{A_0 w}\right) C_0 \tag{2}
\]

\[
C_t = \left(\frac{A_t}{A_0}\right) C_0 \tag{3}
\]

where \(A_0\) and \(A_t\) are the absorbances at zero and \(t\) minutes, respectively, \(C_0\) (mg L\(^{-1}\)) is the initial concentration, \(a\) is the aliquot, and \(w\) is the solid weight (g).

A desorption experiment was performed as follows. Asphaltenes were adsorbed on a silica plate from a 200 mg L\(^{-1}\) solution as described above. After 72 h, the quantity of sample adsorbed was equivalent to 133 mg L\(^{-1}\). The plate was withdrawn, dried, and embedded in pure toluene. The absorbance increase was monitored as usual, and after 50 h no further change was observed. The amount desorbed was equivalent to 4 mg L\(^{-1}\).

Isotherms were measured using the photothermal surface deformation (PSD) method described earlier (10). Briefly, in this method the amount of solute adsorbed on the surface is determined directly. That is, by using a procedure similar to the one described for the kinetic runs (see above), the sample was adsorbed on the silica plate. After the required time, the plate was withdrawn, allowed to dry in a desiccator, and set for analysis. This was performed by using two lasers, one for pumping and the other as the probe. The pumping beam heats the sample and produces a dilatation that is proportional to the sample mass. This is measured by the probe beam.

Signal calibration was performed by comparing with the adsorption measured by transmission measured as described in the kinetic runs.
2.2.3. Area Per Asphaltene Molecule $\sigma_A$

$\sigma_A$ was estimated from Equation 4:

$$\sigma_A = 0.166(MS_S/\nu) \quad (4)$$

Here $S_S$ is the area of the silica in m$^2$/g, $\nu$ is the amount of asphaltene in the first layer in mg g$^{-1}$, $\sigma_A$ is in $\text{Å}^2$ per molecule and $M$ is the molecular weight. When required, an $M$ value near 1000 was used (see below and Tab. 3).

2.2.4. Thermal Lens Method

The thermal lens method has been reported elsewhere (11), so only a brief description will be given here. In the thermal lens, the asphaltene in solution absorbs energy from the excitation beam as usual. Through fast non-radiative relaxation of excited species, part of this energy is then transferred as thermal energy (heat) to the solvent, and a density change within the excitation region is induced. Another laser is used as a probe to monitor the refractive index gradient (thermal lens) so induced, and after the appropriate measurements and calculations the thermal diffusivity $D$ is obtained.

2.2.5. Molecular Weight Methods

The general procedure of asphaltene alkylation with potassium naphthalide and octyl iodine in THF have been described earlier (13,15). Briefly the method can be described as follows. A solution of naphthalene in THF is treated with potassium to form potassium naphthalide. This compound is an ion radical with a high reduction potential. When asphaltenes are added to the solution, they react with the ion radical to form ionic sulfides and carbanions. After this reaction is complete, the alkyl iodine (octyl or methyl iodine) is added to form the alkyl asphaltene.

Molecular weights of samples were measured in nitrobenzene at 100°C using the VPO method described elsewhere (16). The usual range of solution concentrations was employed (more than 1 and less than 10 g L$^{-1}$) and the number average molecular weights $M_n$ were obtained by extrapolation to infinite dilution.

The number $n$ of octyl groups incorporated in the asphaltene per 100 carbons was calculated from the elemental analysis using the equation below.

$$\frac{(H/C)_{OA}}{(H/C)_{A}} = \frac{100(H/C)_{A} + 17n}{100 + 8n} \quad (4-1)$$

In this equation, $(H/C)_{OA}$ and $(H/C)_{A}$ are the hydrogen to carbon ratio corresponding to octylated asphaltenes (OA) and asphaltenes, respectively.
2.2.6. Rate Equations

An adsorption process in two steps, one for diffusion to the surface and the other for adsorption, can be described as follows:

\[ A_m = A_1, \]
\[ A_1 + S \rightarrow A_a. \]

In these equations, \( A_m, A_1, \) and \( A_a \) represent samples in the solvent media, in the interface, and adsorbed on the surface. \( S \) stands for a site on the surface. The first step refers to diffusion from the solvent to the surface, whereas the second refers to adsorption. Let \( k_1 \) and \( k_{-1} \) be the rate constants for the first process and the reverse, and let \( k_2 \) be the adsorption rate constant.

Then, assuming the adsorption to be irreversible (see Fig. 5 and the Desorption section below), the rate would be:

\[ \text{rate} = k_2[A_1][S]. \] (7)

Using the stationary state approximation, for \( A_1 \) Equation (8) is obtained:

\[ k_1[A_m] = k_{-1}[A_1] + k_2[A_1][S] = [A_1](k_{-1} + k_2[S]). \] (8)

Then:

\[ [A_1] = k_1[A_m]/(k_{-1} + k_2[S]). \] (9)

Substitution in 7 leads to Equation 10:

\[ \text{Rate} = k_2k_1[A_m][S]/(k_{-1} + k_2[S]). \] (10)

Equation 10 shows that if the second step is too fast in comparison with the first \((k_2[S] \gg k_{-1})\), the rate equation is:

\[ \text{Rate} = k_1[A_m]. \] (11)

Therefore, a first-order reaction should indicate that diffusion is the slow step. If, on the other hand, the second step is very slow so that \( k_2[S] \ll k_{-1} \), the rate would be:

\[ \text{rate} = k_2k_1[A_m][S]/k_{-1}, \] (12)

or

\[ \text{rate} = k_2K[A_m][S], \] (13)

where \( K \) is the equilibrium constant of the first step.
In this case, the adsorption step would be the slowest path, and a second-order kinetics is obtained.

3. RESULTS

3.1. Precipitation with Silica and PNP

In Figure 2 the precipitation results are shown. In this figure both the amount of precipitate \((p)\) and the amount of PNP adsorbed on the precipitate \((\nu)\) are plotted as a function of the concentration of PNP in solution.

It should be emphasized that solid \(p\) is obtained before extraction of PNP. Solid \(A_2\) was obtained from \(p\), after the extraction of PNP (see section 2.2). Since the PNP adsorbed on \(p\) is small, the amount of \(p\) and \(A_2\) are considered here to be equal.

In Figure 3, these results were plotted in the way of an inverse Langmuir or L-plot, and the following lines were obtained in each case:

\[
\frac{1}{p} = \frac{1.175}{C_{\text{PNP}}} + 0.00012, \quad R^2 = 0.98, \quad (14)
\]

\[
\frac{1}{\nu} = \frac{12.78}{C_{\text{PNP}}} + 0.012, \quad R^2 = 0.97 \quad (15)
\]

Using Equation 14, a maximum of 8.3 g L\(^{-1}\) of precipitate could be expected (inverse of intercept in Eq. 14). Since the asphaltene concentration is 8 g L\(^{-1}\), this is an encouraging result. In any case, the percentages of precipitate are high, suggesting a high percentage of colloidal phase in solution. This is consistent with the 47\% of precipitate material collected for the asphaltene solution in toluene (8 g L\(^{-1}\)) saturated with PNP.

![Figure 2](image_url)

**Figure 2.** Yields of asphaltenes precipitated \((p\), percentage of sample in solution) and co-precipitated PNP \((\nu\), mg per gram of precipitated asphaltene) as function of concentration of PNP in solution \((C_{\text{PNP}})\). Room temperature. \(p\) is in mg/100 ml and \(\nu\) in mg g\(^{-1}\).
Figure 3. Inverse Langmuir type plot corresponding to the results in Figure 2 (see text).

Also for a monolayer coverage (83 mg g\(^{-1}\), inverse of intercept in Eq. 15), an area about 190 m\(^2\)/g for the colloid in solution could be calculated using Equation 1. The area determined for the solid asphaltene in n-heptane using PNT was about 97 m\(^2\)/g (see section 2.1). However, as described in the literature (17), this corresponds to the amount required to saturate the external area of asphaltenes (about 41 mg g\(^{-1}\) for PNT in our case), the actual area being higher. Therefore, the above colloid area appears to be a reasonable value.

It should be mentioned that when this technique was applied to other asphaltenes (see section 2 and Tab. 1 for sample details), no significant formation of precipitate was observed in toluene. However, preliminary tests with the solvent cumene (isopropyl benzene) have afforded large quantities of precipitate in all cases. Thus, this precipitation behavior appears to be a general property of asphaltenes.

Figure 4 shows the precipitation results using crude oils and silica. As shown, the amount of asphaltene precipitated increases with temperature, approaching...
asymptotically the total amount of asphaltene present in the crude. It is interesting to note that in the C-148 case, the usual precipitation by n-heptane addition (see Tab. 1) affords a much lower asphaltene yield than the ones shown in Figure 4. It should be mentioned that no precipitation at all was observed for these crudes in the absence of silica.

### 3.2. Adsorption Results

In Table 2 the values for the apparent first-order kinetic constants \( k \), obtained for the studied solutions and surfaces, are shown. These were obtained by fitting the results to a first-order kinetics. The fittings were quite good for the 5–50 mg L\(^{-1}\) runs. However, for the 200 and 400 mg L\(^{-1}\) runs, the results suggested that the readings at long times should correspond to a slower rate (see Fig. 5).

Figure 6 shows the kinetic results for the desorption experiment. After 3000 min only 0.26% of the material initially adsorbed was desorbed in this experiment (see Experimental).

### 3.3. Adsorption Isotherms

The results of these experiments are shown in Figures 7a–7c. Note that for 18 and 48 h, the isotherms are L-type, whereas the one measured at 96 h shows a tendency toward H-type. This H-type form was finally obtained after 8 days (see Fig. 8). The very slow changes with time of these curves are an important property of these isotherms (see the discussion below).

---

**Table 2.** Apparent First-Order Rate Constant \( k \) for the Adsorption of Toluene Solution of Furrial Asphaltenes

<table>
<thead>
<tr>
<th>Initial Concentration (mg L(^{-1}))</th>
<th>Silica ( k \times 10^3 ) (min(^{-1}))( ^a )</th>
<th>Silica Covered(^b ) ( k \times 10^4 ) (min(^{-1}))( ^a )</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>1.09 ± 0.8</td>
<td>–</td>
</tr>
<tr>
<td>20</td>
<td>1.24 ± 0.3</td>
<td>–</td>
</tr>
<tr>
<td>50</td>
<td>1.12 ± 0.3</td>
<td>4.76</td>
</tr>
<tr>
<td>200</td>
<td>1.62 ± 0.3</td>
<td>5.70</td>
</tr>
<tr>
<td>400</td>
<td>0.79 ± 0.3</td>
<td>4.80</td>
</tr>
</tbody>
</table>

\(^{a}\)Covered with asphaltenes adsorbed after one week from a toluene solution (200 mg L\(^{-1}\) initial conc.). \(^{b}\)At room temperature.
Figure 5. Adsorption kinetics for a toluene solution of Furrial alphanetenes (200 mg L$^{-1}$) obtained on silica at room temperature. Points are experimental and the curve is the fitting to a first order. (Reprinted from (18), with permission.)

Figure 6. Desorption kinetics of Furrial asphaltenes from silica plate at room temperature in toluene. (Reprinted from (18), with permission.)
Figure 7. Adsorption isotherm of toluene solutions of Furrial asphaltenes on silica measured by PSD at room temperature. a) after 18 hours; b) after 48 hours; c) after 96 hours.
3.4. Molecular Weight Values

The $M_n$ values measured in this work are shown in Table 3. The last column was obtained after subtracting the value 113n from the fourth column (113 is the molecular weight of an octyl group). This procedure should give the expected $M_n$ for asphaltenes (fifth column in Tab. 3). That is, the molecular weight corresponding to the free or non-aggregated sample (see below).

The following aspects are noteworthy. (1) There is no correlation between the MA and OA molecular weights and sulfur content (see Tab. 1 for sulfur content). (2) The molecular weight reductions observed after methylation by the present method are either similar (Furrial case) or lower than others observed after methylation with diazomethane or other methylating reagents, where rupture of covalent bonds is not possible (15,16). Hence, these values are not consistent with possible de-polymerization reactions, where the rupture of covalent bonds involving sulfur or any other atoms would lead to low molecular weights.

![Figure 8. Adsorption isotherm for the system: Furrial asphaltene/silica/toluene. Obtained after 8 days at room temperature. Each point is an average of at least three replicates.](image)

Table 3. Number Average Molecular Weight ($M_n$) for Asphaltene and Derivative Samples in Nitrobenzene at 100 °C

<table>
<thead>
<tr>
<th>Samples</th>
<th>A $^a$</th>
<th>MA $^b$</th>
<th>n $^c$</th>
<th>OA $^d$</th>
<th>A (calc) $^e$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Furrial</td>
<td>6050</td>
<td>1100</td>
<td>3.1</td>
<td>1150</td>
<td>920</td>
</tr>
<tr>
<td>DM-153</td>
<td>1300</td>
<td>1050</td>
<td>5.4</td>
<td>1750</td>
<td>832</td>
</tr>
<tr>
<td>VLA-711</td>
<td>1750</td>
<td>1100</td>
<td>5.7</td>
<td>1670</td>
<td>1150</td>
</tr>
<tr>
<td>ALT-17</td>
<td>1420</td>
<td>—</td>
<td>5.9</td>
<td>1150</td>
<td>780</td>
</tr>
</tbody>
</table>

$^a$ asphaltene; $^b$methylated asphaltene; $^c$see Equation 4-1; $^d$octylated asphaltene; $^e$expected $M_n$ value (see text).
4. DISCUSSION

4.1. Molecular Weights

The presence of the long octyl chains and the use of a polar solvent (nitrobenzene) and high temperatures warrant the absence of intermolecular interactions for the OA case. Due to steric hindrance, these long chains would inhibit the formation of aggregates. It is noteworthy that even under the present conditions asphaltene aggregation is apparent, particularly in the case of Furrial (compare first and fifth column in Tab. 3). As described above, Furrial crude oil has asphaltene precipitation problems. Thus, the large tendency for aggregation suggested by the high $M_n$ for this sample is somewhat anticipated.

The results for the methylated derivative ensure that any possible bond breaking leading to reduction of molecular weight could be discarded. As shown previously, the potassium naphtalide could easily break sulfide bonds (13,15). According to the $M_n$ values for the methylated asphaltenes, the sulfides in asphaltenes should be in thiophene rings, in which case the rupture of the corresponding $\text{C-S}$ bonds does not reduce the molecular weight.

The most important conclusion regarding the results in Table 3 is that asphaltene molecular weights should be around 1000 or lower (see column 5 in Tab. 3). These results are in agreement with others reported using mass spectra (19–21) and other techniques (22). Thus, it appears that the long-standing question regarding asphaltene molecular weight has finally found the right answer: asphaltene molecular weights are near to or below 1000 g/mol on average.

4.2. Thermal Lens

As reported earlier, a plot of the thermal diffusivity $D$ versus the logarithm of asphaltene concentration (log C) for toluene solutions afforded a curve with a minimum around $C = 50 \text{ mg L}^{-1}$. A similar plot in THF showed no effect of the concentration on $D$ (11). These results suggested a sequential aggregation of asphaltenes and a threshold for aggregation near the above concentration.

4.3. Precipitation Studies

The L-type isotherms of Figure 2, the corresponding linear plots of Figure 3, and the low solubility in toluene of the recovered precipitate (A2; see section 2.2) are clear evidence for precipitation occurring after the adsorption of PNP on the colloid surface.

Precipitation on the basis of the formation of molecular complexes could be discarded because in such a case, the amount of complex formed $p$ (see Fig. 2),
being the result of a chemical reaction, would be related to $C_{PNP}$ by a straight line. These results strongly suggest that on adsorption, the PNP displaces the dispersing compounds that stabilize the colloid, leading to precipitation and to the low solubility of $A_2$. Thus, as expected, the colloidal fraction of asphaltene ($A_2$) has a low solubility in toluene. When the $A_2$ fraction was added to a toluene solution of $A_3$ (the soluble fraction; see section 2.2) the high solubility of the total asphaltene sample was recovered.

Since the insoluble fraction $A_2$ represents a substantial amount of the sample (47%), the capacity of the soluble fraction $A_3$ for dispersing the colloid should be very high. These aspects are further discussed below, where a more complete description of the colloid is given (see the desorption section below).

A similar argument could be used to account for the flocculation results (see Fig. 4). In the presence of an active surface $S_A$ and the colloid surface $S_C$, the compounds dispersing the colloid ($A_3$; see above) will partition between them, as shown in the Appendix. A higher affinity of $A_3$ for $S_A$ will promote desorption from the colloid surface, reducing the stability of the colloid and promoting flocculation. In the case of crude oils, where the solvency of the media is lower than toluene, precipitation was observed even at $0^\circ$C, and complete precipitation was obtained at high temperatures (see Fig. 4). As expected, the precipitation increases with temperature (see Appendix).

The flocculation results in Figure 4 were performed with quite different crude oil, with small and large asphaltene content, and with and without flocculation problems (see section 2 and Tab. 1). Therefore, the presence of colloids in crude oils appears to be a general characteristic. As suggested by these results, contacting with an active surface will lead to flocculation, regardless of crude oil nature. An important consequence in this case is that there is no need for any composition or pressure change to flocculate asphaltene colloids.

As far as we know, the above results are the first ones in which desorption of dispersing compounds, leading to colloid flocculation, is possible without changing the media conditions. This is particularly the case for the silica experiments. These results provide very convincing evidence for the colloidal nature of asphaltenes, and suggest that any consideration of asphaltenes as molecular dispersion is erroneous and should be abandoned.

### 4.4. Adsorption Studies

Previously the adsorption isotherms on glass plates for Furrial and other asphaltenes without precipitation problems were reported (10). These are shown in Figures 9–11 for Furrial, Jobo, and Hamaca, respectively. These measurements were performed using the PSD method described above.

Several interesting features of these isotherms are discussed below.
Figure 9. Adsorption isotherm for Furrial asphaltenes measured at different times using PSD. Surface, glass plate; solvent, toluene, 25°C. (a) 48 h; (b) 120 h. (Reprinted with permission from (10).)

Figure 10. Adsorption isotherm for Jobo asphaltenes measured at different times using PSD. Surface, glass plate; solvent, toluene, 25°C. (a) 6 h; (b) 31 h; (c) 86 h. (Reprinted with permission from (10).)
Figure 11. Adsorption isotherm for Hamaca asphaltenes (sample PA) measured at different times using PSD. Surface, glass plate; solvent, toluene, 25°C. (a) 6 h; (b) 31 h; (c) 86 h. (Reprinted with permission from (10).)

The first one is related to the first step of these isotherms. As shown in these figures, saturation of the glass surface (first step) occurs for ν values near $2 \times 10^{-4}$ mg/cm$^2$ for the Furrial case. Higher values are apparent for the Hamaca and Jobo samples. The apparent area per molecule of asphaltenes ($\sigma$), in Å$^2$/molec, could be obtained from Equation 16:

$$\sigma = \frac{M_a}{\nu N_A} \times 10^{19},$$

(16)

where $M_a$ is the average molecular weight of the species adsorbed. When the average molecular weight $M_f$, corresponding to the free or non-associated asphaltene ($M_f \approx 1000$; see Tab. 3) and the above $\nu$ values (in mg/cm$^2$) are substituted in Equation 16, the value of 83 Å$^2$/molec is obtained for $\sigma$. Of course, this value is much too small, suggesting that the molecular weight used in Equation 16 is too small. To have an idea of what molecular area to expect for a given molecular weight value, we could use Equation 17:

$$M = \rho \sigma \delta N_A 10^{-24}.$$  

(17)

In this equation, $\rho$ is the density in g/cm$^3$ and $\delta$ is the width of the molecule in Å. Assuming $\rho = 1$, a disk shape for asphaltenes, $M = M_f = 1000$ and $\delta = 3.5$ Å (23), a more reasonable value, around 470 Å$^2$, is obtained for $\sigma$. To obtain this area using Equation 16, $M_a$ should be around 6000 or $M_a = 6 M_f$. 
A similar treatment could be performed with the silica isotherms (see Fig. 7c). After 96 hours, about 24 mg g\(^{-1}\) of asphaltene were adsorbed on the surface (first step). The corresponding \(\sigma\) could be obtained from Equation 4 using the \(S_s\) value determined for silica (13.7 m\(^2\)/g; see Experimental) and the above \(M_f\). In this way, a too-small value for \(\sigma\), around 100 Å\(^2\), is obtained. Again a higher molecular weight should be used to obtain a more reasonable molecular area (\(M_a \approx 3 M_f\)).

These results suggest very strongly that, in the diluted concentration range considered (below 1000 mg L\(^{-1}\), see Figs. 7, 9–11), free asphaltenes, as well as small aggregates, are adsorbed on the glass or silica surfaces. Therefore, as suggested by the calculations in the above paragraphs, the average molecular weight of the sample adsorbed (\(M_a\)) should be higher than the average molecular weight of the free asphaltene (\(M_f\)) (between 3 and 6 times \(M_f\)). Hence, the first adsorption step on glass or silica is the consequence of the adsorption of free molecules and small aggregates of asphaltenes.

As usual, saturation of the surface would be the consequence of consumption of all vacant sites. Probably the surface is active enough so that no discrimination between free or aggregated asphaltenes is made.

It should be mentioned at this point that the solutions were equilibrated by allowing them to stand by for two days before contact with the surfaces (see section 2.2).

It is important to realize that the above results are not consistent with the adsorption of large aggregates in the first step. That would require a larger \(M_a\), possibly 10 \(M_f\) or more. However, such adsorption could be expected for the second or third step (see the above isotherms).

According to the discussion above, these isotherms suggest that aggregation of asphaltenes should begin at very low concentration. This is particularly clear from the silica isotherms, where the first step is completed below 100 mg L\(^{-1}\) (see Fig. 7). These arguments are substantiated by the thermal lens results described above.

Another important feature, common to all samples, relates the step-wise nature of these isotherms (see Figs. 7–11). It should be realized that whatever the effect causing this behavior, this should be one with the property of starting and stopping at certain solution concentration values. When the results for the adsorption on glass were first published, we discussed them in terms of micelle formation, since micelles have the above start–stop property, and several reports on micelle formation at CMC had been reported at the time (10). However, since the above data on thermal lens and the above close analysis of the glass and the silica isotherms are not consistent with any CMC, that micelle hypothesis now appears doubtful or wrong (see also section 1). At this time, the new results described above lead us to suggest that the step-wise isotherms appear to be the consequence of properties of the surface and the presence in the sample of fractions
of different solubility in the media. Certainly, in a mixture containing different fractions in solution, the dependence between solubility and concentration should be step wise. Thus, for a mixture of total concentration $c$, containing compounds $c_1$, $c_2$, $c_3$, etc. with solubilities $s_1$, $s_2$, $s_3$, such that $s_1 < s_2 < s_3$, sequential or step-wise precipitation will occur for any value of $c$ above these solubilities. As suggested by the silica–crude oil results (see Fig. 4), this fractional precipitation would be promoted by the surface.

Thus, the first step would correspond to the adsorption of the sample on the glass surface as described above, whereas the other steps should be the consequence of sequential adsorption according to differences in solubility. Thus, after the first step, mainly determined by properties of the mineral surface, the other steps would follow a sequence determined by solubility differences between the different fractions comprising the asphaltene mixture.

According to the PNP results, asphaltenes are a mixture of compounds with different solubilities. Of course, solubility and adsorption are related phenomena, and thus it could be expected that the less-soluble fraction will be adsorbed first.

It is expected that packing of the adsorbed layers would be a slow process because it occurs on the surface (mostly solid phase). As discussed below, this is likely to be the reason for the very slow changes observed in the isotherms (see Figs. 7, and 9–11). A slower process, other than mere adsorption, is also apparent in Figure 5 at long times (see section 3.2).

4.5. Kinetics

Since a first-order rate constant was obtained, the diffusion of solute to the solid surface should be the slow step (see section 2 and 3.2). If so, changing the surface should have no effect on the order. As shown in Table 2, this is the case when the surface is changed from silica to silica covered by asphaltenes. In order to analyze this point further, let us consider the rate for formation of products in both cases (see Methods).

For the silica case:

$$\frac{d[A_1]}{dt} = k_2 [A_1][S].$$  \hspace{1cm} (18)

For the asphaltene case, different site concentration $[S']$ and adsorption rate constant $k'_2$ should be considered. A similar equation is then obtained:

$$\frac{d[A'_1]}{dt} = k'_2 [A'_1][S'].$$

when $k_2 \gg k_{-1}$ and also $k'_2 \gg k'_{-1}$ diffusion to the surface is the slow step in both cases, and these rates become equal to the rate of disappearance of reactants.
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Then:

\[-d[A_m]/dt = k_2 [A_I] [S].\] (20)

and

\[-d[A_m']/dt = k'_2 [A_I'] [S'].\] (21)

The relation between these rates, \( R \), is:

\[ R = \frac{d[A_m']}{d[A_m]} = \frac{k'_2 [A_I'] [S']}{k_2 [A_I] [S]}, \] (22)

or

\[ R = \frac{k'_1}{k_1} = \frac{k'_2 [A_I'] [S']}{k_2 [A_I] [S]}. \] (23)

Since diffusion is the slow step, the concentration of asphaltenes in the interface should be similar in both cases. Therefore, \([A_I'] \approx [A_I]\) and:

\[ k'_1/k_1 = \frac{k'_2 [S']}{k_2 [S]}. \] (24)

This equation shows that when the surface is changed the rate constant will change in all cases except when \( k'_2 [S'] = k_2 [S] \). Thus, although the rate on both surfaces depends on diffusion, in general their relative values will depend on adsorption.

Taking average values for \( k_1 (1.17 \times 10^{-3}) \) and \( k'_1 (5.08 \times 10^{-4}) \), then \( k'_1/k_1 \approx 0.43 \), or less than 1. (see Tab. 2). This is expected in view of the high activity of the silica (a high value of \( k_2 \)) which usually leads to H-type isotherms in many systems, as is the case in Figure 8. Of course, this fact suggests that the affinity of the asphaltene surface by the asphaltene in solution (measured by \( k'_2 \)) is also very high. Otherwise, a much lower value of \( k'_1/k_1 \) should be expected.

It is quite likely that the high values of \( k_2, k'_2, \) and asphaltene composition combine to produce the precipitation results and step-wise isotherms described above for crude oils and adsorption on glass, respectively. For crude oils, the asphaltene surface adsorbed on silica will promote flocculation by irreversible adsorption of the soluble fractions, which keeps the colloidal phase in solution (see below). For the adsorption results, a high value of \( k'_2 \) reflects the low solubility in toluene of the fraction being adsorbed.

4.6. Desorption and Irreversible Adsorption

The very low desorption of asphaltenes shown in Figure 6 strongly suggests that adsorption is mainly an irreversible process. As described above, the adsorbed asphaltene layer is not a monolayer formed by free asphaltenes. To be consistent with the results, that layer should contain aggregates. Due to the high adsorption capacity of silica, irreversible adsorption could be expected for compounds in
contact with or bonded to the surface. However, this should not be the case for molecules, where such contact is not present (see Fig. 12a).

Thus, the above behavior suggests that after adsorption, the asphaltenes could bond to neighbor molecules in the surface, thereby leading eventually to a network insoluble in toluene (see Fig. 12b). It is plausible that the slow changes with time shown by the isotherms on glass (Figs. 9–11) and silica (Figs. 5 and 7) would be due to the building up of such a network. The following analysis and arguments are useful in this regard.

The average of the adsorption rate constants for the silica case is $1.17 \times 10^{-3} \text{ min}^{-1}$ (see Tab. 2). Thus, the average half time is $t_{1/2} = 591 \text{ min}$. Then, after $4 t_{1/2} = 2366 \text{ min} = 40 \text{ hours}$, 94\% of the sample has been adsorbed or the process is about to finish. However, as shown by the isotherms, important changes apparently occur at much later times (see also Fig. 5).

Thus, a process slower than the rate of adsorption must occur in the surface. Of course, these changes are not due to adsorption because this is the fast step. Therefore, in view of the almost irreversible adsorption described above, these slow changes are likely to be due to the building up on the solid surface of the above insoluble asphaltene network. That is, in a kinetically controlled process the molecules are adsorbed or precipitated. This would be followed by a thermodynamic controlled process, where those molecules packing badly are either expelled or relocated in the layer.

This discussion could also be relevant to the PNP results in particular and to colloidal behavior in general. For instance, the above large differences in solubility found for the fractions A$_1$, A$_2$, and A$_3$ (see section 2.2 and PNP sections) could in part be due to differences in packing rather than to large differences in chemical composition or structure. In fact, no significant differences were found in the elemental analysis of these samples (see Tab. 1).

Since fractions A$_1$ and A$_2$ should come from the colloidal phase, the molecules in that phase could engage in the formation of an insoluble network similar to the one described above. As shown in the experimental part (see solubility and precipitation methods), the usual solubility is recovered when fraction A$_2$ is mixed with

![Figure 12](image-url)  
**Figure 12.** Model to illustrate packing leading to insoluble adsorbed layer. (a) Adsorption of monomer (1) dimers (9-4) and trimers (2-6-5 and 3-8-7). Irreversible adsorption could be expected for molecules 1 to 4, but not for the others. (b) bonding between neighbors (packing) leading to insoluble layer or irreversible adsorption.
fraction $A_3$. Evidently, this solubility behavior is a collective property depending on all species present.

The above results and comments could have important consequences on asphaltene colloidal structure and behavior, particularly when the connection between solubility and packing is made. That is, we expect that low solubility indicates a good ability of the sample to pack.

Thus, colloid particles could be envisaged as being composed by a series of layers in which the capacity to form well-packed or low-soluble networks will decrease from the inside to the periphery (see Fig. 13). Molecules packing badly would be located at the periphery or relegated to the solution. Penetration of this loosely packed region or periphery by the solvent or media would allow solubilization of the colloid (see Fig. 13).

When the molecules in the periphery are removed (as in the case of the silica–crude oil experiments, in which they are removed by the silica) the better packed region, impervious to the solvent, will be exposed. This would lead to the observed flocculation.

An analogous situation could be envisaged for the PNP experiments. Here, the loosely bonded molecules will be displaced by the phenol, leading to precipitation. The $A_2$ sample resulting from extraction of the phenol is well packed, and cannot be dissolved by toluene.

In another experiment a sample of asphaltenes was modified by heating to reduce solubility in toluene. When this sample was used as the adsorption surface to adsorb toluene solutions of asphaltenes, the resulting isotherm was also of the step-wise or multilayer type. Besides, adsorption of resins on asphaltenes also yielded step-wise isotherms when measured in hexane, or toluene–hexane mixtures (24). Thus, the step-wise adsorption phenomena appear to be a general property of asphaltenes and resins. Thus, in this sense, surfaces such as silica or glass are convenient media to observe the phenomena, but they are not responsible for it.

Figure 13. Model for colloidal particle (cross-sectional cut). The core region $c$ is inside the discontinuous curve, whereas the peripheral region is outside this curve. Molecules in $c$ are well packed, whereas those in $P$ are loosely packed. Solvent penetration of $P$ (but not $c$) leads to colloid dispersion.
As was the case for asphaltenes, the two-step isotherm found for the adsorption of resins on asphaltenes, should be due to differences in solubility or packing. That is, the resin molecules adsorbed in the first step would have a better ability to pack than those adsorbed in the second.

In this way we see that the above model can easily be extended to petroleum colloids, to afford a picture consistent with all the experimental evidence described above. As shown in Figure 13 and described above, the core of the particle is formed by a sequence of asphaltene layers in which the ability to pack decreases from the center to the periphery, whereas the solubility has the opposite trend. These asphaltene layers will be followed by layers of resins with the same trend in packing capacity. Since a loose packing is expected for the resins in the periphery section, the solvent or media can easily penetrate or diffuse into this section, providing a favorable increase in mixing entropy. This fact, together with a favorable (negative) enthalpy of mixing, will lead to the negative free energy of mixing required for colloid dispersion.

4.7. Adsorption in the Presence of Surfactants

The results corresponding to this section will be published elsewhere. Here we will comment briefly on some results relevant to the present article.

It is interesting to compare the isotherm on silica with and without surfactant (Figs. 8 and 14). Figure 14 shows the isotherm in the presence of 1% NPE (nonylphenol ethoxylated), a quantity small enough to inhibit the adsorption but not enough to stop it completely. It is apparent that this isotherm resembles the step-wise isotherms on glass described above.

Thus, it is quite likely that by adsorbing on very active sites of the surface, the surfactant reduces the adsorbing capacity of the silica surface, thereby leading to a more selective adsorption. Although the step-wise behavior is somewhat present in the isotherm without surfactant, this is blurred by dispersion in the experimental points (see Fig. 7).

It should be noted that the steps in Figure 14 are displaced to the left when compared with the steps observed for the glass case (see Figs. 9–11). This is likely to be due to the differences in surfaces. Thus, location of the steps appears to be a function of the surface. This is consistent with the hypothesis discussed above, which disregards any major influence of aggregation or micelle formation in solution on the location of the steps (see above).

As shown in Figure 14, more than 1% surfactant would be required to stop asphaltene adsorption effectively. According to the results above and the model depicted in Figure 12b, to reach the silica surface, the surfactant should break many bonds in order to effectively penetrate and dislodge the asphaltene layer. From this analysis is apparent that the best results would be expected in cases
where the number of bond ruptures is keep to a minimum. This would be the case for surfactants capable of dislodging large aggregates from the surface; or, in other words, with a capacity to disperse asphaltene colloids.

Although the 1% of surfactant above appears to be a large quantity compared with the asphaltene present, one should bear in mind that the surface is very active. Perhaps a more fair evaluation of surfactant efficiency would require the use of another surface, such as glass or clay. However, the use of low activity surfaces in general is difficult because the amount adsorbed is so low that special techniques, such as the PSD method described above, must be used to obtain quantitative results.

An alternative method for the convenient evaluation of surfactants could be achieved by treating the silica with enough surfactant so as to obtain a surface activity similar to others more representative of any real situation, such as clays and glass. The similarity could be judged by the corresponding isotherms, such as the case of glass and silica +1% NPE described above.

4.8. Relevance of the Present Results to Precipitation during Oil Production

The above results, discussion, and comments are relevant to the problem of asphaltene precipitation during the production of crude oils. A point we have tried
to underscore is that the published theoretical and experimental treatment of this problem had been focused on the effect of pressure on solubility and had neglected the role of surfaces contacted by the crude during production operations.

There is no doubt that a reduction in pressure will reduce solubility in general and asphaltene solubility in particular. However, as shown above, 47% of Furrial asphaltene is insoluble in toluene, and yet the solubility of the complete mixture in toluene is very high (57 g L\(^{-1}\)). As shown above, the colloidal or insoluble phase in the crude oils examined could also be very high, and yet no precipitation at all is observed at room conditions. That reveals that the dispersion capacity of the soluble compounds or compounds in the external layer of the colloid is very high.

For instance, the crude oil can be diluted with large quantities of poor solvents and no asphaltene precipitation is produced. This is the case for Furrial, where more than one volume of n-heptane is required. Thus, even large reductions in solubility parameter do not necessarily lead to flocculation.

However, the sole contact of any asphaltene solution with an active surface, or the dislodging of the dispersion compounds from the colloids, will lead to substantial precipitation of asphaltenes.

The above comments and results suggest that although low solubility is the reason for colloid formation, it is not necessarily the reason for colloid precipitation or flocculation.

The results and data above point out very clearly that flocculation is caused by the desorption of soluble asphaltenes or soluble resins from the surface of asphaltene or petroleum colloids, respectively.

5. CONCLUSIONS

Use of the reductive alkylation technique, combined with measurements in nitrobenzene at high temperature, showed that the molecular weight of asphaltene is comparatively small, being around 1000 g/mol or smaller. From the analysis of the first layer, corresponding to the step-wise adsorption isotherms on glass and silica, the adsorption of free asphaltenes and small aggregates was proposed. The analysis of results suggests the formation of small aggregates in very diluted toluene solutions (less than 100 mg L\(^{-1}\)). This proposition is in agreement with previous results obtained using the thermal lens technique, which suggests that aggregation in toluene should begin around 50 mg L\(^{-1}\). These data and results are consistent with a sequential aggregation process rather than a process like micelle formation at the CMC.

Precipitation experiments performed with toluene solutions of Furrial asphaltenes using PNP showed the presence of fractions of very low (A\(_1\)) and low (A\(_2\)) solubility in toluene (47% of total sample). This suggests that these fractions
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comprise the colloidal phase. These results were consistent with the displacement by PNP of a layer of loosely packed asphaltene adsorbed on the colloidal particle. Analysis of the results led to adsorption isotherms, which suggests very strongly that PNP displaces the above layer after adsorption on the colloid surface.

Induction by silica of asphaltene precipitation in crude oil was justified in a similar manner by proposing a desorption–adsorption mechanism, that is, desorption of the component in the loosely packed layer adsorbed on the colloid periphery and adsorption of these components on the silica surface.

Kinetic measurements showed that the adsorption rate was first order, suggesting that diffusion of asphaltenes to the silica surface is the slow step. Analysis of results showed that adsorption of asphaltene in toluene solutions, on the asphaltene layer formed on silica, should have a very high adsorption rate, similar to the one for silica. In other words, the affinity of asphaltenes in solution by the sites on the asphaltene surface should be very high.

The very slow changes with time and the negligible desorption from the surface measured for the above isotherms were interpreted as the effect of packing or the building up of a low-solubility network on the surface. This would be achieved by the slow formation and rupture of bonds between neighboring molecules at the surface. Thus, molecules with difficulties to pack, adsorbed by the above kinetically controlled process, are either rejected or relocated in a thermodynamic controlled process.

It was concluded that the step-wise adsorption isotherms found for glass and other surfaces is the result of adsorption, after the first step, of asphaltene fractions with a decreasing ability to pack. In other words, fractions of the lowest solubility, such as A1 above, should have the best ability to pack and hence should be adsorbed in the second step. Then other fractions, next in packing ability, such as A2, will be adsorbed, etc.

Preliminary experiments showed that surfactants were able to dislodge asphaltenes from silica and to compete with them for sites on the surface. Sequential isotherms were also observed in the presence of 1% of NPE for toluene solutions of Furrial asphaltenes. This result suggests that displacement from the surface requires destruction of the packing or layer network.

Based on the step-wise adsorption of asphaltenes, the adsorption and precipitation results of PNP, the formation of the colloidal phase by low soluble compounds or compounds with high ability to pack, and other data in the literature, an improved model for petroleum colloids was proposed. The model would be formed by different layers of compounds in which the ability to pack will decrease from the core to the periphery of the particle.

For asphaltene colloids in toluene, the periphery would be formed by compounds belonging to the asphaltene fraction soluble in toluene, whereas the insoluble fractions would be placed at the core. For petroleum colloids, the asphaltene fraction would be at the core and the resins would be packed at the periphery.
loose packing of the periphery would allow solvent penetration, thus dispersing the colloids. Removal of compounds from the periphery will lead to flocculation. Thus, the important precipitation problem presented during oil production is likely to be due to the above phenomena. Although solubility reductions caused by large pressure drops would increase the insoluble material in the colloidal phase, this is likely to produce a minor effect on precipitation. As shown above, the dispersing compounds in the external colloidal layer are capable of dispersing 47% of insoluble material of Furrial asphaltenes.

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APPENDIX

Change in Standard Chemical Potential For the Transfer of Adsorbate From The Colloid Surface S to a General Surface SG.

The situation can be analyzed as follows. The change in the chemical potential of any adsorbate in surface G from the standard state of very low coverage to surface concentration \( X_G \) is:

\[
\mu_G - \mu_G^\circ = RT \ln X_G.
\]

Similarly, for the colloid surface,

\[
\mu_C - \mu_C^\circ = RT \ln X_C.
\]

In equilibrium the chemical potentials are equal, and the change in standard potentials for the transfer of solute from the colloid surface to surface G is:

\[
\mu_G^\circ - \mu_C^\circ = -RT \ln (X_G / X_C) = -RT \ln K_P.
\]

where \( K_P \) is the partition constant. As this equation shows, the transfer will increase with temperature. This simple treatment suggests that increasing the temperature will lead to a more favorable transfer of dispersing, promoting more flocculation.
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