A structural study of a polymer-surfactant system in dilute and entangled regime: Effect of high concentrations of surfactant and polymer molecular weight

Ahmed Aferni, Moez Guettari, Marwa Kamli, Tahar Tajouri, Alain Ponton

To cite this version:
A structural study of a polymer-surfactant system in dilute and entangled regime: Effect of high concentrations of surfactant and polymer molecular weight

Ahmed El Aferni a, Moez Guettari a,*, Marwa Kamli a, b, Tahar Tajouri a, Alain Ponton c

a University of Tunis, Preparatory Institute for Engineering Studies of Tunis, NMR in Polymers and Composites Laboratory, Tunisia
b Faculty of Sciences of Tunis, Faculty of Mathematical, Physical and Natural Sciences of Tunis, Tunisia
c Laboratory MSC, UMR 7057, Univ Paris Diderot-Paris 7, CNRS, Bât. Condorcet, CC 7056, 75205, Paris, Cedex 13, France

ARTICLE INFO

Article history:
Received 11 July 2019
Received in revised form 8 September 2019
Accepted 8 September 2019
Available online 9 September 2019

Keywords:
Polymer
Surfactant
Conductivity
Viscosity
Newtonian behavior
Pseudo-plastic behavior

ABSTRACT

In the polymer-surfactant system, increasing the concentration of surfactant leads to structural changes in surfactant micelles. This is expected to influence the polymer conformation in the dilute regime where polymer chains are single and in the entangled regime where the polymer system is described by the blob concept. In this report, the effect of large Sodium bis(2-ethylhexyl) sulfosuccinate (AOT), surfactant concentrations on the conformation of two polyvinylpyrrolidone (PVP) molecular weight (Mw = 55000 g/mol and Mw = 360000 g/mol) was discussed. In the first part, in the dilute regime, \( C_{\text{surf}} < C^* \) where polymer chains are considered isolated, the determination of PVP (Mw = 55000 g/mol and Mw = 360000 g/mol) intrinsic viscosity \( [\eta] \) showed a coil to globule transition with the increase of temperature. Electrical conductivity measurements have shown the morphological transition (sphere-cylinder) of surfactant micelles with the increase of surfactant concentrations. The addition of AOT wormlike micelles revealed the expansion of PVP chains for the two molecular weight. In the second part, in PVP entangled regime, \( C_{\text{surf}} = 16C^* \) where overlapped polymer chains form molecular blobs, the addition of AOT wormlike micelles to PVP solutions doesn't affect the Newtonian behavior of PVP (Mw = 55000) and the pseudo-plastic behavior of PVP (Mw = 360000 g/mol). However, AOT surfactant reinforces significantly the dynamic viscosity of PVP \( \eta \) for the two molecular weight. By fitting the PVP viscosity-temperature dependence with Fulcher-Tammann-Vogel law, the results showed that the more the surfactant is added, the stronger the PVP solution becomes, which proves that AOT cylindrical (wormlike) micelles presence enhances the solid character of PVP solutions.

© 2019 Published by Elsevier B.V.

1. Introduction

Efforts to investigate polymer-surfactant systems have attracted the attention of researchers for decades. Researchers have always tried to target the different physicochemical aspects related to materials based on polymer-surfactant systems. Among these aspects, the rheological behavior of these systems was a significant issue for scientists because of the correlation between their structure and their flow behavior [1–5]. The first studies of polymer-surfactant systems have been focused on the mechanisms governing the interaction of neutral polymers such as polyvinylpyrrolidone (PVP), polyethylene glycol (PEG) with anionic surfactants, in particular, the sodium dodecyl sulfate (SDS) in aqueous medium [6–8]. Experimentally, these systems have been analyzed typically by viscometry, conductivity, various spectroscopic techniques such as light scattering Fluorescence molecular probe, Magnetic nuclear resonance [9–13] … The results have always revealed the formation of polymer-surfactant complexes, or the so-called polymer-bound micelles. This polymer-surfactant association was considered because of hydrophobic forces [14–21]. However, for various polymers that have polar groups on their structure, such as PVP ... attractive ion-dipole interaction was found to be an engine for this association [22,23]. These studies have targeted systems with low polymer and surfactant amounts in the vast majority of cases. As a result, there was still a lack of work focusing on large-scale polymer surfactant mixtures. In the latest research [24], we showed that the structure
of PVP in semi-dilute regime is affected by spherical micelles-surfactant complexation. In this context, the aim of this study is to initiate research into the structural properties of large-scale polymer-surfactant mixtures. In fact, for larger polymer concentrations, transition occurred from dilute regime, where every single chain acts separately, to semi-dilute regime, in which the chains are getting closer and interpenetrating and where the concept of correlation length have been introduced by De Gennes to be the key prediction to understand the structure of polymer chains in solution. Otherwise, several works have reported that with more increase of polymer concentration, entanglements between polymers chains leads to the appearance of semi-dilute entangled regime, in which static and dynamic properties of polymer chains are described by other scaling laws. In addition, even for the large surfactant concentrations, the surfactant molecules assembly can have several morphologies (spherical micelles, cylindrical micelles, hexagonal structures ...) that necessarily influence their interactions with the polymer. For this reason, a detailed investigation of polymer-surfactant interaction in various concentration domains is of great importance. Possible Correlation between polymer-surfactant behavior in dilute and concentrated mixtures would be a relevant finding. For this reason, we aim in this paper to study and discuss the interaction of AOT anionic surfactant with a typical neutral polymer, PVP. In our research group, we have investigated intensively the PVP physical properties in binaries solvents mixtures, in which PVP exhibits structural transitions . In addition, we have also analyzed the transport phenomena and the rheological behavior of AOT-based reverse micelles. Indeed, it seems to be expected that the interaction PVP-AOT will affect the rheological behaviors of both AOT and PVP.

This work focus on the study of the effect of AOT wormlike micelles on PVP in the dilute and concentrated regime. Our work consists of 4 steps. Firstly, we use capillary viscometry to develop a conformational analysis of PVP in dilute regime under temperature variation. Secondly, conductivity measurements of AOT in water will be discussed. PVP-AOT system in water will be investigated in dilute regime of PVP with the addition of various AOT concentrations. To extend the study from the polymer dilute regime to its concentrated one, the effect of AOT wormlike micelles on the dynamic viscosity of PVP will be analyzed for two PVP molecular weights.

2. Experimental

2.1. Materials

2.1.1. Sample preparation

Sodium bis (2-ethylhexyl) sulfocinate (AOT) (96%) and PVP (M = 360000 g/mol, 55000 g/mol) were purchased from Sigma Aldrich (USA) and used without any additional purification, deionized water was used as a solvent. An electronic balance (OHUAS AS 200) of precision 0.00001 g was used for weighing the appropriate PVP and AOT quantity. Polymer-surfactant samples were prepared by the addition of PVP (AOT) to a well-defined aqueous solution of PVP.

2.2. Methods

2.2.1. Capillary viscometer

The viscosity measurements were performed with a semi-automatic Visco Clock (Schott Instruments) and an Ubbelohde glass capillary viscometer (K = 0.003 mm².s⁻²). A thermostatic bath type CT72/P was used to control (±0.1 °C) the temperature of the sample. The sample was determined by measuring respectively the flow times of the pure solvent, t₀, and the solution, t. The viscometer was washed with distilled water. The flow times were measured with an accuracy of ±0.01s and corrected by a constant provided by the manufacturer. The corrected values, t₀ and t, were obtained by the following equations:

\[ t₀ = t₀ − \theta \]  \hspace{1cm} (1)

\[ t = t − \theta \]  \hspace{1cm} (2)

2.2.2. Electric conductivity

The electrical conductivity was carried out by a Consort C860 conductimeter with two Platine plate attachments electrodes operating at (9-15) V DC. The cell calibration constant was 1 cm⁻¹. A series of standard solutions of KCl with specific concentrations of known conductivity were used for the calibration. A thermostatic bath type CT72/P was used to control the temperature of the studied samples (±0.01K). The relative accuracy relating to the measures of the conductivity is of 1%.

2.2.3. Rheological measurements

The rheological measurements of polymer solutions were performed by BROOKFIELD RST-CC rheometer. The sample cup FTK-RST and spindle CCT-25 were used for the measurement of the steady viscosities of PVP in water solutions at the shear rate from 100 to 1000 s⁻¹. The steady viscosities of PVP and PVP-AOT solutions were carried out at different temperature from 25 °C to 60 °C. The temperature was monitored (±0.1 °C) by a thermostatic bath type CT72/P.

3. Results and discussions

3.1. PVP conformation in the dilute regime

In the dilute regime of polymer solutions, the determination of polymer intrinsic viscosity has been always a crucial step in the study of polymer dimension evolution under the variation of thermodynamic parameters as well the interaction with other chemical entities. Using capillary viscometry, this part's objective is to analyze the effect of temperature on PVP intrinsic
viscosity in dilute regime for the two selected two molecular weights \((M_w = 55000 \text{ g/mol and } M_w = 360000 \text{ g/mol})\). Fig. 1 depicts the change of reduced viscosity versus PVP concentration \(C_{PVP}\), for two PVP molecular weights (a) \(M_w = 55000 \text{ g/mol}\) and (b) \(M_w = 360000 \text{ g/mol}\). As we can see, the two curves exhibit a slope variation from a well-defined concentration, \(C_{PVP}\), called the overlapping concentration. The fact that \(C_{PVP}\) of PVP \((M_w = 55000 \text{ g/mol})\) takes very huge values in comparison with those taken by PVP \((M_w = 360000 \text{ g/mol})\) (Table 1) seems to be evident result since the more the polymer’s chains are longer the more they overlap earlier. Besides, at 50°C more the polymer’s chains are longer the more they overlap earlier. The fact that \(C_{PVP}\) enhances greatly which brings the chains closer together.

Typically, above \(C_{PVP}\), the polymer-polymer interactions intensify, however, below \(C_{PVP}\), the interchain interactions are so weak. We used the Huggins law [40] to determine the intrinsic viscosity of PVP, \([\eta]_f\) for two different temperatures, 25°C and 50°C as follows:

\[
\eta_{id} = [\eta] + KH[\eta]^2C_{PVP}
\]

(3)

where, \(\eta_{id}\) is the reduced viscosity of PVP in water. \(K_H\) is the Huggins constant which measures the polymer-solvent affinity, its value permits therefore to describe the evolution of polymer conformation. Indeed, when \(K_H < 0.5\), the polymer-solvent interactions are attractive and the solvent is good. However, for the bad solvent case, the polymer-solvent interactions are repulsive if \(0.5 < K_H < 1\). The polymer chains aggregate for high Huggins constant values, \(K_H > 1\). We have exposed the results obtained in Table 1 for the two PVP studied molecular weights.

The reduction of PVP intrinsic viscosity with rising temperature (Table 1) reflects a decrease in polymer chains size. We quantified this reduction by using the following equation to calculate the volume contraction rate \(\tau_{e}\):

\[
\tau_e = 100\frac{[\eta]_{25^\circ C} - [\eta]_{50^\circ C}}{[\eta]_{25^\circ C}}
\]

(4)

The values found are 67.32% and 31.57% respectively for \(M_w = 360000 \) and \(M_w = 55000 \text{ g/mol}\). The contraction of the PVP chains is a sign of a Coil-globule conformation transition of PVP chains under the temperature increase. This can be explained by the change of solvent quality at higher temperature. In addition, the contraction rate is sensitive to polymer molecular weight.

### 3.2. AOT structure in water

It is well known that electric conductivity measurements present a powerful method to detect structural transitions of surfactant assembly in binary mixtures (or ternary systems surfactant/polar (apolar) solvent, microemulsions …) [41,42]. Therefore, we used this technique to detect the behavior of AOT assembly in water for various surfactant concentrations. Fig. 2 shows the AOT-specific conductivity,σ logarithmic plot dependence on AOT concentration, which was performed at 25°C. The curve obtained shows 3 regions with 3 different slopes. A critical AOT concentration separates all regions. This behavior shows the variation of the conductive species in the solution by moving from one regime to another. Indeed, the transition from the first to the second regime is characterized

![Fig. 1. Reduced viscosity versus PVP concentration\(C_{PVP}\), for two PVP masses (a) \(M_w = 55000 \text{ g/mol}\) and (b) \(M_w = 360000 \text{ g/mol}\).](image)

![Fig. 2. Logarithmic plot of AOT specific conductivity versus AOT concentration\(C_{AOT}\) at 25°C.](image)
by the critical micellar concentration of AOT in water, CMC. Its value (CMC = 1.4 g/dm³) is in good agreement with the bibliography values [32]. Above this concentration, the conductive species are spherical micelles of AOT. The slope changes again by increasing the concentration of AOT (7 g/dm³), thus, marking the transition to the third regime in which the conductive species’ morphology varies again. Several literature reports suggest that increasing the concentration of surfactants after CMC can lead to variations in micellar morphology. Micelles adopt cylindrical morphologies (Fig. 2).

3.3. Effect of AOT on PVP conformation in the dilute regime

Previously [23], we have studied the effect of AOT monomers and spherical micelles on the PVP intrinsic viscosity. Results revealed the formation of PVP-AOT complex via electrostatic and hydrophobic forces. The effect of AOT cylindrical micelles on PVP conformation was analyzed by the extrapolating the PVP reduced viscosity in the presence of AOT (Fig. 3). Thus, we have determined the intrinsic viscosity of PVP (M₆ = 55000 g/mol and M₆ = 360000 g/mol) in the presence of 4 high AOT concentrations C_AOT (g/dm³) = 6, 8, 10 and 12. As can be seen from Fig. 4, the PVP intrinsic viscosity increase greatly for the two studied molecular weights (Table 2). Thus, the interaction between PVP chains and AOT micelles leads to the expansion of PVP single chain. We can attribute this effect to the rise of electroviscous effects due to the charged anionic micelles. This increase further confirms the formation of the PVP-AOT complex even between the AOT cylindrical micelles and the polymer chains. We have calculated the expansion rate τₐₑₓᵖ of PVP chains as a function of AOT concentration for the two molecular weights according to the following equation:

\[ \tau_{\text{exp}} = \frac{[\eta |C_{\text{AOT}}] - [\eta]}{[\eta]} \times 100 \]  

(5)

where [\eta] and [\eta |C_{\text{AOT}}] are respectively the intrinsic viscosity of PVP in the absence and the presence of AOT. The results found are listed in Table 2. The expansion rate values prove clearly discern that cylindrical micelles expand more greatly PVP (M₆ = 360000 g/mol) in comparison with PVP (M₆ = 55000 g/mol). In the other side, upon the enhance of AOT concentrations, the intrinsic viscosity of PVP (M₆ = 55000 g/mol) continue its increase. However it decreases slightly for PVP (M₆ = 360000 g/mol). This suggests that the PVP (M₆ = 360000 g/mol) chains are shrinking with the addition of more surfactant. This behavior can be explained by a screening effect because of the increase of counter-ions in solution. The screening effect in polymer-surfactant systems was discussed in previous report [43], it was called “a self-screening effect”. The absence of such screening for PVP (M₆ = 55000 g/mol) suggests that electrostatic interactions between polymer and surfactant are sensitive to the monomers number of polymer chain. Moreover, the values obtained of Huggins constant K_H in the presence of various AOT concentrations show a decrease for M₆ = 360000 g/mol which is a signature of high solvent quality and attractive PVP-solvent interactions. Nevertheless, this trend decrease with the rise of AOT in solution which confirms the screening effect appearance. For M₆ = 55000 g/mol, All the obtained values are, at the vicinity of

---

**Figure 3.** Reduced viscosity of PVP (M₆ = 55000 and M₆ = 360000) in the presence of various AOT concentrations.

**Figure 4.** PVP intrinsic viscosity versus AOT concentration for two PVP mass (M₆ (g/mol) = 55000 and 360000), C_AOT at 25°C.

**Table 2**

Intrinsic viscosities, Huggins constants and expansion rates of PVP (M₆ = 55000 and M₆ = 360000 g/mol) at 25°C.

<table>
<thead>
<tr>
<th>C_AOT (g/dm³)</th>
<th>[η] (dm³/g)</th>
<th>K_H</th>
<th>τₐₑₓᵖ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>M₆ = 360000</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>10.1</td>
<td>0.33</td>
<td>–</td>
</tr>
<tr>
<td>6</td>
<td>44.7</td>
<td>0.04</td>
<td>343.37</td>
</tr>
<tr>
<td>8</td>
<td>40.6</td>
<td>0.07</td>
<td>302.88</td>
</tr>
<tr>
<td>10</td>
<td>35.8</td>
<td>0.16</td>
<td>254.68</td>
</tr>
<tr>
<td>12</td>
<td>37.1</td>
<td>0.12</td>
<td>267.33</td>
</tr>
<tr>
<td>M₆ = 55000</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>1.9</td>
<td>0.34</td>
<td>–</td>
</tr>
<tr>
<td>6</td>
<td>2.3</td>
<td>0.60</td>
<td>25.71</td>
</tr>
<tr>
<td>8</td>
<td>3.2</td>
<td>0.35</td>
<td>72.23</td>
</tr>
<tr>
<td>10</td>
<td>3.3</td>
<td>0.42</td>
<td>73.67</td>
</tr>
<tr>
<td>12</td>
<td>3.5</td>
<td>0.52</td>
<td>73.52</td>
</tr>
</tbody>
</table>
3.4. Rheology of concentrated PVP in water

3.4.1. Effect of temperature

In Fig. 5, we reported the two rheograms, viscosity versus shear rate (100–1000) s\(^{-1}\), of PVP in water for a temperature range (25°C–60°C). The selected PVP concentrations were respectively \(\text{CPVP} = 16\text{C}^* = 100g/dm^3\) for PVP (\(M_w360000 g/mol\)) and \(\text{CPVP} = 16\text{C}^* = 740g/dm^3\) for PVP (\(M_w = 55000 g/mol\)). The results showed that the two selected molecular weights had two different rheological behaviors. As can be seen in Fig. 5, viscosity of PVP with low molecular weight (\(M_w = 55000 g/mol\)) remains constant with the increase of shear rate. This is typical Newtonian behavior that can be described by equation:

\[
\tau = \eta \gamma \tag{6}
\]

where \(\tau\) is the shear stress, \(\gamma\) is the shear rate and \(\eta\) is the apparent viscosity of PVP solution. With the temperature rise, viscosity values decrease significantly from 25 to 45°C and decrease slightly for 50–60°C range. Meanwhile, the viscosity of PVP with a higher molecular weight decreases with the increase in shear rate, which reflects the pseudo-plastic rheological behavior of this molecular weight. Typically, the majority of synthetics polymers have shown a non-Newtonian rheological behavior in water [44]. This is because these fluids have a small flow resistance as the shear rate increases. Mathematically, this behavior can be modeled by a power-law model (Ostwald model) [53]:

\[
\tau = K\gamma^n \tag{7}
\]

\[
\eta = K\gamma^{n-1} \tag{8}
\]

where \(\tau\) is the shear stress (Pa), \(\gamma\) is the shear rate (s\(^{-1}\)), \(n\) is the flow behavior index (dimensionless). The lower \(n\), the more pronounced non-Newtonian flow, and \(K\) is the consistency index (Pa.s\(^n\)). In this rheological behavior, \(K\) traduces the solid aspect in the fluid. Thus, in a particular situation, \(K\) has been confused with the dynamic fluid viscosity studied, \(\eta\).

\[n < 1: \text{Pseudo-plastic behavior} \]

\[n = 1: \text{Newtonian behavior} \]

\[n > 1: \text{Dilatant behavior} \]

For pseudo-plastic fluids, \(n < 1.0\). It should be noted that when \(n = 1\), equation (1) reduces to the Newtonian case. With the temperature increase, viscosity shows a similar trend of PVP (55000 g/mol) (Fig. 5).

The different rheological behavior of PVP for the two molecular weights can be explained by the dependence of the polymer correlation length\(\bar{x}\), on the molecular weight at this concentration regime. De Gennes really expected that chains would begin to entangle at their overlap concentration \(C^*\) [24]. This presumption was affected by the way that there is just an adjustment in power-law exponent for the concentration dependence of viscosity for neutral polymers in good solvent, in going from dilute to semi-dilute solution. All the solution properties in this model were described by the correlation length \(\bar{x}\) depending on the size of the polymer in the dilute regime [45].

\[
\bar{x} = R_g,\text{dilute} \left( \frac{C}{C^*} \right)^{-3/4} \tag{9}
\]

where \(C\) and \(C^*\) are respectively the polymer concentration in the solution and overlap concentration. Given that the radius of gyration of PVP in the dilute regime \(R_g,\text{dilute}\) can be calculated from the Flory-Fox relationship which relies the radius of gyration of neutral polymer to its intrinsic viscosity\(\eta\) as follows [46]:

\[
R_g^2,\text{dilute} = \frac{[\eta]M_w}{\phi} \tag{10}
\]

where \(M_w\) is the polymer molecular weight and \(\phi \approx 3.1 \times 10^24\), is the Flory constant. We have calculated the radius of gyration of two molecular weight of PVP at 25°C and 50°C. Then, the correlation lengths of two PVP molecular weights \(\bar{x}_{360000}\) and \(\bar{x}_{55000}\) for PVP concentration \(C = 16\text{C}^*\) were estimated according to equation (13). The found results were listed in Table 3. We can notice the evident decrease of the correlation length with the molecular weights. Besides, a slight lowering was observed with the temperature rise that can be considered because of the change in solvent quality. It seems important also to note that since the correlation length \(\bar{x}\) is considered also as the blob size at this concentrated regime. Therefore, we can estimate the blob volume \(\bar{x}^3\) as reported in Table 3. It is interesting to highlight that at 50°C the blob volume decrease by 64% for \(M_w (g/mol) = 360000\) and 30% for \(M_w (g/mol) = 55000\). Those contraction rates are in good agreement with

![Fig. 5. Viscosity of PVP dynamic viscosity (\(M_w\) (g/mol) – 55000, 360000) as a function of shear rate at different temperatures.](image)
Table 3

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Molecular Weight (g/mol)</th>
<th>Radius of Gyration (Rg) (nm)</th>
<th>Correlation Length (ξ) (nm)</th>
<th>ηfl(360000)/ηfl(55000)</th>
</tr>
</thead>
<tbody>
<tr>
<td>25</td>
<td>Mw = 360000</td>
<td>83.32</td>
<td>10.41</td>
<td>1128.10</td>
</tr>
<tr>
<td></td>
<td>Mw = 55000</td>
<td>28.72</td>
<td>3.59</td>
<td>46.26</td>
</tr>
<tr>
<td>50</td>
<td>Mw = 360000</td>
<td>59.57</td>
<td>7.44</td>
<td>411.10</td>
</tr>
<tr>
<td></td>
<td>Mw = 55000</td>
<td>25.63</td>
<td>1.20</td>
<td>32.76</td>
</tr>
</tbody>
</table>

Scheme 1. Increased entanglement between chains with polymer concentration.

Table 4

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Consistency Index (K)</th>
<th>Flow Behavior Index (n)</th>
<th>R²</th>
</tr>
</thead>
<tbody>
<tr>
<td>25</td>
<td>0.30</td>
<td>0.88</td>
<td>0.99</td>
</tr>
<tr>
<td>30</td>
<td>0.23</td>
<td>0.90</td>
<td>0.99</td>
</tr>
<tr>
<td>35</td>
<td>0.17</td>
<td>0.93</td>
<td>0.99</td>
</tr>
<tr>
<td>40</td>
<td>0.13</td>
<td>0.94</td>
<td>0.99</td>
</tr>
<tr>
<td>45</td>
<td>0.10</td>
<td>0.96</td>
<td>0.99</td>
</tr>
<tr>
<td>50</td>
<td>0.07</td>
<td>0.99</td>
<td>0.99</td>
</tr>
<tr>
<td>55</td>
<td>0.06</td>
<td>1.01</td>
<td>0.99</td>
</tr>
<tr>
<td>60</td>
<td>0.05</td>
<td>1.02</td>
<td>0.99</td>
</tr>
</tbody>
</table>

The table shows the Rheological Parameters (Consistency Index of Fluid, K, and Flow Behavior Index, n) of PVP in water solutions (100 mg/ml) and the real mean square error, R², at Different Temperatures (T (°C)).

those of PVP intrinsic viscosity [η] in dilute regime as it was founded in section 1 (67.32% and 31.5%).

Otherwise, this supposition was immediately brought up by Grassley et al. [47,48] to be mistaken and it is proved that chain entanglement happens at concentration bigger than C*. It has been called entanglement concentration C_e (Scheme 1). Furthermore, in his work on diffusion of dilute and semi-dilute of PVP solutions, Burchard et al. [49] have studied by dynamic light scattering the diffusive behavior of PVP solutions with concentrations up to C_PVP = 6C*, they proved that at these concentrations range, solutions don’t obey to the scaling equations predicted by de Gennes for semi-dilute regime of neutral polymer solutions. Thus, such results were attributed to the rise of chains-to-chains entanglement. The transition between unentangled and entangled regime affect considerably the solution viscosity [27].

Accordingly, at the studied concentration, C_PVP = 16C*. It seems obvious that the two molecular weights of the PVP show differences in the degree of entanglement. Indeed, this affects PVP viscosity and solutions with two molecular weights, M_w (g/mol) = 55000 and 360000, respond differently under shear stresses.

We have summarized in Table 4 the values of the flow index n and the consistency index K of PVP (360000 g/mol) for the studied temperature range accompanied with the root mean square error, R². The increase of n toward 1 with the rise of temperature reflects a trend to Newtonian behavior. In addition, a decrease in fluid consistency K was clearly discernible. It was found that consistency index K (360000 g/mol) varies in the same way with dynamic viscosity (55000 g/mol) as function of temperature (Fig. 6). The transition toward Newtonian fluid behavior of PVP at higher temperature can be correlated to the contraction of PVP chains at this temperature range and the change in solvent quality from good solvent to bad solvent. Typically, for linear variation with temperature, fluids viscosity is described by Arrhenius law [50]. However, shown as the viscosity versus temperature plots in Fig. 6, variation of both dynamic viscosities, η (M_w = 55000 g/mol) and consistency index, K (M_w = 360000 g/mol) with increase of temperature shows nonlinear behavior, it is found that the temperature dependence of the dynamic viscosity in such cases can be frequently adjusted with the Fulcher-Tammann Vogel-law [51,52] as follows:

\[
\eta = A \exp \left( \frac{\beta}{T - T_0} \right) 
\]

(14)

\[
K = A' \exp \left( \frac{\beta}{T - T_0} \right) 
\]

(15)

where the three adjustable parameters are A (A') (Pa·s) is a pre-exponential factor, β(β') (K) is the strength parameter known also as steepness index, is commonly used to classify forming materials as “strong” or “fragile” [51] and the temperature at zero mobility, T0(T0) (K). The VFTH equation with three parameters serves as the basis for the treatment of viscous systems and is able to model many viscous liquids in a wide temperature range. Fitting parameters at various AOT concentrations for the two PVP molecular weights are listed in Table 5. As we can see, the steepness index B takes very huge values for PVP M_w = 360000 g/mol in comparison with PVP M_w = 55000 g/mol. The more the chain is long the more the concentrated PVP solutions are getting more hardness.

3.4.2. Effect of surfactant

Based on the results already seen for the effect of AOT surfactant on PVP viscosity in the dilute regime, where PVP chains act as individual units and inter-chain interactions are almost absent, we have observed that AOT significantly increases the intrinsic viscosity of PVP, which translates into an expansion of the PVP chain. Otherwise, the situation in the concentrated regime is more
complicated due to the entanglement of the polymer chains. Starting from this known situation, we have measured the dynamic viscosity of PVP in water (Mw (g/mol) = 55000, 360000) under shear rate in the presence of 4 different AOT concentrations, CAOT (g/dm³) = 6 ± 5CMC where surfactant micelles are of spherical form and 8, 10 and 12 ± 10CMC where surfactant micelles become cylindrical. Results are plotted in Fig. 7. It can be clearly seen that for both two molecular weights, PVP viscosity shows enormous grow. Meanwhile, we can discern from Fig. 7 that this increase in viscosity begins from CAOT = 6 g/dm³ for PVP with low molecular weight (Mw (g/mol) = 55000), although, for PVP (Mw (g/mol) = 360000), viscosity exhibits a slight at CAOT = 6 and 8 g/dm³ and a significant rise for CAOT = 10 and 12 g/dm³. This difference between two molecular weights viscosity behaviors can be attributed to the difference in blob size between the two molecular weights as well as the degree of entanglement as discussed in the previous section. Otherwise, in the case of PVP (Mw (g/mol) = 360000), viscosity increase with the addition of more surfactant, this can be explained by a change in the morphology of the micelles from spherical form to cylindrical form. We reported in Fig. 8 the variation of the blob size x values versus AOT concentrations. We discerned two different behaviors for each PVP weight. As a matter of fact, the increase of y for PVP (Mw = 55000 g/mol) with AOT concentration traduce the extension of PVP chains as well the PVP blobs due to the electrostatic repulsions between charged surfactant micelles. Nevertheless, for PVP (Mw = 360000 g/mol), with the addition of more surfactant, PVP chains and blobs show a significant contraction. We considered such behavior as evident result of saturated PVP chains with AOT micelles; so, the more addition of micelles can lead to the appearance of a screening effect of highly charged medium. Thus, these results are consistent with those found in a diluted regime where the electroviscous effects of surfactants are better felt by the big molecular weight.

Surfactant assembly may present morphological transitions at these higher compositions (from spherical to cylindrical micelles) [29–31]. In Fig. 9, we have reported the variation of the consistency index K and the fluid index n versus the surfactant content, it is clear that K increases with the addition of more surfactant, particularly, at CAOT = 8, 10 and 12 g/dm³ where the morphology of AOT assembly changes from spherical to cylindrical. This confirms that surfactant enhances greatly the consistency of PVP solutions. Meanwhile, the addition of surfactant micelles did not result in a

![Fig. 6](image_url) Viscosity of PVP (Mw (g/mol) = 55000) and consistency index of PVP (Mw (g/mol) = 360000) as a function of temperature.

### Table 5

<table>
<thead>
<tr>
<th>CAOT (g/l)</th>
<th>Mw (g/mol) = 55000</th>
<th>Mw (g/mol) = 360000</th>
</tr>
</thead>
<tbody>
<tr>
<td>A'</td>
<td>β'</td>
<td>T₀</td>
</tr>
<tr>
<td>0</td>
<td>0.10</td>
<td>79.45</td>
</tr>
<tr>
<td>6</td>
<td>0.0026</td>
<td>700.26</td>
</tr>
<tr>
<td>8</td>
<td>0.0072</td>
<td>447.88</td>
</tr>
<tr>
<td>10</td>
<td>0.0068</td>
<td>514.29</td>
</tr>
<tr>
<td>12</td>
<td>0.0001</td>
<td>1535.50</td>
</tr>
</tbody>
</table>

![Fig. 7](image_url) Viscosity of PVP dynamic viscosity (Mw (g/mol) = 55000, 360000) as a function of shear rate at different AOT concentrations at 25°C.
significant change of fluidity index values $n$, we can distinguish from Fig. 9 that his value presents very slight decrease as we add more surfactant.

Furthermore, we have plotted in Fig. 10 the variation of dynamic viscosity ($M_w$ (g/mol) = 55000) and the consistency index $K$ ($M_w$ (g/mol) = 360000) versus temperature at the presence of different AOT concentration. By fitting the results with Fulcher-Tammann Vogel-law (Equation (14) and (15)), we have listed the different fit parameters in Table 5. As we can see, with the increase of AOT concentration, parameter $\beta$ increases, this increases the strength of PVP solutions, particularly for $M_w$ (g/mol) = 55000. Otherwise, the enormous values are taken by $\beta$ parameter for $M_w$ (g/mol) = 360000 compared to $M_w$ (g/mol) = 55000 reflect a more solid character of the highest molecular weight, consistent with the trend of the variation of the consistency index $K$. 

Fig. 9. Consistency index $K$ and fluid index $n$ of PVP ($M_w$ (g/mol) = 360000) versus AOT concentrations at 25 °C.

Fig. 10. Viscosity of PVP ($M_w$ (g/mol) = 55000) and Consistency index of PVP ($M_w$ (g/mol) = 360000) as a function of temperature at various AOT concentrations.
4. Conclusion

At low surfactant concentrations, several previous reports have shown a consensus that anionic surfactant interacts with neutral polymers by a kind of complexation between the two entities. Meanwhile, very few studies have provided answers regarding the effect of the polymer conformation and surfactant structures on this interaction. In addition, these systems can behave differently in very concentrated polymer and surfactant regimes from their behavior in dilute regimes. In this paper, the experimental investigation of PVP-AOT system has provided valuable results.

- The conformation analysis in the dilute regime has shown that PVP adopts a Coil conformation for low temperatures and a globular conformation for high temperatures.
- Conductivity measurements have shown the existence of structural transition from spherical micelles to cylindrical micelles (wormlike micelles).
- In dilute regime, it was observed that AOT cylindrical assembly expand greatly the PVP chain dimension for the low molecular weight (Mn = 55000 g/mol), Meanwhile, for higher molecular weight (Mn = 360000 g/mol), PVP chains contract with the addition of AOT cylindrical micelles, the difference between the two molecular weight was attributed to the existence of screening effect for the high molecular weight.
- In the entangled polymer regime, PVP dynamic viscosity has shown different behaviors for the two studied molecular weights. A Newtonian rheological behavior for PVP with lower molecular weights and pseudo-plastic behavior for PVP with higher molecular weight. This was explained by the difference in PVP correlation length between the two molecular weights at this concentration regime.
- In the entangled polymer regime, the addition of AOT cylindrical micelles enhances greatly the PVP dynamic viscosity for both two molecular weights. It was emphasized that PVP solutions acquire a more solid character for both molecular weights with the addition of AOT.

Nevertheless, despite interesting results found, this study can only be initiations to the detailed study of system properties in very high concentration regimes.

Conflicts of interest

Authors declare that there is no conflict of interest.

Acknowledgments

Financial support provided by CNRS (French) - DGRST(Tunisia) project,15SR1301, is gratefully acknowledged.

References

[26] F.B. Wyart, P.G. De Gennes, PHYSICAL JOURNAL E Viscosity at small scales in complex fluids, Transitions de texture et de phases induites par le cisaillement : phases lamellaires et éponges de surfactant Aurélien Leon To cite this version : HAL Id : tel-


A. El Aferni et al. / Journal of Molecular Structure 1199 (2020) 127052