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Role of Charge of Micellar Worms in Modulating Structure and Rheological Properties of Their Mixtures with Nonionic Polymer

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ABSTRACT: The paper describes phase behavior, microstructure, and rheological properties of the mixtures of surfactant micelles of varying negative charge with nonionic hydrophilic polymer poly(vinyl alcohol) (PVA) weakly interacting with the micelles. To reduce the charge of micelles, a cationic surfactant n-octyltrimethylammonium bromide was gradually added to the anionic surfactant potassium oleate. No appreciable interactions between the micelles and polymer were revealed by NMR spectroscopy. At the same time, the system shows a broad range of compatibility, which may be due to surfactant counterions preventing the demixing. The higher the concentration of added polymer, the higher the charge density of micelles necessary to prevent macrophase separation. However, SANS data demonstrate that upon the addition of polymer the intermicellar distances decrease, which suggests that the micelles are expelled from some area filled by PVA macromolecules, indicating a local demixing. A decrease of micellar charge density induces the growth of micelles from short nonoverlapping cylinders to long entangled wormlike chains both in the presence and in the absence of polymer, which is accompanied by an increase of viscosity by up to 5 orders of magnitude. Despite very weak interactions with the micelles, PVA makes their growth in length much more pronounced and forms the entanglements with micellar chains, thereby enhancing significantly the viscoelastic properties. The effect of polymer on micellar length was attributed to polymer–surfactant microsegregation increasing the local surfactant concentration and consequently the length of micelles. These data demonstrate the effectiveness of the addition of weakly interacting polymer for the modulation of rheology of wormlike surfactant micelles, thus opening a new way for manipulating the rheological properties of viscoelastic surfactants.

INTRODUCTION

Wormlike micelles (WLMs) of ionic surfactants can form transient networks imparting viscoelastic properties to aqueous solutions.1–9 These properties are widely exploited for thickening and rheology-control applications in many area including cosmetics, turbulent drag reduction, oil recovery, and so on.10–13 A large variety of applications stimulates an appropriate modification of the viscoelastic properties of WLM solutions.

One of the ways to modify the rheology of WLMs consists in their combination with polymers.10–23 In many cases, addition of polymeric chains enhances the viscoelasticity of WLM solutions.10–18,23 In particular, an increase of viscosity by few orders of magnitude was observed at bridging the WLMs by macromolecules interacting simultaneously with different micelles.10–18 The interactions can be either electrostatic or hydrophobic, or both. The electrostatic attraction occurs when polymer is oppositely charged with respect to the micelles like in the case of cationic, cellulose-based polyelectrolyte, and anionic surfactants sodium dodecylethoxysulfate.10 The hydrophobic interactions usually prevail when uncharged water-soluble polymers with few hydrophobic side11–13 or end14–16 groups capable to penetrate in the surfactant core are added to WLMs. Such a situation was described for uncharged hydrophobically modified (HM) polysaccharide guar with cationic surfactant erucyl bis(hydroxyethyl)methylammonium chloride (EHAC),11 for HM polyacrylamide with EHAC12 and cationic gemini surfactant 1,2-N,N′-bis(dimethyloctadecyl)ethene ammonium bromide,13 and for poly(ethylene oxide) end-capped with alkyl groups interacting with cationic cetyltrimethylammonium tosylate14,15 and cetylpolyridinium chloride/sodium salicylate16 micelles. Hydrophobic attraction also governs the bridging of WLMs by HM polyelectrolytes of similar charge in the presence of salt as was demonstrated for partially hydrolyzed polyacrylamide mixed with anionic WLMs of potassium.
oleate. Both electrostatic and hydrophobic binding to the WLMs are expected for weakly charged HM polyelectrolytes carrying opposite charge with respect to the micelles.

Independently of the nature of polymer–surfactant interactions, it is important to ensure that these interactions are not too strong to violate the geometric balance of the head and tail groups of surfactant (packing parameter) enough to destabilize the cylindrical geometry. Otherwise, it will lead to the disruption of WLMs as a result of cylinder–sphere transition accompanied by the loss of viscoelastic properties or even to the precipitation of the polymer–surfactant complexes.

Recently, it was shown that a pronounced enhancement of rheological properties can be achieved by using a polymer weakly interacting with the micelles, if one can get a homogeneous system with sufficiently high concentration of the components providing a significant interfacial between polymer and micellar chains. This system seems to be very promising since it can be based on a variety of common commercially available water-soluble polymers. To enhance compatibility of the weakly interacting components, the micellar chains should be charged and the ionic strength of solution low, which makes demixing unfavorable, since it will induce a pronounced loss of translational entropy of surfactant solution low, which makes demixing unfavorable, since it will foster a better understanding of the behavior of the mixed systems based on a combination of “living” micellar assemblies and polymeric chains, which can open the way for new practical applications of these industrially important systems.

**EXPERIMENTAL SECTION**

**Materials.** Potassium oleate (potassium cis-9-octadecanoate, C16H35O−C2H5(OCOOK) was purchased from TCI (purity >98%), C8TAB was obtained from ABCR (purity >98%), and PVA with a molar mass 27 000 g/mol (degree of polymerization N = 600) and residual content of acetate units 1.1–1.9% (Mowiol 4-98) was obtained from Aldrich. All chemicals were used without further purification. Distilled deionized water (Millipore Milli-Q) was used for preparation of the samples for phase behavior and rheological studies; D$_2$O obtained from AstraChem (Russia, 99.9% isotopic purity) was used for SANS and NMR measurements.

The critical micelle concentrations of potassium oleate and C$_8$TAB are ~0.9 mM$^{23}$ and 0.14 M$^{-1}$. In all experiments, the concentration of C$_8$TAB never surpassed 0.1 M, so this surfactant did not form micelles itself. The overlap concentration C* of PVA macromolecules is 0.7 monomol/L as determined by viscometric measurements.

**Sample Preparation.** To obtain stock solutions, surfactants were dissolved in water overnight at room temperature. PVA stock solution was made by dissolving it in water for 1 h in an oil bath at 90 °C. Then, the samples were prepared by mixing stock water solutions of surfactants and PVA in appropriate ratios and stirring them with a magnetic stirrer for 1–2 days. After preparation, the samples were left at room temperature for 1–3 weeks before examination.

**Phase Behavior.** For phase behavior studies, the samples were visually inspected 3 weeks after preparation. To determine the composition of each phase, the samples were prepared in D$_2$O and left to phase segregate, and after that the phases were carefully separated from each other by a syringe and examined by $^1$H NMR. For these experiments, the upper highly viscous phase was diluted 1:1 with D$_2$O, whereas the lower phase was studied as is.

**$^1$H Nuclear Magnetic Resonance.** NMR measurements were performed with an Agilent 400 MR spectrometer operating at a frequency of 400 MHz and equipped with an automated triple broadband detector. The samples prepared in D$_2$O were put into standard 5 mm quartz tubes (Norell); the spectra were recorded at 20 °C and accumulated from eight scans. The data were processed using ACD Laboratories 6.0 software, including baseline and reference correction and calculation of peak integrals. Concentration ratios of different compounds in the samples were calculated from the ratio of areas under corresponding peaks.

**Small-Angle Neutron Scattering.** SANS measurements were done at the IBR-2 pulsed reactor, Frank Laboratory of Neutron Physics, Joint Institute for Nuclear Research (Dubna, Russia). A YuM0 spectrometer was used, which has two ring detectors of scattered intensity covering the scattering vectors q dynamical range 0.005–0.7 Å$^{-1}$ due to a two-detector system. The intensity was measured in absolute units (cm$^{-1}$) by using a vanadium standard. Since most of the samples were highly viscous, they were put in the specially constructed mountable cells with parallel quartz plates (beam path 2 mm). The temperature during measurements was controlled by a water-circulating thermostat at 20.0 ± 0.5 °C.

All compounds (surfactants and polymer) were hydrogenated; therefore, the samples were prepared in D$_2$O to obtain high contrast. The scattering length densities of surfactant alkyl tails, polymer, and D$_2$O are equal to $-0.30 \times 10^{-6}$, $0.65 \times 10^{-6}$, and $6.36 \times 10^{-6}$ Å$^{-2}$, respectively. However, the scattered intensity of polymer solutions is 1–2 orders of magnitude lower than that of surfactants, since micelles are much “thicker” objects. Therefore, it was assumed that the scattered intensity is mostly contributed by surfactant micelles.

Primary treatment of the SANS data included corrections for the sample transmission, sample thickness, and electronic noise by the SAS program. Incoherent (background) scattering was subtracted from the data by using a blank solvent (D$_2$O).

Fitting of the scattering curves was performed by the program SasView (http://www.sasview.org/). A part of the scattering curve at
wave vectors \( q \) higher than the structure peak position \( q^* \) (0.06–0.07 Å\(^{-1}\)) was fitted by a form factor of an infinite cylinder (without structure factor). Two fitting parameters were used: radius of the cylinder and background level. Fitting of the scattering curves in the full \( q \)-range was performed by combining a finite cylinder form factor with a Hayter–Penfold potential which accounts for electrostatic repulsion of charged objects.\(^{53}\) The radius of the cylinder and background were fixed, and their values were taken from the high \( q \)-range fitting by an infinite cylinder. In the full \( q \)-range fit, two fitting parameters were used: length of the cylinder and its charge. In the dilute regime, a similar fitting procedure has been recently applied for elliptical charged micelles of cetyltrimethylammonium chloride with sodium salicylate,\(^{34}\) and in our case it allows direct calculation of the length of cylindrical micelles. In the semidilute regime, the mesh size of the entangled micellar network \( \xi \) is obtained from the SANS fit instead of the micellar length, since the micelles are much longer than \( \xi \), so that the largest size of an “inhomogeneity” is the mesh size, and the network is a homogeneous object on the length scales larger than \( \xi \).\(^{35}\)

Rheology. Rheological measurements were carried out with a stress-controlled rotational rheometer Physica MCR 301 (Anton Paar, Austria) as described elsewhere.\(^{30,37}\) In order to study highly viscous samples (with zero-shear viscosity \( \eta_0 > 0.1 \) Pa s), a stainless steel cone–plate (diameter 50 mm, cone angle 1°) was used. For samples with lower viscosity \( (\eta_0 < 0.1 \) Pa s), double-gap coaxial cylinders (mean diameter 26.4 mm, height 40 mm, gap 0.42 mm) were implied. During measurements, the temperature was set at 20.00 ± 0.05 °C by a water-circulating thermostat and Peltier elements. A special measuring cell cover filled with solvent was used to prevent solvent evaporation from the sample. To equilibrate the sample after putting it into the rheometer, it was kept inside the measuring cell for 10–30 min and subjected to low-amplitude (0.5%) oscillatory deformation with a frequency \( \omega \) of 1–10 s\(^{-1}\) and amplitude 0.5–3% prior to investigation. Rheological measurements were carried out at least two times to ensure that they coincide and the sample is equilibrated in the cell.

In oscillatory shear experiments, the angular frequency dependences of the storage \( G'(\omega) \) and loss \( G''(\omega) \) moduli were measured. All measurements were made in the linear viscoelastic regime at the deformation amplitudes of 0.5–3%, at which the storage and loss moduli were independent of deformation, as estimated previously by amplitude sweep tests performed at the frequency of 10 s\(^{-1}\). The plateau modulus \( G_0 \) was determined from the \( G'(\omega) \) dependence at the frequency \( \omega_{\text{m}} \), where \( G' \) gets a minimum value.\(^{38}\) The terminal relaxation time \( \tau \) was estimated as \( \tau = 1/\omega_{\text{m}} \) from the frequency \( \omega_{\text{m}} \), corresponding to the interception of \( G'(\omega) \) and \( G''(\omega) \). In steady shear experiments, the flow curves (variation of viscosity with shear rate) were obtained. From these curves, the values of the zero-shear viscosity \( \eta_0 \) were determined by fitting the rheological data with the Carreau–Yasuda model.

## RESULTS AND DISCUSSION

### Phase Behavior

First, we studied the effect of charge density of surfactant micelles on the phase behavior of potassium oleate/C\(_8\)TAB/PVA mixtures. In these experiments, the concentration of anionic surfactant (potassium oleate) was kept constant at 0.078 M (2.5 wt %), whereas the concentration of cationic surfactant (C\(_8\)TAB) was changed from 0.0078 to 0.1 M (0.2 to 2.5 wt %), which corresponds to the variation of the molar ratio [C\(_8\)TAB]/[potassium oleate] \( \beta \) in the range 0.1–1.3.

The partial phase diagram of potassium oleate/C\(_8\)TAB/PVA mixtures in water is presented in Figure 1. It shows that in the absence of PVA, mixed surfactants form one-phase homogeneous solutions in the whole studied range of [C\(_8\)TAB]/[potassium oleate] molar ratios \( \beta \). In the presence of PVA, solutions are homogeneous at low molar ratios \( \beta \) but phase separate when C\(_8\)TAB content is increased. According to \(^1\)H NMR spectroscopy (Figures 1S and 2S) and elemental analysis data for the demixed systems, the viscous upper phase represents the surfactant solution with some amount of polymer, whereas the lower phase of small viscosity contains only polymer, which indicates the segregation of the components. This is consistent with the results of \(^1\)H NMR studies of model systems (potassium oleate/PVA and C\(_8\)TAB/PVA) presented in Figure 3S. It is seen that the chemical shifts in the spectra of polymer–surfactants mixtures do not differ from those in the spectra of PVA (Figure 1Sb), potassium oleate (Figure 2Sb) and C\(_8\)TAB (Figure 2Sc) taken separately. Therefore, one can conclude that the molecular environment of surfactant protons does not change upon the addition of polymer, which implies that they do not attract each other.

Similar segregative phase separation was observed, for instance, in the mixtures of two slightly incompatible polymers.\(^{40}\) In such systems, two counteracting factors govern the phase behavior: on the one hand, weak repulsive force between the components promotes phase separation and, on the other hand, entropic reasons counteract it. In the present system, the entropic reason counteracting demixing consists in the loss of entropy of surfactant counterions, when they accumulate mainly in one phase.

At low C\(_8\)TAB content, the micelles are strongly negatively charged, and the phase separation is unfavorable for entropic reasons. When the molar ratio [C\(_8\)TAB]/[potassium oleate] \( \beta \) becomes higher, the net charge of micelles decreases alongside with the amount of their counterions, and above a certain point the entropy losses become less important than the repulsion between the micellar and polymer chains, and the phase separation occurs. Note that the WLMs are “living” objects, and the decrease of their charge density is inevitably accompanied by their growth in length.\(^{4,41}\) Indeed, with decreasing charge density the electrostatic repulsion in the corona of micelles becomes smaller, which makes tighter packing of surfactant molecules in the middle cylindrical part of the micelle more favorable in comparison with the semispherical ends, thus reducing the fraction of the end-caps in the system. A smaller fraction of end-caps at the same amount of surfactant suggests longer micelles. Increased length of micelles can also contribute to the demixing, since longer chains are poorer in entropy, but this entropic contribution is much smaller than the entropy of counterions.

From Figure 1 it is seen that the region of phase compatibility narrows with the increase of polymer concentration; i.e., the higher the amount of added PVA, the more
charged micelles can be demixed into a separate phase. This is due to the enhancement of the repulsion between the components. At the largest studied polymer concentration (1.34 monomol/L), the solutions are homogeneous only for highly charged (and short) micelles at very low molar ratios $\beta \leq 0.2$. Further experiments were performed only with homogeneous samples.

**Structure.** To investigate the structural changes, which occur in the potassium oleate/C$_8$TAB/PVA system at varying micellar charge, SANS was used. In the experiments, the concentration of PVA was kept at 0.89 monomol/L (4 wt %), which is slightly higher than its overlap concentration in water ($C^* = 0.7$ monomol/L). The concentration of potassium oleate was equal to 0.078 M, whereas the value of the molar ratio $[C_8$TAB$]/[potassium$ oleate$]$ $\beta$ was varied from 0.1 to 0.45, which corresponds to homogeneous solutions (Figure 1) with strongly negatively charged micelles. Since surfactant micelles in D$_2$O scatter neutrons much stronger than PVA (Figure 4S), the SANS curves in mixed surfactant/polymer systems represent mainly the scattering from the micellar component.

**Effect of Micellar Charge.** Figure 2a shows the evolution of the scattering pattern with decreasing degree of charge of micelles (i.e., increasing amount of added C$_8$TAB) at fixed concentrations of both potassium oleate and PVA. It is seen that in the high q-range (at $q > 0.07 \text{ Å}^{-1}$) the curves superimpose perfectly, indicating that the local structure of surfactant aggregates is the same independently of the micellar charge. In this range, the curves can be fitted by a form factor of a cylinder with the radius of 18.1 ± 0.4 Å (Figure 2a), which is close to the length of oleate tail (19 Å) and is much larger than the length of C$_8$TAB alkyl group (8 Å), suggesting that at the studied molar ratios $\beta = 0.1$–0.45 the radius of an OCS micelle is governed by the size of the longer anionic surfactant tail and does not depend on the content of C$_8$TAB. Similar behavior was previously observed for sodium oleate/C$_8$TAB micelles in the absence of polymer. Fitting of the high q-range of the scattering curves in order to obtain the micellar radii was applied for several charged surfactant–micellar systems.

Thus, according to SANS data, at $\beta$ values under study, the micelles of locally cylindrical shape are always present in potassium oleate/C$_8$TAB/PVA solution, and their radius is not altered by the amount of added C$_8$TAB. At the same time, at low q-values ($q < 0.05 \text{ Å}^{-1}$) reflecting larger scale structural features the scattering curve is strongly affected by the charge of micelles: with decreasing charge density (i.e., with increasing molar ratio $\beta$) the scattered intensity $I$ is greatly enhanced by nearly an order of magnitude (see also the scattering curves in linear representation on Figure 5S). An increase in SANS intensity at low q was previously observed at the screening of the micellar charges with low molecular weight salt or oppositely charged surfactant. It was attributed to the elongation of cylindrical micelles. Indeed, addition of oppositely charged surfactant produces two main effects: (i) the decrease of the net charge of micelles and (ii) the release of counterions increasing the ionic strength of the solution. Both these effects reduce the electrostatic repulsion in the corona of the micelles, thus favoring tighter packing of surfactant molecules in a cylinder at the expense of spherical end-caps. The lower the fraction of end-caps, the longer the micelles. In our work, we show that growth of cylindrical micelles in length with decreasing their charge density occurs also in the presence of PVA. Besides, one should note that decreasing charge of micelles will make the intermicellar repulsive interactions weaker, thereby reducing the contribution of the structure factor. As the structure factor decreases the scattering intensity at low q, the lowering of its contribution will enhance the intensity. So, the increase of scattering intensity at low q can be attributed to two reasons: elongation of micelles and reducing of the intermicellar repulsion. One can expect that the first reason will be important mainly in dilute solutions because in semidilute regime the scattering will characterize the mesh size of the transient network.

Figure 2a demonstrates that at low q the intensity $I$ of experimental curves is smaller than the predicted cylinder form factor, which can be assigned to strong electrostatic repulsion between the similarly charged cylindrical micelles, that is, to the contribution of the structure factor. The scattering curves of surfactant micelles (in the absence of polymer) can be well-fitted in the whole q-range by a model combining a finite cylinder form factor with a Hayter–Penfold potential which accounts for electrostatic repulsion of charged objects (Figure 6S). The fitting gives an estimate of the length of micelles (Table 1S): with increasing $\beta$ from 0.2 to 0.275 (dilute regime) the length of micelles grows from 28 to 40 nm. Note that in this part of the curve we do not observe the scaling dependence $I \sim q^{-1}$ typical for cylindrical objects at $qR \ll 1$ because their form factor is hidden by the contribution of the structure factor.
At intermediate \( q \)-values (\( q = 0.05 - 0.07 \ \text{Å}^{-1} \)) a correlation peak is observed. It is better seen in \( lq \) versus \( q \) representation (Figure 2b) which eliminates the contribution of the form factor in this part of the curve.\textsuperscript{45} The structure peak characterizes the mutual arrangement of similarly charged micelles resulting from electrostatic repulsion between them, which should be rather strong since no salt was added to shield the repulsion. The variation of the position of the peak \( q^* \) with molar ratio \([C_8\text{TAB}]/[	ext{potassium oleate}] \) \( \beta \) for the potassium oleate/C\(_8\)TAB/PVA system is illustrated in Figure 3 (red circles). It is seen that the curve consists of two distinct regions. In the first one (0.1 < \( \beta < 0.25 \)), the lowering of the micellar charge induces a pronounced decrease of \( q^* \), indicating that the intermicellar distances \( 2\pi/q^* \) become much longer, which can be due to the growth of micelles in length.\textsuperscript{46} Indeed, from simple geometrical considerations it can be concluded that in the dilute regime the average distance between longer cylindrical objects should be larger than between the shorter ones made of the same amount of material (the concentration of C\(_8\)TAB added in this range is several times lower than that of potassium oleate). At the molar ratio \( \beta \approx 0.25 \), a sharp inflection is seen at the curve (Figure 3). A similar dependence with an inflection point was observed in the literature for cylindrical micelles of cetyltrimethylammonium tosylate at increasing concentration of surfactant, and the inflection was attributed to the crossover from dilute to semidilute regime.\textsuperscript{46} Although in our system the composition of the surfactant mixture \( \beta \) (instead of the concentration of surfactant) was varied, one can suggest that at \( \beta \approx 0.25 \) the transition to semidilute regime occurs. Thus, the first region on the curve (0.1 < \( \beta < 0.25 \)) corresponds to the dilute regime, whereas the second one (\( \beta > 0.25 \)) to semidilute regime (Figure 3). In semidilute regime, the micelles overlap and their growth soon results in the formation of an entangled network (as will be evidenced later by the rheological data). In the semidilute entangled regime, \( 2\pi/q^* \) roughly corresponds to the mesh size \( \xi \) of the network.\textsuperscript{46} As seen from Figure 3, in the second region (\( \beta > 0.25 \)), \( q^* \) slightly increases with C\(_8\)TAB concentration, indicating the reduction of \( \xi \), i.e., the formation of denser network. Plotting \( q^* \) vs total (potassium oleate and C\(_8\)TAB) surfactant concentration (Figure 3, inset) gives a straight line. The slope of this line is close to the theoretical dependence\textsuperscript{46} for strongly interacting cylindrical objects in semidilute regime: \( q^* \sim C^{0.5} \). Therefore, the observed slight growth of \( q^* \) in the second region can be simply explained by the increase of the total amount of surfactant molecules building the micelles. Thus, the analysis of the SANS data suggests that decreasing micellar charge induces the growth of cylindrical micelles in length in the presence of polymer, which leads finally to the transition from dilute to semidilute regime at \( \beta \approx 0.25 \) and to the formation of a polymer—micellar network.

**Effect of Polymer.** In order to reveal the role of polymer in the microstructure formation, let us compare the scattering curves of potassium oleate/C\(_8\)TAB solutions in the presence and in the absence of PVA at a constant molar ratio \([C_8\text{TAB}]/[	ext{potassium oleate}] \) \( \beta \). As depicted in Figure 4, at intermediate \( q \)-values (0.06 \( \text{Å}^{-1} < q < 0.2 \ \text{Å}^{-1} \)), the curves coincide with each other showing that the local structure of surfactant aggregates is not affected by PVA. A clear effect of polymer is seen at high and at low \( q \)-ranges (Figure 4). At high \( q \)-range (\( q > 0.2 \ \text{Å}^{-1} \)), polymer increases the background intensity of the scattering curve (cf. Figure 4S). In the low \( q \)-range, the effect of polymer is more complicated. Without PVA, the scattered intensity levels off at small \( q \). This may be explained by the formation of an entangled micellar network (which will be evidenced later by the rheological data). The network can be regarded as a “homogeneous” object at length scales larger than its mesh size (i.e., there are no larger inhomogeneities than one mesh), which results in the constant intensity at low \( q \)-range.\textsuperscript{35} At the same time, in the presence of PVA, the intensity is higher, and moreover it grows with decreasing \( q \), which indicates the formation of structures larger than \( 1/q_{\text{min}} = 150 \ \text{nm} \). This is consistent with our recent cryo-TEM data\textsuperscript{39} on potassium oleate/C\(_8\)TAB/PVA mixtures demonstrating the formation of large (several hundreds of nanometers) domains enriched either by WLMs or by polymer, which arise from poor compatibility between polymer and surfactant micelles. These domains seem to be responsible for the enhancement of intensity at low \( q \). Therefore, if we return to the scattering curves at different molar ratios \( \beta \) (Figure 2 and Figure 5S), we can state that in the presence of PVA the rise of low-\( q \) intensity at decreasing charge density of micelles proceeds due to three
reasons (formation of microdomains, increase of micellar length, and reduction of repulsive interactions), whereas in pure surfactant only last two reasons (the growth of micelles and the decrease of repulsion) will contribute to this effect. Indeed, as was discussed above, the scattering curve of the surfactant micelles (without PVA) is well-fitted in the whole q-range by a model combining a finite cylinder form factor with the Hayter—Penfold potential, which accounts for electrostatic repulsion of charged objects (Figure 4 and Figure 6S). Similar fitting of polymer—surfactant system deviates from the experimental scattering curve (Figure 4) at low q-values, indicating additional factor affecting the scattering, which most probably concerns the polymer—surfactant microsegregation with the formation of microdomain structure.

Figure 3 demonstrates the influence of polymer on the dependence of the structure peak position $q^*$ on the molar ratio $[C_8\text{TAB}]/[\text{potassium oleate}]$ $\beta$. It is seen that the curves with and without PVA show similar behavior, suggesting that the general structural evolution of micelles with the decrease of their charge proceeds in the same way in both cases. The most significant effect produced by polymer consists in the shift of the whole curve to higher $q^*$-values, indicating that at all micellar charges under study (i.e., at all $[C_8\text{TAB}]/[\text{potassium oleate}]$ molar ratios $\beta$) the average distance between the micelles decreases when PVA is added. This implies the local concentrating of micelles induced by their segregation from polymeric chains poorly compatible with the micelles. Indeed, as the surfactant micelles cannot occupy some area filled by PVA macromolecules, they need to come closer to each other upon addition of polymer; i.e., their local concentration increases. This should lead to the additional elongation of micellar chains, since the length of wormlike micelles grows with their concentration. Also, Figure 3 shows that in the presence of PVA the minimum on the curve $q^*(\beta)$ is slightly shifted to lower $\beta$, indicating that the overlap concentration $C^*$ of the micelles in polymer—surfactant system is observed for more charged micelles, which may be due to their additional elongation caused by polymer.

Thus, SANS data suggest that in the presence of polymer the local concentration of micelles and their length are higher in comparison with PVA-free micellar solutions and that the inhomogeneities at the scale larger than ca. 150 nm are formed. This is consistent with the behavior described by us recently for potassium oleate/C8TAB/PVA solutions at fixed $\beta = 0.4$ and can be assigned to the local segregation of polymer and surfactant components in the solution.

**Rheology.** In order to relate the structural changes in potassium oleate/C8TAB/PVA solutions with their macroscopic behavior, rheological studies were performed. Figures 5 and 6 show the flow curves and the dynamic rheological data at different $[C_8\text{TAB}]/[\text{potassium oleate}]$ molar ratios $\beta$ and fixed concentration of PVA (0.89 monomol/L), whereas Figure 7 depicts the variation of zero-shear viscosity $\eta_0$ relaxation time $\tau$, and plateau modulus $G_0$ with increasing $\beta$ at different amounts of polymer. In all the experiments the concentration of potassium oleate was equal to 0.078 M.

**Effect of Micellar Charge.** Let us first consider the evolution of rheological data with decreasing micellar charge (i.e., at increasing $\beta$) at constant PVA concentration of 0.89 monomol/L. The corresponding concentration results are presented on Figures 5 and 7a (red curve). From Figure 7a, it is seen that at $\beta = 0.1$ the viscosity is low (0.006 Pa s) and only slightly higher than the viscosity of 0.89 monomol/L PVA solution without surfactants (0.005 Pa s), which is in accordance with the presence of rather short cylindrical micelles. When the molar ratio $\beta$ is raised from 0.1 to 0.23, the viscosity becomes only slightly higher suggesting the elongation of cylindrical micelles in dilute regime. At molar ratio $\beta \approx 0.23$, the rheological behavior changes drastically. The zero-shear viscosity starts to rise sharply (Figure 7a), indicating the transition from dilute to semidilute regime, and finally grows by nearly 4 orders of magnitude, reaching the value of 900 Pa s at $\beta = 0.46$. As seen from flow curves (Figure 5), viscous solutions exhibit shear-thinning behavior.

Dynamic rheological data (Figure 6) demonstrate that with decreasing micellar charge at the addition of C8TAB the solutions progressively gain viscoelastic properties. The intercept point between $G'(\omega)$ and $G''(\omega)$ curves shifts to lower frequency indicating the increase of the relaxation time $\tau$. Note that the present systems do not show simple Maxwell behavior with a single relaxation time. Therefore, the relaxation time determined from the intersection of $G'(\omega)$ and $G''(\omega)$ curves can be regarded as a terminal one. At molar ratios $\beta > 0.38$, one can observe a well-defined high-frequency plateau on $G'(\omega)$ dependence indicating to the formation of an entangled wormlike micellar network. The dynamic rheological data for lower polymer concentration 0.22 monomol/L are quite similar (Figure 7S).

![Figure 5](image-url) Flow curves of 0.078 M potassium oleate aqueous solutions containing 0.89 monomol/L PVA and different amounts of C8TAB corresponding to molar ratios $[C_8\text{TAB}]/[\text{potassium oleate}]$ $\beta$: 0.1 (reverse cyan triangles), 0.24 (gray stars), 0.32 (green triangles), 0.34 (red diamonds), 0.38 (blue circles), and 0.46 (black squares) at 20 °C.

![Figure 6](image-url) Frequency dependences of storage $G'$ (filled symbols) and loss $G''$ (open symbols) moduli for 0.078 M potassium oleate aqueous solutions containing 0.89 monomol/L PVA and different amounts of C8TAB corresponding to molar ratios $[C_8\text{TAB}]/[\text{potassium oleate}]$ $\beta$: 0.32 (green triangles), 0.34 (red diamonds), 0.38 (blue circles), 0.42 (orange reverse triangles), and 0.46 (black squares) at 20 °C.
evolution of rheological properties can be explained by the elongation of micelles upon decreasing their charge leading to the transition of the solution to semidilute regime.

Figures 7a and 7b show that $\eta_0$ and $\tau$ grow up to maximum and slightly decrease after it. The maximum on the viscosity curves versus salt concentration is usually attributed to the formation of branching points on WLMs. Unlike branches in polymers, micellar branches are labile. They can slide along a micelle contour, thus accelerating the stress relaxation and therefore decreasing relaxation time $\tau$ and terminal relaxation time $\tau$ already stay constant or even slightly decrease. As was shown for polymer-free sodium oleate/C8TAB system, in the vicinity of the viscosity maximum the value of $G_\beta$ is independent of the composition of OCS micelles; therefore, one can suggest that the rise of $G_\beta$ with $\beta$ can be due to the increase of the total concentration of surfactants upon addition of C8TAB.

Effect of Polymer. Let us reveal the role of polymer in the evolution of rheological properties with decreasing charge density of micelles by comparing the data at different content of PVA and in the absence of PVA (Figure 7). First, one can note that in the presence of polymer the transition to semidilute regime corresponding to the onset of sharp rise of viscosity (Figure 7a) proceeds at lower $\beta$ (0.23 instead of 0.27 for polymer-free system), which fully agrees with SANS data (Figure 3). As was discussed above, this is due to longer micelles observed in polymer–surfactant system. Second, in the presence of polymer the formation of the network of entanglements reflected in the appearance of the plateau modulus (Figure 6 and Figure 8S) also occurs at smaller molar ratio $\beta$ (0.38 instead of 0.41 for polymer-free surfactants), that is, for more charged micelles. Therefore, polymer decreases the $\beta$ values corresponding to the transition from the unentangled to entangled semidilute regime. Third, one can state that at any $\beta$ value before the maximum polymer increases significantly the viscosity of the system (Figure 7a). For example, at $\beta = 0.4$ the zero-shear viscosity of polymer–surfactant system containing 0.89 monomol/L PVA is 617 Pa s, whereas for surfactant solution it equals only 5 Pa s and for PVA solution 0.005 Pa s. Therefore, the viscosity of the mixed polymer–surfactant system is by more than 2 orders of magnitude higher than the viscosity of its components taken separately. Such a pronounced synergistic effect can be attributed not only to higher length of micelles in the presence of PVA. As was shown recently, the entanglements between polymer and micellar chains can also contribute to enhanced viscosity. Note that the concentration of PVA in the system is near the overlap concentration $C^*$ (0.7 monomol/L) and well below the entanglement concentration $C_e$, which is usually 5–10 times higher than $C^*$. Therefore, the polymer–polymer entanglements can be neglected. Thus, an enhancement of viscosity in semidilute regime in the presence of polymer seems to be due to the elongation of micelles induced by PVA and to the formation of the entanglements between micellar and polymeric chains. For the same reason the polymer increases the terminal relaxation time and the plateau modulus (Figure 7) as well as the $\beta$ values corresponding to the transition from the unentangled to entangled semidilute regime. At the same time, the maximum values of viscosity and terminal relaxation time do not depend on the presence of PVA (Figure 7). Therefore, PVA helps to reach the maximum viscosity and relaxation time at smaller $\beta$, i.e., at higher degree of charging. Thus, the addition of nonionic polymer instead of an ionic compound (for instance, cationic surfactant) may help to obtain a solution of highly charged micelles with high rheological properties, whereas in the absence of polymer the micelles need to be less charged in order to have the same viscosity and elasticity.

Figure 7. Dependences of zero-shear viscosity $\eta_0$ (a), terminal relaxation time $\tau$ (b), and plateau storage modulus $G_\beta$ (c) on the molar ratio [C8TAB]/[potassium oleate] $\beta$ for 0.078 M aqueous solutions of potassium oleate containing C8TAB and different concentrations of PVA: 0 (blue triangles), 0.22 monomol/L (green diamonds), and 0.89 monomol/L (red circles). Dashed black line (a) indicates the zero-shear viscosity of 0.89 monomol/L PVA solution.
CONCLUSIONS

In this paper, we studied the behavior of the solutions of ionic surfactant micelles of variable charge and nonionic polymer PVA. To decrease the degree of charging of micelles, we gradually added a cationic surfactant C12TAB to oppositely charged potassium oleate micelles. We used a combination of rheological measurements exploring the macroscopic properties (viscosity, elasticity, etc.) of the system and SANS revealing the structural rearrangements occurring at the microscopic level. We show that decreasing charge of micelles promotes their elongation and consequently the enhancement of the rheological properties (zero-shear viscosity, terminal relaxation time, and plateau modulus) both in the absence and in the presence of polymer. At a given degree of charging of micelles, the polymer induces the formation of longer micellar chains and of polymer–micellar entanglements and therefore increases significantly the rheological characteristics. We explain the role of polymer in the elongation of micelles by its local segregation from surfactant, which reduces the volume of solution accessible for the micelles, thus increasing their local concentration and promoting their growth in length. This suggestion is confirmed by SANS data, which demonstrate (i) a shift of the structure peak position toward high q-values upon addition of polymer indicating the decrease of intermicellar distance and (ii) an enhancement of low-q scattering pointing to the formation of longer micelles and large-scale inhomogeneities.

The findings reported here demonstrate the effectiveness of the addition of weakly interacting polymer for the modulation of rheology of wormlike surfactant micelles, thus opening a new way for manipulating the rheological properties of viscoelastic surfactants.

ASSOCIATED CONTENT

Supporting Information

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NMR spectroscopy, SANS, and rheology data (PDF)

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Notes
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