Microstructure and Rheology of Lamellar Liquid Crystalline Phases

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We have investigated the microstructure and rheological properties of ternary surfactant mixtures in a salt solution. The surfactants were 6% sodium 4-dodecylbenzenesulfonate, 3% C_{13-15} ethoxylated alcohol with seven ethylene oxide (EO) units and 1% C_{13-15} ethoxylated alcohol with 2, 4, 7, 9, 11, 14, 20, or 25 EO units. The salt solution was 10% nitrilotriacetate-H_2O. Microstructural investigations (electron microscopy, light microscopy, confocal laser microscopy, conductivity measurements, and centrifugation) show that at rest the samples containing the surfactant with 2 EO to 9 EO units are dispersions of lamellar droplets (curved surfactant bilayers). The samples containing the surfactant with 11 EO to 25 EO units show a continuous lamellar structure (sheets of surfactant bilayers) with a small amount of lamellar droplets present. The change in several rheological parameters reflects this change in microstructure. The power law index from flow experiments at low shear rates changes from 0.1 for the lamellar dispersions to 0.4 for the continuous lamellar phases. Similar changes are observed in shear modulus and in the limiting strain for linear viscoelastic behavior. The continuous lamellar phase is converted to droplets by shearing at rates above 1 s^{-1}. The continuous lamellar structures will recover in about a week when the samples are allowed to relax. The nature of the droplets is highly dynamic. Confocal laser microscopy shows small fluctuations in droplet shape on a time scale of about 100 s. This time coincides with a characteristic time of around 100 s pertaining to a (shallow) peak in G''.

1. Introduction

Many (commercially available) surfactants used in detergent products have a tendency, when dissolved in water or a salt solution, to form liquid-crystalline lamellar phases.\(^1\) These lamellar phases consist of stacked bilayers of alternating layers of ordered surfactant molecules and solvent. The surfactants in the bilayers are organized in such a way that the hydrophobic tails of the surfactant molecules are at the center of the lamellae and the hydrophilic parts of the molecules are in contact with the solvent layer. A schematic representation of a lamellar phase is given in Figure 1. The disadvantage of this type of representation is that it is static. It does not do justice to the dynamic nature of these phases, with undulating planes and surfactant molecules migrating in and out of the bilayer.

Lamellar phases are known to exist in at least two configurations. In addition to a planar or continuous lamellar phase, where the bilayers are ordered in sheets,\(^2\) lamellar phases exist with the bilayers ordered in closed concentric shells.\(^3\)–\(^5\) The structural units in the latter phase are often referred to as “onions”, multilamellar vesicles, or lamellar droplets. From previous studies it appears that the onions are stable even at low surfactant concentrations when prepared from mixtures of nonionic and anionic surfactants at sufficiently high salt concentrations.\(^6\)

When the surfactant head groups are well hydrated and the alkyl layer is fluid, these phases can swell considerably, indicating repulsion between the layers. The spacing between the bilayers can be orders of magnitude larger than the typical dimensions of the hydrophilic surfactant head group.\(^7\) This is thought to be caused by thermal undulation of the layers. This swelling behavior can lead to a space-filling arrangement even at low to moderate surfactant concentrations. When, for onions,

\[ G'' \]

Figure 1. Schematic representation of surfactant molecules arranged in bilayers.
the volume fraction of lamellar phase ($\phi_{lm}$) is at or just above the random maximum packing of monodisperse spheres (0.63), it provides an excellent matrix for suspending solid particles, enabling the formulation of, e.g., fabric washing liquids.3,4

Onions and continuous lamellar phases have interesting rheological properties. It has, for instance, been reported that a continuous lamellar phase, prepared from AOT in brine solutions or from SDS, pentanol, water, and dodecane, can be converted to onions by shear9-11 which will relax back to continuous lamellar in a few days to several months, depending on the composition. Strong shear hysteresis in systems containing 5% AOT in water has been reported.12 The hysteresis is thought to be associated with a shear-induced phase transition.

We have studied the microstructure and rheological properties of a typical liquid detergent composition. A small change in the average ethylene oxide chain length of the nonionic surfactant caused transition of onions to a mainly continuous lamellar phase. The liquids exhibit striking non-Newtonian rheological behavior. They show elasticity at small deformations, possibly due to the deformability of the droplets and the interactions between the droplets. Furthermore, the liquids show marked shear thinning.

The research described in this paper was carried out in an industrial research laboratory, using industrial raw materials. Although the use of commercial grade materials may lead to a relatively large scatter in experimental results, the translation to industrial products and processes is as simple and direct as possible.

2. Experimental Section

2.1. Materials. The anionic surfactant used was Marlon AS-3, from Hüls, a commercial preparation of DoB-S acid, which is a mixture of 98.0-98.5% isomerically impure 4-(sec-dodecyl)-benzenesulfonic acid, about 1.5% non-sulfonated organic matter and 0.3% sulfuric acid. The nonionic surfactants were selected from the range of Synperonics from ICI which are C13-15 ethoxylated alcohols, with an average of 2, 4, 7, 9, 11, 14, and 20 moles of ethoxides per molecule. We also used C13-15 ethoxylated alcohol with 25 ethylene oxide monomer units on average (Ukain 102, from UGINE Kuhlmann). It should be stressed that the EO lengths given represent an average value and that the distribution in lengths is very broad.

The salt used was nitritroacetate $\text{HNO}_3$, NTA (Trilon A92, from BASF).

All materials were used without further purification.

Model samples were prepared by dissolving the surfactants in demineralized water under stirring at 50 °C, until a clear isotropic surfactant solution was obtained. As sufficient amount of sodium hydroxide was added to neutralize the acid surfactant. After addition of the NTA, the solution was allowed to cool to room temperature under stirring. Finally the pH of the solution was adjusted to around 8 by adding NaOH solution. The samples were stored for at least 1 month in closed plastic bottles at room temperature before any measurement was made.

The surfactant concentration in the samples was 10% by weight, with an anionic:nonionic weight ratio of 6:4 (based on synthetic conditions). The samples were centrifuged for 16 h at 40 000 $\text{g}$. The height of separated layers, if any, was measured with a ruler. This allows a rough calculation of the volume of the layers.

Dynamic rheological measurements were made with a Bohlin VOR constant shear rate rheometer equipped with a smooth couette geometry (C25, bob diameter 25 mm, gap 0.125 mm) in the linear viscoelastic region. The sample was carefully introduced into the cylindrical geometry, avoiding high rates of deformation as much as possible. Subsequently several frequency sweeps from 10–001 to 10 $\text{Hz}$ were done on the sample. After a maximum of four sweeps, no further systematic changes in $\text{G}'$ and $\text{G}''$ occurred, indicating that the sample had fully relaxed. Data from the fifth and sixth sweep were used. The strain was limited to 0.02, which is in the linear region.

Measurements of viscosity as function of shear rate were made on a Bohlin C5 rheometer, using a constant shear rate software package. The rheometer was equipped with a serrated concentric cylinder setup (bob diameter 25 mm, gap 0.125 mm, grooves of 0.1 mm) to prevent wall slip.3 The same precautions were taken as for the dynamic measurements, and the sample was allowed to relax before any measurements were made. Shear rates ranged from 0.065 to 328 s$^{-1}$ in 40 logarithmically spaced steps. The measuring time was 80 s at the lowest shear rate decreasing in proportional steps to 20 s at the highest shear rate. Two up-down curves were recorded (up and down refer to the direction of change of the shear rates).

For all samples, dynamic measurements were also performed after a certain shear rate had been imposed on the sample for 300 s. The shear rates were 0, 5.8×10$^{-1}$, 5.8×10$^{-1}$, 5.8×10$^{-1}$, 5.8×10$^{-1}$, and 367 s$^{-1}$, respectively. The measurements were performed on the Bohlin VOR. The dynamic moduli of two samples were measured at 1 Hz after each sample was sheared for 300 s at 328 s$^{-1}$ for an extended period of time (about 1 week).
the detection limit of the microscope when they are left to rest for several minutes. The appearance changes from birefringent with some lamellar droplets to platelike, with some lamellar droplets still present. This is the case for the samples 11EO and higher. For the samples 2EO to 9EO no change in appearance with time is observed. The light microscopic image for the sample with 14EO (top right in Figure 2) shows, next to some droplets, large areas of flat, platelike structures, extending over 100 µm or more. The edges of the plates are birefringent. They are either light or dark, depending on the orientation of the sample. When the orientation of the sample under the light microscope is changed, the light edges become dark and vice versa. This means that the observed plates must consist of multiple bilayers since the thickness of a single bilayer is too small (on the order of 10 nm) to be visible by light microscopy. The microstructural information obtained with the confocal laser microscope confirms to a large extent the view developed on the basis of the ordinary light microscopy. The pictures show a filled array of mainly spherical droplets for samples 2EO to 9EO and more extended structures for 11EO to 25EO. We interpret the apparent difference between these large structures under the light microscope and in confocal laser microscopy to be caused by the sample preparation method. In light microscopy, large structures are flattened between the glass surfaces of the slide and cover glass. Therefore, these structures will no longer show internal structure. Also some very large droplets with a typical size of a few micrometers are present. These droplets show Maltese crosses as seen in ordinary light microscopy, because of the radial symmetry present.

The electron micrographs for the samples with 2EO until 9EO confirm that these samples contain many, basically spherical, highly packed droplets. In the case of 4EO the high degree of packing leads to deformation of the droplets into a "honeycomb" like arrangement. Electron micrographs of this type of packing for nearly monodisperse lamellar droplets are shown elsewhere. In the samples 2EO, 7EO, and 9EO the deformation of the droplets is much less noted. We have no explanation for this difference in appearance. The electron micrograph of the sample with 14EO shows continuous lamellar platelike structures. These structures are not perfectly flat but show a curved substructure like in the confocal laser microscopy images. In several places on the photograph the individual surfactant bilayers are visible. It is interesting to see that the continuous lamellar layer terminates at the edge by closure of not more than a few bilayers, showing locally a high curvature. Close examination of all the micrographs reveals that the systems with surfactant EO chain length of 9 and smaller are dispersions of lamellar droplets. For EO chain lengths equal to 11 and larger, the structure changes quite abruptly to primarily continuous lamellar with an occasional droplet.

We also used the confocal laser microscope to record a series of images taken with a constant time interval. When such a sequence, which extended over a period of more than one hour, is played back in a few seconds, one can clearly see the dynamic nature of the samples containing only droplets. The individual droplets show continuous shape fluctuations. The eye is sensitive enough to detect these changes in shape when the pictures are put together in a movie, but it is impossible to see this on static photographs. We estimate the magnitude of the fluctua-

Figure 2. From top to bottom, polarized light (A), confocal laser (B), and electron microscopic (C) pictures of relevant structures observed in 4EO (left column) and 14EO (right column) samples.

The samples were inspected microscopically after shearing at 367 s⁻¹ for 300 s to examine the effect of shear on the microstructure.

3. Results and Discussion

For clarity only a limited number of photographs and curves are presented. The observations made are well represented by the samples 4EO and 14EO (abbreviations explained in section 2.1). We will, however, present results for all samples where appropriate.

3.1. Microstructure. Typical examples of microstructures are given in Figure 2. On the left-hand side, representative images for the sample with 4EO are given, and on the right-hand side, images for the sample with 14EO can be found. The top, middle, and bottom pictures were obtained with light microscopy, confocal laser microscopy, and electron microscopy, respectively.

Polarized light microscopy shows that all samples contain some droplet structures with Maltese extinction crosses. This is a clear indication of the presence of multilamellar droplets. The droplet sizes are above 0.5 µm because they were observed in optical microscopy with a detection limit of around 0.5 µm because of the wavelength of the light used. In these droplets, alternating surfactant–water bilayers are present as concentric spherical shells. Also some birefringence was detected, especially when the sample flows around an entrapped air bubble. This birefringence is associated with the existence of small lamellar droplets, just below the resolution limit of the microscope. The dark and light streaks present in the top left image in Figure 2 indicate the presence of small birefringent structures, just below the detection limit of the microscope.

Some samples show a drastic change in microstructure when they are left to rest for several minutes. The appearance changes from birefringent with some lamellar droplets to platelike, with some lamellar droplets still present. This is the case for the samples 11EO and higher. For the samples 2EO to 9EO no change in appearance with time is observed. The light microscopic image for the sample with 14EO (top right in Figure 2) shows, next to some droplets, large areas of flat, platelike structures, extending over 100 µm or more. The edges of the plates are birefringent. They are either light or dark, depending on the orientation of the sample. When the orientation of the sample under the light microscope is changed, the light edges become dark and vice versa. This means that the observed plates must consist of multiple bilayers since the thickness of a single bilayer is too small (on the order of 10 nm) to be visible by light microscopy. The microstructural information obtained with the confocal laser microscope confirms to a large extent the view developed on the basis of the ordinary light microscopy. The pictures show a filled array of mainly spherical droplets for samples 2EO to 9EO and more extended structures for 11EO to 25EO. We interpret the apparent difference between these large structures under the light microscope and in confocal laser microscopy to be caused by the sample preparation method. In light microscopy, large structures are flattened between the glass surfaces of the slide and cover glass. Therefore, these structures will no longer show internal structure. Also some very large droplets with a typical size of a few micrometers are present. These droplets show Maltese crosses as seen in ordinary light microscopy, because of the radial symmetry present.

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(13) Gulik-Krzywicki, T; Dedieu, J. C.; Roux, D; Degert, C; Laverson, R. Langmuir 1996, 12, 4668.
tions to be, at maximum, about 5% of the size of a typical droplet. On one occasion, for sample 4EO, we observed that the shape fluctuations lead to the breakup of a large droplet into two smaller droplets.

From the electrical conductivity, the volume fraction lamellar phase was calculated using the Bruggeman equation adapted for lamellar dispersions

\[
\left( \frac{\kappa - \kappa_{\text{el}}}{\kappa - \kappa_{\text{lam}}} \right) \left( \frac{\kappa_{\text{el}}}{\kappa_{\text{lam}}} \right)^{(1/3)} = 1 - \phi_{\text{lam}}
\]

in which \( \kappa \) denotes the electrical conductivity of the total system, \( \phi_{\text{lam}} \) is the volume fraction lamellar phase and the subscripts lam and el refer to the dispersed lamellar droplets and the continuous electrolyte solution respectively. For the conductivity of the dispersed lamellar droplets a value of 0.8 mS/cm was used whereas for the continuous salt solution, the conductivity of a salt solution prepared without the surfactants was used.

The volume fraction lamellar phase can also be estimated from the volume of the clear layer that forms after centrifugation. We have assumed that, unlike the case of dispersions of solid particles, the droplets in our systems are deformable enough to facilitate packing to a volume fraction of near one. In some cases, for samples 14EO to 25EO, a very thin isotropic top layer was detected. The layers had a slight yellow color, which is an indication of the presence of a small amount of surfactant in this layer. The original samples showed no such a layer, even after being allowed to stand for more than 6 months at room temperature. The presence of surfactant in the top layer is therefore most likely a result of the high g-force during centrifugation. The results of both methods are listed in Table 1.

Previously it was found that for onion phases the difference in volume fractions between centrifugation and conductivity is at most 10%.\(^3\) The assumption that the lamellar droplets can pack to a volume fraction of 1 may be too optimistic. The maximum attainable packing is probably lower. Despite this, there still exists a rather large discrepancy between the two results. The observed change in microstructure probably explains to a large extent the deviation between the volume fractions calculated from conductivity and from centrifugation. For the calculation of the volume fraction of the dispersed lamellar phase from conductivity, one assumes this phase to be present as lamellar droplets with low conductivity. As soon as the continuous lamellar phase is present, this is no longer the case. When the continuous lamellar layers are positioned perpendicular with respect to the conductivity electrodes, the dispersed phase will have a conductivity that is much higher, probably on the order of the conductivity of the continuous salt solution. The same line of reasoning was followed elsewhere.\(^1\)

We therefore conclude that the volume fraction calculated from conductivity is at best indicative for the volume of lamellar droplets, whereas the volume fraction from centrifugation is the sum of both continuous and droplet lamellar phases.

3.2. Rheology. 3.2.1. Linear Experiments. Two typical examples of dynamic moduli (\( G' \) and \( G'' \)) as function of frequency are given in Figure 3. Again the 4EO data are representative in magnitude and shape of the data for samples 2EO until 9EO whereas 14EO is representative for 11EO until 25EO. Successive runs with fresh samples and new torsion bars give a standard deviation for \( G' \) of 8% and for \( G'' \) of 15%. The larger error in \( G'' \) is inherent to the measuring method when \( G' \) is much larger than \( G'' \), as is the case here.

For the samples 2EO to 9EO one can clearly detect the onset of a broad relaxation transition at low frequencies. The position of this transition shifts to higher frequencies with increasing EO length. From the slope of \( G'' \) as function of frequency after the transition, it can be deduced that this transition is not the result of a single relaxation mechanism, with a single characteristic time. In that case one would expect \( G'' \) to decrease, in a double log plot with a slope of \(-1\). For the sample with 4EO, the characteristic time at which the transition occurs is of the order of 100 s and coincides with the times associated with the fluctuations of the droplet shape as observed with confocal light microscopy. For samples 11EO until 25EO, with a prevailing continuous lamellar phase, this transition is absent. For all samples, the onset of a relaxation process at high frequencies is observed.

The data can be reduced by fitting to a linear viscoelastic model with

\[
G' = G_0 + \sum_{p=1}^{i} \frac{G_p \omega^2 \tau_p^2}{1 + (\omega \tau_p)^2}
\]

and

\[
G'' = \sum_{p=1}^{i} \frac{G_p \omega \tau_p}{1 + (\omega \tau_p)^2}
\]

An apparent plateau in \( G' \) for \( \omega \to 0 \) (\( G_0 \)) was included for curve-fitting purposes although eventually for very low frequencies one expects \( G' \) to drop again with decreasing frequency.
Satisfactory fits were found with \(i = 7\) and with the longest and shortest relaxation times \(r_1\) and \(r_2\) equal to the inverse of the highest and lowest radial frequency used. The remaining \(r_p\) values were logarithmically divided between \(r_1\) and \(r_2\). This leaves eight parameters \(G_0\) to \(G_7\) to be fitted.\(^{15}\) Although in principle an exact solution should be obtainable, since all free parameters are linear, this kind of fit is known to be "ill posed,"\(^{16}\) with many possible solutions. We therefore used a nonlinear curve-fit procedure in which we minimized the logarithmic least-squares sum of \(G''\) and \(G'\). By minimization of the squared sum of the logarithms, no weighing of the data is required to obtain a good fit. Only positive parameter values were allowed. The drawn lines in Figure 3 were calculated with the obtained parameters. The result of the curve-fit is given in Table 2. Note that \(G'\) at the shortest relaxation time \((G_1)\) is almost equal for all samples and does not depend on the morphology of the lamellar phase.

The observed step change in microstructure is clearly shown in the measured dynamic responses. In Figure 4 the change in \(G_0\) and the change in the limiting strain for linear viscoelastic behavior is given as function of EO chain length. Note that an identical step change is seen in \(G_0\) to \(G_7\) in Table 2, indicating that the change is especially noticeable at long times (low frequencies).

The explanation of the magnitude of \(G'\) at high frequencies is not yet clear.\(^{17,18}\) The influence of structure, prestrain, and timescale has to be taken into account, but this has not (yet) been figured out in a complete theory. For the time being, three still relevant estimations can be given. If \(G'\) is determined by the relaxation of multilamellar spherical droplets\(^{17}\)

\[
G' \propto \frac{\sigma_{\text{eff}}}{R} \tag{4}
\]

where \(\sigma_{\text{eff}}\) is the effective surface tension and \(R\) is the radius of the spheres. Here \(\sigma_{\text{eff}} \approx 10^{-5}\) N/m\(^{17}\) and \(R \approx 500\) nm; thus \(G' \approx 20\) Pa, close to the value found for 2EO to 9EO and 11EO to 25EO after shear.

If \(G'\) is determined by interactions between the lamellae, then, in affine deformation, \(G' \sim R\), where \(B\) is the compressibility modulus. Since \(B\) is on order of 1000 Pa,\(^{17,18}\) this much larger than the measured value of \(G'\). The last explanation assumes a prestrain \(\gamma_0\) for asymmetric onions, leading to \(G' \sim B\gamma_0^2\).\(^{18}\) Since \(\gamma_0\) is unknown, this can always be brought into agreement with experimental results.

### 3.2.2. Nonlinear Experiments

Typical examples of viscosity as function of shear rate are given in Figure 5. The curve for sample 4EO is representative for samples up to 9EO. The samples 11EO to 25EO behave as 14EO does. The samples from 2EO to 9EO show no difference between the up-curve and the down-curve, indicating that, in the range of shear rates examined, no detectable, persisting changes in microstructure occur. The samples with 11EO up to 25EO show a noticeable shear/time history. The first up-curve is below the subsequent down-and up-curves, indicating a detectable change in microstructure due to the shear history.

All samples are shear-thinning, with a tendency towards a Newtonian plateau at high shear rates. The data was fitted to the Sisko equation with a nonlinear curve-fit procedure. The Sisko equation is a simplified version of the general Cross\(^{19}\) flow model:

\[
\eta = \eta_\infty + \frac{\eta_0 - \eta_\infty}{1 + \alpha_\gamma^{(1-n)}} \tag{5}
\]

Since we could not detect a first Newtonian plateau \((\eta_0)\), the equation can be simplified. For \(\alpha_\gamma^{(1-n)} \gg 1\), eq 5 reduces to

\[
\eta = \eta_\infty + K_\gamma^{(n-1)} \tag{6}
\]

The parameters in this equation are the viscosity at the high shear rate Newtonian plateau, \(\eta_\infty\), the consistency, \(K\) (/Pa-s\(^n\)) and the power law index \(n\) (/). The drawn lines in Figure 5 are calculated from the fitted parameters. The results of the curve-fit are given in Tables


\(^{17}\) van der Linden, E.: Drogé, J. H. M. Physica A 1993, 193, 439.


\(^{19}\) Cross, M. M. J. Colloid Sci. 1965, 20, 417.
Table 3. Parameters from Curve Fit to Sisko Model, First Flow Curve, Starting at Low Shear Rate (Standard Deviation Given in Parentheses)

<table>
<thead>
<tr>
<th>sample</th>
<th>η₀/Pa·s</th>
<th>K/Pa·s²</th>
<th>power law index, n</th>
</tr>
</thead>
<tbody>
<tr>
<td>2EO</td>
<td>0.061 (0.001)</td>
<td>2.18 (0.014)</td>
<td>0.096 (0.004)</td>
</tr>
<tr>
<td>4EO</td>
<td>0.046 (0.001)</td>
<td>1.65 (0.011)</td>
<td>0.145 (0.004)</td>
</tr>
<tr>
<td>7EO</td>
<td>0.029 (0.001)</td>
<td>1.87 (0.016)</td>
<td>0.144 (0.005)</td>
</tr>
<tr>
<td>9EO</td>
<td>0.030 (0.001)</td>
<td>1.57 (0.014)</td>
<td>0.165 (0.005)</td>
</tr>
<tr>
<td>11EO</td>
<td>0.017 (0.002)</td>
<td>0.74 (0.008)</td>
<td>0.403 (0.006)</td>
</tr>
<tr>
<td>14EO</td>
<td>0.013 (0.002)</td>
<td>0.68 (0.007)</td>
<td>0.431 (0.006)</td>
</tr>
<tr>
<td>20EO</td>
<td>0.018 (0.001)</td>
<td>0.60 (0.005)</td>
<td>0.367 (0.005)</td>
</tr>
<tr>
<td>25EO</td>
<td>0.016 (0.001)</td>
<td>0.53 (0.005)</td>
<td>0.396 (0.005)</td>
</tr>
</tbody>
</table>

Table 4. Parameters from Curve Fit to Sisko Model, Fourth Flow Curve, Starting at High Shear Rate (Standard Deviation Given in Parentheses)

<table>
<thead>
<tr>
<th>sample</th>
<th>η₀/Pa·s</th>
<th>K/Pa·s²</th>
<th>power law index, n</th>
</tr>
</thead>
<tbody>
<tr>
<td>2EO</td>
<td>0.057 (0.001)</td>
<td>2.31 (0.013)</td>
<td>0.123 (0.004)</td>
</tr>
<tr>
<td>4EO</td>
<td>0.040 (0.001)</td>
<td>2.07 (0.014)</td>
<td>0.147 (0.004)</td>
</tr>
<tr>
<td>7EO</td>
<td>0.028 (0.001)</td>
<td>1.95 (0.016)</td>
<td>0.140 (0.005)</td>
</tr>
<tr>
<td>9EO</td>
<td>0.029 (0.001)</td>
<td>1.65 (0.015)</td>
<td>0.151 (0.005)</td>
</tr>
<tr>
<td>11EO</td>
<td>0.031 (0.001)</td>
<td>1.38 (0.014)</td>
<td>0.184 (0.006)</td>
</tr>
<tr>
<td>14EO</td>
<td>0.029 (0.001)</td>
<td>1.19 (0.013)</td>
<td>0.202 (0.007)</td>
</tr>
<tr>
<td>20EO</td>
<td>0.024 (0.001)</td>
<td>0.89 (0.010)</td>
<td>0.221 (0.007)</td>
</tr>
<tr>
<td>25EO</td>
<td>0.023 (0.001)</td>
<td>0.85 (0.010)</td>
<td>0.223 (0.007)</td>
</tr>
</tbody>
</table>

3 (up-curves) and 4 (down-curves). Both η₀ and K decrease with increasing EO chain length up to 11EO. For 14EO to 25EO these values appear to reach a plateau. For samples 2EO to 9EO no large differences are found in the curve-fit parameters between the up- and down-curves. For samples 11EO to 25EO, both η₀ and K are larger after shear. The power law index n is smaller and reduces due to shear from n-values of about 0.4 to 0.2. The power law index as function of EO chain length for an unsheared and a sheared sample is given in Figure 6.

For multilamellar vesicles, Hoffmann et al. found that the yield stress (σ₀) is about 10 times smaller than G'. We found a comparable ratio. The significance of this comparison between a linear and nonlinear quantity comes from the following. The modulus at high frequency, multiplied with the limiting strain for viscoelastic behavior (γₘₐₓ), gives an estimate for the stress at which plastic flow begins. We have compared this quantity (from a plot of G' at 1 Hz as function of strain) to the dynamic yield stress (σ₀) deduced from the flow curves at low rates of shear. The estimate of the dynamic yield stress is only significant when the slope of the flow curve in a double logarithmic plot approaches −1. This is the case for samples 2EO to 9EO. The result of this comparison is given in Figure 7. This figure shows that both quantities are in fair agreement with each other for a given sample, as one would expect if the plastic flow to flow transition takes place via one type of structural change.

Microscopic observation after shear shows that the samples with continuous lamellar phase undergo a transition to droplets. For samples 2EO to 9EO, no change in microstructure could be detected. The microscopic appearance after shear for all samples is well represented by the top left image in Figure 2.

3.2.3. Linear after Nonlinear Experiments. We have further investigated this change in microstructure through shear by shearing at a predefined rate for 300 s followed by a frequency sweep. During preshearing the viscosity of the sample was recorded every 15 s. Even at the lowest shear rate, the viscosity reading was constant after at most 45 s. This indicates that the change in morphology of the lamellar phase due to shear is completed within this time span and occurs quite rapidly. G' at 0.01 Hz, after the cessation of a given shear rate, is plotted in Figure 8 for two samples. Since we anticipate that the change in G' reflects changes in microstructure, we deduce from this graph that there is no large change in the microstructure of the 4EO sample (representative for samples 2EO to 9EO). The microstructure of 14EO (representative for samples 11EO to 25EO) starts to change already at shear rates as low as 1 s⁻¹. The observed initial decrease in G' for 14EO is probably caused by incomplete structure relaxation before the start of the experiment or by alignment of the lamellar phase due to the imposed shear. Note that G' after shear is similar for all samples because of the comparable microstructure of the samples after shear.

We also examined the relaxation of the structure after shear (for 300 s at 367 s⁻¹) by measuring G′ at 1 Hz for an extended period of time. The results are given in Figure 9, as G′/G′₀(at rest). For 4EO we see some initial relaxation possibly caused by the rearrangement of the droplet structure. The sample with 14EO relaxes back to the initial level in about 1 week.

The picture that emerges is that when the lamellar phase is primarily present as lamellar droplets, the power law index is between 0.1 and 0.2. When the structure changes to plate-like continuous lamellar, the power law index increases to 0.35 or 0.4. After shear, the power law index of the latter samples becomes approximately 0.2 due to a reversible change in microstructure from continuous lamellar to lamellar droplets.

4. Conclusions

Lamellar phases, prepared from anionic and nonionic surfactants, can be quite insensitive to small changes in the average EO chain length of the nonionic surfactant used. When to the base composition, containing 6% anionic, 3% 7 EO nonionic, and 10% NTA, was added 1% of EO nonionic with an average EO chain length of 2, 4, 7, or 9, equal lamellar phase morphology and comparable rheological properties were found. The lamellar phase is present as a dispersion of multilamellar droplets (onions). When to the base mixture was added 1% nonionic with an average EO chain length of 11, 14, 20, or 25, mainly continuous lamellar phases with again similar morphology and rheological properties are found.

A drastic and abrupt change in lamellar phase morphology and in rheological properties occurred on going from a 9EO nonionic to an 11EO nonionic surfactant. The change in the limiting strain for linear viscoelastic behavior occurs already at lower EO chain lengths. We have no explanation for this. The change in chemical composition by exchanging 1% of the 9EO nonionic surfactant for 1% of the 11EO nonionic surfactant is very small. It merely means that the distribution of EO chain lengths is shifted marginally. Despite this small change, the morphology changes from dispersed lamellar to primarily continuous lamellar. And as a consequence of that also the linear and nonlinear rheological properties changed considerably. Continuous lamellar phase can be converted to droplets by shear at rates above 1 s⁻¹. The droplets will relax back in about 1 week. Shape fluctuations of lamellar droplets, which were seen using confocal laser microscopy were associated with a relaxation process detected in the measured dynamic moduli.

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