Effect of Polymer Architecture on the Phase Behavior and Structure of Polyelectrolyte/Microemulsion Complexes (PEMECs)

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ABSTRACT: We studied polyelectrolyte/microemulsion complexes (PEMECs) formed by the biopolymer sodium hyaluronate (NaHA) and cationic oil-in-water (O/W) microemulsion droplets. Around equimolar charge conditions, a two-phase region with a liquid−liquid phase separation (coacervate formation) is observed, while mixed complexes are formed at polyelectrolyte excess. The largest complexes are found close to the phase boundary. The detailed structure of these complexes was determined by a combination of static and dynamic light scattering (SLS and DLS), small-angle neutron scattering (SANS), and cryo-transmission electron cryomicroscopy (cryo-TEM). Interestingly, these complexes formed with NaHA are much more elongated compared to previously studied complexes formed with sodium polyacrylate (NaPA). Similar observations were made for another polysaccharide, sodium carboxymethyl cellulose (NaCMC). Apparently, the persistence length of the complexes is largely proportional to the persistence length of the polyelectrolyte, which was determined by analyzing the SANS data with a specifically developed model. Accordingly, the size of the complexes formed with stiff biopolyelectrolytes becomes much larger, increasing with the molecular weight ($M_w$) of the polyelectrolyte. In summary, we conclude that the size and structure of the PEMECs can be controlled by the type of polyelectrolyte, mixing ratio, size of the microemulsion droplets, and $M_w$ of the polyelectrolyte. This means that the polyelectrolyte addition can be used as a key structuring element for the size and shape of such complexes.

INTRODUCTION

Microemulsions classically come in the structural form of oil-in-water (O/W), or water-in-oil (W/O) droplets, or as bicontinuous structures.1,2 These structures are stabilized by the presence of a surfactant (and often also a cosurfactant3,4). Their structural units are typically in the size range of 2–20 nm, which explains their optical transparency. O/W microemulsion systems are particularly promising because of the high solubilization capacity for hydrophobic molecules in aqueous solution, which plays a key role in delivery systems, detergency, decontamination, etc.5–7

Microemulsion droplet systems have been studied to some extent. However, for some applications, one might be interested in modifying their properties and/or superstructures for further purposes. A classical case is the viscosity control, as microemulsions are generally low-viscous liquids, but for many applications, higher viscosities are required. This question is usually addressed by either enhancing the viscosity of the continuous phase8 or cross-linking individual droplets by a polymer with a number of hydrophobic linkers.9,10

Another interesting colloidal system constitutes polyelectrolyte/surfactant complexes (PESCs), which are formed by the ionic assembly of oppositely charged surfactant and polyelectrolyte.11 It has been shown that the electrostatic interactions of charged surfactants with oppositely charged polyelectrolytes can induce a pronounced viscosity increase for certain compositions.12 Often PESCs are studied in the context
of drug delivery,\textsuperscript{13,14} or cosmetical and detergency formulations.\textsuperscript{15} However, such formulations and applications require good solubilization properties, and that is often a major difficulty in PESCs\textsuperscript{16} and accordingly has hampered their further development in this direction. Yet, this problem can become circumvented by starting with ionic O/W microemulsion droplets to be complexed with an oppositely charged polyelectrolyte. Such a system combines the high solubilization capacity of the microemulsion with the mesoscopic structuring effect of the polyelectrolyte.

Recently, a detailed study of the phase behavior and structure of complexes of O/W microemulsion droplets with polyacrylate (NaPA) has been carried out by some of us.\textsuperscript{17} It was observed that such mixtures form self-assembled complexes with elongated structures, where the detailed structure depends markedly on the weight average molecular weight, $M_w$, of the polyelectrolyte, as well as on the size of the microemulsion droplets. These interesting findings can serve as a basis for further fundamental understanding of the assembly conditions. To develop more biofriendly formulations, it is important to explore how a change from the petrol-based and non-biodegradable polyacrylate to a biopolyanion affects the structures formed in polyelectrolyte/microemulsion complexes (PEMECs). In particular, the type of polyelectrolyte (persistence length, charge density, strength of interaction) can be expected to play a key role in the structures formed in these complexes since it is also an important factor in PESCs.\textsuperscript{18}

The development of biofriendly materials is naturally important for applications in biomedicine, food modification, or wastewater treatment, where biocompatibility is of crucial importance.\textsuperscript{19} However, on a more general note, sustainable materials derived from renewable resources are generally needed in all industry sectors to ensure our high standard of living and the well-being of our planet in the future.

To elucidate this question, we studied complexes of the previously investigated O/W microemulsion droplets\textsuperscript{1,24} composed of tetradecyltrimethylammonium oxide (TDMAO), tetradecyl-trimethyl-ammonium bromide (TTAB), 1-hexanol as cosurfactant, and decane as oil, where the formerly employed sodium polyacrylate (NaPA) was substituted by the biopolyanion sodium hyaluronate (NaHA). Both polyelectrolytes differ markedly with respect to their molecular architecture (see Figure 1). Hyaluronic acid is a biopolyelectrolyte with a sugar backbone, i.e., a polymer of disaccharides of D-glucuronic acid and N-acetyl-D-glucosamine, which leads to a much higher persistence length of the NaHA (9 nm) at an ionic strength of 30 mm$^2$\textsuperscript{25} and about twice the value for low ionic strength\textsuperscript{26} compared to NaPA (1.3 nm).\textsuperscript{27} In both cases, the charge is located close to the backbone, but the charge density of the hyaluronate is much lower compared to $\sim 1$ charge/nm) compared to NaPA ($\sim 4$ charges/nm). At the same time, the charged carboxylate group of the NaHA is less accessible to interact strongly with macromons due to its more restricted molecular mobility by being fixed on the polymer backbone. Therefore additional experiments were carried out with the sodium salt of carboxymethyl cellulose (NaCMC), a biopolyelectrolyte with a rigid polysaccharide backbone (6–16 nm persistence length\textsuperscript{28}), but sterically more easily accessible charges (see Figure 1).

In our investigation, we focused on the phase behavior and the mesoscopic structure of the PEMECs formed with hyaluronate at neutral pH ($7 \pm 8$). The main parameters were the mixing ratio of microemulsion droplets and polyelectrolyte, the $M_w$ of the hyaluronate, and the size of the charged microemulsion droplets. The aim has been to deduce systematic correlations between the PEMEC properties and their composition, as well as the comparison to the behavior with NaCMC and with NaPA as polyanions.

### MATERIALS AND METHODS

**Materials.** The surfactants used in the experiments were tetradecyltrimethylammonium oxide (TDMAO), received as a gift from Stepan Company, as a 25% TDMAO solution in water, named Ammonyx M (the solution was freeze-dried before use), and tetradecyltrimethylammonium bromide (TTAB, 99%, Sigma-Aldrich), which was used without further purification. 1-Hexanol (>98%, Merck Schuchardt OHG) was employed as cosurfactant and decane (>98%, Fluka Chemika) as oil to form the microemulsions. The microemulsion droplets were prepared at a constant total surfactant concentration of 100 mM (95 mM TDMAO + 5 mM TTAB). 0, 50, or 75 mM of the cosurfactant hexanol were added, and the droplets were saturated with decane (which means 30, 80, and 200 mM decane, respectively, leading to dispersed microemulsion volume fractions of 3.0, 4.4, and 6.5%). This means that the size of the microemulsion droplets was controlled by the concentration of the cosurfactant.

In this work, we primarily employed biopolyelectrolytes, in particular, the sodium salt of hyaluronic acid (NaHA), which was used in different molecular weights: 51, 150, 360 (all Lifecore), and 800 kDa (SRD). The sodium salts of poly-acrylic acid (NaPA, 60 kDa, Sigma-Aldrich) and carboxymethyl cellulose (NaCMC, 90 and 250 kDa, Sigma-Aldrich) were employed for a generalizing comparison of polyelectrolytes (the $M_w$ values always include the Na$^+$ counterion). All polyelectrolytes were used as received without further purification. The contour lengths (theoretical lengths of the stretched polyelectrolyte chains) are 130, 370, 900, and 2000 nm for the NaHAs1, 150, 360, and 800, respectively, 160 nm for NaPA60, and 200 and 480 nm for NaCMC90 and NaCMC250, respectively. Additional viscosity measurements were performed with NaHAs of 31, 186, 572, and 2073 kDa (all laboratory grade from Contipro, used as received).

Microemulsion droplets and polyelectrolytes were mixed in different charge ratios $z = \frac{-\left[+\right]}{\left[-\right]}$ (where $+\left[\right]$ is given by the TTAB units and $-\left[\right]$ is given by the carboxylate units) while maintaining a constant microemulsion concentration of 100 mM. Samples were named according to composition: the first block indicates the used microemulsion by giving the amount of hexanol, the second block indicates the used polyelectrolyte and the mixing ratio $z$ is given in the end. For example, ME50-NaHA150-0.70 is a sample with medium-sized droplets (50 mM of hexanol) and sodium hyaluronate of 150 kDa, mixed at a charge ratio of $z = 0.70$. The...
employed polyelectrolyte concentrations are above the overlap concentration (calculated for fully extended polymer chains) for all biopolyelectrolyte samples and most NaPA samples. The only exceptions are samples with short NaPA chains and small z ratios (the overlap concentration for NaPA05 is reached at \( z = 0.9 \), and for NaPA15 at \( z = 0.5 \)).

All samples were prepared in Milli-Q water or D_{2}O (>99.5% D, Eurisotop) for the SANS experiments.

### METHODS

#### Static (SLS) and Dynamic (DLS) Light Scattering.

SLS and DLS measurements were performed at TU Berlin simultaneously on an ALV/CGS-3 instrument, with a He–Ne laser with a wavelength of \( \lambda = 632.8 \) nm. Pseudo-cross-correlation functions for DLS were recorded using an ALV 5000/E multiple-goniometer. Diffusion coefficients \( D \) were obtained by plotting the relaxation rates \( \Gamma \) of all measured angles according to \( D = \Gamma / q^{2} \). The SLS data were recorded in the same range of magnitude of the scattering vector \( q \), with

\[
q = \frac{4 m \nu_{s} \sin(\theta/2)}{\lambda}
\]

where \( \eta_{s} \) is the refractive index of the solution and \( \theta \) is the scattering angle. All measurements were carried out at \( 25.0 \pm 0.1 \) °C in a thermostated toluene bath.

For SLS, the mean scattered intensity of the sample at each angle was normalized by the initial laser intensity, the background scattering to the density of the fluid \( \rho \), and \( B \) is a capillary constant. The flow time was averaged from at least three consecutive measurements.

#### Small-Angle Neutron Scattering (SANS).

SANS experiments were carried out on the PAXY instrument of the Laboratoire Léon Brillouin (ILL) in Saclay, France. Measurements were performed at 25 °C at three different configurations with sample-to-detector distances of 1.2, 5, and 6.6 m; collimation lengths of 2, 5, and 6 m; and wavelengths of 4, 4, and 12 Å, respectively, to cover a \( q \)-range of 0.03–5.4 nm\(^{-1}\).

Data reduction was done using the BerSANS software,\(^{30}\) according to the standard procedure.\(^{31}\) The raw intensity data were corrected for the scattering of the background (solvent and sample container), weighted by the respective transmissions. Additionally, the (electronic) background noise was subtracted using a cadmium sample, which absorbs incoming neutrons. The normalization and absolute scaling were done using a 1 mm reference sample of deionized water as an incoherent scatterer. Finally, the two-dimensional (2D) data were radially averaged and converted to one-dimensional (1D) scattering intensities.

Some additional measurements (ME75-NaPA samples in Figure 11) were carried out at D33 at Institut Laue-Langevin (ILL) in Grenoble, France, at sample-to-detector distances of 2(1.2), 7.8, and 12 m; collimation lengths of 2.8, 7.8, and 12 m; and wavelengths of 4.6, 4.6, and 13 Å, respectively, to cover a \( q \)-range of 0.013–61 nm\(^{-1}\). D33 has an additional detector at 1.2 m to measure the large \( q \) simultaneously to the 2 m setting. Data reduction for these data was carried out with LAMP.\(^{32}\)

#### Cryo-TEM.

Cryo-TEM images of selected samples were taken at Technion, Haifa. A small droplet of the solution was placed on a perforated carbon film supported on a TEM copper grid, held by tweezers in a controlled environment vitrification system at 100% relative humidity and 25 °C. It was then blotted by a piece of filter paper, resulting in the formation of 100–300 nm thin films of spanning the holes of a perforated carbon film.\(^{34}\) The specimen was then plunged into a reservoir of freezing liquid ethane, cooled by liquid nitrogen, to ensure its vitriﬁcation. The vitriﬁed specimen was transferred under liquid nitrogen, mounted onto a Gatan 626 cryogenic sample holder, and cooled to –170 °C. All samples were observed under low-dose conditions with an FEI Talos 200C TEM. Images were recorded using novel "phase plates," which convert phase differences between areas of the specimen to amplitude differences, thus enhancing the image contrast without resolution loss.\(^{33}\) The TEM is equipped with a Falcon III direct-imaging camera that allows imaging at very low electron exposure, essential for electron-sensitive specimens such as soft-matter samples. About 20 images were taken from each sample; the images were taken at different sample areas to ensure statistically relevant information.

### RESULTS AND DISCUSSION

#### Phase Behavior of ME/NaHA Complexes.

As a first step, the macroscopic phase behavior of the formed complexes was
studied as a function of the nominal charge ratio $z = [-]/([+] + [-])$ (nominal means that [+] is given by the TTAB units and [-] is given by the carboxylate units) for different molecular weights, $M_w$ of NaHA at room temperature. The pH was checked to be always between 7 and 8, but not controlled by adding acid or base, to avoid adding ions to the mixture. Within this pH range, the acids are largely deprotonated and $z$ should correspond to the actual charge ratio. The phase behavior of the samples was followed by visual inspection at regular time intervals, and documented by photographs, where the most instructive ones are given in Figure S1, nicely showing the phase separation occurring around equimolarity. Samples that phase-separated within an hour were marked as biphasic, and samples that were stable for more than an hour, but not for a week, were called metastable. All samples that still consist of one single phase after 1 week were considered thermodynamically stable, as they also did not change visually during the course of several months.

Figure 2 (left) shows the phase diagram for small microemulsion droplets (ME00, $R = 3.1$ nm) mixed with sodium hyaluronate (NaHA) of different $M_w$ values and for different charge ratios. In this phase diagram, a very asymmetric two-phase region can be observed around the charge equilibrium ($z = 0.5$). While samples at polyelectrolyte excess show one clear and transparent phase, samples in the two-phase region contain a second optically transparent liquid phase that floats on top of a low-viscous water-rich phase, i.e., a coacervate phase of lower density is formed. When the sample is shaken, the two phases mix and the sample appears white. Upon resting the sample, the two phases phase-separate again, typically within a few hours (see Figure S1 for photos of such samples). It is interesting to note that the coacervate phase, floating on top of the other, is very viscous. This is quite different from the previously studied PEMECs with NaPA, where a powdery precipitate was formed, and no viscous coacervate phase was observed. The phase diagrams for larger microemulsion droplets (ME50, $R = 4.1$; ME75, $R = 6.6$ nm) mixed with NaHA are given in Figure S2, which generally show the same features. The single-phase regions at polyelectrolyte excess become somewhat smaller with increasing droplet size, as previously observed for NaPA.

For a direct comparison, the phase diagrams of NaPA and NaCMC with small ME droplets are shown in Figure 2. Even though hyaluronate and polyacrylate are very different polyelectrolytes, a surprisingly similar phase behavior of mixtures with O/W microemulsion droplets is observed (phase diagrams of NaPA with larger droplets are given in Figure S3 for comparison). In both cases, aggregates in the microemulsion charge excess part of the phase diagram are very unstable (precipitation or liquid–liquid phase separation takes place), while long-time stable samples are found at polyelectrolyte excess. It is interesting to note that the upper critical mixing ratio, $z_u$, for forming stable PEMECs is quite similar, ~0.7, for NaPA and NaHA, which means that the colloidal stability here is largely governed by the relative charge ratio of surfactant and polyelectrolyte. For both polyelectrolytes, the phase boundary between unstable and long-time stable samples does not depend much on the $M_w$ value of the polyelectrolyte, but for NaHA, the range of metastable phases within the unstable region is much smaller, i.e., the microscopic phase separation occurs much faster here.

To further generalize our findings, we also employed NaCMC as polyelectrolyte, where the polymer structure and stiffness are similar to those of NaHA, but the charge is attached to the backbone by a flexible spacer (see Figure 1), and the charge density can easily be altered by the degree of substitution (DS). Here, the higher-$M_w$ NaCMC250 has a higher charge density with DS = 1.2, while DS = 0.7 for the shorter NaCMC90. When comparing their phase behavior (see Figure 2, right), it is striking that the NaCMC phase diagram for the small droplets is somewhat different by showing a two-phase region more extended to higher $z$. But the more pronounced effect is seen on the droplet size when mixed with NaCMC (see Figure S4). For samples prepared with NaCMC and medium-large droplets, the monophasic range is much smaller than that with the two other polyelectrolytes, and it almost vanishes for the largest droplets. As mentioned before, the NaCMC charge is not located directly on the PE backbone, but linked with a rather flexible spacer. This spacer makes the charge more accessible for the ME droplets, so droplets can be bound more easily, which increases the strength of interaction, and the complexes then have an increased tendency for phase separation. Apparently, the accessibility of the charge via the flexibility provided by a spacer is very important here.

Interestingly, the nature of the two-phase region depends strongly on the type of PE. For NaPA, a solid powdery phase precipitates, while for NaHA and NaCMC, the second phase is a very viscous gel, i.e., a coacervate phase. Apparently, the extent of hydration in the complex is much higher for the polysaccharides, which is not surprising, given the large number of OH groups on their backbone, which enable stronger hydration. It should be noted that this behavior is quite similar to that observed for complexes of NaHA with TTAB, where a viscous second phase was observed, the difference just being that there the viscous phase precipitated to the bottom. The difference in our case can be explained by the lower density of the microemulsion droplets.

**Structural Characterization of ME/NaHA Complexes. Static and Dynamic Light Scattering (SLS/DLS).** A first structural characterization of the stable single-phase samples at NaHA excess was done by static and dynamic light scattering. DLS measurements show an interesting feature different from the previously studied NaPA samples: Some of the NaHA samples show a bimodal decay (see Figure S5). The tendency for a bimodal distribution function depends on the droplet size and the $M_w$ value of NaHA. The smallest droplets mostly show a bimodal distribution. For ME00, a bimodal relaxation is found for all mixing ratios and all $M_w$ except the shortest NaHA51, which exhibits a monomodal relaxation. For the medium droplet size ME50, only samples with higher-$M_w$ NaHA show a bimodal decay, and only for higher mixing ratios. Finally, for the largest droplets ME75, a bimodal relaxation is no longer found and all measured samples show a simple decay of the correlation function (Figure S6). The slower relaxation mode in the bimodal distribution is nondiffusive (the relaxation rate, $\Gamma$, follows a power law of $\sim q^2$; see Figure S7) and also appears in samples of pure NaHA. It can be attributed to the generally observed “slow mode” of polyelectrolytes, seen in particular at a low ionic strength, which has been attributed to the presence of supramolecular structures or hindered motions of interacting chains. Such a slow mode with $q^2$ scaling has been reported also for hyaluronate before. Because of the lower charge density, NaHA has to be employed in higher weight fractions than NaPA to achieve the same mixing ratios. NaHA also scatters light more strongly than NaPA due to higher $M_w$ per unit
length, so it becomes visible in the measurements at higher concentrations. Even though NaHA is scattering more strongly, it is still a weak scatterer and can be easily concealed by more strongly scattering objects like larger microemulsion droplets. The faster mode observed in DLS belongs to the complexes formed by PE and ME droplets, which are of interest in this study, and thus, only this signal was further analyzed and the slower relaxation mode was neglected.

Both methods, SLS and DLS, show large aggregates close to the phase boundary, which are decreasing in size for larger charge ratios \( z \). As expected, the higher \( M_w \) of the polyelectrolyte and larger ME droplets lead to an increase in complex size. These findings are very similar to the results previously found for ME/NaPA complexes, indicating a generic trend for such systems, independent of the nature of the polyelectrolyte. All light scattering results summarized in Figure S8 show hydrodynamic radii of 25–50 nm close to the phase boundary for the smallest droplets, which increase with the \( M_w \) of the NaHA. Interestingly, the largest values of almost 100 nm are found for medium-size ME droplets (\( R = 4.1 \) nm) and slightly smaller for the largest droplets (\( R = 6.6 \) nm), where the largest complexes of \( R_q \sim 80–100 \) nm are observed for \( M_w \) values of 360 and 800 kDa for NaHA.

The static scattering intensity shows a plateau over the whole \( q \)-range for most samples. Only some samples with high-\( M_w \) NaHA and mixing ratios close to the phase boundary (\( z = 0.7 \)) show an increase in intensity to lower \( q \) (see Figure 3, left, and Figure S9). A plateau in the SLS data means that \( R_q q \ll 1 \), which would be valid for aggregates with \( R_q < 40 \) nm (with \( q_{\text{max}} \) being 0.025 nm\(^{-1}\)), in agreement with the DLS data (one should keep in mind here that for a cylinder of length \( L \) and radius \( R \), it holds: \( R_q^2 = L^2/12 + R^2/2 \); accordingly, this is still valid for cylinders of a length of up to 140 nm). The molecular weight of the complexes can be obtained from the intercept \( I(0) \) of the SLS data (obtained by a Guinier approximation); it shows that close to the phase boundary, values of several million Da are achieved (Table S2). By comparing the molecular weight of the mixed complexes to that of one single ME droplet without interactions (pure microemulsion measured with added salt to suppress interparticle interactions), and neglecting the weak scattering contribution of the polyelectrolyte, an aggregation number \( N_{agg} \) of droplets per complex aggregate can be obtained. This shows that up to 40 droplets may be contained in one complex (Figure 3, right, and Figure S8). The static light scattering results for all NaHA samples are summarized in Table S2. In general, the number of droplets contained in a complex increases with increasing \( M_w \) of the NaHA chain. However, it should be noted here that neglecting the scattering contribution of the NaHA leads to an upper estimate. Similarly, we are not claiming to find finite sized structures in the solution, but at the length scale probed here, we see structures that contain approximately the given numbers of ME droplets.

**Viscosity.** For additional information, the viscosities of a \( z \)-series of medium-sized microemulsion droplets (ME50) and NaHA186 were measured. The viscosity is very low at ME excess and increases strongly once the PE excess regime of the phase diagram is reached. This is simply due to the increased concentrations of NaHA in this regime, which already for the pure NaHA solution leads to a marked viscosity increase (see Figure S10). Figure 4 shows the viscosity ratios of ME/NaHA samples and pure NaHA solutions with the same NaHA concentration, for different charge ratios at PE excess. The viscosity ratios of NaPA and NaCMC samples of \( z = 0.7 \) are also plotted for comparison. As a first statement, it can be pointed out that the added microemulsion does not have a

![Figure 3](https://dx.doi.org/10.1021/acs.macromol.0c00236)

**Figure 3.** Left: SLS intensity vs \( q \) shows a plateau for most samples (small droplets, \( R = 3.1 \) nm). Right: Aggregation numbers \( N_{agg} \) (number of droplets per complex) as obtained from static light scattering (small droplets, \( R = 3.1 \) nm). For data on bigger droplets, see Figures S8 and S9. The last number of the sample name indicates the charge ratio \( z \) of the sample.

![Figure S8](https://dx.doi.org/10.1021/acs.macromol.0c00236)

**Figure S8.** The static light scattering results for all NaHA samples are summarized in Table S2. In general, the number of droplets contained in a complex increases with increasing \( M_w \) of the NaHA chain. However, it should be noted here that neglecting the scattering contribution of the NaHA leads to an upper estimate. Similarly, we are not claiming to find finite sized structures in the solution, but at the length scale probed here, we see structures that contain approximately the given numbers of ME droplets.

![Figure 4](https://dx.doi.org/10.1021/acs.macromol.0c00236)

**Figure 4.** Viscosity ratios of pure polyelectrolytes and PEMECs for different charge ratios \( z \) for medium-sized microemulsion droplets (\( R = 4.3 \) nm).
strong impact on the viscosity of the polyelectrolyte solution. For NaHA186, the largest effect is obtained at $z = 0.7$, close to the phase boundary, where the microemulsion decreases the viscosity compared to the original PE viscosity by a factor of almost 2. This effect becomes much weaker for higher mixing ratios. The viscosity effect is markedly depending on $M_w$ and is about 2 for NaHAS72, while it almost vanishes for NaHA31. For NaCMC250, a similar value to that for NaHAS72 is observed, which may be partly attributed to the similar $M_w$ but may also reflect a stronger attractive interaction due to the more accessible charges of NaCMC.

These results are interpreted by the rather weak electrostatic interactions of the microemulsion droplets with the polyelectrolytes, which is most likely due to the low charge density on the droplet surface. This should result in rather transient aggregates with a short structural relaxation time, thereby not affecting the viscosity significantly. The lower relative viscosity at low $z$ can be explained such that the complexation is most pronounced here, which also leads to a condensation of NaHA chains and thereby to a relatively weaker polyelectrolyte network and correspondingly lowered viscosity. Close to the phase boundary, where large aggregates were observed by DLS and SLS, those weak interactions still have a compacting effect on the hyaluronate, the more so for longer chains (higher $M_w$). When the hyaluronate is bound in aggregates, the charges are screened, and the separate strands can slide next to each other without entanglement. At larger polyelectrolyte excess, the hydrodynamic radii obtained from DLS become smaller, actually approaching those of the free droplets (see Figures 6 and 8). Together with the viscosity measurements, this can be interpreted as a continuous but transient polyelectrolyte network (no change in viscosity), which is decorated with separate ME droplets. Apparently, the transient nature of the interconnection by ME droplets results in a short structural relaxation time and the corresponding low viscosity. In general, it can be concluded that the addition of the ME droplets has a rather small effect on the viscosity of the NaHA network, as previously seen for the addition of pure cationic surfactant at surfactant excess conditions.41 For a comparison, the viscosity effect on NaCMC is more strongly pronounced, confirming a stronger interaction (as seen before in the phase behavior) and a resulting more pronounced compaction of the aggregates.

Small-Angle Neutron Scattering (SANS) and Dynamic Light Scattering (DLS). Although light scattering is a useful tool to determine the overall size of the formed complexes, it is not able to yield detailed information about the shape and internal structure of these complexes because of the limited $q$-range probed and the small size of the microemulsion droplets. For this reason, small-angle neutron scattering (SANS) was employed to characterize the different PEMEC systems with respect to their mesoscopic structure. This was done comprehensively for different $M_w$ values of NaHA and for different size microemulsion droplets, always as a function of the mixing ratio, $z$. Selected SANS data are shown in Figure 5. The full set of scattering curves is shown in Figure S12.

As previously observed for the polycrylate PEMEC,17 the SANS measurements show that the droplets remain unchanged in shape and size when interacting with the hyaluronate. This is confirmed by the unchanged scattering curves at high $q$, especially the form factor minimum (see Figure S13 for zoom into the minimum). The complexing effect of the hyaluronate is seen at low $q$, where the interactions of the droplets are probed. All SANS data show an increase in intensity toward low $q$ that continues up to $q = 0.03$ nm$^{-1}$. Note that no plateau is visible in any of the SANS data of NaHA complexes. In comparison, NaPA samples with small $M_w$ showed a plateau; however, the shortest NaHA chains used in this work are still much longer than the shorter NaPA chains used previously.17

As the differences in the size of the formed complexes become clearer in static light scattering, where the observed $q$-range is at lower $q$ values, the SLS data were scaled according to the contrast differences of SANS and SLS (for details of the calculation and refractive index increments, see SI F) and plotted with the SANS data in Figure 5 for a complete picture. A plateau is observed for most samples in SLS. Unfortunately, the onset of the plateau seems to be exactly in the middle between the two observable length scales of SLS and SANS, and due to the experimental constraints, no overlapping data are available, but by combining these two methods, a good characterization of the PEMECs can still be achieved.

The slope of the intensity increase in the SANS data depends on the mixing ratio, $z$. As previously seen for NaPA, the slope is highest for samples close to the phase boundary (low $z$) and decreases with increasing polyelectrolyte excess, i.e., with increasing electrostatic screening. As an example, this is shown for the smallest microemulsion droplets (ME00, $R = 3.1$ nm) for the shortest ($M_w = 51$ kDa) NaHA in Figure 5 (left) (others are shown in Figure S12). The largest aggregates...
are formed close to the phase boundary (z ∼ 0.6), and the q−2 slope suggests random walk assemblies of microemulsion droplets at this mixing ratio. At z = 0.7, the slope of the intensity is close to q−1, which indicates locally linear arrangements of the microemulsion droplets. At high polyelectrolyte excess (z = 0.9), only a small intensity increase is still visible. Figure 5 (right) compares the SANS data of complexes of small microemulsion droplets at the same mixing ratio (z = 0.7, where linear droplet arrangements were found) with different Mw’s NaHA. Here, only small deviations in the q−1 slope are visible, becoming somewhat steeper with increasing Mw value of NaHA. The bigger the droplet, the smaller is the visible difference between the NaHAs of different Mw’s (see Figure S14 for medium and large droplets). Due to the lack of a plateau, no characteristic length of these linear complexes can be determined from SANS. Simulations of SANS curves based on homogeneous cylinders with different lengths show that the linear PEMEC arrangements extend over more than 100 nm (see Figure S15), but cannot be much larger than the values as seen by the plateau in the SLS data.

It is interesting to note that the local arrangement of ME droplets in the PEMECs apparently is rather independent of the droplet size and the Mw of the NaHA and depends largely on the mixing ratio. For larger ME droplets, the main difference in the SANS data is simply explained by the higher Mw values of these droplets. This result is different from previous measurements with NaPA, where we had found higher aggregation numbers for larger droplets.17

Knowing that the droplets are arranged in locally linear arrangements while retaining their original size and shape, we can now interpret the (fast) diffusion coefficients measured in DLS as the diffusion of cylinders with a diameter of one droplet. The length of one cylinder was calculated using the established expression for the diffusion coefficient by Ortega et al.42 (the radius was taken as that of the microemulsion droplets; more details are given in SI D).

\[
D_1(L) = \frac{1}{3} \frac{k_B T (\ln p + C_p)}{\eta L}
\]

The values given in Figure 6 show that this model predicts the presence of rods in the length range of 50−200 nm, in good agreement with the SANS data (and one should keep in mind that this still corresponds to a value of Rg smaller than 50 nm).

Comparison of PEMECs Formed with Sodium Hyaluronate (NaHA), Sodium Carboxymethyl Cellulose...
(NaCMC), and Sodium Polyacrylate (NaPA). After having analyzed the structural information regarding PEMECs formed with NaHA of different \( M_w \)’s, it is now interesting to compare their structural features with those seen previously for sodium polyacrylate (NaPA). This comparison is interesting since NaPA is a much more flexible polyanion than NaHA with its polysaccharide backbone (intrinsic persistence lengths of 1.3 nm compared to 9 nm for NaHA). Accordingly, we can elucidate the effect of the polymer persistence length on the mesoscopic structure of the PEMECs. In addition, some experiments were performed with the sodium salt of carboxymethyl cellulose (NaCMC) to see how the location of the charge on the polysaccharide affects the complexation behavior, while the backbone and its stiffness ( persistence length of 16 nm) are quite similar to those of NaHA. For NaHA and NaPA, the charge is located close to the backbone, while for NaCMC, it is much more easily accessible, as it is connected to the backbone by a rather flexible spacer (see Figure 1). The charge density of NaCMC depends on the degree of carboxylation, and the two NaCMCs used in this work differ not only in their molecular weight but also in their charge density. The longer NaCMC chain (NaCMC250) also has the higher charge density with 1.2 charges per monomer unit, while the shorter NaCMC90 has a degree of substitution (DS) of 0.7.

Static light scattering was measured for all samples to determine the molecular weight and the aggregation number (Figure 7). The SLS data show the same trend for all polyelectrolytes: the aggregation numbers at a constant charge ratio, \( z \), increase with increasing \( M_w \) of the PE. NaPA/ME complexes have a higher aggregation number than NaHA complexes at a given \( M_w \) of PE, but the aggregation number is highest for NaCMC complexes. In general, \( N_{agg} \) is smaller than that calculated theoretically with the assumption of having one polymer chain complexed stoichiometrically with ME droplets according to the number of charges (given as dotted lines in Figure 7, right). Here, it should be noted that the droplets have a rather low charge density, as the average spacing between the charges on the droplet surface is \( \sim 3.5 \) nm, while that of NaHA is \( \sim 1 \) nm, i.e., a 3.5 times higher value. Under these conditions, it is not possible to fully compensate all PE charges, leading to smaller aggregation numbers. Because of this, the complexes will build up a negative net charge, which is the driving force for their locally elongated form, and which also prevents the complexes from growing very large. This effect is more pronounced with increasing \( M_w \) of the polyelectrolyte, because at the given conditions, the PE concentration is typically above the overlap concentration, which means that the formation of complexes with only one PE chain is unlikely. In addition, only an effective domain size of the structures is visible in light scattering, making it more difficult to give exact aggregation numbers.

The SANS measurements are able to give more information about the shape and composition of the mixed complexes. At first glance, the complexes formed with microemulsion droplets and NaHA, NaCMC, or NaPA look rather similar in SANS. We observe large mixed complexes close to the phase boundary that decrease in size with increasing polyelectrolyte excess (Figures S12 and S17). Figure 8 shows the SANS data for samples of small microemulsion droplets (ME00, \( R = 3.1 \) nm) with different polyelectrolytes at a constant charge ratio of \( z = 0.7 \) (polyelectrolyte excess, close to the phase boundary for NaPA and NaCMC, not so close for NaHA). All of these complexes show a slope between \( q^{-1} \) and \( q^{-1.5} \) in the neutron scattering data, which corresponds to elongated structures. The \( M_w \) values of the polyelectrolytes were chosen so that the stretched chains (contour lengths) are similar in lengths (NaPA60 ~ 160 nm, NaHAS1 ~ 130 nm, NaCMC90 ~ 200 nm). However, it should be noted here that being mixed at the same charge ratio implies that the weight fraction of NaHA (NaCMC90/NaCMC250) is about 4 (3/2) times higher than that of NaPA, due to the lower charge density per mass of NaHA (NaCMC). In Figure S16, the same comparison is done for medium-sized droplets (ME50, \( R = 4.1 \) nm), and the scattering curves look very similar. Figure S17 shows more SANS data of NaCMC complexes, which resemble those of the NaHA in their general behavior.

The direct comparison for a constant mixing ratio of \( z = 0.7 \) shows that the slope for the NaPA samples is generally higher than those for the NaHA or NaCMC ones. While the slopes of the more rigid polysaccharides are very close to a \( q^{-1} \) power law, the slope of the flexible NaPA is \( \sim q^{-1.5} \). This can be attributed to the different stiffnesses of the polyelectrolytes; NaHA and NaCMC are quite rigid due to their backbones being composed of disaccharide units, while NaPA is rather flexible. The stiffer NaHA and NaCMC are more likely to arrange the droplets in a straight cylindrical geometry leading to a \( q^{-1} \) dependence, while the less rigid NaPA forms curved chains of droplets, i.e., assumes the conformation of a semiflexible polymer. Consequently, the low-q region exhibits a higher slope \( (q^{-1.5}) \). The two limits generally known to describe SANS data of similar polymeric structures quantitatively are long straight cylinders with a slope of \( q^{-1} \) or flexible Gaussian coils with a slope of \( q^{-2} \). We can therefore qualitatively conclude that the two polysaccharides employed here form long aggregates with essentially linear arrangements of separate droplets, when interacting with oppositely charged microemulsion droplets. In contrast, the formed aggregates of the flexible NaPA show a semiflexible behavior.

The comparison of the SANS data of NaPA and NaHA samples at \( z = 0.9 \) (data shown in Figure S18), i.e., at very high PE excess, points out another difference. At this high PE excess, the NaPA samples resemble those of single microemulsion droplets without interactions. Apparently, the PE excess at this charge ratio is high enough to stabilize each...
droplet separately. NaHA and NaCMC complexes at the same mixing ratio still show a small intensity increase at very low q. Here, the polyelectrolyte network formation becomes visible, which is much more pronounced for NaHA, in agreement with the highly viscous properties that it shows in aqueous solution.44,45 The small dent visible in these SANS data at q ~ 0.2 nm⁻¹ reminds of a correlation hole, which arises from excluded-volume correlations due to the increased amount of polyelectrolyte.46

Cryo-TEM images were taken of samples of large ME droplets with NaPA and NaHA at PE excess, but close to the phase boundary (z = 0.7 and 0.8) to validate the droplet arrangements suggested by SANS measurements (Figure 9).

While both images show dense packings of droplets that are hard to separate into single aggregates, those images also do not contradict the findings obtained by SANS, which is an important observation. In both images, we can find linear arrangements of droplets. For the NaPA sample, the strands of droplets look more branched and bend than for the NaHA sample, where more extended strings of ME droplets are visible (see indications in red in Figure 9). To be able to interpret the results of two fundamentally different methods in the same way increases the chances that this interpretation represents the real situation substantially.

Further Analysis of the Scattering Data. Further information regarding the SANS data can be obtained by plotting and analyzing the data in the so-called “Holtzer plot,”47 I(q) vs q. For flexible polymers, the Holtzer plot shows a maximum in intensity at low q, which decreases first rapidly and then with a smaller slope. The kink between the two linear parts at qk (transition from coiled to linear structure) can be used to calculate the persistence length lp = 1.9/qk.48 When looking at SANS data of different polyelectrolytes with microemulsion droplets at z = 0.7 in the Holtzer plot, the differences become much more visible than in the simple log–log plot (Figure 10). While NaPA samples follow the classical behavior of polymers, the trend of NaHA samples looks completely different. For the flexible NaPA, a polymer-like behavior is observed with a persistence length lp = 5.81 nm for ME00-NaPA60-0.70, and lp = 9.05 nm for the ME50-NaPA60-0.70 complexes. Interestingly, these lengths correspond almost exactly to the respective droplet diameters, which indicates that dimers of droplets are rather loosely joined. Samples with NaHA at this mixing ratio do not follow the classical Holtzer plot behavior for a flexible chain, so no persistence length can be obtained here. Instead, for the NaHA and NaCMC complexes, one observes the behavior of a stiff rod, and obviously, the persistence length is larger than the experimental observation window of 100 nm. This confirms the previously made assumption that here the ME droplets arrange to straight cylindrical aggregates of a substantial length. The Holtzer plots for NaHA samples closer to the phase boundary (z = 0.6 and 0.65) and at high PE excess are shown in Figure S19A,B.

Another way of comparing the complexes formed by ME droplets with different polyelectrolytes is by fitting the slope, A, of the linear part of the SANS data at small q with I(q) = q⁻⁴⁺Α + B, where A corresponds to the fractal dimension of the formed complexes. Here, a value of A = 1 corresponds to straight cylinders, whereas higher values indicate more bend cylinders. The limit of this model would be a Gaussian coil with A = 2.39 The fitted exponents for z = 0.7 samples are plotted in Figure 11 and show a logarithmic trend with the Mw value of the polyelectrolyte for all systems. The A value of NaPA samples is generally higher than that of NaHA samples at a given mixing ratio, confirming again that much more flexible aggregates are formed by the flexible polyelectrolyte, while a stiff polyelectrolyte arranges the ME droplets in straight rods. In addition, the slope increases systematically with increasing size of the microemulsion droplets. The NaHA samples closer to the

Figure 9. Comparison of cryo-TEM micrographs of large ME droplets (ME75, R = 6.6 nm) mixed with NaPA (left) and NaHA (right) at z-values in PE excess, close to the phase boundary (z = 0.7 for NaHA and 0.8 for NaPA, because a z = 0.7 sample was not stable with large droplets).

Figure 10. Holtzer plots of SANS data of NaHA, NaPA, and NaCMC mixed with small (left) and medium (right)-sized microemulsion droplets. A charge ratio of z = 0.7 was not stable for ME50-NaCMC90 samples.
phase boundary \((z = 0.6 - 0.65)\) showed slopes between 1.6 and 1.1, but no direct comparison to NaPA can be drawn here because NaPA samples were not stable at these mixing ratios.

More analytical description of the scattering data was also made. As a first step, the data were described with a model of homogeneous cylinders, before a more sophisticated model of linearly arranged spheres was attempted. However, both models showed substantial shortcomings in properly describing the scattering data (see SI F). Due to the lack of an appropriate model to describe SANS data of curved droplet chains, a Monte Carlo model was developed by us for this purpose (a comparison of the fit quality of the different models is given in Figure S20, showing the significant improvement by the new model). The model describes chains of \(N\) homogeneous spheres of average radius \(R\), which are separated by an average distance \(d\) between the sphere surfaces. Both \(R\) and \(d\) are polydisperse and randomly picked from normal distributions with standard deviations \((\sigma)\) corresponding to the polydispersity parameters. The chains assume the random conformations of semiflexible polymers of a given persistence length. In practice, this was achieved by sampling random conformations while penalizing high angles between the vectors between each sphere and its two neighbors with a harmonic constraint potential of adjustable strength \(V_0\). In this procedure, nonphysical conformations with overlapping spheres were excluded. To obtain the scattering curves given in Figure 12, at least 5000 conformations were randomly generated in this way. The scattering intensities were then computed by calculating the form factor amplitudes of each sphere, subsequent phase-correct summation based on each set of sphere coordinates using the Debye formula, and averaging over all conformations. The persistence length \(l_p\) was calculated a posteriori from the ensemble-averaged ratio between contour length \(L_C\) and end-to-end distance \(h\), where \(\langle h^2 \rangle = 2L_Cl_p - 2h^2(1 - e^{-L_C/l_p})\) (for more details, see SI F).

With this model at hand, it was possible to quantitatively describe the full set of SANS data of samples with \(z = 0.7\) for different polyelectrolytes (Figure 12). To this end, all microemulsion droplet parameters \((R, \text{polydispersity in } R, \text{volume fraction, contrast})\) were kept constant for all samples. Parameters that depend on the type of polyelectrolyte \((d, \text{polydispersity in } d, V_0)\) were used as global model parameters for all samples with the same PE type. So, the only parameter changing with the \(M_w\) of PE is the number of droplets per aggregate \(N\).

The results show that all data of samples close to the phase boundary (where elongated complexes are found) can be described accurately with the new model of cylindrically arranged droplets with a persistence length of the aggregate that depends on the type of polyelectrolyte. The obtained persistence lengths are \(l_p(\text{NaPA}) \sim 17 \text{ nm}\) and \(l_p(\text{NaHA}) \sim 51 \text{ nm}\), confirming and quantifying the assumptions stated above. Furthermore, the distances between droplet surfaces depend on the type of polyelectrolyte, being \(\sim 2 \text{ nm}\) for NaPA and NaCMC.
~5.5 nm for NaHA complexes, which can be explained by the different charge densities of both polyelectrolytes, which are 4.0 e0/nm for NaPA and 1.0 e0/nm for NaHA (Table 1).

<table>
<thead>
<tr>
<th>sample</th>
<th>R/nm (σ)</th>
<th>d/nm (σ)</th>
<th>N</th>
<th>V0/a</th>
<th>Lc/nm</th>
<th>l/u/nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>ME50-NaPA05-0.70</td>
<td>4.0 (0.6)</td>
<td>2.0 (1)</td>
<td>4</td>
<td>1.0</td>
<td>30</td>
<td>18</td>
</tr>
<tr>
<td>ME50-NaPA15-0.70</td>
<td>4.0 (0.6)</td>
<td>2.0 (1)</td>
<td>5</td>
<td>1.0</td>
<td>40</td>
<td>17</td>
</tr>
<tr>
<td>ME50-NaPA60-0.70</td>
<td>4.0 (0.6)</td>
<td>2.0 (1)</td>
<td>17</td>
<td>1.0</td>
<td>160</td>
<td>16</td>
</tr>
<tr>
<td>ME50-NaPA315-0.70</td>
<td>4.0 (0.6)</td>
<td>2.0 (1)</td>
<td>32</td>
<td>1.0</td>
<td>310</td>
<td>17</td>
</tr>
<tr>
<td>ME50-NaHA51-0.70</td>
<td>4.0 (0.6)</td>
<td>5.5 (3)</td>
<td>6</td>
<td>3.0</td>
<td>68</td>
<td>54</td>
</tr>
<tr>
<td>ME50-NaHA150-0.70</td>
<td>4.0 (0.6)</td>
<td>5.5 (3)</td>
<td>9</td>
<td>3.0</td>
<td>108</td>
<td>52</td>
</tr>
<tr>
<td>ME50-NaHA360-0.70</td>
<td>4.0 (0.6)</td>
<td>5.5 (3)</td>
<td>13</td>
<td>3.0</td>
<td>162</td>
<td>50</td>
</tr>
<tr>
<td>ME50-NaHA800-0.70</td>
<td>4.0 (0.6)</td>
<td>5.5 (3)</td>
<td>15</td>
<td>3.0</td>
<td>190</td>
<td>50</td>
</tr>
</tbody>
</table>

*Roman was kept constant for all ME50 samples. d and V0 were used as global fit parameters for each PE type. Lc and l were calculated from the modeled conformations. Minor variations in l for fixed V0 reflect the influence of the nonoverlap requirement and the stochastic nature of the model.

## CONCLUSIONS

In summary, we studied the structure of polyelectrolyte/micromulsion complexes (PEMECs) of positively charged micromulsion droplets with different negatively charged polyelectrolytes. Three very different PEs were employed: The biopolymer sodium hyaluronate (NaHA) was the main focus of this study, and sodium polyacrylate (NaPA) and the sodium salt of carboxymethyl cellulose (NaCMC) were studied for a systematic comparison. While the molecular architectures of NaHA and NaCMC and also their persistence lengths in aqueous solution are rather similar, the two PEs exhibit very different phase behaviors when interacting with micromulsion droplets. For NaCMC with its more flexibly attached charge, the strength of the interaction with the oppositely charged micromulsion droplets is much larger, and therefore, a phase separation takes place much more easily, and also the viscosity reduction is more marked. In contrast, NaHA shows a similar phase behavior to NaPA even though the PE backbone as well as the charge density are very different. In addition, it is interesting to note that for NaHA and NaCMC, a coacervate is formed, while for NaPA, a simple powdery precipitate is observed. Since the charged group, sodium carboxylate, is the same for all employed PEs, we conclude that the main parameter to influence the phase behavior of such systems is the distance between the charged group and the polyelectrolyte backbone. Other parameters, such as Mw of the PE, nature of the PE backbone, and charge density, play only a secondary role. However, it can be noted that the tendency for precipitation generally increases with increasing droplet size and Mw of the PE. This can be explained such that with increasing size of the micromulsion droplets, their charge density decreases (≈1/R), thereby rendering it easier to compensate the excess charge of the PE. In addition, their surface becomes less curved, which should also make it easier for the PE charges to interact with them.

The structural behavior of stable PEMECs for polyelectrolyte excess is similar for all studied PEs. All systems locally form large elongated structures at PE excess close to the phase boundary, which decrease in size when the PE excess is increased, due to the increased local charging in the complexes. The elongated complexes are formed by separate micromulsion droplets arranged in cylindrical geometries. The ME droplets retain their size and shape when interacting with the polyelectrolytes. The number of droplets per complex depends mainly on the Mw value of the employed polyelectrolyte; it increases proportionally to the latter. The structure of the complexes was studied in detail by small-angle neutron scattering (SANS). It was found that, although though the scattering data of the formed complexes look very similar at first glance, the exact morphology depends strongly on the backbone of the employed polyelectrolyte. For a quantitative analysis, a stochastic model was developed that describes such cylindrically arranged droplets with different persistence lengths of the complexes accurately. The more flexible polyelectrolyte NaPA also forms much more flexible elongated complexes, where the persistence length corresponds to the size of about two micromulsion droplets. Contrarily, stiffer polyelectrolytes like NaHA or NaCMC form much more rigid rodlike aggregates of ~50 nm persistence length when interacting with the micromulsion droplets, as shown in Figure 13. The effectively observed length of these cylindrical

![Figure 13](https://dx.doi.org/10.1021/acsmacrolett.0c00236)
aggregates then was 100–200 nm, while NaPA on this length scale forms structures similar to Gaussian coils. It should also be noted that these aggregates/complexes are not just containing one polyelectrolyte chain, and especially for longer chains (higher $M_c$), they will be interconnected by such chains, thereby forming a transient network, as depicted in Figure 13.

To conclude, it was found that PEMECs can be formed generically with quite different types of polyelectrolytes, and large complexes are formed at polyelectrolyte excess in the vicinity of the phase boundary. The local structure of the complexes is largely controlled by the stiffness of the polyelectrolyte, where quite flexible complexes with short cylindrical units are formed by the flexible PE NaPA, while much stiffer cylindrical arrangements are present for polysaccharide-based PEs like NaHA. The main finding of our work therefore is that the persistence length of the PE becomes translated and amplified into the persistence length of the PEMEC aggregates in a manner similar to that seen before for complexes of lysozyme and hyaluronan. In this manner, PEMECs with tailor-made structures can be designed, to thereby forming a transient network, as depicted in Figure 13.

ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.macromol.0c00236.

Sample compositions, phase behavior, light scattering data, determination of diffusion coefficient of rods, viscosities, description of SANS models, additional SANS data, and list of abbreviations (PDF)

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