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Unexpected observation of an intermediate hexagonal phase upon fluid-to-gel transition: SDS self-assembly in glycerol

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ABSTRACT

Understanding morphological transformations upon temperature-induced mesophase transitions offers mechanistic insights into the self-assembly process. We have recently reported the unexpected formation of a microfibrillar lamellar gel in SDS-glycerol mixtures above a critical gelation concentration (CGC) as low as ~2 wt%. The gel phase comprised a fibrillar structure on the microscale and a lamellar structure on the nanoscale. Here, the nanoscopic structure of the gel as a function of temperature was probed with small-angle neutron scattering (SANS). The gel underwent a gel-to-fluid transition at a critical gelation temperature, \( T_{GC} = 45 \) °C, forming cylindrical micelles at elevated temperatures. Upon cooling, a hexagonal phase formed at ~\( T_{GC} \) evident from the SANS Bragg peaks. This hexagonal phase upon the fluid-to-gel transition sheds light on the gelation mechanism, in which self-assembled SDS micelles undergo a cylindrical-to-lamellar morphological transition via a hexagonal phase. This unprecedented observation also highlights the complexity of self-assembly in nonaqueous hydrogen-bonding rich media.

1. Introduction

Understanding the mechanism of thermotropic mesophase transitions is important to many applications, ranging from industrial formulations to biological processes. For instance, it is relevant to the molecular mechanisms of endocytosis [1] and cryobiology [2], as well as polymorphs of chocolate [3–6] and soaps [7–10] at elevated temperatures during their formulation processes.

The lamellar-to-hexagonal lipid mesophase transition has been widely studied due to its relevance to the molecular deformations involved in the membrane fusion process, in which non-bilayer intermediates could form as postulated in the Stalk model [11]. In particular, phosphatidylylthanolamine (PE) lipids are reported to be prone to the formation of the hexagonal phase in physiological conditions, and are thus implicated in mediating membrane fusion [12,13]. For instance, the kinetics of the reversible lamellar-to-inverse hexagonal phase transition of dihexadecyl phosphatidylethanolamine (DHPE) has been previously probed with X-ray diffraction (XRD), suggesting that the temperature-induced shift in the free energy of the lipid molecule caused its molecular shape to change from a wedge to a rod upon the transition [14–17]. It has also been reported that the tetraoleoyl cardiolipin lamellar phase formed a hexagonal phase in the presence of 3.5 M sodium chloride (NaCl) [18].

Using XRD, Yang and Huang demonstrated the formation of an intermediate ‘stalk’ (an hourglass shape) phase upon fusion of lipid bilayers [19]. The formation of this intermediate phase was predicted mathematically [20], where the energy of the structure was determined by the lipid chain splay and tilt, which is dependent on the membrane composition [21–23]. The formation of the stalk intermediate has also been observed in self-assembled structures of lung surfactant in the presence of a hydrophobic protein (SP-B) [24], conjectured to be relevant to fatal neonatal respiratory diseases due to SP-B deficiency [25].

Temperature-induced phase transitions are common in surfactant systems. For instance, the isotropic to hexagonal phase transition in Triton X100 aqueous media occurs at \( T = 33 \) °C [26,27]. A transition from the lamellar gel phase to micellar and lamellar liquid crystalline...
phases was observed in aqueous solution of anionic sodium lauryl sulfoacetate at $T > 50$ °C [28]. Lysine-based surfactants were reported to undergo a phase transition from micellar aggregates to a gel phase upon cooling below its Krafft temperature due to its suppressed solubility [29].

Molecular dynamic (MD) simulations of sodium dodecyl sulfate (SDS) in aqueous salt solutions (700 mM calcium chloride, CaCl$_2$) showed micelle fission via a stalk intermediate [30], similar to that in the lipid membrane fusion process. However, experimental observations of intermediate structures upon surfactant mesophase transitions have not been reported.

Here, we report that an intermediate hexagonal phase formed in the fluid-to-gel transition upon cooling of 4.4 wt% sodium dodecyl sulfate (SDS)-in-glycerol gel. The supramolecular gel was thermally reversible [31], and below a critical gelation temperature ($T_{GC}$) ~ 45 °C, the gel phase comprised a fibrillar structure on the microscale and a lamellar structure on the nanoscale. Above $T_{GC}$, the fluid phase consisted of cylindrical SDS micelles. Intriguingly, upon cooling from the fluid phase, an intermediate hexagonal phase formed, which has not been previously reported, including in aqueous SDS systems [32,33]. The nanostructures of the fluid-, intermediate-, and gel-phase were probed using small-angle neutron scattering (SANS). The observation of this intermediate phase highlights the complexity of self-assembly in H-bonding non-aqueous media, which is very different from the aqueous medium. Although our focus here is the mesophase phase transition, we note that hydrogels and nanocomposite gels with different intricate nanostructures that are tuneable by external conditions have potential in a wide range of applications (e.g. [34,35]). Thus, understanding the mechanism of the gelation mechanism in our novel nonaqueous lamellar gels is important to their potential future applications.

2. Materials and methods

2.1. Materials

- h-Sodium dodecyl sulfate (SDS) (Sigma-Aldrich, >98.0%) was recrystallised three times from ethanol prior to use, and its purity was checked with $^1$H NMR. h-Glycerol (Fisher Scientific, >98.0%) and d-Glycerol (Sigma-Aldrich, >98.0% and >98.0 atom % D) were used as received. All glycerol-containing (both hydrogenated (h-) and deuterated (d-)) phases and controls were kept sealed from moisture. The gel-like phase was prepared by adding h-SDS to h- or d-glycerol, then incubating and shaking (at 550 RPM) the mixture in a shaker incubator (Stuart SI505) for two hours (2 h) at 60 °C, before equilibrating at room temperature overnight.

2.2. Polarised light microscopy (PLM)

PLM was carried out using an Olympus BX53-P microscope, where the polarisers were crossed at 90° with respect to each other and images were captured using Stream software. PLM measurements were carried out under ambient conditions, using 10, 20, and 40 × magnifications. A 530 nm first order waveplate was placed into the optical path to enhance image contrast.

2.3. Small-angle neutron scattering (SANS)

SANS data was obtained from samples contained in quartz cells with a 2 mm path length on the D11 [36,37] small-angle diffractometer at the Institut Laue-Langevin, ILL, (Grenoble, France). A neutron beam with wavelength ($\lambda = 5.5$ Å with a FWHM of 9%) and three sample-to-detector distances (1.4, 8, and 39 m) were used to obtain a $q$-range of $\sim$0.0016–0.5 Å$^{-1}$, where $q = 4\pi\sin(2\theta)/\lambda$ is the momentum transfer with 2θ the scattering angle. The raw scattering data was corrected for the detector efficiency, sample transmission, and background scattering and converted to scattering cross-section data ($\rho\Delta\Sigma/\rho\Delta\Omega$ vs $q$) using LAMP.

The data was converted to an absolute scale ($\mathrm{cm}^{-1}$) using the secondary calibration standard H$_2$O of 1 mm path length (cross-calibrated against h/d polymer blends sample) with a differential scattering cross section of 0.956 cm$^{-1}$.

2.4. SANS data analysis

The Bragg peaks obtained in the raw scattering profiles were fitted with Gaussian functions using Igor Pro to determine the peak positions ($q$), the peak full width-half maximum ($2\Delta q$), and the coherence length ($L_c$) of the mesophase. The ratio of the peak positions was used to ascertain the surfactant mesophases present in the samples, and the lamellar $d$-spacing was also calculated from the peak position,

$$d = \frac{2\pi}{q}$$

(1)

The coherence length $L_c$, indicative of the structural order in the mesophase, was calculated using the Scherrer equation [38-45]

$$L_c = \frac{2\pi K}{2\Delta q}$$

(2)

where $K$ is a shape factor of order unity.

3. Results and discussion

In a previous paper [31], we have reported the formation of a thermally reversible SDS-in-glycerol lamellar gel above a critical SDS gelation concentration ($c_{GC} \sim 2$ wt%) and below a critical gelation temperature ($T_{GC} \sim 45$ °C; Curve 1 and 4, Fig. 1a). The gel phase consisted of entangled fibres comprising SDS lamellar micro-domains at room temperature, which transformed into cylindrical micelles above $T_{GC}$ (Curve 2, Fig. 1a). The lamellar Bragg peaks noted, previously and here, have a corresponding $d$-spacing, $d = 5.7$ nm. Here, we focus on the self-assembled intermediate structure upon the fluid-to-gel transition. For 4.4 wt% SDS in glycerol, an additional Bragg peak appeared after cooling to 25 °C at $q \sim 0.197$ Å$^{-1}$ (indicated by an arrow on Curve 3, Fig. 2; cf. Table 1), which was not observed before heating (Curve 1, Fig. 2). With respect to the first Bragg peak position $q_0 \sim 0.1$ Å$^{-1}$, this peak was at $\sim 3q_0$, indicative of a hexagonal phase for which the Bragg peak ratio would be 1, $\sqrt{3}$, $\sqrt{4}$ … [46]. This suggests that, upon cooling, the cylindrical micelles observed in the fluid phase above $T_{GC}$ condensed into an intermediate hexagonal phase before transforming into the lamellar phase at thermodynamic equilibrium. We note that the cooling process took place in situ at the SANS beamline over ~30 min at a rate of $\sim 1.5$ °C min$^{-1}$, with a waiting time of 4 h before the measurement, which appeared insufficient to reach the equilibrium structure, probably due to the high viscosity of the hexagonal phase. For comparison, in a previous measurement at ISIS Loq beamline, we observed the equilibrium structure after ~8 h cooling (Curve 4 in Fig. 1) using a similar cooling rate as that at ILL.

The SANS results have confirmed the formation of an intermediate phase in the fluid-to-gel transition upon cooling of the SDS-in-glycerol system. The SANS profile before heating and after overnight (~8 h) cooling were identical, suggesting the thermodynamic equilibrium structure was the lamellar gel phase noted previously [31]. The hexagonal intermediate phase has not been previously reported, whilst the phase diagram of SDS in aqueous media shows that a lamellar-to-hexagonal transition would occur upon cooling [32,33] but at a much higher $c_{GC} \sim 80$ wt%. At $c_{GC} = 4.4$ wt% in water, a micellar phase of spherical or ellipsoidal shape exists [47-49], with no temperature-induced phase transition, in contrast to the behaviour observed here in glycerol. This points to the drastically different self-assembly driving forces in water and in glycerol, despite their similar physical properties such as H-bonding capacity and both being polar, as discussed in Ref. [31].
The formation of intermediate phases adds further details to the proposed mechanism for the gelation of SDS in glycerol at low concentrations – itself an unprecedented observation [31]. Here we have observed that the fluid phase consisting of cylindrical micelles cools via a hexagonal phase before forming the lamellar phase at thermodynamic equilibrium. The fluid-to-gel phase transition can be considered as two steps (Fig. 3): the formation of the hexagonal phase from condensation or packing of the cylindrical micelles, and fusion of the hexagonal phase to form the lamellar phase. The first step is driven by the reduction in the kinetic energy allowing for the intermediate hexagonal phase to be observed.

The second step in the fluid-to-gel transition comprises the hexagonal-to-lamellar transition, where the molecular shape transforms from wedge to rod, noted by Caffrey [14], based on the packing parameter [50, 51]. The metastable hexagonal phase is kinetically stabilised due to the preference of SDS molecular packing in a conical-wedge molecular shape, as per aqueous solutions with a packing parameter \( \sim 0.3 \). The tail volume tends to decrease with increasing temperature, a result of the increased kinetic energy allowing for the opposing holds true upon cooling [52]. The decreasing temperature can thus account for an increase in the tail volume, leading to an increase in the packing parameter, explaining why the structure at thermodynamic equilibrium returns to the lamellar morphology. Another subtle contribution could

allow for the formation of a more ordered phase to retain a favourable free energy. The increase in the solvent viscosity would also reduce the ability of micellar aggregates to diffuse through the solvent in bulk, allowing for the intermediate hexagonal phase to be observed.

The formation of cylindrical micelles cools via a hexagonal phase before forming the lamellar phase at thermodynamic equilibrium. The fluid-to-gel phase transition can be considered as two steps (Fig. 3): the formation of the hexagonal phase from condensation or packing of the cylindrical micelles, and fusion of the hexagonal phase to form the lamellar phase. The first step is driven by the reduction in the kinetic energy allowing for the intermediate hexagonal phase to be observed. The second step in the fluid-to-gel transition comprises the hexagonal-to-lamellar transition, where the molecular shape transforms from wedge to rod, noted by Caffrey [14], based on the packing parameter [50, 51]. The metastable hexagonal phase is kinetically stabilised due to the preference of SDS molecular packing in a conical-wedge molecular shape, as per aqueous solutions with a packing parameter \( \sim 0.3 \). The tail volume tends to decrease with increasing temperature, a result of the increased kinetic energy allowing for the intermediate hexagonal phase to be observed.
derive from the slightly reduced solubility of Na\(^+\) counterions at the lower temperature. Initial cooling would drive more Na\(^+\) to condense on the micelle surface, screening charge repulsion between the headgroups and allowing micelle condensation into the 2D hexagonal phase. Further slow diffusion of Na\(^+\) and condensation on the micelle surface would further shrink the headgroup size. Along with the increased tail volume at the lower T (as discussed above), this would further increase the effective packing parameter, until the lamellar phase is favoured.

Our results here, in combination with our previous observation of the SDS-in-glycerol gel [31], have shown that the gelation mechanism of surfactants in polar-nonaqueous media is more complex than previously thought. Our previous results of SDS in glycerol showed a microfibrillar gel phase with lamellar nanostructure [31] at room temperature, transitioning to a fluid phase consisting of cylindrical aggregates interacting via a Coulombic pair potential. However, here we report the presence of a metastable intermediate phase (Fig. 3), which is kinetically stable as a result of decreasing entropy and increasing bulk viscosity. It is an important consideration in the gelation mechanism, offering insights to further elucidate the self-assembly mechanism of surfactants in nonaqueous H-bonding rich media. This mechanistic understanding has important implications in a variety of applications, for instance in formulations where viscous polar-nonaqueous media are prevalent.

Author contributions

L.M. and W.H.B. designed experiments and secured neutron beam time.
L.M., M.C.S., R.S., and WHB conducted SANS experiments.
L.M. and W.H.B. analysed the SANS data.
R.S., A.J.J., P.B., and W.H.B. supervised L.M.
L.M. wrote the first draft of the manuscript and then co-wrote the subsequent versions of the manuscript with W.H.B., with all the other co-authors contributing to the editing and reviewing of the final draft of the manuscript.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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