



Cite this: *Soft Matter*, 2022, 18, 9133

Received 6th July 2022,
Accepted 15th November 2022

DOI: 10.1039/d2sm00827k

rsc.li/soft-matter-journal

The effects of surfactant and oil chemical structures on self-assembly in apolar media

Adhip Rahman  and Julian Eastoe*

The thermodynamic and chemical structural aspects of surfactant self-assembly in aqueous systems have been much studied. On the other hand, for oil–water interfaces the effects of chemical structures of surfactants and solvents have received less attention. This review focuses on the surfactant chemical effects in low dielectric solvents, in particular formation and properties of surfactant films at oil–water interfaces. For this purpose, reversed micelles (RMs) and water-in-oil (W/O) microemulsions (μ Es) serve as model systems, since electrostatic effects are minimized, allowing a focus on chain architecture of the surfactants and oil solvents themselves. It is noted that chemical structure can have profound effects on stability and self-assembly, suggesting a possibility of identifying unified chemical principles for designing and formulating systems across various thermodynamic conditions.

1. Introduction

Most colloidal systems of interest comprise charged particles in aqueous media; theoretical understanding of colloidal stability of such systems is well established and pertains to systems using high dielectric solvents. The celebrated DLVO theory provides basis for understanding stability in such common systems.^{1a,b} The principal parameter attributed to solvent in DLVO theory is the dielectric constant ϵ , and, of course, the solvent of interest is nearly always water. Hence, it can be argued that in the DLVO framework the solvent merely serves as

a benign medium, the purpose of which is to support charge (*i.e.* accommodating ions). Hence, the effect of solvent chemical structure on stability of aqueous colloidal systems are not explicitly considered.

Consider now an alternative class of colloidal systems utilizing low dielectric oily solvents. These systems are stabilized predominantly by steric interactions, rather than electrostatics as for charged aqueous colloids exhibiting different physical properties. It is true that electrostatic interactions are found in such non-aqueous colloidal systems,^{1c} but these are generally longer range and with weaker interactions compared to aqueous systems. There is a wide variety of different oily solvents with diverse chemical structures, which can be mixed to compose solvent mixtures exhibiting wide ranges of

*School of Chemistry, University of Bristol, Cantock's Close, Bristol BS8 1TS, UK.
E-mail: Julian.Eastoe@bristol.ac.uk*



Adhip Rahman

Adhip Rahman is a PhD student in Bristol – starting in 2019; prior to this, Adhip graduated with an MS (Masters) in Chemistry from University of Dhaka, Bangladesh (and, semi-professionally explored realms of literature and classical music).



Julian Eastoe

Julian Eastoe is Professor of Chemistry at the University of Bristol. His research interests span colloid and interface science, surfactant chemistry and applications of neutron scattering. He was awarded the Rideal Medal from the Royal Society of Chemistry and Society for Chemical Industry in 2007 for “distinction in colloid or interface science”, and the 2015 ECIS-Solvay medal from the European Colloid and Interface Society “for original scientific work of outstanding quality”. He is a Co-editor of the Journal of Colloid and Interface Science.



physical properties and molecular interactions. Such mixtures offer a landscape wider than usually available in aqueous colloids. Hence, for non-aqueous low dielectric colloidal systems variations in solvent chemical structure are expected to be significant considerations. Water-in-oil (W/O) dispersions (emulsions) or microemulsions (μ Es) (thermodynamically stable systems comprising water-immiscible oils) are common classes of non-aqueous colloidal systems. For these systems there must be intimate molecular contact between the stabilizing surfactant (films) and the “continuous phase” external solvents. Therefore, it can be expected that solvent-specific effects may become significant for system phase-stability, structure, and properties. In this review, a fresh perspective on solvent–surfactant specific effects is offered – with a view to improving the understanding of such systems and facilitating their formulation.

Typical surfactants comprise hydrophilic (polar) heads and hydrophobic (non-polar) tail group(s) – such chemical factors lead to their dissolution in aqueous or non-aqueous solvents. Another consequence of the surfactant molecular structure is that surfactants self-assemble in aqueous and non-aqueous media forming micelles, swollen micelles/reverse micelles (RMs) (*i.e.* microemulsions, μ Es) or emulsions. These self-assembly structures are in the size range of between a few tens or hundreds of angstroms (\AA) and have well-defined morphologies (spherical, ellipsoid, cylindrical, disk shaped and so on). In water (or polar media) the hydrophilic heads orient towards the solvent bulk, whereas hydrophobic tails tend to point inwardly, away from the solvent. On the other hand, in oily solvent media reverse micelles (RMs) may form – in which the surfactant tails point outwards towards the solvent and the heads are buried in the reverse micellar core.^{1a,b,2} An important aspect of such self-assembly structures is that they can accommodate substances of opposite polarity relative to the dispersion media inside the RM cores – for example, solubilizing a polar liquid, water. Such ternary systems may be called μ Es if they form spontaneously, are optically transparent and thermodynamically stable dispersions; in such systems, nano domains (or droplets) containing one fluid dispersed in the other – with the two liquids being separated by surfactant monolayers.^{2–5} To be specific, μ Es can be understood as surfactant monolayers thermodynamically stabilizing an otherwise unstable oil–water (O–W) interface where the oil is immiscible with water.^{2,4} As such, surfactants straddle the oil–water interface, and there is necessarily intimate contact between surfactant tail groups and the continuous-phase oil, so that solvent-specific effects can be expected. They show interesting variation in interfacial as well as bulk properties and have been recognized for their applicability in controlling morphology of nanomaterials in the fields of catalysis, personal care, drug delivery, enhanced oil recovery, lubrication, and cleaning technology.^{6–9}

The effect of physical variables such as temperature, pressure, concentration, and composition on surfactant self-assembly in polar or non-polar media has been studied in much detail.^{10,11} While effects of surfactant and solvent chemical structure have been studied for some decades, the

Table 1 Chemical structures of the surfactants and solvents discussed

Molecules	Generic chemical structure
Classes of surfactants	
Bis(2-ethylhexyl)sulfosuccinate salt or Aerosol-OT (M^{n+} = metal ions)	
Dialkyldimethylammonium bromides (C_nC_m DAB)	
Alkyl polyglycol ethers (C_iE_j)	$C_i-O [CH_2CH_2O]_j-H$
Solvents	
Linear alkanes ($n = 0-14$)	
Aromatic hydrocarbons (benzene, toluene and alkylbenzenes)	
Cycloalkanes	

vast majority of these are scattered efforts. Early theoretical models interpreted surfactant self-assembly (micelles and μ Es) entirely from the context of thermodynamics.^{11,12} However, whether surfactant and solvent-chemical architecture exert any significant control over μ E thermodynamic stability and morphology of self-assembled structures has not been considered in exhaustive detail.

This review aims to compile, summarize, and compare the most important results and interpretations related to the chemical effects on surfactant self-assembly in low dielectric solvents. μ Es, water-in-oil μ Es (W/O- μ Es) in particular, are chosen instead of normal emulsions as model colloidal systems for discussion. There are two reasons – (i) the former systems are thermodynamically stable, which is not the case for emulsions, and (ii) W/O systems make it possible to focus entirely on the chemical structure effects of surfactant hydrophobic chains and solvents owing to their intimate contact.^{12a} Surface films of surfactants, wherein the head groups are oriented in intimate contact with diverse solid phases,^{12b} with and without water associated with such head groups are also important but are not discussed in this review. Systems primarily stabilized by the dichain anionic surfactant, Aerosol-OT (AOT) and its analogues, are extensively discussed here; in addition, cationic dialkyldimethylammonium (C_nC_m DAB) and nonionic polyoxyethylene glycol alkyl ether (C_iE_j) surfactants are also considered. Table 1 lists the types of surfactants and nonpolar solvents covered here.

2. Understanding surfactant–solvent thermodynamic stability: phase behaviour, curvature, and interfacial tension

The macroscopic phase-stability of three-component μ Es containing a surfactant, water and an oil is accounted for in terms



of ternary phase diagrams. A typical ternary phase diagram encompasses the whole composition range. For more than three components (hydrotrope or cosurfactant as an additional component) a pseudoternary phase diagram is often used.^{4,13,14} The phase diagram covers all possible phases including non-transparent, non- μ E phases – since not all combinations result in single-phase μ Es. An example of a typical phase diagram of an AOT-based μ Es is given in Fig. 1. The location of different boundaries in a phase diagram depends on the chemical nature of the μ E-components and thermodynamic conditions (temperature and pressure).^{15–18} A phase diagram of a typical three component system may comprise single-phase L_1 - μ E (oil-in-water (O/W)), L_2 - μ E (water-in-oil (W/O)), W/O- μ E phase separated from excess water (Winsor II), O/W- μ E phase-separated from excess oil (Winsor I), and a μ E in equilibrium with excess oil and water (Winsor III).^{3,4}

Normally, surfactant-coated droplets (containing water in W/O- μ Es or oil in O/W- μ Es inside droplets) in these systems are spherical in nature, but under certain circumstances, the μ Es may be viscoelastic – surfactant curvature in such systems is often complex (ellipsoidal, cylindrical or polymerlike).¹⁵ In addition, non- μ E phases (such as gels, liquid crystalline (LC) lamellar, cubic or hexagonal phases) and bicontinuous (L_3) μ Es may also appear.^{19–21}

2.1. Water solubilization and W_{\max}

Before proceeding to the discussion, the difference between RMs and W/O- μ Es should be clarified; RMs form when surfactants self-assemble in nonpolar organic solvents; the core may be “dry”, *i.e.* no water (Section 3.1 will discuss dry RMs in detail) or may have small-quantity of water in which case the water molecules interact strongly with the hydrophilic head-groups.^{22b,e} W/O- μ Es, as pointed out in the introduction, are swollen with water so the water solubilization by the RM cores is much higher; in addition, the core-water resembles regular bulk-water evidenced by spectroscopic studies.²²

A simple and convenient way to understand W/O- μ E macroscopic phase-stability is in terms of water solubilization capacity W (where $W = [\text{water}]/[\text{surfactant}]$). W/O- μ Es droplets can be swollen by water up to a limit; physical variables such as temperature, salinity and cosurfactant affect the extent of W (maximum water solubilization, W_{\max}).^{23a} Shah *et al.*^{23b,c} attempted to understand W/O- μ E phase transitions in terms of the correlation between W_{\max} and O-W interfacial curvature (details can be found in Section 2.2). According to these publications, the balance between the interfacial contribution towards a limiting curvature and droplet-droplet attractive interactions leads to bending of a flat film – leading to μ E-formation. For rigid interfaces (see Section 2.3), the droplet radius in W/O- μ Es does not exceed a threshold “radius of spontaneous curvature, ρ_0 ” (for details, see the later), as exceeding ρ_0 results in phase separation of water from the μ E-droplets (*i.e.* WII-systems).²³ The interface, nevertheless, becomes more flexible if ρ_0 increases (*i.e.* water-solubilization increases in L_2 - μ Es).^{23c} Moreover, on increasing the water content, droplets in L_2 - μ Es tend to achieve a “critical radius”

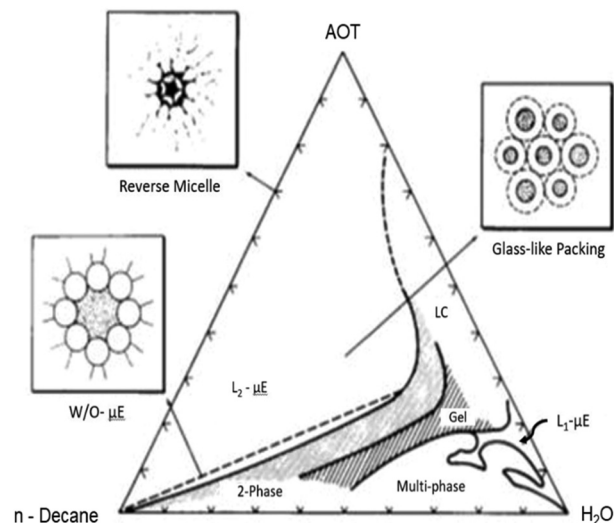


Fig. 1 Ternary phase diagram of AOT/H₂O/*n*-decane μ E along the entire composition range for each of the three components (at room temperature and pressure); the phases are described in the text. Insets show various nanostructural ensembles in the single-phase L_2 - μ E region. The units of the axes are represented as wt%. The figure was reprinted from ref. 20. Copyright 1986, with permission from Elsevier.

– above which macroscopic phase separation occurs. Attempts were made to link solvent chemical-structure with ρ_0 and the critical radius^{23b,c} (Fig. 2). The inference was that ρ_0 (linked to W_{\max}) would be correlated to solvent molecular volume – given no surfactant structural variation has been considered.^{23,24} Some early works assumed that the ability of hydrocarbon solvents, due to variation in their molecular volumes, to penetrate surfactant monolayers would govern W_{\max} by controlling surfactant film-curvature (more on this later) and, thus, droplet-droplet interactions.^{17,25,26} Any correlation of W_{\max} directly to the solvent chemical architecture was, however, unclear.

A more convenient approach to probe W_{\max} as a function of surfactant-solvent chemical variation is through temperature-composition phase-diagrams. This will be covered in Section 2.3 and onwards.

2.2. Packing and surfactant film-bending

The effect of surfactant chemical structure on interfacial film topology was discussed by Israelachvili.²⁷ According to these arguments, surfactant chemical structure was understood in terms of the following parameters:

- Surfactant head-group area (a_h) owing to repulsive (steric and/or electrostatic) interactions between headgroups
- Effective hydrophobic tail volume (v) and extended tail length (l_c) attributed to a balance between attractive hydrophobic interactions and steric interactions between the tails

Overall, the quantity v/a_0l_c , known as the packing parameter (p) is consistent with W/O- μ E formation for $p > 1$. This predicts that for a given surfactant chain length, a small a_0 and high v favours reversed curvature W/O- μ Es. High v can be achieved by means of introducing branching, unsaturation, or double chains into surfactant hydrophobic tail regions.^{4,27} The role



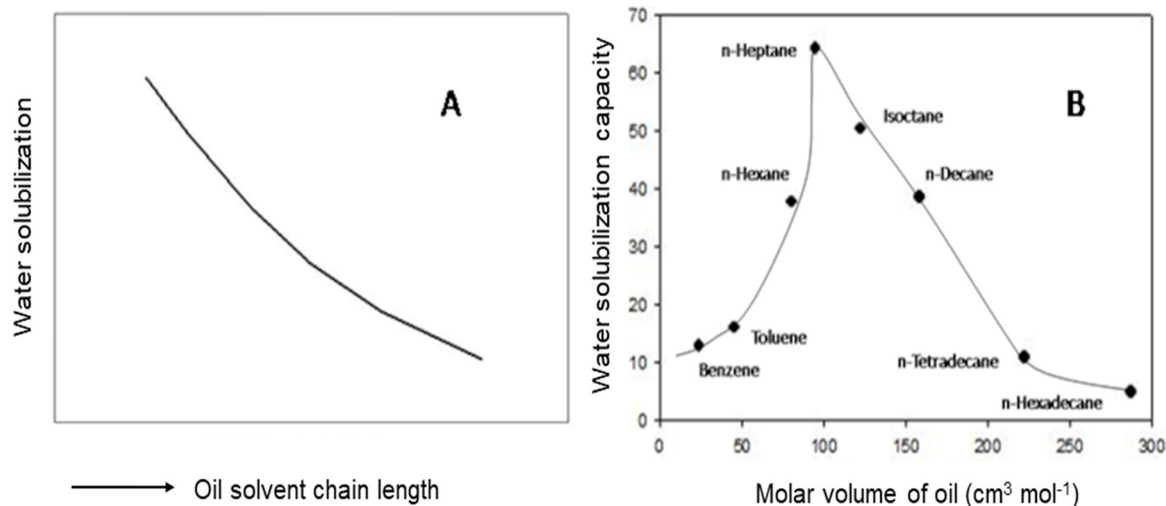


Fig. 2 (A) A schematic illustration showing the relationship between water solubilization limit of L₂-μEs and oil solvent chain length, the arrow along the X-axis indicates increasing (oil chain length); the region below the curve represents L₂-μEs, and above the curve indicates μE-phase separation; (B) maximum water-solubilization (W_{\max}) in AOT-based W/O-μEs as a function of the solvent molar volumes at room temperature and pressure. The scheme in (A) was based on Fig. 2 of ref. 23c. Copyright (1987), with permission from Elsevier. Data for (B) were reprinted (adapted) from ref. 23b. Copyright 1987. Reproduced with permission from American Chemical Society.

of solvent was discussed mostly in the context of the penetration capacity – the greater the solvent penetration the more favoured would be the W/O-μE formation. However, extensive contrast variation small-angle neutron scattering (SANS) studies, designed specifically to assess penetration of oil molecules into the hydrocarbon regions of surfactant films, suggested that solvent penetration is likely to be a subtle effect; moreover, not all types of solvent molecules penetrate through the surfactant tip-regions, and solvent penetration shows a two-way solvent-surfactant chemical-dependency.^{28,29} This aspect will be enlarged on in Section 3.4.

The first realistic model for interpreting properties of surfactant-films at O-W interfaces from the context of film-mechanical properties was developed based on the early theoretical concepts for film-bending and rigidity in liquid crystals by Frank, and later Helfrich for amphiphilic systems.³⁰⁻³² The model considers conformational degrees of freedom of surfactant hydrophobic tails dispersed in a “seemingly” gas-phase, *i.e.* solvent molecules were considered unimportant. Conformational free energy of total film bending can be expressed as

$$\frac{f}{a} = \frac{1}{2}k(2c - c_0)^2 + \bar{\kappa}c^2 = (2\kappa + \bar{\kappa})c^2 - 2kc_0 + \frac{1}{2}kc_0^2 \quad (1)$$

where c_1 and c_2 are the principal film-curvatures, c_0 is the spontaneous (or natural) curvature (for a perfect spherical droplet, $c_1 = c_2 = c_0$) – the curvature formed by a surfactant film at the presence of equal amount of oil and water in a system, a is the area per surfactant molecule, κ is the mean elastic bending constant (or modulus), $\bar{\kappa}$ is the so-called saddle-splay constant (or Gaussian modulus) and c is the mean curvature – which is related to principal radii of curvature ρ_1 and ρ_2 at any

given point on the surfactant film ($\rho_1 = 1/c_1$ and $\rho_2 = 1/c_2$) by

$$c = 0.5 \left(\frac{1}{\rho_1} + \frac{1}{\rho_2} \right) \quad (2)$$

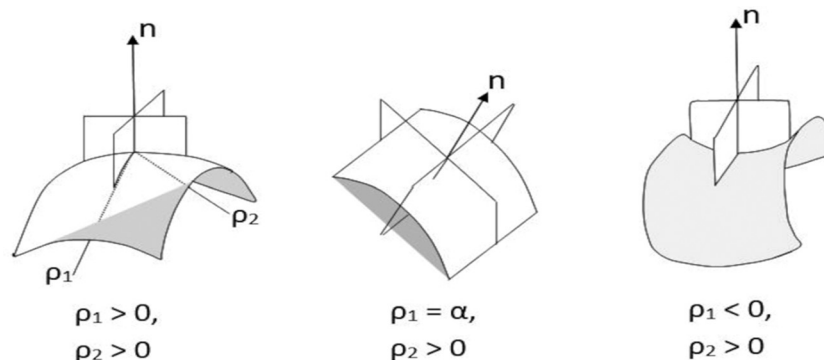
Scheme 1 illustrates various curved planes defined by the abovementioned parameters. This model predicts that, for a monolayer with a given average area per molecule, κ gradually increases with increasing surfactant chain length. However, κ markedly decreases if fractions of the long hydrophobic chains are substituted by shorter chains, whereas $\bar{\kappa}$ is close to zero. This means that the lateral pressure among the longer chains lessens due to shorter chains acting as “spacers”.

The theoretical phase-diagram arising from these considerations, Fig. 3, shows regions of various structural forms in W/O-μEs as a function of ρ/ρ_0 and $\bar{\kappa}/\kappa$, where ρ_0 is spontaneous radius of curvature and ρ is radius of curvature. This is related to c_0 as $\rho_0 = 1/c_0$. ρ has an inverse relationship with mean curvature c and can be expressed by eqn (2) above.⁴ More importantly, ρ may related to the surfactant chemical architecture by

$$\rho = \frac{3\delta\phi}{\phi_s} \quad (3)$$

where ϕ is the volume fraction of the surfactant-coated water-droplets (volume fraction of surfactant + dispersed phase – assuming all surfactant molecules reside at O-W interfaces), δ is surfactant chain length and ϕ_s is surfactant volume fraction. The predicted structural transitions spherical → cylindrical → lamellar involve the decrease of ϕ/ϕ_s or the increase of ϕ_s . Notably, at a given ϕ/ϕ_s the parameter δ signifies that ρ is proportional to the surfactant chain-length (*i.e.* a notable chemical structural effect), so there is a trade-off between the bending constant and curvature.³³ Cates *et al.*³⁴ followed up these ideas, stating that the key to stable μEs over a wide-range





Scheme 1 Depiction of surfactant films for different curved planes (from left to right – convex, cylindrical and saddle-shaped, n is normal vector to the plane at an intersection-point); ρ_1 and ρ_2 are the two radii of curvature.

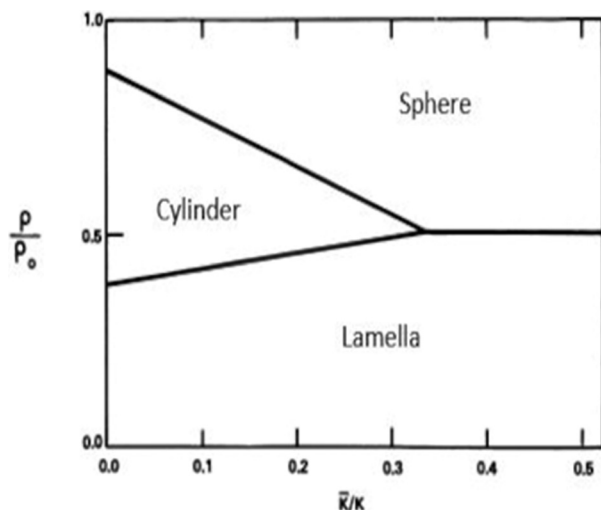


Fig. 3 Theoretical stability regions of lamellar, cylindrical, and spherical phases of μ Es as a function of bending constant ratios ($\bar{\kappa}/\kappa$); all the parameters are described in the text. The figure was reproduced from ref. 33 (S. A. Safran *et al.*, *J. Phys. Lett.*, 1984, **45**, 69) with permission.

of surfactant concentration is to have κ as small as possible (a few $k_B T$, the exact number depends on the balance between κ and c_0); in addition, κ and surfactant chain length are directly proportional. Experimental works^{35,36} on cationic surfactants bearing two linear chains $C_n C_m \text{DAB}$ (Table 1), zwitterionic phosphocholine and nonionic $C_i E_j$ -based μ Es (discussed in Section 2.5)^{75,76} supports this; however, W/O- μ Es with branched chain surfactants might not comply with this model – since the model is for linear chain surfactants.³⁷ An aspect related to this will be discussed later in Sections 2.6 and 3.2.

This model explains a range of cylinder-to-sphere transitions which have been observed in diverse W/O- μ E systems – for example cationic $C_n C_m \text{DAB}$ W/O- μ Es in cyclohexane (cylindrical aggregates until $W = 7$ and spherical aggregates at $W = 10$, $[\text{surfactant}] \approx 0.050 \text{ M}$), AOT containing divalent cations such as Mg^{2+} , Co^{2+} and Ni^{2+} (for the effect of counterions see Section 4) and mixtures of AOT and hydrotropes.^{38–40} The interpretations and models discussed above suggest that there

indeed are intimate links between chemical structure of the components (oils²¹ and surfactants^{26–30}) and phase stability, O-W interfacial curvature and structure of typical W/O- μ Es.

2.3. Chemical-control over phase behaviour: AOT-based μ Es

As it was suggested by earlier theoretical models, solvent penetration capacity into the surfactant hydrophobic tips (di-chain surfactants in particular) could influence μ E-phase stability and structures. However, as mentioned above and below, direct observation of surfactant shells by contrast variation SANS indicated this effect to be of limited significance.^{28,29} Nevertheless, here it is worth revisiting how the solvent chemical architecture affects phase-transitions.

Before discussing anionic AOT-based μ Es, it should be noted that each of the AOT-hydrophobic chains contains eight carbon-atoms and the longer of the two chain fragments contains six carbon-atoms. It was pointed out that the single-phase L_2 - μ Es are more ubiquitous in linear alkane solvents having lower alkane carbon number (ACN).⁴¹ An extensive phase-mapping of AOT/ $\text{H}_2\text{O}/n$ -hexane μ Es showed that in n -hexane, the amount of solubilized water in L_2 - μ E domains can be as high as 40 wt% (at 293 K); on increasing temperature, the L_2 -phase, however, narrowed.^{42,43} Meanwhile, comparison between the L_2 - μ Es in cyclohexane and the aromatic solvent p -xylene showed that the L_2 -domain in cyclohexane was wider than that of in p -xylene.^{44–47} Comparative phase-mapping between isooctane and p -xylene (both having eight carbon-atoms) showed that the L_2 -phase is more extensive in isooctane.^{47–49} Such an apparent disparity in the macroscopic phase-stability due to solvent chemical structure was also demonstrated by Hall *et al.*⁵⁰ in flow birefringence studies. AOT- μ Es in n -decane showed interesting phase behaviour; W_{max} is not significantly high and bicontinuous μ Es occupy a significant portion of the ternary phase diagram; the multiphase regions are narrow, and transitions from L_2 - to LC-phases end in a gel-like phase.^{51,52a} The L_2 -phase further narrowed by replacing n -decane (C_{10}) for n -dodecane (C_{12}).^{52b}

Consideration of temperature with a ternary composition phase diagram leads to a three-dimensional phase-prism, which can become rather difficult to interpret.^{4,23a} A more simplified approach is to reduce the degrees of the 3-D phase



prism into a two-dimensional composition-temperature phase diagram.⁴ In this representation, μ E-composition may be represented by W , and an example phase diagram of AOT W/O- μ Es in n -octane is shown in Fig. 4(A).

It should be noted that throughout this review, the phase-transitions discussed are macroscopic phase-transitions, *i.e.* the phase-changes can be seen by naked eye. In this sense, the macroscopic phase-transitions account for the phase-diagrams shown in Fig. 4. The phase-transitions occur in such systems involve Winsor-type phase transitions⁴ (it is true that beyond the macroscopic phase-realms, μ Es show microscopic phase-transitions – the so-called “second-order phase transition” – as well; this has been quantified by Texter^{53a,b} and Antalek *et al.*^{53c} by droplet-apparent diffusion coefficient measurements in AOT-based W/O- μ Es. It was shown that W/O-droplet to a fractal-like bicontinuous phase transition could be quantified by the order parameter). The funnel-like region covered between the lower and upper temperature phase-boundaries (T_L and T_U , respectively) is of interest as this represents the stability extent (*i.e.* the region in temperature-composition phase diagrams covered by single-phase transparent L_2 -systems) of the single-phase L_2 - μ Es – otherwise termed as the “ L_2 - μ E domain size” for convenience. The lower temperature boundary, T_L , accounts for Winsor II to transparent L_2 - μ E phase transition, whereas the upper-temperature boundary, T_U , accounts for the so-called “cloud-point” phase separation, *i.e.* L_2 -to-cloudy phases. The use of “cloud point” in W/O- μ Es may have different meanings based on appearance of the systems; often when the cloud-point has been reached, the systems turn bluish and more viscous, and over the time, a surfactant-rich phase-separates from the L_2 - μ Es.³⁷ For other systems, close to cloud points the L_2 - μ Es can separate into two

coexisting μ E-phases²⁰ (*i.e.* a critical-type separation, this will be discussed in detail in Section 3.6). In general, the volcano-like L_2 -region normally shifts towards lower temperature on increasing the ACN of linear n -alkane solvents.⁵⁴

The L_2 - μ E domain size and its relationship to chemical variation of hydrophobic tail architecture (with AOT analogue surfactants) was studied by Nave *et al.*^{55–57} In this work, a class of straight chain and branched chain AOT-analogues were custom synthesized (example structures see Fig. 7). For un-branched linear hydrophobic chains, μ Es did not form with the surfactants alone, but rather required short-chain alcohol-co-solvents (*i.e.* four component systems) to stabilize transparent L_2 -phases. This is consistent with the work of Safran *et al.*³³ who outlined the need for sufficient entropic freedom of the hydrophobic chains.⁴⁴ A clear chemical structural effect on temperature-guided phase stability (*i.e.* the L_2 - μ E domain sizes in the temperature-composition phase diagrams) was seen, in general, shortening the surfactant alkyl chains shifted the L_2 -funnel towards higher temperature, and *vice versa*. On the other hand, introduction of chain-branching brought the surfactants into line with normal AOT, and three component μ Es could be stabilized. Interestingly, larger L_2 - μ E domains at higher temperatures were especially noted for surfactants bearing brush-like trimethyl *hedgehog* chain-tips. (Fig. 4B).⁵⁵

A striking and, un-subtle, manifestation of a highly specific surfactant-solvent chemical structural effect was observed. Upon introducing aromatic fragments onto the AOT chain-tips, μ Es only formed in aromatic solvents, such as toluene, but did not form in linear alkane solvents. These “phenyl-tipped” AOT-stabilized W/O- μ Es in toluene were rather less sensitive to temperature – compared to the normal AOT-based μ Es in toluene.⁵⁶ It is notable that the temperature-dependent

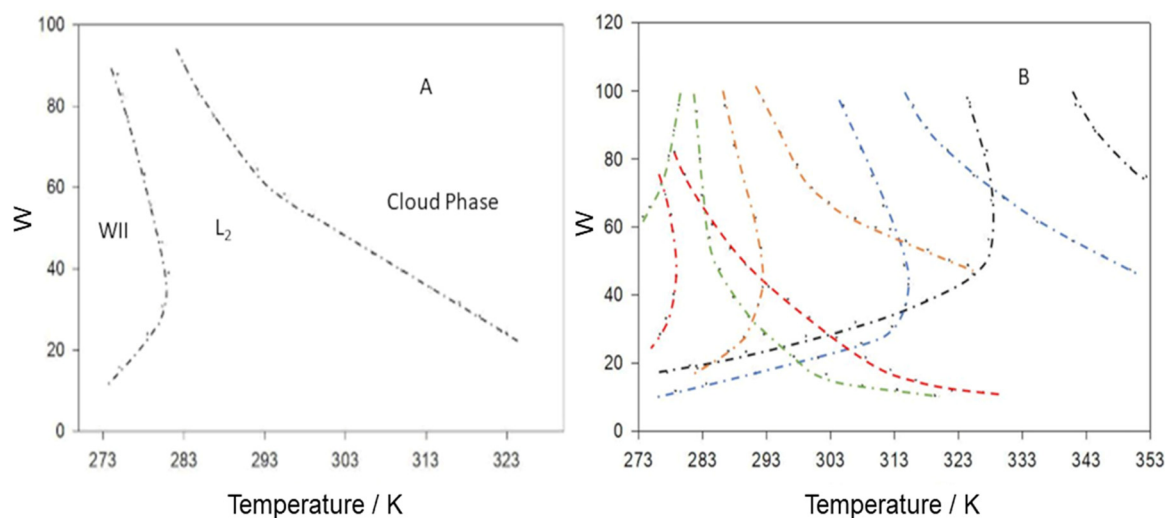


Fig. 4 (A) AOT-based W/O- μ E temperature-composition phase-diagram in n -octane; WII, L_2 and cloudy phases are described in the text, (B) effect of structural variation at the AOT hydrophobic tails – colours indicate AOT analogues: (green) AOT3, (red) AOT5, (orange) AOT1 (*i.e.* normal AOT), (blue) AOT2 and (black) AOT4. Chemical structures of AOT1–AOT5 are shown in Fig. 7. The left-side boundary for each of the phase-funnels represents a WII-to- L_2 transition, whereas right-most boundary represents L_2 -to-cloudy phase (defined below) transition. Pressure: 1 bar, [surfactant] = 0.100 M. Data in (A) were reproduced (adapted) from ref. 23a. Copyright (1991), with permission from Elsevier. Data in (B) were reprinted (adapted) from ref. 55. Copyright 2000. Reproduced with permission from American Chemical Society.



L_2 - μ Es domain sizes could be correlated to the aqueous phase solubility of each of the surfactants.⁵⁵ These experimental observations show the importance of chemical structure for μ E-stability, supporting and amplifying the earlier literature discussed above.^{21,26–30}

2.4. Cationic surfactants

Surfactants based on quaternary dialkyldimethylammonium (C_nC_m DAB) cations (Table 1) having double-chain architecture, like the anionic AOT, have been much studied.^{58–68} The two-tailed structure of these cationics indicate a preference for stabilizing W/O- μ Es – despite them being only very weakly soluble in water as well as hydrocarbon solvents (<1 wt%).^{58,59} Like the AOT-analogues, phase behaviour of C_nC_m DAB- μ Es was observed to be highly oil and surfactant-chain specific;^{58–62} for the common symmetric di-chain surfactant di- C_{12} DAB (DDAB), μ E-formation shows a systematic “oil-specificity”.^{58,61} The onset water content at which μ Es form was ~ 6 wt% for n -hexane, rising to 26 wt% for n -tetradecane; however, W/O- μ Es in n -tetradecane could not be formed at low surfactant concentration. Another study compared n -alkanes and alkenes with the same carbon numbers (C_6 to C_{14}); for each of the pairs, the onset volume fraction of water to form L_2 -phases was lower for the alkene than the corresponding n -alkane.⁶³ A comparison between cyclic and linear n -alkanes showed that the cycloalkane narrows the single-phase region compared to the linear alkane; for cyclohexane, the onset water content for W/O- μ E formation (2 wt%) was found to be much lower compared to its linear chain homologue n -hexane.⁵⁹ In addition, microstructure topology in di- C_{12} DAB-based μ Es was found to be correlated with oil-solvent chemical nature – surfactant films were more rigid in short chain n -alkanes.⁶⁴ The case of aromatic solvents is interesting – since di- C_{12} DAB is highly soluble in toluene (>30 wt%). Unlike n -alkanes or cycloalkanes, in which detectable W/O-spherical droplets could be formed, pulse-field gradient NMR experiments carried out by Olla *et al.*^{65,66} showed no evidence for droplet formation in aromatic solvents such as toluene or fluorinated toluene. Rather, in oil-continuous systems, μ E phase-transitions followed from discrete hydrated molecular clusters to connected bilayers on increasing droplet volume fraction (surfactant + water).⁶⁶ Fluorinated alkanes, on the other hand, induced formation of kinetically stable emulsions, but not thermodynamically stable μ Es.⁶⁶

Surfactant–solvent structure-property relationships for asymmetric chain C_nC_m DAB ($n \neq m$) surfactants were studied in aromatic solvents; uneven chain-lengths ($m > n$)^{67a,b} and incorporation of phenyl-groups in one of the chains^{67b} resulted in W/O- μ Es of W as high as 80 in aromatic hydrocarbons. Moreover, light-scattering and interdroplet exchange kinetics results^{67c} revealed a two-fold effect; for a given surfactant chain-length, inclusion of polar fragments into the aromatic solvent resulted in reduced interdroplet collision-rate constants, for example, in chlorobenzene (on the order of $10^8 \text{ M}^{-1} \text{ s}^{-1}$) compared to non-halogenated benzene ($\approx 10^9 \text{ M}^{-1} \text{ s}^{-1}$). This was found to be equivalent to shortening the hydrophobic chains by around three carbon-atoms, since shortened hydrophobic

tails induced high collision rate and enhanced aggregation in a solvent and experimental condition.

Surfactant chemical-effects with the C_nC_m DAB class was further explored by Warr *et al.*⁶⁸ They observed that the chain-melting temperature of C_nC_m DAB determined whether the surfactant formed μ Es between a given temperature window or not. Chain-melting temperature is related to hydrophobic-chain fluidity and on increasing the chain carbon-number chain-melting temperature increases. For di- C_{12} DAB it was 289 K, whereas for di- C_{14} DAB it was 304 K. Indeed, μ Es formed beyond 304 K in a range of linear hydrocarbons for the latter surfactant, whereas for the former μ Es were found at around 298 K and beyond. The oil specificity with di- C_{12} DAB was also evident, the minimum water content required to form μ Es increases on increasing solvent ACN. The oil-specificity was clearly observed for surfactants with chain-length asymmetry such as C_8C_{16} DAB. For instance, while for C_8C_{16} DAB oil-continuous W/O- μ Es formed in n -hexane, in n -decane the μ Es were bicontinuous with zero mean curvature at O–W interfaces. Nevertheless, literature on cationic μ Es suggests that the surfactant film properties and water solubilization mutually depend on solvent as well as surfactant hydrophobic tail chemical nature – similar to the W/O- μ Es containing anionic AOT and its analogues.

Hence, for two of the common classes of surfactants discussed above, solvent structural specific effects are major features related to μ E-phase stability.

2.5. Nonionics

For historical reasons,^{15,69–73} the phase-behaviour for nonionic C_iE_j (Table 1 for chemical structures) μ Es is presented in a manner different from the ionic μ Es; temperature effects are often studied in terms of Winsor I (O/W- μ E) to surfactant-rich bicontinuous state through to Winsor II (W/O- μ E) transitions. The macroscopic phase-transitions are commonly presented in terms of the so-called “fish-cut like phase-diagram”.¹⁵ A schematic representation can be found in Fig. 1 of ref. 15 – the region covered by the upper-temperature and lower-temperature boundaries corresponds to a three-phase bicontinuous Winsor III- μ E through which transitions from O/W to W/O- μ Es take place. For short chain C_iE_j s (such as C_6E_5), the temperature-boundaries shift towards lower temperature on reducing ACN of linear alkane solvents.⁶⁹ Moreover, on increasing C_j , the three-phase “fish-cut” region requires a lower amount of surfactant. An important factor in these temperature-composition phase diagrams is the phase-inversion temperature (PIT). At a given surfactant concentration, this is the temperature at which a transition from O/W- μ Es to W/O- μ Es takes place.^{70,71} Shinoda *et al.* extensively studied PITs in nonionic systems; in general, the PIT depends on the solubility of a C_iE_i in any given hydrocarbon solvent, and increased solubility of C_iE_j leads to lower PIT.^{71–73} Interestingly, surfactant–solvent chemical effects are seen, for instance if C_iE_j surfactants contain a phenyl group, then the PIT for O/W to W/O transition is lower in aromatic solvents as compared to linear hydrocarbon solvents.^{71,72}



These C_iE_j surfactants are of interest because both the hydrophilic and hydrophobic fragments can be simultaneously and sequentially varied. According to a molecular model proposed by Paunov *et al.*,⁷⁴ a C_iE_j surfactant can be treated as a diblock copolymer containing a hydrophilic (EO) and a hydrophobic block (C_i); the curvature and bending constants of surfactant films arise due to total contributions from both blocks. Therefore, in a C_iE_j - μ E phase (W/O or O/W), interfacial properties of C_iE_j surfactant-films vary in a regular fashion as a function of both C_i and E_j .^{75,76} Extensive oil-water (O-W) interfacial tension (IFT) measurements by Sottmann *et al.*⁷⁵ revealed that the limiting surfactant molecular area at the O-W interface depends strongly on the length of EO fragment (E_j) and is essentially independent of the solvent ACN as well as surfactant chain-length C_i . Despite this, contrast-variation SANS and O-W interfacial tension $\gamma_{o/w}$ measurements by Gradziński *et al.*⁷⁶ revealed that the bending constant ($2\kappa + \bar{\kappa}$) and $\gamma_{o/w}$ increases with surfactant chain-length C_i , so that longer chain C_iE_j -films are more “rigid”. This trend was echoed in studies^{28,29} with cationic C_nC_m DAB W/O- μ Es, as will be discussed in more detail in Section 3.4.

2.6. Very-low oil-water interfacial tension

A widely known application of μ Es is in enhanced recovery (EOR) of crude oils, which is because the thermodynamically stable surfactant monolayers at O-W interfaces reduce O-W interfacial tension ($\gamma_{o/w}$) to very low values – on the order of 10^{-2} to 10^{-4} mN m⁻¹.^{4,77} The history of exploring surfactants at brine-oil interfaces for EOR is extensively covered in the review of Taber.⁷⁷ In fact, in the early years of EOR in the 1970's, the importance of very-low $\gamma_{o/w}$ was recognized to be an important factor and accelerated the development of spinning drop tensiometry (SDT) for experimentally accessing very-low $\gamma_{o/w}$ values.

A standard method for investigating $\gamma_{o/w}$ involves SDT studies with an electrolyte scan using a surfactant aqueous solution $>$ CMC and oil droplet injected into the tensiometer (see Aveyard *et al.*).⁷⁸ Fig. 5 shows a typical IFT dependence of AOT at brine-*n*-heptane interfaces, consistent with Winsor I \rightarrow Winsor III \rightarrow Winsor II- μ Es macroscopic phase transitions.⁵⁷ Since the surfactant is at a concentration \gg CMC, addition of electrolyte changes aqueous-phase surfactant solubility; the minimum $\gamma_{o/w}$ corresponds to the salt concentration (optimal salinity, OS) at which surfactant solubility in both the oil and water is balanced.⁷⁹

In a benchmark series of studies, Binks *et al.* showed the effects of linear solvent ACN;^{78,80,81} on increasing ACN the OS shifts towards higher concentration – meaning equal O-W partitioning needs enhanced “salting out” of AOT. In addition, the minimum in $\gamma_{o/w}$ at the OS tends to increase upon increasing the solvent ACN. Whether this has anything to do with the surfactant hydrophobic chain-length cannot be inferred because only the 2-ethylhexyl chain of AOT has been considered in these works. This observation was correlated to the saddle-splay energy ($\bar{\kappa}$) of AOT films: in Winsor III systems, this energy is positive (ρ_1 and ρ_2 are of opposite signs,

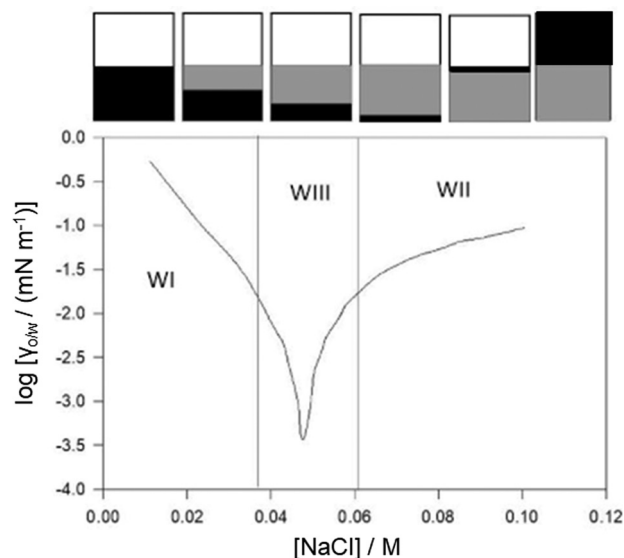


Fig. 5 (top) Cartoon representation of WI \rightarrow WIII \rightarrow WII transitions (white: excess oil, black: μ E-phase and grey: excess water), (bottom) water-*n*-heptane interfacial tension progression as a function of [NaCl], the surfactant is AOT. [AOT] is much higher than the CMC. Temperature: 298 K. Data for (bottom) were reprinted (adapted) from ref. 57. Copyright 2002. Reproduced with permission from American Chemical Society.

Table 2 Optimal salinities (OS) and minimum $\gamma_{o/w}$ for AOT analogues from ref. 57 at brine-*n*-heptane interfaces [surfactant concentration \gg CMCs]; the AOT1–AOT5 chemical structures are shown in Fig. 7

Surfactants	OS/M	$\gamma_{o/w} \times 10^4 / (\text{mN m}^{-1})$
AOT1	0.047	4
AOT2	0.037	6
AOT3	0.062	8
AOT4	0.012	2
AOT5	0.12	10

see Scheme 1 – suggesting presence of saddle-like bicontinuous structures) in *n*-dodecane, close to zero but negative in *n*-decane and negative in *n*-octane (ρ_1 and ρ_2 are of same signs suggesting spheres).⁸² A similar trend was seen for nonionic $C_{12}E_5$ on increasing solvent ACN.⁸³

The surfactant chemical effect on $\gamma_{o/w}$ has also been explored with cationic C_nC_m DABs,³⁵ zwitterionic dichain phosphocholine⁸⁴ and anionic AOT structural analogues.⁵⁷ Two aspects of the AOT hydrophobic architecture were seen to be potentially important in promoting surfactant efficiency at O-W interfaces – (i) brushlike branching at the chain-tips, (ii) absence of branching at the carbon atom immediate to the C–O–C oxygen atom.

Table 2 lists the optimum salinity (OS) and minimum $\gamma_{o/w}$ for various branched AOTs taken from ref. 57 – depicting the chemical effect.^{55,57} For C_nC_m DAB surfactants in W/cyclohexane Winsor II μ Es, $\gamma_{o/w}$ was seen to increase on increasing surfactant hydrophobic chain-length.³⁵ Therefore, for a range of different nonionic, cationic, and anionic surfactants the O-W interfacial properties are notably affected by chemical variation both at the surfactants and solvents.



3. Surfactant–solvent chemical effects on self-assembly

3.1. Dry reverse micelles

As pointed out in Section 2.1, dry reverse micelles (RMs) are surfactant solutions in a non-polar hydrocarbon solvent – without water or with small-quantity of water bound to the surfactant head-groups. Not all surfactants are highly soluble in hydrocarbon solvents – as it was mentioned in Section 2.3. AOT and many of its derivatives are highly soluble and hence these are good model systems to consider. However, for normal AOT, the CMC and aggregation number (n_{agg}) are harder to determine in non-polar media as compared to in aqueous systems.^{85–87} Work of Peri indicated that n_{agg} increases with solvent ACN;⁸⁸ n_{agg} for AOT in *n*-nonane and *n*-dodecane were between 26 and 29 respectively, whereas a separate SANS study suggested n_{agg} to be 22 in *n*-decane.⁸⁹ Pulse-field gradient NMR studies supported the idea that at zero water content, n_{agg} of AOT is of the order of only few molecules.^{53a} In cyclic hydrocarbon solvents such as benzene, *p*-xylene or cyclohexane n_{agg} was found between 20–30.^{47,89,90} Studies of dry-RMs of C_iE_j or DDAB surfactants in nonpolar solvents are few and far between. An infra-red spectroscopic study by Pacynko *et al.*⁹¹ indicated that intermolecular hydrogen bonding among the EO units is unimportant for C_iE_j aggregation in non-polar solvents. In addition, C_iE_j aggregation in linear and cyclic hydrocarbon solvents was observed, but not in solvents such as benzene and CCl_4 .^{92,93} For instance, SANS measurements of Ravey *et al.*⁹³ suggested that based on n_{agg} , solvents for $C_{12}E_4$ could be categorized into three classes (i) long-chain hydrocarbons (*n*-decane and *n*-hexadecane) in giving $n_{\text{agg}} \approx 13$, (ii) medium-chain hydrocarbons (*n*-heptane) in which $n_{\text{agg}} \approx 5$ –8, (iii) polar organics (methanol, CCl_4 *etc.*) in which no significant aggregation was observed. Some hypotheses suggest that the key factor in switching aggregation on/off (*i.e.* aggregation or no aggregation) might be the solvent polarity/dielectric properties.⁹⁴ Hollamby *et al.*⁹⁵ showed that aggregation of AOT and $C_{12}E_5$ can be correlated with “solvent quality” – characterized by solvent Hildebrand and Snyder polarity parameters.

Early SANS measurements indicated lack of micellization of AOT in cyclohexane until the concentration exceeded 0.225 mM; the dry-RM radius (*i.e.* radius covered by surfactant tails and head-groups) in cyclohexane was 12 Å, which was a little smaller compared *n*-decane (15 Å).⁸⁹ Smith *et al.*⁹⁶ applied contrast variation-SANS to investigate aggregation and CMCs of AOT in deuterated cyclohexane, benzene and *n*-dodecane: it was found that the CMCs (≈ 0.09 – $0.13 \text{ mmol kg}^{-1}$) were very similar in these solvents. At $[\text{AOT}] \gg \text{CMCs}$, the dry-RMs in these solvents were spherical (Fig. 6), with radii between 15–16 Å. The aggregation number n_{agg} was essentially constant at ~ 28 between $\sim 0.1 \text{ mmol kg}^{-1}$ (just above CMC) and $\sim 10 \text{ mmol kg}^{-1}$ ($\gg \text{CMC}$) in both *n*-dodecane- d_{26} and cyclohexane- d_{12} ; however, in benzene- d_6 , $n_{\text{agg}} \approx 20$ when the AOT concentration was just above the CMC ($0.09 \pm 0.06 \text{ mmol kg}^{-1}$ in benzene- d_6), and gradually increased to ~ 28 upon increasing the AOT concentration to $\sim 10 \text{ mmol kg}^{-1}$.

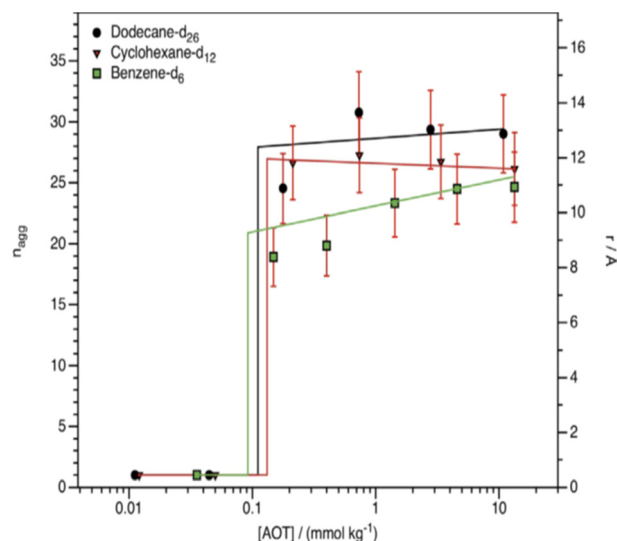


Fig. 6 Aggregation number (n_{agg}) and radii of h-AOT dry-RMs in various organic solvents. Temperature: 298 K. The sudden increase of n_{agg} is consistent with a free molecule to aggregated transition, *i.e.* a CMC. (Reprinted from ref. 96, Copyright (2016), with permission from Elsevier.)

These results indicated that the solvent chemical nature (aromatic solvents especially) might have a subtle effect on AOT-aggregation. Nevertheless, for linear *n*-alkanes it can be said that the solvent ACN has no notable influence on the dry AOT RMs.^{89,93,95–97}

A comparison of AOT aggregation in deuterated *n*-decane- d_{22} and toluene- d_8 solvents by Spehr *et al.*⁹⁸ found the internal RM polar core radii (surfactant heads) to be 3 Å and 2 Å respectively and surfactant shell-thickness ≈ 13 Å. The scattering length density (ρ_s) of solvent-free AOT-shells is ideally $\rho_{\text{shell}} = -0.38 \times 10^{-6} \text{ \AA}^{-2}$; analyses gave ρ_{shell} in toluene- $\text{d}_8 + 1.1 \times 10^{-6} \text{ \AA}^{-2}$ compared to $-0.1 \times 10^{-6} \text{ \AA}^{-2}$ in *n*-decane- d_{22} – suggesting higher penetration of toluene into the surfactant chain-tips compared to *n*-decane. Nevertheless, the general finding based on all the results mentioned above is that solvent chemical identity appears not to have much significant effect on dry-RM aggregation. It is worth noting the differences between these findings and the high degree of sensitivity to phase behaviour for AOT-based μEs nevertheless (see Section 2.3).

3.2. W/O- μEs : surfactant chemical structural effects on interfacial films

The packing parameter model would predict that efficient surfactant packing is a function of surfactant hydrophobic chain length as well as chain-volume – pointing to the importance of sufficiently long hydrophobic chains with branching.^{4,27} This was discussed in Sections 2.2 and 2.3. Theoretical calculations of Nagarajan⁹⁹ emphasized the idea that hydrophobic tails have important effects – adding that the effects can be either “explicit” (modification of hydrophobic tail molecular area) or “implicit” (by satellite-control over surfactant head-group area, a_h). Values listed in Table 3 derived from SANS and surface tension analysis by Nave *et al.*^{55,102} suggest that the surfactant



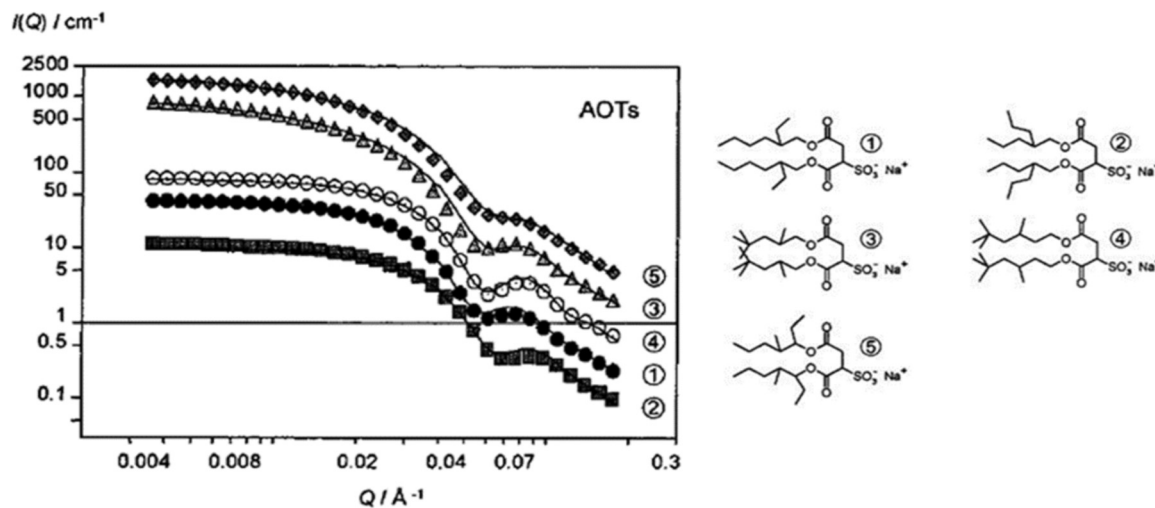


Fig. 7 Surfactant shell-contrast ($D_2O/h\text{-surf}/d\text{-alkane}$: D/H/D; D: deuterated, H: proteated) SANS profiles for W/O- μ Es of normal AOT (AOT1) and AOT-analogues ([surfactant]: 0.050 M, n -heptane- d_{16} as the solvent). Experimental temperatures are listed in Table 3, $W = 30$. Lines are fits to the Schultz core-shell model. Data and fits were multiplied by various factors for better resolution. The figure was reprinted (adapted) with permission from ref. 55. Copyright 2000 American Chemical Society.

Table 3 Properties of normal AOT (AOT1) and its analogue surfactants in aqueous phases and W/O- μ Es (μ E-properties derived from core-shell-drop SANS analyses) described in ref. 55 and 102. For the W/O- μ Es, [surfactant] = 0.100 M, $W = 30$, pressure = 1 bar; the experimental temperatures corresponding to the μ Es are listed below. AOT1–AOT5 structures are depicted in Fig. 7

AOTs	W/O- μ Es ($W = 30$)				
	Aqueous CMC $\pm 0.03/\text{mM}$	Temperature/K	Water-core radius, $R_c \pm 1/\text{\AA}$	Surfactant film thickness, $t_s \pm 1/\text{\AA}$	Headgroup area, a_h (swelling law) ^a $\pm 2/\text{\AA}^2$
1	2.56	295	46	9.1	74
2	3.18	318	41	8.7	79
3	4.36	282	43	8.6	76
4	1.10	323	45	9.4	74
5	7.15	282	41	8.4	86

^a Swelling law: $\alpha(p)R_c^{3v} = 3W\frac{v_w}{a_h} + 3\frac{v_h}{a_h}$ where v_w is volume of a water molecule, v_h is surfactant head-group volume and $\alpha(p) = 1 + 2p^2$, p is droplet polydispersity.

hydrophobic architecture might not influence the surfactant-shell thickness (t_s) and headgroup area (a_h) significantly.⁵⁵ There might be a structure-aggregation correlation in terms of interdroplet attractions, however (Fig. 7);⁵⁵ alkyl-branching at the carbon atom next to the C–O–C group, as it is in AOT3 and AOT5, led to structure factor $S(Q) > 1$ – reminiscent of an Ornstein–Zernike $S(Q)$.¹⁰⁰ Looking back at Fig. 4(B), these two surfactants form single-phase L_2 - μ Es only at low temperatures – and so attractive interactions might become evident at higher temperatures.

In non-polar solvents, the effect of the total carbon number at both cationic and anionic surfactant-tails is believed to be inversely linked to n_{agg} .⁹² Additionally, introducing branching in these hydrophobic fragments reduces n_{agg} .⁹⁰ This pattern is in accordance with the measurements of Frank *et al.*, and CMCs of various linear-chain AOT measured in hydrocarbon solutions as well as aqueous media respectively.^{101,102}

On introducing aromatic phenyl tips into the AOT-hydrophobic tails, the effective hydrophobic-tail area at the

CMC (A_{CMC}) increases.⁵⁶ The limiting aqueous surface tensions (γ_{CMC}) of these surfactants were also higher compared to normal AOT – which was attributed to the polarizable phenyl groups. Adding branching into the phenyl-tipped AOTs reduced γ_{CMC} but increased A_{CMC} . As mentioned before, and importantly, it was observed that these phenyl-tipped AOTs could only form W/O- μ Es in an aromatic solvent such as toluene – effectively mimicking the aromatic nature of the surfactant chain-tips – but not in linear hydrocarbon such as n -heptane. Moreover, inclusion of branched groups in these phenyl-tipped AOTs results in weaker packing at water-toluene interfaces – hence less water uptake inside the droplets. Nevertheless, similar to the linear and branched AOTs mentioned in ref. 55, the phenyl-tipped AOTs also did not notably affect the a_h – showing that the droplet core properties may be dominated by water-headgroup interactions.^{55–57}

Cationic C_nC_m DAB surfactant films at the O–W interfaces were introduced previously in Sections 2.2 and 2.4. in the light of the cylinder \rightarrow spheroid \rightarrow sphere structural evolution on



increasing water content.^{28,38,39} Contrast variation SANS results showed that increasing the C_n -chain length leads to an increase of the cylinder radius, although no clear correlation in terms of cylinder length could be found.²⁸ It was mentioned previously that cyclohexane weakly penetrates the surfactant chains on increasing asymmetry of the two alkyl-tails C_n and C_m .²⁹ (Note, nonionic-film properties have been discussed in Section 2.5.)

3.3. Surfactant-film rigidity

In W/O- μ Es, surfactant monolayers bend towards water – generating negative curvature; on the other hand, in O/W- μ Es the curvature is towards the oil and is said to be positive.⁴ In this context, surfactant film-bending rigidity is synonymous with film-curvature. The loose term “film rigidity”, which is a measure of the energy required for a surfactant monolayer to bend towards a continuous solvent (either oil or water), is a key aspect in understanding μ E-film stability.³¹ Part of the model was discussed in Section 2.3., which relates surfactant monolayer free-energy of bending to curvature c .

For low volume fractions of spherical droplets (ϕ), the free energy of bending of surfactant film per unit area of μ Es (F/A) can be expressed as^{35,76}

$$\frac{F}{A} = \left[2\kappa \left(\frac{1}{\rho} - \frac{1}{\rho_0} \right)^2 + \frac{\bar{\kappa}}{\rho^2} \right] + \left[\left(\frac{k_B T}{4\pi\rho^2} \right) f(\phi) \right] \quad (4)$$

ρ and ρ_0 are the radius and spontaneous radius of curvature as defined before, whereas $f(\phi)$ is a function related to the entropy of mixing of droplets, expressed as

$$f(\phi) = \{\ln(\phi) - 1\}^3 \quad (5)$$

for droplet volume fraction $\phi < 0.1$. This theory connects IFT, which is a macroscopic property, to droplet core-size, a micro(nano)scopic property, through the bending constant ($2\kappa + \bar{\kappa}$), by the following

$$2\kappa + \bar{\kappa} = \gamma_{o/w} R_c^2 - \frac{k_B T}{4\pi} f(\phi) \quad (6)$$

where $\gamma_{o/w}$ is O–W IFT, R_c is the water-core droplet radius. The droplet polydispersity, p , is introduced because droplets experience thermal fluctuations – rendering a size distribution which is linked to film properties. The relationship between p and film-bending is expressed as

$$2\kappa + \bar{\kappa} = \frac{k_B T}{8\pi \left(\frac{\sigma}{R_c} \right)^2} - \frac{k_B T}{4\pi} f(\phi) \quad (7)$$

where $p = \sigma/R_c$ is the droplet size distribution width. Eastoe *et al.* applied the model to cationic $C_n C_m$ DAB-based Winsor II μ Es; on increasing the two C_n and C_m surfactant hydrophobic chain-lengths (droplet shell thickness in other words) a reduction in droplet polydispersity and an increase in the ($2\kappa + \bar{\kappa}$) constant was observed, *i.e.*, surfactant “film rigidity” increased with surfactant chain length (Fig. 8).³⁵ As pointed out in Section 2.5., similar observations were made for the nonionics. The variation of film-bending parameters was attributed to surfactant hydrophobic chain-length; however, the

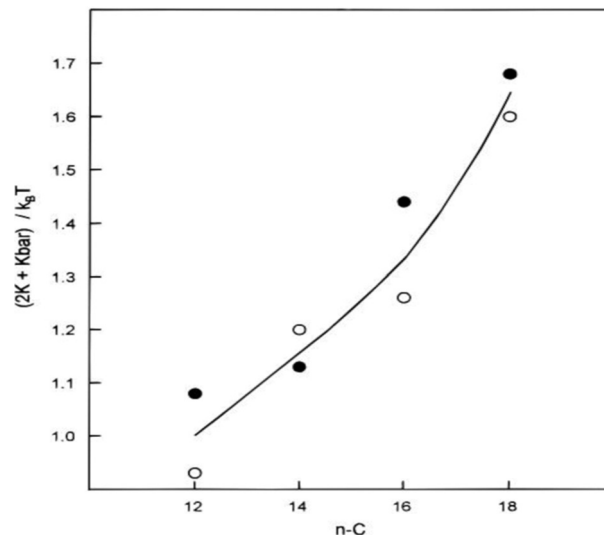


Fig. 8 Variation of surfactant film-bending constant (in units of $k_B T$) as a function of $C_n C_{12}$ DAB W/O- μ Es (solvent: cyclohexane, temperature: 298 K), the line is a guide to the eye. Circles represent calculation of ($2\kappa + \bar{\kappa}$) using eqn (6) (white circles) and eqn (7) (black circles); $\gamma_{o/w}$ was estimated using surface light scattering (SLS), R_c and σ/R_c were obtained by SANS measurements. The figure was reprinted with permission from ref. 35. Copyright 1997 American Chemical Society.

solvent chain-length effect, contrary to the model proposed by Shah *et al.*,^{23b,c} was interpreted to be secondary.^{76,103–105}

Rather, Sottmann *et al.*¹⁰⁵ claimed that for the nonionics, a_h at O–W interface is only dependent on the length of EO fragment and independent of the solvent as well as surfactant hydrophobic chain-length; nevertheless, the film-bending constants increase upon increasing surfactant hydrophobic chain-length. In this respect, data for AOT-based μ Es are scarce, for example, ellipsometry experiments by Binks *et al.*^{80,82} (Section 2.6). To recap, for the AOT, on increasing solvent ACN the tendency to develop saddle-like or bicontinuous structures increases. Surprisingly, fluorescence quenching studies by Almgren *et al.*¹⁰⁶ could not find any such variation – suggesting the film-bending estimation might be exclusively responsive to the experimental technique being employed.

3.4. Solvent chemical-effect on μ E droplet morphology

The ellipsometry data of Binks *et al.*⁸⁰ indicate that solvent chemical nature (*i.e.* the effect of n -alkane ACN variation, presence of aromaticity *etc.*) might have a subtle role in controlling the surfactant-film properties. Regarding solvent molecular penetration into the surfactant monolayers, the key point is that it is subtle – although some solvents might be able to slightly penetrate the surfactant chain-tips, most show only limited effects.^{8,28,29,36} In AOT-based W/O- μ Es, solvent penetration was mostly studied by indirect techniques such as nuclear magnetic resonance (NMR), fluorescence correlation or imaging.^{46,107–110} These results suggest that cyclic molecules (cycloalkanes or aromatic hydrocarbons) are more efficiently incorporated into AOT-monolayers compared to branched or linear alkanes. In addition, IFT measurements by Aveyard *et al.*¹¹¹



suggested that solvent penetration into planar AOT-monolayers at the brine (0.100 M NaCl) solution/oil interface increases with reduction of solvent ACN.

All these methods discussed above were indirect, however, direct and quantitative information on solvent penetration can be obtained from carefully designed and executed contrast-variation SANS measurements.^{17,28,29} Analysis of the SANS data reveals that solvent penetration with symmetric dichain surfactants is not significant – regardless solvent chemical structure. On the other hand, solvent penetration could be detected into monolayers of di-chain surfactants having asymmetric tails: for example, using C_nC_{12} DAB-based W/O- μ Es, on increasing n ($n = 12, 14, 16$ and 18) the measured volume fraction of cyclohexane penetrating into the chain region increases from 0 (no penetration) to no greater than 0.12. This maximum value observed corresponds to approximately one cyclohexane molecule penetrating for every two surfactant molecules.²⁹ Interestingly, these SANS analyses (Table 4) suggested that the a_h can be influenced by changing the solvent chemical nature.

Coming to the second aspect of droplet size and morphology, SANS-results reveal that the effect of n -alkane ACN on water-core radius (R_c) in W/O- μ Es is minimal.^{113–116} The solvent, however, affects droplet polydispersity and the extent of interdroplet attractive interactions (see next section) quite significantly. For instance, neutron spin-echo (NSE) experiments¹¹⁷ suggested that for W/O- μ Es, water-core droplets were less rigid and more polydisperse in linear n -alkanes such as n -hexane compared to cyclohexane ($p = 0.24$ vs. 0.39 and bending constant $K = 0.19 k_B T$ vs. $0.24 k_B T$, $W = 10$). In particular, droplet attractive interactions are minimal in n -hexane to n -decane, but significant for linear n -alkanes with ACN > 10 .^{118,119} Interdroplet interactions can be understood in terms of material exchange through the channels formed due to droplet attraction – time-resolved fluorescence results suggested that n_{agg} as well as interdroplet exchange rate increases on increasing solvent ACN.¹²⁰ Solvent chemical effects on droplet size distribution and surfactant-film geometry were observed by Balakrishnan *et al.*;¹²¹ the size probability distribution tended to be more uneven (less Gaussian-like) as the solvents were progressively changed from n -hexane to n -dodecane.

To summarize Sections 3.2–3.4, the chemical effects on droplet morphology and surfactant-films should be discussed more holistically, *i.e.* the effects may not be due to solely the surfactant or solely the solvent – rather a surfactant–solvent chemical combination might be governing the interfacial and structural properties. For a given set of surfactants, understanding surfactant chemical variation might be relevant considering the solvent identity and *vice versa*. It is seen that the solvent chemical-structure effects are notably manifest in terms of interdroplet attraction parameters, as discussed below.

3.5. Attractive interaction in W/O- μ Es: square-well potential

Shifts in μ E phase-transition boundaries are correlated with the interfacial film bending and droplet–droplet attractive interactions. When droplets in W/O- μ Es start swelling on increasing W and they get closer to one another, interdroplet attractive interactions caused by the overlapping of surfactant hydrophobic tails drive the L_2 -to-cloud point/critical-type phase separation.^{37,112} The attractive interaction between two hard sphere-like surfactant-coated droplets can be interpreted in terms of an attractive square well potential function, expressed as⁹⁷

$$\begin{aligned} U(r) &= \alpha \text{ at } r < \sigma \\ U(r) &= -\epsilon k_B T \text{ at } \sigma < r < r_o \\ U(r) &= 0 \text{ at } r > r_o \end{aligned} \quad (8)$$

where ϵ is the square-well potential depth, which is also considered to be a measure of the tail attractive interactions. Huang *et al.* utilized this expression to analyze the $S(Q)$ of a hard-sphere liquid with an attractive square-well potential proposed by Sharma *et al.*¹²² The so-called attractive interaction parameter (A) can be expressed as

$$A = 8(\lambda^3 - 1)(e^{\epsilon'} - 1) \quad (9)$$

and

$$\epsilon' = \log(\epsilon + 1) \quad (10)$$

Here, the width of the well is given by λ and the range of interaction is expressed as $2\lambda R$ – where R is the “hard-sphere”

Table 4 Properties of cationic C_nC_{12} DAB-films ($n = 12, 14, 16$ and 18) in W/O- μ Es obtained from phase behaviour and SANS analyses²⁹

Surfactants (C_nC_{12} DAB)	Solvents	Maximum water solubilization, W_{max}	Headgroup area, ^a $a_h/\text{\AA}^2$	Surfactant film thickness, ^a $t_s \pm 1/\text{\AA}$	Oil volume fraction ^a ϕ_{oil}
C12–C12	<i>n</i> -Heptane	25	65.8	10.8	0.01
	Cyclohexane	12	54.8	11.0	0
C14–C12	<i>n</i> -Heptane	23	66.3	10.6	0.01
	Cyclohexane	12	59.7	11.7	0.05
C16–C12	<i>n</i> -Heptane	21	66.8	11.5	0.02
	Cyclohexane	11	55.6	12.8	0.07
C18–C12	<i>n</i> -Heptane	19	66.3	11.8	0.02
	Cyclohexane	11	58.9	13.5	0.08

^a Uncertainties: $t_s \pm 1 \text{\AA}$, a_h (swelling law) $\pm 2 \text{\AA}^2$, $\phi_{oil} \pm 10\%$.



droplet radius, ε is the square-well depth and ε' is the normalized square-well depth. It was reported that ε' increased, almost linearly, on increasing R . Values of the attractive interaction parameter A for various AOT-based W/O- μ Es were studied by means of light scattering experiments^{97,125} and by SANS.¹²³ The general consensus is that tail-tail attractions increase with increasing solvent chain length. In terms of the most significant factors for interdroplet attraction, Kaler *et al.*¹²³ emphasized on the relative strength of the short-range tail-tail attractive interactions, rather than long-range van der Waals interactions between droplet water-cores. SANS analyses showed that ε' effectively disappeared when solvent density was 0.692 g cm^{-3} , which is close to the density of solvents such as *n*-heptane and 2-methylheptane – the latter being the closest resemblance to the AOT tail structure.¹²³ Tingey *et al.*¹²⁴ emphasized much on the same factor as well.

3.6. Attractive interactions and critical behaviour

To further extend discussions from the previous sections, the attractive interactions between μ E-droplets is theorized as mutual penetration of surfactant chain-tips. The upper temperature (*i.e.* L_2 -to-cloud point) phase separation can be accounted for by considering interdroplet attractive potential – when ϕ is sufficiently high (10 vol% and onwards for instance);⁹⁷ in W/O- μ Es, at this range the droplets are close enough to one-another due to the surfactant-shell overlapping. Lemaire *et al.*¹¹² interpreted the surfactant-shell interaction (U) as a sum of hard-sphere (U_{hs}) and attractive potential (U_A) as

$$U = U_{\text{hs}} + U_A \quad (11)$$

in which U_A results in due to a sum of interactions among (i) surfactant-shells to surfactant-shells, (ii) surfactant-shells to solvent and, (iii) solvent to solvent. The surfactant-shell interactions, in addition, are treated as a Lennard-Jones interaction energy E_{ij} among all the atoms of two distinct surfactant-shells

$$E_{ij} = 2\varepsilon_{ij}r_{ij}^6/r^6 \quad (12)$$

where ε_{ij} is interaction energy between any two individual atoms and r is the separation distance between atom i and atom j .

In terms of experimental observations,^{55,56,112,126} on increasing temperature the attractive potential reaches to a threshold, initiating phase separation; close to the upper temperature phase-boundary, μ Es show enhanced turbidity suggesting strong attraction among droplets. In AOT-based W/O- μ Es, an interesting phenomenon has been of much attention; here near the phase-boundary, the L_2 - μ E is split to two nearly equal volume μ Es of different μ E-densities.^{97,112,122,125} This is interpreted analogous to liquid-liquid critical separation due to two different types of local structures resulting in a density gradient¹²⁶ within the single-component liquid (such as water). In AOT-in-*n*-decane W/O- μ Es, this separation takes place around 316 K (droplet volume fraction, $\phi = 0.075$).^{20,125} Near the critical temperature (T_c), droplet-hydrodynamic correlation-length increased from 100 Å to several thousand angstroms.¹²⁵

A simplified expression of SANS scattering intensity $I(Q)$ near T_c includes a structure-factor $S(Q)$ ^{4,97}

$$I(Q) = nP(Q)S(Q) \quad (13)$$

where $P(Q)$ is a single-droplet form factor relating to the water-core size and geometry and n is droplet-number density. Through extensive interpretation of SANS data Chen *et al.*,²⁰ Huang *et al.*^{97,125} and Kotlarchyk *et al.*¹²⁷ showed that near the critical point, the $S(Q)$ for AOT-based W/O- μ Es in *n*-decane can be interpreted by an Ornstein-Zernike (OZ) interparticle-potential function¹⁰⁰

$$S(Q) = 1 + \left(\frac{\chi}{1 + Q^2\xi^2} \right) \quad (14)$$

Here, ξ is a correlation length of local droplet-concentration fluctuations and χ is a constant. When there is no, or only an insignificant, attractive interaction then $S(0) = 1$, whereas strong interdroplet attractions lead to increases in the value of $S(Q)$, especially at low Q values. This is observed in AOT-based W/O- μ Es in *n*-decane as the critical boundary has been almost reached on increasing temperature (Fig. 9).²⁰

The extent of attractive interactions near the L_2 -to-cloud point phase-boundary location is governed by temperature as well as solvent.¹²⁸ Earlier (Section 2.3) it was discussed that in AOT-based W/O- μ Es, this boundary shifts to lower-temperature.⁵⁴ The effect of *n*-alkane solvent ACN near this phase-boundary was studied by Toprakcioglu *et al.*,¹²⁸ for $W = 20$ at $[\text{AOT}] = 0.100 \text{ M}$, the critical-behaviour in *n*-decane could be seen around 313 K, whereas in *n*-undecane and *n*-dodecane the critical behaviour is observed at much lower temperatures (295 K and 278 K respectively).

The SANS results explain electrical percolation of droplets in AOT-based W/O- μ Es in various *n*-alkanes.^{129–131} Percolation is interpreted as material exchange (*i.e.* surfactant counterions or dispersed medium) due to droplet clustering owing to the attractive interactions – which results in a sudden rise in electrical conductivity by several orders of magnitude on increasing temperature or ϕ .^{13,129,130} On increasing *n*-alkane ACN, the onset of percolation shifts to lower ϕ and lower temperature in AOT-based W/O- μ Es.^{16,130}

It should be noted that the critical-type OZ behaviour is not exclusive to dichain anionics but was observed in C_iE_j -based μ Es (*n*-octane as solvent) as well.¹³¹ This indicates that regardless of the surfactant chemical identity, the critical behaviour might be a universal phenomenon in W/O- μ Es, but more studies are necessary – especially on dichain cationics. Lemaire *et al.*¹¹² argued about a potential role of solvent molecular volume and their model hinted at a possible effect of surfactant-solvent mutual interaction energies, which are functions of the chemical nature of the constituents.

3.7. W/O μ Es: aromatic hydrocarbons

Aromatic solvents are chemically versatile, differing from their aliphatic counterparts not only in terms of chemical structures, but smaller effective molecular volumes and presence of delocalized electrons. In Section 3.2, it was shown



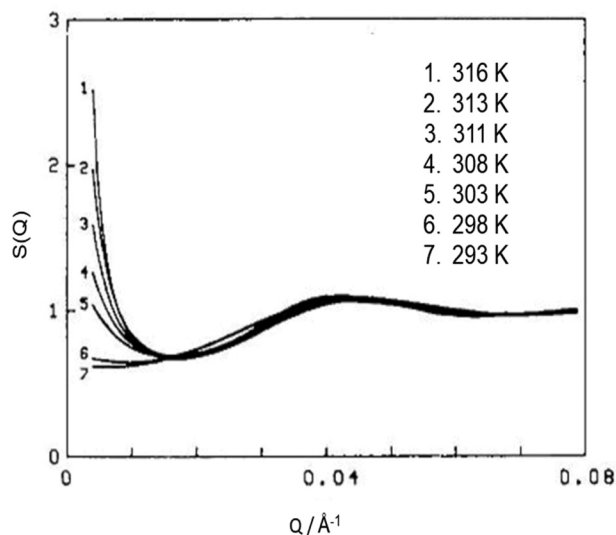


Fig. 9 Mean droplet–droplet structure factor $S(Q)$ depicting interdroplet attractive interactions – the μE is AOT/ D_2O / n -decane (3/5/92 wt%), $\phi = 0.075$, and it is approaching critical temperature (~ 316 K). $S(Q)$ were calculated using a mean-spherical approximation procedure described in ref. 20. The figure was adapted from ref. 20. Copyright (1986), with permission from Elsevier.

that solvent–surfactant chemical matching is important, as observed in phenyl-tipped AOT-based W/toluene μEs .⁵⁶ Literature on ionic W/O- μEs hints that auxiliary aliphatic chains might be important for governing μE -phase behaviour and surfactant film properties.^{96,132–135} Appel *et al.* studied the effect of aromatic solvents on droplet percolation; the temperature corresponding to an onset of percolation in AOT-based μEs decreased on increasing the substituted aliphatic chain-length in aromatic hydrocarbons, this parallels observations of Alexandridis *et al.* for n -alkane solvents on increasing ACN.^{25,132}

NSE and SANS investigations by Spehr *et al.*⁹⁸ found film bending constants in AOT/water/toluene μEs to be higher than in the μEs containing linear- or cycloalkanes. In terms of the water-droplet size and surfactant film-geometry, nevertheless, droplet-exchange kinetics, light scattering and SANS studies pointed out that aromatic solvent-chemical structure, except halogenated aromatics, seldom affects μE -droplet characteristics.^{67c,133–135} The hydrodynamic size of water-droplets in medium-chain aliphatic and aromatic hydrocarbons was compared but no significant variation was found.¹³⁵ The case was similar for dry-RMs, which was noted by Smith *et al.*⁹⁶ (discussed in Section 3.1), as the morphology and radii of these aggregates were insensitive to solvent chemical nature (n -alkanes, cycloalkanes or the aromatics).

3.8. Solvent blends

An interesting approach to tune hydrocarbon solvents, in particular physical properties such as density, is using controlled blends, as shown by Salabat *et al.*¹³⁶ This allows different intermediate solvent properties to be accessed, and the results show clear links between phase behaviour and mixed solvent identity. With blends of n -heptane and toluene, L_2 - μEs

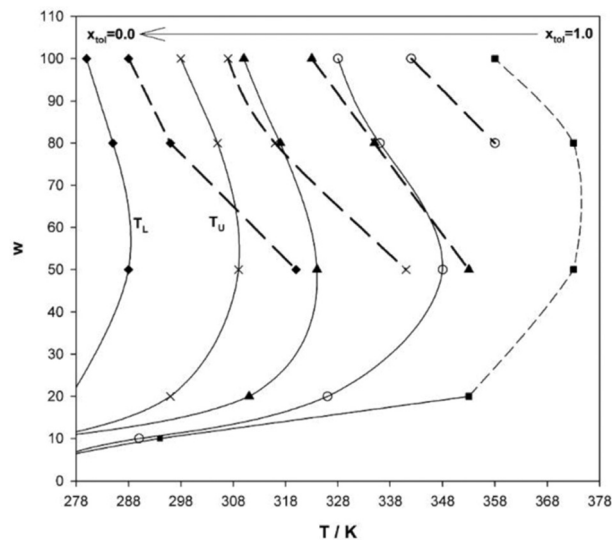


Fig. 10 Phase-behaviour of AOT-based W/O- μEs in pure n -heptane and toluene and their solvent blends at toluene mole-fractions $x_{\text{tol}} = 0.25, 0.50$ and 0.75 (from left to right). $x_{\text{tol}} = 0$ (far left) represents pure n -heptane, $x_{\text{tol}} = 1$ (far right) is pure toluene. [AOT] = 0.100 M, solid lines indicate W11-to- L_2 transition boundary and dashed lines represent L_2 -to-cloud point boundary. Lines are guides to the eye. The figure was reprinted from ref. 136. Copyright (2007), with permission from Elsevier.

domains could be tuned between the two pure-solvent extremes and shifted to higher or lower temperatures by varying solvent composition mole fraction (Fig. 10). The same study showed that the aggregation and colloidal stability in such μEs could also be controlled; translational diffusion coefficients of droplets decreased (*i.e.* “switching off” of interdroplet interactions) on increasing toluene mole fraction. The opposite effect dominates when n -dodecane is used in place of toluene. To characterize the solvent mixtures the effective molar volume of the mixed solvent ($V_{\text{mol}}^{\text{eff}}$) can be defined as –

$$V_{\text{mol}}^{\text{eff}} = X_1 V_{\text{mol}1} + X_2 V_{\text{mol}2} \quad (15)$$

where $V_{\text{mol}1}$ and $V_{\text{mol}2}$ are the molar volumes, and X_1 and X_2 are mole fractions of the two solvents. Myakonkaya *et al.*,¹³⁷ in this regard, introduced a concept of “good” and “bad” solvents for controlling L_2 - μE domain size for a given surfactant (normal AOT in this instance), where “bad” solvents (long-chain hydrocarbons such as n -dodecane) induce droplet attraction and “good” solvents (such as n -heptane) promote, for instance, L_2 - μEs domains having hard-sphere droplets. Apparently, the attractive droplet interactions increase on increasing mole fraction of the solvent with the longer alkyl chain. A matter of interest is the effect of solvent-blends on critical-type phase-separation; in AOT-based W/O- μEs , T_c can be decreased by increasing $V_{\text{mol}}^{\text{eff}}$; this was demonstrated by mixing n -undecane and n -nonane; adding the former increases $V_{\text{mol}}^{\text{eff}}$ of the solvent blend, and by means of this T_c can be reduced to as much as 6 K.

3.9. Supercritical alkanes: oil-density guided self-assembly?

Above the critical pressure (P_c) and temperature (T_c), the gas–liquid boundary for a fluid disappears and a supercritical fluid



will form. Supercritical fluids generally have much lower density and higher diffusivity than normal liquids. Due to being supercritical fluids at room temperature and more compressible than normal liquids, low-density *n*-alkanes (*n*-ethane, *n*-propane and *n*-butane for instance) expand the horizon of stable μ Es. In such low-density supercritical *n*-alkanes, μ E stability is expected to be a strong function of temperature and pressure. It has been observed that on increasing pressure (*i.e.* increasing fluid density) of short-chain supercritical *n*-alkanes the phase transition in AOT-based W/O- μ Es involves a gradual shift from a three-phase system to L₂- μ Es.^{138–140} At high pressures, water solubilization in *n*-propane showed similarity to that of medium-chain *n*-alkanes at ambient conditions.^{139,140} Density is an important factor here: the effect of solvent ACN and pressure on W_{\max} of AOT-based W/O- μ Es can be understood from Table 5 based on studies by Tingey *et al.*,¹²⁴ lowering W_{\max} in isooctane on increasing pressure is worth noting. At 300 bar and 298 K, isooctane has a density of 0.716 g cm⁻³ which is very close to *n*-nonane density at room temperature and pressure.^{141,142} This observation correlates to the L₂-phase shift of AOT W/O- μ Es towards low temperatures on increasing linear hydrocarbon ACN (discussed in Section 2.3). It is clear, that the solvent-density change due to application of pressure is an important parameter, as it enables stabilization of μ Es under different thermodynamic conditions which are not accessible with normal liquid-like solvents at atmospheric pressure.

Fulton *et al.*¹³⁹ reported that the pressure-dependent electrical conductivity of AOT/water/isooctane μ Es at 298 K at a specific AOT concentration (0.036 M) could nearly be mimicked by AOT/water/*n*-propane μ Es at 376 K (103 °C); while the increase of pressure did not affect electrical conductivity of AOT/water/isooctane μ Es, the conductivity of AOT/water/*n*-propane μ Es lowered and appeared to reach closer to the

former as pressure was gradually increased. Meanwhile, phase behaviour and scattering investigations^{116,140} of AOT- μ Es in low density *n*-propane and *n*-butane, as a function of pressure and temperature, found that lowering the pressure from 400 bar to 70 bar pushed L₂- μ Es towards unstable systems. This transition pressure, and the nature of phase-separation, depend on temperature and the surfactant chemical nature. The pressure (and hence solvent density) was used to control interdroplet attraction: lowering pressure, and hence fluid density, promoted growth of droplet clusters, whereas increasing pressure/density led to split the droplets from those clusters. This demonstrated that even in low density *n*-alkanes, it is possible to emulate the phase-boundaries that are often observed in long-chain linear *n*-alkanes. The observations also support SANS results by Kaler *et al.*¹²³ indicating that μ E-droplets formed in supercritical *n*-propane were of similar size to μ E-droplets in regular solvents, for example, *n*-hexane. Nonionic C₈E₅S in near-critical ethane and *n*-propane solvent blends were studied by Beckman *et al.*¹⁴³ by dynamic light scattering; on increasing pressure they found droplet diffusion increased, which was attributed to lowering interdroplet attraction. This is also consistent with the system locating away from the cloud-point phase boundary.

4. The effect of surfactant counterions

So far, the focus on ionic surfactant-chemical architecture was exclusively on the chemical nature of the surfactant-ion (*i.e.* hydrophobic moieties). This section considers the effects of variation in head group counterion in anionic surfactants. Sodium (Na⁺) is the natural counterion for the anionic AOT and its analogues (*i.e.*, Na-AOT), but this counterion can be readily varied by ion exchange techniques. In addition, the “inorganic” nature of the counterion can be changed by substitution with different “organic” quaternary ammonium ions.¹⁵¹ The counterion provides screening of Coulombic repulsion between the like charged head-groups,³⁹ and this effect is important for surfactant self-assembly. In the study by Oshitani *et al.*,¹⁴⁴ electrostatic screening was found to follow the order: K⁺ \approx Rb⁺ > Cs⁺ > Na⁺ > Li⁺. Substituting Na⁺ by monovalent Cs⁺ led to a change of AOT-reverse micellar morphology in *n*-decane from slightly oblate spheroids to disks.¹⁴⁵ The effect of substituting monovalent counterions with divalent ones, such as Mg²⁺, Ca²⁺, Co²⁺, Cu²⁺, Ni²⁺ and Zn²⁺, yields a class of surfactant salts conveniently labeled M^{tt}(AOT)₂ – in which now one counterion associates with two-anionic surfactant-ions. The effects of this variation on droplet morphology, surfactant-film geometry and overall μ E-phases are significant.

Temperature-composition phase-diagrams³⁹ showed that while Na-AOT μ Es in cyclohexane exhibit two-phase boundaries (Winsor II-to-L₂ and L₂-to-cloudy phase), substitution of Na⁺ with Mg²⁺, Ca²⁺, Co²⁺, Cu²⁺ and Zn²⁺ made the W/O- μ Es less temperature-sensitive, and only one phase boundary (L₂ to Winsor II) could be observed.¹⁴⁶ The SANS profiles for M^{tt}(AOT)_n W/O- μ Es at a high dilution region has been represented by a log-log plot.¹⁴⁸ In this representation, at $Q < 0.10 \text{ \AA}^{-1}$, the $I(Q)$ profile

Table 5 Pressure-induced maximum water-solubilization (W_{\max}) of AOT-based W/O- μ Es in nonpolar solvents; [AOT] = 0.087 M, temperature = 298 K¹²⁴

Solvents	Pressure/bar	W_{\max}
Xenon	200	1
	520	5.6
Ethane	200	1
	350	5
	620	10
	840	35
<i>n</i> -Butane	100	45
	300	45
<i>n</i> -Pentane	1	43
	100	43
	300	43
<i>n</i> -Hexane	100	116
	100	105
	300	87
Isooctane	1	74
	100	69
	300	59



can be expressed in terms of a dimensionality (fractal) parameter characterizing the aggregate structure, D , as

$$I(Q) \sim n_p Q^{-D} \quad (16)$$

where n_p is droplet number density. The exponent D is near zero when the counterions were Na^+ , Mg^{2+} and Ca^{2+} – which is consistent with the presence of spherical droplets. However, for $\text{M}^{2+} = \text{Co}^{2+}$, Ni^{2+} and Zn^{2+} SANS profiles showed $D \approx -1.0$, indicating rigid rod-like aggregates. SANS profiles when $\text{M}^{2+} = \text{Ni}^{2+}$ and Zn^{2+} at higher surfactant concentration showed more interesting features. For instance, in the absence of any added water ($W = 0$), the dry RMs were spherical, whereas, between $W = 3$ and 9.75, well-defined rods were formed. As W was increased further, the rod-like structures evolved into less distorted spheroids. In $\text{Co}(\text{AOT})_2$ based μEs , the rod-lengths gradually decreased from 400 Å to 250 Å on increasing W from 5 to 10. Nevertheless, a sphere-to-cylinder transition was observed between $W = 0$ –10 and between $W = 10$ –25 the cylinders again turned into spheres.^{39,147,148} In general, monovalent counterions induce spherical curvature of interfacial surfactant-films, whereas divalent – especially transition metal-ions – induce cylindrical RMs (Fig. 11).^{39,147–150} The physical interpretation was based on the capacity of the large transition metal ions to form hydrated coordination covalent complexes with water – resulting in reduced interactions between the oppositely charged surfactant anions and lower screening of repulsions between the SO_3^- head groups.

Another related class of AOT surfactants includes symmetrical cations such as quaternary ammoniums $[\text{R}_4\text{N}]^+$ (e.g. tetra-*n*-ethylammonium).¹⁵¹ In $W/\text{O}-\mu\text{Es}$, it was observed that the NH_4^+ ion induced spherical droplets, similarly as seen for the Na^+ ion.^{152a} On the other hand, substituting the hydrogen

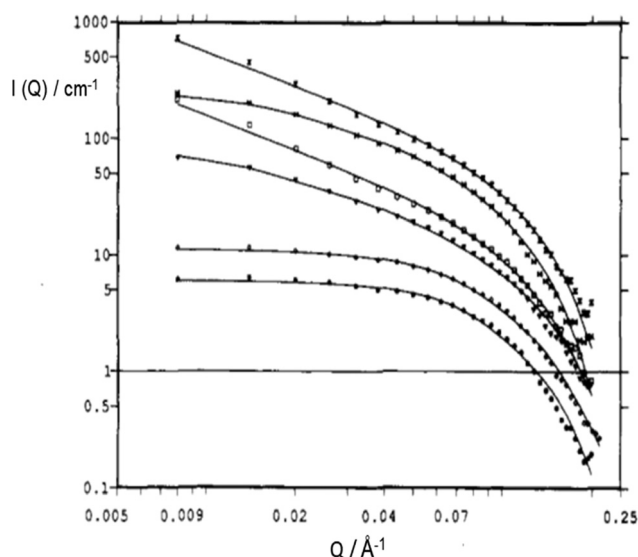


Fig. 11 SANS profiles of $\text{M}^{2+}(\text{AOT})_n$ $W/\text{O}-\mu\text{Es}$ in cyclohexane- d_{12} ($W = 5$, [surfactant] = 0.075 M); (●) Mg^{2+} , (▽) Co^{2+} , (□) Ni^{2+} , (◇) Cu^{2+} , (⊗) Zn^{2+} , (△) Cd^{2+} . Solid lines are fits for rigid rod; data were multiplied by arbitrary numbers for better visual. The figure was reprinted (adapted) with permission from ref. 149. Copyright 1993 American Chemical Society.

atoms of the NH_4^+ with bulky alkyl groups (tetra-*n*-ethylammonium) induced cylindrical micelles at low W and gave a cylinder-to-sphere transition on increasing W – reminiscent of the transition-metal AOTs.^{152a,b} Interestingly, the counterion can be exchanged by even more exotic chemical moieties; for instance, when the Na^+ ion of AOT is exchanged with polymerizable units such as [2-methacryloyloxy]ethyl trimethylammonium cations and the surfactant has been dispersed to polar oils such as methyl methacrylate (MMA), L_2 $W/\text{O}-\mu\text{Es}$ could be formed.^{152c} Similar ion exchange could be carried out for cationic C_{12}DAB surfactants by exchanging the Br^- ion with a more bulky MMA^- ion – albeit the L_2 - μE region is not as broad compared to the former systems.^{152c,d}

Here, the discussions imply that while the surfactant tails as well as solvents significantly affect the phase-behaviour, structural properties, and O- W IFT of $W/\text{O}-\mu\text{Es}$ (Sections 2 and 3), surfactant counterions also have profound control over the phase appearance and droplet morphology. In particular, the cylinder \rightarrow sphere transition, as it was observed for cationic $\text{C}_n\text{C}_m\text{DAB} - \mu\text{Es}$ (Section 2.2),^{28,29,38} is a phenomenon associated with counterion variation for dichain anionics as well. The landscape for exploring this aspect is still wide-open, especially for AOT analogues – since the hydrophobic tails in this class of surfactants can be manipulated to a larger degree due to its flexible synthetic routes,³⁷ for example by incorporating silicon atoms into the tails.¹⁵³

5. General discussion and conclusion

As examples of model thermodynamically stable colloidal systems, it is clear that stability and phase behaviour of $W/\text{O}-\mu\text{Es}$ are clearly linked to surfactant and solvent chemical structures. In terms of phase stability alone, these chemical effects are manifested by the relative position of single-phase μEs within temperature-composition phase diagrams. Chemical effects can be seen more obviously at the macroscopic level (*i.e.* visual means), whereas, on a microscopic local structural level chemical effects are more subtle. It is observed that – (i) enhanced hydrophobicity in the surfactant chains (*i.e.* longer hydrophobic tails and/or presence of branching – including branching at the chain-tip), may result in better surfactant partitioning at an O- W interface – evidenced by lowering the minimum $\gamma_{\text{O/W}}$ values compared to surfactants with less branching or shorter hydrophobic chain-length, (ii) the role of solvent is subtle but significant – solvent chemical structure dictates the L_2 $W/\text{O}-\mu\text{Es}$ phase domains on the temperature-composition landscape (phase diagrams); in addition, depending on the surfactant chemical nature, the solvent might control droplet-droplet attractive interactions, (iii) furthermore, chemical variation in the surfactant-ions and counterions (for ionic surfactants) or in the hydrophobic and hydrophilic moieties (both ionic and nonionic surfactants) renders notable changes in μE -phase stability. Of particular note, especially for double tail anionic surfactants, chain branching seems to be important, since it allows sufficient surfactant conformational chain freedom, which is needed for efficient μE -formation. The degree of importance of



chain branching is, however, not completely understood, and synthesis of surfactants with different chain-branching motifs might shine light on this matter. On the other hand, using linear, more ordered, hydrophobic chains tends to limit conformational degrees of freedom and this is detrimental to μ E-formation.

An important question remains: what are the actual chemical structural factors required for efficient μ E-formation? Is there a universal feature, such as matching of solvent and surfactant-hydrophobic tail density/molecular volume that can be correlated to chemical matching/mismatching? Although these remain open-ended issues, it should be possible to develop a universal surfactant-solvent framework. Clearly, these all go back to thermodynamic and geometric aspects of surfactant films at O-W interfaces, and a recent review¹⁵⁴ is very instructive in this respect. This missing link may be considered as the “holy grail”, accounting for μ E-formation, stability, and structure from a chemical viewpoint. Furthermore, since W/O- μ Es represent ideal systems for probing surfactant-solvent effects, this would have widespread general applications in colloid science, especially for dispersion in non-aqueous and low dielectric solvents. Molecular dynamics (MD) simulations have potential for unraveling these local rules at surfactant-solvent interfaces, especially, since surfactant tip-solvent interactions can be probed which are dominated by van der Waals interactions. A likely advance might be the quantification of surfactant tail-solvent mixing entropies by MD simulations. Attempts for surfactant and solvent chemical unification have been made before, by Peck *et al.*,¹⁵⁵ based on classical thermodynamics. In that model, surfactant tail-solvent mixing energy was modelled based on the Flory interaction parameter (χ),¹⁵⁶ which can then be related to corresponding solubility parameters (δ). In the end however, the interactions were interpreted based on solvent penetration into the surfactant shell, for which careful SANS experiments (see discussions in Sections 2.4 and 3.4) have shown to be only limited.

It is important to remember that these are “fluid” systems; therefore, in W/O- μ Es the boundaries between the surfactant hydrophobic tips and solvent molecules experience local fluctuations over time.¹⁵⁷ It might be possible by MD simulations to “see” these ever-evolving boundaries (*i.e.* chemical configurations) or “feel” (*i.e.* electronic properties, charge and polarizability distribution) to shed light on chemical structural effects. Examples of all-atom as well as coarse-grained modelling of W/O- μ Es, have been emerging recently.^{158–161} Clearly a close link up between experiments and computational modelling is desired, and this might be the ultimate tool for conclusive understanding of the mutual surfactant-solvent chemical effects at fluid interfaces.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

A. R. thanks Commonwealth Scholarship Commission (CSC) of the Foreign, Commonwealth and Development Office (FCDO),

UK for a PhD scholarship. A. R. thanks the Commonwealth Scholarship Commission (CSC) – which is funded by the Foreign, Commonwealth and Development Office (FCDO), UK – for a PhD scholarship.

References

- (a) J. N. Israelachvili, *Intermolecular and surface forces*, Academic Press (Elsevier), Massachusetts, 2011; (b) Y. Moroi, *Micelles (Theoretical and Applied Aspects)*, Springer Science + Business Media, New York, 1992; (c) G. N. Smith and J. Eastoe, *Phys. Chem. Chem. Phys.*, 2013, **15**, 424.
- J. Eastoe, in *Colloid science principles, methods and applications*, ed. T. Cosgrove, Wiley, New York, 2nd edn, 2010, ch. 4, pp. 61–90.
- I. Danielsson and B. Lindman, *Colloids Surf.*, 1981, **3**, 391.
- J. Eastoe, *Surfactant chemistry*, Wuhan University Press, China, 2005.
- D. Langevin, *Acc. Chem. Res.*, 1988, **21**, 255.
- E. Acosta, J. H. Harwell, J. F. Scamehorn and D. A. Sabatini, in *Handbook for cleaning/decontamination of surfaces*, ed. I. Johansson and P. Somasundaran, Elsevier B. V., Amsterdam, 1st edn, 2007, ch. 7, vol. 1, pp. 831–884.
- (a) A. K. Ganguli, A. Ganguly and S. Vaidya, *Chem. Soc. Rev.*, 2010, **39**, 474; (b) S. Eriksson, U. Nylen, S. Rojas and M. Boutonnet, *Appl. Catal., A*, 2004, **265**, 207; (c) M. Boutonnet, S. Lodgberg and E. Svensson, *Curr. Opin. Colloid Interface Sci.*, 2008, **13**, 270.
- V. C. Santana, F. D. Curbelo, T. N. Castro Dantas, A. A. Dantas Neto, H. S. Albuquerque and A. C. Garnica, *J. Pet. Sci. Eng.*, 2009, **66**, 117.
- C. W. Pouton, *Adv. Drug Delivery Rev.*, 1997, **25**, 47.
- R. G. Laughlin, *The aqueous phase behavior of surfactants*, Academic Press, San Diego, 1994.
- J. N. Israelachvili, D. J. Mitchell and B. W. Ninham, *J. Chem. Soc., Faraday Trans.*, 1976, **72**, 1525.
- (a) E. Ruckenstein and J. C. Chi, *J. Chem. Soc., Faraday Trans.*, 1975, **71**, 1690; (b) G. N. Smith, S. Alexander, P. Brown, D. A. Gillespie, I. Grillo, R. K. Heenan, C. James, R. Kemp, S. E. Rogers and J. Eastoe, *Langmuir*, 2014, **30**, 3422.
- S. P. Moulik and B. K. Paul, *Adv. Colloid Interface Sci.*, 1998, **78**, 99.
- P. A. Winsor, *Trans. Faraday Soc.*, 1948, **44**, 376.
- T. Hellweg, *Curr. Opin. Colloid Interface Sci.*, 2002, **7**, 50.
- M. Kahlweit, R. Strey and G. Busse, *J. Phys. Chem.*, 1990, **94**, 3881.
- G. J. McFann and K. P. Johnston, *J. Phys. Chem.*, 1991, **95**, 4889.
- A. Skauge and P. Fotland, *SPE Res. Eng.*, 1990, **5**, 601.
- L. E. Scriven, *Nature*, 1976, **263**, 123.
- S. H. Chen, *Phys. B*, 1986, **137**, 183.
- W. Meier, *Langmuir*, 1996, **12**, 6341.
- (a) B. Bagchi, *Water in Biological and Chemical Processes*, Cambridge University Press, Cambridge (UK), 2013;



- (b) H. MacDonald, B. Bedwell and E. Gulari, *Langmuir*, 1986, **2**, 704; (c) M. Valero, F. Sanchez, C. Gomez-Herrera and P. Lopez-Cornejo, *Chem. Phys.*, 2008, **345**, 65; (d) K. P. Johnston, K. L. Harrison, M. J. Clarke, S. M. Howdle, M. P. Heitz, F. V. Bright, C. Carlier and T. W. Randolph, *Science*, 1996, **271**, 624; (e) N. Zhou, Q. Li, J. Wu, J. Chen, S. Weng and G. Xu, *Langmuir*, 2001, **17**, 4505.
- 23 (a) J. Eastoe, B. H. Robinson, D. C. Steytler and D. Thorn-Leeson, *Adv. Colloid Interface Sci.*, 1991, **36**, 1; (b) M. J. Hou and D. O. Shah, *Langmuir*, 1987, **3**, 1086; (c) R. Leung and D. O. Shah, *J. Colloid Interface Sci.*, 1987, **120**, 320.
- 24 D. Liu, J. Ma, H. Cheng and Z. Zhao, *Colloids Surf., A*, 1998, **143**, 59.
- 25 P. Alexandridis, J. F. Holzwarth and T. A. Hatton, *J. Phys. Chem.*, 1995, **99**, 8222.
- 26 R. P. Bagwe and K. C. Khilar, *Langmuir*, 2000, **16**, 905.
- 27 J. N. Israelachvili, *Colloids Surf., A*, 1994, **91**, 1.
- 28 (a) J. Eastoe, J. Dong, K. J. Hetherington, D. Steytler and R. K. Heenan, *J. Chem. Soc., Faraday Trans.*, 1996, **92**, 65; (b) J. Eastoe, K. J. Hetherington, D. Sharpe, J. Dong, R. K. Heenan and D. C. Steytler, *Langmuir*, 1996, **12**, 3876.
- 29 A. Bumajdad, J. Eastoe, R. K. Heenan, J. R. Lu, D. Steytler and S. Egelhaaf, *J. Chem. Soc., Faraday Trans.*, 1998, **94**, 2143.
- 30 F. C. Frank, *Discuss. Faraday Soc.*, 1958, **25**, 19.
- 31 W. Helfrich, *Z. Naturforsch., C: J. Biosci.*, 1973, **28**, 693.
- 32 I. Szleifer, D. Kramer, A. Ben-Shaul, W. M. Gelbart and S. A. Safran, *J. Chem. Phys.*, 1990, **92**, 6800.
- 33 S. A. Safran, L. A. Turkevich and P. Pincus, *J. Phys. Lett.*, 1984, **45**, 69.
- 34 M. E. Cates, D. Andelman, S. A. Safran and D. Roux, *Langmuir*, 1988, **4**, 802.
- 35 J. Eastoe, D. Sharpe, R. K. Heenan and S. Egelhaaf, *J. Phys. Chem. B*, 1997, **101**, 944.
- 36 J. Eastoe, K. J. Hetherington, D. Sharpe, D. C. Steytler, S. Egelhaaf and R. K. Heenan, *Langmuir*, 1997, **13**, 2490.
- 37 S. Nave, PhD doctoral thesis, University of Bristol, 2001.
- 38 J. Eastoe, *Langmuir*, 1992, **8**, 1503.
- 39 J. Eastoe, G. Fragneto, B. H. Robinson, T. F. Towey, R. K. Heenan and F. Leng, *J. Chem. Soc., Faraday Trans.*, 1992, **88**, 461.
- 40 M. H. Hatzopoulos, J. Eastoe, P. J. Dowding and I. Grillo, *J. Colloid Interface Sci.*, 2013, **392**, 304.
- 41 O. Ghosh and C. A. Miller, *J. Phys. Chem.*, 1987, **91**, 4528.
- 42 C. La Messa, L. Coppola, G. A. Ranieri, M. Terenzi and G. Chidichimo, *Langmuir*, 1992, **8**, 2616.
- 43 W. Sager, W. Sun and H. F. Eicke, *Prog. Colloid Polym. Sci.*, 1992, **89**, 284.
- 44 H. Kunieda and K. Shinoda, *J. Colloid Interface Sci.*, 1979, **70**, 577.
- 45 K. Sato, Y. Hotta, T. Nagaoka, M. Yasuoka and K. Watari, *J. Mater. Sci.*, 2006, **41**, 5424.
- 46 C. A. Martin and L. J. Magid, *J. Phys. Chem.*, 1981, **85**, 3938.
- 47 P. Ekwall, L. Mandell and K. Fontell, *J. Colloid Interface Sci.*, 1970, **33**, 215.
- 48 B. Tamamushi and N. Watanabe, *Colloid Polym. Sci.*, 1980, **258**, 174.
- 49 A. M. Bellocq, *Langmuir*, 1998, **14**, 3730.
- 50 A. C. Hall, E. Tekle and Z. A. Schelly, *Langmuir*, 1989, **5**, 1263.
- 51 S. H. Chen, S. L. Chang and R. Strey, *J. Chem. Phys.*, 1990, **93**, 1907.
- 52 (a) J. Rouviere, J. M. Couret, M. Lindheimer, J. L. Dejardin and R. Mammy, *J. Chim. Phys.*, 1979, **76**, 289; (b) J. Peyrelasse and C. Boned, *J. Phys. Chem.*, 1985, **89**, 370.
- 53 (a) J. Texter, B. Antalek and A. J. Williams, *J. Chem. Phys.*, 1997, **106**, 7869; (b) J. Texter, *Colloids Surf., A*, 2000, **167**, 115; (c) B. Antalek, A. J. Williams, J. Texter, Y. Feldman and N. Garti, *Colloids Surf., A*, 1997, **128**, 1.
- 54 P. D. Fletcher, A. M. Howe and B. H. Robinson, *J. Chem. Soc., Faraday Trans.*, 1987, **83**, 985.
- 55 S. Nave, J. Eastoe, R. K. Heenan, D. C. Steytler and I. Grillo, *Langmuir*, 2000, **16**, 8741.
- 56 S. Nave, A. Paul, J. Eastoe, A. R. Pitt and R. K. Heenan, *Langmuir*, 2005, **21**, 10021.
- 57 S. Nave, J. Eastoe, R. K. Heenan, D. C. Steytler and I. Grillo, *Langmuir*, 2002, **18**, 1505.
- 58 F. D. Blum, S. Pickup, B. W. Ninham, S. J. Chen and D. F. Evans, *J. Phys. Chem.*, 1985, **89**, 711.
- 59 S. J. Chen, D. F. Evans, B. J. Ninham, D. J. Mitchell, F. D. Blum and S. Pickup, *J. Phys. Chem.*, 1986, **90**, 842.
- 60 K. Fontell, A. Ceglie, B. Lindman and B. J. Ninham, *Acta Chem. Scand.*, 1986, **40**, 247.
- 61 D. F. Evans, D. J. Mitchell and B. W. Ninham, *J. Phys. Chem.*, 1986, **90**, 2817.
- 62 M. Monduzzi, F. Caboi, F. Larche and U. Olsson, *Langmuir*, 1997, **13**, 2184.
- 63 B. W. Ninham, S. J. Chen and D. F. Evans, *J. Phys. Chem.*, 1984, **88**, 5855.
- 64 I. S. Barnes, S. T. Hyde, B. W. Ninham, P. J. Derian, M. Drifford and T. N. Zemb, *J. Phys. Chem.*, 1988, **92**, 2286.
- 65 M. Olla and M. Monduzzi, *Langmuir*, 2000, **16**, 6141.
- 66 M. Olla, M. Monduzzi and L. Ambrosone, *Colloids Surf., A*, 1999, **160**, 23.
- 67 (a) R. E. Verrall, S. Milioto and R. Zana, *J. Phys. Chem.*, 1988, **92**, 3939; (b) A. Jada, J. Lang and R. Zana, *J. Phys. Chem.*, 1990, **94**, 381; (c) A. Jada, J. Lang, R. Zana, R. Makhloufi, E. Hirsch and S. J. Candau, *J. Phys. Chem.*, 1990, **94**, 387.
- 68 G. G. Warr, R. Sen, D. F. Evans and J. E. Trend, *J. Phys. Chem.*, 1988, **92**, 774.
- 69 M. Kahlweit, R. Strey, P. Firman and D. Haase, *Langmuir*, 1985, **1**, 281.
- 70 S. Friberg, I. Lapczynska and G. Gillberg, *J. Colloid Interface Sci.*, 1976, **56**, 19.
- 71 K. Shinoda and H. Arai, *J. Phys. Chem.*, 1964, **68**, 3485.
- 72 K. Shinoda and H. Arai, *J. Colloid Interface Sci.*, 1967, **25**, 429.
- 73 K. Shinoda, *J. Colloid Interface Sci.*, 1967, **24**, 4.
- 74 V. N. Paunov, S. I. Sandler and E. W. Kaler, *Langmuir*, 2000, **16**, 8917.
- 75 T. Sottmann and R. Strey, *J. Chem. Phys.*, 1997, **106**, 8606.
- 76 M. Gradzielski, D. Langevin and B. Farago, *Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys.*, 1996, **53**, 3900.



- 77 J. J. Taber, *Pure Appl. Chem.*, 1980, **52**, 1323.
- 78 R. Aveyard, B. P. Binks, S. Clark and J. Mead, *J. Chem. Soc., Faraday Trans.*, 1986, **82**, 125.
- 79 K. S. Chan and D. O. Shah, *J. Dispersion Sci. Technol.*, 1980, **1**, 55.
- 80 B. P. Binks, H. Kellay and J. Meunier, *Europhys. Lett.*, 1991, **16**, 53.
- 81 H. Kellay, B. P. Binks, Y. Hendriks, L. T. Lee and J. Meunier, *Adv. Colloid Interface Sci.*, 1994, **49**, 85.
- 82 H. Kellay, J. Meunier and B. P. Binks, *Phys. Rev. Lett.*, 1993, **70**, 1485.
- 83 R. Aveyard, B. P. Binks, T. Lawless and J. Mead, *J. Chem. Soc., Faraday Trans.*, 1985, **81**, 2155.
- 84 J. Eastoe and D. Sharpe, *Langmuir*, 1997, **13**, 3289.
- 85 A. Kitahara, T. Kobayashi and T. Tachibana, *J. Phys. Chem.*, 1962, **66**, 363.
- 86 C. J. O'Connor, T. D. Lomax and R. E. Ramage, *Adv. Colloid Interface Sci.*, 1984, **20**, 21.
- 87 H. F. Eicke, *Top. Curr. Chem.*, 1980, **87**, 85.
- 88 J. B. Peri, *J. Colloid Interface Sci.*, 1969, **29**, 6.
- 89 M. Kotlarchyk, J. S. Huang and S. H. Chen, *J. Phys. Chem.*, 1985, **89**, 4382.
- 90 K. Kon-no and A. Kitahara, *J. Colloid Interface Sci.*, 1971, **35**, 636.
- 91 W. F. Pacynko, J. Yarwood and G. J. Tiddy, *J. Chem. Soc., Faraday Trans.*, 1989, **85**, 1397.
- 92 R. Zana, *Colloids Surf., A*, 1997, **123–124**, 27.
- 93 J. C. Ravey, M. Buzier and C. Picot, *J. Colloid Interface Sci.*, 1984, **97**, 9.
- 94 E. Ruckenstein and R. Nagarajan, *J. Phys. Chem.*, 1980, **84**, 1349.
- 95 M. J. Hollamby, R. Tabor, K. J. Mutch, K. Trickett, J. Eastoe, R. K. Heenan and I. Grillo, *Langmuir*, 2008, **24**, 12235.
- 96 G. N. Smith, P. Brown, C. James, S. E. Rogers and J. Eastoe, *Colloids Surf., A*, 2016, **494**, 194.
- 97 J. S. Huang, S. A. Safran, M. W. Kim, G. S. Grest, M. Kotlarchyk and N. Quirke, *Phys. Rev. Lett.*, 1984, **53**, 592.
- 98 T. Spehr, B. Frick, I. Grillo, P. Falus, M. Muller and B. Stuhn, *Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys.*, 2009, **79**, 031404.
- 99 R. Nagarajan, *Langmuir*, 2002, **18**, 31.
- 100 L. S. Ornstein and F. Zernike, *Proct. Sect. Sci. K. Med. Akad. Wet.*, 1914, **17**, 793.
- 101 S. G. Frank and G. Zografis, *J. Pharm. Sci.*, 1969, **58**, 993.
- 102 S. Nave, J. Eastoe and J. Penfold, *Langmuir*, 2000, **16**, 8733.
- 103 M. Gradzielski, D. Langevin, T. Sottmann and R. Strey, *J. Chem. Phys.*, 1997, **106**, 8232.
- 104 F. Sicoli, D. Langevin and L. T. Lee, *J. Chem. Phys.*, 1993, **99**, 4759.
- 105 T. Sottmann, R. Strey and S. H. Chen, *J. Chem. Phys.*, 1997, **106**, 6483.
- 106 M. Almgren, R. Johannsson and J. C. Eriksson, *J. Phys. Chem.*, 1993, **97**, 8590.
- 107 J. P. Cason, M. E. Miller, J. B. Thompson and C. B. Roberts, *J. Phys. Chem. B*, 2001, **105**, 2297.
- 108 F. M. Menger and G. Saito, *J. Am. Chem. Soc.*, 1978, **100**, 4376.
- 109 A. Maitra, *J. Phys. Chem.*, 1984, **88**, 5122.
- 110 E. Keh and B. Valeur, *J. Colloid Interface Sci.*, 1981, **79**, 465.
- 111 R. Aveyard, B. P. Binks, P. Cooper and P. D. Fletcher, *Adv. Colloid Interface Sci.*, 1990, **33**, 59.
- 112 B. Lemaire, P. Bothorel and D. Roux, *J. Phys. Chem.*, 1983, **87**, 1023.
- 113 L. Arleth and J. S. Pedersen, *Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys.*, 2001, **63**, 061406.
- 114 B. H. Robinson, C. Toprakcioglu, J. C. Dore and P. Chieux, *J. Chem. Soc., Faraday Trans.*, 1984, **80**, 13.
- 115 M. Hirai, R. K. Hirai, M. Sanada, H. Iwase and S. Mitsuya, *J. Phys. Chem. B*, 1999, **103**, 9658.
- 116 J. Eastoe, W. K. Young, B. H. Robinson and D. C. Steytler, *J. Chem. Soc., Faraday Trans.*, 1990, **86**, 2883.
- 117 C. L. Kitchens, D. P. Bossev and C. B. Roberts, *J. Phys. Chem. B*, 2006, **110**, 20392.
- 118 H. Kellay and J. Meunier, *J. Phys.: Condens. Matter*, 1996, **8**, 49.
- 119 M. J. Hou, M. Kim and D. O. Shah, *J. Colloid Interface Sci.*, 1988, **123**, 398.
- 120 J. Lang, A. Jada and A. Malliaris, *J. Phys. Chem.*, 1988, **92**, 1946.
- 121 S. Balakrishnan, N. Javid, H. Weingartner and R. Winter, *ChemPhysChem*, 2008, **9**, 2794.
- 122 R. V. Sharma and K. C. Sharma, *Phys. A*, 1977, **89**, 213.
- 123 E. W. Kaler, J. F. Billman, J. L. Fulton and R. D. Smith, *J. Phys. Chem.*, 1991, **95**, 458.
- 124 J. M. Tingey, J. L. Fulton and R. D. Smith, *J. Phys. Chem.*, 1990, **94**, 1997.
- 125 J. S. Huang, *J. Chem. Phys.*, 1985, **82**, 480.
- 126 A. M. Cazabat, D. Langevin, J. Meunier and A. Pouchelon, *J. Phys. Lett.*, 1982, **43**, L89.
- 127 M. Kotlarchyk, S. H. Chen and J. S. Huang, *Phys. Rev. A: At., Mol., Opt. Phys.*, 1983, **28**, 508.
- 128 C. Toprakcioglu, J. C. Dore, B. H. Robinson, A. Howe and P. Chieux, *J. Chem. Soc., Faraday Trans.*, 1984, **80**, 413.
- 129 S. H. Chen, J. Rouch, F. Sciotino and P. Tartaglia, *J. Phys.: Condens. Matter*, 1994, **6**, 10855.
- 130 I. Chakraborty and S. P. Moulik, *J. Colloid Interface Sci.*, 2005, **289**, 530.
- 131 M. Gradzielski, D. Langevin, T. Sottmann and R. Strey, *J. Chem. Phys.*, 1996, **104**, 3782.
- 132 M. Appel, T. Spehr, R. Wipf and B. Stuhn, *J. Colloid Interface Sci.*, 2012, **376**, 140.
- 133 L. Garcia-Rio, A. Godoy and P. Rodriguez-Dafonte, *Eur. J. Org. Chem.*, 2006, 3364.
- 134 R. A. Day, B. H. Robinson, J. H. Clarke and J. V. Doherty, *J. Chem. Soc., Faraday Trans.*, 1979, **75**, 132.
- 135 J. H. Clarke, J. D. Nicholson and K. N. Regan, *J. Chem. Soc., Faraday Trans.*, 1985, **81**, 1173.
- 136 A. Salabat, J. Eastoe, K. J. Mutch and R. F. Tabor, *J. Colloid Interface Sci.*, 2008, **318**, 244.
- 137 O. Myakonkaya, J. Eastoe, K. J. Mutch, S. Rogers, R. Heenan and I. Grillo, *Langmuir*, 2009, **25**, 2743.
- 138 R. W. Gale, J. L. Fulton and R. D. Smith, *J. Am. Chem. Soc.*, 1987, **109**, 920.



- 139 J. L. Fulton and R. D. Smith, *J. Phys. Chem.*, 1988, **92**, 2903.
- 140 J. Eastoe, D. C. Steytler, B. H. Robinson and R. K. Heenan, *J. Chem. Soc., Faraday Trans.*, 1994, **90**, 3121.
- 141 A. A. Padua, J. M. Fareleira, J. C. Calado and W. A. Wakeham, *J. Chem. Eng. Data*, 1996, **41**, 1488.
- 142 W. J. Tay and J. M. Trusler, *J. Chem. Thermodyn.*, 2018, **124**, 107.
- 143 E. J. Beckman and R. D. Smith, *J. Phys. Chem.*, 1990, **94**, 3729.
- 144 J. Oshitani, S. Takashina, M. Yoshida and K. Gotoh, *Langmuir*, 2010, **26**, 2274.
- 145 E. Y. Shew, P. L. Nostro, G. Capuzzi and P. Baglioni, *Langmuir*, 1999, **15**, 6671.
- 146 C. M. Dunn, B. H. Robinson and F. J. Leng, *Spectrochim. Acta, Part A*, 1990, **46**, 1017.
- 147 J. Eastoe, G. Fragneto, D. C. Steytler, B. H. Robinson and R. K. Heenan, *Physica B*, 1992, **180**, 555.
- 148 J. Eastoe, D. C. Steytler, B. H. Robinson, R. K. Heenan, A. N. North and J. C. Dore, *J. Chem. Soc., Faraday Trans.*, 1994, **90**, 2497.
- 149 J. Eastoe, T. F. Towey, B. H. Robinson, J. Williams and R. K. Heenan, *J. Phys. Chem.*, 1993, **97**, 1459.
- 150 J. Tanori, T. G. Krzywicki and M. P. Pileni, *Langmuir*, 1997, **13**, 632.
- 151 P. Brown, C. Butts, R. Dyer, J. Eastoe, I. Grillo, F. Guittard, S. Rogers and R. K. Heenan, *Langmuir*, 2011, **27**, 4563.
- 152 (a) J. Eastoe, B. H. Robinson and R. K. Heenan, *Langmuir*, 1993, **9**, 2820; (b) R. E. Riter, E. P. Undiks and N. P. Levinger, *J. Am. Chem. Soc.*, 1998, **120**, 6062; (c) J. Texter, L. Ge, T. H. Mourey and T. G. Bryan, *Langmuir*, 2004, **20**, 11288; (d) F. M. Pavel and R. A. Mackay, *Langmuir*, 2000, **16**, 8568.
- 153 A. Czajka, C. Hill, J. Peach, J. C. Pegg, I. Grillo, F. Guittard, S. E. Rogers, M. Sagisaka and J. Eastoe, *Phys. Chem. Chem. Phys.*, 2017, **19**, 23869.
- 154 S. Prevost, M. Gradzielski and T. Zemb, *Adv. Colloid Interface Sci.*, 2017, **247**, 374.
- 155 D. G. Peck, R. S. Schechter and K. P. Johnston, *J. Phys. Chem.*, 1991, **95**, 9541.
- 156 P. J. Flory, *J. Chem. Phys.*, 1941, **9**, 660.
- 157 H. Reiss, W. K. Kegel and J. Groenwold, *Ber. Bunsen-Ges.*, 1996, **100**, 279.
- 158 E. Negro, R. Latsuzbaia, A. H. de Vries and G. M. Koper, *Soft Matter*, 2014, **10**, 8685.
- 159 C. A. Bearchell and D. M. Heyes, *Phys. Chem. Chem. Phys.*, 2001, **3**, 5255.
- 160 Y. Fu, S. Xiao, S. Liu, Y. Chang, R. Ma, Z. Zhang and J. He, *Langmuir*, 2022, **38**, 3129.
- 161 B. Lukanov and A. Firoozabadi, *Langmuir*, 2016, **32**, 3100.

