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Minimum surfactant concentration required for inducing self-shaping of oil droplets and competitive adsorption effects†

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Surfactant choice is key in starting the phenomena of artificial morphogenesis, the bottom-up growth of geometric particles from cooled emulsion droplets, as well as the bottom-up self-assembly of rechargeable microswimmer robots from similar droplets. The choice of surfactant is crucial for the formation of a plastic phase at the oil–water interface, for the kinetics, and for the onset temperature of these processes. But further details are needed to control these processes for bottom-up manufacturing and understand their molecular mechanisms. Still unknown are the minimum concentration of the surfactant necessary to induce the processes, or competing effects in a mixture of surfactants when only one is capable of inducing shapes. Here we systematically study the effect of surfactant nature and concentration on the shape-inducing behaviour of hexadecane-in-water emulsions with both cationic (CTAB) and non-ionic (Tween, Brij) surfactants over up to five orders of magnitude of concentration. The minimum effective concentration is found approximately equal to the critical micelle concentration (CMC), or the solubility limit below the Krafft point of the surfactant. However, the emulsions show low stability at the vicinity of CMC. In a mixed surfactant experiment (Tween 60 and Tween 20), where only one (Tween 60) can induce shapes we elucidate the role of competition at the interface during mixed surfactant adsorption by varying the composition. We find that a lower bound of ~75% surface coverage of the shape-inducing surfactant with C14 or longer chain length is necessary for self-shaping to occur. The resulting technique produces a clear visual readout of otherwise difficult to investigate molecular events. These basic requirements (minimum concentration and % surface coverage to induce oil self-shaping) and the related experimental techniques are expected to guide academic and industrial scientists to formulations with complex surfactant mixtures and behaviour.

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1. Introduction

Bottom-up fabrication promises to replace energy- and material-wasteful top-down processes such as lithography with sustainable,

scalable processes, though the types of shapes and control over them have been limited.^{1–3} The recent discovery of artificial morphogenesis can lead to scalable manufacturing of particles of a large variety of regular geometric shapes, simply by cooling down emulsion droplets.^{4–6} These include polymer particles^{7,8} in the shapes of polyhedra, hexagons, rhomboids, triangles, rods, and fibres. The mild conditions and lack of subtractive steps promise a sustainable and efficient process for fabrication by bottom-up growth. We have control of the size of these shapes over orders of magnitude, from as small as 50 nm to over 1 mm.⁷ On a fundamental level, the process is started by the formation of a few layers of plastic crystal (rotator phase) in the oil just near the water interface, which exert stress opposing the interfacial tension and generate anisotropic shapes in an established sequence.^{9,10} The formation of the plastic crystal phase depends critically on the molecular properties of the surfactant used. Alternative mechanisms have been discussed in the literature. Garcia-Aguilar *et al.*

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proposed an elastic surface model;¹¹ we compared this hypothesis to ours.¹²

We have described the length of the surfactant tail needed to induce the transition, and have classified the behaviour of many classes of surfactants according to their ability to generate the shapes at a constant surfactant concentration.⁵ We have successfully modelled the shape evolution sequence as a competition between the fundamental thermodynamic driving forces for the formation of the plastic crystal phase and the opposing increase of surface energy due to interfacial tension.^{9,10} We have recently also shown that the phenomenon is quite general, with many classes of oils showing this behaviour,⁵ and also that even oils that do not by themselves undergo such changes can be induced to do so in mixtures with other oils.¹³ The phenomenon was demonstrated also for water-in-oil emulsions and at flat water/oil interfaces.^{14,15} We have only recently developed the tools of molecular dynamics to simulate liquid–solid and solid–solid crystal–rotator phase transitions that may allow soon to reveal the molecular details of this process.¹⁶ Still, many details of the mechanism remain unknown¹³ and behaviour is hard to predict. For example, small changes in surfactant structure or difference between surfactant tail and oil, have caused the emergence of self-assembled swimmers.¹⁴

One of the open questions is about the minimum concentration of surfactant and the type of packing at the surface that is necessary for the plastic crystal templating to occur.^{17,18} There are even more complicated questions of how mixtures of surfactants would affect the packing on the surface and rotator phase formation.

In this report, we conduct systematic experiments into the minimum surfactant concentration necessary to induce the self-shaping. Two classes of surfactants are studied: non-ionic families Tween (Tween 40 and Tween 60) and Brij (Brij S10, S20, C10 and Brij58) and cationic hexadecyltrimethylammonium bromide (C₁₆TAB). For all the surfactants we show the minimum concentration necessary to induce the transformation, with or without salt in the solution, is close to the critical micellar concentration (CMC) (Fig. 1). Near the CMC the droplets can only transform to the initial shapes of the prescribed shape evolution sequence. At higher concentrations, they transform to the end of the sequence (Fig. 1). We also perform competitive adsorption experiments with binary mixtures of surfactants, only one of which can induce self-shaping.

2. Experimental section

2.1. Materials

As dispersed oily phase in the studied emulsions, we used the linear alkane hexadecane (C₁₆), obtained from Sigma-Aldrich with purity $\geq 99\%$, melting temperature $T_m = 18\text{ }^\circ\text{C}$, used as received, without further purification.

To stabilize the emulsions, we used a variety of nonionic and ionic surfactants. The nonionic surfactants were from two main groups Tween and Brij surfactants. Polyoxyethylene sorbitan monoalkylates (Tween surfactants, C_nSorbEO₂₀) differed in the

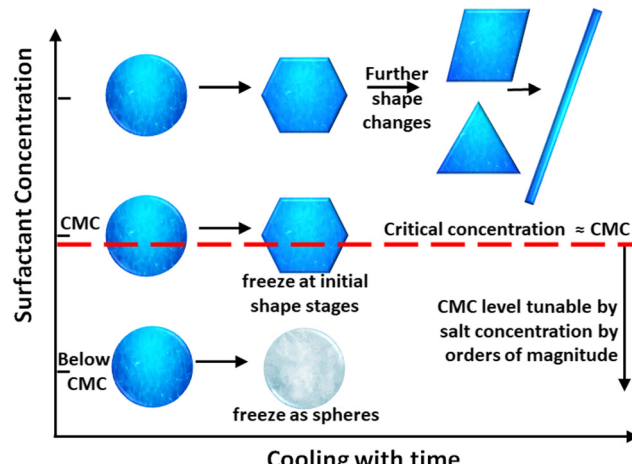


Fig. 1 Schematic representation of the experiment. Hexadecane emulsions were prepared in aqueous surfactant solution. Upon cooling, when below the CMC (or below solubility, in the case of CTAB), droplets freeze as spheres. At or near CMC they exhibit shape changes on cooling. CMC and therefore surfactant packing on the surface is important, as the same absolute concentration of surfactant that could not induce shape changes, does in the presence of salt where CMC is lower. At high concentrations ($>$ several times CMC), droplets deform all the way along the shape-change sequence we have established for the process of artificial morphogenesis.⁴

length of their alkyl chain, n , varied between 12 and 18 (Tween 20, Tween 40, and Tween 60); the number of oxyethylene units, m , is ~ 20 . The polyoxyethylene alkyl ethers (Brij surfactants, C_nEO_m) were with $n = 16$ or 18 carbon atoms and $m = 10$ or 20 oxyethylene units. We used a cationic surfactant with hexadecyl tail – C₁₆TAB. Detailed information for the purity, HLB values of the nonionic surfactants and surfactant producers is presented in the ESI,† Table S1. All surfactants were used as received. For experiments in the presence of salt we used NaBr, a product of Sigma with purity $\geq 99.5\%$.

As a fluorescent dye, we used the oil soluble Solvent Green 5 (diisobutyl perylenedicarboxylate), produced by TCI Chemicals. The addition of the fluorescent dye brings two main benefits: (1) helps to observe some finer structures and eliminates the stains on the background, (2) acts as an indicator in the membrane emulsification process (see Methods below), distinguishing droplets from small air bubbles. The concentration of fluorescent dye in C₁₆ drops is 0.1 wt%.

All aqueous solutions were prepared with deionized water, which was purified by an Elix 3 module (Millipore, USA).

2.2. Methods

2.2.1. Preparation of the initial emulsions. The initial alkane-in-water emulsions were prepared by three different experimental procedures. In the first we prepare each emulsion at the specified concentration, though the emulsion stability is poorer. In the second we prepare a stable emulsion at high concentration and dilute it as necessary. In the third procedure, the double syringe technique was applied for the preparation of emulsions.



Experimental procedure 1: preparation of polydisperse emulsion droplets in target solution concentration. First, we prepared an aqueous surfactant solution with fixed surfactant concentration. Afterwards, we added the alkane phase into the aqueous phase. The alkane-in-water emulsions by the first experimental procedure were prepared by homogenizing the disperse phase on a magnetic stirrer for 15 minutes and after that, a specimen of the studied emulsion was placed in a capillary for the optical observations. In this procedure, the experiments with the emulsions were performed without further dilution of the droplets. The droplet size range is 2–80 μm .

Experimental procedure 2: preparation of monodisperse emulsion droplets in stock solution to be diluted later. The second set of emulsions were prepared by a laboratory Microkit membrane emulsification module from Shirasu Porous Glass Technology (SPG, Miyazaki, Japan). The oil phase was passed through tubular glass membranes with outer diameter of 10 mm and working area of approximately 3 cm^2 . This method allows production of drops with relatively narrow drop size distribution.^{19–22} Membranes with mean pore diameter of 5 μm were used for preparing initial emulsions resulting in drop diameters of around 15–20 μm . There are several advantages of the membrane emulsification method compared with classical emulsification: (1) narrow size distribution of droplets, (2) little heat generated due to no high pressure/shear stress required, (3) the energy input is quite low while the energy efficiency is high.

In this experimental procedure, we kept the surfactant concentration sufficiently high during preparation of the monodisperse emulsion drops, *i.e.* 1.5 wt% for the non-ionic surfactants and 0.5 wt% for the cationic surfactant. After that, we diluted the alkane-in-oil emulsion with distilled water until the desired surfactant concentration was reached. In addition, we stirred gently the sample for 15 minutes to allow for surfactant redistribution between the solution and droplets.

The main difference between these two experimental procedures is the initial surfactant concentration in which the C_{16} droplets are emulsified.

Experimental procedure 3: preparation of monodisperse emulsion droplets using double syringe technique. Emulsions stabilized by Brij surfactants were prepared by making an aqueous surfactant solution at the final concentration, adding the required amount of oil, and passing the mixture between two syringes, connected by a double-sided Luer-Lock 18 gauge micro-emulsifying needle (stainless steel, 18G*2-7/8 inches with reinforcing bar, Cadence Science, Inc.). The syringe sizes were 5 cm^3 plastic syringes with Luer Lock connectors, and care was taken to remove all air from the system, including the needle, before emulsification. The mixture was passed at high speed through the needle between the two syringes. 15–50 passes were made, depending on the surfactant concentration and the resulting emulsion was used as obtained, without further dilution.

2.2.2. Optical observations of the drop shape transformations. For microscope observations, a specimen of the studied emulsion was placed in a capillary with rectangular cross section: 50 mm

length, 1 mm width, and 0.1 mm height. The capillary was enclosed within a custom-made metal cooling chamber, with optical windows for microscope observation. The chamber temperature was controlled by cryo-thermostat (JULABO CF30, Cryo-Compact Circulator) and measured close to the emulsion location, using a calibrated thermo-couple probe with an accuracy of ± 0.2 $^\circ\text{C}$. The initial temperature for cooling droplets was 20 $^\circ\text{C}$. The thermo-probe was inserted in one of the orifices of the aluminum thermostatic chamber and mounted in the position where a capillary with the emulsion sample would be normally placed for microscope observations. In the neighboring orifice, the actual capillary with the emulsion sample was placed. The correct measurement of the temperature was ensured by calibrating the thermo-couple with a precise mercury thermometer in the respective range of temperatures measured. Furthermore, we always observe melting of the frozen particles at temperatures within ± 0.2 $^\circ\text{C}$ of the melting temperature of the bulk oil, reported in the literature.

We also prepared another set of experiments in which the temperature of the sample is controlled by a Linkam cooling system for regulating temperature and cooling rate. The temperature limits and cooling rate can be set up on the control panel, with temperature ranging from -25 $^\circ\text{C}$ to 120 $^\circ\text{C}$ and cooling rate ranging from 0.01 K min^{-1} to 20 K min^{-1} .

The optical observations were performed with AxioImager.M2m (Zeiss, Germany) and Olympus BX51 microscopes. Long-focus objectives 20 \times , 40 \times , 50 \times , 60 \times and 100 \times were used. For the experimental observations performed in transmitted, cross-polarized white light, we used λ (compensator) plate, situated after the sample and before the analyzer, at 45° with respect to both the analyzer and the polarizer. Under these conditions the liquid background and the fluid objects have typical magenta color, whereas the birefringent areas appear brighter and may have intense colors.^{23,24} For the experiments with emulsion droplets containing a fluorescent dye a reflective light was used.

2.2.3. Interfacial tension (IFT) measurement. The pendant drop tests were performed using Kruss Tensiometer (DSA 100) to measure the IFT and estimate the CMC of the surfactants. A calibration test was performed by surface tension measurement of deionized water before the start of each set of analysis. A glass cuvette made of optical glass and a hooked needle (0.74 mm diameter) were used to measure IFT between hexadecane/aqueous phases.

3. Results and discussion

3.1. Surfactant concentration effects on self-shaping

We present results obtained with C_{16} emulsion droplets stabilized by different surfactants, with different surfactant concentrations, and in the presence and absence of salt. In Section 3.2 we describe the surfactant concentration effects for cationic surfactant on the self-shaping phenomena. In Section 3.3 we describe these effects for non-ionic surfactants. The competitive adsorption effects on self-shaping in mixtures of nonionic surfactants are presented in Section 3.4.



3.2. Concentration effects for cationic surfactant

We set out to find the minimum surfactant concentration for the cationic surfactant cetyltrimethylammonium bromide ($C_{16}\text{TAB}$) that induces the self-shaping behaviour of the hexadecane (C_{16}) droplets. For ionic surfactants, both solubility limit and CMC are greatly affected by concentration of the added salt.^{25,26} Therefore, we determined the minimum required surfactant concentration for self-shaping of the droplets both in the absence and in presence of salt. We performed one set of experiments with surfactant in pure water and another set with the same surfactant in the presence of 100 mM NaBr.

$C_{16}\text{TAB}$ is below its Krafft temperature in both cases. Thermal hysteresis is, however, very common for this surfactant – it can exist as metastable micelles for days even at concentrations 20 times higher than the solubility limit, 10 °C below the Krafft temperature.²⁷ To determine the CMC/solubility value in each case, we measured the interfacial tension of the hexadecane/water interface as a function of the surfactant concentration. According to the data from the literature, the CMC of $C_{16}\text{TAB}$ at room temperatures in the absence of salt is around 0.98 mM and in presence of 100 mM NaBr is ~ 0.04 mM.²⁸ The solubility of the surfactant is difficult to measure, but at the Krafft temperature (26.6 °C in the absence of salt),²⁷ it is equal to CMC, therefore, the solubility product is $K_s = [C_{16}\text{TA}^+][\text{Br}^-] = (0.98 \text{ mM})^2$. At $[\text{Br}^-] = 100 \text{ mM}$, this corresponds to solubility $[C_{16}\text{TA}^+] = K_s/[\text{Br}^-] = 0.01 \text{ mM}$. The results that we obtained from the breakpoint of the measured interfacial tension *vs.* $[C_{16}\text{TA}^+]$ are in better agreement with the CMC values: 0.99 mM for

the system without salt and 0.03 mM in the presence of 100 mM NaBr.

Below the breakpoint (be it solubility limit or CMC), the C_{16} emulsion drops do not transform their shape at all and freeze into spherical shape at temperatures between 8–10 °C, see Fig. 2A for $[C_{16}\text{TAB}] = 0.75 \text{ mM}$, no salt. Similar results were obtained with 0.8 and 0.9 mM $C_{16}\text{TAB}$. At 1.0 mM, we are able to observe drop shape transformations, see Fig. 2B, where emulsion drops with $d_{\text{ini}} \approx 10\text{--}25 \mu\text{m}$ reached the initial 3D shapes of the transformation, and only transformed to the next flat platelet stage if we decreased the cooling rate to $\approx 0.2 \text{ K min}^{-1}$. When we increase the surfactant concentration 10 times (10 mM $C_{16}\text{TAB}$), we are able to observe formation of flat platelets which are extruding long thin fibres, see Fig. 2C. Droplets with smaller initial drop size diameter ($< 10 \mu\text{m}$) are able to reach the final stages of the evolutionary scheme forming ellipsoidal droplets connected with long thin fibres even at 1 mM concentrations.

Similarly, in the case of salt added, we first observed deformations around the interfacial tension breakpoint, though in this case at much lower surfactant concentrations, because the addition of salt decreases both CMC and solubility of the surfactant. At surfactant concentration below 0.04 mM no shape changes are observed, while in the range between 0.04 mM and 0.2 mM, we are able to observe deformations only in the small droplets ($d_{\text{ini}} \leq 7 \mu\text{m}$), see Fig. 2D. At higher concentrations ($\geq 0.3 \text{ mM}$, prepared by heating and cooling back to room temperature), the bigger droplets also start to transform, see Fig. 2E and F. The obtained frozen particles are

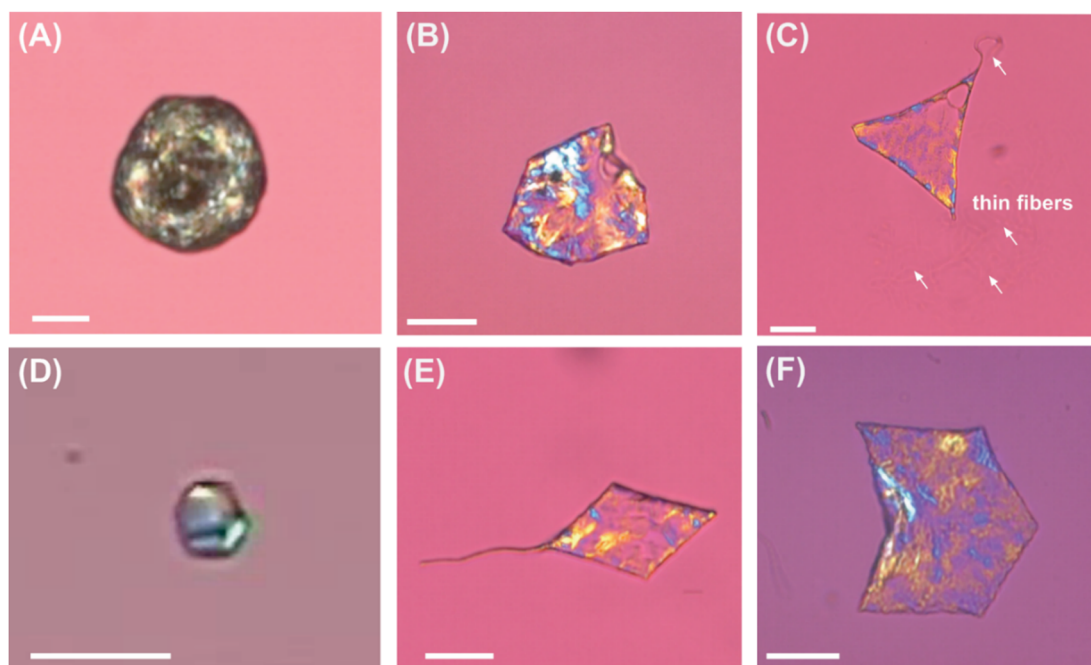


Fig. 2 Microscopic pictures of frozen C_{16} particles stabilized by $C_{16}\text{TAB}$ with different surfactant concentrations. (A)–(C) In the absence of salt, $\text{CMC} \approx 0.98 \text{ mM}$: (A) 0.75 mM, (B) 1.0 mM and (C) 10 mM of $C_{16}\text{TAB}$. By increasing the surfactant concentration, we increase the probability of formation of different flat shapes. At 0.75 mM $C_{16}\text{TAB}$, droplets freeze into a spherical shape, while at 10 mM $C_{16}\text{TAB}$ we observe formation of flat shapes, which are extruding long thin fibres. Cooling rate 0.44 K min^{-1} . Scale bar $20 \mu\text{m}$. (D)–(F) In the presence of salt (100 mM NaBr), $\text{CMC} \approx 0.04 \text{ mM}$: (D) 0.05 mM, (E) 0.4 mM and (F) 0.5 mM $C_{16}\text{TAB}$. Cooling rate 0.44 K min^{-1} . Scale bar $20 \mu\text{m}$.



identical to those in the absence of NaBr. For example, close to CMC/solubility limit, the drops are only able to transform their shape to the initial stages of the evolutionary scheme, forming regular polyhedral and other faceted 3D shapes. At higher surfactant concentration (≥ 0.4 mM C₁₆TAB, Fig. 2B), the drops are able to transform their shape into flat platelets (rhomboid, triangular and hexagonal prisms) extruding long thin fibres. With this series of experiments we showed that by the addition of an electrolyte to the water phase, we are able to reduce the concentration of the ionic surfactant necessary to produce shapes from 1 mM to 0.04 mM (~ 25 fold).

3.3. Surfactant concentration effects for non-ionic surfactants

Tweens were chosen for these experiments as they are edible, biocompatible and are the most common commercial non-ionic class of surfactants. They are polysorbate molecules each containing a hydrophilic head group of oligo (ethylene glycol) chains and a hydrophobic tail of fatty acid ester moiety (Table S1, ESI†).²⁹ They are used as emulsifiers and solubilizers from pharmaceuticals to food products, including bread, cake mixes, salad dressings, shortening oil and chocolate. In addition, they are often used in cosmetics to solubilize essential oils into water-based products. Both Tween 40 (approx. 90%

C₁₆SorbEO₂₀ + 10% C₁₈SorbEO₂₀) and Tween 60 (approx. 50% C₁₆SorbEO₂₀ + 50% C₁₈SorbEO₂₀) have been shown previously to induce self-shaping in hexadecane emulsion droplets.

In our general classification of shape-inducing surfactants⁵ these two fall into Group B. For this group, shape transformations start around the oil melting temperature, *i.e.* $T_d \approx T_m$. They are able to form very strong (thick) interfacial layers and typically form hexagonal and other platelets, as well as relatively stiff rods. These include formation of perforated fluid platelets that maintain their outer shape integrity after the thin liquid film in their middle spontaneously breaks. All previously reported results for polysorbate surfactants are obtained at fairly high surfactant concentration (1.5 wt% surfactant concentration, which is 11.75 mM for Tween 40 and 11.56 mM for Tween 60)^{4,5,30–32} and the minimum concentration that induces transitions is unknown.

The Brij-class alcohol ethoxylates C₁₈EO₁₀ (Brij S10), C₁₈EO₂₀ (S20), C₁₆EO₁₀ (C10), and C₁₆EO₂₀ (Brij58) are well suited to align efficiently at liquid interfaces, to support stabilization of dispersed phases, commonly oil-in-water emulsions. They also induce shape transformations and were previously⁵ classified as Group A (C₁₈EO₁₀, C₁₈EO₂₀, C₁₆EO₁₀; induce shapes above the melting temperature of the oil) and Group C (C₁₆EO₂₀, like C₁₆TAB, induces shapes below the melting temperature).

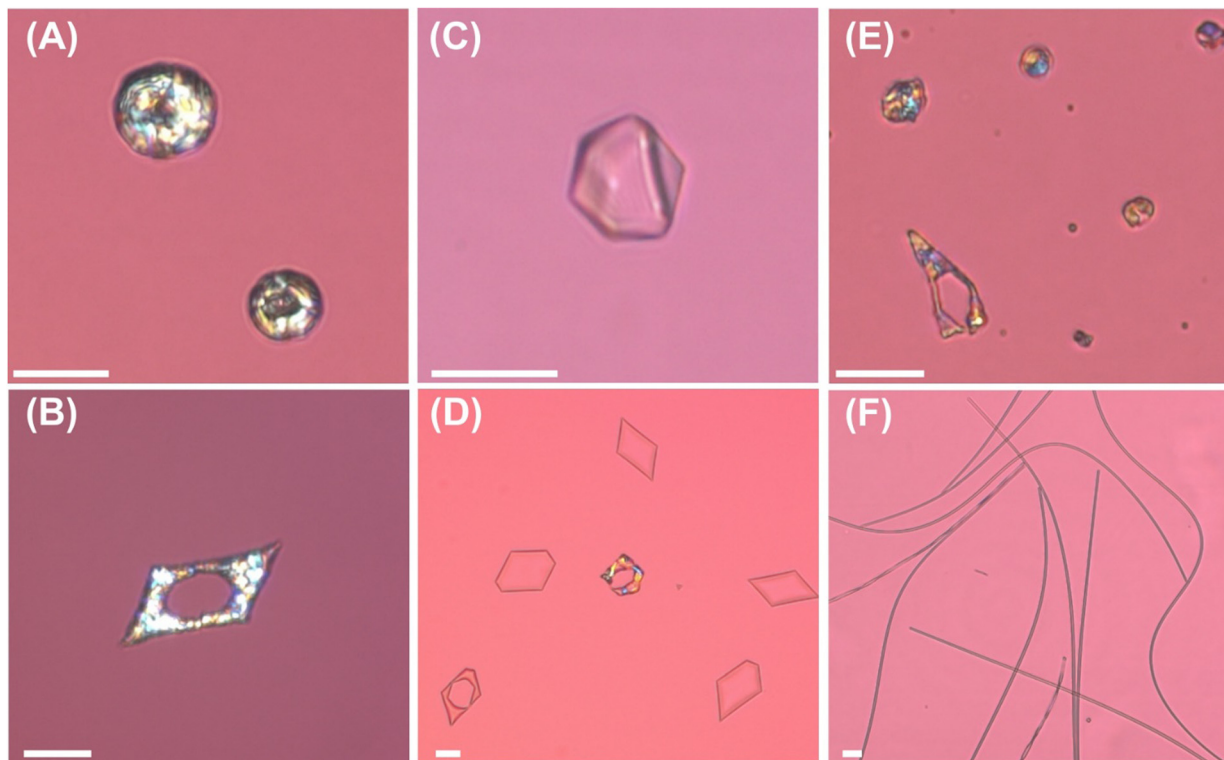


Fig. 3 Drop shape transformations for emulsions prepared from C₁₆ oil drops stabilized by Tween 40 (C₁₆SorbEO₂₀) by two different procedures. Images on the first row are obtained with emulsion droplets prepared by adding the oil phase into the aqueous surfactant solution with fixed surfactant concentration. The images on the second row are obtained with emulsion droplets prepared by membrane emulsification method in 1.5 wt% surfactant solution and then diluted with water to the desired surfactant concentration, $d_{ini} \approx 15 \mu\text{m}$. Microscope images of (A) frozen spherical drops; (B) frozen, perforated parallelogram flat particle; (C) fluid deformed particle which freeze into the same shape; (D) flat parallelogram particles – the particle in the centre of the picture is frozen; (E) frozen faceted 3D particles; (F) frozen long fibres. The total surfactant concentration in the emulsion is (A and B) 0.01 mM and (C–F) 0.1 mM. The cooling rate is (A–D) 0.42 K min^{−1} and (E and F) 0.25 K min^{−1}. Scale bars: 20 μm .



We measured the interfacial tension on the hexadecane/water interface as a function of the surfactant concentration at fixed temperature. The values we determined for CMC are 0.022 mM for Tween 40 and 0.011 mM for Tween 60. These are close to values reported in the literature at this temperature (0.033 mM for Tween 40 and 0.0167 mM for Tween 60),³³ though neither of the values is very reliable, and do change for both different suppliers and batches. While the values are not exact, they are by almost 3 orders of magnitude lower than the concentrations at which the self-shaping was previously studied (0.022 mM of Tween 40 is ~ 500 times more dilute than the initial 1.5 wt% sample). Just like for the other surfactants, we expect we would see shape changes near these values of CMC. And the CMC we obtained by surface tension are 0.069 mM for Brij S20, 0.0408 mM for Brij 58, 0.126 mM for Brij C10, 0.61 mM for Brij S10.

3.4. Experiments with Tween 40 surfactant

The procedure for emulsion preparation has a significant effect on the drop shape transformations, see Fig. 3. For emulsions prepared by the first experimental procedure, at surfactant concentration of about 0.01 mM, which is about half the value of the measured CMC (0.022–0.033 mM), the C_{16} droplets all freeze into spherical shapes at 12 °C, Fig. 3A. If we increase the surfactant concentration up to 0.03–0.04 mM, the drops start to transform their shape at around 18 °C, but still only reach the initial stages from the evolutionary scheme, including formation of regular polyhedra and other faceted 3D shapes.^{4,5,9,10} In the concentration range from 0.03 to 0.1 mM, the transformation and obtained shapes remain the same, see Fig. 3C. There is no significant effect from the cooling rate – even at rate as low as ~ 0.05 K min⁻¹, no further shapes appear, Fig. 3E. At higher surfactant concentrations, ≥ 0.2 mM, we observe formation of flat shapes, which can elongate further into rods if we apply a lower cooling rate or perform the experiment with smaller emulsion droplets, $d_{ini} \leq 15$ μ m. For this case, it is known that the C_{16} drops stabilized by $C_{16}\text{SorbEO}_{20}$ surfactant typically transform their shape to hexagonal or other platelets and relatively stiff rods with diameter $\approx 5 \pm 1$ μ m. Note the variance between the shapes of the droplets on the photographs Fig. 3D and E: it is due to the effect of the size of the droplets on the rate of shape transformations, and probably some temperature inhomogeneities.

The results from the second experimental procedure are presented on the second row of Fig. 3. In this case, we observe formation of flat platelets even at 0.01 mM Tween 40, see Fig. 3B, as opposed to the emulsion droplets prepared by first procedure, see Fig. 3A. The observed differences in the behaviour of the alkane droplets indicate that the denser absorption of surfactant at higher concentration is maintained upon dilution in procedure #2, *i.e.* no complete equilibration was reached. This could be either due to the presence of a barrier to desorption or (more likely) the difference in kinetics of exchange.

We also analysed the effect of the cooling rate, see Fig. 3D and E. It can be seen that depending on the rate of cooling, the droplets are able to reach different stages of the evolutionary

scheme, *i.e.* slower cooling rates ensures more time for the drops to transform their shape and to elongate into rods and ellipsoidal droplets connected by thin fibres. It's shown that the cooling rate does not affect the order of the shape change sequence.

3.5. Experiments with Tween 60 surfactant

Additional experiments with Tween 60 ($C_{16}\text{SorbEO}_{20} + C_{18}\text{SorbEO}_{20}$) allowed us to analyse the effect of the surfactant chain-length. CMC from our interfacial tension isotherm measurements was 0.011 mM, close to literature values for the CMC = 0.017–0.026 mM.

By the first experimental procedure, in which we do not perform an additional dilution of the surfactant solution, the minimum surfactant concentration (~ 0.01 mM) necessary to observe drop shape transformations is somewhat below the measured CMC value. At this surfactant concentration, the droplets have enough surfactant molecules on their surfaces to template the plastic crystal and start the self-shaping with formation of corrugated polyhedral, which upon further cooling freeze in this shape. However, the system cannot proceed to the next stages of the self-shaping evolutionary scheme, even at much slower cooling rate of 0.05 K min⁻¹, which is known to induce changes more easily.

Further increasing the concentration of the surfactant leads to an increased probability for formation of different shapes. In the concentration range between 0.01 and 0.05 mM, the C_{16} drops are able to transform their shapes only to regular and corrugated polyhedra and irregular 3D shapes, see Fig. 4A. The later stages of the evolutionary scheme, such as flat platelet shapes (hexagons, triangles, rhombus), rods and thin fibres, can be reached by increasing the concentration of Tween 60 above 0.1 mM at around 18 °C, see Fig. 4B and C. Similarly to the experimental results obtained with Tween 40, the experiments with emulsions prepared by the 2nd experimental procedure yielded lower minimum concentrations necessary to produce self-shaping compared to the 1st procedure. The procedure also affected significantly the final-stage shapes which could be obtained at a given surfactant concentration. Whereas the C_{16} drops stabilized by 1.1×10^{-3} mM Tween 60 (1.5×10^{-4} wt%) and prepared by the first experimental procedure do not transform and freeze into spherical shape at 12 °C, the drops prepared by the second experimental procedure (with the same final surfactant concentration) transform their shape all the way to flat platelets of triangles and parallelograms. The minimum concentration to induce shapes was much lower with the second procedure, $\sim 2 \times 10^{-4}$ mM (3×10^{-5} wt%), see Fig. 4D–F. In this case the drops could reach only the initial stages of the evolutionary scheme (formation of polyhedra or hexagons) upon cooling and eventually freeze in this shape, as shown in Fig. 4E.

To analyse whether the final shapes were the result of a fast cooling rate or of surfactant concentration effects, we decreased the cooling rate 4-fold – from 0.2 K min⁻¹ to 0.05 K min⁻¹, which normally provides sufficient time for all the droplet deformation. Still, similar results were obtained: the hexagon was the furthest shape in the evolution tree formed before droplet freezing, see Fig. 4F. Below surfactant concentration of



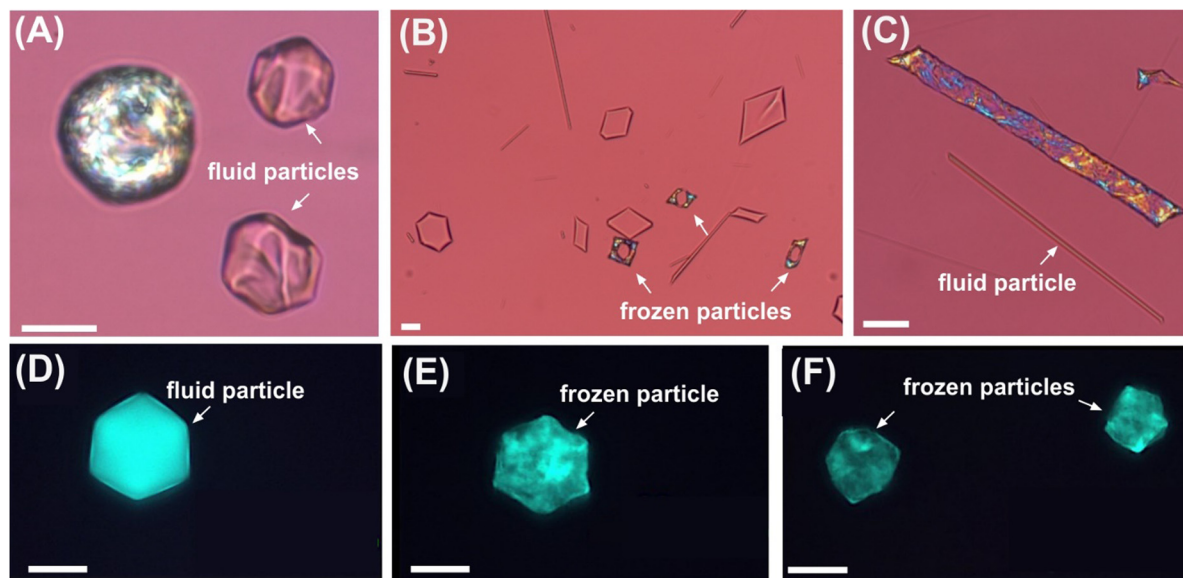


Fig. 4 Drop shape transformations for emulsions of hexadecane drops in water stabilized by Tween 60. Images on the first row are obtained with emulsions prepared by adding the oil phase into the aqueous surfactant solution with fixed surfactant concentration (procedure 1). The images on the second row are obtained with emulsion droplets prepared by membrane emulsification method in 1.5 wt% surfactant solution and then diluted with water to the desired surfactant concentration (procedure 2). Droplets $d_{ini} \approx 15 \mu\text{m}$. Microscope images of (A) corrugated polyhedra particles; (B) flat shapes; (C) rods; (D) fluid deformed particle; (E) frozen deformed hexagon particle; (F) frozen deformed particles. The total surfactant concentration in the emulsion is (A–C) 0.05 mM, (B and C) 1 mM, and (D–F) 2.0×10^{-4} mM. The cooling rate is (A–C) 0.42 K min^{-1} , (D and E) 0.2 K min^{-1} , and (F) 0.05 K min^{-1} . Images in (A–C) are made in transmitted polarized light, while these in (D–F) are in reflected light. Scale bars: 20 μm .

1.1×10^{-4} mM, *i.e.* 1.5×10^{-5} wt%, the process of self-shaping stops completely and all the alkane droplets freeze into a spherical shape. In this case, packing of the molecules is not tight enough, and the plastic phase cannot form, or cannot counteract the corresponding higher surface tension of the droplets at these reduced concentrations, so no self-shaping phenomenon can be observed.

The analysis yields a narrow range for the critical concentrations necessary to induce shapes. For emulsions prepared by

the experimental procedure 1, the critical concentration is just below the CMC (0.01 mM *vs.* CMC ~ 0.011 mM). When diluting from an emulsion with concentrated surfactant (second procedure), the critical concentration in solution is $50\times$ lower.

3.6. Experiments with Brij surfactants

Emulsions stabilized by Brij surfactants were prepared by double syringe technique (drops with $d_{ini} \approx 5\text{--}25 \mu\text{m}$). The

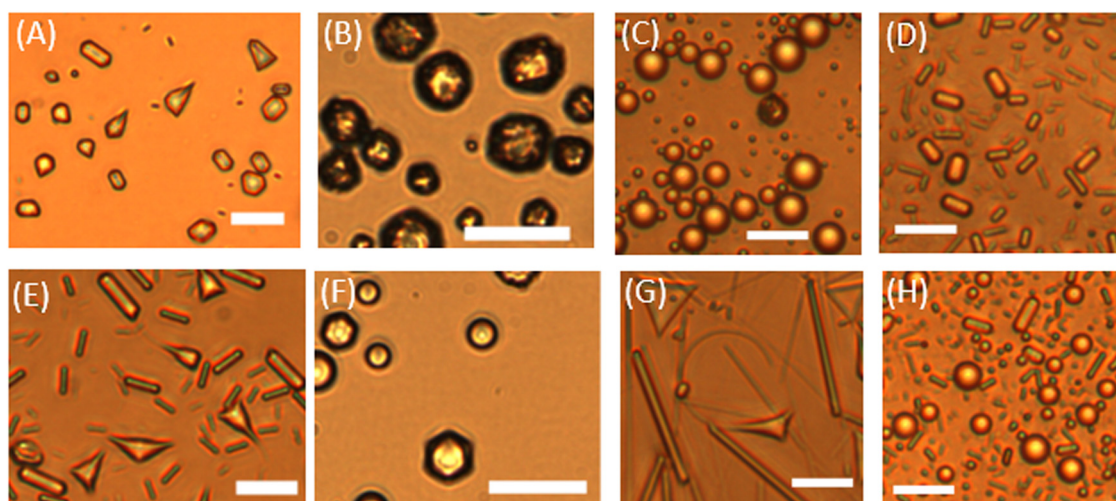


Fig. 5 Drop shape transformations for emulsions of hexadecane drops in water stabilized by (A and E) Brij S20, (B and F) Brij 58, (C and G) Brij C10, (D and H) Brij S10. Images on the first row are obtained from samples with surfactant concentration just below the CMC. The images on the second row are obtained from samples with surfactant concentration just above the CMC. Initial temperature = 20 °C, final temperature = 11 °C and cooling rate is 0.5–1 K min $^{-1}$. Scale bar 20 μm .



shape change of droplets in Brij surfactant emulsions were analyzed at four different concentrations, two of which were closely above and below the CMC, whilst the other two were far away from the CMC (measurements for different Brij surfactants in Fig. S2, ESI†).

Our observations confirm that the concentration at which shape changes first appear is close to the CMC, usually around $0.7 \times \text{CMC}$ the droplets changed shaped at around at $10\text{--}12\text{ }^\circ\text{C}$ (Fig. 5 and Table S3, ESI†). Fig. S3–S7 in the ESI† show the droplets freezing/shape changes at different surfactant concentrations. These include the formation of rechargeable droplet swimmers³⁴ shown in Fig. S3 and S7 (ESI†), in the presence of fluorescent dyes. Such minimal systems are of interest both in origin of life studies³⁵ and in growing bottom-up robotics.^{36,37}

3.7. Competitive adsorption effects on self-shaping in surfactant mixtures

Most commercial surfactants contain significant amounts of impurities, and many industrial formulations include more than one surfactant molecule on purpose. It is therefore important to investigate the effect of the composition of a mixture on the self-shaping behaviour of droplets. We chose the non-ionic mixture of the long-tailed Tween 60 ($\text{C}_{16}\text{SorbEO}_{20} + \text{C}_{18}\text{SorbEO}_{20}$), which has the ability to self-shape C_{16} drops, and the short-tailed

Tween 20 (roughly 40% $\text{C}_{12}\text{SorbEO}_{20}$, 40% $\text{C}_{14}\text{SorbEO}_{20}$, 20% $\text{C}_{16}\text{SorbEO}_{20}$), which does not.^{4,5} Keeping the total concentration of surfactant constant (1.5 wt%), we explored the effect of the balance in the mixture on the ability of droplets to self-shape.

First, a 1:1 wt/wt surfactant mixture was prepared at a total concentration of 1.5 wt%. Hexadecane droplets of 15 to 20 μm were cooled at a constant rate of 0.2 K min^{-1} , as in the previous experiments. Fig. 6A depicts the three different states of surfactant molecules as found in our system – in micelles, a small fraction existing free in solution, and adsorbing and packing at the surface of oil drops. Due to its shorter chain length, Tween 20 cannot effectively template the formation of the alkane rotator phases and disrupts the templating from Tween 60. However, Tween 60 has higher surface activity, so the fraction of Tween 60 at the interface is much larger than the 1:1 ratio in the aqueous solution.

To determine the concentration of $\text{C}_{12}\text{SorbEO}_{20}$ on the drop surface we assume that the components of Tween 20 and Tween 60 mix ideally in the micelles. Under this assumption, accounting for the composition of the two surfactants, and based on measured CMC and adsorption coefficient values, we find that the fraction of Tween 20 in the bulk composition is close to the fraction in the interfacial composition. This means that the large fraction of long chain surfactants in Tween 20

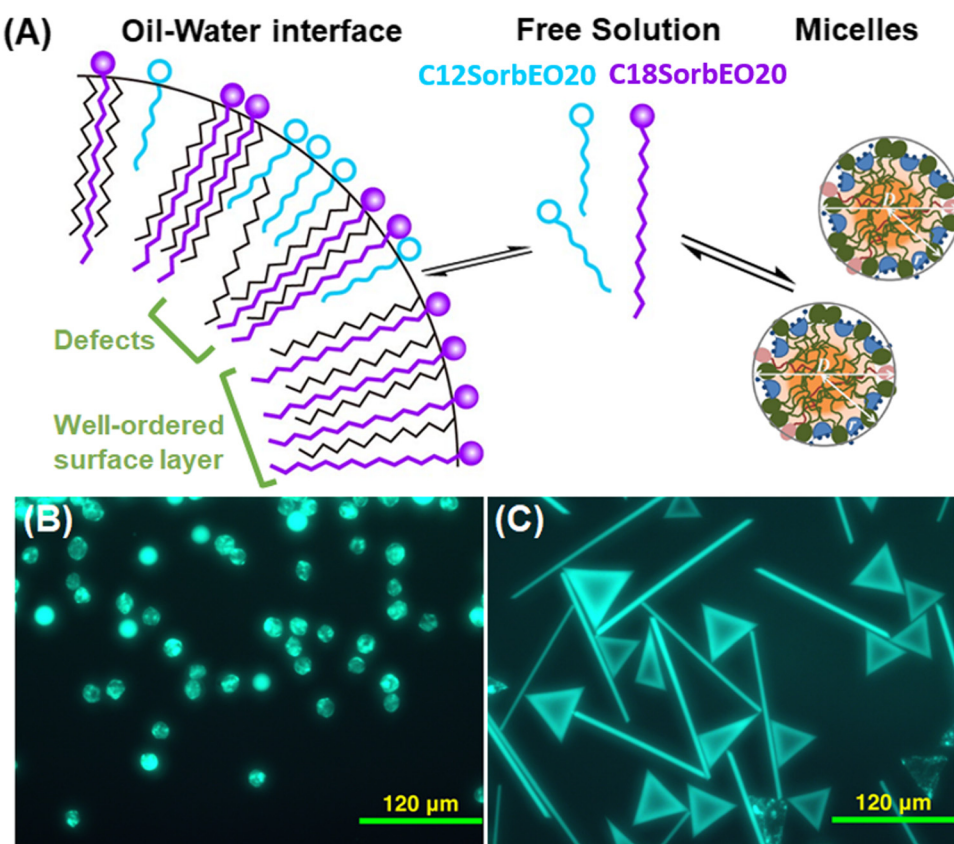


Fig. 6 (A) Schematic illustration of a mixed surfactant layer containing $\text{C}_{18}\text{SorbEO}_{20}$ and $\text{C}_{12}\text{SorbEO}_{20}$ at the oil/water interface. $\text{C}_{18}\text{SorbEO}_{20}$ may favorably interdigitate with C_{16} alkane molecules to form a well-ordered layer and template the formation of further alkane rotator phase layers beneath it. However, $\text{C}_{12}\text{SorbEO}_{20}$ cannot – due to short chain length, this surfactant produces defects in the surface structure. (B) 34% $\text{C}_{12}\text{SorbEO}_{20}$ occupation. (C) 26% $\text{C}_{12}\text{SorbEO}_{20}$ occupation.



Table 1 Influence of mixtures of surfactants with different tail lengths (at constant total 1.5 wt% concentration) on the appearance of shapes upon cooling of hexadecane emulsion droplets

Wt ratio in aqueous solution [Tween 60]/[Tween 20]	Tween 60 (%)	Concentration of $C_{12}\text{SorbEO}_{20}$ in mM	% surface occupation by $C_{12}\text{SorbEO}_{20}$	Appearance of self-shaping
1:73	1.35	4.8	40	No
1:6.6	13.2	4.2	37	No
1:3.28	23.4	3.7	34	No
1:1	50	2.4	26	Yes
2:1	66.7	1.6	19	Yes

contributes to the effective adsorption on the liquid interface and the apparently low CMC of the Tween 20 – see ESI.† We also found that the self-shaping process starts when the bulk fraction of $C_{12}\text{SorbEO}_{20}$ is lower than 20%, see Table 1.

4. Conclusions

The experiments above show the minimum concentration of surfactant necessary to induce shape formation is around the surfactants' CMC. We also verify it is not the absolute concentration in solution that is important, but rather the coverage on the water/oil interface. We do this by performing experiments with ionic surfactants in the presence of salts where concentrations that previously could not induce shaping suddenly do, because the presence of salt drastically lowers the CMC, and again we find the critical concentration for inducing shapes is around CMC.

We also perform experiments with a mixed set of non-ionic surfactants with the same heads but one with a tail similar to the alkane oil (Tween 60, $C_{16}\text{SorbEO}_{20}$ + $C_{18}\text{SorbEO}_{20}$) that induces shapes and one with a much shorter one does not (Tween 20). In experiments we vary the bulk surfactant fraction of $C_{12}\text{SorbEO}_{20}$ from 40% to 19% and found that at 26% surface coverage and below of $C_{12}\text{SorbEO}_{20}$ we observe self-shaping.

In the context of surface coverage of a surfactant in an experiment with a simple visual readout (as in Fig. 6B and C) this approach would allow us to receive accurate information about molecular orientation and packing on the surface (and the formation of rotator phase beneath the surface layer), which was previously only possible to find out by much more expensive, rare, and time-consuming techniques. One could then easily and effectively switch on and off the shape transformations of oil droplets by controlling solution concentrations. Such precise guidance of interfacial layer composition would enable further fundamental experiments into the molecular basis for inducing such phase transitions.

The discovery of minimum requirements needed for shaping droplets has potential for scalable manufacturing of anisotropic structures from monomer droplets,⁷ and development of multi-functionality by sequential interpenetration,³⁸ e.g., supercapacitance,³⁹ programmable actuation,³⁸ and self-sensing.⁴⁰ Bottom-up, sustainable growth of active particles could be enabled by the minimal surfactant and mixed surfactant criteria established here.

Author contributions

J. F. and S. K. S. conceived the experiments for minimal surfactants concentration to induce the shapes. In discussions with ST we designed a wider set of experiments including many that were supervised by S. T. and performed by Zh. V. E. L. and E. N. worked on the Brij surfactant's part and analysed the results. T. W. helped J. F. in the emulsification and microscope setups. R. S. and S. T. theoretically analysed the description of mixed monolayer experiments. J. F. and S. K. S. co-wrote the manuscript. J. F., S. S. and Zh. V. created the figures. S. T. and R. S. critically edited the manuscript. All authors contributed to the discussion of results and commented on the manuscript.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

- 1 L. Hong, S. Jiang and S. Granick, Simple method to produce janus colloidal particles in large quantity, *Langmuir*, 2006, **22**, 9495–9499.
- 2 L. Cademartiri and G. A. Ozin, Ultrathin nanowires – A materials chemistry perspective, *Adv. Mater.*, 2009, **21**, 1013–1020.
- 3 S. K. Smoukov, *et al.*, Scalable liquid shear-driven fabrication of polymer nanofibers, *Adv. Mater.*, 2015, **27**, 2642–2647.
- 4 N. Denkov, S. Tcholakova, I. Lesov, D. Cholakova and S. K. Smoukov, Self-shaping of oil droplets via the formation of intermediate rotator phases upon cooling, *Nature*, 2015, **528**, 392–395.



5 D. Cholakova, N. Denkov, S. Tcholakova, I. Lesov and S. K. Smoukov, Control of drop shape transformations in cooled emulsions, *Adv. Colloid Interface Sci.*, 2016, **235**, 90–107.

6 S. Guttman, *et al.*, How faceted liquid droplets grow tails, *Proc. Natl. Acad. Sci. U. S. A.*, 2015, **113**, 10–13.

7 I. Lesov, *et al.*, Bottom-Up Synthesis of Polymeric Micro- and Nanoparticles with Regular Anisotropic Shapes, *Macromolecules*, 2018, **51**, 7456–7462.

8 O. Marin, *et al.*, Journal of Colloid and Interface Science Self-faceting of emulsion droplets as a route to solid icosahedra and other polyhedra, *J. Colloid Interface Sci.*, 2019, **538**, 541–545.

9 P. A. Haas, R. E. Goldstein, S. K. Smoukov, D. Cholakova and N. Denkov, Theory of Shape-Shifting Droplets, *Phys. Rev. Lett.*, 2017, **118**, 1–5.

10 P. A. Haas, D. Cholakova, N. Denkov, R. E. Goldstein and S. K. Smoukov, Shape-shifting polyhedral droplets, *Phys. Rev. Res.*, 2019, **1**, 23017.

11 I. García-Aguilar, P. Fonda, E. Sloutskin and L. Giomi, Faceting and Flattening of Emulsion Droplets: A Mechanical Model, *Phys. Rev. Lett.*, 2021, **126**, 38001.

12 P. A. Haas, R. E. Goldstein, D. Cholakova, N. Denkov and S. K. Smoukov, Comment on ‘faceting and Flattening of Emulsion Droplets: A Mechanical Model’, *Phys. Rev. Lett.*, 2021, **126**, 259801.

13 D. Cholakova, Z. Valkova, S. Tcholakova, N. Denkov and S. K. Smoukov, “Self-Shaping” of Multicomponent Drops, *Langmuir*, 2017, **33**, 5696–5706.

14 S. R. Liber, *et al.*, Polyhedral Water Droplets: Shape Transitions and Mechanism, *J. Am. Chem. Soc.*, 2020, **142**, 8672–8678.

15 M. M. Knock, G. R. Bell, E. K. Hill, H. J. Turner and C. D. Bain, Sum-frequency spectroscopy of surfactant monolayers at the oil–water interface, *J. Phys. Chem. B*, 2003, **107**, 10801–10814.

16 S. A. Burrows, I. Korotkin, S. K. Smoukov, E. Boek and S. Karabasov, Benchmarking of Molecular Dynamics Force Fields for Solid–Liquid and Solid–Solid Phase Transitions in Alkanes, *J. Phys. Chem. B*, 2021, **125**, 5145–5159.

17 D. Cholakova, N. Denkov, S. Tcholakova, Z. Valkova and S. K. Smoukov, Multilayer Formation in Self-Shaping Emulsion Droplets, *Langmuir*, 2019, **35**, 5484–5495.

18 D. Cholakova and N. Denkov, Rotator phases in alkane systems: In bulk, surface layers and micro/nano-con filaments, *Adv. Colloid Interface Sci.*, 2019, **269**, 7–42.

19 G. T. Vladisavljević and R. A. Williams, Recent developments in manufacturing emulsions and particulate products using membranes, *Adv. Colloid Interface Sci.*, 2005, **113**, 1–20.

20 N. Christov, *et al.*, Capillary mechanisms in membrane emulsification: oil-in-water emulsions stabilized by Tween 20 and milk proteins, *Colloids Surf., A*, 2002, **209**, 83–104.

21 C. Charcosset, I. Limayem and H. Fessi, The membrane emulsification process—a review, *J. Chem. Technol. Biotechnol.*, 2004, **79**, 209–218.

22 T. Nakashima, M. Shimizu and M. Kukizaki, Membrane Emulsification by Microporous Glass, *Key Eng. Mater.*, 1992, **61–62**, 513–516.

23 R. H. Newton, J. P. Haffee and M. W. Ho, Polarized light microscopy of weakly birefringent biological specimens, *J. Microsc.*, 1995, **180**, 127–130.

24 S. T. Hyde Identification of lyotropic liquid crystalline mesophases, in *Handbook of applied surface and colloid chemistry*, 2001, vol. 2, pp. 299–332.

25 D. D. Miller, L. J. Magid and D. F. Evans, Fluorescence quenching in double-chained surfactants. 2. Experimental results, *J. Phys. Chem.*, 1990, **94**, 5921–5930.

26 G. D. Noudeh, M. Housaindokht and B. S. F. Bazzaz, The Effect of Temperature on Thermodynamic Parameters of Micellization of Some Surfactants, *J. Appl. Sci.*, 2007, **7**, 47–52.

27 C. L. A. Mesa and M. Terenzi, Studies on Krafft Point Solubility Solutions, *Thermochim. Acta*, 1988, **137**, 143–150.

28 H. Nakahara, O. Shibata and Y. Moroi, Examination of Surface Adsorption of Cetyltrimethylammonium Bromide and Sodium Dodecyl Sulfate, *J. Phys. Chem. B*, 2011, **115**, 9077–9086.

29 B. A. Kerwin, Polysorbates 20 and 80 used in the formulation of protein biotherapeutics: Structure and degradation pathways, *J. Pharm. Sci.*, 2008, **97**, 2924–2935.

30 B. M. Ocko, *et al.*, Surface freezing in chain molecules: Normal alkanes, *Phys. Rev. E*, 1997, **55**, 3164–3182.

31 N. Denkov, D. Cholakova, S. Tcholakova and S. K. Smoukov, On the Mechanism of Drop Self-Shaping in Cooled Emulsions, *Langmuir*, 2016, **32**, 7985–7991.

32 S. Tcholakova, *et al.*, Efficient self-emulsification via cooling-heating cycles, *Nat. Commun.*, 2017, **8**, 1–11.

33 M. E. Mahmood and D. A. F. Al-Koofee, Effect of Temperature Changes on Critical Micelle Concentration for Tween Series Surfactant, *Glob. J. Sci. Front. Res. Chem.*, 2013, **13**, 1–7.

34 D. Cholakova, *et al.*, Rechargeable self-assembled droplet microswimmers driven by surface phase transitions, *Nat. Phys.*, 2021, **17**, 1050–1055.

35 I. Ivanov, S. K. Smoukov, E. Nourafkan, K. Landfester and P. Schwille Origin of life from a maker’s perspective – focus on protocellular compartments in bottom-up synthetic biology, 2022, ArXiv, DOI: [10.48550/arxiv.2207.07225](https://arxiv.org/abs/2207.07225).

36 S. K. Sustainably Smoukov Grown: The Underdog Robots of the Future in Mazzolai *et al.* Roadmap on Soft Robotics, *Multifunct. Mater.*, 2022, arxiv:2206.10306.

37 B. Mazzolai, *et al.*, Roadmap on Soft Robotics: multifunctionality, adaptability and growth without borders, *Multifunct. Mater.*, 2022, DOI: [10.1088/2399-7532/ac4c95](https://doi.org/10.1088/2399-7532/ac4c95).

38 A. Khaldi, C. Plesse, F. Vidal and S. K. Smoukov, Smarter Actuator Design with Complementary and Synergetic Functions, *Adv. Mater.*, 2015, **27**, 4418–4422.

39 K. D. Fong, T. Wang, H. K. Kim, R. V. Kumar and S. K. Smoukov, Semi-Interpenetrating Polymer Networks for Enhanced Supercapacitor Electrodes, *ACS Energy Lett.*, 2017, **2**, 2014–2020.

40 T. Wang, *et al.*, Electroactive polymers for sensing, *Interface Focus*, 2016, **6**, 20160026.

