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Bile Salts Caught in the Act: From Emulsification to Nanostructural Reorganization of Lipid Self-Assemblies

Amin Sadeghpour^{1,2}, Michael Rappolt¹, Shravasti Misra^{3,4,5} and Chandrashekar V. Kulkarni^{3*}

¹*School of Food Science and Nutrition, University of Leeds, Leeds, LS2 9JT, United Kingdom*

²*Empa, Swiss Federal Laboratories for Materials Science and Technology, Center for X-ray Analytics, Dübendorf, 8600, Switzerland*

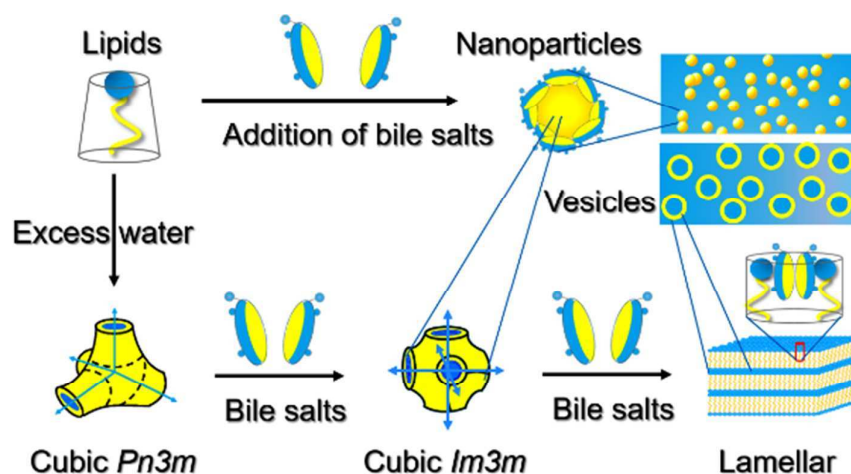
³*School of Physical Sciences and Computing, University of Central Lancashire, Preston, PR1 2HE, United Kingdom*

⁴*Department of Biosciences and Bioengineering, Indian Institute of Technology Bombay, Mumbai, 40076, India*

⁵*Department of Biology and Biochemistry, University of Houston, Science Center, Houston, TX 77204, United States of America (current address)*

*Corresponding authors: C.V. Kulkarni: E-mail cvkulkarni@uclan.ac.uk, Tel: +44-1772-89-4339, Fax: +44-1772-89-4981

TOC Graphics



Abstract

Bile salts (BS) are important for digestion and absorption of fats and fat-soluble vitamins in the small intestine. In this work, we scrutinized, with small angle X-ray scattering (SAXS), the crucial functions of bile salts beyond their capacity for interfacial stabilization of submicron sized lipid particles. By studying a wide compositional range of BS-lipid dispersions using two widely applied lipids for drug-delivery systems (one a monoglyceride being stabilizer-sensitive and the other an aliphatic alcohol being relatively stabilizer-insensitive), we identified the necessary BS to lipid ratios for guaranteeing full emulsification. A novel *ad hoc* developed global small angle-X-ray scattering analysis method revealed that the addition of BS hardly changes the bilayer thicknesses in bicontinuous phases, while a significant membrane thinning is observed in the coexisting fluid lamellar phase. Furthermore, we show that BS strongly decreases the average critical packing parameter. At increasing BS concentration, the order of phases formed are (i) the bicontinuous diamond cubic ($Pn3m$), (ii) the bicontinuous primitive cubic ($Im3m$) followed by (iii) the fluid lamellar phase (L_α). These distinctive findings on BS driven 'emulsification' and 'membrane curvature reduction' provide new molecular scale insights for the understanding of the interfacial action of bile salts on lipid-assemblies.

Introduction

Food fats contain an assortment of different lipid types that self-assemble under *excess of water* conditions into various complex structures including (i) lyotropic liquid crystalline (LLC) phases based on bilayers (these include the fluid lamellar L_α and gel lamellar L_β phase as well as the bicontinuous cubic phases with the space groups $Pn3m$ and $Im3m$); (ii) discontinuous LLC phases based on *spherical micelles* (cubic $Fd3m$ phase) and *rod-like micelles* (H_2 phase); and (iii) reverse micelles on their own i.e. L_2 phase¹⁻³ (Figure 1).

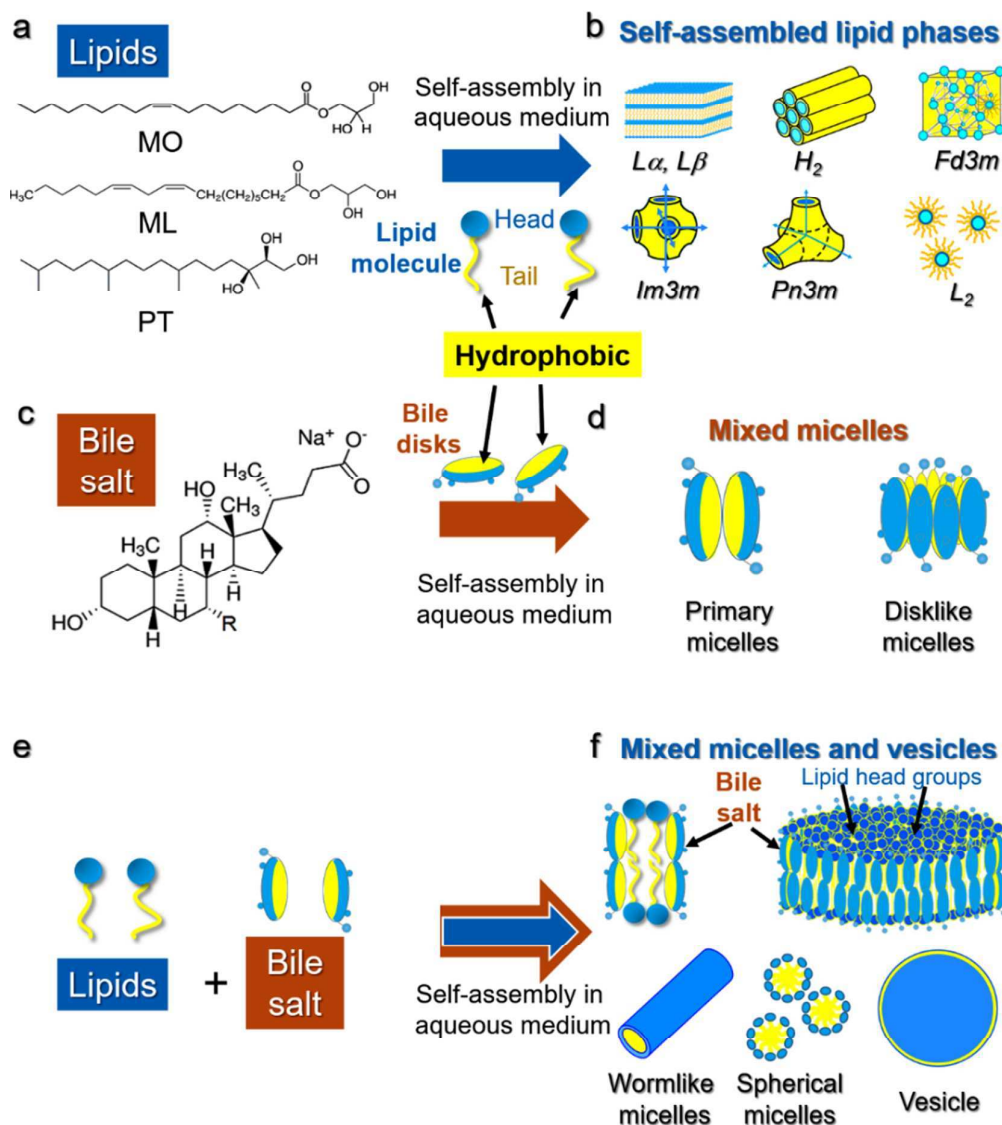


Figure 1. Chemical structures of a) lipids: monoolein (MO) and monolinolein (ML) (main constituents of the commercial lipid source Dimodan U/J (DU) and phytantriol (PT), and c) the common bile salt (BS) sodium deoxycholate. Schematic drawings depict the head-tail structures of lipids and disk-like shapes of bile salts. b) Edible lipids self-assemble into a remarkable range of nanostructures in an aqueous medium³⁻⁴, for example, cubic $Pn3m$ and $Im3m$, lamellar L_α and L_β as well as inverse hexagonal H_2 , inverse micellar L_2 and inverse micellar cubic $Fd3m$ phases. d) Bile

1 salts, in aqueous solution induce the formation of mixed micelles⁵⁻⁶; whereas BS and lipids
2 together e) tend to form mixed structures including spherical, disk-like and worm-like micelles as
3 well as stabilized unilamellar vesicles f).⁷⁻¹²

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6 The lamellar phases exist as fluid- L_{α} , gel- L_{β} or crystalline- L_c polymorphs being distinguished by
7 their type of lipid packing as well as by their degree of bilayer fluidity². On the other hand,
8 common non-lamellar phases comprise the inverse hexagonal (H_2) phase and inverse bicontinuous
9 cubic phases¹³ with space groups $Im3m$, $Pn3m$ and $Ia3d$. These cubic phases usually exhibit a very
10 high viscosity (in the range of 10^4 - 10^5 Pa·s)¹⁴ rendering them problematic for various applications,
11 but also contributing to their lower digestibility as compared to other LLC phases.¹⁵⁻¹⁹ Further,
12 disk-shaped aggregates also termed as bicelles (a portmanteau word created from 'bilayer' and
13 'micelles') can be produced. They are composed of long-chain lipids that make up their planar
14 region and either detergent or short-chain lipids are chosen to form their rim; other LLC phases
15 concern the micellar cubic $Fd3m$ and sponge (L_3) phases²⁰⁻²². Similar to the inverse bicontinuous
16 cubic phases, also the latter two structures display highly viscous agglutinated physical forms,
17 potentially hampering their accessibility by digestive molecules. Digestion media in the
18 gastrointestinal (GI) tract help to fragment and emulsify these fatty globules into smaller particles,
19 thereby increasing the interfacial area, which in turn assists the digestion process^{7, 12, 23-25}. Apart
20 from gastric and pancreatic lipase being active in the stomach and small intestine, respectively,²⁶
21 bile salts (BSs) are secreted into the lumen of the **small intestine** and play an important role in fat
22 digestion and absorption²⁷.

23
24
25 Bile salts are biological surfactants, synthesized in the liver, which enhance the solubility of non-
26 polar molecules including lipids⁵ (Figure 1a). Main representatives of bile salts are sodium cholate
27 (NaC) and sodium deoxycholate (NaDC) molecules (Figure 1c) with **air-water** interfacial tensions of
28 $52 \text{ mN}\cdot\text{m}^{-1}$ and $44 \text{ mN}\cdot\text{m}^{-1}$ at concentrations of 0.01 mol/Kg , respectively²⁸. The low interfacial
29 tensions **with respect to the one from water** cause BS to aggregate into various micellar structures
30 including primary or disk-like micelles (Figure 1d) above their critical micellar concentration
31 (CMC).⁵⁻⁶ Bile salts' molecular structure deviates strongly from the classical head-tail structure
32 (Figure 1a) of common surfactant molecules, as they exhibit planar shapes with the hydrophobic
33 and hydrophilic faces on either sides⁵ (Figure 1c). Together with lipids, bile salts tend to form
34 mixed micelles and vesicles in dilute solutions⁷⁻¹² (Figure 1f). Structural studies suggest that the
35 formation of spherical, worm-like and disk-like micelles (Figure 1f) are preferred over stabilizing
36 vesicles, where bile salts are located in the lipid head-group region, while the hydrophobic tails are
37 shielded from aqueous medium^{9, 11-12, 29}. These particular BS-lipid interactions primarily contribute

1 to the emulsification of lipidic food lumps^{7, 12, 23-25}, as studied in this work with the emulsification
2 capacity of BS on bicontinuous cubic phases.
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6 Several research groups have investigated the role of bile salts in digesting lipids by simulating
7 and/or modelling the different digestive juice conditions, both, in the presence and absence of bile
8 salts^{7, 11, 24, 30-31}. Numerous other studies concentrated on *in-vivo* investigations reporting on
9 possible causes for the reduction in fat absorption,³² highlighting the role of mean droplet sizes
10 and surface charge densities at different stages of digestion,³³ and several groups have been
11 studying structural variations in micro-emulsions under simulated gastrointestinal conditions.^{34, 35}
12 Studies also demonstrated an intensification of the lipolysis process in emulsion system, e.g. milk
13 fats, upon interactions with bile salts.³⁶ On the one hand, such interactions play a critical role in
14 controlling the fat absorption, but on the other hand, are important to optimize the absorption of
15 lipid soluble bioactives. Finally, the influence of fat stabilizing proteins and/or polysaccharides on
16 the fat absorption has been investigated in the presence of bile salts in simulated digestive
17 systems.³⁷
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27 In this study, we investigate a wide range of BS-lipid compositions, but without adding any other
28 digestive molecules. Here two widely applied lipids for drug delivery, namely Dimodan-U/J (DU)
29 and phytantriol (PT), were employed to form non-lamellar liquid crystalline phases (bicontinuous
30 cubic phases),^{16, 38} whose detailed structural analysis upon interaction with BS has not been
31 reported in the literature yet. Nevertheless, a wide range of studies has published on the influence
32 of bile salts with other lipid self-assemblies. For instance, Gustafsson et al. have reported on the
33 phase behavior and formation of lamellar and cubic liquid crystalline phases in aqueous mixtures
34 of monooleate and bile salts.³⁹ Various different lipid phase diagrams and structural changes upon
35 interactions with different surfactants including bile salts have been intensively investigated over
36 the last two decades.⁴⁰⁻⁴⁶
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45 We note, based on recent nutritional reports, that our studied non-lamellar self-assembled phases
46 can occur and promote the digestion of food products such as human breast milk or mayonnaise
47 and aid the absorption of oil-soluble food compounds and various nutraceuticals^{31, 47-49}. In
48 addition, monoglycerides and aliphatic alcohols are receiving a growing interest in the formulation
49 of smart food and novel drug delivery systems,^{30, 50-52} because as by-products of hydrolysis of
50 common triglycerides, in particular monoglycerides are cheap and at the same time accepted
51 food-grade materials.
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Owing to an *ad hoc* developed new global small angle-X-ray scattering analysis procedure (based on previous work in this field of LLC nanostructural analysis⁵³⁻⁵⁷), we were able to obtain nanostructural details from not only the bicontinuous cubic and fluid lamellar phases, but moreover, able to extract structural information in the phase coexistence regime. In particular, we are able to provide new insights on the architecture of these LLC phases in the presence of BSs and provide detailed information BS-lipid interactions on the molecular scale.⁵⁸⁻⁶⁰ Rheological measurements support our data on the macroscopic scale. The results demonstrate that the bile salts do not only contribute in the emulsification of lipids, but they also possess 'membrane curvature power' to convert, for instance, the complex bicontinuous cubic phases into vesicles. While different digestion processes of lipids have been widely investigated^{7, 24, 30-31, 48}, a systematic study on 'emulsifying role' of pure bile salts and the concomitant significance of 'interfacial curvature reduction' of monoglyceride and aliphatic alcohols assemblies is presented for the first time.

Materials and Methods

Materials: Dimodan U/J (DU) containing 96% distilled monoglycerides, mainly monoolein and monolinolein, and smaller amounts of diglycerides and free fatty acids, was kindly supplied by Danisco (Brabrand, Denmark). The other monoglyceride source, namely phytantriol (PT) was a gift from DSM Nutritional Products Europe (provided by the local distributor Adina Pharma, UK). The bile salt mixture containing sodium cholate (NaC) and sodium deoxycholate (NaDC) was purchased from Sigma-Aldrich (UK). All chemicals were used without further purification. All samples were prepared using Milli-Q (Millipore, UK) water.

Preparation of bile salt-lipid emulsions: A wide range of BS-lipid dispersions were prepared by weighing appropriate amounts of BS and lipid, respectively, i.e. all concentrations refer to weight (wt%). The ratio between bile salt and lipid is defined by the parameter β as:

$$\beta = \frac{\text{wt of bile salt}}{\text{wt of lipid}} \times 100 \quad (1)$$

For instance, to prepare 5 wt% emulsion, 500 mg of molten lipid (DU or PT) was transferred into an empty glass vial and diluted with a 9.5 g aqueous solution of bile salts. An ultra-sonication probe (Sonics & Materials Vibra-Cell VCX750, Jencons, UK) with a 30% amplitude for the duration of 5 minutes in pulse mode (1 s pulse and 1 s delay) was employed to prepare BS-lipid dispersions.

1 The stability of dispersions was monitored visually by assessing the homogeneity against phase
2 separation. Dispersions with visible phase separation were registered as 'unstable', while
3 homogeneous mixtures were reported as 'stable' emulsions (Figure 2). BS-lipid emulsions were
4 prepared varying the lipid concentration in the range of 5 to 20 wt% and the BS concentration in
5 the range of 0 to 1 wt%.

10
11 **Rheological measurements:** The viscosity measurements were conducted using a Bohlin
12 rheometer (Malvern Instruments Ltd., Worcestershire, England, U.K), with a cone and plate type
13 geometry. Apparent viscosity was measured at shear rates in the range of 0.2-200 Pa·s⁻¹ using
14 continuous shear, with a 30 s delay time and a 30 s integration time at 25 °C.

18
19 **Dynamic Light Scattering:** Particle size distributions (normalized by volume) were measured using
20 dynamic light scattering technique (Zetasizer Nano ZS, Malvern Instruments, UK).

23
24 **Small angle X-ray scattering experiments:** Small angle X-ray scattering (SAXS) technique was used
25 for the analysis of liquid crystalline nanostructures. The SAXSpace instrument (Anton Paar, Graz,
26 Austria), utilized for these studies is equipped with a sealed-tube Cu anode X-ray generator. It was
27 operated at 40 kV and 50 mA and chilled by a closed water circuit. The line-focus camera (Anton
28 Paar, Graz, Austria) uses Cu-K_α radiation with a wavelength $\lambda=0.154$ nm. For current experiments,
29 the minimum accessible scattering vector value, q_{min} , was 0.05 nm⁻¹ ($q = (4\pi/\lambda)\sin\theta$, where 2θ is
30 the scattering angle). Silver behenate with a known lamellar spacing of 5.84 nm⁶¹ was used to
31 calibrate the scattering vector modulus q . In order to identify the precise position of the primary
32 beam and the transmission correction of the scattering profiles, a semitransparent beam stop is
33 used.

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41 Reusable vacuum tight quartz capillary (Anton Paar, Graz, Austria) with an outer diameter of 1 mm
42 was used to study fluid samples. For gel-like samples, a vacuum-tight paste cell (Anton Paar, Graz,
43 Austria) sealed with thin Kapton foils was used. The temperature was controlled by a remote
44 controlled sample stage (TCStage 150, Anton Paar, Graz, Austria) with a precision of 0.1 °C. The
45 SAXSpace is equipped with a Mythen micro-strip X-ray detector (Dectris Ltd, Baden, Switzerland).
46 Three separate recordings each with an exposure time of 600 sec were averaged to obtain the
47 final scattering profile.

53
54 The scattering patterns were corrected with respect to the position of the primary beam using the
55 SAXStreat software (Anton Paar, Graz, Austria). The relative intensity of scattering data was

further corrected using the transmittance of the direct X-ray beam ($2\theta = 0$). The background scattering from empty cells and water was subtracted. All standard corrections were applied using the SAXSQuant software (Anton Paar, Graz, Austria).

Theoretical models: The theoretical scattering curves were calculated by introducing a new model combining the scattering intensities from lamellar (I_L) and cubic (I_Q) phases with f representing the phase fraction of lamellar structures (equation 2),

$$I_t(q) = f I_L(q) + (1 - f)I_Q \quad (2)$$

The scattering intensities of lamellar structures were calculated based on the methods described in details elsewhere.⁵³⁻⁵⁴ Briefly, the method considers a double-Gaussian model as the electron density profile in real space and its Fourier transform as the form factor, ($F(q)$). In this global fitting procedure, the structure factor, $S(q)$, is calculated applying the *Modified Caillé Theory (MCT)*⁵⁵. The overall scattering from multilamellar and positionally uncorrelated membranes is obtained by:

$$I_L(q) = |F(q)|^2 S(q)/q^2 + N_u |F(q)|^2 / q^2 \quad (3)$$

where N_u is the scaling constant for the diffuse scattering contribution from single bilayer.

The reflections and relative peak amplitudes for $Im3m$ (primitive) and $Pn3m$ (diamond) cubic phases were calculated according to a model described by Garstecki *et al.*⁵⁶⁻⁵⁷ In this model the reflections are calculated by the following formula:

$$I_{hkl}^{(mod)}(L) = \mathcal{M}_{hkl} \left[\frac{F_{hkl}^S \sin(\alpha_{hkl}\pi(h^2+k^2+l^2)^{1/2} L/a)}{a^2 \alpha_{hkl}\pi(h^2+k^2+l^2)^{1/2}} \right]^2 \quad (4)$$

in which the \mathcal{M}_{hkl} is the multiplicities for each reflection, L/a is the dimensionless thickness of the bilayer separating the two continuous phases, a is the lattice parameter and α_{hkl} is a correction factor and $\frac{F_{hkl}^S}{a^2}$ are the normalized structure factors. This model also takes into account the experimental broadening of the reflections using a Gaussian distribution function.

$$I_{hkl}^{(exp/mod)}(q) = \frac{1}{\sigma_r(2\pi)^{1/2}} \sum_{hkl} \exp\left[-\frac{(q_{hkl}-q)^2}{2\sigma_r^2}\right] I_{hkl}^{(mod)} \quad (5)$$

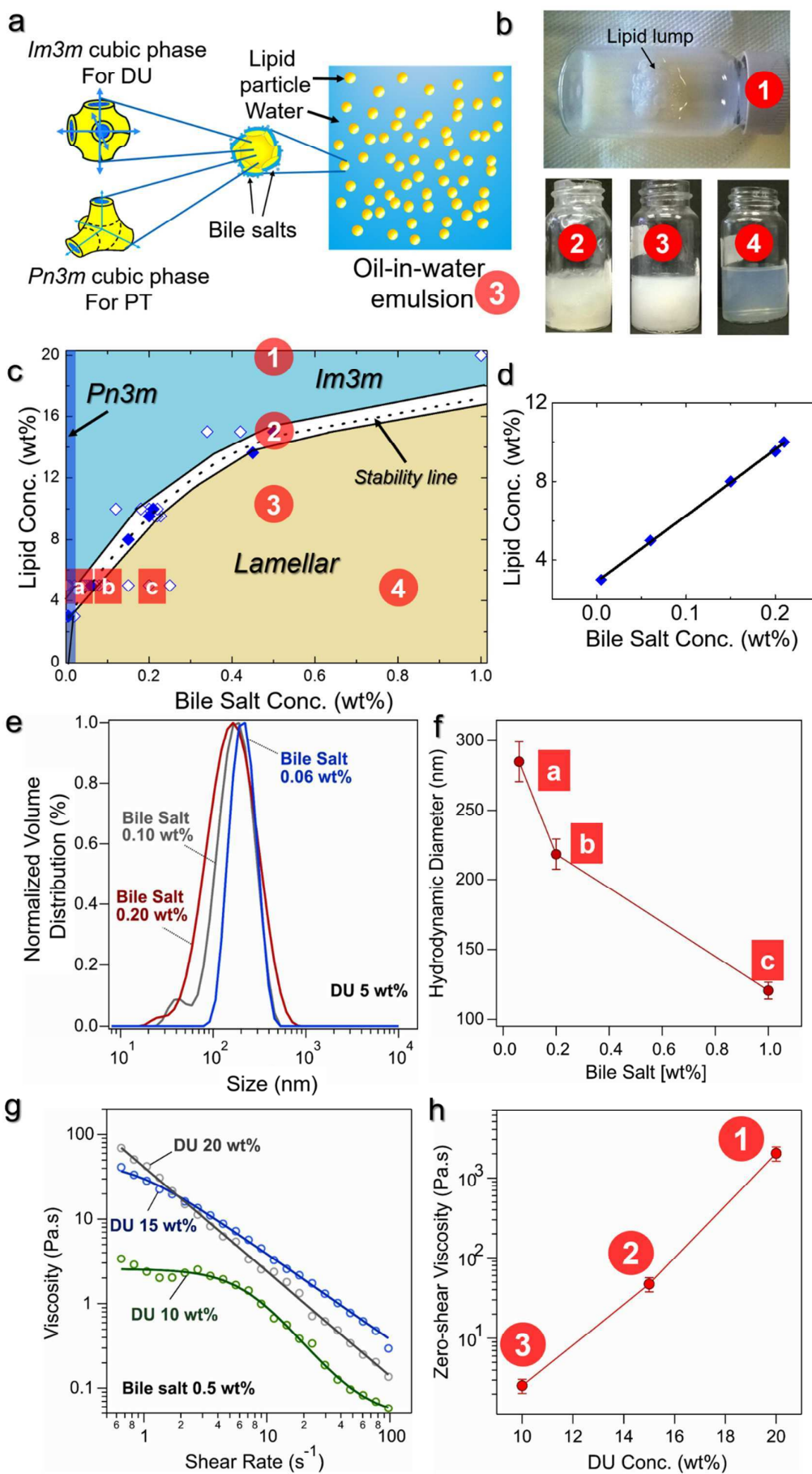
where σ_r is related to the standard deviation of the experimental Bragg peaks. We have applied the global fitting analysis using a Particle Swarm Optimization (PSO) approach⁶² in order to avoid local minima fitting solutions. Finally, the refined form factor values of the cubic phases (F_{hkl})

1
2 obtained from our global fits, were used to reconstruct the 3D electron density maps by applying
3 standard Fourier transform procedures⁶³⁻⁶⁴. Based on the global data analysis methods for SAXS
4 data (Eq. 2 to 5), the bilayer as well as the water-layer thicknesses in the lamellar and cubic phases
5 could be estimated.
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11 Results and Discussion

12 Emulsification of lipid self-assemblies into discrete particles

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15 By preparing a range of emulsions of DU and PT (stabilizer-sensitive and none-sensitive,
16 respectively), we verified the emulsifying role of bile salts. Both lipids, DU¹⁶ and PT³⁸ are known to
17 form gel-like and highly viscous bicontinuous cubic *Pn3m* phase in excess water. A variety of
18 interfacial stabilizers⁶⁵⁻⁶⁶ have been used to disperse viscous lipid phases and produce fluid
19 emulsions targeting their applications for biotechnological industries⁶⁶⁻⁷⁰. In this manner, the
20 thermodynamically equilibrated cubic bulk phase are converted into kinetically stabilized discrete
21 submicron-sized particles commonly termed as 'cubosomes'⁷¹. In this study, we were able to
22 create such dispersions by using a range of BS concentrations. No additional stabilizer was
23 required. However, we noticed that a certain BS to lipid ratio is necessary for obtaining
24 homogeneous and stable emulsions. At very low bile salt concentration, for instance, 0.06 wt% BS
25 and the DU concentration being >5 wt%, the resultant liquid crystalline phases do not disperse very
26 well. These phases are insoluble and thus remain completely separated from the aqueous bulk
27 phase in the form of lipid lumps (Figure 2b,c: ①). Dispersions displaying this appearance were
28 designated as *unstable* emulsion (Figure 2c; blue region). In contrast, at high BS concentrations,
29 e.g. ≥ 0.06 wt% and a DU concentration of 5 wt%, the liquid crystalline phase was dispersed well
30 and attained a '*stable emulsion*' (Figure 2b,c: ③,④; ochre region). Such low bile salt
31 concentrations (0.06 to 1.0 wt%) required for stabilization, clearly reveals the strong emulsifying
32 property of BS.
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1 **Figure 2.** a) Interfacial stabilization of lipid particles by bile salts into an oil-in-water emulsion. The
2 particles contain *Im3m* or *Pn3m* cubic phases in the cores of these lipid particles for DU and PT,
3 respectively. b) DU-BS mixtures (indicated in panel c) phase diagram) are designated as follows: ①
4 unstable emulsion displaying a lipid lump in a phase separated aqueous solution, ② pseudoplastic
5 emulsions with a white-yellowish appearance, ③ stable milky fluid emulsion, and ④ translucent
6 emulsion at high BS concentration, attributed to the strong presence small unilamellar vesicles
7 and/or micelles. c) A phase diagram describing an emulsification behavior of DU for various lipid-
8 bile salt concentrations. A dotted line indicates the border between unstable and stable emulsion
9 (homogeneous) regions supported by the measurements taken (blue diamonds). In a small
10 concentration regime, this boarder displays a linear behavior as shown in d). The cubic *Pn3m*,
11 cubic *Im3m* and Lamellar phases shown in the corresponding regions were determined from SAXS
12 analysis. e) Normalized volume distribution data for DU based emulsions (5 wt%) for various BS
13 concentrations monitored using the DLS technique. f) The hydrodynamic diameter values
14 (obtained from DLS) indicate the decrease in particle size of DU-based emulsions as a function of
15 BS concentration (for points a, b, c, see phase diagram in panel c). g) Rheological behavior of DU-
16 BS emulsions with the solid lines displaying fits to the Carreau model⁷². h) The increase in zero
17 shear viscosity with increasing lipid concentration is displayed.

22
23 A range of compositions for DU or PT with BS resulted in the formation of homogeneous
24 emulsions, which remained stable over the examined period of a month. For a generalized
25 comparison, we use the previously defined parameter β (see Eq. 1 in the Materials and Methods
26 section) as the BS to lipid weight ratio given in percent. We note that stable emulsions were
27 evident for mixtures with β -values ranging from 1.25% to 6%. Using visual inspections of the
28 samples (at least three samples prepared under the same conditions), a clear stabilization-
29 boundary could be determined (Figure 2c: dotted line). The boundary line follows a linear trend up
30 to 10 wt% DU (Figure 2d). Here the β -values required for full stabilization, display an asymptotic
31 behavior approaching the value $\beta = 2.35\%$. This signifies that at about 10 wt% lipid, the β -value has
32 reached a point of saturation, equal to say, the lipid particles sizes have reached a local minimum
33 here. At higher lipid concentrations, however, the β -values start to increase again (non-linear
34 regime of the lipid to BS weight function) (Figure 2c). In other words, even larger BS to lipid ratios
35 are required for stabilizing emulsions with DU contents above 10 wt% (Figure 2c).

36
37 Self-assembled nanostructured phases in the stable DU emulsions were determined by small angle
38 X-ray scattering (SAXS) and their detailed analysis will be discussed further with Figures 3, 5 and 6,
39 here we shall only illustrate their appearance within the evaluated phase diagram. At 0.06 wt% BS
40 for 5 wt% DU a stable bicontinuous cubic *Im3m* phase dispersion was detected, which upon
41 increase in BS concentration, converted into a stabilized lamellar L_{α} phase dispersion (Figure 2c:
42 point a to b to c), and as can be expected with increasing amount of stabilizer, the overall particle
43 sizes decrease significantly with increasing BS-concentration (Figure 2e, f). The particle size, as

1 determined by DLS technique (Figure 2e) in the stable region stayed in the range of 120 to 280 nm.
2 Noteworthy, stable emulsions prepared from PT, required lower concentrations of bile salts as
3 compared to DU; for instance, 15 wt% DU required at least 0.5 wt% BS, whereas the same
4 concentration of PT needs only 0.2 wt% BS. PT-based emulsions also displayed lower particle sizes
5 in the range of 120 to 190 nm (Figure S1 in Supporting Information). The particle size
6 (hydrodynamic diameter) for DU emulsions decreased with increasing the BS concentration
7 (Figure 2f) depicting the fact that greater β -values at a given lipid concentration lead to a better
8 emulsification, i.e., increasing the surface to volume ratio of the particles, hence, decreasing the
9 overall particle sizes. The same trend was observed for PT-based systems (Figure S1); relatively
10 greater accumulations of bile salts at the lipid-water interface led to the stabilization of smaller
11 discrete lipid particles. Thus, an important outcome of the lipid emulsification process driven by
12 bile salts is that highly viscous cubic phases get fragmented into smaller discrete particles, which in
13 turn, can be easily assessed by digestive molecules as compared to sticky and insoluble lipid
14 lumps.

15 At low lipid concentrations (5 wt% DU) stable non-viscous dispersion were observed (Figure 2b,c:
16 ③④), however, at higher DU concentrations the viscosity was seen to clearly build up in these
17 samples. Especially >10 wt% DU emulsions displayed a highly viscous appearance (Figure 2b:
18 ①②), which contrasts the fluid consistency observed for the low lipid concentration emulsions.
19 This behavior was confirmed by rheological studies at 25 °C (0.5 wt% BS varying DU from 10 to 20
20 wt%) revealing a pseudoplastic behavior of these emulsions (Figure 2g, h). The viscosity profiles
21 (Figure 2g) display a decay in viscosity as a function of shear rate, which is an indication of shear
22 thinning or pseudoplastic behavior; such viscosity behavior fits well with the Carreau model⁷²⁻⁷³
23 defined by solid lines in Figure 2g. The zero-shear viscosities deduced using this model represent a
24 clear increase at high lipid concentrations (Figure 2h). Such increase in shear viscosity with the
25 increase in lipid concentrations is common for self-assemblies and has been reported earlier⁷⁴. We
26 note that shear thinning properties of liquid crystalline material-based emulsions are desirable for
27 formulation processing in pharmaceutical and cosmetic industries⁷⁵. However, the pseudoplastic
28 behavior was observed only for DU based emulsions; on the contrary, all PT based emulsions could
29 be dispersed without difficulty (no lipid lumps formed), i.e. their consistency was fluid even at high
30 lipid concentrations (>10 wt%). Their optical appearance was milky white and only at very low PT
31 concentrations, it was translucent to shiny. From this entirely fluid consistency of the PT-samples
32 and from their optical inspection, one can anticipate their Newtonian fluid behavior. This

1 demonstrates that the increasing amounts of non-dispersed *Im3m* phase assemblies are mainly
2 responsible for the augmenting pseudoplastic behavior with increasing DU concentrations (also
3 represented by the lumpy appearance of sample ① in Figure 2b). Similarly, the viscosity build-up
4 caused by an entanglement of worm-like mixed micelles in lipid-bile salt mixtures has been
5 reported earlier.^{8, 29, 76}
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10 11 12 **Influence of bile salts on the nanoscale architecture of self-assembled** 13 **nanostructures** 14 15

16 As mentioned in the introduction, it is well documented that DU and PT form the *Pn3m* cubic
17 phase in excess water in bulk (non-dispersed state).^{16, 38} The nanostructural analysis of BS
18 stabilized emulsions formulated from DU reveals that BS molecules not only interact with the
19 surface of the submicron sized particles and stabilize them, but also induce changes in the
20 interfacial membrane curvature leading to phase transitions as explained below. The type and
21 nanostructural changes of the internal lipid self-assembly of various dispersed DU particles (lipid
22 concentrations = 5, 10, 15 and 20 wt%) were analyzed by small angle X-ray scattering (SAXS)
23 (Figure 3a). At 20 wt% lipid ($\beta = 2.5\%$), the *Im3m* cubic phase with a lattice parameter of 15.5 nm
24 was observed, rather than the cubic *Pn3m* phase as expected from bulk DU-water systems (Figure
25 3a). The same phase transition, i.e., from *Pn3m* to *Im3m*, has been observed for various DU based
26 systems using the 'gold standard' block-copolymer Pluronic® F-127 and various other types of
27 stabilizers⁷⁷⁻⁷⁸. Reducing the lipid concentration, from 20 down to 15, 10 and 5 wt%, meaning
28 increasing the relative BS concentration in the emulsion (increasing β -values), leads the gradual
29 disappearance of cubic phase structures (peaks indicated by arrows in Figure 3a).
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41 The SAXS analysis on PT based emulsions confirms the presence of the cubic *Pn3m* phase instead,
42 as originally observed for the bulk PT-water system³⁸, for all (lipid concentrations = 5, 10, 15 and
43 20 wt%) samples stabilized by 0.2 wt% BS (Figure 3b). The absolute lattice parameters of the *Pn3m*
44 phase, however, increased from 6.75 to 7.23 nm as the BS-lipid ratio was increased (β value from
45 1 to 4%).
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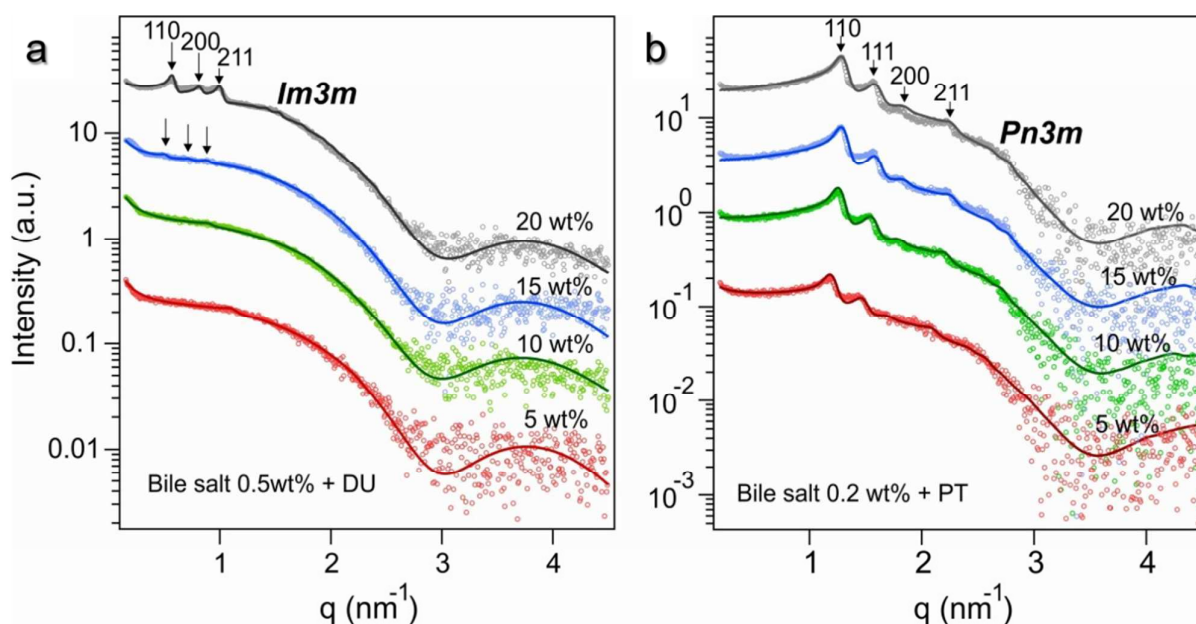


Figure 3. SAXS patterns for a) DU and b) PT based emulsions stabilized by bile salts. The global analysis of scattering curves was performed by fitting of the curves with a model provided by Garstecki and Holyst⁵⁶⁻⁵⁷ (for cubic phases) combined with a planar bilayer model from Pabst and co-workers^{54, 79}. The solid lines represent the optimized model fits attained by Particle Swarm Optimization⁴⁸. These studies demonstrate that lamellar (L_{α}) phase co-exists with cubic phases and the broad background scattering mainly originates from the form factor contribution of lamellar phases. The cubic $Im3m$ phase (indicated by arrows for Bragg diffractions with Miller indices of 110, 200 and 211) for DU disappears when the bile salt to lipid ratio is increased, while the cubic $Pn3m$ phase (with Bragg peaks of miller indices 110, 111, 200 and 211) persists in the case of PT based emulsions.

The novel global analysis of scattering curves (see Methods section) was performed by fitting each with the models provided by Garstecki and Holyst⁵⁶⁻⁵⁷ (for cubic phases) combined with a planar bilayer model description based the modified Caillé theory (for lamellar phase)⁵⁵. It revealed the co-existence of lamellar phase along with $Im3m$ and $Pn3m$ phases as observed correspondingly for DU and PT emulsions. Drawing an interim conclusion, we have clearly shown that the BS is not only an *efficient emulsifier*, but at the same time also *promotes less curved lipid/water interfaces*. This is experimentally confirmed (i) by the formation of the least-curved bicontinuous $Im3m$ phase after the addition of BS to DU (originally displaying the cubic $Pn3m$ phase) (Figure 2c: point a), (ii) by the transformation of the $Im3m$ phase into the L_{α} phase at higher BS concentrations (Figure 2c: point a to b to c, and Figure 3a), and (iii) in the case of PT, by the lattice parameter increase of the $Pn3m$ phase as a function of BS concentration (Figure 3b, 6f).

Remarkably, other bile salts such as sodium taurochenodeoxycholate (NaTCDC) have been reported earlier to stabilize egg yolk lecithin (phosphatidylcholine) (EYPC) based vesicles at low BS concentrations, while inducing worm-like cylindrical mixed micelles at higher concentrations⁹. These mixed micelles were investigated with small-angle neutron scattering and displayed semi-flexible rods with the persistence length of about 18-20 nm (Kuhn length 36-40 nm). Similarly also glycochenodeoxycholate (GCTC) was studied by the same group.¹⁰ Again, GCTC induced the formation of worm-like mixed micelles at higher BS concentrations, which would further grow with increasing GCTC concentration.

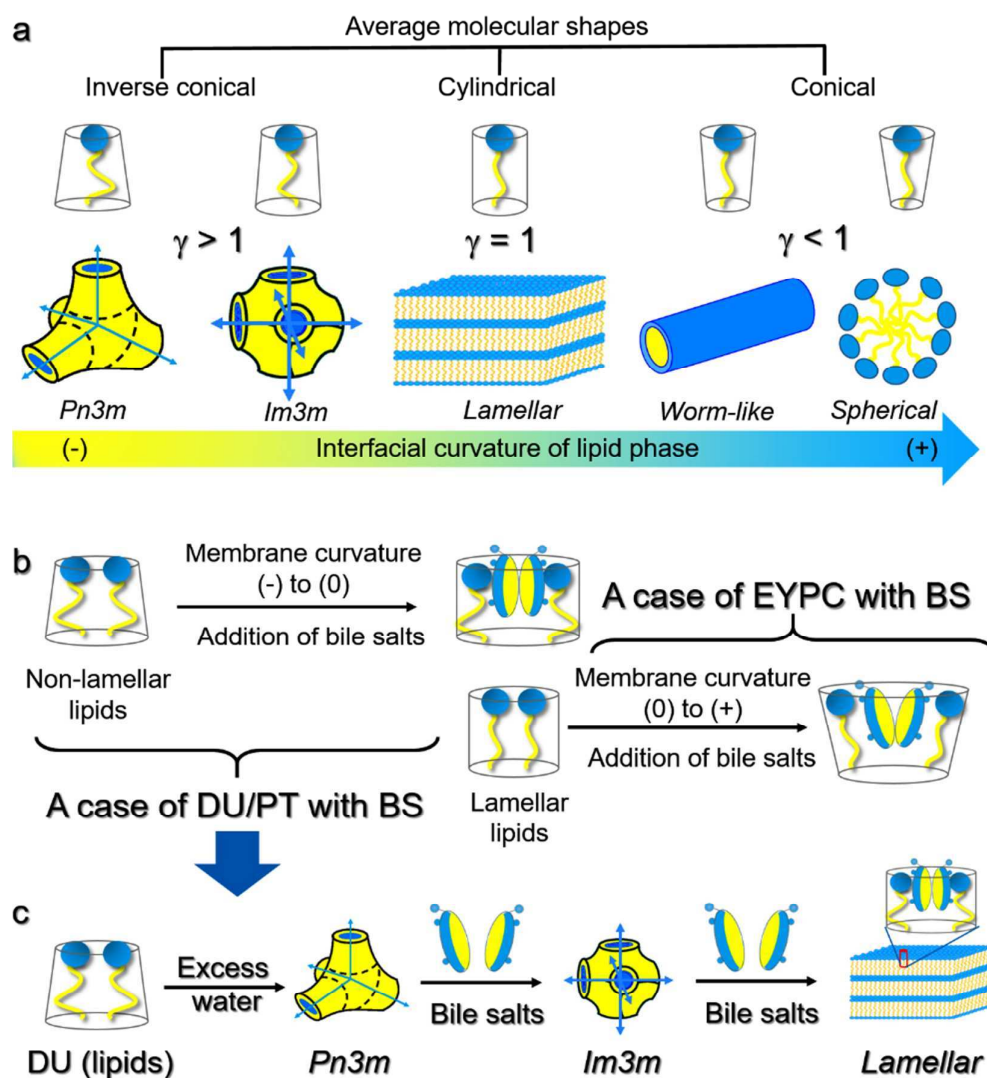


Figure 4. a) Average molecular shapes adopted by common lipids and corresponding self-assembled nanostructures observed. Inverse conical molecules (with $\gamma > 1$) form inverse nanostructures exhibiting negative interfacial curvature, whereas conical molecules (with $\gamma < 1$) form phases with positive interfacial curvature; cylindrical molecules commonly self-assemble into planar (lamellar) phases. b) In case of DU/PT and BS mixtures (studied here), BS molecules reside in the head group regions thereby altering the inverse conical shapes into more cylindrical shapes.

1 Similarly, the cylindrical shape of the molecules like EYPC is transformed into conical shape due to
2 BS effect.⁹⁻¹⁰ c) In case of DU, the $Pn3m$ phase formed by pure lipid in water is converted into
3 $Im3m$ cubic phase, which upon further addition of bile salts converts into lamellar phase indicating
4 systematic deviation from inverse conical to cylindrical molecular shape.
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7 Although our applied bile salt mixture, and the pure NaTCDG and GCTC from the above mentioned
8 studies do exhibit differing side groups, their 'membrane curvature power' is qualitatively the
9 same. In all cases, the bile salts have driven the lipidic membrane systems towards positive
10 curvature (Figure 4): that is, for DU we observed a $Pn3m$ to $Im3m$ to L_α phase conversion with
11 increasing BS concentration (Figure 2c: point a to b to c), and for PT an increase in the lattice
12 parameter of $Pn3m$ phase (Figure 3b, 6f), and for EYPC the L_α phase was found to get converted
13 into normal cylindrical-shaped micelles.⁹⁻¹⁰
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20 On a molecular level this observation is best explained with the related critical packing parameter
21 (CPP) usually denoted by γ , which according to Israelachvili⁸⁰ is

$$\gamma = \frac{v}{a_0 l_c} \quad (6)$$

22 where v is the hydrophobic volume of the lipid, a_0 is the interfacial area of the hydrophilic head
23 group and l_c is the critical chain length, which is the maximum effective length that the
24 hydrophobic chains can assume. Bicontinuous cubic phases are formed with molecules that have a
25 γ of above 1, which in case of monoolein (main component in DU) molecules is 1.30⁶⁴. Note, the
26 relatively small head group of monoglycerides (relatively small a_0) and the unsaturated oleic chain
27 (relatively short l_c) render the monoolein molecular shape to be inversely cone-shaped promoting
28 negative interfacial curvatures, or in other words, leading to inverse phase assemblies (water-in-oil
29 phases) (Figure 4). These lipids are sometimes referred to as non-lamellar lipids as they form non-
30 lamellar phases. The influence of BS on PT assemblies was least efficient, since only small changes
31 towards positive curvature were observed (increased lattice parameter of the conserved $Pn3m$
32 phase upon addition of BS). This becomes immediately understood by taking into account the
33 bulky hydrocarbon chain given in PT (additional methyl groups along the chain lead to an overall
34 increase in the chain volume). Last, NaTCDG and GCTC added to EYPC membranes in the L_α phase
35 with a γ of about 1, were reported to induce the formation of normal worm-like micelles having a γ
36 in the range of 0.33 to 0.50⁹⁻¹⁰ (Figure 4). Thus, we can recapitulate that bile salts do drive lipid
37 self-assemblies towards aggregates with a lower γ ; i.e., towards positive interfacial curvatures
38 (towards oil-in-water phases) (Figure 4).
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To investigate this BS-induced interfacial curvature trend further and understand the effect of bile salts in more details, we performed another set of experiments, where the lipid content was kept constant and the BS concentration was increased (Figure 5). The BS concentration was varied between 0.06 and 1.0 wt% for emulsions prepared from 5 wt% DU or PT. The cubic $Im3m$ phase with lattice parameter of 12.8 nm was observed at 0.06 wt% BS for DU emulsions (Figure 5a). By further increasing BS concentration to 0.1 wt% or above, the cubic phase gradually disappeared and undergone a rearrangement to lamellar L_α phase (Figure 5a). We note, primarily only unilamellar bilayers were formed and then, at BS concentrations about 0.2 wt%, a bilayers correlation peak arises, indicating the formation of multilamellar bilayer stacks (inset of Figure 5a). The disappearance of cubic phase and concomitant formation the L_α phase were computed as a function of the β -value (BS/lipid ratio in %); the volume fraction of the $Im3m$ phase is shown in Figure 5c. At about $\beta = 3\%$ (BS = 0.15 wt%) half of the cubic phase has converted into the L_α phase.

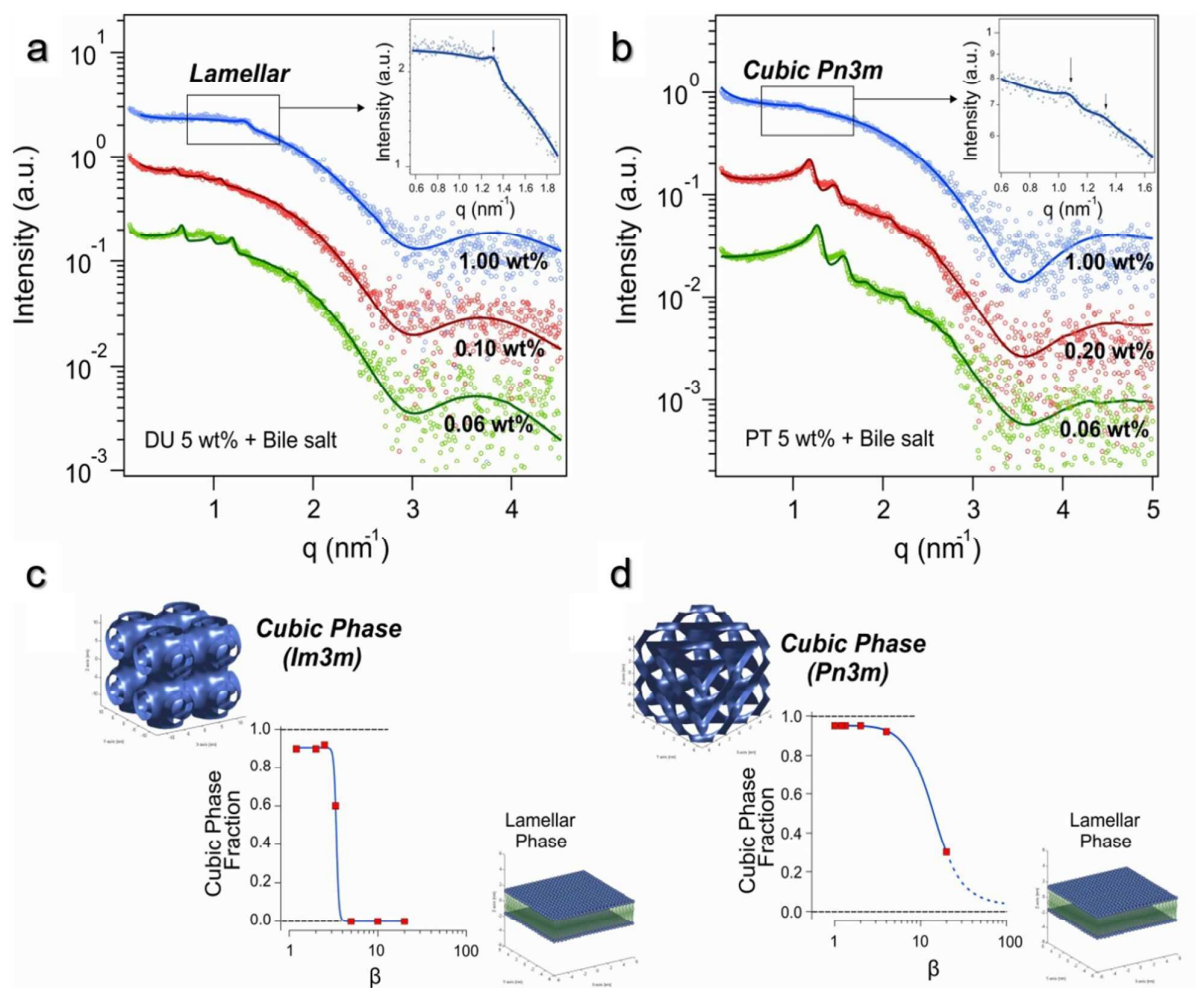


Figure 5. SAXS patterns for 5 wt% a) DU and b) PT based emulsions stabilized by various concentrations of bile salts. The solid lines represent the globally best fits to the data (for details

1 see methods section). Relative intensities of cubic phase peaks decrease with increasing bile salt
2 concentration. The insets in panel a) and b) do show traces of multilamellar vesicles and of the
3 *Pn3m* phase, respectively. Bile salt driven transformation of cubic to lamellar phases via
4 coexistence regime is depicted by the computed fractions of cubic phases as function of β for DU
5 and PT in the corresponding panels c) and d).
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10 Similarly, the experiments conducted on PT indicate the coexistence of lamellar phase with cubic
11 *Pn3m* phase. However, the fraction of the cubic phase decreases slower: only at about $\beta = 10\%$ (BS
12 = 0.5 wt%) half of the cubic phase has converted into the L_α phase, which can be seen from the
13 stronger Bragg peak intensities of the *Pn3m* phase as compared to the *Im3m* phase (Figure 5b) and
14 from the turnover function in Figure 5d. We note, the phase transitions from bicontinuous cubic to
15 vesicles or even mixed micelles are anticipated to be beneficial for digestion purposes. This is
16 because, the cubic phase remains insoluble in water even at a very high dilution levels, whereas
17 vesicles and mixed micelles are more easily accessible by different lipases⁸¹. Thus, the nanoscale
18 organization of lipid self-assemblies, from complex inverse structures (water-in-oil phases)
19 towards vesicle formation and normal micelles (spherical, wormlike and disc-like), endowed by
20 bile salts, aids lipid digestion and absorption.
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32 Nanoscale changes in the bilayer structure: thickness, molecular shape and 33 interfacial curvature 34 35

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37 The global fine analysis of scattering profiles allows to deduce estimates for the bilayer
38 thicknesses, both for the cubic phases as well as for the lamellar phase. Note, for the L_α phase a
39 simple 2-Gaussian model⁸² was refined during the fitting procedure and the results are displayed
40 in Figure 6a and b. For 5 wt% DU and 0.06 wt% BS based emulsion, the bilayer thickness for
41 lamellar phase was estimated to be 3.1 nm. Similarly, for 5 wt% PT and 0.05 wt% BS sample, the
42 bilayer thickness in lamellar phase was determined to be about 2.7 nm. Its lower value is explained
43 throughout the shorter chain length of PT, when compared to oleic acid in monoolein, which is the
44 main glycerol component in DU (C14 versus C18:1). The bilayer thickness in the cubic phases is
45 determined by a simple one-component slip model⁸³, which gave within errors the same bilayer
46 thicknesses as for the coexisting lamellar bilayers. Nevertheless, it is interesting to note that, the
47 membrane thickness in both DU and PT based lamellar phases reduces by increasing BS
48 concentration, while the retrieved bilayer thicknesses in cubic phases remain within given error
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1 margins constant (data not shown). This notion directs to the fact that in both cases BS is actually
2 interacting with the head group region and causing an increase in the area per lipid, a_0 . On the
3 other hand, the chain volume, v , is expected to remain constant at a given temperature. For the L_α
4 phase (cylindrical molecular shape; $\gamma = 1$), this actually means, that an effective shortening of the
5 chain length l_C can be expected with increasing BS concentration ($\beta_i < \beta_f$). Note, you may also
6 express this in the simply relationship: $v_i = v_f$ or $a_{oi} \cdot l_{Ci} = a_{of} \cdot l_{Cf}$, and with $a_{oi} < a_{of}$ follows $l_{Ci} > l_{Cf}$. For
7 the cubic phases, the arguments on effective chain length changes, Δl_C , are expected to show an
8 opposite effect.
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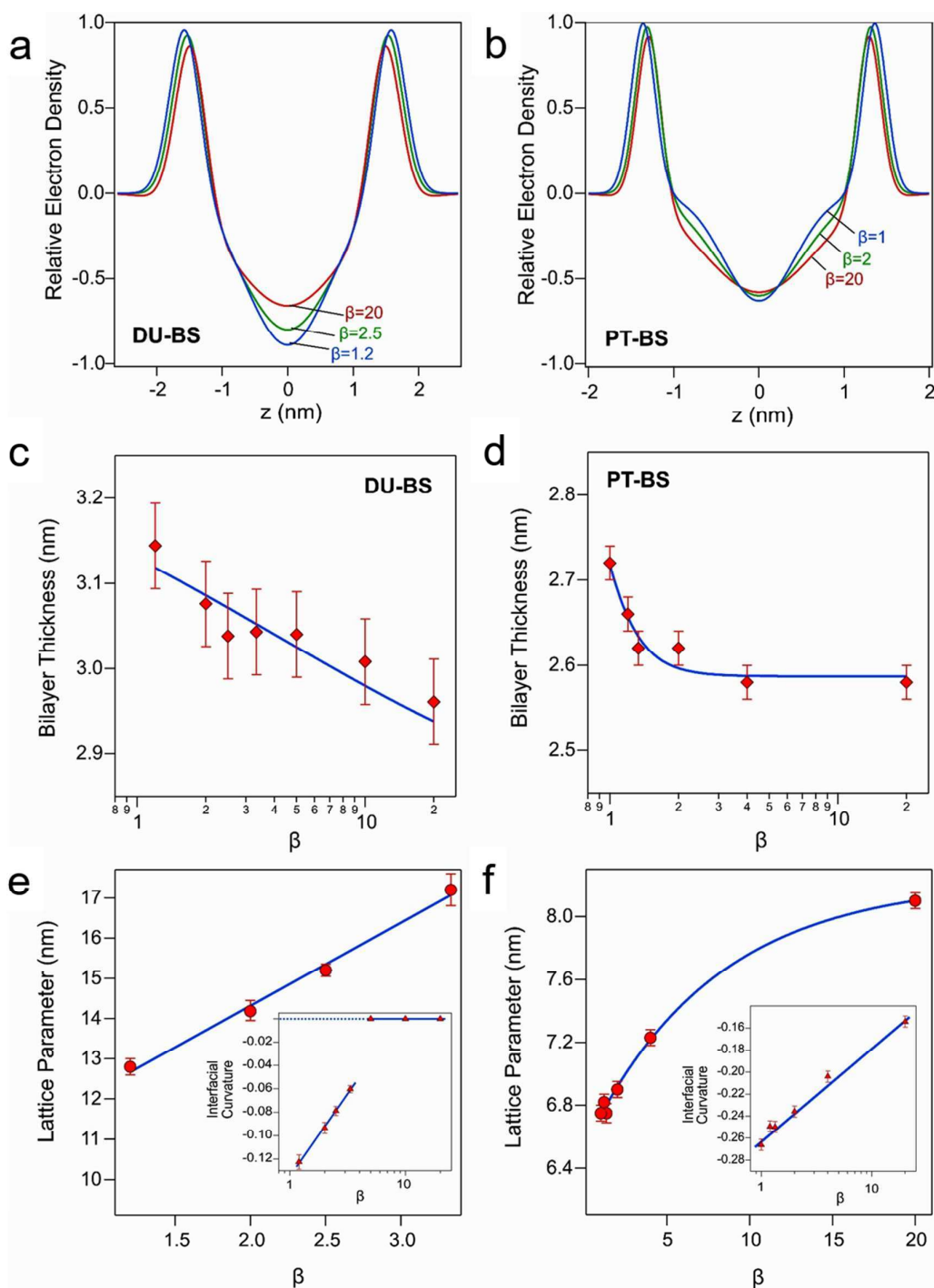


Figure 6. Computed bilayer electron density profiles for the lamellar phases for a) DU and b) PT emulsions for varying bile salt concentrations. Plots c) for DU and d) for PT indicate that the bilayer thickness decreases with bile salt to lipid ratio. Lattice parameters increase as a function of BS concentration for both e) the cubic $Im3m$ phases in DU and f) the cubic $Pn3m$ phase for PT. Insets of e) and f) correspondingly elucidate the increase in the mean interfacial curvature as the bile salt concentration increases.

As argued before, lipid molecular packing follows a simple trend: the addition of BS always leads to a reduction in the γ value. In cubic phases, an increase in a_0 makes molecular shape more

1 straight, cylindrical-like, i.e. leading to an reduction in γ (1.30 towards 1.00) (Figure 4). Again, we
2 can expect the lipid chain volume, v , in the cubic phases not to change significantly, when
3 increasing the BS concentration. Thus, a reduction in γ , should actually lead a reduced chain-splay,
4 particularly at the terminal ends of the lipid chains. Consequently, l_c values in the bicontinuous
5 cubic phases are expected to increase with augmenting BS concentration. However, the given low-
6 resolution data of the cubic phases in this work only allow a rough estimate of the bilayer
7 thickness and minor bilayer thickness trends are not possible to be deduced with the Garstecki-
8 Holyst^{42, 43} model in this case.

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16 Figures 6e and f display the lattice parameter trends of the $Im3m$ and $Pn3m$ phase, which both
17 increase with BS content, i.e. both cubic phases swell (reduction of the interfacial curvature
18 modulus). Accordingly, on a molecular level, we estimate the γ value to change from 1.25 ($Im3m$)
19 at low BS concentrations to 1.00 (lamellar) at high BS concentration for DU, whereas the γ value
20 changes from 1.46 ($Pn3m$) to 1.00 (lamellar) for PT based self-assemblies. Note, here we estimated
21 the γ by Hyde's ansatz⁸⁴ applying $\gamma = 1 + H_i l + G_i l^2 / 3$, in which H_i is the mean interfacial
22 curvature modulus, G_i denotes the Gaussian interfacial curvature modulus and l the lipid length (=
23 half the bilayer thickness).

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25 In summary, the induction of positive (or less negative) average mean interfacial curvature by bile
26 salts, confirms the molecules' tendency to mostly occupy the head group (or interfacial) region.
27 Inverse bicontinuous cubic phases are known to exhibit slightly negative mean curvatures,
28 whereas it is close to zero for planar lamellar phases.⁵¹ The insets in Figure 6e and f represent the
29 variation of mean interfacial curvature $\langle H_i \rangle$ calculated with,

$$\langle H_i \rangle = 2\pi\chi l / S_i \quad (7)$$

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31 where χ is the Euler characteristic and S_i is the area at the interface integrated over a single
32 monolayer ($S_i = S_0 a^2 + 2\pi\chi l^2$). a is the lattice parameter and l is the lipid length.⁸⁵

33 Conclusion and Perspectives

34 Strong emulsifying power of bile salts (BS) is clearly demonstrated via interfacial stabilization of
35 fragmented oily and rather hydrophobic lipid residues into submicron-sized particles. Relative
36 hydrophilicity and consequential solubility of lipids was elevated by BS-shielding of the self-
37 assembled inverse lipid nanostructures that are otherwise less soluble in water (Figure 2b).
38 Furthermore, the effective surface to volume ratio of bulk cubic phases was improved due to the

1 formation of discrete submicron (< 300 nm) particles (Figure 2e). The particle size was clearly
2 reduced with increasing the BS concentration depicting efficient emulsion stabilizing role of bile
3 salts (Figure 2f). The overall viscosity ($\sim 10^4$ - 10^5 pa·s)¹⁴ of bulk cubic phases was drastically reduced
4 by transforming them into dispersions. However, some degree of viscosity was raised again at
5 higher concentrations of lipid, especially in case of DU, but it was not as high as for the original
6 bulk cubic phases ($\sim 10^2$ pa·s) (Figure 2g). Such an increase in viscosity can be attributed to the
7 non-dispersed fractions of viscous cubic phases. To summarize, the bile salts act as interfacial
8 stabilizers for oil-in-water emulsions prepared from lipid cubic phases.
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10 Another important role of BS, portrayed in this work, is its influence on the type of lipid
11 nanostructure; at increasing BS concentrations highly complex cubic phases were transformed into
12 vesicles. Having applied an *ad hoc* developed novel global SAXS fitting procedure, we were able to
13 underpin the mechanism behind these phase transitions at a molecular level. Increasing bile salt
14 concentrations, the obtained decrease in the bilayer thickness in lamellar phases (Figures 6), and
15 simultaneous increase in the area per lipid, a_0 indicate that BS prefers to interact in the lipid head-
16 group region. This was also illustrated by a BS-driven change in the molecular shapes (inverse cone
17 to cylinder) adopted by cubic phases (Figures 4). Results reveal that the BS significantly decreases
18 the average critical packing parameter, γ , i.e. bile salts *drive the interfacial membrane curvature*
19 *towards positive values* (Figures 4a, 6e, f). These results are in good agreement with the literature
20 reports, where cylindrically shaped molecules were shown to adopt conical shapes due to the
21 interaction of BS molecules and the interfacial curvature modulus of the resultant phase changes
22 from zero to positive values.⁸⁻¹⁰ Formation of higher positive curvature lipid assemblies evidently
23 means an enhancement of accessible hydrophilic interface area, and therefore leading to a greater
24 propensity to disperse or solubilize them in aqueous **digestion media**.
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42 **This work** highlights the importance of the '*dual role of bile salts*' in fat digestion. Not only, that fat
43 globules get efficiently emulsified by bile salts, but inverse lipid assemblies (oil-in-water phases)
44 are driven - depending on the initial lipid composition - to form either BS-stabilized vesicles or
45 normal BS-lipid mixed micelles, which are both more readily accessible in subsequent digestion
46 **steps and enhance lipid absorption**.
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52 Acknowledgment

53 We acknowledge Amin Farshchi for his support on rheological measurements.
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Author contributions

CVK planned the project and experiments. SM and AS performed experiments. SM, AS, MR and CVK analyzed and interpreted results. AS and MR developed *ad hoc* global SAXS analysis method. AS, MR and CVK wrote the manuscript.

Competing interests

We declare no conflict of interest regarding the work and the manuscript.

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