Effect of Bile Salts on the Interfacial Dilational Rheology of Lecithin in the Lipid Digestion Process

Aicha Mekkaoui¹,², Yang Liu¹, Pingping Zhang¹, Sana Ullah¹,², Ce Wang¹,²*, and Baocai Xu¹,²*

¹ School of Light Industry, Beijing Technology and Business University, Beijing 100048, CHINA
² Beijing Advanced Innovation Center for Food Nutrition and Human Health, Beijing Technology and Business University, Beijing 100048, CHINA

Abstract: The effects of bile salts on the emulsifier adsorption layer play a crucial role in lipid digestion. The current study selected sodium cholate (NaCh) and lecithin as model compounds for bile salts and food emulsifiers, respectively. The interface dilational rheological and emulsification properties of NaCh and lecithin were carried out. The results showed that the NaCh molecules could quickly diffuse from the bulk to interface, which broke the tightly-arranged interfacial layer of lecithin and enhanced the viscoelasticity of interfacial film. As a result, the interfacial adsorption layer, which was originally dominated by the slow relaxation processes within the interface, was transformed into one controlled by the fast molecular diffusion exchange. This accelerated the exchange of materials between the bulk and interface, thereby creating suitable conditions for the interfacial adsorption of lipases, which promoted the digestion process. These results provided a mechanism for the promotion of lipid digestion by bile salts from the perspective of interfacial viscoelasticity and relaxation processes. A deeper understanding of the interfacial behavior of bile salts with emulsifiers would provide a basis for the rational design of interfacial layer for modulating lipid digestion.

Key words: bile salt, lecithin, emulsion, lipid digestion, interfacial dilational rheology

1 Introduction

During the last decade, the occurrence of obesity and other long-term chronic diseases, such as cardiovascular disease, diabetes and certain cancers, is continuously increasing worldwide, which is mainly due to the changing lifestyles and patterns of food consumption. High-fat diet is a major contributor, causing the imbalance of energy intake and appearance of unhealthy conditions¹. The regulation of bioavailability of dietary fats by controlling lipid digestion is believed as an effective approach to prevent the increasing obesity and other related diseases². Currently, increasing studies are focusing on the reduction of rate and degree of lipids digestion using functional foods, for effective reduction of lipid absorption and promotion of satiety³⁻⁶.

In brief, the lipid digestion mainly occurs in gastrointestinal tract. When the digested food is transported into small intestine, the lipids are mixed with pancreatic juice, containing bicarbonate of soda, bile salts, phospholipids, pancreatic lipase, and co-lipase. As a result of interfacial activity, the bile salts facilitate the emulsification of lipids by adsorbing to the surfaces of droplets and contribute to the binding of enzymes to lipid substrates⁷. For the lipid emulsions stabilized with emulsifiers before entering small intestine, the interfacial films prevent lipases from coming into close proximity to lipids. In this case, the bile salts compete and displace the pre-adsorbed amphiphiles at the interface, which makes it possible for lipases to bind to the surface of lipid droplets. Then, the hydrolysis of triacylglycerols takes places under the catalytic action of pancreatic lipase/co-lipase⁸. After that, the products of lipid digestion, including diacylglycerol/monoacylglycerol (DAG/MAG) and free fatty acid (FFA), and the components dissolved in lipids, such as oil-soluble vitamins, phospholipids, and cholesterol, are removed from the interface by the bile salts and form mixed micelles or vesicles, which then pass through the mucous layer that covers the epithelium walls and finally reach to enterocyte cells⁹⁻¹⁰. It can be seen that bile salts play a significant role in the interfacial process of lipid digestion.

As a class of amphiphilic molecules secreted by human body, bile salts are derived from cholesterol and have a steroidal core structure¹¹. Unlike classical amphiphiles that
consist of hydrophilic head-groups and hydrophobic tails, bile salts exhibit planar polarity. Their hydrophobic surface lies on the convex side of the rigid steroid ring structure with angular methyl groups at positions C-18 and C-19, while the hydrophilic surface lies on the concave side with one, two, or three hydroxyl groups and a carboxyl group conjugated with taurine, glycine or other amino acids. Due to this unique structure, they are less effective in terms of lowering the interfacial tension (IFT) than many other surfactants, but more effective in terms of competing with other amphiphiles at interface. This has been proved by detection of changes in zeta-potential, IFT, or interfacial dilational modulus before and after the addition of bile salts. The designing of an appropriate interface adsorption film, inhibiting the competitive adsorption of bile salts, is a well-accepted feasible approach for the regulation of lipid digestion. However, it is still unclear how the bile salts displace emulsifier at the oil-water interface, because the intermolecular interactions and relaxation processes during the replacement cannot be well explained simply by monitoring the changes of zeta-potential, IFT, or interfacial dilational modulus. But, this is important for the rational design of emulsification system for the regulation of lipid digestion. A better understanding of the interfacial processes responsible for the displacement of emulsifiers by bile salts is still needed.

According to the literature, the measurement of the dilational rheological properties of interface can provide a credible means for the detection of its viscoelasticity and relaxation processes, reflecting the interaction of molecules at the interface. The use of this technique in physiological interfacial issues has now been receiving increasing attention. The microscopic interfacial behaviors of amphiphiles can be inferred by detecting the dependence of interfacial viscoelasticity on the concentration of amphiphiles, interfacial oscillating frequency, and interfacial pressure. Therefore, it was used to study the interfacial dilational rheological behaviors of emulsifiers in the presence of bile salts in the present work.

Lecithin is one of the most widely used food emulsifiers, containing two non-polar fatty acids, which are esterified into the backbone of glycerol with a polar phosphate head-group attached to a hydrophilic residue, such as inositol, choline, serine or ethanolamine. As zwitterionic surfactants, lecithin can effectively lower oil-water IFT and stabilize oil-in-water emulsions. In the current study, soybean lecithin was used as a representative of exogenous food emulsifiers. On the other hand, despite the varieties, all the bile salts behave qualitatively in a similar manner. Herein, sodium cholate (NaCh) was used as an endogenous bile salt. The interactions between lecithin and bile salts have attracted increasing attention because of their ability to form mixed micelles that facilitate the digestion of lipids in intestine. However, studies on the interfacial inter-action between these two amphiphiles are rarely reported.

The emulsification and interfacial behaviors of lecithin, NaCh, and their mixture were studied in this work. The interfacial behaviors were assessed by the IFT and interfacial dilational rheology at the oil-aqueous interface using oscillating drop method. The emulsification behaviors were estimated by the distribution of droplet size and destabilization index. Finally, the in vitro lipid digestion of lecithin-stabilized emulsion was carried out in the absence and presence of NaCh, respectively. Remarkably, the presence of NaCh greatly altered the interfacial viscoelasticity of lecithin, which provided a mechanism of the promotion of lipid digestion by bile salts from the perspective of interfacial dilatation rheology.

2 Materials and Methods

2.1 Materials

NaCh hydrate was obtained from Beijing Budweiser Technology CO., Ltd. Soybean lecithin was obtained from Tokyo Chemical Industry CO., Ltd. The molecular structures of NaCh and Soybean lecithin are shown in Scheme 1. Medium-chain triacylglycerols (MCT) were obtained from Shanghai Yuanye Bio-Technology Co., Ltd. Tris(hydroxymethyl)methyl aminomethane was obtained from Shanghai Macklin Bio-Technology Co., Ltd. Hydrochloric acid (HCl) and NaOH were obtained from Sinopharm Chemical Reagent Co., Ltd. Sodium chloride (NaCl) and calcium chloride (CaCl₂) were obtained from Beijing J&K Scientifc Ltd. Lipase was obtained from Shanghai Yuanye Bio-Technology Co., Ltd. Ultrapure water with a resistivity of 18.2 MΩ·cm was used for the preparations of solutions. n-Decane, with a purity of 99.5%, was obtained from Beijing Budweiser Technology CO., Ltd. All the other chemicals used in this study were of analytical grade.

2.2 Interfacial tension and dilational rheological measurements

The measurements of IFT and interfacial dilational rheology were taken using an oscillating drop tensiometer ODG-20 (DataPhysics Instruments GmbH, Germany). This instrument mainly composed of a computer-controlled syringe with a needle for droplet formation, quartz cuvette,

![Scheme 1](Image)

**Scheme 1** Molecular structures of NaCh and lecithin.
CCD camera, light source, and a frame grabber. The interface was created by injecting an oil droplet from a reversed stainless steel needle into the aqueous phase in cuvette. A sinusoidal periodic compression and expansion of the interface area (ΔA) was performed at a selected amplitude (ΔA/A, 10%) at a frequency of 0.1 Hz. The image of the droplet was taken with the CCD camera. IFT (γ) was calculated by fitting the Laplace equation to the shape of the droplet. As primarily defined by Gibbels, the dilational modulus (ε) can be expressed as given in Eq. (1).

\[ \varepsilon = -\frac{d\gamma}{d \ln A} \] (1)

The interfacial dilational modulus can also be expressed as a complex number, which contains a real portion (storage modulus), representing the elastic energy kept in the interface, and an imaginary portion (loss modulus) resulted from interfacial relaxation processes, which match to the elasticity (\varepsilon_r) and viscosity (\varepsilon_\eta = \omega \eta d), respectively, as given in Eq. (2).^\text{40}

\[ \varepsilon = \varepsilon_r + i \omega \eta d \] (2)

When the adsorption of amphiphiles at interface reached equilibrium, the interfacial oscillations at different frequencies (0.005~0.1 Hz) were carried out to study the interfacial viscoelasticity.

All the measurements were performed at 30°C. The complex composition of MCT makes it difficult to analyze the interface behavior of amphiphilic molecules. Therefore, n-decane was selected as simulation oil in this experiment. Lecithin was dissolved in the n-decane as oil phase, while NaCh was dissolved in water as aqueous phase. It should be noted that, for the lecithin oil solution-NaCh aqueous solution systems, the concentration of NaCh in aqueous phase and that of lecithin in oil phase were equal, and the concentrations in the results and discussion part were equal to the sums of the concentration of NaCh in the aqueous phase and the concentration of lecithin in the oil phase.

2.3 Preparation of emulsion

NaCh was dissolved in water at different concentrations (0.1 wt%, 0.5 wt%, 1.0 wt%, and 1.5 wt%) by stirring gently until complete dissolution. The pH of aqueous solutions was adjusted to 7, according to the condition of small intestine. The oil phase was prepared by dispersing lecithin in MCT and stirring until complete dissolution. The oil and aqueous phases were mixed with a volume ratio of 2.8 (O/W) using a high-shear mixer (15-081, Ningbo Scientz Biotechnology Co., Ltd) at a speed of 8000 rpm for 3 min to form coarse emulsion. The emulsion was then homogenized using a Panda PLUS processor (GEA Niro Soavi Corporation, Italy) at the operational pressure of 800 MPa for five times.

2.4 Determination of droplet size

The droplet size of emulsions was determined using a dynamic light scattering instrument (Zetasizer Nano ZS, Malvern, U.K.). In order to avoid multiple scattering effects, the samples were diluted 100 times with ultrapure water prior to taking measurements. The droplet sizes were calculated from the average of 5 readings taken using the same sample. All the measurements were carried out at least in duplicate with three prepared samples.

2.5 Stability of emulsion

Multiple light scattering measurements were taken for the evaluation of the stability of emulsion using a commercial apparatus, Turbiscan (Formulation, L’Union, France). The freshly prepared emulsion samples were placed in cylindrical quartz tubes and the measurements were taken at 30°C for 24 h. The dispersion of emulsion was analyzed using two detectors (0° from the incident beam-the transmitted light, and 135° from the incident beam-the backscattered light). The turbiscan stability index (TSI), the sum of all the variations in the intensity of backscattering light at different locations of the test tube, could reflect the stability of emulsions. This could be calculated using Eq. (3) as following:^\text{41}

\[ TSI = \sum_{j} | \text{scan}_{j}(h_j) - \text{scan}_{j}(h_j) | \] (3)

where \text{scan}_{ref} and \text{scan}_j are the initial backscattering value and backscattering at a given time point, and h_j is a given height in the measuring cylindrical glass tube. In general, the higher the TSI value, the less stable the emulsion is.

2.6 Analysis of in vitro lipid digestion

This study used a modified form of a previously-described in vitro digestion model.\textsuperscript{42} The samples, containing 175 mg MCT, were placed in a glass beaker containing 30 mL buffer solution (10 mM Tris-HCl, pH 7.0) and incubated at 37°C for 15 min. Then, 2.5 mL of NaCh, 1.5 mL of CaCl\textsubscript{2} solution and 1 mL of NaCl solution were added to all the samples (except for control group to which NaCh was not added) with continuous stirring and the pH was adjusted to 7. Then, 1.5 mL of freshly prepared pancreatic lipase suspension was added to the mixture. The final digestion medium contained 175 mg lipid, 150 mM NaCl, 20 mM/L CaCl\textsubscript{2}, 5 mg/mL NaCh, and 1.6 mg/mL pancreatic lipase. A pH-stat automatic titration unit (916 Ti-Touch Metrohm, Switzerland) was used to automatically monitor the pH and maintain it at 7.0 by titrating appropriate amount of NaOH solution to neutralize the release of any free fatty acids (FFAs). The volume of NaOH added to the samples was recorded. The percentage of FFAs released was calculated using Eq. (4), as following:

\[ \text{FFA(%) = 100 \times \frac{V_{\text{NaOH}} \times m_{\text{NaOH}} \times M_{\text{NaOH}}}{W_{\text{fat}} \times 2}} \] (4)

where, \(V_{\text{NaOH}}\) is the volume of NaOH required for neutralization.
ing the FFA released (mL); $m_{NaOH}$ is the molarity of NaOH solution used (0.1 mol/L); $W_{lipid}$ is the total weight of oil present initially in the reaction vessel; and $M_{lipid}$ is the average molecular weight of MCT (509.44 g/mol).

2.7 Statistical analysis
All the experiments were performed in duplicates, with at least three measurements being made per sample.

3 Results and Discussion
3.1 Interfacial dilational rheology
3.1.1 Dynamic interfacial tension and dilational modulus
The dynamic IFTs and dilational moduli for the interfac-

![Dynamic IFTs and dilational moduli for the interfaces of pure oil-NaCh aqueous solution (A, B), lecithin oil solution-pure water (C, D), and lecithin oil solution-NaCh aqueous solution (E, F).](image)

es of pure oil-NaCh aqueous solution, lecithin oil solution-pure water, and lecithin oil solution-NaCh aqueous solution are presented in Fig. 1. Generally, when a new interface is created, the amphiphiles in the bulk phase start to adsorb at the interface, leading to the decrease in IFT. As the interfacial concentration of amphiphiles increases, the deformation in interface causes higher IFT gradient, leading to the increase in interfacial dilational modulus. When the amphiphiles reach an equilibrium adsorption, both the IFT and dilational modulus become constants. The dynamic IFTs and interfacial dilational moduli for the three systems in this study followed similar trends as that of general amphiphiles. At low concentrations, the diffusion rate of amphiphiles from the bulk to interface was low, so the IFT decreased slowly. With the increase in the concentration of

![Fig. 1 Dynamic IFTs and dilational moduli for the interfaces of pure oil-NaCh aqueous solution (A, B), lecithin oil solution-pure water (C, D), and lecithin oil solution-NaCh aqueous solution (E, F).](image)
amphiphiles, their diffusion also accelerated. For NaCh at higher concentrations ($\geq 1 \times 10^{-5}$ mol/L), IFTs and dilational moduli could quickly reach equilibrium values. But for lecithin at high concentration ($1 \times 10^{-5}$ mol/L $\sim 1 \times 10^{-3}$ mol/L), the IFTs still decreased slowly during the interfacial adsorption, indicating slower diffusion rate than that of NaCh. But lecithin could reduce the final IFT to a lower level than that of NaCh at the same concentrations. For the interface between the lecithin oil solution and the NaCh aqueous solution, the IFT could be further decreased while still maintaining a rapid adsorption rate.

3.1.2 Concentration dependence of interfacial tension and dilational modulus

The equilibrium IFTs and interfacial dilational moduli for the three systems at different concentrations are presented in Fig. 2. As shown in Fig. 2(A), the IFTs between lecithin oil solutions and pure water were lower than those of between pure oil and NaCh aqueous solutions at the same concentrations. This indicated that lecithin had higher interfacial activity. Moreover, Fig. 2(B) also shows that the dilational moduli for lecithin were also dramatically higher than those of NaCh.

As is known to all, NaCh exhibits planar polarity. Given this special structure, NaCh molecules tend to lie on the interface with the concave side facing the aqueous phase while the convex side faces the oil phase. Therefore, it is unlikely for the NaCh molecules to be closely packed and thereby exhibit weak intermolecular interactions. As to lecithin, the major component, phosphatidylcholine, is composed of a polar head-group containing phosphocholine and glycerol residue. Its polar group bears both positive and negative charges, which may generate strong electrostatic attraction. Moreover, the long hydrocarbon chains of phospholipids enhance van der Waals forces between hydrophobic groups. The high dilational modulus and low IFT indicated the formation of a compact arrangement of adsorption film under strong intermolecular interactions.

At the interface between lecithin oil solution and NaCh aqueous solution, both the lecithin and NaCh tended to adsorb to the interface. As shown in Fig. 2(A), the IFT dropped to lower values than individual lecithin and NaCh. For example, at the concentration of $1 \times 10^{-5}$ mol/L, the individual NaCh and lecithin decreased IFTs to 21.2 mN/m and 18.8 mN/m, respectively, while that of lecithin oil solution and NaCh aqueous solution decreased to 7.7 mN/m. This signified that NaCh was mixed adsorbed with lecithin at interface and they showed a good synergistic effect in decreasing IFT. Tung et al. used infrared spectroscopy for studying the interactions between bile salts and lecithin in oil phase. They found hydrogen bonding interactions between phosphate groups of lecithin and hydroxyl groups of NaCh played a significant role in the formation of reverse wormlike micelles. The hydrogen bonding interactions might also be a driving force for their assembly at the interface. Njauw et al. reported that the bile salts could be adsorbed onto phospholipid monolayers and decrease the IFT to lower level than that determined by the bile salts alone. This suggested a certain synergistic interaction of bile salts with the adsorbed-lipid layer.

The amount of amphiphiles adsorbed at interface increases with the increase in bulk concentration until it reaches critical micelle concentrations (CMC). Therefore, it enhances intermolecular interactions at the interface and causes higher IFT gradients during the interfacial compression and expansion. On the other hand, increase in the bulk concentration of amphiphiles also increases the molecular diffusion exchange between the bulk and interface, which can quickly reduce the IFT gradient and lead to a decrease in the dilational modulus. Generally, in the low concentration range, the increase in the amount of amphiphiles adsorbed at the interface is the decisive factor that determines the interfacial dilational modulus. In the high concentration range, the diffusion exchange of amphiphile molecules between the bulk and interface is enhanced and becomes dominant in determining the dilational modulus. Under these two opposite effects, the dilational modulus increases in the range of low concentration and decreases in the range of high concentration, and a maximal dilational

Fig. 2 Equilibrium IFTs (A) and interfacial dilational moduli (B) at the n-decane-aqueous interfaces as a function of concentration for NaCh, lecithin, and NaCh-lecithin mixed adsorption layers.
amphiphiles had generated an adsorption film with high dilational modulus. This unique variation indicated that, even the synergy between these two amphiphiles. However, with the increase in oscillation frequency, the interface deformation rate exceeds the characteristic frequencies of relaxation processes. Therefore, no relaxation processes occur and the dilational viscosity is also close to 0. In the moderate frequency range, the viscosity spectrum will exhibit maxima at the characteristic frequencies of the relaxation processes. As shown in Fig. 3(B), the dilational viscosity of individual NaCh increased monotonically with the increase in frequency, meaning that the maximum dilational viscosity would appear at a frequency higher than the highest oscillation frequency (0.1 Hz) used in this study. This indicated that there was a rapid relaxation process at or near the NaCh-adsorbed interfacial film, such as the diffusion exchange of amphiphilic molecules. For individual lecithin, as shown in Fig. 3(D), the dilational viscosity decreased gradually with the increase in oscillation frequency, meaning that the maximum dilational viscosity would appear at a frequency lower than the lowest oscillation frequency used in this study (0.005 Hz). This suggested that the slow relaxation processes, such as re-orientation or rearrangement, played a decisive role in determining the interfacial dilational properties. However, the mixed adsorption layer of lecithin with NaCh showed an increasing trend for dilational viscosity with the increase in oscillation frequency within the tested frequency range. This means that, like single NaCh adsorption layer, the characteristic frequency of dominate relaxation process at the mixed adsorption layer was higher than the highest oscillating frequency (0.1 Hz) used in the present work. As compared to the individual lecithin layer, the relaxation processes of the lecithin-NaCh composite interface have been accelerated significantly, which enhanced the interfacial viscoelasticity.

3.1.4 Interfacial dilational modulus as a function of interfacial pressure

As reported, for irreversibly adsorbed interfacial films, such as proteins and amphiphiles with strong intermolecular interactions, their interfacial properties are dominated by the amount and intermolecular interactions of the amphiphile molecules at the interface, which can be quantitatively expressed as their interfacial pressure. Therefore, the dilational modulus at various adsorption time and bulk...
concentrations coincides on a single dilational modulus-interfacial pressure curve. On the contrary, the interfacial properties of conventional surfactants are mainly dominated by the molecular diffusion exchange between the bulk and interface at high concentrations. Therefore, at a fixed interfacial pressure, higher the concentration of surfactant in bulk phase, faster will be the diffusion exchange of surfactant molecules between the bulk and interface. As a result, the IFT gradient caused by oscillation can be quickly reduced, leading to lower interfacial dilational modulus. That is, the dilational modulus-interfacial pressure curves of different concentrations are scattered instead of overlapping.

The dilational modulus vs interfacial pressure curves for individual NaCh and lecithin, and their interfacial mixture at various concentrations are presented in Fig. 4. As shown in Fig. 4(A), the curves of NaCh at different concentrations could be divided into two groups. At low NaCh concentrations ($\leq 1 \times 10^{-5}$ mol/L), these curves coincided with each other, indicating that the diffusion exchange of NaCh molecules between bulk and interface was inappreciable. However, with the increase of amphiphile concentration, the diffusion exchange became increasingly significant. When the NaCh concentration was higher than $1 \times 10^{-5}$ mol/L, the curves started to separate from being overlapped, suggesting that the diffusion exchange of amphi-

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**Fig. 3** Interfacial dilational elasticity and viscosity for NaCh (A, B), lecithin (C, D), and NaCh-lecithin mixed film (E, F) as a function of frequency.
philes between the interface and bulk solution began to dominate the nature of the adsorption layer. However, for the individual lecithin, as shown in Fig. 4(B), the dilational modulus vs interfacial curves were well-overlapped, indicating that the interfacial properties were dominated by the in-interface processes, while the diffusion exchange was negligible. The main relaxation processes of the lecithin interfacial films might only be the slow processes within the interface, such as the reorientation and rearrangement of amphiphile molecules, which are consistent with the low characterization frequencies as inferred from Fig. 3(D). As to the NaCh-lecithin mixed layers, the curves of dilational modulus vs. interfacial pressure at different concentrations were separated from each other, implying the fast relaxation process, diffusion exchange, has become the dominant factor, which further confirmed the speculation in Section 3.1.3.

### 3.2 Emulsion properties

In order to evaluate the emulsion properties, NaCh and lecithin were dissolved in water and MCT, respectively, and the aqueous and oil phases were mixed to prepare the NaCh-lecithin emulsions. The concentration of NaCh ranged from 0 to 1.5 wt%, while that of lecithin was fixed at 0.5 wt%. The emulsion stabilized only by NaCh was also prepared and used as a control. Figure 5(A) presents the effects of NaCh on the droplet size of the lecithin-stabilized emulsions. The size of emulsion droplets stabilized only with 0.5 wt% lecithin was around 600 nm. NaCh-stabilized emulsions had smaller droplets with the average size of around 160 nm at the same concentration. When NaCh and lecithin were introduced into water and oil phases separately to prepare emulsions, the droplets size was comparable to that of NaCh-stabilized emulsions. With the increase of NaCh concentration from 0.1 wt% to 1.5 wt%, a moderate decrease in the mean droplet size was observed. As discussed before, NaCh exhibited faster diffusion behavior from the bulk to interface. As a result, an adsorption layer was quickly formed on the interface during emulsification, thereby contributing to the better emulsifying ability of NaCh. Even in the presence of lecithin, NaCh played a leading role in the process of emulsification.

Figure 5(B) shows the destabilization kinetics of emulsions stabilized with the individual NaCh and lecithin, and their interfacial mixtures. Figure 6 shows the backscattering light intensity (BS) of lecithin and NaCh emulsion as a function of time and sample height. It can be seen from Fig. 6(A) that the BS curves kept almost unchanged with either the time or the sample height for the emulsion stabilized by lecithin. As discussed in 3.1, lecithin formed a
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Fig. 5 Oil droplets size (A) and stability of emulsion (B) stabilized with NaCh and/or lecithin.

Fig. 6 Variation of BS of O/W emulsions stabilized by (A) lecithin, (B) NaCh, and (C, D) NaCh-lecithin mixed layer.

tightly arranged film on the interface, thereby exhibiting better emulsion stability. On the contrary, due to the loose arrangement of NaCh molecules at the interface, the emulsion stabilized by NaCh was less stable as shown in Fig. 5 (B). Figure 6 (B) shows that the BS of NaCh emulsion decreased uniformly at different sample heights, indicating the increase of average droplet size due to the flocculation and coalescence of emulsion droplets. As to the emulsions stabilized by lecithin-NaCh mixtures shown in Figs. 6C and 6D, the BS still decreased uniformly for the entire sample height range, meaning that destabilization of emulsion was mainly due to droplet flocculation and coalescence. For the lecithin-NaCh mixed adsorbed layers, higher the concentration of NaCh in bulk, greater would be the influence of bile salts on the interface films and worse would be the stability of emulsions.

3.3 In vitro digestion fate of emulsions

The in vitro digestion fate of emulsions stabilized by lecithin was measured and the digestion kinetics is shown in Fig. 7. During the in vitro digestion in the absence of NaCh, the lecithin molecules covered the oil-aqueous interface and formed a compact adsorption film, which greatly hindered the molecular diffusion exchange between the bulk and the interface. Therefore, it blocked the contact of lipase with triacylglycerols. As a result, the digestion rate was dramatically slow and digestion extend was as low as 40 %. However, when NaCh was added during the digestion, due to the fast diffusion exchange of NaCh between bulk and interface, it quickly entered the interface and formed a mixed adsorption layer with lecithin. As shown in the results of interface dilatational rheology in Section 3.1, the viscoelasticity of interface increased, making the material exchange between the bulk and interface easier to take places. This created suitable conditions for the binding of lipases to lipid droplets. Therefore, the lipid digestion became faster and the digestion extend was significantly promoted.
layer to modulate lipid digestion. This would provide a basis for the rational design of interfacial behaviors, emulsion properties, and lipid digestion from the perspective of interfacial viscoelasticity and relaxation processes. NaCh and lecithin were selected as model compounds for bile salts and food emulsifiers, respectively. The interfacial dilational rheological behaviors, emulsion properties, and in vitro digestion fate of lipids were characterized. The experimental results showed that the lecithin alone formed a tightly-arranged elastic layer at the interface, thereby reducing the diffusion exchange of molecules between the bulk and interface. This interfacial feature helped to form a stable interfacial film, which enhanced the stability of emulsion and hindered the accessibility of lipases to lipids. Due to the special polar surface structure of NaCh, they could not form tightly-arranged layers at the interface. However, the diffusion exchange of NaCh molecules between bulk and interface was significant, suggesting that they could quickly enter the interface and replace the lecithin molecules. As a result, the lecithin interfacial adsorption layer, which was originally dominated by the slow relaxation processes within the interface, was transformed into one controlled by molecular diffusion exchange process. This transition accelerated the material exchange between the bulk and interface, thus creating suitable conditions for the interfacial adsorption of lipases, and promoting the digestion process. These results provided a deeper understanding of the interfacial behavior of bile salts with emulsifiers, which would provide a basis for the rational design of interfacial layer to modulate lipid digestion.

4 Conclusions

The current study focused on the role of bile salts in lipid digestion from the perspective of interfacial viscoelasticity and relaxation processes. NaCh and lecithin were selected as model compounds for bile salts and food emulsifiers, respectively. The interfacial dilational rheological behaviors, emulsion properties, and in vitro digestion fate of lipids were characterized. The experimental results showed that the lecithin alone formed a tightly-arranged elastic layer at the interface, thereby reducing the diffusion exchange of molecules between the bulk and interface. This interfacial feature helped to form a stable interfacial film, which enhanced the stability of emulsion and hindered the accessibility of lipases to lipids. Due to the special polar surface structure of NaCh, they could not form tightly-arranged layers at the interface. However, the diffusion exchange of NaCh molecules between bulk and interface was significant, suggesting that they could quickly enter the interface and replace the lecithin molecules. As a result, the lecithin interfacial adsorption layer, which was originally dominated by the slow relaxation processes within the interface, was transformed into one controlled by molecular diffusion exchange process. This transition accelerated the material exchange between the bulk and interface, thus creating suitable conditions for the interfacial adsorption of lipases, and promoting the digestion process. These results provided a deeper understanding of the interfacial behavior of bile salts with emulsifiers, which would provide a basis for the rational design of interfacial layer to modulate lipid digestion.

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