DYNAMIC PROCESSES IN SURFACTANT STABILIZED EMULSIONS

Authors: Krassimir D. Danov, Peter A. Kralchevsky, and Ivan B. Ivanov

Laboratory of Chemical Physics Engineering*, Faculty of Chemistry, University of Sofia, Sofia 1164, Bulgaria

TABLE OF CONTENTS

I. Introduction2
II. Dynamics of surfactant adsorption monolayers
 III. Mechanisms of coalescence
IV. Hydrodynamic interactions and drop coalescence
V. Interpretation of the Bancroft rule
 VI. Kinetics of coagulation in emulsions
VII. Summary

^{*} Formerly Laboratory of Thermodynamics and Physicochemical Hydrodynamics

I. INTRODUCTION

The process of emulsification usually takes place under essentially dynamic conditions. It is accompanied with the creation of new drops (new phase boundary) between the two liquids and with frequent collisions between the drops. Their instantaneous size distribution is a result of a competition between two oppositely directed processes: (i) breaking of the drops into smaller ones by the shear strain and (ii) coalescence of the newly formed drops into larger ones upon collision. If *surfactant* is present, it tends to adsorb at the surface of the drops and thus to protect them against coalescence. The rate of surfactant adsorption should be large enough to guarantee obtaining a sufficiently high coverage of the oil-water interface during the short period between two drop collisions. Therefore, an important parameter characterizing a given surfactant as emulsifier is its characteristic *adsorption time* τ_1 ; the latter can vary by many orders of magnitude depending on the type of surfactant, its concentration and the presence or absence of added non-amphiphilic electrolyte (salt) in the aqueous phase. In Section II.B we demonstrate how to quantify τ_1 for both ionic and nonionic surfactants.

The adsorbed surfactant molecules counteract the drop coalescence in two manners [1,2]. The presence of surfactant gives rise to repulsive surface forces (of either electrostatic, steric or oscillatory-structural origin) between the drops, thus providing a *thermodynamic* stabilization of the emulsion; see also Refs. [3,4]. Moreover, the adsorbed surfactant reduces (or completely removes) the tangential mobility of the drop surfaces and in this way markedly decelerates the interdroplet collisions; this is known as *kinetic* stabilization [1]. The latter is related to the Marangoni effect, i.e. to the appearance of gradients of adsorption and interfacial tension along the surfaces of two colliding droplets (see Fig. 1):



Fig. 1. Schematic presentation of the zone of contact of two approaching emulsion drops. The convective outflow of liquid from the gap between the drops drags the surfactant molecules along the two film surfaces; **j** and **j**_s denote the bulk and surface diffusion fluxes of surfactant.

$$\nabla_s \sigma = -\frac{E_G}{\Gamma_1} \nabla_s \Gamma_1 \tag{1}$$

here ∇_s is surface gradient operator, σ is surface tension, Γ_1 is surfactant adsorption and E_G is the Gibbs (surface) elasticity; expressions for estimating E_G can be found in Section II.A below.

In the case of low interfacial coverage with surfactant, the collision of two emulsion drops (step A \rightarrow B in Fig. 2) usually terminates with their coalescence (step B \rightarrow C in Fig. 2). The merging of the two drops occurs when a small critical distance between their surfaces, h_c is reached. Sometimes, depending on the specific conditions (larger drop size, attractive surface forces, smaller surface tension, etc. – see e.g. Ref. 2), the approach of the two drops could be accompanied with a deformation in the zone of their contact (step B \rightarrow D in Fig. 2); in this way a *liquid film* of almost uniform thickness h is formed in the contact zone. This film could also have a critical thickness h_c of rupture; in fact, the film rupture is equivalent to drop coalescence, see step D \rightarrow C in Fig. 2 The mechanisms of coalescence and the theoretical evaluation of h_c are considered in Section III.



Fig. 2. Possible consequences from a collision between two emulsion drops. Step A \rightarrow B: the two drops approach each other under the action of a driving force F; the viscous friction, accompanying the expulsion of liquid from the gap between the two drops, decelerates their approach. Step B \rightarrow C: after reaching a given *critical distance* between the two drop surfaces coalescence takes place. Step B \rightarrow D: after reaching a given threshold distance, h_{inv} , between the two drop surfaces, called the *inversion thickness*, the spherical drops deform and a film is formed in the zone of their contact. Step D \rightarrow C: The film, intervening between the two drops, thins and eventually breaks after reaching a certain *critical thickness*; then the two drops coalesce.

The driving force of the drop-drop collisions (**F** in Fig. 2) can be the Brownian stochastic force, the buoyancy force, some attractive surface force (say, the van der Waals interaction); in stirred vessels important role play the hydrodynamic (incl. turbulent) forces. The mutual approach of two emulsion drops (step $A \rightarrow B$ in Fig. 2) is decelerated by the *viscous friction* due to the expulsion of the liquid from the gap between the drops. If a doublet of two drops (Fig. 2D) is sufficiently stable, it can grow by attachment of additional drops; thus aggregates of drops (flocs) are produced.

If the stirring of an emulsion is ceased, there is no longer generation of new droplets, but the opposite processes of drop flocculation and/or coalescence continue. After some period of time this will lead to the appearance of sufficiently large flocs and/or drops, for which the gravitational force is stronger than the Brownian force; this will lead to a directional motion of the drops/flocs upwards (creaming) or downwards (sedimentation), depending on whether the buoyancy force or the drop weight prevails.



Fig. 3. Experimental data for creaming in xylene-in-water emulsions. The volume of the transparent "serum" left below the creaming emulsion, scaled with the total volume of the liquid mixture, is plotted against the time elapsed after ceasing the agitation. The emulsion is stabilized with β -lactoglobulin, whose concentrations, corresponding to the separate curves, are shown in the figure. The empty and full symbols denote, respectively, "coarse" emulsion (mean drop size 5 μ m) and "fine" emulsion (mean drop size 0.35 μ m).

As an illustrative example, Fig. 3 shows the occurrence of the creaming in an oil-inwater emulsion stabilized by the protein β -lactoglobulin – data from Ref. [5]. The rise of the boundary between the lower transparent aqueous phase (serum) and the upper turbid emulsion phase is recorded as a function of time; in particular, the ratio of the volume of the serum to the total volume (turbid + transparent phase) is plotted vs. time. Two types of emulsions are used in these experiments: "coarse" and "fine" emulsion of average drop size 5 μ m and 0.35 μ m, respectively. In Fig. 3 one sees, that in the *fine* emulsion creaming is not observed (the volume of the separated serum is identically zero). In contrast, there is creaming in the *coarse* emulsion, which starts some time after the initial moment (the ceasing of agitation); this period of time is necessary for "incubation" of sufficiently large flocs, which are able to emerge under the action of the buoyancy force. The stabilizing effect of β -lactoglobulin is manifested as an increase of the "incubation period" with the rise of protein concentration.

The theoretical description of the mutual approach and coalescence of two emulsion drops is subject of Section IV; the Bancroft rule about emulsification is interpreted and generalized in Section V; the kinetics of flocculation is considered in Section VI, where the size of the aggregates needed for the creaming to start is estimated.

II. DYNAMICS OF SURFACTANT ADSORPTION MONOLAYERS

A. Gibbs (Surface) Elasticity

1. Nonionic surfactant solutions

Let us consider the boundary between an aqueous solution of a nonionic surfactant and the oil phase. We choose the dividing surface to be the equimolecular dividing surface with respect to water. Then the Gibbs adsorption equation takes the form [6,7]

$$d\sigma = -kT\Gamma_1 d\ln c_1 \tag{2}$$

where the subscript "1" denotes the nonionic surfactant: c_1 and Γ_1 are its bulk concentration and adsorption; *k* is the Boltzmann constant and *T* is temperature. The surfactant adsorption isotherms, expressing the connection between Γ_1 and c_1 are usually obtained by means of some molecular model of the adsorption. The most popular is the Langmuir [8] adsorption isotherm,

$$\frac{\Gamma_1}{\Gamma_\infty} = \frac{Kc_1}{1 + Kc_1} \tag{3}$$

which stems from a lattice model of localized adsorption of *non-interacting* molecules [9]. In Eq. (3) Γ_{∞} is the maximum possible value of the adsorption ($\Gamma_1 \rightarrow \Gamma_{\infty}$ for $c_1 \rightarrow \infty$). On the other hand, for $c_1 \rightarrow 0$ one has $\Gamma_1 \approx Kc_1$; the adsorption parameter *K* characterizes the surface activity of the surfactant: the greater *K* the higher the surface activity.

Table 1.	The	most	popular	surfactant	adsorption	isotherms	and t	he 1	respective	surface	tension
isotherms	c_{1s}	is the	subsurfa	ace concen	tration of th	ne surfactar	nt mol	lecu	les.		

	Surfactant adsorption isotherms			
	(for nonionic surfactants: $c_{1s} \equiv c_1$)			
Henry	$Kc_{1s} = \frac{\Gamma_1}{\Gamma_{\infty}}$			
Freundlich	$Kc_{1s} = \left(\frac{\Gamma_1}{\Gamma_F}\right)^{1/m}$			
Langmuir	$Kc_{1s} = \frac{\Gamma_1}{\Gamma_{\infty} - \Gamma_1}$			
Volmer	$Kc_{1s} = \frac{\Gamma_1}{\Gamma_{\infty} - \Gamma_1} \exp\left(\frac{\Gamma_1}{\Gamma_{\infty} - \Gamma_1}\right)$			
Frumkin	$Kc_{1s} = \frac{\Gamma_1}{\Gamma_\infty - \Gamma_1} \exp\left(-\frac{2\beta\Gamma_1}{kT}\right)$			
van der Waals	$Kc_{1s} = \frac{\Gamma_1}{\Gamma_{\infty} - \Gamma_1} \exp\left(\frac{\Gamma_1}{\Gamma_{\infty} - \Gamma_1} - \frac{2\beta\Gamma_1}{kT}\right)$			
	• Surface tension isotherm $\sigma = \sigma_0 - kTJ + \sigma_d$ (for nonionic surfactants: $\sigma_d \equiv 0$)			
Henry	$J = \Gamma_1$			
Freundlich	$J = \frac{\Gamma_1}{m}$			
Langmuir	$J = -\Gamma_{\infty} \ln \left(1 - \frac{\Gamma_1}{\Gamma_{\infty}} \right)$			
Volmer	$J = \frac{\Gamma_{\infty}\Gamma_1}{\Gamma_{\infty} - \Gamma_1}$			
Frumkin	$J = -\Gamma_{\infty} \ln \left(1 - \frac{\Gamma_1}{\Gamma_{\infty}} \right) - \frac{\beta \Gamma_1^2}{kT}$			
van der Waals	$J = \frac{\Gamma_{\infty}\Gamma_1}{\Gamma_{\infty} - \Gamma_1} - \frac{\beta\Gamma_1^2}{kT}$			

Table 1 contains the 6 most popular surfactant adsorption isotherms, those of Henry, Freundlich, Langmuir, Volmer [10], Frumkin [11], and van der Waals [9]. For $c_1 \rightarrow 0$ all other isotherms (except that of Freundlich) reduce to the Henry isotherm. The physical difference between the Langmuir and Volmer isotherms is that the former corresponds to a physical model of *localized* adsorption, whereas the latter – to *non-localized* adsorption. The Frumkin and van der Walls isotherms generalize, respectively, the Langmuir and Volmer isotherms for the case, when there is interaction between the adsorbed molecules; β is a parameter, which accounts for the interaction. In the case of van der Waals interaction β can be expressed in the form [12,13]:

$$\beta = -\pi kT \int_{r_0}^{\infty} \left[1 - \exp\left(-\frac{u(r)}{kT}\right) \right] \approx -\pi \int_{r_0}^{\infty} u(r) r dr$$
(4)

where u(r) is the interaction energy between two adsorbed molecules and r_0 is the distance between the centers of the molecules at close contact. The comparison between theory and experiment shows that the interaction parameter β is important for air-water interfaces, whereas for oil-water interfaces one can set $\beta = 0$ [14,15]. The latter fact, and the finding that $\beta > 0$ for air-water interfaces, leads to the conclusion that β takes into account the van der Waals attraction between the hydrocarbon tails of the adsorbed surfactant molecules across air (such attraction is missing when the hydrophobic phase is oil). Note, however, that even for oilwater interface one could have $\beta < 0$ if some non-electrostatic repulsion between the adsorbed surfactant molecules takes place, say steric repulsion between some chain branches of amphiphilic molecules with more complicated structure.

What concerns the parameter *K* in Table 1, it is related to the standard free energy of adsorption, $\Delta f = \mu_1^{(0)} - \mu_{1s}^{(0)}$, which is the energy gain for bringing a molecule from the bulk of the water phase to a diluted adsorption layer [3,16]:

$$K = \frac{\delta_1}{\Gamma_{\infty}} \exp\left(\frac{\mu_1^{(0)} - \mu_{1s}^{(0)}}{kT}\right)$$
(5)

Here δ_1 is a parameter, characterizing the thickness of the adsorption layer, which can be set (approximately) equal to the length of the amphiphilic molecule. Let us consider the integral

$$J \equiv \int_{0}^{c_{1}} \Gamma_{1} \frac{dc_{1}}{c_{1}} = \int_{0}^{\Gamma_{1}} \Gamma_{1} \frac{d\ln c_{1}}{d\Gamma_{1}} d\Gamma_{1}$$
(6)

The derivative $d \ln c_1 / d\Gamma_1$ can be calculated for each adsorption isotherm in Table 1 and then the integration in Eq. (6) can be carried out analytically [17]. The expressions for J thus obtained are also listed in Table 1. The integration of the Gibbs adsorption isotherm, Eq. (2), along with Eq. (6), yields [17]

$$\sigma = \sigma_0 - kTJ \,, \tag{7}$$

which in view of the expressions for J in Table 1 presents the surfactant adsorption isotherm, or the two-dimensional (surface) equation of state.

As mentioned in the Introduction, an important thermodynamic parameter of a surfactant adsorption monolayer is its Gibbs (surface) elasticity. The physical concept of surface elasticity is the most transparent for monolayers of *insoluble* surfactants, for which it has been initially introduced by Gibbs [18,19]. The increments $\Delta \sigma$ and $\Delta \Gamma_1$ in the definition of Gibbs elasticity,

Table 2. Expressions for the Gibbs elasticity of adsorption monolayers (valid for both nonionic and ionic surfactants), which correspond to the various types of isotherms in Table 1.

Type of surface tension isotherm	Gibbs elasticity E_G
Henry	$E_G = kT\Gamma_1$
Freundlich	$E_G = kT \frac{\Gamma_1}{m}$
Langmuir	$E_G = kT\Gamma_1 \frac{\Gamma_{\infty}}{\Gamma_{\infty} - \Gamma_1}$
Volmer	$E_G = kT\Gamma_1 \frac{\Gamma_{\infty}^2}{\left(\Gamma_{\infty} - \Gamma_1\right)^2}$
Frumkin	$E_{G} = kT\Gamma_{1} \left(\frac{\Gamma_{\infty}}{\Gamma_{\infty} - \Gamma_{1}} - \frac{2\beta\Gamma_{1}}{kT} \right)$
van der Waals	$E_G = kT\Gamma_1 \left[\frac{\Gamma_{\infty}^2}{\left(\Gamma_{\infty} - \Gamma_1\right)^2} - \frac{2\beta\Gamma_1}{kT} \right]$

$$E_G = -\Gamma_1 \lim_{\Delta \Gamma_1 \to 0} \frac{\Delta \sigma}{\Delta \Gamma_1} = -\Gamma_1 \left(\frac{\partial \sigma}{\partial \Gamma_1} \right)_T$$
(8)

correspond to variations in surface tension and adsorption during a real process of interfacial dilatation. Expressions for E_G , corresponding to various adsorption isotherms, are shown in Table 2. As an example, let us consider the expression for E_G , corresponding to the Langmuir isotherm; combining results from Tables 1 and 2 one obtains

$$E_G = \Gamma_{\infty} kTKc_1$$
 (for Langmuir isotherm) (9)

One sees that for Langmuirian adsorption the Gibbs elasticity grows linearly with the surfactant concentration c_1 . Since the concentration of the monomeric surfactant cannot exceed the critical micellization concentration, $c_1 \le c_{\text{CMC}}$, then from Eq. (9) one obtains

$$E_G \le \left(E_G\right)_{\max} = \Gamma_{\infty} kTKc_{CMC} \qquad (\text{for Langmuir isotherm}) \tag{10}$$

Hence one could expect higher elasticity E_G for surfactants with higher c_{CMC} ; this conclusion is consonant with the experimental results [20].

The Gibbs elasticity characterizes the lateral fluidity of the surfactant adsorption monolayer. For high values of the Gibbs elasticity the adsorption monolayer at a fluid interface behaves as tangentially immobile. Then, if two oil drops approach each other, the hydrodynamic flow pattern, and the hydrodynamic interaction as well, is the same as if the drops were solid particles, with the only difference that under some conditions they could deform in the zone of contact. For lower values of the Gibbs elasticity the Marangoni effect appears, see Eq. (1), which can considerably affect the approach of the two drops. These aspects of the hydrodynamic interactions between emulsion drops are considered in Section IV below.

In the case of a soluble nonionic surfactant the detected increase of σ in a real process of interfacial dilatation can be a pure manifestation of surface elasticity only if the period of dilatation, Δt , is much shorter than the characteristic *relaxation time of surface tension* τ_{σ} , $\Delta t \ll \tau_{\sigma}$ [21]. Otherwise the adsorption and the surface tension would be affected by the diffusion supply of surfactant molecules from the bulk of solution toward the expanding interface. The diffusion transport tends to reduce the increase of surface tension upon dilatation, thus apparently rendering the interface less elastic and more fluid. The initial condition for the problem of adsorption kinetics involves an "instantaneous" ($\Delta t \ll \tau_{\sigma}$) dilatation of the interface. This "instantaneous" dilatation decreases the adsorptions Γ_i and the subsurface concentrations c_{is} of the species (the subsurface is presumed to be always in equilibrium with the surface), but the bulk concentrations $c_{i\infty}$ remain unaffected [22-24]. This initially created difference between c_{is} and $c_{i\infty}$ further triggers the diffusion process. Now, let us inspect closer how this approach is to be extended to the case of ionic surfactants.

2. Ionic surfactant solutions

The thermodynamics of adsorption of *ionic* surfactants is more complicated due to the presence of long-range electrostatic interactions in the system. Let us consider a boundary between two immiscible fluid phases (say, water and oil), which bears some electric charge owing to the adsorption of charged amphiphilic molecules (ionic surfactant). The charged surface repels the *coions*, i.e. the ions having charge of the same sign, but it attracts the *counterions*, which bear charge of the opposite sign (Fig. 4). Thus an electric double layer (EDL) appears, that is a non-uniform distribution of the ionic species in the vicinity of the charged interface [25].



Fig. 4. Electric double layer formed in a vicinity of an adsorption monolayer of ionic surfactant. The diffuse layer contains free ions involved in Brownian motion, while the Stern layer consists of adsorbed (immobilized) counterions. Near the charged surface there is an accumulation of counterions and a depletion of coions.

The conventional model of the EDL stems from the works of Gouy [26], Chapman [27] and Stern [28]. According to this model the EDL consists of two parts: (i) *adsorption layer* and (ii) *diffuse layer*, see Fig. 4. The adsorption layer includes surfactant molecules, which are

immobilized (adsorbed) at the phase boundary, as well as bound counterions, which form the Stern layer. The diffuse layer consists of free ions in the aqueous phase, which are involved in Brownian motion and influenced by the electric field of the charged interface. The boundary, separating the adsorption from the diffuse layer, called the *Gouy plane*, can be used as a Gibbs dividing surface between the two neighboring phases [15]. The electric potential varies across the EDL: $\psi = \psi(x)$. The boundary values of $\psi(x)$ are $\psi(x=0) = \psi_s$ at the Gouy plane (at the interface) and $\psi(x\to\infty) = 0$ in the bulk of solution. At equilibrium, the subsurface concentrations of the ionic species, c_{is} , are related to the respective bulk concentrations, $c_{i\infty}$, by means of the Boltzmann distribution [25]:

$$c_{is} \equiv c_i \ (x=0) = c_{i\infty} \exp\left(-\frac{z_i e \psi_s}{kT}\right),\tag{11}$$

i = 1, 2, 3, ..., N. Here *e* is the electronic charge, z_i is the valency of the *i*-th ion. The Gibbs adsorption equation can be presented in the form [15,17,29-31]:

$$d\sigma = -kT \sum_{i=1}^{N} \widetilde{\Gamma}_{i} d \ln c_{i\infty} \qquad (T = \text{const})$$
(12)

Equations (11) and (12) are rigorous in terms of activities of the ionic species, rather than in terms of concentrations. For simplicity, here we set the activities equal to the concentrations, which is a good approximation for ionic strengths below 0.1 M; see Refs. [14,15,17] for details. In Eq. (12) $\tilde{\Gamma}_i$ denotes the adsorption of the *i*-th component; $\tilde{\Gamma}_i$ represents the surface excess of component "*i*" with respect to the *uniform* bulk solution. For an ionic species $\tilde{\Gamma}_i$ is a total adsorption, which includes contributions Γ_i and Λ_i , respectively from the adsorption layer (adsorbed surfactant + counterions in the Stern layer) and the diffuse layer, which are defined as follows [17,29-31]

$$\Lambda_{i} \equiv \int_{0}^{\infty} [c_{i}(x) - c_{i\infty}] dx , \qquad \Gamma_{i} \equiv \widetilde{\Gamma}_{i} - \Lambda_{i}$$
(13)

Using the theory of EDL and Eq. (13) one can prove that the Gibbs adsorption equation, Eq. (12), can be represented in the following equivalent form [17]

$$d\sigma_a = -kT \sum_{i=1}^{N} \Gamma_i d \ln c_{is} \qquad (T = \text{const})$$
(14)

where $\sigma_a = \sigma - \sigma_d = \sigma_0 - kTJ$ is the contribution of the *adsorption* layer into the surface tension (*J* is the same as in Eq. 6 and Table 1), and σ_d is the contribution of the *diffuse* layer [17,29]:

$$\sigma_d = -\frac{\varepsilon}{4\pi} \int_0^\infty \left(\frac{\partial \psi}{\partial x}\right)^2 dx \tag{15}$$

 ε is the dielectric permittivity of the aqueous phase. The integrand in Eq. (15) represents the anisotropy of the Maxwell electric stress tensor, which contributes to the interfacial tension in accordance with the known Bakker formula [32-34]. The comparison between Eqs. (12) and (14) shows that the Gibbs adsorption equation can be expressed either in terms of σ , $\tilde{\Gamma}_i$ and $c_{i\infty}$, or in terms of σ_a , Γ_i and c_{is} . The total surface tension is

$$\sigma = \sigma_a + \sigma_d \tag{16}$$

Note that σ_d represents a non-local, integral contribution of the whole diffuse electric double layer, whereas σ_a is related to the two-dimensional state of the adsorbed surfactant ions and bound counterions (Fig. 4).

Let us consider a solution of ionic surfactant, which is a symmetric *z*:*z* electrolyte, in the presence of additional non-amphiphilic *z*:*z* electrolyte (salt); here $z \equiv z_1 = -z_2 = z_3$. We assume that the counterions due to the surfactant and salt are identical. For example, this can be a solution of sodium dodecyl sulfate (SDS) in the presence of NaCl. We denote by $c_{1\infty}$, $c_{2\infty}$ and $c_{3\infty}$ the bulk concentrations of the surface active ions (1), counterions (2), and coions (3), respectively. For the special system of SDS with NaCl $c_{1\infty}$, $c_{2\infty}$ and $c_{3\infty}$ are the bulk concentration of the DS⁻, Na⁺ and Cl⁻ ions, respectively. The requirement for the bulk solution to be electroneutral implies $c_{2\infty} = c_{1\infty} + c_{3\infty}$. The binding of *coions* due to the non-amphiphilic salt is expected to be equal to zero, $\Gamma_3 = 0$, because they are repelled by the similarly charged interface [17]. However, $\Lambda_3 \neq 0$; hence $\widetilde{\Gamma}_3 = \Lambda_3 \neq 0$. The difference between the adsorptions of surfactant ions and counterions determines the surface charge density, $\rho_s = ez(\Gamma_1 - \Gamma_2)$. For the considered system Eq. (11) can be presented in the form

$$c_{is} = c_{i\infty} \exp[(-1)^i \phi_s], \qquad \qquad \phi_s \equiv \frac{z e \psi_s}{kT}, \qquad (17)$$

(i = 1, 2, 3). Note that the dimensionless surface electric potential ϕ_s thus defined is always positive, irrespective of whether the surfactant is cationic or anionic.

Let us proceed with the definition of Gibbs elasticity for an adsorption monolayers from *ionic* surfactant. The main question is whether or not the electric field in the EDL should be affected by the "instantaneous" dilatation of the interface, $-\Delta\Gamma_1$, which is involved in the definition of E_G , see Eq. (8). This problem has been examined in Ref. 35 and it has been established that a variation of the electric field during the initial instantaneous dilatation leads to results, which are unacceptable from a theoretical viewpoint. The latter conclusion is related to the following two facts: (i) The speed of propagation of the electric signals is much greater than the characteristic rate of diffusion. (ii) Even a small initial variation in the surface charge density ρ_s immediately gives rise to an electric potential, which is linearly increasing with the distance from the interface (potential of a planar wall). Consequently, a small initial perturbation of the interface would immediately affect the ions in the *whole* solution; of course, such an initial condition is physically unacceptable. In reality, a linearly growing electric field could not appear in an ionic solution, because a variation of the surface charge density would be immediately suppressed by exchange of counterions, which are abundant in the subsurface layer of the solution. The theoretical equations suggest the same [35]: to have a mathematically meaningful initial condition of *small* perturbation for the diffusion problem, the initial dilatation must be carried out at constant surface charge density ρ_s ; for details see the Appendix in Ref. [35]. Thus the following conclusion has been reached: the initial sudden interfacial dilatation, which is related to the definition of Gibbs elasticity of a soluble ionic surfactant, must be carried out at $\rho_s = \text{const.}$ From Eq. (16) one obtains [36]

$$(d\sigma)_{\rho_s} = (d\sigma_a)_{\rho_s} + (d\sigma_d)_{\rho_s}$$
(18)

An interfacial dilatation at constant ρ_s does not alter the diffuse part of the EDL, and consequently, $(d\sigma_d)_{\rho_s} \equiv 0$, see Eq. (15). Since [17],

$$\sigma_a = \sigma_0 - kTJ , \qquad (19)$$

the expressions for *J* in Table 1 show that σ_a depends only on Γ_1 at constant temperature. Then the definition of Gibbs elasticity of nonionic adsorption layers can be extended to ionic adsorption layers in the following way [36]:

$$E_G \equiv -\Gamma_1 \left(\frac{\partial \sigma}{\partial \Gamma_1}\right)_{T,\rho_s} = -\Gamma_1 \left(\frac{\partial \sigma_a}{\partial \Gamma_1}\right)_T \qquad (\text{ionic surfactant}) \qquad (20)$$

The definition of Gibbs elasticity given by Eq. (19) corresponds to an "instantaneous" $(\Delta t \ll \tau_{\sigma})$ dilatation of the *adsorption* layer (that contributes to σ_a) without affecting the *diffuse* layer and σ_d . The dependence of σ on Γ_1 for nonionic surfactants is the same as the dependence of σ_a on Γ_1 for ionic surfactants, cf. Eqs. (7) and (19). Then Eqs. (8) and (20) show that the expressions for E_G in Table 2 are valid for *both* nonionic and ionic surfactants. The effect of the surface electric potential on the Gibbs elasticity E_G of an ionic adsorption monolayer is implicit, through the equilibrium surfactant adsorption Γ_1 , which depends on the electric properties of the interface. To illustrate this let us consider the case of Langmuir adsorption isotherm for an *ionic* surfactant [17]

$$Kc_{1s} = \frac{\Gamma_1}{\Gamma_{\infty} - \Gamma_1}, \qquad \qquad K \equiv K_1 + K_2 c_{2s}$$
(21)

where K_1 and K_2 are constants. Note that the above linear dependence of the adsorption parameter K on the subsurface concentration of counterions, c_{2s} , can be deduced from the equilibrium exchange reactions, which describe the adsorption of surfactant ions and counterions, see Ref. [37]. Combining the respective expression from Table 2 with Eq. (21) we obtain $E_G = \Gamma_{\infty} kTKc_{1s}$. Further, having in mind that $K \equiv K_1 + K_2 c_{2s}$, we substitute Eq. (17) to derive

$$E_G = \Gamma_{\infty} kT c_{1\infty} [K_1 \exp(-\phi_s) + K_2 c_{2\infty}]$$
 (for Langmuir isotherm) (22)

Equation (22) reveals the effect of salt on E_G : when the salt concentration increases, $c_{2\infty}$ also increases, whereas the (dimensionless) surface potential ϕ_s decreases, see Fig 5 in Ref. 17; then Eq. (22) predicts an increase of E_G with the rise of salt concentration.

A numerical illustration of the latter prediction is given in Fig. 5 below. The Gibbs elasticity is calculated with the help of Eq. (22), i.e. the Langmuir isotherm, using the values of K_1 , K_2 and Γ_{∞} determined in Ref. [17] from the fit of experimental data due to Tajima et al [38,39] for sodium dodecyl sulfate (SDS). The surface potential ϕ_s is computed as a function of the surfactant and salt concentrations using steps 2–6 of the calculation procedure described in Section 9.2 of Ref. [17] with $\beta = 0$. As seen in Fig. 5, E_G increases with the rise of surfactant (SDS) concentration. Moreover, for a fixed surfactant concentration one observes a strong

increase of E_G with the increase of NaCl concentration. To understand this behavior of E_G we notice that according to Table 2 E_G depends explicitly only on Γ_1 at fixed temperature *T*. Hence the influence of surfactant and salt on the Gibbs elasticity E_G can be interpreted as an increase of the surfactant adsorption Γ_1 with the rise of both surfactant and salt concentrations.



Fig. 5. Plot of the Gibbs (surface) elasticity E_G vs. the surfactant (SDS) concentration, $c_{1\infty}$. The four curves correspond to four fixed NaCl concentrations: 0, 20, 50 and 115 mM; E_G is calculated by means of Eq. (22) using parameters values determined from the best fit of experimental data in Ref. 17.

B. Characteristic Time of Adsorption

1. Nonionic surfactant solutions

The characteristic time of surfactant adsorption at a fluid interface is an important parameter for surfactant-stabilized dynamic systems like emulsions. Sutherland [22] derived an expression describing the relaxation of a *small* dilatation of an initially equilibrium adsorption monolayer from soluble *nonionic* surfactant (diffusion control):

$$\frac{\sigma(t) - \sigma^{(e)}}{\sigma(0) - \sigma^{(e)}} = \frac{\Gamma_1(t) - \Gamma_1^{(e)}}{\Gamma_1(0) - \Gamma_1^{(e)}} = \exp\left(\frac{t}{\tau_1}\right) \operatorname{erfc}\left(\sqrt{\frac{t}{\tau_1}}\right)$$
(23)

where *t* is time,

$$\tau_1 = \frac{1}{D_1} \left(\frac{\partial \Gamma_1}{\partial c_1} \right)^2 \tag{24}$$

is characteristic relaxation time and D_1 is surfactant diffusivity; here and hereafter the superscript "(*e*)" denotes the equilibrium value of the respective parameter; erfc(*x*) is the complementary error function [40-42]. Using the asymptotics of the latter function for x >> 1 one obtains

$$\frac{\sigma(t) - \sigma^{(e)}}{\sigma(0) - \sigma^{(e)}} = \frac{\Gamma_1(t) - \Gamma_1^{(e)}}{\Gamma_1(0) - \Gamma_1^{(e)}} = \sqrt{\frac{\tau_1}{\pi t}} + O(t^{-3/2}) \qquad (t >> \tau_1, \text{ small perturbation})$$
(25)

Equation (25) is often used as a test to verify whether the adsorption process is under diffusion control: data for the dynamic surface tension $\sigma(t)$ are plotted vs. $1/\sqrt{t}$ and it is checked if the plot complies with a straight line; the extrapolation of this line to $1/\sqrt{t} \rightarrow 0$ is used to determine the equilibrium surface tension $\sigma^{(e)}$ [23,43].

In the experiment one often deals with *large* initial deviations from equilibrium; for example, such is the case when new oil-water interface is formed by breaking of larger emulsion drops during emulsification. In the case of large perturbation there is no general analytical expression for the dynamic surface tension $\sigma(t)$ since the adsorption isotherms (except that of Henry, see Table 1) are nonlinear. In this case one can use either computer solution [44,45] or to apply the von Karman's approximate approach [46,47]. Analytical asymptotic expressions for the long time ($t \gg \tau_1$) relaxation of surface tension of a *nonionic* surfactant solution was obtained by Hansen [48]:

$$\sigma(t) - \sigma^{(e)} = \left(\Gamma_1^{(e)}\right)^2 \frac{kT}{c_1^{(e)}} \sqrt{\frac{1}{\pi D_1 t}} \qquad (t \to \infty); \tag{26}$$

When deriving Eq. (26), the surfactant adsorption at the initial moment was set zero, $\Gamma_1(0) = 0$. Equation (26) has been verified, utilized and generalized by many authors [24,49-53]. With the help of Eqs. (2), (8) and (24) one can represent Eq. (26) in the following equivalent form

$$\sigma(t) - \sigma^{(e)} = E_G \sqrt{\frac{\tau_1}{\pi t}} + O(t^{-3/2}) \qquad (t \gg \tau_1, \text{ large perturbation}) \qquad (27)$$

where, as usual, E_G denotes Gibbs elasticity. The comparison of Eqs. (25) and (27) shows that the relaxation of surface tension is characterized by the *same* relaxation time τ_1 , irrespective of whether the interfacial perturbation is large or small. (The same conclusion is valid also for ionic surfactants, see below). For that reason the relaxation time can be considered as a general kinetic property of the adsorption monolayer [36].

2. Ionic surfactant solutions

In the case of *ionic* surfactants the existence of a diffuse electric double layer essentially influences the kinetics of adsorption. The process of adsorption is accompanied with a progressive increase of the surface charge density and electric potential. The charged surface repels the new-coming surfactant molecules which results in a deceleration of the adsorption process [54]. The theoretical studies on dynamics of adsorption encounter difficulties with the non-linear set of partial differential equations, which describes the electro-diffusion process [55].

Another important effect, which adds to the complexity of the problem, is the *adsorption (binding) of counterions* at the conversely charged surfactant headgroups in the adsorption layer, see Fig. 4. The adsorbed (bound) counterions form the Stern layer, which affects strongly the adsorption kinetics of ionic surfactants insofar as up to 70-90% of the surface electric charge could be neutralized by the bound counterions [17,56-58]. The addition of non-amphiphilic electrolyte (salt) in the solution increases the occupancy of the Stern layer. It turns out that in the case of ionic surfactants (with or without salt) there are two adsorbing species: the surfactant ions and the counterions. The adsorption of counterions can be described by means of the Stern isotherm [6,17,28]. It is worthwhile noting that the counterion binding enhances the adsorption of surfactant [17]; formally, this appears as a linear increase of the surfactant adsorption parameter *K* with the rise of the subsurface concentration of counterions, c_{2s} , see Eq. (21).

In a recent articles [35,36] the problem about the kinetics of adsorption from an ionic surfactant solution was addressed in its full complexity, including the time evolution of the electric double layer, the effect of added salt and the counterion binding. Analytical solution was found only in the asymptotic cases of *small* and *large* initial deviations from equilibrium and *long* times of adsorption. Thus generalizations of Eqs. (25) and (27) for the case of ionic surfactants was obtained (see below). An interesting result is that the electrostatic interaction leads to the appearance of three distinct characteristic relaxation times, those of surfactant adsorption τ_1 , of counterion adsorption (binding) τ_2 and of surface tension relaxation τ_{σ} . In

particular, the relaxation of surfactant and counterion adsorptions, Γ_1 and Γ_2 , under electrodiffusion control, is described by the equation

$$\frac{\Gamma_i(t) - \Gamma_i^{(e)}}{\Gamma_1(0) - \Gamma_1^{(e)}} = \sqrt{\frac{\tau_i}{\pi t}} + O(t^{-3/2}) \qquad (t \to \infty, i = 1, 2)$$
(28)

where τ_1 and τ_2 are given by a generalized version of Eq. (24), which can be found in Refs. [35,36] together with the procedure of calculations. The relaxation of interfacial tension of *ionic* surfactant solutions is given again by Eqs. (25) and (27), in which τ_1 is to be replaced by τ_{σ} defined as follows [36]:

$$\sqrt{\tau_{\sigma}} = (1+w)\sqrt{\tau_1} - w\sqrt{\tau_2} - \frac{2qw\lambda}{\kappa} \tanh\frac{\phi_s^{(e)}}{4}$$
(29)

where κ is the Debye parameter,

$$\kappa^{2} \equiv \frac{4\pi z^{2} e^{2} \sum_{i} c_{i\infty}}{\varepsilon kT};$$
(30)

$$w = \frac{2kT\Gamma_{1}^{(e)}}{E_{G}} \tanh \frac{\phi_{s}^{(e)}}{2}, \qquad \lambda = \frac{1 + \zeta\Gamma_{2}^{(e)}/\Gamma_{1}^{(e)}}{1 + \zeta}, \qquad \zeta = \exp(-\phi_{s}^{(e)}/2)$$
(31)

The latter expression for the parameter λ corresponds to the case of large perturbations; for *small* perturbations one simply has $\lambda \equiv 1$ [36]. The computations show that in the case of large perturbations λ is close to 1, and therefore the relaxation time is not sensitive to the magnitude of perturbation.

As an illustration, in Fig. 6 we show the calculated dependence of the relaxation times τ_1 , τ_2 and τ_{σ} on the surfactant concentration. As in Fig. 5, we have used the values of K_1 , K_2 and Γ_{∞} determined in Ref. [17] from the fit of experimental data due to Tajima et al [38,39] for sodium dodecyl sulfate (SDS). All necessary equations and the procedure of calculation are described in Ref. [36] for the case of large perturbations. The range of surfactant and salt concentrations correspond to the *non-micellar* surfactant solutions studied experimentally in Refs. [38,39]. In Figs. 6a and 6b one notices the wide range of variation of the relaxation times, which is from 2 to 6 orders of magnitude. For example, the relaxation time of surface tension, τ_{σ} , drops from about 40 s for 10^{-5} M SDS concentration down to $\approx 4 \times 10^{-5}$ s for 10^{-3} M SDS. (see Fig. 6a). In addition, one sees that systematically $\tau_2 < \tau_1 < \tau_{\sigma}$; the difference between these three relaxation times can be greater than one order of magnitude for the lower surfactant concentrations, especially in the case without added electrolyte (Fig. 6b). One can conclude that

the terms proportional to w in Eq. (29), which give rise to the difference between τ_1 and τ_{σ} , play an important role, particularly for solutions of lower ionic strength. Figure 6 demonstrates that the approximation $\tau_{\sigma} \approx \tau_1$, which is widely used in literature, is applicable only for the higher surfactant concentrations, for which $\tau_{\sigma} \rightarrow \tau_1$. Note also that for a given surfactant concentration t_2 is always smaller than τ_1 and τ_{σ} , that is the adsorption of counterions relaxes faster than does the adsorption of surfactant ions and the surface tension.



Fig. 6. Ionic surfactant solution: relaxation times of interfacial tension, τ_{σ} , of surfactant adsorption, τ_1 , and of counterion adsorption (binding), τ_2 , calculated in Ref. [36] as functions of surfactant (SDS) concentration, $c_{1\infty}$, using parameters values determined from the best fit of experimental data in Ref. [17]. (a) SDS solutions with 115 mM added NaCl; (b) SDS solutions without added NaCl.

The physical importance of these results is related to the fact, that the coalescence of drops at the early highly-dynamic stages of emulsion production is expected to be sensitive to the degree of saturation of the newly-created interfaces with surfactant, and correspondingly, to the relaxation time of surfactant adsorption. The surfactant transport is especially important when the emulsion is prepared from non-preequilibrated liquid phases. In such a cases one can observe dynamic phenomena like the *cyclic dimpling* [59,60] and *osmotic swelling* [61], which bring about additional stabilization of the emulsions, see also Refs. [1, 62].

3. Micellar surfactant solutions

Emulsions are often prepared from micellar surfactant solutions. As known, above a given critical micellization concentration (CMC) surfactant aggregates (micelles) appear inside the surfactant solutions. At rest the micelles exist in equilibrium with the surfactants monomers in the solution. If the concentration of the monomers in the solution is suddenly decreased, the micelles release monomers until the equilibrium concentration, equal to CMC, is restored at the cost of disassembly of a part of the micelles [63,64].

The dilatation of the surfactant adsorption layer leads to a transfer of monomers from the subsurface to the surface, which causes a transient decrease of the subsurface concentration of monomers. The latter is compensated by disintegration of a part of the micelles in the subsurface layer. This process is accompanied by a diffusion transport of surfactant monomers and micelles due to the appearance of concentration gradients. In general, the micelles serve as a powerful source of monomers which is able to quickly saturate the surface of the newly created emulsion drops. In this way, the presence of surfactant micelles strongly accelerates the kinetics of adsorption.

The theoretical model by Anianson and Wall [65-68] describes the micelles as polydisperse aggregates, whose growth or decay happens by exchange of monomers. The general theoretical description of the diffusion in such a solution of polydisperse aggregates taking part in chemical reactions (exchange of monomers) is a heavy task; nevertheless, it has been addressed in several works [69-72]. The relaxation of surface tension of a micellar solution at small initial deviation from equilibrium can be described by the following expression derived in Ref. [70]:

$$\frac{\sigma(t) - \sigma^{(e)}}{\sigma(0) - \sigma^{(e)}} = \frac{\Gamma_1(t) - \Gamma_1^{(e)}}{\Gamma_1(0) - \Gamma_1^{(e)}} = \frac{1}{g_1 - g_2} \left[E(g_1, \tau) - E(g_2, \tau) \right] \exp\left(-\frac{t}{\tau_m}\right)$$
(32)

where $E(g,\tau) = g \exp(g^2 \tau) \operatorname{erfc}(g\sqrt{\tau})$, $\tau = t/\tau_d$, $g_{1,2} = [1 \pm \sqrt{1 + 4\tau_d / \tau_m}]/2$,

$$\tau_m = \left[K_d \left(1 + m^2 c_m^{(e)} / c_1^{(e)} \right) \right]^{-1}, \quad \tau_d = \left(\partial \Gamma_1 / \partial c_1 \right)^2 / D_1.$$
(33)

 τ_m and τ_d are the characteristic relaxation times of micellization and monomer diffusion, see Ref. [73]; for the sake of estimates τ_d can be identified with τ_1 as given by Eq. (24); K_d is rate constant of micelle decay; as earlier, the index "*e*" refers to the equilibrium state and *m* is the average micelle aggregation number. In the absence of micelles $\tau_d / \tau_m \rightarrow 0$; then $g_1 = 1$, $g_2 = 0$, and Eq. (32) reduces to Eq. (23), as it should be expected. One can estimate the characteristic time of relaxation in the presence of micelles by using the following combined expression:

$$\tau_{\sigma} \approx \frac{4\tau_d}{\left(1 + \sqrt{1 + 4\tau_d / \tau_m}\right)^2} \tag{34}$$

According to the latter expression $\tau_{\sigma} \approx \tau_m$ for $\tau_d \gg \tau_m$, and $\tau_{\sigma} \approx \tau_d$ for $\tau_d \ll \tau_m$.

Equation (32) is applicable only for small perturbations. An approximate analytical approach, which is applicable for both small and large deviations from equilibrium, is developed in Ref. [47].

III. MECHANISMS OF COALESCENCE

A. Mechanisms of Rupture of Emulsion Films

1. Thermodynamic and kinetic factors preventing coalescence

Often the contact of two emulsion drops is accompanied with the formation of a liquid film between them. The rupture of this film is equivalent to coalescence of the drops, that is step D \rightarrow C in Fig 2. Figure 7 shows schematically the zone of contact of two emulsion drops of different radii, R_1 and R_2 ($R_1 < R_2$). For the sake of simplicity we assume that the two drops are made of the same liquid and have the same surface tension σ . The film formed in the contact zone has radius R and thickness h. The interaction of the two drops across the film leads to the appearance of additional *disjoining pressure* Π inside the film, which in general depends on the film thickness: $\Pi = \Pi(h)$, see e.g. Refs. [2-4, 62]. Positive Π corresponds to repulsion between the two film surfaces (and the two drops), whereas negative Π corresponds to attraction between them. The presence of disjoining pressure gives rise to a difference between the tension of the film surfaces, σ^{f} , and the interfacial tension σ of the droplets. The force balance at the contact line reads [62,74,75]

$$\sigma^{f} = \sigma \cos \alpha \,; \tag{35}$$

 α is contact angle, which is related to the disjoining pressure Π as follows [62,76]

$$\cos\alpha = 1 + \frac{1}{2\sigma} \int_{h}^{\infty} \Pi(h) dh$$
(36)

Since $\cos \alpha < 1$, a necessary condition to have contact angle is the integral in Eq. (36) to be negative; for emulsion drops this can be ensured by the long-range van der Waals attraction, which dominates Π for the larger *h*, see Fig. 8a. Geometrically α appears as the angle subtended between the tangents to the film and drop surfaces at the contact line (Fig. 7). At *equilibrium* (no applied external force) the radius of the film between the two drops is determined by the equation

$$R = \frac{2\sigma \sin \alpha}{\Pi} \cos \theta \approx \frac{2\sigma \sin \alpha}{\Pi} ; \qquad \qquad \cos \theta \equiv \left(1 - R^2 / R_f^2\right)^{1/2} \approx 1 \qquad (37)$$

where R_f is the curvature radius of the film. Equation (37) follows from Eqs. (144) and (179) in Ref. [75]. One sees that the greater the contact angle α , the larger the equilibrium film radius R. On the other hand, for $\alpha = 0$ the equilibrium film radius R is also zero and there are no equilibrium doublets or larger aggregates (flocs) of emulsion droplets.



Fig. 7. Sketch of a film between two non-identical emulsion drops of radii R_1 and R_2 . The film thickness and radius are denoted by *h* and *R*, respectively; α is contact angle, P_1 , P_2 and P_3 denote the pressure in the respective liquid phases. If the two drops have different radii, as it is in Fig. 7, the film between them is curved. The balance of the pressures applied per unit area of the two film surfaces can be expressed by means of versions of the Laplace equation [75]

$$\frac{2\sigma^{f}}{R_{f}} = P_{1} - (P_{3} + \Pi) \qquad \text{(upper film surface)} \tag{38}$$
$$\frac{2\sigma^{f}}{R_{f}} = (P_{3} + \Pi) - P_{2} \qquad \text{(lower film surface)} \tag{39}$$

where P_1 and P_2 are the pressures inside the respective drops (Fig. 7); P_3 is the pressure in the continuous phase; the effect of disjoining pressure is equivalent to an increase of the pressure within the film, which is $P_3 + \Pi$. To obtain Eqs. (38) and (39) we have neglected some very small terms, of the order of h/R_f , see Ref. [75] for details. To determine R_f we apply the Laplace equations for the two drops (Fig. 7):

$$\frac{2\sigma}{R_1} = P_1 - P_3, \qquad \frac{2\sigma}{R_2} = P_2 - P_3$$
(40)

Combining Eqs. (36) and (38)–(40) one determines the curvature radius of the film:

$$\frac{1}{R_f} = \frac{1}{2\cos\alpha} \left(\frac{1}{R_1} - \frac{1}{R_2} \right)$$
(41)

If the two drops have identical size $(R_1 = R_2)$, then Eq. (41) yields $1/R_f \rightarrow 0$, i.e. the film between the drops is flat, as it should be expected.

The disjoining pressure Π is the major *thermodynamic* stabilizing factor against drop coalescence. Stable equilibrium state of a liquid film can exist if only the following two conditions are satisfied [3]:

$$\Pi = P_{\rm A}$$
 and $\left(\frac{\partial \Pi}{\partial h}\right)_{\Pi = P_{\rm A}} < 0$ (42)

Here P_A is the pressure difference applied across the surface of the film, which in view of Eqs. (38) and (39) can be expressed in the form

$$P_{A} = P_{1} - P_{3} - \frac{2\sigma^{f}}{R_{f}} = \frac{2\sigma^{f}}{R_{f}} + P_{2} - P_{3} = \sigma \left(\frac{1}{R_{1}} + \frac{1}{R_{2}}\right)$$
(43)

At the last step we have used also Eqs. (35), (40) and (41). For two identical drops $R_f \rightarrow \infty$, and then P_A reduces to the capillary pressure of the drops: $P_A = 2\sigma/R_1 = 2\sigma/R_2$. The condition $\Pi = P_A$, see Eq. (42), means that at equilibrium the disjoining pressure Π counterbalances the pressure difference P_A applied across the film surface. In addition, the condition $\partial \Pi/\partial h < 0$ guarantees that the equilibrium is *stable* (rather than unstable).



(a) Electrostatic DLVO Barrier

(b) Multiple Barriers Due to Micelles

Fig. 8. Typical plots of disjoining pressure Π vs. film thickness *h*; *P*_A is the pressure difference applied across the film surface, see Eq. (43); the equilibrium states of the liquid film correspond to the points in which $\Pi = P_A$. (a) DLVO-type disjoining pressure isotherm $\Pi(h)$; the points at $h = h_1$ and h_2 correspond to primary and secondary films, respectively; Π_{max} is the height of a barrier due to the electrostatic repulsion between the film surfaces. (b) Oscillatory structural force between the two film surfaces caused by the presence of surfactant micelles (or other monodisperse colloidal particles) in the continuous phase; $\Pi(h)$ exhibits multiple decaying oscillations; the stable equilibrium films with thickness h_0 , h_1 , h_2 and h_3 correspond to stratifying films containing 0, 1, 2 and 3 layers of micelles, respectively, see Fig. 9.

As an illustration, Fig. 8a shows a typical DLVO-type disjoining pressure isotherm $\Pi(h)$, see Refs. [3,4,62] for more details. There are two points, $h = h_1$ and $h = h_2$, at which the condition for stable equilibrium, Eq. (42), is satisfied. In particular, $h = h_1$ corresponds to the so called *primary film*, which is stabilized by the electrostatic (double layer) repulsion. The addition of electrolyte to the solution may lead to a decrease of the height of the electrostatic barrier, Π_{max} [3,4]; at high electrolyte concentration it is possible to have $\Pi_{\text{max}} < P_A$; then primary film does not exist. Note, however, that the increase of electrolyte concentration may

lead also to a shift of the maximum toward smaller thicknesses and to an increase of the barrier Π_{max} . Therefore, primary films could be observed even at relatively high salt concentrations.

The equilibrium state at $h = h_2$ (Fig. 8a) corresponds to a very thin *secondary film*, which is stabilized by the short-range Born repulsion. The secondary film represents a bilayer of two adjacent surfactant monolayers; its thickness is usually about 5 nm (slightly greater than the doubled length of the surfactant molecule) [77]. Secondary films can be observed in emulsion flocs and in creamed emulsions.



Fig. 9. The spots of lower thickness in stratifying liquid films correspond to a local decrease of the number of micelle layers in the colloid-crystal-like structure of surfactant micelles formed inside the liquid film. The appearance of spots could be attributed to the condensation of vacancies in that structure [82].

The situation is more complicated when the aqueous solution contains surfactant micelles, which is a common experimental and practical situation. In such a case the disjoining pressure isotherm $\Pi(h)$ can exhibit multiple decaying oscillations, whose period is close to the diameter of the micelles (Fig. 8b); for details see e.g. Ref. [78]. The condition for equilibrium liquid film, Eq. (42), can be satisfied in several points, denoted by h_0 , h_1 , h_2 and h_3 in Fig. 8b; the corresponding films contain 0, 1, 2 and 3 layers of micelles, respectively. The transitions between these multiple equilibrium states represent the phenomenon *stratification*, see Fig. 9 and Refs. [78-91]. The presence of disjoining pressure barriers, which are due either to the electrostatic repulsion (Fig. 8a), or to the oscillatory structural forces (Fig. 8b), has a stabilizing effect on liquid films and emulsions [2].

The existence of a stable equilibrium state (see Fig. 8) does not guarantee that a draining liquid film can "safely" reach this state. Indeed, hydrodynamic instabilities, accompanying the drainage of liquid, could rupture the film before it has reached its thermodynamic equilibrium state [1]. There are several *kinetic stabilizing factors*, which suppress the hydrodynamic instabilities and decelerate the drainage of the film, thus increasing its lifetime. Such a factor is the *Gibbs (surface) elasticity*, E_G , of the surfactant adsorption

monolayers, see Section II.A above; it tends to eliminate the gradients in adsorption and surface tension and damps the fluctuation capillary waves. At higher surfactant and salt concentrations the Gibbs elasticity is also higher and it renders the interface tangentially immobile, see Fig. 5. The *surface viscosity* also impedes the drainage of water out of the films because of the dissipation of a part of the kinetic energy of the flow within the surfactant adsorption monolayers (see Section IV below). The *surfactant adsorption relaxation time* (see Section II.B) is another important kinetic factor. If the adsorption relaxation time is small enough, a dense adsorption monolayer will cover the newly formed emulsion drops during the emulsification and will protect them against coalescence upon collision. In the opposite case (slow adsorption kinetics) the drops can merge upon collision and the emulsion will be rather unstable.

2. Mechanism of film breakage

The role of the emulsion stabilizing (or destabilizing) factors can be understood if the mechanism of film breakage is known. Several different mechanisms of rupture of liquid films have been proposed, which are briefly described below.

The *capillary-wave mechanism* has been proposed by de Vries [92] and extended in subsequent studies [93-98, 2, 24], see Fig. 10a. The conventional version of this mechanism is developed for the case of monotonic attraction between the two surfaces of a liquid film (say, van der Waals attraction). Thermally excited fluctuation capillary waves are always present at the film surfaces. With the decrease of the average film thickness, h, the attractive disjoining pressure enhances the amplitude of some modes of the fluctuation waves. At a given critical value of the film thickness, h_c , corrugations on the two opposite film surfaces can touch each other and then the film will break [97]. The same mechanism takes place also in the case of slightly deformed emulsion drops. If the emulsion drops are quiescent, only the thermodynamic and geometric factors determine the critical thickness; indeed the finite area of the drops (films) imposes limitation on the maximum length of the capillary waves (see Sections III.B and III.C). When the breakage happens during the drainage of the emulsion film (during the approach of the emulsion drops), then the critical thickness is also affected by various hydrodynamic factors (see Section IV for details).

The mechanism of *film rupture by nucleation of pores* has been proposed by Derjaguin & Gutop [99] to explain the breaking of very thin films, built up from two attached monolayers

of amphiphilic molecules. Such are the secondary foam and emulsion films and the bilayer lipid membranes. This mechanism was further developed by Derjaguin & Prokhorov [100,101,3], Kashchiev & Exerowa [102-104], Chizmadzhev et al. [105-107], and Kabalnov & Wennerström [108]. The formation of a nucleus of a pore (Fig. 10b) is favored by the decrease of the surface energy, but it is opposed by the edge energy of the pore periphery. The edge energy can be described (macroscopically) as a line tension [100-104], or (micromechanically) as an effect of the spontaneous curvature and bending elasticity of the amphiphilic monolayer [108]. For small nuclei the edge energy is predominant, whereas for larger nuclei the surface energy gets the upper hand. Consequently, the energy of pore nucleation exhibits a maximum at a given *critical* pore size; the larger pores spontaneously grow and break the film, while the smaller pores shrink and disappear.



Fig. 10. Mechanisms of breakage of liquid films. (a) *Fluctuation-wave mechanism*: the film rupture results from growth of capillary waves enhanced by attractive surface forces [92]. (b) *Pore-nucleation mechanism*: it is expected to be operative in very thin films, virtually representing two attached monolayers of amphiphilic molecules [99]. (c) *Solute-transport mechanism*: if a solute is transferred across the two surfaces of the liquid film due to gradients in solute chemical potential, then Marangoni instability could appear and break the film [109].

A third mechanism of liquid-film breakage is observed when there is a *transport of solute across the film*, see Fig. 10c. This mechanism, investigated experimentally and theoretically by Ivanov et al. [109-111], was observed with emulsion systems (transfer of alcohols, acetic acid and acetone across liquid films), but it could appear also in some asymmetric oil-water-air films. The diffusion transport of some solute across the film leads to the development of Marangoni instability, which manifests itself as a forced growth of capillary waves at the film surfaces and eventual film rupture. Note that Marangoni instability can be caused by both mass and heat transfer [112-114].

A fourth mechanism of film rupture is the *barrier mechanism*. It is directly related to the physical interpretation of the equilibrium states in Fig. 8. For example, let us consider an electrostatically stabilized film of thickness h_1 (Fig. 8a). Some processes in the system may lead to the increase of the applied capillary pressure P_A . For instance, if the height of the column of an emulsion cream increases from 1 cm to 10 cm, the capillary pressure in the upper part of the cream increases from 98 Pa up to 980 Pa owing to the hydrostatic effect. Thus P_A could become greater than the height of the barrier, Π_{max} , which would cause either film rupture (and coalescence), or transition to the stable state of secondary film at $h = h_2$ (Fig. 8a). The increase of the surfactant adsorption density stabilizes the secondary films. In addition, the decrease of Π_{max} decreases the probability the film to rupture after the barrier is overcome. Indeed, the overcoming of the barrier is accompanied with a violent release of mechanical energy accumulated during the increase of P_A . If the barrier is not too high, the film could survive the transition.

The overcoming of the barrier can be facilitated by various factors. Often the transition happens trough the formation and expansion of spots of lower thickness within the film, rather than by a sudden decrease of the thickness of the *whole* film. Physically this is accomplished by a nucleation of spots of sub- μ m size, which resembles a transition with a "tunnel effect", rather than a real overcoming of a barrier. A theoretical model of spot formation in stratifying films by condensation of vacancies in the structure of ordered micelles (vacancy mechanism), has been developed in Ref. [82], see Figs. 8b and 9. The nucleation of spots makes the transitions less violent and decreases the probability for film breakage. The increase of the applied capillary pressure P_A facilitates the spot formation and the transition to state with lower film thickness; this has been established by Bergeron and Radke [85], who experimentally obtained

portions of the stable branches of the oscillatory disjoining-pressure curve (Fig. 8b) for foam films. Oscillatory disjoining pressure curves due to reverse micelles in an oily phase were directly measured by Parker et al. [86] by using a version of the surface force apparatus. Marinova et al. [91] investigated the stabilizing role of the oscillatory disjoining pressure in oil-in-water emulsions which contain surfactant micelles in the aqueous phase.

Below we present in more details the predictions of the capillary-wave mechanism.

B. Critical Thickness of Quiescent Emulsion Films

Let us first consider a quiescent emulsion film, say the film between two drops within a floc or cream. At a given sufficiently small thickness of the film, termed the *critical thickness* [92-97,115], the attractive surface forces prevails and causes growth of the thermally-excited capillary waves. This may lead to either film rupture or transition to thinner secondary film. Two modes of film undulations have been distinguished: symmetric (squeezing, peristaltic) and antisymmetric (bending) modes; it is the symmetric mode which is related with the film breakage/transition. The critical thickness, $h = h_c$, of a film having area πR^2 is the value of h, can be estimated from the equation [94]:

$$\frac{2R^2}{\sigma} \left(\frac{\partial \Pi}{\partial h} \right) = j_1^2 = 5.783..., \tag{44}$$

where j_1 is the first zero of the Bessel function J_0 ; as usual, σ denotes surface tension. Equation (44) has been derived in a different manner by Vrij [94], Ivanov *et al.* [95], and Malhotra & Wasan [116]. It is obvious that Eq. (44) can be satisfied only for positive $\partial \Pi / \partial h$. In the special case of van der Waals interaction one is to substitute $\partial \Pi / \partial h = A_H / (2\pi h^4)$, where A_H is the Hamaker constant; then from Eq. (44) it follows that the critical thickness increases with the increase of the film radius R, i.e. the films of larger area break easier (at a greater thickness) than the films of smaller area. Note that the effect of surfactant on the tangential mobility of the interface, which involves the surface elasticity, viscosity and diffusion, does not affect the form of Eq. (44), and correspondingly, the critical thickness h_c . The surfactant affects Eq. (44) and h_c only indirectly, through the values of σ and $\partial \Pi / \partial h$. These conclusions are valid only for quiescent films, which do not thin during the development of instability.

When an aqueous film is stabilized by an ionic surfactant, then the stability problem becomes more complicated due to the electrostatic interactions between the charged film surfaces [117]. Electrolyte films surrounded by dielectric were initially studied by Felderhof [118], who examined the stability of an equilibrium infinite plane-parallel film surrounded by vacuum. Sche and Fujnaut [119] extended the Felderhof's analysis to account for the effect of surface shear viscosity and surface elasticity. In these works the electrostatic (double-layer) component of disjoining pressure Π was involved in the theory and a quasi-static approximation was employed to describe the electrostatic interaction [117-119]. In other words, it has been assumed that the ions immediately acquire their equilibrium distribution for each instantaneous shape of the film. The electric field has been computed by solving the Poisson-Boltzmann equation for the respective instantaneous charge configuration. This quasi-static approximation, which neglects the electro-diffusion fluxes, leads to a counterpart of Eq. (44) in which the total disjoining pressure Π includes an electrostatic component. The latter leads to $\partial \Pi / \partial h < 0$ at the equilibrium state ($h = h_1$ in Fig. 8a) and then Eq. (44) has no positive root for $h = h_c$; that is the film should remain stable for infinitely long time in agreement with the conventional DLVO theory [3]. On the other hand, if electrolyte is added at sufficiently high concentration, the double-layer repulsion is suppressed and the liquid films rupture under the action of the van der Waals force, see Ref. [120] and Eq. (86) below.

In reality, aqueous films stabilized with ionic surfactant, without electrolyte, also do rupture, especially at surfactant concentrations below the CMC. The latter fact cannot be explained in the framework of the quasistatic approximation [117-119]; this is still an opened problem in the theory of liquid-film stability.

C. Critical Distance between Quiescent Emulsion Drops

Let us consider two emulsion drops of different radii, R_1 and R_2 , like those depicted in Fig. 7 but without the formation of a film between them, i.e. R = 0. In this case the gap between the two drops represents a liquid film of uneven thickness. The frequently used lubrication approximation [121] is not applicable to description of the fluctuation capillary waves on the drop surfaces because this approximation presumes infinite interfacial area and does not impose the natural upper limits on the capillary wavelength, originating from the finite size of the drops. On the other hand, it is possible to solve the problem by means of the usual spherical coordinates, locating the coordinate origin in the center of one of the two drops. We consider the case in which effects of surface electric charge are negligible and the interaction between the drops (the disjoining pressure) is dominated by the van der Waals attraction. The critical distance between the two drops can be determined from a thermodynamic requirement, viz. the fluctuation of the local disjoining pressure in the narrowest zone of the gap to be equal to the fluctuation of the capillary pressure of the drops. (For shorter distances the fluctuation of the attractive disjoining pressure will prevail and will initiate film rupture.) This requirement leads to the following equation:



Fig. 11. Plot of the critical distance between two quiescent drops, h_c , vs. the mean drop radius, *a*, calculated by means of Eq. (48) for three values of the Hamaker constant $A_{\rm H}$.

$$\frac{1}{\sin\theta} \frac{d}{d\theta} \left(\sin\theta \frac{d\zeta}{d\theta} \right) + \left\{ 2 + \frac{A_{\rm H}a^2\cos^2\theta}{\pi\sigma} \left[h + 2a(1-\cos\theta) \right]^{-4} \right\} \zeta = 0 , \qquad (45)$$

Here ζ is the fluctuation in the drop shape, θ is the polar angle of the spherical coordinate system, *h* is the shortest distance between the two drop surfaces, $A_{\rm H}$ is the Hamaker constant and

$$a = 2R_1 R_2 / (R_1 + R_2) \tag{46}$$

is the mean drop radius. We used the following two boundary conditions: (i) $d\zeta/d\theta = 0$ at $\theta = 0$, i.e. at the narrowest region of the gap; (ii) $\zeta = 0$ for $\theta = \pi/2$, that is far from the gap zone. The value $h = h_c$, corresponding to the *greatest* eigenvalue of the spectral problem, Eq. (45), gives the critical distance between the two drops. Note that the effect of surfactant on the tangential mobility of the interface, which involves the surface elasticity, viscosity and diffusion, does not affect the form of Eq. (45), and correspondingly, the critical distance h_c . We found the greatest eigenvalue numerically. The results for the critical distance as a function of the drop radius a and the Hamaker constant $A_{\rm H}$ are shown in Fig. 11; for the interfacial tension we used the value $\sigma = 30$ mN/m. One sees that the critical distance is of the order of dozens of nanometers and that it increases with the rise of both $A_{\rm H}$ and a.

Note however, that if the two drops are not quiescent, but instead approach each other, the critical distance is influenced by the hydrodynamic interactions – see the next Section.

IV. HYDRODYNAMIC INTERACTIONS AND DROP COALESCENCE

First, we consider the hydrodynamic interactions between two emulsion drops, which remain spherical when the distance between them decreases (Section IV.A); This is the transition $A \rightarrow B$ in Fig. 2. Second, we consider the thinning of the film formed between two emulsion drops (Section IV.B): this is stage D in Fig. 2. In both cases the effect of surfactant is taken into account and the critical distance (thickness) for drop coalescence is quantified.

A. Interaction of Spherical Emulsion Drops

1. Limiting cases of low and high surface mobility

The solution of the problem about the hydrodynamic interaction between two rigid spherical particles, approaching each other across a viscous fluid, was obtained by Taylor [122]. Two spherical emulsion drops of *tangentially immobile* surfaces (due to the presence of dense surfactant adsorption monolayers) are hydrodynamically equivalent to the two rigid particles considered by Taylor. The hydrodynamic interaction is due to the dissipation of kinetic energy when the liquid is expelled from the gap between the two spheres. The resulting friction force decreases the velocity of the two spherical drops proportionally to the decrease of the surface-to-surface distance h in accordance with the equation [122]

$$V_{\rm Ta} = \frac{2h}{3\pi\eta a^2} (F - F_s) \qquad (\text{tangentially immobile surfaces}) \qquad (47)$$

Here *a* is the mean drop radius defined by Eq. (46); *F* is the external force exerted on each drop, F_s is the surface force originating from the intermolecular interactions between the two drops across the liquid medium. When the range of the latter interactions is much smaller than the drop radii, then F_s can be calculated by means of the Derjaguin approximation [3,4]

$$F_s = \pi a \int_h^\infty \Pi(h) dh \tag{48}$$

where, as before, Π is disjoining pressure.

If the surface of an emulsion drop is mobile, it can transmit the motion of the outer fluid to the fluid within the drop. This leads to a circulation of the fluid inside the drop and influences the dissipation of energy in the system. The problem about the approach of two nondeformed spherical drops or bubbles in the *absence of surfactants* has been investigated by many authors [123-132]. A number of solutions, generalizing the Taylor equation (47), have been obtained. In particular, the velocity of central approach of two spherical drops in pure liquid, $V_{\rm p}$, is related to the Taylor velocity $V_{\rm Ta}$, defined by means of a Padé-type expression derived by Davis *et al.* [131]

$$V_{\rm p} = V_{\rm Ta} \frac{1 + 1.711\xi + 0.461\xi^2}{1 + 0.402\xi}, \qquad (\text{no surfactant})$$
(49)

$$\xi = \frac{\eta_{\text{out}}}{\eta_{\text{in}}} \sqrt{\frac{a}{2h}}$$
(50)

where, as usual, *h* is the closest surface-to-surface distance between the two drops; η_{in} and η_{out} are the viscosities of the liquids inside and outside the drops. In the limiting case of solid particles one has $\eta_{in} \rightarrow \infty$, $\xi \rightarrow 0$ and then Eq. (49) reduces to the Taylor equation, Eq. (47). Note that in the case of *close* approach of two drops ($h \rightarrow 0$, $\xi >> 1$) the velocity V_p is proportional to \sqrt{h} . This implies that the two drops can come into contact (h = 0) in a finite period of time ($\tau < \infty$) under the action of a given force, *F*, because the integral expressing the lifetime [97]

$$\tau = \int_{h_c}^{h_{in}} \frac{dh}{V}$$
(51)

(with $V = V_p$) is convergent for $h_c = 0$; h_{in} is the surface-to-surface distance at the initial moment t = 0. In contrast, in the case of immobile interfaces ($\xi << 1$) Eq. (47) gives $V_{Ta} \propto h$ and $\tau \rightarrow \infty$ for $h_c \rightarrow 0$. Moreover, the counterbalancing of the external force by the surface force, i.e. $F - F_s = 0$, implies $V_{Ta} = V = 0$ and $\tau \rightarrow \infty$ (equilibrium state) irrespective of whether the drop surfaces are tangentially mobile or immobile.

It has been established both theoretically and experimentally [133,134] that if the surfactant is dissolved only in the drop phase, the film formed between two emulsion drops (Fig. 2D) thins just as if surfactant is missing. Likewise, one can use Eq. (49) to estimate the velocity of approach of two emulsion drops when surfactant is contained only in the drop phase [2].

2. Effects of surface elasticity, viscosity and diffusivity

When surfactant is present in the continuous phase at not too high concentration, then the surfactant adsorption monolayers, covering the emulsion drops, are tangentially mobile, rather than immobile. The adsorbed surfactant can be dragged along by the fluid flow in the gap between two colliding drops thus affecting the hydrodynamic interaction between them. The appearance of gradients of surfactant adsorption are opposed by the Gibbs elasticity, surface viscosity, surface and bulk diffusion. Below we consider the role of the enumerated factors on the velocity of approach of two emulsion drops.

If the driving force *F* (say the Brownian or the buoyancy force) is small compared to the capillary pressure of the droplets, the deformation of two spherical droplets upon collision will be only a small perturbation in the zone of contact. Then the film thickness and the pressure within the gap can be presented as a sum of a non-perturbed part and a small perturbation. Solving the resulting hydrodynamic problem for low (negligible) interfacial viscosity, an analytical formula for the velocity of drop approaching, V = -dh / dt, can be derived [121]:

$$\frac{V}{V_{\text{Ta}}} = \frac{h_s}{2h} \left[\frac{d/h+1}{d/h} \ln(d/h+1) - 1 \right]^{-1},$$
(52)

where *a* is the mean drop radius defined by Eq. (46); V_{Ta} is the Taylor velocity, Eq. (47); the other parameters are defined as follows:

$$d = \frac{h_s}{1+b},\tag{53}$$

$$h_s = \frac{6\eta_{\text{out}} D_{1,s}}{E_G},\tag{54}$$

$$b = \frac{3\eta_{\text{out}}D_1}{E_G} \left(\frac{\partial c_1}{\partial \Gamma_1}\right)^{(e)} = \frac{3\eta_{\text{out}}D_1}{\Gamma_1^{(e)}} \left(\frac{\partial c_1}{\partial \sigma}\right)^{(e)}$$
(55)

As usual, the subscript (*e*) denotes that the respective quantity should be estimated for the equilibrium state; the dimensionless parameter *b* accounts for the effect of *bulk diffusion*, whereas h_s has a dimension of length and takes into account the effect *surface diffusion*. In the limiting case of very large *Gibbs elasticity* E_G (tangentially immobile interface) the parameter *d* tends to zero and then Eq. (52) yields $V \rightarrow V_{Ta}$, as it should be expected [121,135,136].

If the effect of *surface viscosity* is taken into account, then Eq. (52) can be expressed in the generalized form [137,138]:

$$\frac{V}{V_{\rm Ta}} = \Phi_{\rm v}(h; h_s, d, s_{\rm v}) \tag{56}$$

where Φ_v is termed the *mobility factor (function*); the dimensionless parameter s_v takes into account the effect of surface viscosity:

$$s_{\rm v} \equiv \frac{\eta_s}{6a\eta_{\rm out}} , \qquad \eta_s \equiv \eta_{\rm sh} + \eta_{\rm dil}$$
(57)

Here η_{sh} and η_{dil} are, respectively, the interfacial shear and dilatational viscosities. In fact, Eq. (52) gives an analytical expression for the mobility factor Φ_v in the case when $s_v \ll 1$, i.e. the effect of surface viscosity can be neglected. However, if the effect of surface viscosity is essential, there is no analytical expression for Φ_v ; in this case a numerical procedure for computation of Φ_v has been developed [137,138]. Table 3 contains asymptotic expressions for Φ_v . A general property of Φ_v is

$$\Phi_{\rm v} = 1$$
 for $E_G \to \infty$ (tangentially immobile surfaces) (58)

It is important to note that the surface viscosity parameter s_v appears only in the combinations $s_v h_s / h = \eta_s D_{1s} / (hE_G a)$ and $s_v b$, see Table 3. Then, in view of Eqs. (53)-(55), it follows that the surface viscosity can influence the mobility factor Φ_v only if either the Gibbs elasticity, E_G , or the drop radius, a, or the gap width, h, are small enough.

To illustrate the dependence of the mobility function Φ_v on the concentration of the surfactant in the continuous phase, in Fig. 12 we present theoretical curves calculated in Ref. [138] for the nonionic surfactant Triton X100, for the ionic surfactant sodium dodecyl sulfate (SDS) + 0.1 M NaCl, and for the protein Bovine serum albumin (BSA). The parameters values, used to calculated the curves in Fig. 12, are listed in Table 4. Γ_{∞} and *K* are parameters of the Langmuir adsorption isotherm used to describe the dependence of surfactant adsorption, surface tension and Gibbs elasticity on the surfactant concentration, see Tables 1 and 2. As

before, we have used the approximation $D_{1s} \approx D_1$ (surface diffusivity equal to the bulk diffusivity). The surfactant concentration in Fig. 12 is scaled with the reference concentration c_0 , which is also given in Table 4; for Triton X100 and SDS + 0.1 M NaCl c_0 is chosen to coincide with the critical micellization concentration (CMC). The driving force, F, was taken to be the buoyancy force for dodecane drops in water. The surface force F_s is identified with the van der Waals attraction; the Hamaker function $A_H(h)$ was calculated by means of Eq. (86), see below. The mean drop radius in Fig. 12 is $a = 20 \,\mu\text{m}$. As seen in the figure, for such small drops $\Phi_v \approx 1$ for Triton X100 and BSA, i.e. the drop surfaces turn out to be tangentially immobile in the whole investigated concentration range. On the other hand, Φ_v becomes considerably greater than 1 for the lowest SDS concentrations, which indicates increased mobility of the drop surfaces.

Asymptotic cases	Mobility factors, Φ_v and Φ_p		
d	$\Phi_{v} = (b+1) / \left[1 - \frac{d}{3h} + \frac{8s_{v}b}{3(b+1)} \left(b + \frac{d}{h} \right) \right]$		
$\frac{a}{h} \ll 1$	$\Phi_{p} = \frac{\Phi_{v}}{b+1} \left[1 - \frac{2d}{3h} + \frac{16s_{v}b}{3(b+1)} \left(b + \frac{2d}{h} \right) \right]$		
d	$\Phi_{v} = \frac{h_s}{2h} \left[\frac{4s_v h_s}{3h} - 1 - 8s_v + \ln \frac{d}{h} \right]$		
$\frac{a}{h} >> 1$	$\Phi_{\rm p} = \frac{2h\Phi_{\rm v}}{h_s} \left(1 + \frac{8s_{\rm v}h_s}{3h}\right)$		

 Table 3. Asymptotic
 expressions for the mobility factors from Ref. [138]

Table 4. Parameters for the solutions of Triton X100, SDS + 0.1 M NaCl, and BSA [138].

	Triton X100	SDS + 0.1 M NaCl	BSA
Γ_{∞}	$1.75 \times 10^{-6} \text{ mol/m}^2$	$3.75 \times 10^{-6} \text{ mol/m}^2$	$0.38 \times 10^{-6} \text{ mol/m}^2$
1/K	$4.9 \times 10^{-6} \text{ kg/m}^3$	$1.4 \times 10^{-3} \text{ kg/m}^3$	$0.124 \times 10^{-6} \text{ kg/m}^3$
D_1	$2.6 \times 10^{-10} \text{ m}^2/\text{s}$	$6.0 \times 10^{-10} \text{ m}^2/\text{s}$	$1.0 \times 10^{-10} \text{ m}^2/\text{s}$
\mathcal{C}_0	0.4 kg/m^3 (CMC)	0.33 kg/m ³ (CMC)	0.6 kg/m^3
η_s	1.0×10 ⁻⁶ mPa s	1.0×10^{-6} mPa s	5.0×10^{-3} mPa s


Fig. 12. Theoretical dependence of the mobility factor Φ_v on the surfactant concentration c_1 , calculated in Ref. [144] for the nonionic surfactant Triton X100, ionic surfactant SDS + 0.1 M NaCl, and the protein BSA; the curves for Triton X100 and BSA coincide. The mean drop radius is $a = 20 \mu m$ and the film thickness is $h \approx 10 nm$; the other parameters values are listed in Table 4.

3. Formation of pimple

Let us consider two spherical emulsion drops approaching each other, which interact through the van der Waals attractive surface force. Sooner or later interfacial deformation will occur in the zone of drop-drop contact. The calculations [138] show that if the drop radius *a* is greater than c.a. 80 μ m, the drop interfaces bend inwards (under the action of the hydrodynamic pressure) and a "*dimple*" is formed in the contact zone; soon the dimple transforms into an almost plane-parallel film (Fig. 2D). In contrast, if the drop radius *a* is smaller than c.a. 80 μ m, then at a given surface-to-surface distance $h = h_p$ the drop surface in the contact zone bends outwards and a "*pimple*" forms due to the van der Waals attraction, see the inset in Fig. 13. Correspondingly, h_p is called the pimpling distance. Since the size of the drops in an emulsion is usually markedly below 80 μ m, we will consider here only the formation of pimple.

The formation of pimple has been found out by Yanitsios & Davis [139] in computer calculations for emulsion drops from pure liquids, without any surfactant. Next, by means of numerical calculations Cristini et al. [140] established the formation of pimple for emulsion

drops covered with insoluble surfactant in the case of negligible surface diffusion; their computations showed that a rapid coalescence took part for $h < h_p$. A complete treatment of the problem for the formation of pimple was given in Ref. [138], where the effects of surface and bulk diffusion of surfactant, as well as the surface elasticity and viscosity, were taken into account, and analytical expressions were derived.



Fig. 13. Calculated dependence of the pimple thickness, h_p , on the surfactant concentration, c_1 , for emulsion films formed from aqueous solutions of SDS + 0.1 M NaCl, Triton X100, and BSA; the used parameters values are listed in Table 4. The inset illustrates the shape of the drop surfaces in the zone of contact.

The origin of the pimple formation is the fact that the van der Waals disjoining pressure, $\Pi \propto 1/h^3$, grows faster than the hydrodynamic pressure with the decrease of *h*. For a certain distance, $h = h_p$, Π counterbalances the hydrodynamic pressure [138]:

$$\frac{F - F_s}{\pi a h_p} \Phi_p(h_p; b, d, s_v) + \Pi(h_p) = 0 \quad , \tag{59}$$

where Φ_p is the mobility factor for the pressure. Further, for shorter distance between the drops, $h < h_p$, the pimples spontaneously grow until the drop surfaces touch each other and the drops coalesce. The pimple formation at $h = h_p$ can be interpreted as an onset of instability *without* fluctuations.

Analytical asymptotic expressions for the pressure mobility factor, Φ_p , can be found in Table 3. In general, Φ_p is to be calculated numerically. In the case of tangentially immobile surfaces of the drops Eq. (59) yields a very simple formula for the pimpling distance [138]:

$$h_{\rm p} = \sqrt{\frac{aA_{\rm H}}{12F}}$$
 (tangentially immobile surfaces) (60)

In the more complicated case of mobile drop surfaces Eq. (59) has to be solved numerically. Fig. 13 shows calculated curves for the dependence of h_p vs. the surfactant concentration; the used parameter values are the same as for Fig. 12, see Table 4. Since the surfaces of the drops with BSA and Triton X100 are tangentially immobile, the respective pimpling distance is practically constant (independent of surfactant concentration), and given by Eq. (60). The effect of surface mobility shows up for the emulsions with SDS + 0.1 M NaCl, for which the pimpling distance h_p is greater (Fig. 13). These calculations demonstrate that h_p is typically of the order of 10 nm.

If the pimpling distance is greater than the critical distance, $h_p > h_c$, then the pimpling will be the reason for coalescence. On the other hand, if $h_c > h_p$, then the coalescence will be caused by the fluctuation capillary waves, see the next subsection.

4. Transitional and critical distance

As already mentioned, when two emulsion drops approach each other, the attractive surface forces promote the growth of fluctuation capillary waves in the contact zone. At a given, sufficiently small surface-to-surface distance, called the transitional distance, h_t , the waves with a given length (usually the longest one) begin to grow; this is a transition from stability to instability. During the growth of the waves the gap width continues to decrease, which leads to destabilization and growth of waves with other lengths. Finally, the surfaces of the two drops touch other due to the enhanced interfacial undulations, and coalescence takes place. The latter act corresponds to a given mean surface-to-surface distance, called the critical thickness, h_c . The difference between the transitional and critical distance, $h_t > h_c$, is due to the fact that during the growth of the capillary waves the average film thickness continues to decrease, insofar as the drops are moving against each other driven by the force $F - F_s$. In the simpler case of immobile drops ($F - F_s = 0$), considered in Section III.C, one has $h_t = h_c$.

A general equation for determining h_t , which takes into account the effect of surface mobility, has been reported in Ref. [136]:

$$\left[F - F_s\right]\Psi(\overline{d}) = \frac{\pi a^2 h_t^3}{32\sigma} \left[\left(\frac{\partial \Pi}{\partial h}\right)^2 - \frac{8\sigma}{a} \left(\frac{\partial^2 \Pi}{\partial h^2}\right) \right]_{h=h_t}, \qquad (61)$$

where

$$\overline{d} = \frac{d}{h} , \qquad \Psi(\overline{d}) = \frac{(\overline{d}+2)\overline{d}^2}{4(\overline{d}+1)^2 [(\overline{d}+1)\ln(\overline{d}+1) - \overline{d}]}$$
(62)

The function $\Psi(\overline{d})$ accounts for the effect of the surface mobility. For large interfacial elasticity one has $\overline{d} \rightarrow 0$, see Eq. (53); then $\Psi \rightarrow 1$ and Eq. (61) acquires a simpler form, corresponding to drops of tangentially immobile interfaces. In the other limit, small interfacial elasticity, one has $\overline{d} \gg 1$ and in such a case $\Psi \propto 1/\ln \overline{d}$, i.e. Ψ decreases with the increase of \overline{d} , that is with the decrease of $E_{\rm G}$. Numerical solution of this problem is reported in Ref. [24]. The effect of the interfacial viscosity on the transitional distance, which is neglected in Eq. (61), is examined in Ref. [141]. It is established therein that the critical distance, $h_{\rm c}$, can be with about 10 % smaller than $h_{\rm t}$.



Fig. 14. Dependence of the transitional distance between two drops, h_t , on the surfactant concentration, c_1 , calculated with the help of Eq. (61) for three values of the mean drop radius a.

The dependence of the transitional distance h_t on the surfactant concentration, calculated with the help of Eq. (61), is shown in Fig. 14; the three curves correspond to three fixed values of the mean drop radius a. The calculations are carried out for the system with SDS + 0.1 M NaCl in the aqueous phase, see Table 4; the oil phase is dodecane. One sees that the increase of the surfactant concentration leads to a decrease of the transitional thickness, which corresponds to a greater stability of the emulsion against coalescence. Physically this is related to the damping of the fluctuation capillary waves by the adsorbed surfactant. Moreover, the transitional thickness for two approaching drops increases with the decrease of the drop radius a (Fig. 14), which is exactly the opposite to the tendency for quiescent drops in Fig. 11 (we recall that $h_t = h_c$ for quiescent drops). The difference can be attributed to the strong dependence of the buoyancy force F on the drop radius a (such an effect is missing for the quiescent drops).

The comparison between Figs. 13 and 14 shows that for the emulsion with SDS + 0.1 M NaCl one has $h_t > h_p$. In other words the theory predicts that in this emulsion the drops will coalesce due to the fluctuation capillary waves, rather than owing to the pimpling.

If the coalescence is promoted by the van der Waals attractive surface force, from Eq. (61) one can deduce asymptotic expressions for h_t , corresponding to tangentially immobile drop surfaces ($\Psi = 1$) [136]:

$$h_{\rm t} = \left[a^2 A_{\rm H}^2 / (128\pi\sigma F) \right]^{1/5}, \qquad \text{for } F >> F_s \qquad (63)$$

$$h_{\rm t} = [5aA_{\rm H} / (12F)]^{1/2}, \qquad \text{for } F << F_s$$
 (64)

In particular, if F is the buoyancy force, then $F \propto a^3$ and for small droplets ($F \ll F_s$) one obtains $h_t \propto 1/a$, i.e. the critical thickness markedly increases with the decrease of the droplet radius.

B. Interaction between Deforming Emulsion Drops

1. Drops of tangentially immobile surfaces

In this section we consider the case, in which a liquid film is formed in the zone of contact of two emulsion drops, see Fig. 7. Such configuration appears between drops in flocs and in concentrated emulsions, including creams.

In a first approximation, one can assume that the viscous dissipation of kinetic energy happens mostly in the thin liquid film intervening between two drops. (In reality, some energy dissipation happens also in the transition zone between the film and the bulk continuous phase.) Then, if the drop interfaces are tangentially immobile (owing to adsorbed surfactant), then the velocity of approach of the two drops can be estimated by means of the Reynolds formula for the velocity of approach of two parallel solid discs of radius R, equal to the film radius [142]:

$$V_{\rm Re} = \frac{2h^3 F_{\rm tot}}{3\pi\eta_{\rm out}R^4} \tag{65}$$

As usual, here h is the film thickness; F_{tot} is the total force exerted on a drop [2]:

$$F_{\rm tot} = F - F_{\rm s} - F_{\rm def} \tag{66}$$

As before, F is the applied external force (buoyancy, centrifugal force, Brownian force, etc.); F_s is the surface force of intermolecular origin, which for deformable drops can be expressed in the form [2,143]

$$F_s = -\frac{dW}{dh} , \qquad W(R,h) \approx \pi R^2 f(h) + \pi a \int_h^\infty f(\hat{h}) d\hat{h} , \qquad (67)$$

where

$$f(h) = \int_{h}^{\infty} \Pi(\hat{h}) d\hat{h}$$
(68)

is the interaction free energy per unit area of a plane-parallel liquid film; W_s is the drop-drop interaction energy due to surface forces, which is a sum of contributions from the planar film and the transition zone film-bulk liquid; for R = 0 Eq. (67) reduces to Eq. (48). Finally, F_{def} is a force originating from the deformation of the drop interfaces [2]:

$$F_{\rm def} = -\frac{d(W_{\rm dil} + W_{\rm bend})}{dh} , \qquad (69)$$

where W_{dil} is the work of interfacial dilatation [143-145]

$$W_{\rm dil} = \sigma \frac{\pi R^4}{2a^2} + \frac{1}{2} E_G \left(\frac{\pi R^4}{2a^2} \right)^2 + \dots, \text{ for } \left(\frac{r}{a} \right)^2 << 1,$$
(70)

and W_{bend} is the work of interfacial bending [146]

$$W_{\text{bend}} = -2\pi r^2 B_0 / a, \quad (r/a)^2 << 1$$
 (71)

 $B_0 = -4k_cH_0$ is the interfacial bending moment; H_0 is the so called spontaneous curvature and k_c is the interfacial curvature elastic modulus.

Initially the two approaching drops are spherical. The deformation in the zone of contact begins when the surface-to-surface distance reaches a certain threshold value, called the

inversion thickness, h_{inv} . One can estimate the inversion thickness from the simple expression $h_{inv} = F/(2\pi\sigma)$, see e.g. Refs. (98,121). The generalized form of the latter equation, accounting for the contribution of the surface forces, reads [136]:

$$h_{\rm inv} = \frac{F}{2\pi\sigma} + \frac{a}{2\sigma} [h_{\rm inv} \Pi(h_{\rm inv}) - f(h_{\rm inv})]$$
(72)

The inversion thickness can be determined by solving numerically Eq. (72).

A generalized expression for the velocity V = -dh/dt, which takes into account the energy dissipation in both film and the transition zone film-bulk liquid, has been derived in Refs. [2,147]:

$$\frac{1}{V} = \frac{1}{V_{\rm Re}} + \frac{1}{V_{\rm Ta}} + \frac{1}{\sqrt{V_{\rm Re}V_{\rm Ta}}}$$
(73)

where the Taylor velocity, V_{Ta} , and the Reynolds velocity, V_{Re} , are defined by means of Eqs. (47) and (65). For $R \rightarrow 0$ (nondeformed spherical drops) Eq. (73) reduces to $V = V_{\text{Ta}}$. On the contrary, for $h \rightarrow 0$ one has $1/V_{\text{Ta}} \ll 1/V_{\text{Re}}$, and then Eq. (73) yields $V \rightarrow V_{\text{Re}}$. Substituting Eqs. (47) and (65), and assuming $F \gg (F_s + F_{\text{def}})$ one can bring Eq. (73) in the form [147]

$$\frac{V_{\rm Ta}}{V} = 1 + \frac{R^2}{ha} + \frac{R^4}{h^2 a^2}$$
(74)

One sees that $V \rightarrow V_{\text{Ta}}$ for $R^2/(ha) \ll 1$. If the external force *F* is predominant, then $R^2 \approx aF/(2\pi\sigma)$, $h_{\text{inv}} \approx F/(2\pi\sigma)$ and it follows that $R^2/a \approx h_{\text{inv}}$ [97,135]; the substitution of the latter equation in Eq. (74) yields

$$\frac{V_{\text{Ta}}}{V} \approx 1 + \frac{h_{\text{inv}}}{h} + \frac{h_{\text{inv}}^2}{h^2}$$
(75)

Equation (75) shows that for $h \le h_{inv}$ the velocity V becomes considerably smaller than V_{Ta} .

2. Effect of surface mobility

When the surfactant is soluble only in the *continuous* phase (we will call such a system "System I", see Fig. 15), it turns out that the respective rate of film thinning V_I is affected by the surface mobility mainly by through the Gibbs elasticity E_G , just as it is for foam films [97,121]:

$$\frac{V_{\rm I}}{V_{\rm Re}} \approx 1 + \frac{1}{\varepsilon_{\rm f}}, \qquad \frac{1}{\varepsilon_{\rm f}} = \frac{6\eta_{\rm I}D_{\rm Is}}{hE_{\rm G}} + \frac{3\eta_{\rm I}D_{\rm I}}{\Gamma_{\rm I}} \left(\frac{\partial c_{\rm I}}{\partial\sigma}\right)^{(e)}$$
(76)



Fig. 15. Two complementary types of emulsion systems obtained by a mere exchange of the continuous phase with the disperse phase. The surfactant is assumed to be soluble only in Liquid 1. (a) Liquid 2 is the disperse phase; (b) Liquid 2 is the continuous phase.

Here $\varepsilon_{\rm f}$ is the so called *foam parameter*; $\eta_{\rm l}$ is the viscosity in the surfactant-containing phase (Liquid 1 in Fig. 15); the influence of the transition zone film - bulk liquid is not accounted for in Eq. (76). Note that the bulk and surface diffusion fluxes (see the terms with $D_{\rm 1s}$ and $D_{\rm 1}$ in the latter equation), which tend to damp the surface tension gradients and to restore the uniformity of the adsorption monolayers, accelerate the film thinning (Fig. 1). Moreover, since $D_{\rm 1s}$ in Eq. (76) is divided by the film thickness *h*, the effect of surface diffusion dominates that of bulk diffusion for small values of the film thickness. On the other hand, the Gibbs elasticity $E_{\rm G}$ (the Marangoni effect) decelerates the thinning. Equation (76) predicts that the rate of thinning is not affected by the circulation of liquid in the droplets, i.e. System I really behaves as a foam system.

It was established theoretically [97,133] that when the surfactant is dissolved in the *drop* phase (System II in Fig. 15) it remains uniformly distributed throughout the drop surface during the film thinning and interfacial tension gradients do not appear. This is due to a powerful supply of surfactant, which is driven by the convective diffusion from the bulk of the drops toward their surfaces. For that reason, the drainage of the film surfaces is not opposed by surface tension gradients and the rate of film thinning, $V_{\rm II}$, is the same as in the case of pure liquid phases [97,133]:

$$\frac{V_{\rm II}}{V_{\rm Re}} \approx \frac{1}{\varepsilon_{\rm e}} \approx \frac{\eta_1 \delta}{\eta_2 h} \approx \left(\frac{108\pi \eta_1^3 R^4}{\rho_2 \eta_2 h^4 F}\right)^{1/3} \tag{77}$$

Here ε_e is called emulsion parameter, δ is the thickness of the hydrodynamic boundary layer inside the drops, ρ_2 and η_2 are the mass density and dynamic viscosity of Liquid 2, which does not contain dissolved surfactant. The validity of Eq. (77) was confirmed experimentally [134].

The only difference between the two systems in Fig. 15 is the exchange of the continuous and drop phases. Assume for simplicity that V_{Re} is the same for both systems. In addition, usually $\varepsilon_{\text{f}} \approx 0.1$ and $\varepsilon_{\text{e}} \approx 10^{-2} - 10^{-3}$. Then from Eqs. (76) and (77) one obtains [97,121,133]

$$\frac{V_{\rm II}}{V_{\rm I}} \approx \frac{\varepsilon_{\rm f}}{\varepsilon_{\rm e}} \approx \frac{0.1}{10^{-2} \text{ to } 10^{-3}} \approx 10 \text{ to } 100 \qquad (\text{deformed drops}) \tag{78}$$

Hence, the rate of film thinning in System II is much greater than that in System I. Therefore, the location of the surfactant has a dramatic effect on the thinning rate and thereby – on the drop life time. Note also that the interfacial tension in both systems is the same. Hence, the mere phase inversion of an emulsion, from Liquid 1-in-Liquid 2 to Liquid 2-in-Liquid 1 (Fig. 15), could change the emulsion life time by orders of magnitude. As discussed in Section V below, the situation with the interaction in Taylor regime (between spherical, nondeformed drops) is similar. These facts are closely related to the explanation of the Bancroft rule for the stability of emulsions (see Section V) and the process of chemical demulsification, see Ref. [1].

Equations (76) and (77) do not take into account the hydrodynamic interactions across the transition zone around the film, which can be essential if the film radius R is relatively small. In the latter case the effect of surface viscosity becomes important for System I. In this case Eq. (76) can be presented in a more general form [137]:

$$\frac{V_{\rm I}}{V_{\rm Re}} = \Phi_{\rm v}(h;b,d,R,s_{\rm v}) , \qquad (79)$$

where Φ_v is a mobility function. In Ref. [137] a general, but voluminous, analytical expression for Φ_v is derived in the form of infinite series expansion; it accounts for the effects of surface elasticity, surface viscosity, bulk and surface diffusion. In some special cases this infinite series can be summed up and closed expressions for Φ_v can be obtained. Such is the case when the effect of the surface viscosity is negligible, $s_v \rightarrow 0$; the respective expression for Φ_v reads [137]:

$$\frac{1}{\Phi_{v}} = \frac{1}{1+b+h_{s}/h} + \frac{2h}{N_{R}^{4}h_{s}} \left\{ \left[\frac{h}{h_{s}} (1+b)(1-N_{R}^{2}) + 1 \right] \ln \left[1 + \frac{h_{s}}{h(1+b)} \right] + N_{R}^{2} - 1 \right\}, \quad (80)$$

where the dimensionless parameter $N_R \equiv R/\sqrt{ah}$ accounts for the effect of the film radius. In the case of emulsion drops $N_R \approx 1$; however, if experiments with emulsion films are performed in the experimental cell of Scheludko & Exerowa [148,149], which allows independent control of *R*, then one usually has $N_R \gg 1$. {The original experiments in Refs. [148,149] have been carried out with foam films, but a similar technique can be applied to investigate emulsion films, see e.g. Refs. [91, 150–158]} In the limit of large plane-parallel film, $N_R \gg 1$, Eq. (80) reduces to the result of Radoev *et al.* [159]: $V_1/V_{Re} = 1 + b + h_s/h$ (effect of the transition zone negligible). For insoluble surfactants the parameter *b* in Eq. (80) must be set equal to zero.

Under certain experimental conditions, like those in Ref. [60], the motion of surfactant along an oil-water interface represents a flow of a two-dimensional *incompressible viscous* fluid. In such a case Eq. (79) acquires the following specific form [137]:

$$\frac{1}{V_{\rm I}} = \frac{1}{V_{\rm Ta}} + \frac{1}{V_{\rm Re}} + \frac{1}{\sqrt{V_{\rm Ta}V_{\rm Re}}} - \frac{1}{2s_{\rm v}} \left(\frac{1}{V_{\rm Ta}} + \frac{1}{V_{\rm Re}} + \frac{1}{\sqrt{V_{\rm Ta}V_{\rm Re}}} + \frac{N_R^2}{3\sqrt{V_{\rm Ta}V_{\rm Re}}} \right)$$
(81)

Equation (81) is a truncated power expansion for $s_v >> 1$. In the limit of tangentially immobile interfaces ($s_v \rightarrow \infty$) Eq. (81) reduces to Eq. (73).

To illustrate the effects of various factors on the velocity of approach of two deforming emulsion drops (Fig. 15a) we used the general expression from Ref. [137] (the infinite series expansion) to calculate the mobility factor Φ_v ; results are shown in Figs. 16 and 17. First of all, in Fig. 16 we illustrate the effects of bulk and surface diffusion. For that reason $\Phi_v \equiv V_1/V_{Re}$ is plotted vs. the parameter *b*, related to the bulk diffusivity, for various values of h_s/h ; h_s is related to the surface diffusivity, see Eq. (54). If the hydrodynamic interaction were operative only in the film, then one would obtain $V_1/V_{Re} \ge 1$. However, all calculated values of V_1/V_{Re} are smaller than 0.51 (Fig. 16); this fact evidences for a significant effect of the hydrodynamic interactions in the transition zone around the film. Moreover, in Fig. 16 one sees that for b > 10the mobility factor Φ_v is independent of the surface diffusivity. On the other hand, for b < 10 a considerable effect of surface diffusivity shows up: the greater the surface diffusivity effect, h_s/h , the greater the interfacial mobility factor Φ_v . For the upper curve in Fig. 16 the interfacial mobility is determined mostly by the effect of surface viscosity, s_v , which is set equal to 1 for all curves in the figure.



Fig. 16. Effect of the surface diffusion parameter, h_s/h , on the variation of the mobility factor, $\Phi_v \equiv V_I/V_{Re}$, with the bulk diffusion parameter, *b*, for fixed $s_v = 1$ and $N_R = 1$.



Fig. 17. Effect of the surface viscosity parameter, s_v , on the variation of the mobility factor, $\Phi_v \equiv V_I/V_{Re}$, with the bulk diffusion parameter, *b*, for fixed $h_s/h = 1$ and $N_R = 1$.

To illustrate the effect of surface viscosity, s_v , in Figure 17 we have plotted the mobility factor $\Phi_v \equiv V_I/V_{Re}$ vs. *b* for three different values of s_v . For the higher surface viscosities, $s_v = 1$ and 5, the mobility factor is $V_I/V_{Re} < 1$, which again indicates a strong hydrodynamic interaction in the transition zone around the film. For the lowest surface viscosity, $s_v = 0.1$, the mobility factor is sensitive to the effect of bulk diffusion, characterized by *b*: for b > 3 we have $V_I/V_{Re} > 1$, i.e. we observe a considerable rise in the interfacial mobility (Fig. 17).

3. Critical thickness of the film between two deforming drops

As already mentioned, the transition from stability to instability occurs when the thickness of the gap between two colliding emulsion drops decreases down to a "transitional" thickness h_t . For $h_t > h > h_c$ the film continues to thin, while the instabilities grow, until the film ruptures at the critical thickness $h = h_c$.

Equation (61) above determines the transitional distance between two spherical emulsion drops. An analogue of this equation for the case of two deformed drops (Fig. 15a) has been obtained in the form of a transcendental equation [2,136]:

$$\frac{2h_{\rm t}+d}{h_{\rm t}+d} = \frac{h_{\rm t}R^2[\Pi'(h_{\rm t})]^2}{8\sigma[2\sigma/a - \Pi(h_{\rm t})]}, \qquad \Pi' \equiv \frac{\partial\Pi}{\partial h}$$
(82)

Equation (82) shows that the disjoining pressure significantly influences the transitional thickness h_t . The effect of surface mobility is characterized by the parameter d, see Eq. (53); in particular, d = 0 for tangentially immobile interfaces. Equation (82) is valid for $\Pi < 2\sigma/a$, i.e. when the film thins and ruptures before reaching its equilibrium thickness, corresponding to $\Pi = 2\sigma/a$; cf. Eqs. (42), (43) and (59).

The calculation of the transitional thickness h_t is a prerequisite for computing the critical thickness h_c , which can be obtained as a solution of the equation [95,96]

$$h_{\rm c}^2 = \frac{2kT}{I(h_{\rm c},h_{\rm t})} \exp\left[\frac{I(h_{\rm c},h_{\rm t})}{4\sigma}\right]$$
(83)

where $I(h_t,h_c)$ stands for the following function

$$I(h_{\rm c}, h_{\rm t}) = \Pi'(h_{\rm t}) R^2 \int_{h_{\rm c}}^{h_{\rm t}} \frac{\Pi' dh}{\Phi_{\rm v}(h) [2\sigma / a - \Pi(h)]}$$
(84)

In the special case of tangentially immobile interfaces and large film (negligible effect of the transition zone) one has $\Phi_v(h) \equiv 1$, and the integration in Eq. (84) can be carried out [95]:

$$I(h_{\rm c}, h_{\rm t}) = \Pi'(h_{\rm t}) R^2 \ln \left[\frac{2\sigma / a - \Pi(h_{\rm c})}{2\sigma / a - \Pi(h_{\rm t})} \right]$$
(85)

Note that Eqs. (82)-(85) hold not only for an emulsion film formed between two oil drops, but also for a foam film intervening between two gas bubbles. In Fig. 18 we compare the prediction of Eqs. (82)-(84) with experimental data for h_c vs. R, obtained by Manev et al. [120] for foam films formed from aqueous solution of 0.43 mM SDS + 0.1 M NaCl. The mobility factor $\Phi_v(h)$ was calculated by using the exact expression (the infinite series) from Ref. [137]. Parameters such as σ , E_G , Γ_1 and $\partial \Gamma_1/\partial c_1$, see Eqs. (53)-(55), are obtained from the experimental fit in Ref. [17], in the same way, as the numerical data in Fig. 5 have been obtained (see Section II.A.2). The disjoining pressure was attributed to the van der Waals attraction: $\Pi = -A_H/(6\pi h^3)$. To account for the effect of the electromagnetic retardation on the dispersion interaction, we used an expression proposed by Russel et al. [160]:

$$A_{\rm H}(h) = \frac{3h_P \nu}{4\pi} \frac{\left(n_o^2 - n_w^2\right)^2}{\left(n_o^2 + n_w^2\right)^{3/2}} \int_0^\infty \frac{\left(1 + 2\tilde{h}z\right)\exp\left(-2\tilde{h}z\right)}{\left(1 + 2z^2\right)^2} dz$$
(86)

Here $h_p = 6.63 \times 10^{-34}$ J.s is the Planck constant, $v \approx 3.0 \times 10^{15}$ Hz is the main electronic absorption frequency; n_o and n_w are the refractive indices of the non-aqueous and aqueous phases; for a foam film $n_o = 1$ and $n_w = 1.333$; the dimensionless thickness \tilde{h} is defined by the expression

$$\widetilde{h} = n_w \left(n_o^2 + n_w^2 \right)^{1/2} \frac{2\pi v h}{c} , \qquad (87)$$

where $c = 3.0 \times 10^{10}$ cm/s is the speed of light. For small thickness $A_{\rm H}$, as given by Eqs. (86)-(87), is constant, whereas for large thickness *h* one obtains $A_{\rm H} \propto h^{-1}$. The solid line in Fig. 18 was calculated with the help of Eqs. (82)-(87) *without* using any adjustable parameters; one sees that there is an excellent agreement between this theoretical model and the experiment.

The dot-dashed line in Fig. 18 shows the prediction of the theoretical model by Malhotra and Wasan [116]. Our calculations showed that for the specific surfactant and salt concentrations (0.43 mM SDS + 0.1 M NaCl) the interfaces are almost tangentially immobile. Moreover, in these experiments the film radius R is sufficiently large, which allows one to

neglect effects of the transition zone, i.e. to ignore the last two terms in Eq. (73). Consequently, the difference between the model from Ref. [116] and the experimental data (Fig. 18) cannot be attributed to the latter two effects (interfacial mobility and transition zone), which have not been taken into account in Ref. [116]. The main reasons for the difference between the output of Ref. [116] and the experiment are that (i) these authors have, in fact, calculated h_t and identified it with h_c , and (ii) a constant value of A_H has been used, instead of Eq. (86), i.e. the electromagnetic retardation effect has been neglected. It is interesting to note that the retardation effect turns out to be important in the experimental range of critical thicknesses, in this specific case: 25 nm $< h_c < 50$ nm.



Fig. 18. Critical thickness, h_c , vs. radius, R, of a foam film formed from aqueous solution of 0.43 mM SDS + 0.1 M NaCl: comparison between experimental points, measured by Manev et al. [120], with our theoretical model based on Eqs. (82)-(87) (the solid line) and the model by Malhotra & Wasan [116] (the dashed line).

V. INTERPRETATION OF THE BANCROFT RULE

A simple rule connecting the emulsion stability with the surfactant properties was formulated by Bancroft [161]. The Bancroft rule states that "in order to have a stable emulsion the surfactant must be soluble in the continuous phase". Most of the emulsion systems obey this rule, but some exclusions have been also found [162]. The results about the drop-drop

interactions, presented in Section IV, allow one to give a semi-quantitative interpretation of the rule and the exclusions [163,1,2].

According to Davies and Rideal [6], both types of emulsions (water-in-oil and oil-inwater) are formed during the homogenization process, but only the one with lower coalescence rate survives. If the initial drop concentration for the two emulsions (System I and System II, see Fig. 15) is the same, the corresponding coalescence rates for the two emulsions will be (approximately) proportional to the respective velocities of film thinning, $V_{\rm I}$ and $V_{\rm II}$ [163]:

$$\frac{\text{Rate I}}{\text{Rate II}} \approx \frac{V_{\text{I}}}{V_{\text{II}}}$$
(88)

A. Case of Deforming Drops

In the case of *deforming* drops, using Eqs. (65), (76) and (77) one can represent Eq. (88) into the form [1,163]

$$\frac{\text{Rate I}}{\text{Rate II}} \approx (486\rho_2 D_{1s}^3)^{1/3} \left(\frac{h_{\text{c,II}}^3}{h_{\text{c,II}}^2}\right)^{1/3} \left(\frac{\eta_2}{R^2}\right)^{1/3} \frac{2\sigma/a - \Pi_{\text{I}}}{E_G (2\sigma/a - \Pi_{\text{II}})^{2/3}}$$
(89)

where $h_{c,I}$ and $h_{c,II}$ denote the critical thickness of film rupture for the two emulsion systems in Fig. 15; Π_I and Π_{II} denote the disjoining pressure of the respective films. To obtain Eq. (89) we have also used the estimate $F_{tot} \approx (2\sigma/a - \Pi)R^2$, see Ref. [149]. The product of the first three multipliers in the right-hand side of Eq. (89), which are related to the *hydrodynamic* stability, is c.a. $8 \times 10^{-5} \text{ dyn}^{2/3} \text{ cm}^{-1/3}$ for typical parameter values [1]. The last multiplier in Eq. (89) accounts for the *thermodynamic* stability of the two types of emulsion films. Many conclusions regarding the type of the formed emulsion can be drawn from Eq. (89) [1,62,163]:

In thick films the disjoining pressures, Π_{I} and Π_{II} , are zero, and then the ratio in Eq. (89) will be very small. Consequently, emulsion I (surfactant soluble in the continuous phase) will coalesce much more slowly than emulsion II; hence emulsion I will survive. Thus we get an explanation of the empirical Bancroft rule. The emulsion behavior in this case is controlled mostly by the hydrodynamic factors, i.e. the factors related to the *kinetic* stability.

The disjoining pressure, Π , can substantially change, and even reverse, the behavior of the system if it is comparable by magnitude with the capillary pressure, $2\sigma/a$. For example if $(2\sigma/a - \Pi_{II}) \rightarrow 0$ at finite value of $(2\sigma/a - \Pi_{I})$, then the ratio in Eq. (89) may become much larger than unity, which means that System II will become *thermodynamically* stable. This fact

can explain some exclusions from the Bancroft rule, like that established by Binks [162]. Moreover, a large stabilizing disjoining pressure is operative in emulsions with high volume fraction of the disperse phase, above 95% in some cases [164].

The Gibbs elasticity, $E_{\rm G}$, favors the formation of emulsion I (Fig. 15a), because it slows down the film thinning. On the other hand, increased surface diffusivity, $D_{\rm 1s}$, decreases this effect, because it helps the interfacial tension gradients to relax thus facilitating the formation of emulsion II.

The film radius, *R*, increases, whereas the capillary pressure, $2\sigma/a$, decreases with the rise of the drop radius, *a*. Therefore, larger drops will tend to form emulsion I, although the effect is not very pronounced, see Eq. (89). The difference between the critical thicknesses of the two emulsions affects only slightly the rate ratio in Eq. (89), although the value of h_c itself is important.

The viscosity of the surfactant-containing phase, η_1 , does not appear in Eq. (89); there is only a weak dependence on η_2 . This fact are consonant with the experimental findings about a negligible effect of viscosity, see Ref. [6], p. 381 therein.

The interfacial tension, σ , affects directly the rate ratio in Eq. (89) through the capillary pressure, $2\sigma/a$. The addition of electrolyte would affect mostly the electrostatic component of the disjoining pressure, see Fig. 8a, which is suppressed by the electrolyte; the latter has a destabilizing effect on O/W emulsions. In the case of ionic surfactant solutions the addition of electrolyte rises the surfactant adsorption and the Gibbs elasticity (see Fig. 5), which favors the stability of emulsion I.

Surface active additives (such as cosurfactants, demulsifiers, etc.) may affect the emulsifier partitioning between the phases and its adsorption, thereby changing the Gibbs elasticity and the interfacial tension. The surface active additive may change also the surface charge (mainly through increasing the spacing among the emulsifier ionic headgroups) thus decreasing the electrostatic disjoining pressure and favoring the W/O emulsion. Polymeric surfactants and adsorbed proteins increase the steric repulsion between the film surfaces; they may favor either of the emulsions O/W or W/O depending on their conformation at the interface and their surface activity.

The temperature affects strongly both the solubility and the surface activity of non-ionic surfactants [165]. It is well known that at higher temperature non-ionic surfactants become

more oil-soluble, which favors the W/O emulsion. These effects may change the type of emulsion formed at the phase inversion temperature (PIT) [166]. The temperature effect has numerous implications, some of them being the change of the Gibbs elasticity, E_G , and the interfacial tension, σ .

B. Case of Spherical Drops

Equation (89) was obtained for deforming emulsion drops, i.e. for drops which can approach each other at a surface-to-surface distance smaller than the inversion thickness h_{inv} , see Eq. (72). Other possibility is the drops to remain *spherical* during their collision, up to their eventual coalescence at $h = h_c$; in such a case the expressions for V_I and V_{II} , which are to be substituted in Eq. (88), differ from Eqs. (76) and (77).

Let us first consider the case of System II (surfactant inside the drops, Fig. 15b) in which case the two drops approach each other like drops from pure liquid phases (if only the surface viscosity effect is negligible). Therefore, to estimate the velocity of approach of such two aqueous droplets one can use the following approximate expression, which directly follows from Eq. (49) for $\xi >> 1$:

$$V_{\rm II} / V_{\rm Ta}^{\rm (II)} \approx 0.811 \frac{\eta_2}{\eta_1} \sqrt{\frac{a}{h}}$$
 (90)

(For the system from Fig. 15b one is to set $\eta_{out} = \eta_2$ and $\eta_{in} = \eta_1$.) On the other hand, the velocity V_I of droplet approach in System I can be expressed by means of Eq. (52). Note that the Taylor velocities for System I and System II, $V_{Ta}^{(I)}$ and $V_{Ta}^{(II)}$, are different because of differences in viscosity and droplet-droplet interaction, see Eq. (47). Then combining Eqs. (47), (52), (88) and (90) we arrive at the following criterion for formation of emulsion of type I or II [2]:

$$\frac{\text{Rate I}}{\text{Rate II}} \approx 1.233 \ \frac{h_s}{2h} \sqrt{\frac{h}{a}} \left[\frac{\overline{d}+1}{\overline{d}} \ln(\overline{d}+1) - 1 \right]^{-1} \frac{(F-F_s)_{\text{I}}}{(F-F_s)_{\text{II}}}$$
(91)

where $\overline{d} \equiv d/h$; see Eqs. (53)-(55) about the definition of *d*, *b* and *h_s*. In the case of large surface (Gibbs) elasticity, $E_G \gg 1$, one has $\overline{d} \ll 1$; hence, one can expand the logarithm in Eq. (91) to obtain [2]:

$$\frac{\text{Rate I}}{\text{Rate II}} \approx 1.233 \sqrt{\frac{h_c}{a}} \frac{1+b}{1-\overline{d}/3+O(\overline{d}^2)} \frac{(F-F_s)_{\text{I}}}{(F-F_s)_{\text{II}}}$$
(92)

Here we have substituted $h = h_c$, which is fulfilled at the moment of coalescence. For typical emulsion systems one has $a \gg h_c$, and then Eq. (92) yields Rate I/Rate II << 1; therefore System I (with surfactant in the continuous phase, Fig. 15a) will survive. This prediction of Eq. (92) for spherical drops is analogous to the conclusion from Eq. (89) for deformable drops. Both these predictions essentially coincide with the Bancroft rule and are valid for cases, in which the *hydrodynamic* stability factors prevail over the *thermodynamic* ones. The latter become significant close to the equilibrium state, $F_s \approx F$, and could bring about exclusions from the Bancroft rule, especially when $(F - F_s)_{II} \rightarrow 0$. The following conclusions, more specific for the case of *spherical* drops, can be also drawn from Eqs. (91) and (92):

For larger droplets (larger *a*) the transitional distance h_t (and the critical distance h_c as well) is smaller, see Figure 14. Then it follows from Eq. (91) that the difference between the coalescence rates in Systems I and II will become larger [2]. On the contrary, the difference between Rate I and Rate II decreases with the reduction of the droplet size *a*, which is accompanied with an increase of the critical thickness h_c . Note that this effect of *a* cannot be derived from the criterion for deforming drops, Eq. (89).

The effect of the bulk viscosity is not explicitly present in Eq. (92), although there could be some weak implicit dependence through the parameters d and b, see Eqs. (53) and (55). This conclusion agrees with the experimental observations about a very weak dependence of the volume fraction of phase inversion on the viscosity of the continuous phase [6].

The increase of the bulk and surface diffusivities, D_1 and D_{1s} , which tend to damp the surface tension gradients, leads to an increase of the parameters *b* and *d*, which decreases the difference between Rate I and Rate II, see Eq. (53), (55) and (92). In contrast, the increase of the Gibbs elasticity, E_G , leads to decrease of *d* and thus favors the survival of System I. These are the same tendencies as for deforming drops (Section V.A). In the limit of tangentially *immobile* interfaces ($E_G \rightarrow \infty$) one has d = 0 and b = 0 and the criterion (92) further simplifies [2]:

$$\frac{\text{Rate I}}{\text{Rate II}} \approx \frac{V_{\text{I}}}{V_{\text{II}}} \approx 1.233 \sqrt{\frac{h_{\text{c}}}{a}} \frac{\left(F - F_{s}\right)_{\text{I}}}{\left(F - F_{s}\right)_{\text{II}}} \qquad (E_{\text{G}} \rightarrow \infty)$$
(93)

The effect of surface viscosity, η_s , is neglected when deriving Eqs. (91)-(93). Based on the hydrodynamic equations one can estimate that this effect is really negligible when [2]

$$\frac{\eta a^2}{\eta_s h} >> 1, \tag{94}$$

where η stands for the bulk viscosity, which is assumed to be of the same order of magnitude for the liquids inside and outside the drops. If for a certain system, or under certain conditions, the criterion, Eq. (94), is not satisfied, one can expect that the surface viscosity will suppress the interfacial mobility for both Systems I and II. Then the difference between Rate I and Rate II will be determined mostly by thermodynamic factors, such as the surface force F_s .

Although Eqs. (89) and (91) lead us to some more general conclusions than the original Bancroft rule (e.g. the possibility for inversion of the emulsion stability due to disjoining pressure effects), we neither claim that Bancroft rule, or its extension based on Eqs. (89) and (91), have general validity, nor that we have given a general explanation of the emulsion stability. The coagulation in emulsions is such a complex phenomenon, influenced by too many different factors, that according to us any attempt for formulating a general explanation (or criterion) is hopeless. Our treatment is theoretical and as every theory, it has limitations inherent to the model used and therefore is valid only under specific conditions. It should not be applied to system where these conditions are not fulfilled. The main assumptions and limitations of the model are [2]: the fluctuation-wave mechanism for coalescence is assumed to be operative (see Fig. 10); the surfactant transfer onto the surface is under diffusion or electrodiffusion control; parameter b defined by Eq. (55) does not account for the demicellization kinetics for $c_1 > CMC$; the effect of surface viscosity is not taken into account in Eqs. (89) and (91). Only *small* perturbations in the surfactant distribution, which are due to the flow, have been considered; however, under strongly non-equilibrium conditions (like turbulent flows) we could find that new effects come into play, which may significantly alter the trend of the phenomenon.

VI. KINETICS OF COAGULATION IN EMULSIONS

A. Types of Coagulation in Emulsions

The coagulation in an emulsion is a process, in which the separate emulsion drops merge to form larger drops (coalescence) and/or assemble into flocs (flocculation), see Fig. 2. If

the films intervening between the drops in a floc are unstable, their breakage is equivalent to coalescence, see step $D\rightarrow C$ in Fig. 2. In other words, the coagulation in an emulsion includes flocculation and coalescence, which could occur as parallel or consecutive processes.

Various *experimental methods* for monitoring the kinetics of coagulation in emulsions have been developed, such as the electroacoustic method [167], the direct video enhanced microscopic investigation [168], and the ultrasonic attenuation spectroscopy [169].

To a great extent the occurrence of coagulation is determined by the energy, W(R,h), of interaction between two drops. Equation (67), which defines W(R,h), can be applied to any type of surface force (irrespective of its physical origin) if only the range of action of this force is much smaller than the drop radius *a*. In Ref. [2] one can find theoretical expressions for the components of *W* stemming from various surface forces: electrostatic, van der Waals, ionic correlations, hydration repulsion, protrusion and steric interactions, oscillatory structural forces, etc.

If the two drops remain spherical during their interaction (i.e. there is no film in the contact zone and consequently R = 0), then W depends only on a single parameter, W = W(h); as usual, h is the surface-to-surface distance between the two drops. When the approach of the two drops is accompanied with the formation and expansion of a film in the contact zone (Fig. 7), then one can characterize the interaction with $\overline{W}(h)$, which is obtained by averaging of W(R,h) over all configurations with various R at fixed h, see Ref. [143].

The shape of W(h), or $\overline{W}(h)$, qualitatively resembles that of $\Pi(h)$, see Fig. 6. In particular, if only electrostatic and van der Waals interactions are operative, the shape of the dependence W = W(h) resembles Fig. 6a, where an electrostatic barrier is present. The coagulation is called "fast" or "slow" depending on whether that electrostatic barrier is less than kT or higher than kT. In addition, the flocculation is termed "reversible" or "irreversible" depending on whether the depth of the primary minimum (that on the left from the barrier in Fig. 6a) is comparable with kT or much greater than kT. The driving forces of coagulation can be the following.

(i) The *body forces*, like gravity and centrifugation force, cause rising or sedimentation of the droplets depending on whether their mass density is smaller or greater than that of the continuous phase. Since drops of different size move with different velocities, they are subjected to frequent collisions leading to drop aggregation or coalescence, called *orthokinetic coagulation*.

(ii) The *Brownian stochastic force* dominates the gravitational body force for droplets, which are smaller than c.a. 1 μ m. Thus the Brownian collision of two droplets becomes a prerequisite for their flocculation and/or coalescence, which is termed *perikinetic coagulation*.

(iii) The heating of an emulsion produces *temperature gradients*, which in their own turn cause termocapillary migration of the droplets driven by thermally excited gradients of surface tension [170-172]:

$$\nabla_s \sigma = -\frac{E_T}{T} \nabla_s T \qquad \qquad E_T \equiv -\left(\frac{\partial \sigma}{\partial \ln T}\right)_{\Gamma_1}$$
(95)

Here ∇_s is the surface gradient operator and E_T is the coefficient of interfacial thermal elasticity, cf. Eq. (1). The drops moving with different thermocapillary velocities can collide and flocculate or coalesce; this is the *thermal coagulation*.

B. Kinetics of Irreversible Coagulation

1. Basic equations

The kinetic theory of the fast irreversible coagulation was first developed by Smoluchowski [173,174] and later extended to the case of slow and reversible coagulation. In any case of coagulation the general set of kinetic equations reads [175]:

$$\frac{dn_k}{dt} = \frac{1}{2} \sum_{i=1}^{k-1} a_f^{i,k-i} n_i n_{k-i} - n_k \sum_{i=1}^{\infty} a_f^{k,i} n_i + q_k \qquad (k=1, 2, ...)$$
(96)

where t is time, n_1 denotes the number of single drops per unit volume; n_k is number of the flocs of k drops (k=2,3,...) per unit volume; $a_f^{i,j}$ (i,j = 1, 2, 3,...) are rate constants of flocculation, see Fig. 19; q_k denotes flux of flocs of size k which are products of other processes, different from the *flocculation* itself [say, the reverse process of floc disassembly, or the droplet coalescence, see Eqs. (116) and (120) below]. Analogously to flocculation, the *coalescence* in emulsions can be considered as a kind of irreversible coagulation [176-179]

In the special case of *irreversible* coagulation one has $q_k \equiv 0$. The first term in the righthand side of Eq. (96) is the rate of formation of *k*-flocs by merging of two smaller flocs, whereas the second term expresses the rate of disappearance of *k*-flocs due to their incorporation into larger flocs. The total concentration of flocs (as kinetically independent units), n, and the total concentration of the constituent drops (including those in flocculated form), n_{tot} , are given by the expressions

$$n = \sum_{k=1}^{\infty} n_k, \qquad \qquad n_{\text{tot}} = \sum_{k=1}^{\infty} k n_k \tag{97}$$

The rate constants in Eq. (96) can be expressed in the form

$$a_f^{i,j} = 4\pi D_{i,j}^{(0)} \left(R_i + R_j \right) E_{i,j}$$
(98)

where $D_{i,j}^{(0)}$ is the relative diffusion coefficients for two flocs of radii R_i and R_j , and aggregation number *i* and *j*, respectively; $E_{i,j}$ is the collision efficiency [180,181]. Below we give expressions for $D_{i,j}^{(0)}$ and $E_{i,j}$ applicable to the various types of coagulation.



Fig. 19. Examples for elementary acts of flocculation according to the Smoluchowski scheme; $a_f^{i,j}$ (*i*,*j* = 1, 2, 3, ...,) denote the respective rate constants of flocculation.

The Einstein approach to the theory of diffusivity D gives the following expression,

$$D = kT/B, \qquad B \equiv F/V; \qquad (99)$$

B is friction coefficient, *V* is the velocity acquired by a given particle under the action of an applied net force *F*. For a solid sphere of radius R_0 one has $B = 6\pi\eta R_0$. For a liquid drop *B* is given by the equation of Rybczynski [182] and Hadamar [183]:

$$B = 2\pi\eta R_0 \frac{3\eta_d + 2\eta}{\eta_d + \eta} \tag{100}$$

where η_d is the viscosity inside the drop and η is the viscosity of the continuous phase. The combination of Eqs. (99) and (100) yields the following expression for the relative diffusivity of two isolated Brownian droplets of radii R_i and R_j :

$$D_{i,j}^{(0)} = \frac{kT}{2\pi\eta} \frac{\eta_d + \eta}{3\eta_d + 2\eta} \left(\frac{1}{R_i} + \frac{1}{R_j} \right)$$
 (perikinetic coagulation) (101)

The limiting case $\eta_d \rightarrow 0$ corresponds to two bubbles, whereas in the other limit, $\eta_d \rightarrow \infty$, Eq. (101) describes two solid particles or two liquid drops of tangentially immobile surfaces.

When the drop relative motion is driven by a *body force* or by the *thermocapillary migration* (rather than by selfdiffusion), Eq. (101) is no longer valid. Instead, in Eq. (98) one has to formally substitute the following expression for $D_{i,j}^{(0)}$, see Rogers and Davis [184]

$$D_{i,j}^{(0)} = \frac{1}{4} \left(R_i + R_j \right) |\mathbf{v}_i - \mathbf{v}_j| \qquad \text{(orthokinetic coagulation)} \tag{102}$$

Here \mathbf{v}_j denotes the velocity of a floc of aggregation number *j*. Physically Eq. (102) accounts for the fact that the drops/flocs of different size move with different velocities under the action of the body force. In the case of gravity driven flocculation \mathbf{v}_j is the velocity of a rising/sedimenting particle, which for a drop of tangentially immobile surface is given by the Stokes formula,

$$v_j = v_{bf} \equiv 2g \,\Delta \rho R_j^2 / (9\eta)$$
 (sedimentation velocity) (103)

see e.g. Ref. [16]; here g is the acceleration due to gravity and $\Delta \rho$ is the density difference between the two liquid phases.

In the case of *thermal coagulation*, the drop velocity \mathbf{v}_i is given by the expression [185]

$$\mathbf{v}_{j} = \mathbf{v}_{\text{tm}} \equiv \frac{2R_{j}E_{T}\lambda}{(3\eta_{d} + 2\eta)(\lambda_{d} + 2\lambda)}\nabla(\ln T) \qquad (\text{thermocapillary velocity}) \quad (104)$$

where the thermal conductivity of the continuous and disperse phases are denoted by λ and λ_d ; the interfacial thermal elasticity E_T is defined by Eq. (95).

The *collision efficiency* $E_{i,j}$ in Eq. (98) accounts for the interactions (of both hydrodynamic and intermolecular origin) between two colliding drops. The inverse of $E_{i,j}$ is

called the *stability ratio* or the *Fuchs factor* [186] and can be expressed in the following general form [3,180]

$$\Phi_{i,j} = \frac{1}{E_{i,j}} = 2 \int_{0}^{\infty} \frac{\beta(s)}{(s+2)^2} \exp\left[\frac{W_{i,j}(s)}{kT}\right] ds, \qquad s \equiv \frac{2h}{R_i + R_j}$$
(105)

$$\beta = \left(2\pi\eta a \frac{3\eta_d + 2\eta}{\eta_d + \eta}\right)^{-1} B(s) \tag{106}$$

As usual, *h* is the closest surface-to-surface distance between the two drops; *a* is defined by Eq. (46); $W_{i,j}(s)$ is the energy of *non-hydrodynamic* interactions between the drops, see Eq. (67); $\beta(s)$ accounts for the *hydrodynamic* interactions; B(s) is the drop friction coefficient. For $s \rightarrow \infty$ one obtains $\beta \rightarrow 1$, since for large separations the drops obey the Rybczynski-Hadamar equation (100). In the opposite limit, $s \ll 1$, i.e. close approach of the two drops, $B(s) \equiv F/V$ can be calculated from either Eq. (47), (49), (52) or (56) depending on the specific case. In particular, for $s \ll 1$ one has $\beta \propto s^{-1/2}$ for two spherical droplets of tangentially *mobile* surfaces, whereas $\beta \propto 1/s$ for two drops of tangentially *immobile* surfaces (or two solid particles). In the latter case the integral in Eq. (105) seems to be divergent. To overcome this problem it is usually accepted that for the smallest separations $W_{i,j}$ is dominated by the van der Waals attraction, i.e. $W_{i,j} \rightarrow -\infty$ for $s \rightarrow 0$, and consequently, the integrand in Eq. (105) tends to zero for $s \rightarrow 0$.

The Fuchs factor $\Phi_{i,j}$ is determined mainly by the values of the integrand in a vicinity of the electrostatic maximum (barrier) of $W_{i,j}$, cf. Fig. 6a, since $W_{i,j}$ enters Eq. (105) as an exponent. By using the method of the saddle point Derjaguin [3] estimated the integral in Eq. (105):

$$\Phi_{i,j} \equiv \frac{1}{E_{i,j}} \approx \left[\frac{8\pi kT}{-W_{i,j}''(s_m)}\right]^{1/2} \frac{\beta(s_m)}{(s_m+2)^2} \exp\left[\frac{W_{i,j}(s_m)}{kT}\right]$$
(107)

Here s_m denotes the value of *s* corresponding to the maximum. One sees that the higher the barrier, $W_{i,j}(s_m)$, the smaller the collision efficiency, $E_{i,j}$, and the slower the coagulation.

The infinite set of Smoluchowski equations (96) was solved by Bak and Heilmann [187] in the particular case when the flocs cannot grow larger than a given size; an explicit analytical solution was obtained by these authors.

2. Special results

For imaginary drops, which experience neither long-range surface forces ($W_{i,j}=0$), nor hydrodynamic interactions ($\beta = 1$), Eq. (105) yields a collision efficiency $E_{i,j}=1$ and Eq. (98) reduces to the Smoluchowski [173,174] expression for the rate constant of the *fast irreversible* coagulation. In this particular case Eq. (96) represents an infinite set of nonlinear differential equation. If all flocculation rate constants are the same and equal to a_f , the problem has an exact analytical solution [173,174]:

$$n = \frac{n_0}{1 + a_f n_0 t/2} , \qquad n_k = n_0 \frac{\left(a_f n_0 t/2\right)^{k-1}}{\left(1 + a_f n_0 t/2\right)^{k+1}} , \qquad (k = 1, 2, ...)$$
(108)

The total average concentration of the drops (in both singlet and flocculated form), n_{tot} , does not change and is equal to the initial number of drops, n_0 . Unlike n_{tot} , the concentration of the flocs, n, decreases with time, while their size increases. Differentiating Eq. (108) one obtains

$$\frac{dn}{dt} = -\frac{a_f}{2}n^2, \qquad \frac{d\overline{V}}{dt} = \frac{a_f}{2}\phi_0, \qquad \overline{V} \equiv \frac{\phi_0}{n}$$
(109)

where \overline{V} is the *average volume per floc* and ϕ_0 is the initial volume fraction of the constituent drops. Combining Eqs. (98) and (109) one obtains the following result for *perikinetic* (*Brownian*) coagulation:

$$\frac{\overline{V}}{V_0} = 1 + \frac{t}{t_{\rm Br}}, \qquad t_{\rm Br} = \frac{R_0^2}{3\phi_0 D_0 E_0}$$
(110)

where $V_0 = 4\pi R_0^3/3$ is the volume of a constituent drop of radius R_0 , $t_{\rm Br}$ is the characteristic time of the coagulation process in this case, E_0 is an average collision efficiency and D_0 is an average diffusion coefficient. Equation (110) shows that for fast irreversible coagulation \overline{V} increases linearly with time.

In contrast, \overline{V} is not a linear function of time for *orthokinetic* coagulation, except in the limit of short times. When the flocculation is driven by a *body force*, i.e. in case of sedimentation or centrifugation, one obtains [181]

$$\frac{\overline{V}}{V_0} = \left(1 - \frac{t}{3t_{\rm bf}}\right)^{-3}, \qquad t_{\rm bf} = \frac{2R_0}{3\phi_0 v_{\rm bf} E_0}$$
(111)

where t_{bf} is the characteristic time in this case and v_{bf} is an average velocity of floc motion, which can be expressed by means of Eq. (103) if the body force is the gravitational one.

If the orthokinetic coagulation is driven by the *thermocapillary migration*, the counterpart of Eq. (111) reads [181]

$$\frac{\overline{V}}{V} = \exp\left(\frac{t}{t_{\rm tm}}\right), \qquad t_{\rm tm} = \frac{2R_0}{3\phi_0 v_{\rm tm}E_0} , \qquad (112)$$

where v_{tm} is an average velocity of thermocapillary migration, see Eq. (104), and t_{tm} is the respective characteristic time. Note that $D_0 \propto R_0^{-1}$, $v_{bf} \propto R_0^2$ and $v_{tm} \propto R_0$, cf. Eqs. (99) and (104). Then from Eqs. (110)-(112) it follows that the three different characteristic times exhibit different dependencies on drop radius: $t_{Br} \propto R_0^3$, $t_{bf} \propto R_0^{-1}$, while t_{tm} is independent of R_0 . Hence, the Brownian coagulation is faster for the smaller drops, the body force induced coagulation is more rapid for the larger drops, whereas the thermocapillary driven coagulation is not sensitive to the drop size.

Using the Stokes-Einstein expression for the diffusivity D_0 and Eq. (110) one obtains

$$t_{\rm Br} = \frac{2\pi\eta R_0^3}{\phi_0 kTE_0}$$
(113)

On the other hand, the combination of Eqs. (103) and (111) yields

$$t_{\rm bf} = \frac{3\eta}{\phi_0 R_0 E_0 g \Delta \rho} \tag{114}$$

Let us consider the quantity

$$\chi(R_0) = \frac{t_{\rm bf}}{t_{\rm bf} + t_{\rm Br}} = \frac{1}{1 + 3R_0^4 / (5R_{\rm cr}^4)}, \qquad R_{\rm cr} = \left(\frac{9kT}{10\pi g\Delta\rho}\right)^{1/4}$$
(115)

For $R_0 < R_{cr}$ Eq. (115) yields $\chi(R_0) \approx 1$, that is $t_{Br} \ll t_{bf}$ and the Brownian flocculation is much faster than the orthokinetic flocculation. In contrast, for $R_0 > R_{cr}$ Eq. (115) yields $\chi(R_0) \approx 0$, that is $t_{bf} \ll t_{Br}$ and the orthokinetic flocculation is much more rapid than the Brownian flocculation. At $R_0 = R_{cr}$ a sharp transition from Brownian to orthokinetic flocculation takes place; R_{cr} corresponds to the inflection point of the dependence $\chi = \chi(R_0)$. Since the orthokinetic flocculation happens through a directional motion of the particles, then R_{cr} can be considered as a *threshold radius of the flocs* needed for the creaming (or sedimentation) to begin. With $\Delta \rho = 0.1$ g/cm³ and T = 298 K from Eq. (115) one calculates $R_{cr} = 1.05$ µm. It turns out that the threshold size for creaming is around 1 μ m. This conclusion is consonant with the experimental data in Fig. 3, which show that emulsions with $2R_0 = 5 \mu$ m do cream, whereas those with $2R_0 = 0.35 \mu$ m do not.



Fig. 20. (a) Elementary act of splitting of a floc, containing i+j constitutive drops, to two smaller flocs containing, respectively, *i* and *j* constitutive drops. (b) The coalescence transforms a floc composed of *k* drops into a floc containing *i* drops (i < k). The rate constants of the respective processes are $a_f^{i,j}$ and $a_c^{k,i}$ (*i*, *j*, k = 1, 2, 3, ...).

C. Kinetics of Reversible Flocculation

If the dept of the primary minimum (that on the left from the maximum in Fig. 6a) is not so great, i.e. the attractive force which keeps the drops together is weaker, then the formed flocs are labile and can disassemble into smaller aggregates. This is the case of *reversible flocculation* [3]. For example, a floc composed of *i*+*j* drops can be split on two flocs containing *i* and *j* drops. We denote the rate constant of this *reverse* process by $a_r^{i,j}$ (see Fig. 20a). In the present case both the straight process of flocculation (Fig. 19) and the reverse process (Fig. 20a) take simultaneously place. The kinetics of aggregation in this more general and complex case is described by the Smoluchowski set of equations, Eq. (96), where one is to substitute

$$q_{1} = \sum_{i=1}^{\infty} a_{r}^{1,i} n_{i+1}, \quad q_{k} = \sum_{i=1}^{\infty} a_{r}^{k,i} n_{i+k} - \frac{1}{2} n_{k} \sum_{i=1}^{k-1} a_{r}^{i,k-i} \quad (k = 2,3,...)$$
(116)

Here q_k is the rate of formation of *k*-flocs in the process of disassembly of larger flocs minus the rate of decay of the *k*-flocs. As before, the total number of constituent drops, n_{tot} , does not change. However, the total number of the flocs, *n*, can either increase or decrease depending on whether the straight or the reverse process prevails. Summing up all equations in (96) and using Eq. (116) one derives the following equation for *n*:

$$\frac{dn}{dt} = \frac{1}{2} \sum_{i=1}^{\infty} \sum_{j=1}^{\infty} \left(a_r^{i,j} n_{i+j} - a_f^{i,j} n_i n_j \right)$$
(117)

A general expression for the rate constants of the reverse process was obtained by Martinov and Muller [188]:

$$a_r^{i,j} = \frac{D_{i,j}^{(0)} E_{i,j}}{Z_{i,j}} \frac{1}{\left(R_i + R_j\right)^2}$$
(118)

Here $Z_{i,j}$ is the so called *irreversible factor*, which is defined as follows

$$Z_{i,j} = \frac{1}{8} \int_{W_{i,j} < 0} (s+2)^2 \exp\left[-\frac{W_{i,j}(s)}{kT}\right] ds$$
(119)

The integration in Eq. (119) is carried out over the region around the primary minimum, where $W_{i,j}$ takes negative values, cf. Fig. 6a. In other words, $Z_{i,j}$, is determined by the values of $W_{i,j}$ in the region of the primary *minimum*, whereas $E_{i,j}$ is determined by the values of $W_{i,j}$ in the region of the electrostatic *maximum*, cf. Eqs. (107) and (119). When the minimum is deeper, $Z_{i,j}$ is larger and the rate constant in Eq. (118) is smaller. Moreover, Eqs. (107) and (118) show that the increase of the height of the barrier also decreases the rate of the reverse process. The physical interpretation of this fact is the following: to detach a drop from a floc, the drop has to first emerge from the well and then to "jump" over the barrier (cf. Fig. 6a).

As an illustration in Fig. 21 we show theoretical curves about the rate of flocculation calculated in Ref. [62]. The curves are computed by solving numerically the set of Eqs. (96), (116) and (117). To simplify the problem the following assumptions have been used [62]: (i) the Smoluchowski assumption that all rate constants of the *straight* process are equal to a_{f} ; (ii) flocs, containing more than M drops cannot decay; (iii) all rate constants of the *reverse* process are equal to a_r ; (iv) at the initial moment only single constituent drops of concentration n_0 are available. In Fig. 21 we present the calculated curves of n_0/n , vs. the dimensionless time, $\tau = a_f n_0 t/2$, for a fixed value M = 4 and various values of the ratio of the rate constants of the

straight and the reverse process, $u = 2a_r/(n_0a_f)$. Note that *n* is defined by Eq. (97). The increase of n_0/n with time means that the concentration *n* of the flocs decreases; i.e. the emulsion contains smaller number of flocs, but their size is larger. Consequently, larger n_0/n corresponds to a larger degree of flocculation. It is seen that for the short times of flocculation ($\tau \rightarrow 0$) all curves in Fig. 21 touch the Smoluchowski distribution (corresponding to u = 0), but for the longer times one observes a reduction in the degree flocculation, which smaller for the curves with larger values of *u* (larger rate constants of the reverse process). The "*S-shaped*" curves in Fig. 21 are typical for the case of *reversible* flocculation; curves of similar shape have been obtained experimentally [3,168,189].



Fig. 21. Plot of the inverse dimensionless concentration of flocs, n_0/n , vs. the dimensionless time, $\tau = a_f n_0 t/2$, for M = 4 and various values of the dimensionless ratio $u = 2a_r/(n_0a_f)$; a_r and a_f are the rate constants of the reverse and the straight process. Theoretical curves for reversible flocculation from Ref. [62].

D. Kinetics of Simultaneous Flocculation and Coalescence

In the case of *pure* flocculation considered above the total number of constituent drops, n_{tot} , does not change, see Eq. (97). In contrast, if coalescence is present, in addition to the flocculation, then n_{tot} decreases with time [6]. Hartland and Gakis [190], and Hartland and Vohra [191] developed a model of coalescence, which relates the life-time of single films to the rate of phase separation in emulsions of comparatively large drops (c.a. 1 mm) in the *absence*

of surfactant. The effect of surfactant (emulsifier) was taken into account by Lobo *et al.* [192], who quantified the process of coalescence within an already creamed or settled emulsion containing drops of size less than 100 μ m. Danov *et al.* [175] generalized the Smoluchowski scheme of flocculation to account for the fact that the droplets within the flocs can coalesce to give larger droplets, as illustrated in Fig. 20b. In this case, in the right-hand side of Eq. (96) one has to substitute [175]

$$q_1 = \sum_{i=2}^{\infty} a_c^{i,1} n_i, \qquad q_k = \sum_{i=k+1}^{\infty} a_c^{i,k} n_i - n_k \sum_{i=1}^{k-1} a_c^{k,i} \qquad (k = 2,3,...)$$
(120)

where $a_c^{k,i}$ is the rate constant of transformation (by coalescence) of a floc containing k droplets into a floc containing *i* droplets (see Fig. 20b). The resulting floc is further involved in the flocculation scheme, which thus describes the interdependence of flocculation and coalescence. In this scheme the total coalescence rate, $a_{c,tot}^i$, and the total number of droplets, n_{tot} , are related as follows [175]

$$\frac{dn_{\text{tot}}}{dt} = -\sum_{i=2}^{\infty} a_{c,\text{tot}}^{i} n_{i}, \qquad a_{c,\text{tot}}^{i} = \sum_{k=1}^{i-1} (i-k) a_{c}^{i,k} \qquad (i=2,3,...)$$
(121)

To determine the rate constants of coalescence, $a_c^{k,i}$, Danov *et al.* [147] examined the effects of the droplet interactions and the Brownian motion on the coalescence rate in dilute emulsions of μ m and sub- μ m sized droplets. The processes of film formation, thinning, and rupture were included as consecutive stages in the scheme of coalescence. Expressions for the interaction energy due to various DLVO and non-DLVO surface forces between two deformed droplets were obtained [143].

Average models for the total number of droplets have been also proposed [193,194]. The average model of van den Tempel [193] assumes linear structure of the flocs. The coalescence rate is supposed to be proportional to the number of contacts within a floc. To simplify the problem van den Tempel has used several assumptions, one of them being that the concentration of the single droplets, n_1 , obeys the Smoluchowski distribution, Eq. (108) for k=1. The model of Borwankar *et al.* [194] employs some assumptions, which make it more applicable to cases, in which the flocculation (rather than the coalescence) is slow and is the rate determining stage. This is illustrated by the curves shown in Fig. 22, which are calculated for the same rate of *coalescence*, but for two different rates of *flocculation*. For relatively high rates of flocculation (Fig. 22a) the predictions of the three theories differ, but the model of

Borwankar *et al.* [194] gives values closer to that of the more detailed model by Danov *et al.* [175] For very low values of the flocculation rate constant, a_f , for which the coalescence is not the rate determining stage, all three theoretical models, Refs. [175,193,194], give results for n_{tot}/n_0 vs. time, which almost coincide numerically (Fig. 22b).

Finally, it is worthwhile noting that the simultaneous flocculation and coalescence in emulsions could be also accompanied with adsorption of amphiphilic molecules on the drop surfaces [195]; this possibility should be kept in mind when interpreting experimental data.



Fig. 22. The total number of constituent drops in a flocculating emulsion, n_{tot} , decreases with time, t, because of a parallel process of coalescence. The curves are calculated for the following parameter values: initial number of constituent drops $n_0=10^{12} \text{ cm}^{-3}$; coalescence rate constant $k_c^{2,1} = 10^{-3} \text{ s}^{-1}$. Curve 1 is numerical solution of Eq. (121). Curves 2 and 3 are the result predicted by the models of Borwankar *et al.* [194] and van den Tempel [193], respectively. The values of the flocculation rate constant are: (a) $a_f = 10^{-11} \text{ cm}^3/\text{s}$; (b) $a_f = 10^{-16} \text{ cm}^3/\text{s}$.

VII. SUMMARY

The surfactants play a crucial role in emulsification and emulsion stability. A first step in any quantitative study on emulsions should be to determine the equilibrium and dynamic properties of the oil-water interface, such as interfacial tension, Gibbs elasticity, surfactant adsorption, counterion binding, surface electric potential, adsorption relaxation time, etc. Useful theoretical concepts and expressions, which are applicable to ionic, nonionic and micellar surfactant solutions, are summarized in Section II.

The emulsion drops in flocs and creams are separated with thin liquid films, whose rupture leads to coalescence and phase separation. At equilibrium the area of the films and their contact angle are determined by the surface forces (disjoining pressure) acting across the films (Section III.A.1). Several ways of breakage of these emulsion films have been established: capillary-wave mechanism, pore-nucleation mechanism, solute-transport mechanism, barrier mechanism, etc. (Section III.A.2).

Experimental and theoretical results evidence that the capillary wave mechanism is the most frequent reason for the coalescence of both deformed and spherical emulsion drops. For a certain critical thickness (width), h_c , of the film (gap) between two emulsion drops the amplitude of the thermally excited fluctuation capillary waves begins to grow, promoted by the surface forces, and causes film rupture. The capillary waves can bring about coalescence of two spherical emulsion drops, when the distance between them becomes smaller than a certain critical value, which is estimated to be about 10 - 50 nm (see Section III.C).

The interactions of two emulsion drops, and their theoretical description, become more complicated if the drops are moving against each other, instead of being quiescent. In such a case, which happens most frequently in practice, the hydrodynamic interactions come into play (Section IV). The velocity of approach of two drops and the critical distance (thickness) of drop coalescence are influenced by the drop size, disjoining pressure, bulk and surface diffusivity of surfactant, Gibbs elasticity, surface viscosity, etc. If attractive (negative) disjoining pressure prevails, then "pimples" appear on the opposite drop surfaces in the zone of contact; thus the drop coalescence can be produced by growth and merging of these "pimples" (Section IV.A.3). Alternatively, the drop coalescence can be produced by the growth of fluctuation capillary waves; the theory of the respective critical thickness is found to agree excellently with available experimental data (Section IV.B.3).

The finding that the hydrodynamic velocity of mutual approach of two emulsion drops is much higher when the surfactant is dissolved in the drop phase (rather than in the continuous phase) provides a natural explanation of the Bancroft rule in emulsification (Section V). A generalized version of the Bancroft rule is proposed, Eqs. (89) and (91), which takes into account the role of various thermodynamic and hydrodynamic factors. For example, the existence of considerable repulsive (positive) disjoining pressure may lead to exclusions from the conventional Bancroft rule, which are accounted for in its generalized version.

The knowledge about the individual acts of drop-drop collision is a prerequisite for development of a kinetic theory of such collective phenomena as flocculation/coalescence and phase separation. The cases of fast and slow, perikinetic and orthokinetic, irreversible and reversible flocculation are considered in Section VI. Special attention is paid to the case of parallel flocculation and coalescence. Much work remains to be done in order to built up an united theory including both individual drop interactions and collective phenomena in emulsions.

Acknowledgments. Financial Support from Inco-Copernicus project No. IC15 CT98 0911 of the European Commission is gratefully acknowledged. The authors are indebted to Ms. Mariana Paraskova and Mr. Vesselin Kolev for their help in the preparation of the figures.

REFERENCES

- 1. I. B. Ivanov and P. A. Kralchevsky, Colloids Surf. A, 128: 155 (1997).
- 2. I. B. Ivanov, K. D. Danov, and P. A. Kralchevsky, Colloids Surf. A, 152: 161 (1999).
- 3. B. V. Derjaguin, *Theory of Stability of Colloids and Thin Films*, Plenum Press: Consultants Bureau, New York, 1989.
- 4. J. N. Israelachvili, Intermolecular & Surface Forces, Academic Press, London, 1992.
- V. M. Mikova, *Investigation of Emulsions Stabilized by β-Lactoglobulin*, M. Sc. Thesis, (T. D. Gurkov, Supervisor), Faculty of Chemistry, University of Sofia, 1999.
- 6. J. T. Davies and E. K. Rideal, *Interfacial Phenomena*, Academic Press, London, 1963.
- 7. P. C. Hiemenz and R. Rajagopalan, *Principles of Colloid and Surface Chemistry*, Marcel Dekker, New York, 1997.
- 8. I. Langmuir, J. Amer. Chem. Soc. 15: 75 (1918).
- 9. T. L. Hill, *An Introduction to Statistical Thermodynamics*, Addison-Wesley, Reading, MA, 1962.
- 10. M. Volmer, Z. Physikal. Chem. 115: 253 (1925).
- 11. A. Frumkin, Z. Physikal. Chem. 116: 466 (1925).
- 12. L. D. Landau and E. M. Lifshitz, *Statistical Physics, Part 1*, Pergamon Press, Oxford, 1980.
- 13. T. D. Gurkov, P. A. Kralchevsky, and K. Nagayama, Colloid Polym. Sci. 274: 227 (1996).
- 14. E. H. Lucassen-Reynders, J. Phys. Chem. 70: 1777 (1966).
- 15. R. P. Borwankar and D. T. Wasan, Chem. Eng. Sci. 43: 1323 (1988).
- 16. E. D. Shchukin, A. V. Pertsov, and E. A. Amelina, *Colloid Chemistry*, Moscow University Press, Moscow, 1982 [in Russian].
- 17. P. A. Kralchevsky, K. D. Danov, G. Broze, and A. Mehreteab, Langmuir 15: 2351 (1999).
- 18. J. W. Gibbs, The Scientific Papers of J. W. Gibbs, vol.1, Dover, New York, 1961.
- 19. E. H. Lucassen-Reynders, in *Encyclopedia of Emulsion Technology, Vol. 4* (P. Becher, Ed.), Marcel Dekker, New York, 1996, p. 63.
- 20. Y. Tian, R. G. Holt, and R. Apfel, J. Colloid Interface Sci. 187: 1 (1997).
- 21. A. Ozawa, A. Minamisawa, K. Sakai, and K. Takagi, *Jpn. J. Appl. Phys.* 33: L-1468 (1994).
- 22. K. Sutherland, Austr. J. Sci. Res. Ser. A 5: 683 (1952).
- 23. S. S. Dukhin, G. Kretzschmar, and R. Miller, *Dynamics of Adsorption at Liquid Interfaces*; Elsevier, Amsterdam, 1995.

- 24. K. D. Danov, P. A. Kralchevsky, and I. B. Ivanov, in *Handbook of Detergents* (G. Broze, Ed.), Marcel Dekker, New York, 1999, p. 303.
- 25. J. Th. G. Overbeek, in *Colloid Science, Vol. 1* (H. R. Kruyt, Ed.), Elsevier, Amsterdam, 1953; *J. Colloid Sci. 8*: 420 (1953).
- 26. G. Gouy, J. Phys. Radium 9: 457 (1910).
- 27. D. L. Chapman, Phil. Mag. 25: 475 (1913).
- 28. O. Stern, Ztschr. Elektrochem. 30: 508 (1924).
- 29. S. Hachisu, J. Colloid Interface Sci. 33: 445 (1970).
- D. G. Hall, in *The Structure, Dynamics and Equilibrium Properties of Colloidal Systems*, (D. M. Bloor and E. Wyn-Jones, Eds.) Kluwer, Dordrecht, 1990; p. 857.
- 31. D. G. Hall, Colloids Surf. A, 90: 285 (1994).
- 32. G. Bakker, Kapillarität und Oberflächenspannung, in: *Handbuch der Experimentalphysik, Band 6*, Akademische Verlagsgesellschaft, Leipzig, 1928.
- 33. S. Ono, S. Kondo, Molecular Theory of Surface Tension in Liquids, in: *Handbuch der Physik*, vol. 10 (S. Flügge, Ed.), Springer, Berlin, 1960; p. 134.
- 34. J. S. Rowlinson and B. Widom, *Molecular Theory of Capillarity*, Clarendon Press, Oxford, 1982.
- 35. K. D. Danov, P. M. Vlahovska, P. A. Kralchevsky, G. Broze, and A. Mehreteab, *Colloids Surf. A*, 156: 389 (1999).
- 36. K. D. Danov, V. L. Kolev, P. A. Kralchevsky, G. Broze, and A. Mehreteab, *Langmuir 16*: 2942 (2000).
- 37. V. V. Kalinin, and C. J. Radke, Colloids Surf. A, 114: 337 (1996).
- 38. K. Tajima, M. Muramatsu, and T. Sasaki, Bul. Chem. Soc. Japan 43: 1991 (1970).
- 39. K. Tajima, Bul. Chem. Soc. Japan 43: 3063 (1970).
- 40. E. Janke, F. Emde, and F. Lösch, F. *Tables of Higher Functions*, McGraw-Hill, New York, 1960.
- 41. M. Abramowitz and I. A. Stegun, *Handbook of Mathematical Functions*, Dover, New York, 1965.
- 42. G. A. Korn and T. M. Korn, Mathematical Handbook, McGraw-Hill, New York, 1968.
- 43. G. Loglio, E. Rillaerts, and P. Joos, Colloid Polym. Sci. 259: 1221 (1981).
- 44. R. Miller, Colloid Polym. Sci. 259: 375 (1981).
- 45. Y. M. Rakita, V. B. Fainerman, and V. M. Zadara, Zh. Fiz. Khim. 60: 376 (1986).
- 46. P. A. Kralchevsky, Y. S. Radkov, and N. D. Denkov, J. Colloid Interface Sci. 161: 361 (1993).

- 47. K. D. Danov, P. M. Vlahovska, T. S. Horozov, C. D. Dushkin, P. A. Kralchevsky, A. Mehreteab, and G. Broze, *J. Colloid Interface Sci.* 183: 223 (1996).
- 48. R. S. Hansen, J. Chem. Phys. 64: 637 (1960).
- 49. R. van den Bogaert and P. Joos, J. Phys. Chem. 83: 2244 (1979).
- 50. E. Rillaerts and P. Joos, J. Chem. Phys. 96: 3471 (1982).
- 51. R. Miller and G. Kretzschmar, Adv. Colloid Interface Sci. 37: 97 (1991).
- 52. V. B. Fainerman, A. V. Makievski, R. Miller, R. Colloids Surf. A, 87: 61 (1994).
- 53. L. K. Filippov, J. Colloid Interface Sci. 164: 471 (1994).
- 54. A. Bonfillon, F. Sicoli, and D. Langevin, J. Colloid Interface Sci. 168: 497 (1994).
- 55. C. MacLeod and C. J. Radke, Langmuir 10: 2965 (1994).
- 56. A. W. Cross, G. G. Jayson, J. Colloid Interface Sci. 162: 45 (1994).
- 57. S. B. Johnson, C. J. Drummond, P. J. Scales, S. Nishimura, Langmuir 11: 2367 (1995).
- 58. R. G. Alargova, K. D. Danov, J. T. Petkov, P. A. Kralchevsky, G. Broze, A. Mehreteab, *Langmuir 13*: 5544 (1997).
- 59. O. D. Velev, T. D. Gurkov, and R. P. Borwankar, J. Colloid Interface Sci. 159: 497 (1993).
- 60. K. D. Danov, T. D. Gurkov, T. D. Dimitrova, and D. Smith, *J. Colloid Interface Sci. 188*: 313 (1997).
- 61. O. D. Velev, T. D. Gurkov, I. B. Ivanov, and R. P. Borwankar, *Phys. Rev. Lett.* 7: 264 (1995).
- 62. P. A. Kralchevsky, K. D. Danov, and N.D. Denkov, Chapter 11 in "*Handbook of Surface and Colloid Chemistry*" (K. S. Birdi, Ed.), CRC Press, Boca Raton, 1997; p. 333.
- 63. M. J. Rosen, Surfactants and Interfacial Phenomena, Wiley, New York, 1989.
- 64. J. Clint, Surfactant Aggregation, Chapman & Hall, London, 1992.
- 65. E. A G. Aniansson and S. N. Wall, J. Phys. Chem. 78: 1024 (1974).
- 66. E. A. G. Aniansson and S. N. Wall, J. Phys. Chem. 79: 857 (1975).
- 67. E. A. G. Aniansson and S. N. Wall, in: *Chemical and Biological Applications of Relaxation Spectrometry* (E. Wyn-Jones, Ed.), Reidel, Dordrecht, 1975; p. 223.
- 68. E. A. G. Aniansson, S. N. Wall, M. Almgren, H. Hoffmann, I. Kielmann, W. Ulbricht, R. Zana, J. Lang, and C. Tondre, *J. Phys. Chem.* 80: 905 (1976).
- 69. C. D. Dushkin and I. B. Ivanov, Colloids Surf. 60: 213 (1991).
- 70. C. D. Dushkin, I. B. Ivanov, and P. A. Kralchevsky, Colloids Surf. 60: 235 (1991).
- 71. B. A. Noskov, Kolloidn Zh. 52: 509 (1990).
- 72. B. A. Noskov, Kolloidn Zh. 52: 796 (1990).
- 73. G. C. Kresheck, E. Hamori, G. Davenport, and H. A. Scheraga, *J. Am. Chem. Soc.* 88: 264 (1966).
- 74. P. A. Kralchevsky and I. B. Ivanov, Chem. Phys. Lett., 121: 111 (1985).
- 75. I. B. Ivanov and P. A. Kralchevsky, in *Thin Liquid Films* (I. B. Ivanov, Ed.), Marcel Dekker, New York, 1988; p. 49.
- 76. I. B. Ivanov and B. V. Toshev, Colloid. Polym. Sci. 253: 593 (1975).
- 77. J. A. de Feijter and A. Vrij, J. Colloid Interface Sci. 70: 456 (1979).
- 78. P. A. Kralchevsky and N. D. Denkov, Chem. Phys. Lett. 240: 385 (1995).
- 79. A. D. Nikolov, D. T. Wasan, P. A. Kralchevsky, and I. B. Ivanov, in: *Ordering and Organisation in Ionic Solutions* (N. Ise and I. Sogami, Eds.), World Scientific, Singapore, 1988; p. 302.
- 80. A. D. Nikolov and D. T. Wasan, J. Colloid Interface Sci. 133: 1 (1989).
- 81. A. D. Nikolov, P. A. Kralchevsky, I. B. Ivanov, and D. T. Wasan, *J. Colloid Interface Sci. 133*: 13 (1989).
- 82. P. A. Kralchevsky, A. D. Nikolov, D. T. Wasan, and I. B. Ivanov, *Langmuir 6*: 1180 (1990).
- 83. A. D. Nikolov, D. T. Wasan, N. D. Denkov, P. A. Kralchevsky, and I. B. Ivanov, *Prog. Colloid Polym. Sci.* 82: 87 (1990).
- 84. D. T. Wasan, A. D. Nikolov, P. A. Kralchevsky, and I. B. Ivanov, *Colloids Surf.* 67: 139 (1992).
- 85. V. Bergeron and C. J. Radke, Langmuir 8: 3020 (1992).
- 86. J. L. Parker, P. Richetti, P. Kékicheff, and S. Sarman, Phys. Rev. Lett. 68: 1955 (1992).
- 87. M. L. Pollard and C. J. Radke, J. Chem. Phys. 101: 6979 (1994).
- 88. X. L. Chu, A. D. Nikolov, and D. T. Wasan, Langmuir 10: 4403 (1994).
- 89. X. L. Chu, A. D. Nikolov, and D. T. Wasan, J. Chem. Phys. 103: 6653 (1995).
- 90. E. S. Basheva, K. D. Danov, and P. A. Kralchevsky, Langmuir 13: 4342 (1997).
- 91. K. G. Marinova, T. D. Gurkov, T. D. Dimitrova, R. G. Alargova, and D. Smith, *Langmuir* 14: 2011 (1998).
- 92. A. J. Vries, Rec. Trav. Chim. Pays-Bas 77: 44 (1958).
- 93. A. Scheludko, Proc. K. Akad. Wetensch. B, 65: 87 (1962).
- 94. A. Vrij, Disc. Faraday Soc. 42: 23 (1966).
- 95. I. B. Ivanov, B. Radoev, E. Manev, and A. Scheludko, *Trans. Faraday Soc.* 66: 1262 (1970).
- 96. I. B. Ivanov and D. S. Dimitrov, Colloid Polymer Sci. 252: 982 (1974).

- 97. I. B. Ivanov, Pure Appl. Chem. 52: 1241 (1980).
- 98. P. A. Kralchevsky, K. D. Danov, and I. B. Ivanov, Thin Liquid Film Physics, in: *Foams* (R. K. Prud'homme and S. A. Khan, Eds.), Marcel Dekker, New York, 1995; p.1.
- 99. B. V. Derjaguin and Y. V. Gutop, Kolloidn. Zh. 24: 431 (1962).
- 100. B. V. Derjaguin and A. V. Prokhorov, J. Colloid Interface Sci. 81: 108 (1981).
- 101. A. V. Prokhorov and B. V. Derjaguin, J. Colloid Interface Sci. 125: 111 (1988).
- 102. D. Kashchiev and D. Exerowa, J. Colloid Interface Sci. 77: 501 (1980).
- 103. D. Kashchiev and D. Exerowa, Biochim. Biophys. Acta 732: 133 (1983).
- 104. D. Kashchiev, Colloid Polymer Sci. 265: 436 (1987).
- 105. Y. A. Chizmadzhev and V. F. Pastushenko, in: *Thin Liquid Films* (I. B. Ivanov, Ed.), M. Dekker, New York, 1988; p. 1059.
- 106. L. V. Chernomordik, M. M. Kozlov, G. B. Melikyan, I. G. Abidor, V. S. Markin, and Y. A. Chizmadzhev, *Biochim. Biophys. Acta* 812: 643 (1985).
- 107. L. V. Chernomordik, G. B. Melikyan, and Y. A. Chizmadzhev, *Biochim. Biophys. Acta* 906: 309 (1987).
- 108. A. Kabalnov and H. Wennerström, Langmuir 12 (1996) 276.
- 109. I. B. Ivanov, S. K. Chakarova, and B. I. Dimitrova, Colloids Surf. 22: 311 (1987).
- 110. B. I. Dimitrova, I. B. Ivanov, and E. Nakache, J. Dispers. Sci. Technol. 9: 321 (1988).
- 111. K. D. Danov, I. B. Ivanov, Z. Zapryanov, E. Nakache, and S. Raharimalala, in: *Proceedings of the Conference of Synergetics, Order and Chaos* (M. G. Velarde, Ed.), World Scientific, Singapore, 1988; p. 178.
- 112. C. V. Sterling and L. E. Scriven, AIChE J. 5: 514 (1959).
- 113. S. P. Lin and H. J. Brenner, J. Colloid Interface Sci. 85: 59 (1982).
- 114. T. D. Gurkov, K. D. Danov, N. Alleborn, H. Raszillier, and F. Durst, J. Colloid Interface Sci. 198: 224 (1998).
- 115. J. Lucassen, van den Temple, A. Vrij, and F. Th. Hesselink, *Proc. Konkl. Ned. Akad. Wet. B73*: 109 (1970).
- 116. A. K. Malhotra and D. T. Wasan, Chem. Eng. Commun. 48: 35 (1986).
- 117. Ch. Maldarelli and R. K. Jain, in: *Thin Liquid Films: Fundamental and Applications* (I. B. Ivanov, Ed.), M. Dekker, New York, 1988; p. 497.
- 118. B. U. Felderhof, J. Chem. Phys. 49: 44 (1968).
- 119. S. Sche and H. M. Fijnaut, Surface Sci. 76: 186 (1978).
- 120. E. D. Manev, S. V. Sazdanova, and D. T. Wasan, J. Colloid Interface Sci. 97: 591 (1984).

- 121. I. B. Ivanov, and D. S. Dimitrov, in *Thin Liquid Films* (I. B. Ivanov, Ed.), M. Dekker, New York, 1988, p. 379.
- 122. P. Taylor, Proc. Roy. Soc. (London) A108: 11 (1924).
- 123. E. Rushton, G.A. Davies, Appl. Sci. Res. 28: 37 (1973).
- 124. S. Haber, G. Hetsroni, and A. Solan, Int. J. Multiphase Flow 1: 57 (1973).
- 125. L. D. Reed and F. A. Morrison, Int. J. Multiphase Flow 1: 573 (1974).
- 126. G. Hetsroni and S. Haber, Int. J. Multiphase Flow 4: 1 (1978).
- 127. F. A. Morrison, L. D. Reed, Int. J. Multiphase Flow 4: 433 (1978).
- 128. V. N. Beshkov, B. P. Radoev, and I. B. Ivanov, Int. J. Multiphase Flow 4: 563 (1978).
- 129. D. J. Jeffrey and Y. Onishi, J. Fluid Mech. 139: 261 (1984).
- 130. Y. O. Fuentes, S. Kim, and D. J. Jeffrey, Phys. Fluids 31: 2445 (1988).
- 131. R. H. Davis, J. A. Schonberg, and J. M. Rallison, Phys. Fluids A1: 77 (1989).
- 132. X. Zhang and R. H. Davis, J. Fluid Mech. 230: 479 (1991).
- 133. T. T. Traykov and I. B. Ivanov, Int. J. Multiphase Flow 3: 471 (1977).
- 134. T. T. Traykov, E. D. Manev, and I. B. Ivanov, Int. J. Multiphase Flow 3: 485 (1977).
- 135. I. B. Ivanov, B. P. Radoev, T. Traykov, D. Dimitrov, E. Manev, and C. Vassilieff, in: *Proceedings of the International Conference on Colloid and Surface Science* (E. Wolfram, Ed.), Vol. 1, Akademia Kiado, Budapest, 1975, p. 583.
- 136. K. D. Danov and I. B. Ivanov, Critical Film Thickness and Coalescence in Emulsions, *Proc. Second World Congress on Emulsions*, Paris, 1997, Paper No. 2-3-154.
- 137. K. D. Danov, D. S. Valkovska, and I. B. Ivanov, J. Colloid Interface Sci. 211: 291 (1999).
- 138. D. S. Valkovska, K. D. Danov, and I. B. Ivanov, Colloids Surf. A, 156: 547 (1999).
- 139. S. G. Yiantsios and R. H. Davis, J. Colloid Interface Sci. 144: 412 (1991).
- 140. V. Cristini, J. Blawzdziewicz, and M. Loewenberg, J. Fluid Mech. 366: 259 (1998).
- 141. D. S. Valkovska, K. D. Danov, and I. B. Ivanov, Colloids Surf. A 156: 547 (1999).
- 142. O. Reynolds, Phil. Trans. Roy. Soc (London) A177: 157 (1886).
- 143. K. D. Danov, D. N. Petsev, N. D. Denkov, and R. Borwankar, J. Chem. Phys. 99: 7179 (1993).
- 144. N. D. Denkov, P. A. Kralchevsky, I. B. Ivanov, and C. S. Vassilieff, *J. Colloid Interface Sci. 143*: 157 (1991).
- 145. N. D. Denkov, D. N. Petsev, and K. D. Danov, Phys. Rev. Lett. 71: 3327 (1993).
- 146. D. N. Petsev, N. D. Denkov, and P. A. Kralchevsky, J. Colloid Interface Sci. 176: 201 (1995).

- 147. K. D. Danov, N. D. Denkov, D. N. Petsev, and R. Borwankar, *Langmuir 9*: 1731 (1993).
- 148. A. Scheludko and D. Exerowa, Kolloid-Z. 165: 148 (1959).
- 149. A. Scheludko, Adv. Colloid Interface Sci. 1: 391 (1967).
- 150. O. D. Velev, A. D. Nikolov, N. D. Denkov, G. Doxastakis, V. Kiosseoglu, G. Stalidis, *Food Hydrocolloids*, 7: 55 (1993).
- 151. O. D. Velev, T. D. Gurkov, S. K. Chakarova, B. I. Dimitrova, I. B. Ivanov and R. P. Borwankar, *Colloids & Surfaces A*, 83: 43 (1994).
- 152. O. D. Velev, G. N. Constantinides, D. G. Avraam, A. C. Payatakes, and R. P. Borwankar, *J. Colloid Interface Sci.* 175: 68 (1995).
- 153. K. G. Marinova, T. D. Gurkov, O. D. Velev, I. B. Ivanov, B. Campbell and R. P. Borwankar, *Colloids Surf. A, 123-124*: 155 (1997).
- 154. K. P. Velikov, O. D. Velev, K. G. Marinova, G. N. Constantinides, J. Chem. Soc., Faraday Trans., 93: 2069 (1997).
- 155. O. D. Velev, K. D. Danov and I. B. Ivanov, J. Dispersion Sci. Technology 18: 625 (1997).
- 156. K. G. Marinova, T. D. Gurkov, T. D. Dimitrova, R. G. Alargova and D. Smith, *Progress in Colloid & Polymer Sci.*, 110: 245 (1998).
- 157. O. D. Velev, B. E. Campbell, R. P. Borwankar, Langmuir 14: 4122 (1998).
- 158. T. D. Gurkov, K. G. Marinova, A. Zdravkov, C. Oleksiak, and B. Campbell, *Prog. Colloid* & *Polym. Sci.* 110: 263 (1998).
- 159. B. P. Radoev, D. S. Dimitrov, and I. B. Ivanov, Colloid Polym. Sci. 252: 50 (1974).
- 160. W. B. Russel, D. A. Saville, and W. R. Schowalter, *Colloidal Dispersions*, Cambridge University Press, 1989; p. 155.
- 161. W. D. Bancroft, J. Phys. Chem. 17: 514 (1913).
- 162. B. P. Binks, Langmuir 9: 25 (1993).
- 163. K. D. Danov, O. D. Velev, I. B. Ivanov and R. P. Borwankar, "Bancroft Rule and Hydrodynamic Stability of Thin Films and Emulsions", in: *Proc. First World Congress on Emulsion*, Paris, 1993.
- 164. H. Kunieda, D. F. Evans, C. Solans, and Yoshida, Colloids Surf. A, 47: 35 (1990).
- 165. M. J. Schick, Nonionic Surfactants: Physical Chemistry, M. Dekker, New York, 1986.
- 166. K. Shinoda and S. Friberg, *Emulsions and Solubilization*, Wiley, New York, 1986.
- 167.E. E. Isaacs, H. Huang, and A. J. Babchin, *Colloids Surf.* 46: 177 (1990). *Colloids Surf. A*, 123/124: 195 (1997).
- 168. O. Holt, O. Seather, and N. A. Mishchuk, Colloids Surf. A, 123/124: 195 (1997).
- 169. K. Demetreades and D. J. McClements, Colloids Surf. A, 150: 45 (1999).

- 170. F. Feuillebois, J. Colloid Interface Sci., 131: 267 (1989).
- 171. R. M. Merritt and R. S. Subramanian, J. Colloid Interface Sci. 131: 514 (1989).
- 172. K. D. Barton and R. S. Subramanian, J. Colloid Interface Sci. 133: 214 (1989).
- 173. M. von Smoluchowski, Phys. Z. 17: 557 (1916).
- 174. M. von Smoluchowski, Z. Phys. Chem. 92: 129 (1917).
- 175. K. D. Danov, I. B. Ivanov, T. D. Gurkov, and R. P. Borwankar, J. Colloid Interface Sci. 167: 8 (1994).
- 176. J. L. Klemaszewski, K. P. Das, and J. E. Kinsella, J. Food Sci. 57: 366 (1992).
- 177. Ph. T. Jaeger, J. J. M. Janssen, and W. G. M. Agterof, Colloids Surf. A, 85: 255 (1994).
- 178. E. Dickinson and A. Williams, Colloids Surf. A, 88: 317 (1994).
- 179. A. J. F. Sing, A. Graciaa, and J. L. Salager, Colloids Surf. A, 152: 31 (1999).
- 180. X. Zhang and R. H. Davis, J. Fluid Mech. 230: 479 (1991).
- 181. H. Wang and R. H. Davis, J. Colloid Interface Sci. 159: 108 (1993).
- 182. W. Rybczynski, Bull. Intern. Acad. Sci. (Cracovie) A (1911).
- 183. J. S. Hadamar, Comp. Rend. Acad. Sci. (Paris) 152: 1735 (1911).
- 184. J. R. Rogers and R. H. Davis, Mettal. Trans. A21: 59 (1990).
- 185. N. O. Young, J. S. Goldstein, and M. J. Block, J. Fluid Mech. 6: 350 (1959).
- 186. N. A. Fuchs, Z. Phys. 89: 736 (1934).
- 187. T. A. Bak and O. Heilmann, J. Phys. A: Math. Gen. 24: 4889 (1991).
- 188. G. A. Martinov and V. M. Muller, in *Research in Surface Forces, Vol. 4* (B. V. Derjaguin, Ed.) Plenum Press: Consultants Bureau, New York, 1975; p. 3.
- 189. I. M. Elminyawi, S. Gangopadhyay, and C. M. Sorensen, J. Colloid Interface Sci. 144: 315 (1991).
- 190. S. Hartland and N. Gakis, Proc. Roy. Soc. (London) A369: 137 (1979).
- 191. S. Hartland and D. K. Vohra, J. Colloid Interface Sci. 77: 295 (1980).
- 192. L. A. Lobo, I. B. Ivanov, and D. T. Wasan, AIChE J. 39: 322 (1993).
- 193. M. van den Tempel, *Recueil* 72: 433 (1953).
- 194. R. P. Borwankar, L. A. Lobo, and D. T. Wasan, Colloids Surf. 69: 135 (1992).
- 195. R. Hogg, Colloids Surf. A, 146: 253 (1999).