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1. Introduction

Turbulent emulsification is a fundamentally complex phenomenon where intermittent turbulent structures interact with a viscous interface,¹⁻⁸ typically in the presence of surface-active emulsifiers (added or naturally occurring).⁶⁻⁸ The emulsifiers interact and modify the interface by several different mechanisms, each with its own timescale.⁹⁻¹⁴ Turbulent drop breakup is of considerable industrial interest for the processing of colloidal food systems such as emulsions,¹⁵⁻¹⁷ since it is the dominating mechanism in high-energy density emulsification devices such as high-pressure homogenizers and rotor-stator mixers.^{3,18-22}

Empirical investigations suggest that if the emulsifier is supplied at a sufficiently high concentration, turbulent drop breakup dominates the emulsification process and the resulting emulsion drop diameter is controlled by the physical properties of the two phases, the interfacial tension delivered by the emulsifier (γ) and the dissipation rate of turbulent kinetic energy of the flow (ε), as described by a Weber number, defined as the ratio between disruptive stress and the cohesive

Emulsifier adsorption kinetics influences drop deformation and breakup in turbulent emulsification

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Turbulent drop breakup is of large importance for applications such as food and pharmaceutical processing, as well as of substantial fundamental scientific interest. Emulsification typically takes place in the presence of surface-active emulsifiers (natural occurring and/or added). Under equilibrium conditions, these lower the interfacial tension, enabling deformation and breakup. However, turbulent deformation is fast in relation to emulsifier kinetics. Little is known about the details of how the emulsifier influences drop deformation under turbulent conditions. During the last years, significant insight in the mechanism of turbulent drop breakup has been reached using numerical experiments. However, these studies typically use a highly simplistic description of how the interface responds to turbulent stress. This study investigates how the limited exchange rate of emulsifier between the bulk and the interface influences the deformation process in turbulent drop breakup for application-relevant emulsifiers and concentrations, in the context of state-of-the-art single drop breakup simulations. In conclusion, if the Weber number is high or the emulsifier is supplied at a concentration giving an adsorption time less than 1/10th of the drop breakup time, deformation proceeds as if the emulsifier adsorbed infinitely fast. Otherwise, the limited emulsifier kinetics delays breakup and can alter the breakup mechanism.

Laplace pressure. Assuming that the drop is large in comparison to the smallest eddies of the turbulence, the Weber number is given by,

We =
$$\frac{2 \cdot \rho_{\rm C} \cdot \varepsilon^{2/3} \cdot d_0^{5/3}}{\gamma}$$
, (1)

(where $\rho_{\rm C}$ denotes continuous phase density and d_0 denotes the initial drop diameter). Moreover, the resulting drop diameter can be accurately predicted by a viscosity-modified Kolmogorov–Hinze model.^{7,22–27}

If the emulsifier concentration is below a critical limit, however, re-coalescence dominates²⁸⁻³² and the emulsification process results in the drop diameter which is able to achieve a sufficiently high surface coverage. For an emulsifier soluble in the continuous phase, the critical concentration forming the demarcation between the two regimes is,^{6,7}

$$c^* = \frac{\phi_{\rm D}}{(1 - \phi_{\rm D})} \Gamma^* \cdot \frac{6}{d_{32}},$$
 (2)

where Γ^* is the surface load required for monolayer coverage (the surface load beyond which the interfacial tension does not decrease substantially), ϕ_D is the volume fraction of disperse phase and d_{32} is the final surface-area weighted mean diameter.

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During the last decade, in addition to experimental single drop breakup visualization studies,^{14,33-38} substantial advances in understanding turbulent drop and bubble breakup has been reached using simulation-based techniques. These studies combine direct numerical simulation (DNS) with highly resolved interface tracking.^{10-12,39-46} The simulations based on DNS offer an unprecedented ability to describe the turbulent interaction free of modelling assumptions and at a high spatial and temporal resolution. However, from a colloidal chemistry perspective, the studies are based on rather crude descriptions of how the interface responds to stress and how this is related to the presence of an emulsifier. Typically, the interface is described using an interfacial tension, assumed to be uniform and constant in time. This is equivalent to assuming that the exchange of emulsifier between bulk and interface is infinitely fast,⁴¹ *i.e.*, as soon as the interface starts to deform due to a turbulent interaction, adsorption is fast enough to restore the original surface load instantly. Moreover, this implies that the state-of-the-art in these simulations neglect adsorption kinetics, and all effects of interfacial rheology. Some numerical studies have been performed including the Marangoni effects resulting from an uneven distribution of surface-active material, however only for a highly idealized system.^{47,48}

It is problematic that the numerical literature on drop breakup relies on assuming an instantaneous emulsifier exchange, especially so in the studies including coalescence.^{42,43,49,50} However, assuming an instantaneous emulsifier exchange is also problematic when studying pure breakup, since the timescales of turbulent deformation and turbulently driven adsorption are generally considered to be similar.^{7,9} This suggests that adsorption often have difficulties keeping up with deformation, especially when considering larger polymeric emulsifiers [p. 82].⁹ When adsorption is not much faster than deformation, surface load decreases upon deformation, leading to an increase in interfacial tension, thus, increasing the drop's resistance to deformation. Moreover, the magnitude of this additional resistance to deformation differs between emulsifiers as characterized by their Gibbs elasticity,^{51,52}

$$E = \frac{\mathrm{d}\gamma}{\mathrm{d}\ln\Gamma}.$$
 (3)

Furthermore, at least for low-molecular weight surfactants, with high interfacial mobility, localized deformation gives rise to spatial gradients in surface load driving additional restorative Marangoni flow. Protein emulsifiers, on the other hand, often display substantial effects of interfacial rheology such as interfacial shear and dilatational viscosity and interfacial shear and dilatational elasticity.^{53,54}

From previous investigations, it is well known that the adsorption kinetics and interfacial rheology brought about by the properties imposed by the emulsifier is essential to understand coalescence,^{32,51,55} as well as to understand slow laminar deformation.^{56,57} The effect emulsifiers have on turbulent breakup (beyond the effect on the equilibrium value of the interfacial tension) is less well understood. One reason is that it is difficult-theoretically impossible, some argue¹⁹-to build an

experimental system allowing direct visualization with highspeed imaging that correctly scales both adsorption/deformation timescales and hydrodynamics. Thus, simulation-based techniques are required to understand these phenomena.

The aim of the present contribution is to investigate how the limited exchange rate of emulsifier between the bulk and the interface influences the deformation process in turbulent drop breakup. Our focus is on settings corresponding to oil-in-water emulsions with application-relevant properties and concentrations of emulsifiers, in the context of state-of-the-art single drop breakup simulations. Application we aim to understand in the long run include processing of milk, mayonnaise, and dressings (food technology), and processing of intravenous emulsions (pharmaceutical technology).

In this first step it is assumed that the effect of the emulsifier is in lowering the interfacial tension in relation to the increase in surface load brough about by adsorption, as described by the adsorption kinetics and surface equation of state of the system. It is assumed that the emulsifier is surface active as soon as it arrives at the interface (neglecting unfolding/structural change processes sometimes necessary when arriving at the interface), that the effect of interfacial shear and/or dilatational viscosity does not have time to appear, and that the emulsifier distributes itself instantly across the interface (i.e., interfacial tension is assumed to be spatially homogeneous). Such an instant re-distribution across the interface has been argued to be a relevant assumption in (e.g., food) emulsification due to longitudinal waves acting to counter spatial gradients in surfactant concentrations.9 Under these assumptions, the dynamic effect of the emulsifier is described by a limited number of parameters (see Section 2.3). Thus, the general approach of this study is to analyse how drop deformation and breakup is influenced under application-relevant combinations of these parameters, using numerical drop breakup experiments as the methodological framework.

Theory and computational methods

2.1 Single drop breakup simulation methodology

Numerical breakup experiments have been used extensively in fluid mechanics literature to study turbulent breakup.^{10–12,40–48} A more comprehensive description of our methodology can be found elsewhere.⁴⁰ In brief, we inject a single spherical drop into a cuboidal domain with periodic boundaries and containing a developed, forced, isotropic and homogeneous turbulence (Taylor Reynolds number, $\text{Re}_{\lambda} = 32$). By solving the Navier–Stokes equations coupled to a MTHIC volume-of-fluid (VOF) scheme describing the interface^{42,58} the evolution of the flow-field and of the drop is followed with high temporal and spatial resolution (spatial resolution corresponding to 41 cells across the initial drop – see ref. 40 for a mesh-dependence study). The temporal resolution is $0.001\tau_{\eta}$, where τ_{η} denotes the Kolmogorov timescale of the turbulent flow,

$$\tau_{\eta} = \sqrt{\frac{\nu_{\rm C}}{\varepsilon}} \tag{4}$$

(and $\nu_{\rm C}$ denotes kinematic viscosity of the continuous phase). For reference, $\tau_{\eta} \sim 30$ ns for conditions typical of high-pressure homogenization of a low-volume fraction o/w emulsion (*i.e.*, $\varepsilon \sim 10^9$ m² s⁻³, $\nu_{\rm C} \sim 10^{-6}$ m² s⁻¹).

Note that the methodology entails solving for full two-way coupling between the continuous phase turbulent flow and the interface. The volume fraction of disperse phase is set low (1.7%) ensuring that drop filaments does not start interacting with other parts of the drop through the periodic boundaries.

Simulations are based on the CaNS implementation,⁵⁹ which has been extensively tested and validated in previous studies. For emulsification under conditions like those investigated in the present study, good agreement has previously been found between simulations and high-speed breakup visualizations⁶⁰ as well as between simulations and emulsification experiments.⁶¹

Traditionally, when neglecting emulsifier adsorption dynamics and interfacial rheology, three factors are expected to control breakup: the Weber number (We), the disperse to continuous phase viscosity ratio (μ_D/μ_C), and the disperse to continuous phase density ratio (ρ_D/ρ_C). Empirical studies also suggest that the ratio of the initial drop-diameter to the Kolmogorov lengthscale (d_0/η) plays a role, at least in the turbulent inertial regime.^{22,40} In the majority of the simulations in the present study, we set We = 5, $\mu_D/\mu_C = 20$, $\rho_D/\rho_C = 0.9$ and $d_0/\eta = 22$. From an application perspective, this represents the smallest viscous oil drops breaking in an emulsification device.⁴⁰ To compare the behaviour of small *versus* large drops, we also run a sequence of tests with We = 30, corresponding, approximately, to the behaviour of the largest drops entering an emulsification device.^{40,41}

Note, however, that this direct translation of We to drop size is only possible when considering single emulsifier systems. The different emulsifiers studied (see Section 2.3) give rise to different value of the interfacial tension. Thus, We = 5 corresponds to a smaller drop diameter (in physical units) when considering an emulsifier with a higher interfacial pressure.

2.2 Modelling simultaneous adsorption and deformation/ breakup

Under quiescent conditions, such as in a dynamic pendant drop experiment, adsorption is either diffusion controlled as described by a Ward–Tordai model^{62,63} or approximately 'barrier-limited', for example as described by a Langmuir–Hinshelwood model.^{62,64,65}

Turbulent emulsification is far from quiescent, however, and turbulent 'eddies' will lead to a substantial increase in the collision rate between drops, particles and/or molecules,^{66–68} leading to an increase in the effective adsorption rate.⁶⁹ Assuming turbulent breakup to occur in the inertial regime (*i.e.*, the drop diameter is large in comparison to the Kolmogorov length-scale of the turbulence), we model the adsorption rate as the collision rate between the emulsion drop (diameter d_0) and a sphere with a volume-equivalent diameter of the emulsifier (d_E), multiplied with an collision efficiency, taken to vary linearly with

$$(1 - \Gamma/\Gamma^*)$$
;^{64,68-7}

$$\frac{\mathrm{d}\Gamma}{\mathrm{d}t} = 0.272\pi \cdot \varepsilon^{\frac{1}{3}} \cdot (d_0 + d_\mathrm{E})^{\frac{7}{3}} \cdot \frac{c(t)}{\pi \cdot d_0^2} \cdot \left(1 - \frac{\Gamma(t)}{\Gamma^*}\right) \tag{5}$$

when $\Gamma < \Gamma^*$ and 0 elsewise. In eqn (5), the equivalent emulsifier diameter, $d_{\rm E}$, will be either that of the individual molecular aggregate (protein emulsifier) or that of the micelle (low molecular weight surfactant).

From the adsorption rate model in eqn (5) we define the adsorption timescale, τ_{ads} , as the time required to reach a surface load of $\Gamma = 0.99\Gamma^*$, following an instantaneous 5% increase in interfacial area (see Appendix A for derivation):

$$\tau_{\rm ads} = 5.7 \cdot \Gamma^* \cdot c_0^{-1} \cdot \varepsilon^{-1/3} \cdot d_0^{-1/3}, \tag{6}$$

(where c_0 denote the initial bulk concentration of emulsifier). Note that the right-hand-side of eqn (6) scales identical with previous suggestions for turbulent driven adsorption timescales.^{9,67}

To integrate this adsorption model into the single drop deformation simulation framework outlines above (Section 2.1), a multi-scale modelling approach is used. Each drop is assumed to enter the emulsification device after a preemulsification stage; hence, it is in equilibrium with the emulsifier-solution having an initial surface load, Γ_0 , given by the equilibrium adsorption isotherm of the respective emulsifier (see Section 2.3). The DNS-framework is used for advancing the flow-field and the interface in time. Every $\Delta t = 0.01\tau_{\eta}$ (chosen sufficiently small not to influence how deformation evolves over time and representing approximately 10 time-steps in the DNS/VOF), the simulations are stopped, and the surface load in step k, Γ_k , is updated from mass conversion based on the adsorption rate (eqn (5))⁶⁴

$$\Gamma_k = \Gamma_{k-1} \cdot \frac{A_{k-1}}{A_k} + \frac{\mathrm{d}\Gamma}{\mathrm{d}t_k} \Delta t.$$
(7)

The first term in eqn (7), describes the change in surface load due to deformation/relaxation, and the second term in eqn (7) describes the contribution from adsorption. From Γ_k , the interfacial tension in that time-step is calculated from the surface equation of state for the emulsifier system (Section 2.3), and the DNS/VOF is run for another Δt .

Depletion of emulsifier bulk concentration is described using a simple mass balance in the limit of a low volumefraction of disperse phase,⁶⁴

$$c(t) = c_0 - \left(\Gamma(t) \cdot A(t) - \Gamma_0 \cdot A_0\right) \cdot \frac{6 \cdot \phi_{\mathrm{D}}}{\pi D_0^3}.$$
 (8)

Note that this methodology allows for a time-varying interfacial tension but assumes that there are no spatial gradients in the surface load across the interface. Thus, once adsorbed, the emulsifiers are assumed to instantly spread across the interface. According to Walstra and Smulders⁹ this is a reasonable assumption for turbulent emulsification due to the redistributing effect of longitudinal waves.

2.3 Identification of realistic emulsifier parameters

Under the simplifying assumptions outlined above (Section 1), each emulsifier (in combination with a particular choice of disperse and continuous phases liquids) can be described by a limited number of parameters: the initial bulk concentration of the emulsifier (c_0), the hydrodynamic diameter of the emulsifier/micelle (d_e), the equilibrium isotherm (describing the initial surface load corresponding to c_0) and the surface equation of state (relating the interfacial tension to the surface load at each point in time). In particular, the surface equation of state includes information about the interfacial tension for the pure interface (γ_{max}), the interfacial tension when the surface is fully covered (γ_{min}), and the surface load corresponding to this 'fully covered' state (Γ^*).

As seen from the definition of We in eqn (1) (and from viscosity-corrected Kolmogorov–Hinze theory^{7,22–27}), an emulsifier providing a lower interfacial tension will result in smaller drops. In this study, however, we want to compare the effect of different emulsifiers at the same Weber number (since We is the primary quantity controlling the turbulent deformation and breakup). Thus, since We is kept constant when comparing between emulsifiers, it is not the absolute value of the interfacial tension that matters but how much interfacial tension increases following a decrease in surface load from the initial value (which is typically equal to Γ^* since the drop has been subjected to pre-emulsification in the presence of emulsifier at a concentration higher than the critical micelle concentration).

Fig. 1 displays normalized surface equations of state for three different emulsifier systems: (i) the non-ionic triblock copolymer Pluronic F-68 (F68) at the vegetable oil/water interface, (ii) the anionic low-molecular weight surfactant sodium dodecyl sulfate (SDS) at the dodecane/water interface; and (iii) the milk protein β -lactoglobulin (BLG) at the tetradecane/ water interface. All three emulsifiers are soluble in the continuous aqueous phase, and commonly used as model emulsifiers

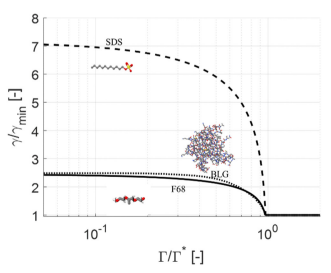


Fig. 1 Interfacial tension (normalized to its minimum value, γ_{min} , reached at full surface coverage) as a function of surface load (Γ/Γ^*), based on literature data for SDS^{72,74} BLG⁷³ and F68.⁶⁴

for studying emulsification. (Physical properties, equilibrium adsorption isotherms, and surface equations of state are taken from literature and are summarized in Table 1.) The horizontal axis in Fig. 1 displays the normalized surface load, Γ/Γ^* , indicating how close the surface load is to its maximum value. The vertical axis shows the interfacial tension scaled with its value when the surface load approaches Γ^* (*i.e.*, interfacial tension at 'full coverage').

In Fig. 1, first, note that BLG and F68 result in similar normalized interfacial tension curves, despite that they correspond to two molecularly dissimilar emulsifiers (protein vs. non-ionic block copolymer) operating on two molecularly dissimilar disperse phase liquids (hydrocarbon vs. vegetable oil). More specifically, the rate at which the interfacial tension increases with a small decrease in surface load is similar near $\Gamma \sim \Gamma^*$ for these two systems. The third system (SDS) gives rise to a substantially faster increase in interfacial tension when decreasing the surface load. This follows from that the Gibb's elasticity of the BLG and F68 are similar (90 and 88 mN m^{-1} , as calculated from their surface equations of state and eqn (3)) and lower than for SDS (125 mN m^{-1}). Also note that SDS can, if the surface load is substantially reduced, lead to a substantially larger relative increase in interfacial tension compared to its initial (nominal) value (factor 7 for SDS as compared to a factor 2.5 for BLG and F68).

In this study, we use the three emulsifier systems from Fig. 1 and Table 1 (F68, SDS and BLG) as illustrative cases to study how adsorption kinetics influences turbulent deformation and breakup. The choice of these three is based on (a) availability of high-quality experimental data and surface equations of states, (b) that these three represent the three main classes of emulsifiers found in food applications, and (c) that they allow us to investigate the effect of Gibbs elasticity.

Each emulsifier is investigated at two concentrations ('high' and 'low'), chosen to be in the range relevant for emulsification applications. For BLG, the 'high' concentration is that achieved when preparing an emulsion with same BLG-concentration as in raw milk and 'low' is one tenth of this – which is still twice the concentration above which surface load is approximately independent of bulk concentration under equilibrium conditions.⁷³ For F68 and SDS, 'high' concentration is 20 times the critical micelle concentration (CMC) and 'low' is a concentration twice the CMC (the 'high' concentration is still well below the aqueous solubility limit for the respective emulsifier).

Deformation gives rise to an expansion of the drop interface, which lowers the surface load and drives adsorption (through eqn (5)). Note that the adsorption rate is zero once the drop is fully covered by emulsifier ($\Gamma = \Gamma^*$). However, deformation is typically not monotonic.^{10,40,45,75} For a relaxing drop, to which adsorption has taken place during deformation, this might lead to surface load increasing beyond the threshold/maximum surface load Γ^* (typically the magnitude of the effect is small, *i.e.*, $\Gamma < 1.05\Gamma^*$). Since low-molecular weight surfactants typically adsorb reversibly, SDS was assumed to instantly desorb back to $\Gamma = \Gamma^*$ if this occurred. For the larger emulsifiers

Table 1 Interfacial properties of the three studies emulsifier systems

	Triblock copolymer (F68)	Sodium dodecyl sulfate (SDS)	β -Lactoglobulin (BLG)
Emulsion	Vegetable oil in water	Dodecane in water	Tetradecane in water
Temperature	25 °C	20 °C	22 °C
Molecular weight of emulsifier (<i>M</i>) [kg mol ^{-1}]	8.4^{64}	0.288	$18.3^{81,82}$
Hydrodynamic diameter $(d_{\rm E})$ [nm]	12 ⁷⁹ (micelle)	4 ⁸⁰ (micelle)	7 ⁸³ (dimer)
Isotherm parameter (Γ^*) mol m ⁻²⁻	$1.4 imes 10^{-6.64}$	5.2×10^{-672}	0.11×10^{-673}
Equilibrium adsorption isotherm	Langmuir isotherm ⁶⁴	Two states model ⁷²	Multiple-state model ^{73,82,8}
Interfacial tension at $\Gamma = 0(\gamma_{max}) [mN m^{-1}]$	19.6 ⁶⁴	51.3^{72}	51.3 ^{73,84}
Interfacial tension at $\Gamma = \Gamma^{*}(\gamma_{\min}) [\text{mN m}^{-1}]$	8.0^{64}	7.18^{72}	$20.6^{73,82}$
Surface equation of state	Szsyzkowski/Frumkin equation ⁶⁴	Two states model ⁷²	Multiple-state model ^{73,82,84}
Critical micelle concentration (c_{CMC}) [mol m ⁻³]	0.48^{64}	8.3 ⁷²	_ 1
Technically relevant low conc. (c_{low}) [mol m ⁻³]	$2 \cdot c_{\rm CMC}$	$2 \cdot c_{\rm CMC}$	1.7
	20·c _{CMC}	$20 \cdot c_{\rm CMC}$	0.17

(F68 and BLG) adsorption is irreversible^{76,77} and $\Gamma > \Gamma^*$ is allowed in the simulations.

Using the numerical data on our three emulsifier systems, we can now see more quantitatively that the turbulently driven adsorption rate is substantially faster than both the Brownian/ diffusion driven adsorption given by,^{71,78}

$$\frac{\mathrm{d}\Gamma}{\mathrm{d}t}\Big|_{\mathrm{diffusion}} = \frac{2 \cdot k_{\mathrm{B}} \cdot T}{3\mu_{\mathrm{C}}} (d_0 + d_{\mathrm{e}}) \cdot \left(\frac{1}{d_0} + \frac{1}{d_{\mathrm{e}}}\right) \cdot \frac{c(t)}{\pi \cdot d_0^2} \cdot \left(1 - \frac{\Gamma(t)}{\Gamma^*}\right),\tag{9}$$

(where $k_{\rm B}$ is Boltzmann's constant and *T* is absolute temperature), and the barrier-limited adsorption^{64,67} at these conditions. This is illustrated in Fig. 2 displaying the adsorption rate in a non-dimensionalized form, $d\Gamma/dt/(c_0 \cdot u_\eta)$, where u_η is the Kolmogorov micro-scale velocity. For F68, turbulent adsorption is a factor 3.7×10^4 faster than diffusion-controlled adsorption and a factor 2.4×10^5 faster than barrier-limited Langmuir–Hinshelwood adsorption (using adsorption parameters for F68 as suggested by Maindarkar *et al.*⁶⁴) at $\Gamma = 0.95\Gamma^*$. This can be compared to applying

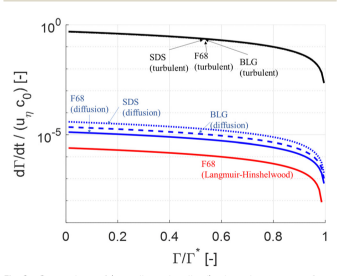


Fig. 2 Comparison of (non-dimensionalized) adsorption rate as a function of surface load. Comparing turbulently driven adsorption (black line, eqn (5)) to Brownian diffusion driven transport (blue line⁷⁸) and Langmuir–Hinshelwood limited adsorption (red line^{62,64}).

Levich $[p. 218]^{67}$ rough scaling which suggests that turbulence results in roughly a 10^6 times faster collision rate than Brownian diffusion under these conditions.

Note that diffusion is insignificant in comparison to turbulent driven adsorption despite that F68 is a relatively small polymer. Moreover, as seen in Fig. 2, the turbulently driven adsorption rate is almost independent of which emulsifier we consider since the velocity of the emulsion drop is the dominant factor driving drop-emulsifier collisions (*i.e.*, since $d_0 + d_E \sim d_0$ in eqn (5)). For SDS turbulent adsorption is a factor 1.3×10^4 faster than diffusion and for BLG, the turbulent adsorption is a factor 2.0×10^4 faster than diffusion (at $\Gamma = 0.95\Gamma^*$).

3. Results and discussion

3.1 Deformation and breakup neglecting emulsifier adsorption dynamics

Fig. 3 displays how a We = 5 drop-based on the standard assumption of instant exchange of emulsifier between bulk and interface deforms in one example flow realization (data displayed as an iso-surface contour of VOF = 0.5, flow realization L, see drop library in ref. 41). The drop starts to deform directly upon injection and is rapidly flattened. Starting at $t/\tau_{\eta} = 5$, a segment of the drop is further pulled out (downwards). As time progresses, this segment is deformed, forming a bulb at the end of the segment. Over time, the neck connecting the two is thinned out and the neck is critically deformed (*i.e.*, deformed to the extent that breakup is now deterministic even if it was removed from the turbulence⁴¹) at $\tau_{def}/\tau_{\eta} = 9.9$. Initial breakup (first detachment of a fragment⁸⁵) occurs at $\tau_{break}/\tau_{\eta} = 15.3$.

Fig. 4(A) displays the total interfacial area of the drop as a function of time, as a measure of the global extent of deformation, up to the point of initial breakup (marker). The drop deforms monotonously and has reached a value of 1.8 times the initial area when breaking, see black solid line ('Instant') in Fig. 4(A). Fig. 4(B) and (C) display the evolution of the normalized surface load and the normalized interfacial tension, respectively. Note how these are constant under the assumption of instant exchange of emulsifier between bulk and interface.

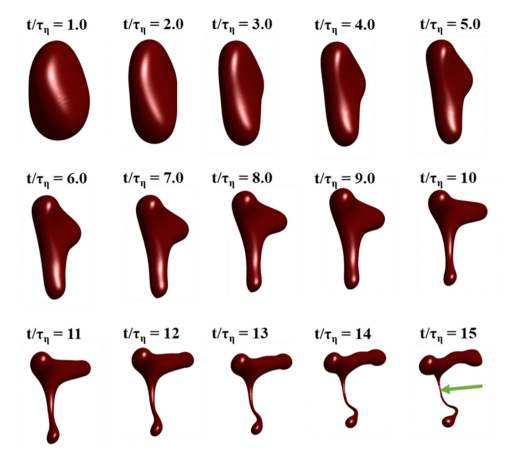


Fig. 3 Deformation and breakup sequence for the drop when assuming instant exchange of emulsifier between bulk and interface. Arrow displays point of first detachment. (Isosurfaces of VOF = 0.5. We = 5. Flow realization L.)

3.2 Effect of emulsifier dynamics on deformation

Fig. 5 displays drops injected in the same flow realization as in Fig. 3 (i.e., at the same Weber number and viscosity ratio) but when using dynamically adsorbing/desorbing emulsifiers at different concentrations to achieve these conditions. First note, that if the emulsifier concentration is sufficiently high (e.g., BLG/high in Fig. 5), the drop deformation is almost indistinguishable from that seen under instant exchange between bulk and interface ('Instant' in Fig. 5). Also note that BLG/high corresponds to using β -lactoglobulin as the emulsifier, under concentrations comparable to that in milk, *i.e.*, we can achieve an adsorption rate independent deformation under conditions relevant in technical applications. The surface load does decrease initially due to the fast initial deformation as the drop encounters the turbulence (see drop colour-scale in Fig. 5 and plot in Fig. 4(B)). This also leads to an initial increase in interfacial tension of the drop (see Fig. 4(C)). However, with a high emulsifier concentration (BLG/high in Fig. 5), the adsorption rate is sufficiently fast to restore the surface load and, thus, lower the interfacial tension back to its initial value before deformation has ceased. Consequently, deformation and breakup are not limited by the adsorption under these conditions.

If the emulsifier is used at a lower concentration, however, this significantly alters the deformation process, including morphology at breakup, deformation time and breakup time. This is exemplified by using the same emulsifier at the lower (but still technically relevant concentration), see 'BLG/low' in Fig. 5. With the lower concentration, the emulsifier adsorption is slower, and, thus, the surface load decreases more (reaching as low as $\Gamma/\Gamma^* = 0.88$ at $t/\tau_{\eta} = 2.2$), leading to a higher interfacial tension ($\gamma/\gamma_{min} = 1.4$), decreasing the effective Weber number by 40%. This increase in interfacial tension brings an increased stabilization. Thus, the segment which is pulled out from the drop under conditions of instant adsorption ($t/\tau_{\eta} > 5$, Fig. 3), is halted and relaxes back when the emulsifier concentration is too low ('BLG/low' in Fig. 5). Breakup is observed even at the lower concentration, but it occurs later ($\tau_{\text{break}}/\tau_n = 31.2$).

3.3 Emulsifier type and Gibbs elasticity

How susceptible the deformation process is to the emulsifier depend on which emulsifier is used. BLG and F68 behaves similarly (Fig. 4); when used at technically relevant 'high' concentrations ('BLG/high' and 'F68/high' in Fig. 4), the adsorption is fast enough to restore surface load and, consequently, breakup occurs at approximately the same time as when assuming instant exchange of emulsifier between bulk and interface. At the 'low' (but still technically relevant) concentrations, both BLG and F68 give rise to a delayed breakup,

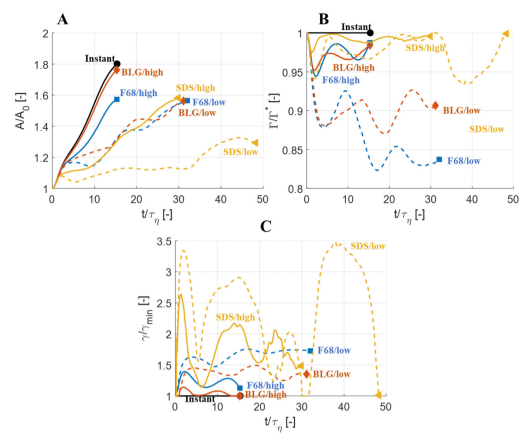


Fig. 4 Evolution of (A) interfacial area, A/A_0 ; (B) surface load, Γ/Γ^* ; and (C) interfacial tension, γ/γ_{min} , comparing the case of instant exchange of emulsifier between bulk and interface (black, solid) to the three emulsifiers at low and high concentration. (We = 5. Flow realization L.) Marker display the state of initial breakup.

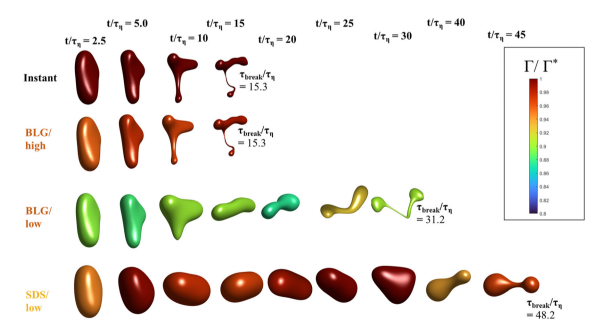


Fig. 5 Comparing how drop deformation and surface load evolves over time for different emulsifiers and concentrations. (Iso-surfaces of VOF = 0.5, coloured by surface load.) Time of initial breakup, τ_{break} , supplied for each case as reference (We = 5. Flow realization L).

occurring at $\sim 31\tau_{\eta}$. To understand why these two emulsifiers behave similarly, it is also interesting to note that they give rise to

similar Gibbs elasticities, *i.e.*, the interfacial tension responds similarly to a small change in surface load (see eqn (3)).

The low weight molecular emulsifier (SDS), however, is different. The adsorption rate is relatively high (as indicated by the relatively short adsorption time, even at the 'low' concentration for SDS), resulting in a high surface load ('SDS/ low' in Fig. 5 and 4(B)). The higher Gibbs elasticity imposed by SDS (see Fig. 1), however, gives even a slight reduction in surface load a large impact on the interfacial tension ('SDS/ low' in Fig. 4(C)). For the low SDS concentration case, the interfacial tension rapidly increases to 3.3 times its initial value, thus decreasing the effective Weber number encountered by the drop from 5 to 1.5. The high Laplace pressure causes the drop to quickly relax from the initial deformation (Fig. 5). Also note that deformation is delayed, even if using this high Gibbs elasticity emulsifier at a high concentration (SDS/high); despite that adsorption rapidly loads the expanding interface with new emulsifier, the slight and short-lived depletion of the adsorbed layer stabilizes the drop (Fig. 4).

The high Gibbs elasticity drop ('SDS/low') does eventually break, but it does so in a slow process. After the initial deformation has been halted by the high interfacial tension, it proceeds by deforming slowly (dA/dt is low in Fig. 4(A)). During this slow deformation, the drop continues to accumulate emulsifier through adsorption to the deformed interfaceas the interface deforms, Γ/Γ^* remains below one, resulting in a continuous driving force for adsorption, which increases the total adsorbed amount, $\Gamma(t) A(t)$, over time. Shortly after $t/\tau_n \sim$ 30, some turbulent structures appear to interact more forcefully with the drop, increasing the deformation rate (see Fig. 4(A)). The drop is unable to break directly upon this extra stress (since interfacial tension increases rapidly in response, as seen in Fig. 4(C)). However, the turbulent structures allow the drop to relax in such a way so that it forms the necessary thinning neck, while simultaneously reducing its total interfacial area (cf. Fig. 4(A) and 'SDS/low' in Fig. 5). Due to the slow accumulation of emulsifier during the first $30\tau_{\eta}$, the drop can now achieve a relatively high surface load even in the critically deformed state (see colour-scale of 'SDS/low' in Fig. 5), allowing the drop to break.

3.4 Adsorption time

The observations on how emulsifier concentration influences the deformation process can be quantified in terms of adsorption time (cf. eqn (6)). Using the same emulsifier at a higher concentration reduces the adsorption time (*i.e.*, by increasing c_0 in eqn (6)). Fig. 6(A) displays the maximum reduction in surface load, min(Γ/Γ^*), during the initial deformation ($t < 10\tau_n$), comparing different emulsifiers and concentrations thereof, plotted versus the adsorption time imposed by using different emulsifier concentration. Starting with a single emulsifier (F68, blue squares), the shorter is the adsorption time, the smaller is the maximum reduction in the surface load. The curve shows a sigmoidal shape, approaching no depression in surface load if the adsorption time is short in comparison to the Kolmogorov timescale (τ_n) , and levelling off at min $(\Gamma/\Gamma^*) \approx 0.7$ if adsorption time is long in comparison to τ_{η} . Note that if $\tau_{ads}/\tau_{\eta} > 5$, the adsorption is sufficiently slow for adsorption to be negligible,

and the surface load evolution is almost entirely determined by how fast the interface is deformed by the external turbulent stress. As before, BLG and F68 behaves similarly, with $\min(\Gamma/\Gamma^*)$ falling approximately on the same master curve.

The corresponding maximum increase in interfacial tension during the initial deformation, $\max(\gamma/\gamma_{\min})$, can be seen in Fig. 6(B). Note that the two emulsifiers with approximately the same Gibbs elasticity (F68 and BLG) (disks and squares) fall approximately on the same curves in Fig. 6. The high interfacial elasticity emulsifier (SDS), however, gives a somewhat lower relative reduction in surface load at the same adsorption time (*i.e.*, $\min(\Gamma/\Gamma^*)$), see Fig. 6(A), and a higher relative increase in interfacial tension (*i.e.*, $\max(\gamma/\gamma_{\min})$), see Fig. 6(B).

The effect of emulsifier dynamics on breakup time can also be better understood in terms of adsorption time, see Fig. 6(C). Starting with BLG and F68 (low interfacial elasticity), the breakup time is approximately equal to that obtained by assuming instant exchange of emulsifier between bulk and interface if (and only if) $\tau_{ads} < 1.4\tau_{\eta}$. This corresponds to an adsorption time that is less than 0.1 of the breakup time. Thus, if the emulsifier is used at a high-enough concentration so to achieve an adsorption time that is one order magnitude lower than the breakup time, the turbulent deformation process is independent of emulsifier dynamics. If not, there is a delay in breakup.

From previous investigations, it is well-known that using an emulsifier concentration less than the level indicated by eqn (2) makes turbulent coalescence dominate turbulent breakup.^{6,7,28–31} However, the critical concentration required to prevent coalescence is low unless using exceedingly high volume-fractions of disperse phase, *i.e.*, ~1/100 $c_{\rm cmc}$ for F68 according to eqn (2). This is substantially lower than the concentration required to ensure that the deformation process is uninfluenced by the rate at which emulsifier is exchanged between bulk and interface.

3.5 Generalization across flow initializations

Turbulent flows are highly stochastic, and each drop subjected to it will take a different trajectory and interact with different turbulent structures.^{10,36,40,46,60} Looking at a larger collection of viscous drops ($\mu_D/\mu_C = 20$) subjected to conditions of We = 5, we see mainly two types of deformation processes:⁴¹ some drops-upon being injected in the turbulence-encounter sufficiently forceful turbulent stress to immediately go into a monotonous deformation that eventually leads to breakup (*cf.* Fig. 3). Other drops go through a stochastic number of oscillations before eventually reaching a deformation sequence that eventually break them (see Fig. 7 for an example). This behaviour has also been seen in similar investigations across a larger number of flow realizations in independent studies.^{11,45}

As a test of how the results from the investigations above generalize between flow realizations, Fig. 8 (open markers) displays the results of how maximum surface load suppression (Fig. 8(A)), minimum interfacial tension (Fig. 8(B)) and breakup time (Fig. 8(C)), depends on adsorption timescale, comparing the previously investigated flow realization leading to

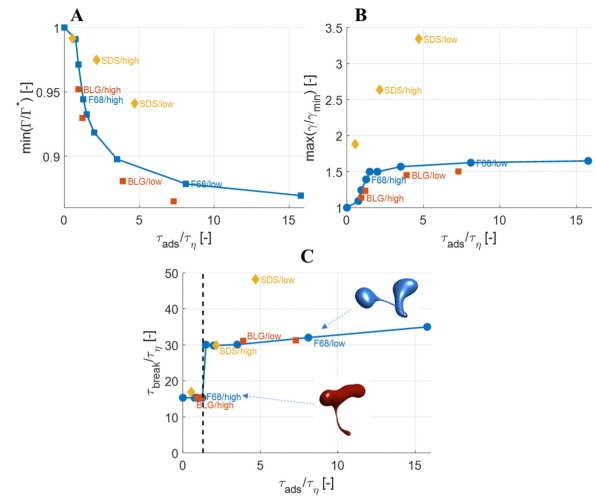


Fig. 6 (A) Surface load suppression, min(Γ/Γ^*); (B) interfacial tension increase, max(γ/γ_{min}); and (C) breakup time, τ_{break} , as functions of the adsorption time, τ_{ads} , imposed by the different emulsifiers at varying concentrations. Insets in C displays drop morphology at breakup for one case (F68/high) where the emulsifier imposes a sufficiently fast adsorption to make emulsifier exchange fast (independent of emulsifier dynamics) and one case (F68/slow) where adsorption is slow (dependent on emulsifier dynamics) (We = 5. Flow realization L).

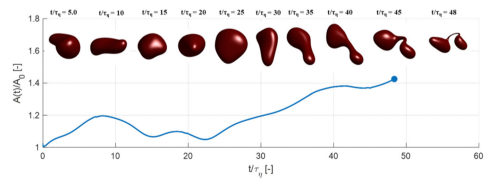


Fig. 7 Deformation and breakup sequence for a drop displaying oscillations before reaching breakup, assuming instant exchange of emulsifier between bulk and interface (Isosurfaces of VOF = 0.5. We = 5. Flow realization D).

monotonous deformation (labelled 'L'), to the flow realization in Fig. 7, leading to oscillatory behaviour (labelled 'D'). Solid markers show results from the flow realization used in previous sections ('L'). As seen in Fig. 8(A) and (B), the maximum initial reduction in surface load and the resulting maximum increase in interfacial tension is similar between the two flow realizations.

More interestingly, Fig. 8(C) displays how breakup time depends on the emulsifier concentration (*via* its effect on the

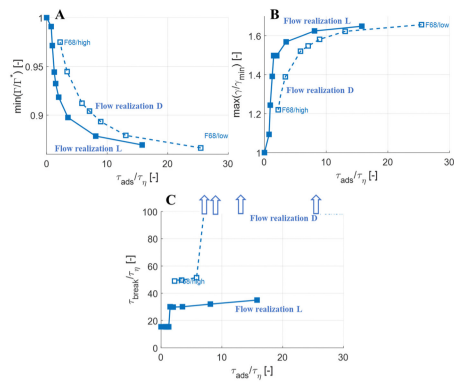


Fig. 8 Surface load suppression (A), interfacial tension increase (B) and breakup time as a function of the adsorption timescale, τ_{ads} , comparing F68 between flow realization D and L. Arrows in (C) illustrate that the breakup time is larger than $100\tau_{\eta}$ at these combinations of τ_{ads} and flow realization (We = 5).

imposed adsorption timescale), comparing the two flow realizations. As discussed above, for the previously discussed flow realization ('L'), breakup is unaffected if emulsifier is supplied at a concentration to ensure that $\tau_{ads} < 0.1 \tau_{break}$ (solid markers), and breakup is delayed if the emulsifier concentration is below this threshold. For the flow realization where breakup is preceded by oscillations (flow realization 'D', open markers in Fig. 8(A)), a similar effect is seen, but the delay is long enough to move it beyond the end-time in the simulations (corresponding, approximately, to the time spent in the emulsification device, $t/\tau_n \approx 100^{40}$). In Fig. 8(C), these conditions where the imposed adsorption time is sufficiently long for drops not to break before $t/\tau_{\eta} \approx 100$ are illustrated using arrows. In terms of breakup time, this critical adsorption time required for fast exchange is longer for the second flow realization ($\tau_{ads} < 0.14 \tau_{break}$, as compared to τ_{ads} < 0.11 τ_{break} for the first flow realization), but of the same order of magnitude, $\tau_{ads} < 0.1 \tau_{break}$.

The results above suggest that for drops with We = 5, the assumption of an instant exchange of emulsifier between bulk and interface, which is imposed in the majority of previous numerical drop breakup studies, is only realistic if using an emulsifier concentration that is high enough to ensure that the adsorption time is a decade lower than the breakup time.

3.6 Effect at higher Weber number

The focus thus far has been on low Weber numbers (*e.g.*, characteristic of the smallest drops breaking in an emulsification device). For a large drop (illustrated here by We = 30),

the adsorption dynamics has less of an effect on the deformation and breakup behaviour (Fig. 9). With a slowly adsorbing emulsifier, the drop deforms less than if adsorption was instant (Fig. 9(A)) but break at almost the same time (see markers in Fig. 9(A)). At low emulsifier concentrations (SDS/low and F68/low), the adsorption is slow, and the surface load decreases initially—for F68/low Γ/Γ^* reaches as low as 0.6 (Fig. 9(B))—which leads to a large increase in interfacial tension (Fig. 9(C)). However, even with interfacial tension reaching 4.5 times higher than the equilibrium value at its peak (SDS/low, see Fig. 9(C)), the effective Weber number obtained by replacing the equilibrium value of interfacial tension with the timedependent one from Fig. 9(C), is still large (We \geq 11), indicating that the disruptive turbulent stresses still substantially outweigh the stabilizing Laplace pressure. The results even suggest that the high-We drop breaks marginally earlier if subjected to a slowly adsorbing emulsifier giving a high Gibbs elasticity (SDS/low breaks $0.8\tau_{\eta}$ earlier than when assuming instant emulsifier exchange, Fig. 9(A)-(C)), most likely due to that the transient increase in interfacial tension hinders the excessive deformation in regions further from the neck which acts as a drain of turbulent kinetic energy (cf. ref. 44).

The observation that large drops (*i.e.*, drops with large We) are less sensitive to emulsifier dynamics, also explains why emulsification experiments see relatively little effect of emulsifier concentration on the drop size distribution beyond the concentration required to prevent coalescence during emulsification.^{7,28-30} Whereas a considerable amount of turbulent coalescence will

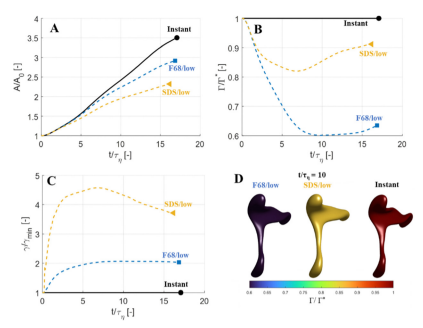


Fig. 9 Evolution of drop deformation (A), surface load (B) and interfacial tension (C), for a drop with a high Weber number (We = 30), comparing instant emulsifier exchange (black solid) to SDS and F68 at low concentrations. (D) Drop morphology (isosurface of VOF = 0.5) and surface load (colour-scale) of drops at t/τ_{η} (We = 30, flow realization L).

drastically increase the number of large drops which shows up as a marked translation of the drop-size distribution and volume-based average diameters, a delay in the deformation process which only acts on small drops have less of an effect on the drop size distribution, thus making it more difficult to identify using emulsification experiments.

3.7 Non-uniform surface load, Marangoni stresses and interfacial rheology

In the present study, following Walstra and Smulders,9 it is assumed that once adsorbed to the interface, the emulsifiers distribute themselves evenly across the interface. However, the rate at which emulsifiers redistributes on the interface is not understood in detail. If redistribution over the interface is slower than the deformation/relaxation, this would lead to gradients in surface load (and, consequently, in interfacial tension) across the interface, which drives tangential Marangoni stresses that act by further pulling the interface together.^{51,52,55,56} Thus, surface load gradients will also act to delay the deformation process. Large emulsifiers such as proteins will generally require additional time, once arrived at the interface, to undergo the conformational changes ('unfolding') required to reach more energetically favourable conformations. This process can take several orders of magnitude longer than the time a drop spends in the turbulent zone of an emulsification device.^{86,87} Moreover, these emulsifiers typically give rise to interfacial rheology expected to further affect the deformation process.^{53,54} Since surface active material is typically present, both at drop and bubble interfaces during turbulent deformation and breakup in technically relevant and naturally occurring systems, further investigations continuing to include more of these phenomena in single drop breakups simulations is of great importance to advance the field as well as

for advancing our understanding of turbulent drop and bubble breakup.

4. Conclusions

If the Weber number is sufficiently high or if the emulsifier concentration is high enough to give an adsorption timescale which is a decade lower than the breakup time, then the breakup process is identical to that obtained when assuming instant exchange of emulsifier between bulk and interface (as in current state-of-the-art drop simulation literature). For a small drop (We = 5) with relatively high viscosity ($\mu_D/\mu_C = 20$), using a lower emulsifier concentration than this delays breakup. If the time spent in the turbulence is short enough, this delay can result in that the drop does not have time to break before exiting (as seen for the flow realization where the drop oscillates before entering the stage of critical deformation).

Note that both these conditions (deformation not influenced by emulsifier dynamics and deformation delayed by emulsifier dynamics) are found within the span of technically relevant emulsifier concentrations.

The Gibbs elasticity imposed by the emulsifier plays an important role. For the emulsifier imposing a higher interfacial elasticity, the breakup is delayed longer when using a low emulsifier concentration. Moreover, for emulsifiers imparting higher Gibbs elasticity, breakup of limiting drops (*i.e.*, drops with low Weber number) is only possible through a slow deformation process where emulsifier is accumulated allowing critical deformation to be reached without a rapid global increase in interfacial area. Thus, the kinetics imposed by the emulsifier does not only delay breakup but might also influence the mechanism of breakup.

Author contributions

AH: conceptualization, writing – original draft; methodology; analysis; visualization. LN: conceptualization, writing – review & editing.

Conflicts of interest

There are no conflicts to declare.

Appendix A: adsorption time

Consider an emulsion drop in equilibrium with a continuous bulk concentration of emulsifier, c_0 , which is above the critical micelle concentration, CMC, so that the drop has a surface load Γ^* . Assume that at t = 0 the drop is suddenly deformed so that its interfacial area increases by 5%. Since the deformation is too fast for adsorption to compensate, the surface load decreases to an amount given by mass conservation across the interface, *i.e.*, $\Gamma(t = 0) = \Gamma^*/1.05$.

The time necessary for the surface load to increase to 99% of the critical surface load, denoted τ_{ads} , can be obtained from integration of the adsorption rate (eqn (5)):

$$\int_{\Gamma=\frac{1}{1.05}\Gamma^*}^{0.99\Gamma^*} \frac{1}{1-\Gamma(t)/\Gamma^*} d\Gamma = \int_{t=0}^{\tau_{ads}} 0.272\pi \cdot \varepsilon^{\frac{1}{3}} \cdot (d_0 + d_E)^{\frac{7}{3}} \cdot \frac{c(t)}{\pi \cdot d_0^2} dt.$$
(A.1)

Since the volume fraction of disperse phase is low, the bulk concentration will remain approximately constant, *i.e.*, $c(t) \equiv c_0$. Moreover, $d_0 + d_E \approx d_0$, since the emulsifier (or micelle thereof) is small in comparison to emulsion drops in these applications. Thus, the equation can be simplified to,

$$\Gamma^* \cdot \ln\left(\frac{1-1/1.05}{1-0.99}\right) = 0.272 \cdot c_0 \cdot \varepsilon^{1/3} \cdot d_0^{1/3} \int_{t=0}^{\tau_{\text{ads}}} dt, \qquad (A.2)$$

which is equivalent to,

$$\tau_{\rm ads} \approx 5.7 \cdot c_0^{-1} \cdot \Gamma^* \cdot \varepsilon^{-1/3} \cdot d_0^{-1/3}.$$
 (A.3)

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