Level-set simulations of a 2D topological rearrangement in a bubble assembly: effects of surfactant properties

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A liquid foam is a dispersion of gas bubbles in a liquid matrix containing surface active agents. Their flow involves the relative motion of bubbles, which switches neighbours during a so-called topological rearrangement of type 1 (T1). The dynamics of T1 events, as well as foam rheology, have been extensively studied, and experimental results point to the key role played by surfactants in these processes. However, the complex and multiscale nature of the system has so far impeded a complete understanding of the mechanisms at stake. In this work, we investigate numerically the effect of surfactants on the rheological response of a 2D sheared bubble cluster. To do so, a level-set method previously employed for simulating two-phase flow has been extended to include the effects of the surfactants. The dynamical processes of the surfactants —diffusion in the liquid and along the interface, adsorption/desorption at the interface— and their coupling with the flow —surfactant advection and Laplace and Marangoni stresses at the interface— are all taken into account explicitly. Through a systematic study in Biot, capillary and Péclet numbers which characterise the surfactant properties in the simulation, we find that the presence of surfactants can affect the liquid/gas hydrodynamic boundary condition (from a rigid-like situation to a mobile one), which modifies the nature of the flow in the volume from a purely extensional situation to a shear. Furthermore, the work done by surface tension (the 2D analogue of the work by pressure forces), resulting from surfactant and interface dynamics, can be interpreted as an effective dissipation, which reaches a maximum for Péclet number of order unity. Our results, obtained at high liquid fraction, should provide a reference point, to which experiments and models of T1 dynamics and foam rheology can be compared.

Key words: Bubble dynamics - Foams - Interfacial Flows (free surface)

1. Introduction

Liquid foams are concentrated dispersions of gas in a liquid matrix. They belong to the material class of complex fluids, characterised by their multiscale structure, and their rheological properties have been widely investigated, for example in the two pioneering...
paper series of Princen and Kraynik (Princen 1983, 1985; Princen & Kiss 1986, 1989; Kraynik et al. 1991). A commonly used empirical description for foam rheology is the Herschel-Bulkley relationship (Cantat et al. 2010; Cohen-Addad et al. 2013). Foam behaviour is complicated further by these being out of equilibrium systems that evolve with time, due to different mechanisms such as gravity liquid drainage, bubble coarsening and coalescence (Cantat et al. 2010).

Several numerical and analytical methods have been utilized to attempt to link their properties at the local scale to the macroscopic rheological behavior (Buzza et al. 1995; Cantat 2011; Denkov et al. 2008; Tcholakova et al. 2008; Besson et al. 2008; Cohen-Addad et al. 2013). Among these, large or multi-scale simulations may describe the foam either as bubble or soft sphere assemblies (Durian 1995, 1997; Rognon et al. 2010; Seth et al. 2011; Sexton et al. 2011), or as networks of films and Plateau borders (Kern et al. 2004; Cantat 2011; Saye & Sethian 2013). However, to be fully accurate, such simulations require a full description of local responses and timescales, which is still missing.

One essential feature of a liquid foam is that the liquid matrix is filled with surface-active molecules (i.e. surfactants), which adsorb at interfaces, and whose primary role is to stabilise the liquid films separating the bubbles by inducing nanometric range repulsion between the interfaces (Israelachvili 2010). Macroscopic behaviour of foams is strongly affected by the surfactant nature. For instance, liquid transport through foams is limited either by dissipation in the Plateau borders or in the nodes (Durand et al. 1999) and depends on interfacial boundary conditions and hence on surfactant dynamics (Lorenceau et al. 2009; Cohen-Addad et al. 2013). Regarding foam rheology, shear stress in flow (Tcholakova et al. 2008) can be modified by changing the nature of surfactants; viscoelastic measurements have shown that relaxation timescales (Krishan et al. 2010; Costa et al. 2013) are also affected. Finally, the foam stability, characterised either by a critical volume fraction or by a critical capillary pressure, can be modified (Biance et al. 2011; Rio & Biance 2014).

Foam flow occurs through relative motion of deformable bubbles. As depicted in figure 1, this process involves switching of neighbours, which is referred to as a topological rearrangement of type 1 (T1) (Höhler & Cohen-Addad 2005). T1 dynamics has been largely studied experimentally in model systems such as bubble clusters (Biance et al. 2009), 2D foams (Durand & Stone 2006), soap film architecture (Hutzler et al. 2008; Petit et al. 2015) and 3D foams (Le Merrer et al. 2012, 2013). It is affected mainly by the amount of liquid in the foam, the viscosity of the liquid and the nature of the surfactant. T1 involves the flow and the stretching of a thin liquid film, a process which has been extensively characterised experimentally (Seiwert et al. 2013; Petit et al. 2015; Champougny et al. 2015) and was found to depend on the nature of the surfactants. Elongation of the film is observed in some cases whereas shear in the contacting meniscus appears in some others, as already predicted for small deformations in Buzza et al. (1995). On the theoretical side, analytical prediction of T1 dynamics when neglecting bulk flow has also been performed, taking into account surfactant transport along the interface (Durand & Stone 2006; Satomi et al. 2013).

Surfactants not only affect the static properties of interfaces, but also their dynamical ones, described through the interfacial stress. They generate intrinsic surface dissipation but also alter the liquid flow by changing the hydrodynamic boundary condition at the interface. Surfactants can diffuse along the interface and in the bulk, and can partially adsorb and desorb (Langevin 2014), with many consequences, such as the generation of an elastic solutal Marangoni stress at the interface and a viscous one (Lucassen & van den Tempel 1972). Such interfacial Marangoni stresses influence the liquid flows, which
in turn modify the surfactant distribution, making this coupled non-linear problem a complex one.

The effects of surfactants on flows involving bubbles have already been established in a variety of situations. Perhaps the oldest one is the sedimenting drop or rising bubble, a problem dating back to the early days of surface rheology (Edwards et al. 1991), and which is still being explored (Bel Fdhila & Duineveld 1996; Cuenot et al. 1997). By immobilising the interface and decreasing the velocity, the presence of surfactants not only affects the individual behavior of a bubble but has also macroscopic consequences on the turbulence structure (Takagi & Matsumoto 2011). Deformation and break-up of droplets or bubbles in elongational or shearing flows is also sensitive to the presence of surfactants (Stone 1994), as exemplified by the "tip streaming" phenomenon, where a thin liquid thread is drawn from the drop tips (Eggleton et al. 2001). Interfacial boundary conditions influenced by surfactants are also known to affect film coating (Park 1991; Ou Ramdane & Quéré 1997; Scheid et al. 2010; Champougny et al. 2015) or the similar problems of bubbles sliding along a rigid wall (Ratulowski & Chang 1990; Cantat 2013) or foam wall slip (Denkov et al. 2005, 2006). Finally, surfactants may affect the draining process of films, revealing the importance of surface elasticity (Sonin et al. 1993), or resulting in dimpled profiles (Breward & Howell 2002).

Theoretical investigations of surfactant effects relying on analytical or semi-analytical methods generally assume a fixed geometry (e.g. Schwalbe et al. 2011) or lubrication approximation (e.g. Scheid et al. 2010; Cantat 2013). Only numerical approaches can handle the large deformations and topological changes that often occur in bubble or drop dynamics. A number of methods have been developed to do so, which fall in two distinct classes. The interface tracking methods, such as boundary integral (Pozrikidis 2001), front-tracking (Tryggvason et al. 2001), immersed boundary (Mittal & Iaccarino 2005) schemes, all involve an explicit representation of the interface with dedicated grid or point sets, which allows for high accuracy, but makes it more difficult to handle topological changes, in particular for three-dimensional systems. In interface-capturing methods, such as volume-of-fluid (VOF), level-set and diffuse-interface methods, the representation of the interface is only implicit, with the benefit that arbitrary changes in interface shape can be treated with no further complication. Most such numerical methods have been extended to account for the presence of soluble or insoluble surfactants (see Teigen et al. 2011; Dieter-Kissling et al. 2015, for an overview).

In this work, our goal is to unveil the role of surfactants on the dynamics of a T1 event, a situation which has not been considered so far, and to investigate the mechanisms governing the rheology: surfactant diffusion along the interfaces and in the bulk, bulk/interface exchanges, viscous shear. This is only possible if the full surfactant distribution can be tracked in the bulk and along the interfaces, and requires numerical simulations. We
use a level-set approach (e.g., Sussman et al. 1994; Sethian 1999; Osher & Fedkiw 2003) extended to account for the presence of surfactants. Our configuration is the minimal one, both in terms of scale and dimensionality: we consider a unit cell of a semi-periodic arrangement in two dimensions. Our aim is to relate the local microscopic properties of the surfactants to the macroscopic foam rheology: for monodisperse crystalline foams, this intermediate situation is representative of a macroscopic foam; for disordered foams, it may still provide insight on the dominant dissipation mechanisms at play, and could be used to refine local ingredients for a multiscale approach.

The article is organized as follows. We first present in section 2 the equations governing the flow of the bubble assembly in presence of surfactants as well as the main dimensionless parameters and the configuration considered. Section 3 then briefly describes our level-set method and numerical implementation. Finally, we report in section 4 our results on T1 events, together with a specific discussion. A particular attention will be paid to the coupling between bulk flow and interfacial stress.

2. Shear of a bubble assembly: problem formulation

2.1. Configuration studied

The initial configuration, depicted in figure 2, consists of four hemicircular bubbles arranged on a hexagonal lattice and separated by a center-to-center distance $2H/\sqrt{3}$, where $H$ is the domain height. Considering the symmetry of the system, simulations were also done on a cluster of two half-bubbles only, which simply corresponds to one half of the domain represented in figure 2. Unless indicated otherwise, the liquid fraction of the system, defined as liquid area divided by total area, was set to $\psi_l = 30\%$. This large liquid fraction, much closer to bubbly liquids than to real foam, has been chosen to ensure numerical resolution of the resulting liquid film between the bubbles. The shear is imposed by prescribing a velocity $+U$ (resp. $-U$) for the top (resp. bottom) plate. The contact lines, where solid meets both liquid and gas, are assumed to be pinned and thus move at the plate velocity. The plates are impermeable to fluids and surfactants, with zero flux across them. Periodic boundary conditions are applied in the lateral direction.

2.2. Equations and relevant parameters

2.2.1. Flow dynamics and interfacial stress

The flow in the liquid and in the gas is governed by the full Navier-Stokes equations. Assuming that both fluids are incompressible and denoting by $\mathbf{u}$ the velocity, $\rho$ and $\mu$
the local density and viscosity, the equations read

\begin{align}
\nabla \cdot \mathbf{u} &= 0, \\
\rho \left( \frac{\partial \mathbf{u}}{\partial t} + (\mathbf{u} \cdot \nabla) \mathbf{u} \right) &= -\nabla p + \nabla \cdot \left( \mu (\nabla \mathbf{u} + (\nabla \mathbf{u})^T) \right) = \nabla \cdot \mathbf{\sigma},
\end{align}

with \( \mathbf{\sigma} = -p\mathbf{I} + \mu (\nabla \mathbf{u} + (\nabla \mathbf{u})^T) \) the stress tensor. Note that the viscosity and density depends on the position, since they are different in the liquid and in the gas. A no-slip boundary condition applies for the velocity at the plates. At a gas-liquid interface with surface tension \( \gamma \), the stress jump is (Pozrikidis 2011)

\[ [\mathbf{\sigma} \cdot \mathbf{n}] = -\gamma C \mathbf{n} - \nabla_s \gamma. \tag{2.3} \]

Here \([X] = X_{\text{liquid}} - X_{\text{gas}}\) is the jump of quantity \( X \) across the interface, \( \mathbf{n} \) is the normal unit vector pointing toward the liquid, \( C = -\nabla \cdot \mathbf{n} \) is the interface curvature, and \( \nabla_s = \mathbf{I}_s \cdot \nabla \) (with \( \mathbf{I}_s = \mathbf{I} - \mathbf{n} \otimes \mathbf{n} \) the surface identity tensor) is the surface gradient. The first term on the right-hand side of (2.3) represents the Young-Laplace normal stress jump across the interface whereas the second term corresponds to the tangential solutal Marangoni stress.

We adopt herein a one-fluid formulation (e.g., Brackbill et al. 1992), wherein the governing equations for both fluids and the stress jump condition are combined into one set of governing equations. This is facilitated by introducing the Dirac function \( \delta_T \) featuring the interface, and the interfacial stress tensor \( \mathbf{T}_s = \gamma \delta_T \mathbf{I}_s \). We use here the simplest possible form of the interfacial stress tensor, but more complex models involving complex surface shear or dilatational rheology (Sagis 2011; Erni 2011) could be described with this approach. The one-fluid formulation valid in the entire domain can then be written as

\[ \rho \left( \frac{\partial \mathbf{u}}{\partial t} + (\mathbf{u} \cdot \nabla) \mathbf{u} \right) = \nabla \cdot (\mathbf{\sigma} + \mathbf{T}_s) = \nabla \cdot \mathbf{\sigma} + \gamma C \delta_T \mathbf{n} + (\nabla_s \mathbf{\gamma}) \delta_T. \tag{2.4} \]

The singular terms added on the right-hand side represents the singular contribution arising from taking the divergence of the discontinuous stress (Teigen et al. 2011): Since \( \nabla \delta_T \) is normal to the interface, we indeed have \( \nabla \cdot \mathbf{T}_s = \gamma C \delta_T \mathbf{n} + (\nabla_s \mathbf{\gamma}) \delta_T = -\delta_T [\mathbf{\sigma} \cdot \mathbf{n}] \).

In dimensionless coordinates, the mass conservation reduces to \( \nabla \cdot \mathbf{u} = 0 \) and the conservation of momentum to:

\[
\hat{\rho} \left( \frac{\partial \mathbf{\hat{u}}}{\partial \hat{t}} + (\mathbf{\hat{u}} \cdot \nabla) \mathbf{\hat{u}} \right) = -\hat{\nabla} \hat{p} + \frac{1}{\text{Re}} \hat{\nabla} \cdot \left( \hat{\mu} (\hat{\nabla} \mathbf{\hat{u}} + (\hat{\nabla} \mathbf{\hat{u}})^T) \right) \\
+ \frac{1}{\text{Ca}_0 \text{Re}} \left[ \hat{\gamma} \hat{C} \mathbf{n} \delta_T + \nabla_s \hat{\gamma} \delta_T \right],
\]

where the characteristic parameters are the shear velocity \( U \) and the box height \( H \), yielding the typical timescale \( T = H/U \). The characteristic viscosity and density \( \mu_l \) and \( \rho_l \) are those of the liquid phase. The pressure is made non-dimensional with \( \rho_l U^2 \). \( \delta_T \) is the dimensionless Dirac function used to feature the interface. Regarding the interfacial tension, \( \hat{\gamma} = \gamma / \gamma_0 \) where \( \gamma_0 \) is the bare liquid/gas surface tension. We also introduced the Reynolds number defined as \( \text{Re} = \rho_l U H / \mu_l \), which compares inertial and viscous stresses, and a capillary number \( \text{Ca}_0 = \mu_l U / \gamma_0 \) which compares viscous and surface tension forces.
2.2.2. Model of surfactant dynamics and adsorption

To completely describe the dynamics of the fluids, one must fully account for the transport of surfactants in the liquid and along the interface. In the former, it reads

$$\frac{\partial F}{\partial t} + \nabla \cdot (uF) = D_F \nabla^2 F,$$

(2.6)

where $F$ is the volume concentration of surfactant and $D_F$ its diffusion coefficient in the bulk. At interfaces, we define the surface concentration of surfactants, denoted herein by $f$. This is also governed by diffusion and advection; the balance equation developed by Wong et al. (1996) is written here in a fixed reference frame. Upon supposing $f$ to be extended off interfaces as a constant along the interface normal, the balance equation for $f$ can be written as (Pereira et al. 2007; Teigen et al. 2009):

$$\frac{\partial f}{\partial t} + \nabla_s \cdot (uf) = D_f \nabla^2_s f + j,$$

(2.7)

with $D_f$ the diffusion coefficient along the interface. The source term $j$ accounts for the exchange of surfactants between the interface and the bulk and is assumed to be given by

$$j = r_a F^*(f_\infty - f) - r_d f,$$

(2.8)

with $r_a$ and $r_d$ the adsorption and desorption coefficient respectively and $F^*$ the bulk surfactant concentration in the vicinity of the interface. Regarding boundary conditions, the following equality applies at the interface

$$D_F \nabla F \cdot n = -j,$$

(2.9)

while at the plates, the no-flux condition ensuring surfactant conservation imposes

$$n_{pl} \cdot \nabla F = 0, \quad t \cdot \nabla f = 0,$$

(2.10)

where $n_{pl}$ is the vector normal to the plates and $t$ the vector tangential to the interface (see figure 2). Finally, the interfacial stress and surface tension depend strongly on the amount of surfactants adsorbed at the interface. To link the surface tension to the surface concentration of surfactants at the interface $f$, a simple choice is the Langmuir equation of state (Langevin 2014)

$$\gamma(f) = \gamma_0 \left[ 1 + \frac{RT f_\infty}{\gamma_0} \ln \left( 1 - \frac{f}{f_\infty} \right) \right],$$

(2.11)

with $R$ the ideal gas constant, $T$ the temperature and $f_\infty$ the surface concentration at saturation.

These equations can be recast in dimensionless form. First, the convection-diffusion of surfactants yields

$$\frac{\partial}{\partial \tilde{t}} (H_\epsilon \tilde{F}) + \tilde{\nabla} \cdot (H_\epsilon \tilde{F} \tilde{u}) = \frac{1}{Pe_F} \tilde{\nabla} \cdot (H_\epsilon \tilde{\nabla} \tilde{F}) - h \delta_f \tilde{\gamma},$$

(2.12)

with $Pe_F = UH/D_F$ the bulk Péclet number, which compares advection and diffusion. The adsorption depth $h = f_{eq}/(HF_{eq})$, with $F_{eq}$ and $f_{eq}$ the volume and surface concentrations at equilibrium, compares the amount of surfactant at the surface and in the bulk. Note that the Heaviside function $H_\epsilon$ has been introduced to account for the fact that the surfactants are only present in the liquid phase. The transport equation

\[\text{Note that an equivalent expression is } h = r_a f_\infty (1 - \chi)/(r_d H), \text{ which involves } r_a.\]
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along the interface reads

\[ \frac{\partial}{\partial t}(\tilde{f}\delta_r) + \nabla \cdot (\tilde{f}\delta_r \tilde{\mathbf{u}}) = \frac{1}{P_e_f} \nabla \cdot (\delta_r \nabla \tilde{f}) + \delta_r \tilde{j}, \]  

(2.13)

with \( P_e_f = U H / D_f \) the interface Péclet number and with the dimensionless source term

\[ \tilde{j} = Bi \left[ \frac{\chi}{1 - \chi} F_s \left( \frac{1}{\chi} - \tilde{f} \right) - \tilde{f} \right]. \]  

(2.14)

Here \( Bi = r_d H / U \) is the so-called Biot number, which compares the timescale of surfactant desorption with convection. Finally, the adimensionalized equation of state is

\[ \tilde{\gamma}(\tilde{f}) = 1 + \beta \ln(1 - \chi \tilde{f}), \]  

(2.15)

where \( \beta = RT f_\infty / \gamma_0 \) governs the sensitivity of surface tension versus surfactant concentration and \( \chi = f_{eq} / f_\infty \) corresponds to the ratio of surfactant concentration at equilibrium and at saturation.


3.1. The level-set function and the numerical scheme

3.1.1. Level-set method

The governing equations presented above must be supplemented by a method to determine the evolution of interfaces. For this purpose, we used a level-set scheme already developed in the case of multiphase flows in previous work (e.g., Sussman et al. 1994; Osher & Fedkiw 2003). It consists in the introduction of a level-set distance function noted \( \phi(x, t) \) whose sign defines the location of each phase:

\[ \phi(x, t) = \begin{cases} d & \text{in the liquid}, \\ -d & \text{in the gas}, \\ 0 & \text{along interface } \Gamma. \end{cases} \]  

(3.1)

where \( d \) is the closest distance to the interface. It allows to define the evolution of interface location versus time. The level-set function is advected by the flow at a velocity \( \mathbf{u} \) and then satisfies at the interface:

\[ \frac{\partial \phi}{\partial t} + \mathbf{u} \cdot \nabla \phi = 0. \]  

(3.2)

The fluid properties at each location then directly depend on the value of the level-set function and are defined as

\[ \rho(\phi) = \rho_l H_\varepsilon(\phi) + \rho_g (1 - H_\varepsilon(\phi)), \]  

(3.3)

\[ \mu(\phi) = \mu_l H_\varepsilon(\phi) + \mu_g (1 - H_\varepsilon(\phi)), \]  

(3.4)

with \( \rho_l, \rho_g, \mu_l, \mu_g \) respectively the liquid (gas) density and viscosity. \( H_\varepsilon(\phi) \) is a smoothed Heaviside function defined as

\[ H_\varepsilon(\phi) = \begin{cases} 0 & \text{if } \phi < -\varepsilon, \\ \frac{1}{2} \left[ 1 + \frac{\phi}{\varepsilon} + \frac{1}{\pi} \sin(\pi \phi / \varepsilon) \right] & \text{if } |\phi| \leq \varepsilon, \\ 1 & \text{if } \phi > \varepsilon. \end{cases} \]  

(3.5)

\[ 2\varepsilon \] thus corresponds to the width of the smooth interface. The Dirac function introduced above to feature the interface also directly depends on this Heaviside function by \( \delta \nabla \mathbf{n} = \nabla H_\varepsilon \). The normal vector is obtained as \( \mathbf{n} = \nabla \phi / |\nabla \phi| \) and the interface curvature as \( C = \)
In order to accurately determine the interface curvature and normal vector, and to keep the interface thickness nearly constant, a reinitialisation stage is included. Motivated by the results of a comparative study of various reinitialisation schemes (Solomenko et al. 2017), we use the interface-preserving algorithm of Sussman & Fatemi (1999). Thus, after the solution of (3.2) has been advanced over a timestep to yield a level-set function $\phi = \phi_0$, this is corrected by solving

$$\frac{\partial \phi}{\partial \tau} + \text{sgn}(\phi_0)(|\nabla \phi| - 1) = \lambda \delta(\phi_0) |\nabla \phi_0|$$  \hspace{1cm} (3.6)

over the pseudo time variable $\tau$, subject to the initial condition $\phi = \phi_0$; the coefficient $\lambda$ is chosen such as to preserve the volume of each fluid over any fixed volume of the two-phase flow (see Sussman & Fatemi 1999, for details).

### 3.1.2. Numerical implementation

The transport equations of surfactants were implemented in an already existing and validated level-set code (O’Náraigh et al. 2014), with some further improvements. This uses a standard projection method; the spatial discretisation is on a Marker-and-Cell (MAC) grid, with velocity components defined at cell faces and scalar quantities defined at cell centers. The momentum source term is discretised on the MAC grid. The temporal discretisation of the momentum equation involves a Crank-Nicolson scheme for diagonal viscous terms and third-order Adams-Bashforth scheme for convective terms and off-diagonal viscous terms; second-order central differences were used for the spatial discretisation of these terms. For temporal discretisation of the transport equation of the level-set function, a third-order Adams-Bashforth scheme was used; a fifth-order Weighted-Essentially-Non-Oscillatory (WENO) scheme was used for the spatial discretisation. At the reinitialisation stage (3.6), a second-order Runge-Kutta scheme was used for the temporal discretisation, and fifth-order WENO for spatial discretisation. Results of basic tests of the computational method without surfactants can be found elsewhere (Solomenko et al. 2017; O’Náraigh et al. 2014). The transport equations for surfactants have been implemented following Teigen et al. (2009), in the same manner as the viscous terms in the momentum equations.

The validation of the numerical scheme for flows with surfactants has been achieved by studying a droplet at rest and under shear, and by quantitative comparison with numerical results obtained by Teigen et al. (2009) with a different (diffuse-interface) method. We have also found in tests for static drops that surfactants did not amplify spurious currents. Those two points are elaborated in Appendix A. Finally, the mesh size $\Delta z$ is typically $H/200$ to $H/100$ and the total interface width $2\epsilon$ is chosen to be $3\Delta z$. Both were checked to have little effect on simulation results.

### 3.2. Potential pitfalls

#### 3.2.1. Surfactant leakage

The amount of surfactants is fixed in our configuration, since the plates are impermeable walls. From the no-flux condition (2.10), we obtain for our specific configuration that:

(i) at the liquid-plate boundary, the vertical flux of surfactant is zero: $\nabla F \cdot e_z = \partial_z F = 0$;

(ii) at the contact line, the tangential flux is zero: $\nabla f \cdot t = 0$, which can be rewritten as $\partial_z f = \frac{\partial_x f}{\partial_x z} \partial_z f$.

However, it turns out that these simple conditions led to a loss in the total amount of surfactants (~ 1% per bubble rearrangement). We thus constrained the boundary...
conditions to impose \( \partial_z f = \frac{\partial z}{\partial x} \frac{\partial}{\partial x} f \) and \( \partial_z F = \frac{\partial z}{\partial x} \frac{\partial}{\partial x} F \) at the contact line, and both \( \partial_z F = 0 \) and \( \partial_z f = 0 \) at the liquid-plate and gas-plate boundaries. We also consider that at the triple line, the bulk/interface exchange is inhibited by the presence of the wall \((j = 0)\). While still satisfying the physical boundary conditions, these conditions lead to a five-fold reduction of the surfactant leakage, which we consider acceptable. Besides, relaxing the last condition gives similar results, both on surfactant leakage and on the macroscopic force (section 4.1).

3.2.2. Pinning the contact lines

Besides the no-slip condition for fluid at the wall, the contact lines were pinned to the plates. Boundary conditions for the level-set function were implemented through prescription at ghost cells; at any time, the positions of contact lines being known, the level-set function was prescribed in a thin layer around the contact line accordingly. Not accounting for this, and merely relying on the no-slip condition instead, resulted in some deviations of the contact line position with respect to the plate displacement (1.3% per T1).

3.3. Parameter range

All dimensionless numbers are set to unity unless stated otherwise, implying that none of the physical effects is neglected. \( \chi \) is set to 0.3 to ensure that the value of surface tension with and without surfactants correspond to a reasonable system. We explore capillary numbers in the 0.02-0.3 range, Péclet numbers in the 0.1-100 range and Biot numbers in the 0.1-10 range. The viscosity and density ratios between the liquid and the gas are both set to 10. We also consider that surface and bulk Péclet numbers are equal \((Pe_F = Pe_f = Pe)\). Thus, our parameter ranges do not necessarily coincide with those expected in typical systems: for instance, the capillary number in experiments is usually smaller. Though supercomputing resources were used, our ability to explore parameter space is limited by the significant computing time required. We note, however, that with \( Ca, Bi \) and \( Pe \) numbers covering one or two decades, their influence can be clearly identified. Finally, it should be noted that the present numerical method and configuration are also relevant to describe neighbour-switching dynamics in emulsions composed of oil droplets in water (Seth et al. 2011). In this case, the ratio between the density of inner and outer fluids is close to 1, while the viscosity ratio can be either smaller or larger than 1.

4. Numerical results and discussion

In this section, we present the results of numerical simulations and discuss their physical interpretation. We conducted a parametric study that focuses mainly on the influence of the capillary number \( (i.e. \) the shear rate), the Péclet number \( (i.e. \) the ability of surfactant to diffuse both in bulk and along the interface) and the Biot number \( (i.e. \) the ability of bulk/surface exchange for surfactants). To reduce computation time, most of the simulations have been performed on half the domain of figure 2. All quantities shown below are for this two-bubble system.

4.1. Time variation of the system and forces

The temporal evolution of the interfaces is shown in figure 3 for parameters typical of our simulations \((Ca = 0.1, Bi = 0.1 \) and \( Pe = 1, \) or case B in table 1). One can observe that T1 processes indeed occur, and that a stationary regime is reached after two switches
Figure 3. Temporal evolution of the bubble configuration for \(Ca = 0.1\), \(Bi = 0.1\) and \(Pe = 1\) (case B in table 1). Lines correspond to the liquid-gas interface, i.e. \(\phi = 0\). The first snapshot corresponds to \(t = 1.82\) and the time interval between two successive snapshots is \(\Delta t = 0.19\).

Figure 4. Forces applied by the fluids on the rigid plates for \(Ca = 0.2\): solid lines: bottom plate, dotted lines: minus the forces on the top plate. From top to bottom: The orange lines correspond to the capillary force \(F_{cap}\), the black lines to the total force \(F_{tot} = F_{cap} + F_{vg} + F_{vl}\), the blue lines to the viscous force in the gas \(F_{vg}\) and the red lines to the viscous force in the liquid \(F_{vl}\). (a) Without surfactant, (b) with surfactants at \(Bi = 10\) and \(Pe = 1\).

\((t = 2/\sqrt{3} \approx 1.2\) in reduced variables). To be more quantitative, we turn to the forces exerted on the plates. The tangential force per unit width \(F_{tot}\) exerted by the fluids on the bottom plate can be computed as the integral over the plate of \(\mathbf{e}_x \cdot (\sigma + T_s) \cdot \mathbf{e}_z\). Whether surfactants are present or not, the total force \(F_{tot}\) can be separated in three contributions defined as follows:

(i) The capillary force at the contact lines \(F_{cap} = -\sum_{\text{contact lines}} \gamma n_z \text{sgn}(n_x)\), with \(n\) the normal to the interface,

(ii) The viscous drag force from the gas, \(F_{vg} = \int_{\text{gas-plate}} \mu_g (\partial u_x/\partial z) dx\),

(iii) The viscous drag force from the liquid, \(F_{vl} = \int_{\text{liquid-plate}} \mu_l (\partial u_x/\partial z) dx\).

The forces exerted by the fluids on the top plate are obtained by applying a minus sign to the formulas above. All those forces are represented in figure 4 as a function of time in a dimensionless form, i.e. they are normalized by the equilibrium surface tension \(\gamma_0\) in the surfactant-free case and \(\gamma_{eq}\) in the presence of surfactants. As a reminder, \(\gamma_{eq}\) is linked to \(\gamma_0\) by the Langmuir equation of state (2.11) and with our parameter values \(\gamma_0/\gamma_{eq}\) is equal to 1.554. Similarly, capillary numbers are defined with respect to equilibrium surface tension.

Whether surfactants are present or not, the variations of the force with time exhibit common features. First, after one \(T_1\), the forces reach a stationary state and periodically oscillate around a mean value. This oscillation coincides with the \(T_1\) process. In the following, we do not examine the transient regime and focus only on the steady state. Second, forces computed on the bottom and top plates coincide, as expected from symmetry. Finally, the contribution of the capillary force is largely dominant over the two viscous components.

4.2. Macroscopic rheology: force versus velocity

We now examine the dependence of the mean total force on the capillary number. As visible in figure 5, \(\langle F_{tot} \rangle\) increases with \(Ca\) and \(Pe\), but decreases with \(Bi\). Large
Figure 5. Mean total force $\langle F_{\text{tot}} \rangle$, divided by the equilibrium surface tension $\gamma_{\text{eq}}$ as a function of the capillary number $Ca$. The different symbols correspond to: black triangles: no surfactants; red squares: $Pe = 1$, $Bi = 10$; blue circles: $Pe = 1$, $Bi = 0.1$; orange diamonds: $Pe = 100$, $Bi = 10$. The solid line corresponds to $\langle F_{\text{tot}} \rangle/\gamma_{\text{eq}} \sim Ca$, the dashed line to $\langle F_{\text{tot}} \rangle/\gamma_{\text{eq}} \sim Ca^{2/3}$ and the dotted line to $\langle F_{\text{tot}} \rangle/\gamma_{\text{eq}} \sim Ca^{1/2}$.

Péclet numbers imply that surfactants are significantly advected by the flow, while small Biot numbers involves slow adsorption compared to the flow timescale. In both cases, we expect a less homogeneous distribution of surfactants on the interface, hence the existence of Marangoni stresses, and a larger force. The origin of this force increase is discussed below. Now, for fixed Biot and Péclet numbers, the force appears to vary as a power law of the capillary number $\langle F_{\text{tot}} \rangle/\gamma_{\text{eq}} \sim Ca^n$, with an exponent $n$ between 0.5 and 1. Large $Pe$ seems to yield a smaller exponent, though the investigated range in $Ca$ is too narrow to reach a clear-cut conclusion. Sublinear scalings are common in phenomena coupling viscous flows and surface tension, like the Landau-Levich problem (Landau & Levich 1942) or the sliding of bubbles against a wall (Bretherton 1961; Aussillous & Quéré 2002; Hodges et al. 2004). However, as underlined by Cantat (2013), several sublinear contributions can superimpose, making it difficult to identify the main dissipation mechanisms in the problem.

4.3. Velocity field, viscous dissipation and surfactant distribution

To get insight in the mechanisms at stake during the T1 process, we now characterise the local quantities. We do so for the four illustrative cases A-D given in table 1: they all have $Ca = 0.1$ but they differ in their Biot and Péclet numbers. The velocity fields, symbolized by arrows, are shown in figure 6. A derived quantity is the local rate of viscous dissipation defined as (see also Appendix B)

$$D_{v,\text{loc}} = \nabla \cdot \left[ \mu (\nabla u + (\nabla u)^T) \right],$$

(4.1)

which is shown in figure 7. Finally, the concentration in surfactant is plotted in the bulk (figure 8) and along the interface (figure 9). The snapshots are taken at the following instants: time of the third minimum ($t_1$), average value ($t_2$), maximum ($t_3$) and average value ($t_4$) of the capillary force as defined in figure 4. For convenience of discussion,
Table 1. Parameters for the illustrative cases considered in figures 6, 7, 8 and 9.

<table>
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<th>Surfactants</th>
<th>Bi</th>
<th>Pe</th>
<th>Σ</th>
<th>⟨D_v⟩</th>
<th>⟨D_s⟩</th>
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<td>1</td>
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<td>10</td>
<td>100</td>
<td>1.4</td>
<td>19.1</td>
<td>1.8</td>
</tr>
</tbody>
</table>

Figure 6. Velocity field at times \(t_1\), \(t_2\), \(t_3\) and \(t_4\) (from left to right) defined in the text and in figure 4 for the cases A, B, C and D (from top to bottom) described in table 1. Arrows represent velocity vectors. The green dotted circle indicate the zone of extensional flow, while the red dashed circles show sheared films.

we identify the three distinct types of film (see figure 2): the stretched film, which increases in thickness, the squeezed film, which decreases in thickness, and the adjacent film separating bubbles laterally.

Our main observations are as follows:

(i) In the absence of surfactant, the velocity field is of elongational nature (see encircled stretched film of case A at instant \(t_3\) in figure 6), the velocity vector being almost normal to the interface. The viscous dissipation is significant in the stretched film, but remains low in the squeezed film (see figure 7).

(ii) The interfacial profile of surfactant, that is the surface concentration \(f(s)\) along the
the curvilinear coordinate \( s \), barely evolves with time during the bubble rearrangement. This is shown in figure 9a for case B, but applies more generally.

(iii) At intermediate Péclet number \( (Pe = 1) \), the effect of the Biot number which characterises the timescale of bulk/interface surfactant exchange (adsorption/desorption) is investigated by comparing case B \( (Bi = 0.1) \) and case C \( (Bi = 10) \). The influence of the Biot number remains limited when comparing the velocity profile and viscous dissipation distribution, which are quite similar in the two situations. Looking now at the surfactant distribution along the interface, we see in figure 9b that it is not homogeneous. This implies that Marangoni stresses are generated at the interface but they are not sufficient to induce a rigid-like behaviour of the interfaces for the flow (see below). Besides, the inhomogeneities in surfactant distribution are more pronounced at the interface (resp. in the bulk) for small (resp. large) \( Bi \). This can be understood as follows. At small \( Bi \), the bulk/interface exchanges are too slow to occur during the course of a rearrangement, while at large \( Bi \), the gradient in interfacial concentration leads to surfactant desorption (resp. adsorption) in the zone enriched (resp. depleted) in surfactants, which smoothes interfacial gradients but induces bulk inhomogeneities.

(iv) The effect of Péclet number is investigated by comparing cases C \( (Pe = 1) \) and D \( (Pe = 100) \). In the latter, new features appear in the flow. The direction of the velocity seems to be more parallel to the interface, the viscous dissipation is not only located in the stretched film but also in the squeezed one. The shear occurring in the latter (see squeezed film of case D at time \( t_1 \) and \( t_4 \) in figure 6) can only be supported by Marangoni stresses at the interface, where we consistently observe large gradient in
Figure 8. Bulk surfactant concentration $F$ at times $t_1$, $t_2$, $t_3$ and $t_4$ (from left to right) defined in the text and in figure 4 for the cases B, C and D (from top to bottom) described in table 1. Note that for clarity, the $F$-scale has been truncated to values below 1.7, while $F$ actually takes larger values in case D.

Figure 9. (a) Interfacial surfactant concentration $f$ as a function of the curvilinear coordinate $s$ (defined in figure 2 and in the inset) for case B (see table 1) at instants $t_1$ (pink dash-dotted line), $t_2$ (solid black line), $t_3$ (red dashed line), $t_4$ (blue dotted line) defined in the text and in figure 4. Inset: snapshot of the interfacial surfactant distribution ($f$ is color-coded) at instant $t_3$. (b) $f$ as a function of $s$ at instant $t_3$ for cases B (red dashed line), C (solid black line) and D (blue dotted line) (see table 1).

These large interfacial gradients also lead to large bulk inhomogeneities in surfactant distribution. It seems that in this case, shearing of the liquid is the main mechanism of foam flow. This last point echoes recent experimental observations in another geometry where switching from elongational profile to shearing has been observed varying the surfactant properties (Petit et al. 2015) or deformation rate (Seiwert et al. 2013).
4.4. Surface or bulk dissipation

4.4.1. Bulk dissipation

Our qualitative observations indicate that tuning the surfactant properties (desorption rate, diffusivity) not only affects the elongation of the interfaces and the surfactant distribution, but also the nature of flow, by changing the interfacial boundary condition from mobile to rigid-like interface. To put this point on a quantitative basis, we first calculate the total viscous dissipation $D_v = \int_V D_{v,loc} dV$, with $D_{v,loc}$ given by (4.1). The non-dimensional and temporally averaged viscous dissipation rate $\langle \tilde{D}_v \rangle$ is reported in figure 10 as a function of $Pe$ for $Ca = 0.1$ and $Bi = 1$. It exhibits an increase by 60% in our range of Péclet number at fixed capillary number (i.e., the shear rate). This is only possible if the nature of the flow is fundamentally changing. For shear to exist in the films, the interfacial stress needs to be large enough to sustain the viscous stress at the interface. To verify that this is actually the case, we introduce the ratio

$$\Sigma = \frac{h_{\text{min}}|\nabla_s \gamma|_{\text{max}}}{\mu U}.$$  \hspace{1cm} (4.2)

Here, $h_{\text{min}}$ is the minimum thickness reached during the T1 by the squeezed or stretched films. $\Sigma$ thus compares the maximum value of Marangoni stress $|\nabla_s \gamma|_{\text{max}}$ along the interface to the typical viscous stress for sheared films $\mu U/h_{\text{min}}$. $\Sigma$ is zero for (mobile) stress-free interfaces, while it should be of order 1 for rigid-like interfaces sustaining shear. Values of $\Sigma$ are reported for A-D cases in table 1, and are plotted as a function of the Péclet number in figure 11a. One can indeed observe that the surface stress becomes comparable to the viscous stress as soon as Péclet number reaches 2, in full agreement with our previous observations.

We now turn to the influence of the capillary number on the viscous dissipation (figure 12a). This plot shows that the (time averaged) dimensionless viscous dissipation $\langle \tilde{D}_v \rangle$ decreases with the capillary number. This may seem counter-intuitive as one expects the dissipation to increase with the velocity, but it should be kept in mind that the actual viscous dissipation corresponds to $\langle D_v \rangle = \rho_l U^3 H \langle \tilde{D}_v \rangle = \mu_l U^2 \langle \tilde{D}_v \rangle$ since we have $Re = 1$. 

**Figure 10.** Temporally averaged viscous dissipation $\langle \tilde{D}_v \rangle$ (red circles) and injected power $\langle \tilde{P}_{\text{inj}} \rangle$ (black squares) as a function of the Péclet number for $Ca = 0.1$ and $Bi = 1$. The dashed line corresponds to the injected power $\langle \tilde{P}_{\text{inj}} \rangle$ for the case without surfactants.
Figure 11. (a) Ratio $\Sigma$ of Marangoni to viscous shear stress, as defined by (4.2), as a function of $Pe$ for the simulations of figure 10. (b) Ratio $\langle \tilde{D}_s \rangle / \langle \tilde{D}_v \rangle$ of the effective surface dissipation over viscous dissipation, as a function of Péclet number, for the same simulations.

Figure 12. (a) Average viscous dissipation $\langle \tilde{D}_v \rangle$ as a function of $Ca$ for different Péclet and Biot numbers (same symbols as in figure 5). (b) Ratio $\langle \tilde{D}_s \rangle / \langle \tilde{D}_v \rangle$ of the effective surface dissipation over the bulk viscous dissipation, as a function of $Ca$ (same symbols as in figure 5).

This quantity indeed increases with the capillary number but slower than $Ca^2$, as in the problem of a bubble sliding against a plane (Cantat 2013). This is a consequence of the sublinear behavior of the total force as a function of $Ca$.

Finally, based on our estimation of the viscous dissipation, we can compare the relative contributions of inertia and viscosity in the flow. As explained in Appendix C, and although the Reynolds number is fixed to unity, we have found that the role of inertia remains limited in our simulations, presumably because of the confinement of the liquid films.

4.4.2. Work by surface tension

We have shown that the liquid flow is of a different nature when the surfactant properties are modified and that at high Péclet numbers, the distribution of interfacial surfactants becomes increasingly inhomogeneous, and the viscous dissipation increases. Now, if we compute the injected power $P_{\text{inj}}$ (as the plate velocity times the applied force), it is apparent in figure 10 that $\langle P_{\text{inj}} \rangle$ may be larger than the average viscous dissipation. The difference reveals the work done by surface tension $D_s = \int_{\Gamma} \gamma (\nabla_s \cdot \mathbf{u}) dS$, as discussed in Appendix B. This term can be seen as the 2D analogue of the work done by pressure forces in compressible fluids. Numerically, the evaluation of this interfacial quantity is delicate. However, in steady state, the variation of kinetic energy over a rearrangement period should be zero, so that the time-averaged work by surface tension $\langle D_s \rangle$ can be readily calculated as $\langle P_{\text{inj}} \rangle - \langle D_v \rangle$. Looking at $\langle D_s \rangle$ as a function of $Pe$, we see in figure 10
that it tends to zero with vanishing Pe (in this case, the interface is homogeneous, and
the surface tension constant, as in the absence of surfactants), but that otherwise, this
contribution is positive. This suggests that the work by surface tension may be seen as
an effective dissipation on the surfactant-covered interface (hence our notation $D_s$). Note
that the interfaces considered here have no intrinsic surface viscosities and thus lack
the corresponding additional term. However, even without this effect, it is of interest to
investigate which contribution, surface or viscous, dominates the dissipation.

Figure 11b shows the ratio $\langle \tilde{D}_s \rangle / \langle \tilde{D}_v \rangle$, as a function of the Péclet number for $Bi = 1$ and
$Ca = 0.1$. One can observe a maximum ($\approx 0.5$) reached at intermediate Péclet numbers
($\approx 1$). This result is reminiscent of the famous model of Lucassen & van den Tempel
(1972) for interfacial viscoelasticity: assuming instantaneous adsorption/desorption but
diffusion in the bulk, it predicts that the interfacial loss modulus—which characterises
the viscous response of the interface—vanishes at both high and small frequencies
(analogous to large and small Péclet number respectively), but goes through a maximum
at intermediate frequencies. Figure 12b shows the dependence of $\langle \tilde{D}_s \rangle / \langle \tilde{D}_v \rangle$ as a function
of the capillary number $Ca$ for different $Bi$ and $Pe$ parameters. The values of $\langle \tilde{D}_s \rangle$ and
$\langle \tilde{D}_v \rangle$ for the illustrative cases A-D are also reported in table 1. No clear tendency is visible
as the capillary or Biot numbers are changed but the effect of Péclet number evidenced
above is recovered†.

To recapitulate, the key point is that for all systems considered here, the viscous
dissipation is always dominant. This conclusion holds even if the pattern of dissipation
is changing with the surfactant properties. At low Péclet number, the interfaces are not
able to sustain the viscous stress and the flow is mainly elongational. At intermediate
Péclet number, the ratio between surface and bulk dissipation reaches a maximum.
This is the threshold above which interfacial stress sustains viscous shear, and viscous
dissipation starts to increases with the Péclet number (figure 10). These results are
supported by recent experiments in a model system (Petit et al. 2015) where the flow
in the film is observed to be of elongational or shear types depending on the surfactant
nature. However, the resulting picture presented here is very different from other scenarii
previously put forward in the literature for dissipation. In particular, Tcholakova et al.
(2008) showed that surface viscoelasticity can dominate the rheology and the total
dissipation in macroscopic foams, while Durand & Stone (2006) and Bianca et al. (2009)
also concluded that it plays the major role in their experiments of T1 dynamics. These
point to the role of intrinsic surface viscosities, known to be large in some experimental
systems (Golemanov et al. 2008).

As a final remark, we recall that the liquid fraction in our system is high ($\psi_l = 30\%$),
contrary to experiments; we anticipate that the dominant source of dissipation may
change at low liquid content. To test this point, we performed additional simulations at
$Pe = 1$, $Bi = 1$ and $Ca = 0.1$, for various liquid fractions in the range $\psi_l = 17 - 40\%$.
These results are shown in figure 13: We find that the injected power $\langle \tilde{P}_{inj} \rangle$ decreases
with increasing liquid fraction $\psi_l$, while the viscous dissipation $\langle \tilde{D}_v \rangle$ barely changes. This
indicates that the effective surface dissipation $\langle \tilde{D}_s \rangle$ indeed becomes larger for drier foams,
as shown in figure 13b. However, the liquid fraction range remains limited and additional
numerical developments are necessary to reach realistic liquid fractions, as explained in
the conclusion.

† Note that points for the no-surfactant system are also shown in figure 12b: in this case,
the effective surface dissipation should be zero, as the surface tension is uniform. We attribute
the comparatively small scatter of the numerical points around the zero value to inaccuracy in
the calculation of viscous dissipation due to the non-negligible width of the interfaces in our
simulations.
4.5. Film thickness and film rupture

Finally, we examine how the liquid distribution is modified by the coupling between flow and surface tension effects. An observation of the interfaces in the illustrative cases A-D (see figure 6 for instance) indicates that the thickness of liquid films depends on the parameter set. To analyse this point quantitatively, we extract for each simulation the minimum thickness of sheared films $h_{\text{min}}$ (which separate top bubbles from bottom ones), as well as $h_{\text{min,adj}}$ the minimum thickness between adjacent (top-top or bottom-bottom) bubbles (see figure 2). Figure 14 reports both thicknesses as functions of $Ca$ and $Pe$. We observe that $h_{\text{min}}$ increases with $Ca$, as in the problem of bubbles sliding along a rigid plane (Cantat 2013). Meanwhile, due to volume conservation, $h_{\text{min,adj}}$ is reduced. Besides, for larger $Pe$, $h_{\text{min}}$ also increases, as expected for more rigid-like interfaces. These observations have implications for the (in)stability of foams against coalescence. It has been shown (Biance et al. 2011) that foam collapse, i.e. the occurrence of coalescence avalanches, can be triggered by bubble rearrangements. The suggested mechanism is that some liquid films might become extremely thin in the course of the T1 event, leading to its breaking. Our results may provide information on the probable location of film breaking, which might prove valuable in further studies of foam collapse.
5. Conclusion

We presented numerical simulations of T1 events in a 2D semi-periodic system of sheared bubbles. The level-set method employed fully accounts for the coupling between the flow dynamics—inhomogeneous Navier-Stokes equations—and the dynamics of surfactants—bulk and interfacial diffusion, as well as bulk/interface exchanges.

Our approach offers a detailed view of all local fields in the course of a bubble rearrangement and enables us to investigate the relative importance of diffusion, desorption/adsorption, viscous effects and surface tension through a parametric study of Péclet, Biot and capillary numbers. A key result of our simulations is that surfactant inhomogeneities are able to build up Marangoni stresses at the liquid-gas interfaces, with deep consequences on the nature of viscous dissipation, including the appearence of shear flow in the liquid films between the bubbles. We also identify regimes where the work by surface tension leads to an effective surface dissipation, which is reminiscent of the effective surface viscosity developed in the classical model of surface dilatational viscoelasticity of Lucassen & van den Tempel (1972). Besides, we show that as the capillary number increases, the mechanical response of the system is strongly affected by the surfactant dynamics, thus providing microscopic insight to understand foam or emulsion rheology. Finally, our detailed view of liquid distribution in films may provide hints on the possible location for film rupture, and subsequent foam collapse.

Perspectives of this work are many, but are all motivated by bringing our simulated systems closer to real foams. So far, the limitations of our system are the following: (i) the geometry is two-dimensional; (ii) the density and viscosity ratios between the liquid and the gas have been set to 10, which implies that the effect of the gas is not negligible as we would expect in real foams; (iii) we only consider the case of bubbles pinned against a solid wall, which hinders the flow and surfactant dynamics as compared to bubbles in the bulk of a foam; (iv) we consider a very wet system, with unrealistically thick films. Regarding the first two points: extension to three-dimensional systems and more realistic density and viscosity values is straightforward but will prove demanding in terms of computation time. The third point can be addressed by imposing some pseudo-periodic boundary conditions in the vertical direction—accounting for the opposite velocities between top and bottom edges. In contrast, reaching the low liquid fractions characteristic of actual foams will require new developments, including additional terms to prevent bubble coalescence upon contact. In actual foams, coalescence is prevented by non-hydrodynamical forces, such as disjoining pressure (Israelachvili 2010) (including van der Waals, electrostatic or steric interactions), which are short-ranged and thus computationally expensive. These non-hydrodynamic forces are also expected to be crucial in bubble coalescence, which would deserve further studies to understand how foam collapse can be triggered by bubble rearrangements. Finally, future work will also account for intrinsic surface dilatational and shear viscosities. This would allow to assess the relative importance of these contributions in foam rheology, and open the way to a microscopic understanding of the link between foam macroscopic behavior and interfacial properties.

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Figure 15. Simulations of a sheared droplet: (a) Droplet shape (black) and bulk surfactant concentration (colors) at $t = 2H/U$ for $Bi = 20$, $Pe_F = 8$ and all other parameters set to values similar to Teigen et al. (2011). (b) Interfacial surfactant concentration $f$ as a function of the normalized curvilinear coordinate $s/s_{\text{max}}$ for three parameter sets corresponding to increasing inhomogeneities: (blue) $Bi = 20$ and $Pe_F = 8$; (red) $Bi = 2$ and $Pe_F = 80$; (black) $Bi = 0.2$ and $Pe_F = 8$. Solid lines are results from our level-set simulations, while the dashed lines were obtained by Teigen et al. (2011) using a diffuse-interface approach. The coordinate $s$ increases clockwise and its origin is shown by the white cross in (a).

Appendix A. Validation of the numerical method

Several validations have been conducted on the numerical method. First, as regards numerical stability, we first investigated the generation of spurious currents (Solomenko et al. 2017), using as a test configuration a bubble at rest. Not only were those currents found of negligible magnitude, with a relative velocity variation below $10^{-3}$ but the presence of surfactants actually results in a damping of these unwanted velocity fluctuations. Second, we observed with the initial implementation a significant drift in the total amount of surfactants; the problem was overcome by adjusting the boundary conditions on the plate as described in Sec. 3.2.1. Finally a quantitative comparison was made to the results of Teigen et al. (2011) obtained with a diffuse-interface method. The configuration is close to ours, with a single bubble sheared by the motion of bottom and top plates (figure 15a). Shown in figure 15b is the surfactant concentration $f$ as a function of the position $s$ along the interface for several values of the Biot and Péclet numbers. Our curves are in close agreement with those of Teigen et al. (2011).

Appendix B. Viscous dissipation and work by surface forces

In the following, we detail the different contributions to dissipation in our system. The total variation of kinetic energy $K$ is given by

$$\frac{dK}{dt} = \int_V \rho u \cdot \frac{Du}{Dt} dV. \quad (B1)$$

Substituting the acceleration term from the generalized Navier-Stokes equation (2.4) with the total stress tensor $T_{\text{tot}} = \sigma + T_s$, we obtain

$$\frac{dK}{dt} = \int_V u \cdot (\nabla \cdot T_{\text{tot}}) dV, \quad (B2)$$

or

$$\frac{dK}{dt} = \int_V \nabla \cdot (u \cdot T_{\text{tot}}) dV - \int_V (\nabla u : T_{\text{tot}}) dV. \quad (B3)$$
Using Green-Ostogradsky theorem, the first term can be rewritten as an integral over the volume boundary $S$, here the top and bottom plates:

$$
\int_V \nabla \cdot (u \cdot T_{\text{tot}}) dV = \int_S u \cdot T_{\text{tot}} \cdot n dS = \int_{S,\text{top}} UT_{\text{ext,top}} dS - \int_{S,\text{bot}} UT_{\text{ext,bot}} dS, \tag{B 4}
$$

with $T_{\text{ext}}$ the stress exerted on the fluid at the plates and $U$ the plate velocity. Since the total stress includes a bulk contribution (pressure and viscous stress) and a surface one (interfacial stress tensor), this results for the top plate in a viscous contribution, which reads per unit width,

$$
UF_v = \int_{S,\text{top}} \mu \frac{\partial u_z}{\partial z} dx, \tag{B 5}
$$

and a capillary one

$$
UF_{\text{cap}} = \sum_{\text{contact lines}} U \gamma \cos(\theta), \tag{B 6}
$$

with $\theta$ the angle between the normal to the interface and the $z$-axis. The two above equations are used in sections 4.1 and 4.2 to compute the forces at the wall. Note also that equation (B 4) corresponds to the instantaneous injected power $P_{\text{inj}}$.

We now focus on the second term in the right-hand side of (B 3), which represents the dissipation rate in the system,

$$
\mathcal{D} = \int_V (\nabla u : T_{\text{tot}}) dV. \tag{B 7}
$$

Using the expression for the total stress $T_{\text{tot}}$ and the fact that the flow is incompressible yields

$$
\mathcal{D} = \mathcal{D}_v + \mathcal{D}_s = \int_V \nabla u : \left[ \mu (\nabla u + (\nabla u)^T) \right] dV + \int_V \nabla u : T_{s} dV. \tag{B 8}
$$

Whereas the first term $\mathcal{D}_v$ corresponds to usual viscous dissipation as computed in section 4.4.1, the second term $\mathcal{D}_s$ is a contribution from the liquid-gas interfaces which can be rewritten as

$$
\mathcal{D}_s = \int_{\Gamma} \nabla u : (\gamma I_s) dS = \int_{\Gamma} \gamma (\nabla_s \cdot u) dS. \tag{B 9}
$$

$\mathcal{D}_s$ can be interpreted as the rate of change of surface energy (Dangla 2012) or as the instantaneous work done by surface tension, a 2D analogue of the work done by pressure forces in compressible fluids. In general, this quantity can take either positive values (e.g. if interfaces are being stretched) or negative values (e.g. for homogeneous $\gamma$ if interfaces are being compressed). Note, however, that if surface dilatational viscosity $\mu_s$ were incorporated in the expression of the interfacial stress tensor $T_s$ (Erni 2011; Sagis 2011), this would contribute to $\mathcal{D}_s$ as $\int_{\Gamma} \mu_s (\nabla_s \cdot u)^2 dS$ which is always positive, and indeed corresponds to a true dissipation term.

As the shear-driven T1 events considered in our simulations quickly reach a steady state, we also consider the energy balance (B 3) after time-averaging over one period. Since the kinetic energy is constant on average ($\langle (dK/dt) \rangle = 0$), this reduces to

$$
\langle P_{\text{inj}} \rangle = \langle \mathcal{D}_v \rangle + \langle \mathcal{D}_s \rangle, \tag{B 10}
$$

which we use in the present study to estimate the average surface work $\langle \mathcal{D}_s \rangle$. For interfaces with constant surface tension $\gamma_0$, $\langle \mathcal{D}_s \rangle = \gamma_0 \langle \int_{\Gamma} (\nabla_s \cdot u)^2 dS \rangle$ would vanish as the variation in total length averages to zero. However, for surfactant-laden interfaces, we found that $\langle \mathcal{D}_s \rangle$ can be significant in magnitude ($\approx \langle \mathcal{D}_v \rangle/2$), and is generally positive, which we interpret as an effective surface dissipation.
Appendix C. Influence of inertia

Our simulations were conducted for $Re = \mu U H / \rho l = 1$. The Reynolds number compares inertial and viscous effects, so that inertia is not negligible a priori. However, in our configuration, viscous flows are mostly confined in the liquid films which separate bubbles, and viscosity may thus dominate inertia. To quantify this point in the illustrative case A (no surfactants, $Ca = 0.1$, see table 1), we consider the total kinetic energy of the system $K = \int dV \rho |u|^2$, which is reported in dimensionless form as a function of time in figure 16. As expected, we find that, after a transient, the kinetic energy oscillates as rearrangements proceed. The average kinetic energy is $\langle \tilde{K} \rangle \approx 0.3$ whereas the oscillation amplitude is of order 0.1 and its period $\approx 0.6$, from which we obtain a rough estimate of $d\tilde{K}/dt \approx 0.2$. To assess the importance of inertia compared to viscous forces, we finally compare this value to the average viscous dissipation in the same simulation, which is $\langle \tilde{D}_v \rangle = 8.9$ (table 1): as a first approximation, the effect of inertia can be neglected in the present simulations.

REFERENCES


Level-set simulations of a 2D topological rearrangement in a bubble assembly


