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Kernel functions to flotation bubble size distributions

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ABSTRACT

Flotation modelling has advanced from deterministic single particle-bubble models into using such models to solve flotation systems by using modern computational techniques. The step from a single particle–single bubble event to multiple events taking place in the large computational volume like a flotation cell poises the challenge of handling bubble and particle distributions in all computational cells.

The estimation of bubble size has either been omitted (constant size) or has been lately estimated by a population balance approach. The physical performance of flotation is exessively determined by the bubble size distribution (BSD). Therefore, the bubble size distribution estimate is crucial for modelling. Although the BSD can be measured, the underlying effects of different variables causing changes in break-up and coalescence rates producing changes in the measured BSD’s are not well understood. This paper discusses the profound effects frothers have on both the coalescence and break-up of gas bubbles.

Depending on the bubble surface stiffness caused by frother adsorption, the drainage rate of fluid between two approaching bubbles is very different. Frothers like DF250 and Pentanol have a higher coalescence rate than frothers like DF250 and NF240.

Break-up is shown to be a function of the dynamic surface tension, not the static surface tension. Fast adsorbing frothers (DF200) have at very short time scales a higher rate of break-up.

The paper suggests a division of frothers into two distinct classes for modelling purposes. Those with fast adsorption and desorption, which leave the gas-air interface mobile and those frothers that by slower adsorption and desorption create stiff interfaces. The effects in real systems may be more varied. The modelling of subtler frother effects will not substantially improve modelling quality.

1. Introduction

Flotation modelling has the challenge to combine several physico-chemical phenomena into a concise model framework. The main body of modelling has been related to the well-known first order reaction model. During the years this approach with its additions and improvements has proven to be a good simple engineering model to be fitted with batch flotation data results e.g. a plug flow reactor.

\[
dC \frac{dt}{dt} = -kC.
\] (1)

The rate constant k can be obtained experimentally for any given steady-state condition. However, to formulate the dependence of the rate constant from process variables has turned out to be difficult. There have been attempts to relate the rate constant to both local and global parameters (Jameson et al., 1977) and to link the rate constant to the probabilistic bubble-particle encountering by the time averaged bubble horizontal interface flux (termed bubble surface area flux $S_b$) (Gorain et al., 1995a,b, 1996, 1999). To make the challenge more tractable, the total process has been generally divided into probabilistic sub-processes as outlined first by Gaudin (1932) and in more detail by Sutherland (1948). The “total probability of flotation” consists of the sub-process probabilities e.g. particle-bubble collision, attachment and stability (detachment)

\[
k = P_1 S_b
\] (2)

The flotation rate $k$ is then modelled as a product of the “total probability of flotation” and the frequency that bubbles and particles interact (come so close to each other for the above mentioned sub-processes to take place e.g. collide).

\[
Z P_l = Z P_l P_r P_i = k
\] (3)

There are several deterministic models for the particle-bubble collision (Gaudin, 1932; Sutherland, 1948; Yoon and Lutrell, 1989;
Population balance models can be solved either by the method of classes, where the NDF is discretized into a number of classes. Each class leads to an equation considering all the processes that will affect the units of the said class. The issue rising is mainly the bubble breakage and bubble coalescence. Each class would need its own kernel functions. The benefit is the natural reconstruct of the NDF. It is, however, computationally very expensive and time consuming to be used in flotation simulation. The other method to solve the bubble PBM is the method of quadrature of moments (QMOM) (McGraw, 1997; Marchisio and Fox, 2005; Bhutani, 2016). In the method the moments of the NDF are solved from the available transported moments. As is discussed by Bhutani (2016), for an approximate estimate of NDF four moments is often sufficient. For this approach a set of kernel functions would be sufficient. This, however, requires robust kernels.

Bubble break-up and coalescence kernels have been studied extensively for bubble columns (Prince and Blanch, 1990a; Luo and Svendsen, 1996; Martinez-Bazan et al., 1999, 2010; Lehr and Mewes, 2001; Wang et al., 2003; Zhao and Ge, 2007; Liao and Lucas, 2009, 2010; Solvik and Jakobsen, 2015 among others). These studies have performed with electrolytes. They differ in several important aspects from flotation systems with varying chain length surface active reagents (frothers) and solids.

2. Bubble break-up kernels

The Martinez-Bazan et al. (2010) model assumes that a pair of bubbles can be formed, when the stresses caused by turbulence are larger than the stresses opposing deformation at the length scales corresponding to the mother bubble. There exists always a critical bubble size at each level of turbulent intensity that cannot be broken up. This critical size is expressed as

$$h_\text{crit} = \epsilon^{-2/3} \left( \frac{12 \lambda_s}{\beta \rho} \right)^{1/3},$$

(8)

where $\epsilon$ is the turbulent energy dissipation, $\lambda_s$ surface tension (without surfactant), $\rho$ the density and $\beta$ a constant. The probability of break-up is (Martinez-Bazan et al., 2010) for a bubble with a volume $V$ is as follows

$$P_V(V) \propto \left( \frac{1}{2} \beta (V h_b)^{3/2} \right) \left( 1 - V^{1/3} - \Lambda^{2/3} \right) \left( 1 - V^2 - \Lambda^{4/3} \right),$$

(9)

where $h_b$ is the mother bubble diameter, $V$ its volume and $\Lambda$ the ratio between the mother bubble size $h_b$ and the critical bubble size $h_\text{crit}$. The obtained bubble size distribution is (Martinez-Bazan et al., 2010):

$$f^*(b^*) = \frac{b^{2/3} (b^{2/3} - \Lambda^{2/3}) (1 - b^{3/2} - \Lambda^{4/3})}{\int_{\text{max}}^{\text{min}} b^{2/3} (b^{2/3} - \Lambda^{2/3}) (1 - b^{3/2} - \Lambda^{4/3}) db^*},$$

(10)

If surfactants are added to the solution in dilute concentrations their adsorption on the gas-water interphase will change the surface tension in a linear way

$$\lambda - \lambda_s = \Gamma \cdot \sigma,$$

(11)

where $\lambda_s$ is the surface tension without the surfactant. $\Gamma$ is the surface concentration of the surfactant (mass of surfactant per unit area of surface). Following the linear relationship, one can write for a small change in the surfactant to have an effect on the surface tension as follows (Stone and Leal, 1990)

$$\lambda' - \lambda_s = (1 - \varphi).$$

(12)

This will change the critical bubble size of Eq. (8) to

$$h_\text{crit} = \epsilon^{-2/3} \left( \frac{12 \lambda_s (1 - \varphi)}{\beta \rho} \right)^{1/3}.$$

(13)

As Martinez-Bazan et al. (2010) point out, the parameter $\Lambda$ in Eq.
can be interpreted as the inverse of the mother bubble Weber number

\[ \frac{1}{W_e} = \frac{\rho b_0^{2/3} \lambda^{1/3}}{12\pi}. \] (14)

The critical Weber number is given as 2.2 for this model (Martinez-Bazan et al., 1999). The Weber number will decrease also with a factor \((1 - \phi)\).

Grau et al. (2005) measured the surface tensions at very low concentrations in equilibrium conditions showing the expected larger decrease in surface tension of “stronger” frothers DF250 and DF1012 and a much lower decrease with DF200 (Fig. 11 of Grau et al.). However high-speed photos taken from outside of a 300 mm impeller indicate that the bubble size at the rotor is much finer with the DF200 (Fig. 1).

Jávor (2014) showed the need to use dynamic surface tension values in evaluating the correction factor in Eq. (13) instead of static values (Fig. 2). Of the different frothers tested DF200 decreased the critical bubble size with about 10%, while the “stronger” frothers NF240 and DF 250 had only a marginal 2–3% decrease at very short bubble lifetimes. What is to be noted is the temporal change in NF 240 and 250 frothers. As bubbles age, the critical bubble size starts to decrease. As an outcome in a flotation cell is that after the imminent breakage of the bubbles close to the impeller, the probability of further bubble breakage diminishes rapidly. This indicates that bubble breakage kernels of the source term (Eq. (5)) need to be used only in cases, where the critical bubble size is above the limit in Eq. (13). The results of Jávor (2014) indicate that the assumption for the velocity field to be independent from the internal coordinates (Eq. (5)) may not hold. There are also indications that in a turbulent field the inertia of attached bubbles creates additional breakage mechanisms (Omelka et al., 2009).

3. Bubble coalescence kernels

In pure water gas bubbles have a tendency to coalesce as has been observed in flotation systems by Finch et al. (2008). Bubble coalescence takes place in three steps (Prince and Blanch, 1990a). Bubbles need to collide (e.g. come close enough for liquid to get trapped between the bubbles). In the second phase the liquid drains away until it reaches a critical thickness. As that thickness is reached the van der Waals forces cause the film to break. There is a large body of literature discussing bubble coalescence (Marrucci and Nicodemo, 1967; Sager et al., 1976; Derjaguin and Churaev, 1978; Prince and Blanch, 1990b; Marrucci, 1969; Craig et al., 1993a, 1993b; Christenson and Yaminsky, 1995; Deschenes et al., 1998; Marcelja, 2000; Henry et al., 2007; Christenson et al., 2008; del Castillo et al., 2011 and others). In a simplified model we can multiply the collision rate with a success rate (Eq. (15)) in order to get a coalescence birth kernel for CFD modelling.

The simple equation for coalescence success rate Coulaloglou (1975) suggested is

\[ P_c(b, h) = \exp \left( -\frac{t_{\text{drainage}}}{t_{\text{contact}}} \right) \] (15)

It relates the time the bubbles spend at a distance where the drainage of the film takes place \(t_{\text{contact}}\) to the time it takes for the film drain to a thickness where van der Waals forces rupture the thinned film \(t_{\text{drainage}}\). Machon et al. (1997) added a boundary condition

\[ P_c = 0, \quad \text{if} \quad t_{\text{drainage}} < t_{\text{contact}}. \] (15b)

The drainage is governed by the following equations (Chesters and Hofman, 1982; Carnie et al., 2005), where \(p\) is hydrodynamic pressure, \(\Pi\) disjoining pressure, \(\lambda\) surface tension and \(\mu\) dynamic viscosity (see notations in Fig. 3).

\[ \Delta p = (p + \Pi) - \frac{\lambda}{r} \left( \frac{\partial h}{\partial r} \right). \] (16)
If the bubble surface is immobile then \( u(z,r)z=h = 0 \) and the result is a parabolic Poiseuille flow

\[
\frac{\partial h}{\partial t} = \frac{1}{r} \frac{\partial}{\partial r} \left[ r \int_{0}^{h(r,t)} u(z,t)dz \right].
\]

\[
\frac{\partial u}{\partial z} = \frac{\partial p}{\partial r}.
\]

For a fully mobile surface \( u(z,r)z=h \neq 0 \) leads to a boundary condition of \( \frac{\partial u}{\partial z} = 0 \). The final outcome is a plug flow profile with a four times faster rate of bubble distance \( (h) \) diminishing.

\[
\frac{\partial h}{\partial t} = \frac{1}{3\mu r} \frac{\partial}{\partial r} \left( h^3 \frac{\partial p}{\partial r} \right).
\]

The real situation is more complex due to surfactant concentration gradients and resulting Marangoni stresses. However, such details are not rate determining in flotation CFD modeling (Finch et al., 2008). As a first estimate we can use viscoelasticity to define the flow regime (Fig. 4). The surface active frothers DF250 and PPG425 show marked viscoelasticity, while DF200 and 1-Pentanol had viscoelasticity values below the detection limit (dilatation elasticity method was employed for viscoelasticity measurement that has a relatively low reproducibility in test solutions where the measured viscoelasticity does not rise above 1.5 mN/m). For DF250 and PPG425 Eq. (19) should be used and for DF200 and 1-Pentanol Eq. (20) is to be used.

If Eqs. (19) and (20) are taken to represent the different frothers depending on their viscoelastic behavior and their drainage rate we can see in Fig. 5 a very marked difference in the relative coalescence rates computed using the Coulaloglou equation (Eq. (15)).

Horn et al. (2011) have mapped the effects of bubble approach velocity and NaCl concentrations on coalescence (Partially reproduced as Fig. 6). Fig. 6 is intentionally left without axis values; as such data for commercial frothers is not available. The slow viscous drainage represents the immobile frother surfaces (DF250 and PPG 425) and the rapid inertial drainage represents the mobile surfaces (DF200 and 1-Pentanol). The study of Horn et al. (2011) indicate that there exists a critical speed of approach \( V_c \), which has an impact on the relative immobility of the surfaces (Yaminsky et al., 2010).
where $\Delta \gamma$ is the surface tension gradient from the bubble centreline onwards. The equation does not take into consideration the different behaviour of frothers, like their effects on surface viscoelasticity. The equation, however, indicates the existence of a critical bubble approach velocity as also has been considered by Kirkpatrick and Lockett (1974).

As surfactants are adsorbed onto the surface the increased rigidity of the surfaces decreases the velocity difference between the fluid and the bubble substantially. Clift et al. (1978) showed (Fig. 7.3) in compiling the results from several researchers that for one and two agent (Fig. 7b) made the interface rigid within milliseconds reducing any change in rising velocity. Whereas the more surface-active frothers, like their effects on surface viscoelasticity, decrease the velocity difference within the bubbles. Eq. (25) may not capture the physical complexity of bubble-bubble collisions to a degree sufficient for modelling.

We can write the coalescence kernel (Eq. (7.3)) by combining Eqs. (15) and (25) to get

$$\beta(b, b_1) = P_{co}P(b, b_1) = P_{co} \exp(-\alpha \xi(V, b, b_1)),$$

where $\alpha = 1$ for mobile surfaces and $\alpha = 4$ for immobile surfaces, $\xi(V, b, b_1)$ the ratio between drainage time and contact time in a pure system at an approach velocity $V$.

4. Discussion

Computational modelling of flotation has drawn much of its methods and algorithms from the works dealing with bubbly flows (e.g. Lehr and Mewes, 2001; Marchisio and Fox, 2005; McGraw, 1997; Marrucci, 1969 and several others). These models have used information of bubble behaviour from experimental work done in electrolyte systems (e.g. Liao and Lucas, 2010; Yaminsky et al., 2010; Horn et al., 2011). Frothers are different in several aspects from electrolytes. The basic difference is that frothers are adsorbed onto the surface and change the surface “stiffness” as discussed above. The hysteresis between adsorption and desorption in a flow field is one of the most important features (Jávor, 2014) differentiating the frother effects from electrolyte effects.

The film drainage model used in computational modelling determines the coalescence efficiency (and therefore the bubble size distribution) from the characteristic time scales of surface contact and drainage. The former is strongly determined by fluid dynamic while the latter is by bubble surface rigidity (Lee and Hodgson, 1968). By linking the bubble surface mobility to characterisable surface property that can be easily measured in dynamic water-surfactant solution can help to get a step closer to a physical based model. The above presented approach is an attempt of integrating frother properties into the modelling that could bring more realistic bubble size distributions estimates.

5. Conclusions

The direct use of kernel equations derived from pure systems or from studies performed with electrolytes (notably NaCl) lead to wrong estimates of the bubble size at any spatial volume modelled. The characteristics of frothers have not been taken into consideration in the modelling of bubble behaviour. For bubble break–up it is proposed to take the frother effects into account by a correction factor dependent on the dynamic surface tension (Eqs. (12) and (13)). It is argued that
break-up kernel computations are really needed only at volumes with high turbulent energy dissipation.

It has been shown that frothers can be divided into two major categories (for modelling purposes) based on their effects on viscoelasticity. Frothers like DF200 will have a coalescence drainage behaviour best described by a plug flow of liquid, while frothers like DF250 produce a parabolic Poiseuille flow. This difference in behaviour can be simplified to be taken into account in the coalescence kernel only by a constant factor of four (Eq. (26)). It is acknowledged that there exists a critical bubble-bubble approach velocity, but equations derived for pure systems do not capture the phenomena taking place in flotation in such a way that they could be used in developing the kernel models.

As a simplified outcome for flotation CFD modelling, we propose two modelling regimes depending on the properties of the used frother for the time being.

For future improvements of flotation modelling more frother behaviour derived data is needed to “fill” the framework presented as Fig. 6 (Horn et al., 2011; Yaminsky et al., 2010).

References