Abstract

The buoyancy-driven interaction between a free droplet and solid wall is simulated. The presented model makes use of the continuum-surface-force (CSF) framework to couple an interface-capturing technique to a lower-dimensional lubrication reduction of the thin-film momentum equation. Droplet-scale fluid transport was modelled using an axisymmetric level-set method, including a reinitialisation procedure. Through comparison of model results with experimental data from the literature, we demonstrate the effectiveness of this approach in simulating the challenging problem of small-length-scale drainage in computational fluid dynamics (CFD) modelling. Film-thickness profiles were modelled up until the onset of asymmetric film-drainage. Results presented will aid in the future improvement of simulation techniques for droplet interactions in liquid–liquid systems.

Introduction

A detailed understanding of droplet–interface and droplet–droplet interactions is required in the design and optimisation of multi-phase industrial processing equipment. Droplet collisions, however, pose unique experimental and numerical challenges due to the large differences in length scales present.

Experimental studies have focused on (i) droplet–wall [5], (ii) droplet–home-phase [2] and (iii) droplet–droplet [4] collision scenarios. Using a capillary, Hartland [6] generated free droplets of known volume that were allowed to fall under buoyancy and collide with a solid surface. Profiles of the film thickness were obtained via tracing magnified photographs of the collision sequence. The refractive indices of the two fluids were matched to minimise optical distortion. A positive pressure was reasoned to exist near the centre of the film in order to drive viscous film drainage, consistent with the observed “dimpling” of the film-thickness profiles. Similar droplet–wall systems have been studied using high-speed video and interferometric techniques [11, 5].

Modelling studies have applied film-thickness based adaptive mesh refinement and moving-mesh interface-tracking methods [14]. To capture the collision dynamics completely, a discretisation mesh is required to span scales from the droplet diameter (typically of order 1–10 mm) to the thickness of the draining film, which can be as low as 10–100 nm prior to the onset of wetting. Alternative approaches have included the coupling of droplet-scale dynamics, via force balances [11] or experimental imaging [12], to semi-analytical thin-film drainage models [3].

In this study, we demonstrate the efficacy of a multi-scale modelling approach that alleviates the need for computationally taxing mesh refinement. A level-set method is coupled to a concurrent numerical solution of the Reynolds lubrication equations using the finite-volume solver arb [9]; the coupling is achieved via incorporating the film pressure into the continuum–surface–force (CSF) framework of Brackbill et al. [1]. A volume-of-fluid based version of the modelling framework has previously been applied to Leidenfrost [10] and binary coalescence [13] dynamics in liquid–gas spray systems. Here, we demonstrate its applicability to droplet–wall collisions in an emulsion system. Direct comparisons are made to the buoyancy-driven collision experiments of Hartland [6].

Numerical Method

Droplet-scale Level-set Model

A level-set method [15] is used to capture multiphase fluid motion: the level-set function, $s$, is defined via

$$s(x) = \begin{cases} d(x), & \text{for } x \in \text{disperse phase} \\ 0, & \text{on interface} \\ -d(x), & \text{for } x \in \text{continuous phase} \end{cases}$$

where $d(x)$ is the minimum normal distance from location $x$ to the interface. Level-set advection is undertaken using

$$\frac{\partial s}{\partial t} + \nabla \cdot (s \mathbf{u}) = 0,$$

where $\mathbf{u}$ is the fluid velocity. The incompressible Navier–Stokes equations are solved in conservation form:

$$\frac{\partial (\rho \mathbf{u})}{\partial t} + \nabla \cdot (\rho \mathbf{u} \mathbf{u} + \rho \mathbf{I}) + \nabla \cdot \mathbf{f}_b = \rho g$$

$$\nabla \cdot \mathbf{u} = 0$$

where the viscous stress tensor, $\tau$, is given by

$$\tau = -\mu \left[ \nabla \mathbf{u} + (\nabla \mathbf{u})^T \right],$$

and $\mathbf{f}_b$ is a CSF source term incorporating the interfacial tension and film pressure forces [1, 10, 13]:

$$\mathbf{f}_b = (\gamma \mathbf{c} + \rho_{\text{film}}) \hat{n}$$

Here, $\gamma$, $\mathbf{c}$ and $\rho_{\text{film}}$ are the interfacial tension, local interface curvature and subgrid film pressure, respectively. The volume-averaged volume fraction is then given by $\phi(x) = H(s(x))$, where $H$ is a smoothed Heaviside function. The fluid density, $\rho$, and viscosity, $\mu$, can be related to $\phi$ via

$$\rho = \phi (\rho_d - \rho_c) + \rho_c$$

$$\mu = \phi (\mu_d - \mu_c) + \mu_c$$

where subscripts “d” and “c” refer to the disperse and continuous phase. Level-set reinitialisation is applied for locations separated further than $\epsilon$ from the interface (figure 1).

Film-scale Lubrication Model

The lubrication equations [10]

$$\frac{\partial \rho_{\text{film}}}{\partial z} = 0$$

Subgrid CFD Film-drainage Modelling: Application to Buoyancy-driven Droplet–wall Collisions in Emulsions

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and
\[
\frac{\partial^2 \nu_r}{\partial z^2} = \frac{1}{\mu_c} \frac{\partial p_{\text{film}}}{\partial r}
\]
(10)
are solved on a one-dimensional grid (figure 1a). The radial film
velocity, \(v_r\), is subject to a no-slip boundary condition on the
solid wall,
\[
(v_r)_{z=0} = 0,
\]
and a matched-velocity constraint on the lower droplet interface,
\[
(v_i)_{z=h} = (u_i)_{z=h},
\]
(12)
where \(h(r)\) is the local film thickness. The local film volumetric
flow rate, \(Q\), is determined via
\[
Q = 2\pi rh\bar{v}_r,
\]
(13)
where \(\bar{v}_r\) is the \(z\)-averaged local radial velocity from the solution
to equation (10). A volume-conservation equation can then be
deﬁned on all cell faces,
\[
A_i \frac{\partial h}{\partial t} = Q_{\text{in}} - Q_{\text{out}},
\]
(14)
where \(A_i\) is the annular cell area and \(Q_{\text{in}}\) and \(Q_{\text{out}}\) refer to the
volumetric flow rates into and out of a cell, respectively, via its
tables (figure 1a). We interpolate cell-centred film variables \((h, p_{\text{film}})\)
to the faces using an integral interpolation method based on an analytical solution to equation (10). Reflection
boundary conditions are applied on the \(r = 0\) centreline, and the
film pressure is set to \(p_{\text{film}} = 0\) on the outer boundary.

The droplet-scale motion equations, equations (3) and (4), are solved together with the volume-conservation film equation,
equation (14), using the finite-volume solver \(arb\) [9]. Parameter values are given in table 1.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>0.10 mL</th>
<th>0.25 mL</th>
<th>0.50 mL</th>
</tr>
</thead>
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<tr>
<td>(D) (mm)</td>
<td>5.76</td>
<td>7.82</td>
<td>9.85</td>
</tr>
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<td>(z_0/D)</td>
<td>0.60</td>
<td>0.60</td>
<td>0.70</td>
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<td>(\varepsilon/\Delta r)</td>
<td>38</td>
<td>39</td>
<td>36</td>
</tr>
<tr>
<td>(\mu_d) (kg/m(^3))</td>
<td>1.58 (\times 10^3)</td>
<td>1.58 (\times 10^3)</td>
<td>1.55 (\times 10^3)</td>
</tr>
<tr>
<td>(\rho_c) (kg/m(^3))</td>
<td>1.07 (\times 10^3)</td>
<td>1.07 (\times 10^3)</td>
<td>1.07 (\times 10^3)</td>
</tr>
<tr>
<td>(\mu_d) (Pa s)</td>
<td>17.5</td>
<td>17.5</td>
<td>20.0</td>
</tr>
<tr>
<td>(\mu_c) (Pa s)</td>
<td>13.7</td>
<td>13.7</td>
<td>13.7</td>
</tr>
<tr>
<td>(\gamma) (mN/m)</td>
<td>24.0</td>
<td>24.0</td>
<td>24.2</td>
</tr>
</tbody>
</table>

Table 1. Simulation parameters corresponding to the 0.10, 0.25 and 0.50 mL droplet-volume experimental cases of Hartland [6].

Results and Discussion

Each experimental case of Hartland [6] was investigated, corre-
sponding to droplet volumes, \(V\), of 0.10, 0.25 and 0.50 mL. To highlight the satisfactory agreement between model and experimental results at the small scale of film drainage, time sequences of the film-thickness, \(h(r,t)\), and a characteristic film-pressure profile, \(p_{\text{film}}(r)\) are given. To demonstrate similarly good agreement at the larger droplet scale, a characteristic cross-sectional droplet profile is presented.

Film-scale Validation

Film-thickness time sequences are shown in figure 2. Simu-
lation results agree well with the corresponding experimental profiles until the onset of asymmetric drainage, which cannot be accounted for using the present axisymmetric model. For each

collision case, the earliest profile presented in [6], at nominal
t = 0 s, displays ﬂattening consistent with a droplet–wall inter-
action. As the initial droplet release heights, \(z_0\), are not known
from [6], we use an optimised \(z_0\) such that the film thickness
deforms to the same extent, at the correct distance from the wall,
as the \(t = 0\) s experimental proﬁle. Optimum \(z_0\) results are given
in table 1 and all simulation times have been translated to be
consistent with the quoted experimental times. Simulations of
freely accelerating droplets, with no wall present, indicate that
the deformation present in the \(t = 0\) s experimental proﬁle cannot
be attributed to viscous drag. The film-thickness shapes at ter-
minal velocity are shown in ﬁgure 2, denoted “terminal proﬁle”.
Results from additional simulations (not shown) undertaken using
perturbed, aspherical droplets of equivalent volume indicated
that evolution of the ﬁlm proﬁle is not highly sensitive to the
initial droplet shape, consistent with the numerical results of [8].

Time series for the centreline, \(h_0\), and barrier rim, \(h_t\), ﬁlm thick-
ness are shown in ﬁgure 3 for the 0.1 mL case. The present model
grees well with experimental results for both the centreline and rim thicknesses; note that the model of [8] underestimates the rim thickness at early times in the sequence.

An analysis of the model ﬁlm pressure is given in ﬁgure 4, in
which the simulation pressure is shown at \(t = 120\) s for the
0.25 mL case. Using experimental ﬁlm-thickness proﬁles as inputs, Hartland [7] predict corresponding film pressures via a
numerical method: limits of variation in the derived ﬁlm pres-
sures for all experimental times later than \(t = 90\) s are shown

Figure 1. Sketch of modelling approach: (a) droplet-scale domain mesh
and subgrid model, (b) level-set reinitialisation scheme.
in figure 4. Also shown is the predicted film pressure for an equilibrium film of uniform, zero thickness [7]. The simulated film pressure at $t = 120\text{s}$ was found to be of the same magnitude as the equilibrium pressure, though slightly lower than the predicted pressure bounds of [7]. Note that the simulation film pressure trends smoothly to zero as the film thickness increases: the film pressure of [7] approaches zero at a predetermined film-radius, 3.96 mm in this case.

**Droplet-scale Validation**

Model and experiment droplet-scale cross sections for the 0.25 mL droplet are shown in figure 5. Note that the experimental image (figure 5a) shows the droplet silhouette, though the centre cross-section and inner dimple are still visible inside the droplet, as fluids of matched refractive index were used. The shape of the simulation cross-section (figure 5b) agrees well with that of the experiment, validating model performance at the droplet scale.

**Figure 2.** Film-thickness profiles, $h$, at times corresponding to the experimental data of Hartland [6]. Simulation and experimental results are in good agreement until the onset of asymmetric film drainage. Note that reverse flow was observed in the 0.25 mL experiment, which cannot be captured with the present model implementation.

**Figure 3.** Film thickness at the centreline, $h_0$, and at the rim, $h_r$, for the 0.1 mL droplet case.

**Figure 4.** Simulation film pressure, $p_{\text{film}}$, and results of Hartland’s [7] analysis for the 0.25 mL droplet case.

**Figure 5.** Droplet-scale cross section results for the 0.25 mL case at $t = 60\text{s}$: (a) experiment [6] and (b) simulation. For illustration, the simulation results have been reflected in the rz-plane. Image ©1951–1971 AIChE, reproduced with permission.
Model Limitations

The current model implicitly assumes a fully mobile interface, hence the fluid shear stress and velocity are taken to be continuous across the two-phase boundary. We note that Hartland [6] concluded an immobile, zero-shear-stress interface was present at early experimental times, before transitioning to a mobile interface. Though not investigated in the present study, early-time influence of interface mobility could be explored via modification of the subgrid boundary condition, equation (12). Experimental complications also increased the uncertainty of the immobility finding [6]: reverse flow occurred into the film at early times (see figure 2b), decreasing the net drainage rate, while tilting of the droplet occurred at late times, increasing the drainage rate. Due to axisymmetric constraints, late-time tilting, and associated lateral motion, cannot be accounted for in the present model: the model is primarily applicable to smaller-diameter droplets, for which the tilting effect is less pronounced [6].

As per [6], electrostatic forces have been assumed to be negligible for the film thicknesses studied. For modelling of smaller film thickness with appreciable electrostatic forces, a disjoining-pressure treatment of the electric double layer terms should be included in equation (6).

In applying the CSF model [1], via equation 6, the droplet-scale mesh has been extended below the wall z-location (figure 1a). Viscous continuous phase interaction with the wall is not accurately accounted for, as fluid elements can pass directly through the wall, until reaching the outflow condition on the bottom of the domain. Accounting for these interactions by modifying conventional CFD wall boundary conditions to incorporate film-drainage equations, of a similar form to equation 14, is the focus of current research.

Conclusions

The buoyancy-driven collision between a free droplet and wall was modelled for a highly viscous liquid–liquid system. Droplet-scale dynamics were captured using an implicit level-set method with reinitialisation, while thin-film drainage dynamics were modelled using a coupled subgrid-scale volume-conservation equation.

Model predictions agree favourably with the experimental results of Hartland [6] for all droplet-volume cases considered. Model results were found to be more accurate, when tracking the barrier-rim thickness, than the numerical model of Hartland [8]. Droplet-scale cross-sectional profiles were shown to match those of the original experiment for the 0.25 mL volume case, while the radial pressure profile was found to asymptotically approach the predicted equilibrium profile of Hartland [7].

As per previous models [8], axial symmetry has been assumed meaning that the model cannot reproduce the tilting and non-uniform drainage behaviour that is exhibited at late times in the experimental data. Ongoing research focuses on addressing limitations of the model implementation including (i) extension of lubrication model from a linear to planar system of equations, enabling the simulation of glancing collisions and asymmetric film drainage, and (ii) accounting for wall-effects on the continuous-phase velocity field.

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References