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The influence of surface tension gradients on drop coalescence

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We present the results of a combined experimental and numerical investigation of the coalescence of a drop with a liquid reservoir of a miscible but distinct fluid. Particular attention is given to elucidating the influence on the coalescence process of a surface tension difference between drop and reservoir. Drops are gently deposited on the surface of the reservoir, and so coalesce with negligible initial vertical velocity. Depending on the drop size and reservoir composition, partial or total coalescence may occur. Three distinct regimes, depending on the reservoir to drop surface tension ratio, $R_s$, are identified and delineated through both experiments and numerics. If $R_s<0.42$, droplets are ejected from the top of the drop, while satellite droplets are left in its wake. For $0.42\leq R_s<0.93$, only total coalescence is observed. When $R_s\approx 0.93$, partial coalescence is increasingly favored as the reservoir’s surface tension increases. © 2009 American Institute of Physics. [DOI: 10.1063/1.3177339]

I. INTRODUCTION

The coalescence of a droplet into a reservoir of like fluid has been studied in the context of emulsions,1 raindrop formation2 and, more recently, microfluidic devices.3,4 The mixing that results from drop coalescence has received particular attention owing to its importance in both natural and industrial settings.5,6 The advent of fast cameras and powerful numerical simulations has shed new light on the details of droplet coalescence. In particular, the phenomenon of partial coalescence, first reported almost 50 years ago,7,8 has received considerable recent attention.

When a drop is gently placed on a reservoir of the same fluid, gravity draws the drop downward and eventually the drop and reservoir interfaces become sufficiently close (~100 nm) that attractive Van der Waals forces initiate coalescence. The resulting fluid neck that joins them rapidly opens up; moreover, in the inertia-dominated regime of interest here, capillary waves propagate away from the initial point of contact. Two very different outcomes may follow: Either total coalescence occurs and the drop merges entirely with the reservoir, or only a fraction of the drop liquid merges with the reservoir and a daughter drop is left behind. For partial coalescence to occur, the horizontal constriction of the interface, driven by the surface tension on the sides of the drops, must overcome the vertical collapse driven by the curvature at the top of the drop. When capillary waves are able to reach the drop’s summit before being damped, they interfere with the vertical collapse and allow the horizontal pull of surface tension to prevail and cause pinch-off of the interface.9 The daughter drop left behind by partial coalescence is projected downward, bounces on the reservoir surface, and eventually comes to rest before undergoing a similar process, yielding a coalescence cascade. When the daughter drop becomes sufficiently small, viscous effects come into play and total coalescence occurs.

In the inertially dominated regime relevant to liquids of low viscosity10 determined that the dominant forces at play were surface tension and inertia, and the characteristic time scale of coalescence $\tau=(\rho R^3/\sigma)^{1/2}$, where $R$ is the drop radius, $\rho$ is its density, and $\sigma$ is the surface tension. The critical role of capillary waves in partial coalescence was identified by Chen et al.11 (2006) and criteria for partial coalescence in air-liquid systems were presented by Blanchette and Bigi0n12 in 2006. A mechanism for partial coalescence relying on the presence of surfactants was also presented.12 Subsequent experimental studies have elucidated further details of the coalescence process including the existence of a viscously dominated partial coalescence regime.13–15

We examine here the influence on the coalescence process of a surface tension difference, as may arise from differences in temperature or composition, between drop and reservoir. The influence of Marangoni effects, or equivalently surface tension gradients,16,17 on droplet coalescence has received very little attention. Coalescence in the presence of an initially uniform concentration of surfactants was investigated18,19 with a particular focus on residence times and on the evolution of the thin film separating the drop from the reservoir, but no systematic surface tension gradients were introduced. It was later demonstrated that drop coalescence may be delayed by a temperature gradient between drop and reservoir20,21 since the resulting Marangoni stresses may serve to resist the drainage of the intervening air layer. The early stages of the coalescence of a drop held by a tube and of composition different from that of the underlying reservoir were recently visualised using an extremely fast camera.22 However, the coalescence of a freestanding, initially quiescent drop in the presence of Marangoni stresses has yet to be investigated. In particular, the conditions under which the coalescence is partial or total have yet to be determined. A direct application of inhomogeneous drop coales-
cence was recently suggested as a sorting mechanism for drops on a vertically vibated reservoir.\(^{23}\) Drops larger than a given threshold undergo partial coalescence while smaller drops bounce indefinitely on the interface.\(^{24}\)

In Sec. II, we present the results of our experimental study. The governing equations are developed in Sec. III, and our numerical model described in Sec. IV. In Sec. V, we present the results of our numerical investigation, and in Sec. VI we use them to rationalize the observations reported in Sec. II. Our principal conclusions are presented in Sec. VII along with a discussion of directions for future research.

II. EXPERIMENTS

We examine the coalescence of a droplet brought by gravity into contact with a liquid reservoir of different compositions. To highlight the effects of surface tension variations while minimizing those of density and viscosity, we used water-ethanol mixtures for both the drop and reservoir fluids. This also ensured that coalescence took place in an inertially dominated regime so that the early neck expansion remained stable to the a previously reported instability.\(^{25}\)

A schematic representation of the experimental apparatus is shown in Fig. 1. Drops were generated by manually pressing on a syringe, located 6 mm above the reservoir, and detached when their weight overcame surface forces. The radius, \(R\), of the resulting drops typically ranged between 1.2 and 1.5 ± 0.1 mm, depending on their composition. The coalescence of the initial drop typically lasted approximately 0.01 s and each coalescence event was recorded using a 3000 frames/s high-speed camera (Phantom V7.5). When partial coalescence occurred, up to five generations of daughter droplets were observed but only the first coalescence was examined, as the composition of the daughter droplet and the underlying bath could not be accurately determined thereafter. The container was inclined to avoid difficulties associated with focusing across the glass. To eliminate contamination and temperature gradients, the solutions were replenished after each experiment.

To reduce the number of parameters influencing the coalescence process, we focus on drops that are virtually immobile at the onset of coalescence and for which the reservoir interface is nearly flat. While the former condition is relatively easily met when the composition of the drop and reservoir are identical, we found that drops of different compositions tended to initiate coalescence before coming to rest. After leaving the syringe, such drops often coalesced upon first contact with the reservoir or while rebounding, thus initiating coalescence with a significant downward or upward velocity, respectively. This short rest time may be due to the condensation of ethanol vapor into either the drop or reservoir, thereby creating on the interface an area of lower surface tension. Tangential flow away from the point of contact then tends to drive air out of the gap between drop and reservoir, and reduce the time the drops spend above the interface before coalescence.\(^{26}\) Obtaining reproducible initial conditions was thus proved challenging and limited the scope of our experimental observations. The drop and reservoir fluids were made of either pure water, pure ethanol, or one of five ethanol-water mixtures, as shown in Table I. The viscosity was calculated based on the results of Ref. 27 and the density measured with an Anton–Parr densitometer. The surface tension was measured using the Wilhelmy method with a Kruss K10 tensiometer.

We consider a drop of density \(\rho_1\), viscosity \(\mu_1 = \rho_1 \nu_1\), and surface tension \(\sigma_1\), overlying a reservoir of density \(\rho_2\) and viscosity \(\mu_2\), and most importantly with different surface tensions \(\sigma_2\). The surrounding air has density \(\rho_{\text{gas}}\). The compositional diffusivity of alcohol in the alcohol-water solutions comprising the drop and reservoir fluid is given by \(\kappa\). To focus on the effects of surface tension gradients, we assume in our analysis that \(\mu_1 = \mu_2 = \mu_\ldots\) and \(\rho_1 = \rho_2 = \rho_\ldots\) and find that four dimensionless numbers characterize the system,

\[
\begin{align*}
\text{Bo} &= \frac{(\rho_1 - \rho_{\text{gas}})gR^2}{\sigma_1}, \\
\text{Sc} &= \frac{\nu_1}{\kappa}, \\
\text{Oh} &= \frac{\mu_1}{\sqrt{\rho_1 R \sigma_1}}, \\
R_\sigma &= \frac{\sigma_2}{\sigma_1}.
\end{align*}
\]

The Bond number, \(\text{Bo}\), prescribes the relative magnitudes of the gravitational potential and surface energies of the drop. The Schmidt number, \(\text{Sc}\), prescribes the relative magnitudes of the diffusivities of momentum and composition. The Ohnesorge number, \(\text{Oh}\), prescribes the relative importance of

<table>
<thead>
<tr>
<th>Mix No.</th>
<th>Ethanol fraction (%)</th>
<th>Surface tension (mN/m)</th>
<th>Density (g/cm³)</th>
<th>Viscosity (cP)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>100 ± 0.5</td>
<td>22.0 ± 0.1</td>
<td>0.7851 ± 10⁻⁴</td>
<td>1.10 ± 0.02</td>
</tr>
<tr>
<td>1</td>
<td>60</td>
<td>27.2</td>
<td>0.8895</td>
<td>2.24</td>
</tr>
<tr>
<td>2</td>
<td>55</td>
<td>27.7</td>
<td>0.9010</td>
<td>2.29</td>
</tr>
<tr>
<td>3</td>
<td>50</td>
<td>28.8</td>
<td>0.9125</td>
<td>2.33</td>
</tr>
<tr>
<td>4</td>
<td>40</td>
<td>30.3</td>
<td>0.9342</td>
<td>2.34</td>
</tr>
<tr>
<td>5</td>
<td>30</td>
<td>33.4</td>
<td>0.9535</td>
<td>2.13</td>
</tr>
<tr>
<td>6</td>
<td>0</td>
<td>73.0</td>
<td>1.000</td>
<td>0.893</td>
</tr>
</tbody>
</table>
Ohnesorge numbers, and qualitatively similar to the coalescence was also partial for drops of sufficiently small $R$ in the range of $0.3 \leq R \leq 0.42$, a new type of partial coalescence arose. For $R = 1$, the coalescence was total and in $a$, $d$ a downward moving daughter drop was formed. In $c$ the coalescence was total and in $d$ a small daughter drop was ejected upward.

FOR $R = 1$, we observed a coalescence cascade analogous to that reported in Refs. 7–10, see Fig. 2(b). For $R > 1$, the coalescence was also partial for drops of sufficiently small Ohnesorge numbers, and qualitatively similar to the $R = 1$ case, Fig. 2(a). The resulting daughter drop appeared slightly larger than for $R = 1$ case and itself proceeded to undergo partial coalescence. The entire coalescence cascade showed little variation from the $R = 1$ case.

More complex behavior was observed for $R < 1$. For $R > 0.94$, partial coalescence still developed in a manner similar to the $R = 1$ case. However, for $0.42 < R < 0.94$, total coalescence was always observed, see Fig. 2(c). For $R < 0.42$, a new type of partial coalescence arose [Fig. 2(d)]. The early stages resembled those of the classic partial coalescence, with the lower neck rapidly opening, the formation of capillary waves on the interface and the vertical stretching of the drop. The stretching, however, was seen to be much more pronounced than when $R = 1$. The top of the drop formed a nearly cylindrical column that rose under the influence of the Marangoni stresses, while fluid in the lower regions of the drop drained into the lower reservoir. Eventually, the column of fluid ejected a droplet from its summit. This upward ejection was a robust, repeatable effect. The ejected droplet was approximately 2.5–3.5 times smaller in radius than the mother drop, with smaller drops arising from larger surface tension differences. Unlike daughter drops normally seen in partial coalescence, which are moving downward at the time of pinch-off, the daughter droplet was ejected vertically upward, and its ejection speed increased with surface tension contrast. In the most extreme setup considered [Fig. 2(d)], a water drop on a pure ethanol bath, the droplet rose up to the height of the syringe (6 mm above the reservoir). The ejection also gave rise to smaller satellite droplets that formed between the column of fluid and the ejected droplet. Contrary to daughter droplets formed by the classic partial coalescence of drops surrounded by gas, the ejected droplets, once brought downward by gravity, did not bounce on the reservoir surface, but instead coalesced immediately with the underlying bath.

It is worth emphasizing that such ejection events do not result from converging capillary waves that are known to generate microjets in various circumstances, including bursting bubbles, oscillating satellite drops, drops impacting a solid surface, liquid crater collapse, and the coalescence of drops at large Bond numbers. Rather than converging waves, here the ejection is generated by the converging tangential motion along the interface. This motion entrains a significant quantity of fluid, which leads to the formation of daughter droplets of size comparable to that of the mother drop.

III. GOVERNING EQUATIONS

To model our experiments, we consider a drop of liquid 1 slowly coming into contact with a reservoir of liquid 2 while surrounded by air (Fig. 3). The two liquids are miscible with identical density, $\rho$, and viscosity, $\mu$, but may have different surface tensions. To describe this multiphase system, we first introduce a concentration function, $C_0$, that describes the relative fraction of liquid at a given point, with $C_0 = 0$ in the air and $C_0 = 1$ in either liquid. The position of the interface, $S$, determines the value of $C_0$ at any point. We also introduce a concentration function, $C_1$, describing the
proportion of liquid 1 relative to liquid 2 at a point, with 

\[ C_1 = 0 \text{ in pure liquid 2 and } C_1 = 1 \text{ in pure liquid 1}. \]

We may write expressions for the viscosity \( \mu(C_0) \) and density \( \rho(C_0) \) at every point in the domain. The surface tension, \( \sigma(C_1) \), is only defined on the interface and is therefore independent of \( C_0 \). Following several earlier authors,\textsuperscript{33–35} we model the interface as a forcing term, \( F_\text{s} \), which we add to the momentum equation of the Navier–Stokes equations. Assuming all fluids to be incompressible, the governing equations for our system are

\[ \rho (\vec{u}_i + \vec{u} \cdot \nabla \vec{u}) = -\nabla p + \nabla \cdot \{ \mu [\nabla \vec{u} + (\nabla \vec{u})^T] \} + \rho g + F_\text{s}, \tag{5} \]

\[ \nabla \cdot \vec{u} = 0, \tag{6} \]

\[ S_i = \vec{u}, \tag{7} \]

\[ (C_1)_i + \vec{u} \cdot \nabla C_1 = \nabla \cdot (\kappa \nabla C_1), \tag{8} \]

where \( \vec{u} \) is the velocity field, \( p \) is the pressure, \( f_i \) denotes the time derivative of \( f \), \( \kappa \) is the compositional diffusivity, which generally depends on \( C_1 \), and the interface position is defined by \( S \). Equations (5) and (6) describe the evolution of the momentum and mass of incompressible fluids. Equation (7) states that the interface moves with the flow and that no evaporation or condensation is taking place. Finally, Eq. (8) describes the evolution of the concentration \( C_1 \) under the combined influence of fluid advection and molecular diffusion.

The forcing term \( F_\text{s} \) is a surface delta function that is nonzero only on the interface,\textsuperscript{34–36}

\[ F_\text{s} = \delta(S) \nabla \cdot [(\vec{n} - \vec{n}^T \vec{n}) \sigma] = \delta(S)(\nabla S \sigma + 2 \kappa_\text{m} \vec{n} \sigma), \tag{9} \]

where \( \vec{n} \) is a unit normal to the surface, \( \vec{n} - \vec{n}^T \vec{n} \) is a projection operator, \( \nabla S \) is a gradient operator defined on the interface, \( \kappa_\text{m} = \frac{1}{2} \nabla \cdot \vec{n} \) is the mean curvature of the interface, and \( \delta(S) \) is defined such that

\[ \int \int \int_V \delta(S) f(\vec{x}) dV = \int \int_S f(\vec{x}) dS \]

for any smooth test function \( f(\vec{x}) \). The forcing term therefore captures the normal component of curvature-induced stress, as well as the tangential component resulting from surface tension gradients. Using such a formulation is equivalent to enforcing normal and tangential stress conditions on the interface, but has proven to be much more efficient to implement in simulations.

We nondimensionalize the governing equations using the drop radius, \( R \), as a length scale, \( \tau = (\rho R^2 / \sigma_1)^{1/2} \), as a time scale, with \( \sigma_1 \) the drop surface tension of the drop, to find

\[ \rho' (\vec{u}_i + \vec{u} \cdot \nabla \vec{u}) = -\nabla p + \text{Oh} \ \nabla \cdot \{ \mu' [\nabla \vec{u} + (\nabla \vec{u})^T] \} + \rho' \ \text{Bo} + \delta(S) \nabla \cdot [(\vec{n} - \vec{n}^T \vec{n}) \sigma'], \tag{10} \]

\[ \nabla \cdot \vec{u} = 0, \tag{11} \]

\[ S_i = \vec{u}, \tag{12} \]

\[ (C_1)_i + \vec{u} \cdot \nabla C_1 = (\text{Oh}/\text{Sc}) \nabla \cdot (\kappa' \nabla C_1), \tag{13} \]

where all variables are now dimensionless and \( \mu' = \mu / \mu_1 \),

\[ \rho' = \rho / \rho_1, \ \kappa' = \kappa / \kappa_1, \ \text{and } \sigma' = \sigma / \sigma_1, \]

and the dimensionless numbers \( \text{Bo}, \text{Oh}, \) and \( \text{Sc} \) are as defined in Sec. II. For simplicity, the hydrostatic pressure of the gas was incorporated into the pressure term so that \( p \) now represents a dynamic pressure. Note that Eq. (8) describes the evolution of miscible fluids as well as that of liquids of variable temperature, if \( C_1 \) stands for the liquid temperature and \( \kappa \) for the thermal diffusivity.

IV. NUMERICAL SIMULATIONS

The core of the numerical simulations used here is analogous to those described by Blanchette and Bigioni\textsuperscript{32} (2009). A fixed staggered grid is used to compute the velocity, concentration, stress, and pressure fields. The projection method is used to compute the pressure\textsuperscript{37} and first order corrections are included near the interface to take into account the discontinuity of the pressure across the interface.\textsuperscript{35} The interface is tracked using tracer particles that move with the fluid, and cubic splines are used to interpolate between the markers. We consider an axisymmetric cylindrical domain corresponding to a container closed at the top and bottom with the drop aligned on its axis. The boundary conditions imposed on the system are those of no-slip on, and zero flux through, the solid walls, which are positioned sufficiently far from the drop that they do not affect the coalescence process. Our initial conditions consist of a spherical drop at rest just at the point of contact with a horizontal surface: The two surfaces are initially connected by a neck. We follow the evolution of the drop until either the surface comes within a given threshold value of the symmetry axis, at which point we assume that pinch-off occurs, or total coalescence is achieved. A validation of simulations based on the same numerical scheme, but with constant surface tension, was presented in Ref. 32.

We now describe the treatment of the second liquid and its impact on fluid properties and related computations, which have not been described elsewhere. We use linear relationships between the gaseous and liquid fractions, \( C_0 \) and \( C_1 \), and the density, viscosity, and surface tension. For the density and viscosity we have

\[ \rho' = \rho_\text{gas}(1 - C_0) + C_0 \rho_1 \]

\[ \mu' = \mu_\text{gas}(1 - C_0) + C_0 \mu_1 \]

where subscripts \( l \) and “gas” indicate properties of the pure liquid and of the gas, respectively. In the case of surface tension, only the concentration in the liquid phase is taken into account,

\[ \sigma' = \sigma_\text{gas}(1 - C_1) + C_1 \sigma_1 / R_\text{d}(1 - C_1) + C_1. \]

To keep the number of parameters to a manageable level while highlighting the effect of the surface tension gradients, we assume that the compositional diffusivity, \( \kappa \), is independent of \( C_1 \).

Blanchette and Bigioni\textsuperscript{32} (2009) found that air plays a negligible dynamical role in the system evolution and that
overestimating the air density and viscosity did not affect the coalescence outcome provided they remained significantly less than those of the liquid. Moreover, the density and viscosity discontinuities across the interface render the use of the exact air density and viscosity difficult, as mass and energy conservation can only be achieved through the use of an extremely fine resolution.

To allow for faster computations, we therefore fix the air density and viscosity to be 10% those of the drop. While the concentration of air, $C_0$, is determined solely from the position of the interface, the liquid concentration, $C_1$, is advanced in time according to Eq. (8). The advection term is computed via an upwinding scheme while the diffusive term is obtained via central differences. We also impose the condition $0 \leq C_1 \leq 1$ directly. Zero-flux boundary conditions are imposed on the solid walls.

To calculate the surface forcing term, we first introduce markers that locate the position of the interface. The velocity of the markers is determined by a bilinear interpolation of the velocity of the surrounding nodes. The position of the markers is then advanced in time using a simple Euler method. Cubic spline interpolations provide an estimate of the interface position between markers. The average surface forcing term per grid cell is then estimated by integrating the interface position between markers. The average surface markers is then advanced in time using a simple Euler the velocity of the surrounding nodes. The position of the interface, the liquid concentration, $C$, and the interface, $\tilde{I}$, from the bulk of the drop and reservoir to the interface. Because our configuration is axially symmetric, the surface force term reduces to

$$\left( F_3 \right)_{i,j} = \int_{s_1}^{s_2} \sigma' \cdot \left( \tilde{I} - \hat{n} \cdot \tilde{n} \right) n \cdot \nabla \sigma' ds$$

$$= \int_{s_1}^{s_2} \partial \sigma' \frac{dr}{ds} + \frac{\sigma' \partial y}{r} ds + i \frac{\partial \sigma'}{r} ds$$

where $\sigma'$ is the surface tension variation, $r$ is the distance to the symmetry axis, $y$ is the vertical coordinate, and $s$ is the arc length. $s_1$ and $s_2$ are the intersection points between the curve and the $(i,j)$th grid cell, and $i$ is a unit vector tangent to the curve. The last integral is evaluated numerically by discretizing the interface between markers. Note that $(F_3)_{i,j} = 0$ when the curve does not intersect the $(i,j)$th cell.

V. NUMERICAL RESULTS

A. Validation

In order to focus on the effects of variable surface tension, we kept the Schmidt number constant at $Sc = \nu_1 / \kappa_1 = 50$ to model systems where diffusion of concentration is much slower than that of momentum, such as ethanol-water solutions. Note that using a larger $Sc$ is more computationally demanding and yielded nearly identical results. We also set the Bond number, which characterizes the importance of gravity, to $Bo=0$, and initiate coalescence when the drop is spherical and the interface is flat. These conditions provide a good approximation of our experiments, where the surface tension variations are dominant.

We begin by verifying the conservation of mass and energy at various resolutions [Fig. 4(a)]. While the liquid mass is well conserved even at fairly low resolutions, the conservation of energy requires higher resolutions. In the presence of spatial surface tension variations for miscible liquids of equal density, and neglecting gravitational effects, the energy evolution was shown to obey

$$\frac{d}{dt} (K + SE) = -\Phi + \int_S \frac{D\sigma}{Dt} dS,$$  \hspace{1cm} (14)

where $K$ and $SE$ are the total kinetic and surface energy of the fluids, respectively, $\Phi$ is the rate of energy dissipated through the action of viscosity, and the last term accounts for the transfer from internal to surface energy, as the diffusion of $C_1$, from the bulk of the drop and reservoir to the interface, affects the local surface tension.

![Figure 4](http://phf.aip.org/phf/copyright.jsp)

**FIG. 4.** Time evolution of (a) the liquid mass and (b) the total energy calculated at different resolutions. (c) Evolution of the different energy components enumerated in Eq. (14). System parameters are $R_e=1.17$, $Oh=0.01$, $Bo=0$, and $Sc=50$. 

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larger energy transfers, up to 40% for a ratio of higher resolutions. The conservation of total energy worsens layer under the interface, accurately modeling them requires Because such transfers take place across a thin boundary showed a similar transfer. Larger values of liquid 1 shown in Fig. 4 /H20849. FIG. 5. /H20849 Color online. The evolution of the various energy components is examined in sequence with (a) $R_\sigma=0.92$, (b) $R_\sigma=1$, and (c) $R_\sigma=1.5$. Top row: the concentration of liquid 1 (red) and 2 (blue). Row 2: the vertical velocity (red is upward, blue downwards). Row 3: the radial velocity (red indicates radial flow away from the center, blue toward the center). Row 4: the pressure (red is high and blue is low). Bottom row: the magnitude of the tangential stress (red is high and blue low).

The evolution of the various energy components is shown in Fig. 4(c). Initially, the excess surface energy (relative to that of a flat surface of the liquid of lowest surface tension) accounts for all the available energy. At early times, surface energy is transferred to kinetic energy and dissipated through the action of viscosity at approximately equal rates. Closer to pinch-off, the surface area of the drop and associated surface energy increase slightly, while the kinetic energy decreases. The increase in the concentration of liquid 1 at the interface due to diffusive effects results in a significant transfer of the available energy, up to 15%, from the internal energy of the bulk fluid into surface energy. Simulations performed with various Schmidt numbers, Sc=μ/ρκ, all showed a similar transfer. Larger values of $R_\sigma$ lead to even larger energy transfers, up to 40% for a ratio of $R_\sigma=1.5$. Because such transfers take place across a thin boundary layer under the interface, accurately modeling them requires higher resolutions. The conservation of total energy worsens with increasing $R_\sigma$, but it remained better than 90% for surface tension ratios as large as $R_\sigma=1.5$ at the resolution used throughout our study, N=1024. In the regime 0.66≤$R_\sigma$≤1.5, our simulations are therefore sufficiently accurate to provide a reliable description of the drop coalescence.

We show in Fig. 5 snapshots of the concentration, velocities, pressure, and tangential stress fields for different values of the surface tension ratio. Although all the plots correspond to the same nondimensional time, the evolution of the coalescence is more advanced for drops of larger $R_\sigma$. In all cases, the stresses are confined to a thin region near the interface. The most important difference to note between the different values of $R_\sigma$ is the tangential motion present along the interface, inward in sequence (a) where $R_\sigma<1$ and outward in sequence (c) where $R_\sigma>1$. We note that the region of high pressure present near the tip of the drop is largest in the case $R_\sigma<1$, which is likely to accelerate the downward motion of the drop fluid, an observation we investigate further in Sec. V B.

B. Coalescence process

The evolution of the coalescence process is shown in Figs. 6–9 for $R_\sigma=\sigma_2/\sigma_1$ equal to 0.92, 0.967, 1, and 1.5, respectively. In the online version, the drop and reservoir liquids and the outside gaseous phase are initially maroon, blue, and green, respectively, while they are pale gray, dark

FIG. 6. (Color online) Simulation of the coalescence of a drop of larger surface tension than that of the reservoir, $R_\sigma=\sigma_2/\sigma_1=0.92$. Each image shown is 0.04 nondimensional time units apart. Despite the low Ohnesorge number, Oh=0.001, the surface tension variations prevent partial coalescence.

FIG. 7. (Color online) Simulation of the coalescence of a drop with slightly larger surface tension than the reservoir, $R_\sigma=\sigma_2/\sigma_1=0.9667$. Each image shown is 0.04 nondimensional time units apart and Oh=0.001. Partial coalescence is observed, although the drop is smaller than it would be in the absence of the surface tension mismatch.
gray, and white in the print version, with regions of mixed liquids appearing as white. The Ohnesorge number was chosen sufficiently small, Oh=0.001, to ensure that viscous dissipation is negligible. The radius of the computational domain was chosen to be sufficiently large, 6R, to ensure that boundaries played no role in the coalescence process; in particular, it was much larger than the domain shown in Figs. 6–9.

We compare the coalescence of liquids of identical surface tension to that of liquid of uneven surface tensions. Whenever \( R_a \neq 1 \), the region of higher surface tension pulls on the interface and generates a tangential flow that partially coats the surface of the liquid of largest surface tension with the second fluid. Zooming in on the interface at the point where the drop and reservoir liquids meet clearly shows the tangential motion (Fig. 10), and the tiny eddy it generates just below the interface. To provide a local measure of the Reynolds number resulting from such tangential flows, \( Re_i \), we estimate the thickness of the fluid layer entrained by the tangential surface flow as \( \delta \sim \sqrt{\mu_i \tau / \rho} \), with \( \tau = (\rho R^2 / \sigma)^{1/2} \) the characteristic coalescence time scale, and the tangential velocity by balancing characteristic tangential Marangoni and viscous stresses to find \( U \sim \Delta \rho / \mu \). We thus estimate \( Re_i = U \delta \rho / \mu i = Oh^{-1} \Delta \sigma / \sigma \), which in the cases shown in Fig. 10 yields \( Re_i \sim 50 \). Even over such a small length scale, tangential motion may thus generate vigorous vortices.

As can be seen by comparing the final images of Figs. 7–9, the coalescence process is much more sensitive to the surface tension ratio for \( R_a < 1 \) than for \( R_a > 1 \). For drops such that \( R_a > 1 \), the tangential motion causes the drop fluid to spread over the reservoir, a process which does not directly influence the competition between pinch-off and vertical collapse. Coalescence then progresses in a manner very similar to when \( R_a = 1 \). However, when \( R_a < 1 \), the tangential motion along the interface brings fluid of lower surface tension up the sides of the drop. The interface driving the horizontal constriction therefore has a diminished surface tension. The horizontal constriction is then slowed (Fig. 7), and the vertical collapse dominates, expelling fluid downward. If \( R_a \) is sufficiently close to 1, the sides of the drop are eventually covered with the larger surface tension fluid, and pinch-off may then occur. The resulting partial coalescence thus occurs at a later time and the resulting daughter drop is smaller and made up nearly entirely of the same fluid as the mother drop. A drop with \( R_a < 1 \) also causes the vortex ring created as fluid is expelled into the bath to detach from the interface. Vortex detachment can also occur in homogeneous coalescence but our simulations show that it requires a significantly smaller Ohnesorge number.

We also performed simulations with Schmidt numbers \( Sc = \nu / \kappa \) ranging from 1 to 50, which encompass the regime where temperature variations cause surface tension variations, the Schmidt number then playing the role of the Prandtl number. The observed coalescence showed very little dependence on \( Sc \) and the main features described above were reproduced. We also attempted to simulate coalescence in conditions resembling those of the last sequences of pictures shown in Fig. 2, with \( R_a < 0.5 \). In this regime, we found that accurately representing the dynamics of the thin layer of outer fluid required far higher resolutions, and we could not investigate numerically the regime where drop ejections were observed in our experiments.

**VI. DISCUSSION**

The influence of surface tension variations on drop coalescence is summarized in Fig. 11. Partial coalescence in the ordinary sense is only possible when \( R_a > 0.93 \). In that regime, we find a critical value of Oh below which partial coalescence occurs [Fig. 11(a)]. To determine numerically the value of Oh, we fix all other parameters and systematically vary the Ohnesorge number, using a bisection method to accurately determine Oh. In regions where the slope of Oh as a function of the surface tension becomes large, we reverse the procedure by fixing Oh and varying the surface tension equal to that of the reservoir, \( R_a = \sigma_l / \sigma_i = 1.5 \). Each image shown is 0.04 nondimensional time units apart and Oh=0.001. Tangential motion along the interface is away from the drop, which prevents the detachment of the vortex ring.
tension ratio. When the surface tension of the reservoir is less than that of the drop, partial coalescence is hindered and \( \text{Oh} \) quickly goes to zero. However, using a drop of weaker surface tension than the reservoir only slightly increases \( \text{Oh} \). Our simulations have not yielded any partial coalescence for \( \text{Oh} > 0.04 \), regardless of the choice of \( R \).

For \( R < 1 \), quite surprisingly, \( \text{Oh} \) increases nonmonotonically with \( R \). Our observations suggest that viscous effects act to both damp capillary waves and slow the spread of the low surface tension fluid at the interface. At very low values of Oh, the hindering of tangential flow by viscous effects is dominant, and increasing Oh therefore facilitates partial coalescence. However, for \( \text{Oh} > 0.02 \), the damping of capillary waves by viscous effects becomes dominant, and so favors total coalescence.

The daughter drop size and velocity are strongly affected when \( R < 1 \). Figure 11(b) shows the daughter drop radius and velocity at a fixed Ohnesorge number \( \text{Oh}=0.001 \) for given surface tension ratios. We first note that the daughter drop radii are only weakly affected by surface tension variations. The smallest drops are generated when \( R = 0.93 \) and have radii as small as 0.47. Figure 11(b) shows the droplet fluid’s average downward velocity at the moment of pinch-off. The time of pinch-off, not shown, follows a curve similar to that of the daughter drop velocity, but with far less variation, with pinch-off times of 1.94, 1.63, and 1.71 at surface tension ratios of \( R = 0.93, 1 \), and 1.5, respectively (see Figs. 7–9). Longer pinch-off times allow the daughter droplets to gather more downward momentum before being released to bounce on the reservoir surface, hence the similarity of the pinch-off time and velocity curves. We note that drops of larger surface tension than the reservoir can reach velocities more than twice that of a drop of the same composition as the reservoir, and thus potentially bounce up to four times higher.

A striking feature of partial coalescence is the coalescence cascade that occurs as the daughter droplets become progressively smaller. We proceed by discussing how this cascade is affected by a compositional difference between drop and bath. We observed coalescence cascades for drops having surface tension smaller than that of the reservoir (\( R > 1 \)). Figure 9 illustrates that typically, only the first coalescence event of the cascade would be influenced by the surface tension mismatch. Unless the daughter drop drifted sideways significantly (which does occasionally happen), the second coalescence event would be between a drop and reservoir of local composition identical to that of the drop. As the first daughter drop size is only weakly affected by the surface tension gradient, earlier descriptions of partial coalescence are therefore also relevant when \( R > 1 \).

Daughter drops issued from the partial coalescence of drops with \( R < 1 \) appear to have a composition identical to that of the mother drop. However, surface tension gradients favor the detachment from the interface of the vortex ring formed during coalescence; thus, any daughter drop would encounter a reservoir with a surface tension nearly equal to that of the original reservoir (see Fig. 7). The lower critical Ohnesorge number of coalescence associated with \( R < 1 \), combined with the progressively smaller size of the daughter droplets, thus restricts the coalescence cascade to a very narrow parameter regime.

The presence of surface active reagents (surfactants) on the interface also alters surface tension, and varying concentrations generally lead to surface tension gradients. Because such molecules, when insoluble, are surface bound, expansion or contraction of the interface affects their concentration, thus contributing an effective elasticity to the interface, an aspect not considered here. We note that the most important contraction of the interface occurs in the early stages of coalescence, at the opening of the neck. If surfactants are present, such a contraction will increase their concentration, and so typically reduce surface tension. In light of the current results, we anticipate that surfactant-laden interfaces would therefore tend to favor total coalescence, as the horizontal pull of surface tension would be weakened by the larger concentration of surfactants accumulating around the neck of the drop. As a caveat, we note that such a prediction remains a conjecture.
VII. CONCLUSION

We have presented the results of a combined experimental and numerical investigation of drop coalescence into a reservoir in the presence of Marangoni stresses. Vigorous tangential motions generate eddies lying tangent to the interface. Drops of smaller surface tension than the reservoir are only weakly affected by the tangential motion due to the surface tension mismatch. Conversely, the conditions under which drops of larger surface tension undergo partial coalescence are strongly restricted by the intrusive surface motion of the reservoir fluid. The surface tension along the sides of the drop is then diminished, and the total coalescence driven by the curvature at the top of the drop is favored. When partial coalescence did occur ($R_c > 0.93$), the size of the resulting daughter drops increased with increasing $R_c$. The converging surface flow arising for $R_c < 1$ favored the detachment from the surface of the vortex ring formed during coalescence, while the diverging flow arising for $R_c > 1$ hindered its detachment.

We also discovered a different type of droplet ejection prompted by the convergence of tangential surface motion driven by Marangoni stresses. This type of ejection differs from the typical partial coalescence in that it generates significantly smaller drops, with approximately a tenth radius of the mother drop. Our numerical methods were insufficiently refined to replicate such ejections, but we are hopeful that improved methods will soon be able to accurately model the thin boundary layer then present near the interface, and thus to capture drop ejection resulting from converging surface flows.

Our results provide a description of drop coalescence when drops and reservoirs are not necessarily identical, isothermal fluids. In particular, we may now explain the observation of Ref. 9 that partial coalescence was more difficult to observe in a repeatable manner using water as the reservoir and drop liquid. Water left standing quickly becomes contaminated with dust particles that decrease its surface tension. In contrast, a newly formed drop is likely to have a clean surface with a relatively high surface tension. The effective surface tension ratio is therefore likely to satisfy $R_c < 1$, which restricts the conditions under which partial coalescence may arise. While the same effects are likely to come into play for fluids of lower surface tension such as oils, they will be less significant owing to the relative insensitivity of such fluids to surface contamination.

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