Coalescence preference and droplet size inequality during fluid phase segregation

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Abstract – Using molecular dynamics simulations and scaling arguments, we investigate the coalescence preference dynamics of liquid droplets in a phase-segregating off-critical, single-component fluid. It is observed that the preferential distance of the product drop from its larger parent, during a coalescence event, gets smaller for large parent size inequality. The relative coalescence position exhibits a power-law dependence on the parent size ratio with an exponent $q \approx 3.1$. This value of $q$ is in strong contrast with earlier reports 2.1 and 5.1 in the literature. The dissimilarity is explained by considering the underlying coalescence mechanisms.

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Introduction. – When two liquid droplets come in contact with each other they form a liquid bridge and the composite structure finally relaxes to a single big drop—a kinetic process known as coalescence. In recent years this phenomenon has gained significant research attention for a wide variety of natural systems including collision and coalescence of droplets on a solid surface [1–5], coalescence of water drops in water [6], collision of rain drops [7], magneto coalescence of ferrofluidic drops [8], etc. Most of the studies on droplet coalescence sought to understand the growth of liquid bridge, and the effects of contact angle. On the other hand, an intriguing yet rarely explored feature of coalescence processes is the so-called coalescence preference: the product drop (bubble) which emerges from the coalescence of two different-sized parent droplets (bubbles) tends to be placed closer to its larger parent. Such a preferential positioning is caused by the Laplace pressure [9] difference between the parents. A smaller parent has higher Laplace pressure compared to the larger one and hence the merged product is formed closer to its larger parent. While this general trait is conceivable, understanding of the spatial and temporal properties of the coalescence preference effect is still in its infancy and very recent [10,11]. Specifically, the questions of “how close” to the larger parent the merged drop forms and its dependence on the parent size ratio are unsettled issues [10] and call for future studies. The microscopic mechanism of the coalescence preference also remains poorly understood.

In this paper, using extensive molecular dynamics (MD) simulations we investigate the droplet coalescence preference in phase-segregating fluids which are rendered thermodynamically unstable via a sudden change of temperature. In recent years, droplet coalescence and growth in phase-segregating [12,13] fluids has gained huge momentum because of the underlying rich physics originating from the combined effects of hydrodynamics and diffusion field [14–16], and their temperature and dimensionality dependence [17]. However, the coalescence preference (CP) dynamics in such systems has never been investigated at all. On the other hand, fluids coarsening [18,19] via droplet/bubble coalescence are very common in nature. To the best of our knowledge, this is the first study of coalescence preference effects in a system close to phase transition. Apart from having fundamental importance such studies also hold particular relevance in industrial applications. For example, the stability of emulsions/foams which is a crucial factor for pharmaceutical and petroleum industries is controlled by droplet/bubble coalescence [20]. For an improved prediction and control of emulsion stability it is necessary to understand the geometric preference and the microscopic mechanism of rearrangement during coalescence [21].
Typically, the proximity of the merged drop to its larger parent is dictated by the relative coalescence position \(a_L/a_S\) (see fig. 1), where \(a_L\) and \(a_S\) are the distances of the centres of the larger and smaller parent, respectively, to the projected centre of the coalesced drop on the line joining the two parents. As evident from recent experiments [10,11], the location of the product drop depends on the parent size ratio \(R_L/R_S\) in a power-law fashion

\[
\left(\frac{a_L}{a_S}\right) \sim \left(\frac{R_L}{R_S}\right)^{-q},
\]

\(R_L\) and \(R_S\) being the radii of the larger and the smaller parent, respectively. The CP exponent \(q\) determines the closeness of the merged drop to the larger parent. For a fixed parent size ratio, a higher \(q\) means the product forms much closer to the larger parent.

As will be demonstrated, our results on phase-segregating fluids reveal that with increasing size ratio of two coalescing droplets the product drop forms much closer to the larger parent. Although this general trend is similar to earlier reports [10,11] on droplets/bubbles, the “extent” of closeness of the merged drop to the bigger parent and the underlying mechanism of coalescence preference, as observed in our study, are different from previous reports. In particular, the observed exponent \(q \approx 3.1\) is in strong contrast with earlier findings \(q \approx 2.1\) and 5.1. This difference has been attributed to the underlying coalescence mechanism. For phase-segregating fluids where the coalescing drops undergo inelastic sticky collisions [22] the CP dynamics is governed by the formation of the product drop at the centre-of-mass location of the parents. This leads to a value \(q \approx 3.0\). Whereas, higher values of \(q\) in [11] were explained to be due to the coalescence motion controlled by the release of surface energy.

We first explain the CP dynamics here. Figure 1 illustrates schematically one coalescence event during fluid phase separation, where the centres of the smaller, larger parent and the product drop are marked by S, L, P, respectively. In order to describe the relative position of the product drop we define \(a_L\) (\(a_S\)) as the distance between the larger (smaller) parent centre and the point on the line linking the two parent centres closest to the product centre. Using geometric considerations [10] the relative position \(a_L/a_S\) can be directly measured from the locations of the drop centres as

\[
\left(\frac{a_L}{a_S}\right) = \frac{(a^2 - b^2 + c^2)}{(a^2 + b^2 - c^2)},
\]

where \(a, b, c\) are defined in fig. 1. Equation (2) involves only the pairwise distance among the three droplets and, therefore, the knowledge on the droplet positions suffices for the calculation of \(a_L/a_S\). From earlier experiments [10,11], \(a_L/a_S\) is known to exhibit an algebraic dependence on the parent size ratio as dictated by eq. (1). Note that for equal-size droplets the product always forms exactly at the centre of the two parents, i.e., \(a_L/a_S = 1\). The preferential positioning is observed only when the parent droplets are of unequal size. It should be noted here that the universality of the exponent \(q\) is till date questionable. For densely packed microbubbles \(q = 2.06 \pm 0.33\) [10]. For free bubbles \(q = 5.09 \pm 0.43\) and for free droplets \(q = 4.33 \pm 0.54\) [11]. The discrepancy between these high and low density values has been attributed to the blocking effect due to neighboring bubbles.

As will be discussed later, our molecular dynamics results on \(q\) for phase-separating fluids is markedly different from the above-mentioned findings. This is explained within the framework of centre-of-mass theory.

**Model and simulation.** – For a single-component fluid a monodisperse model is considered whose particles interact via the Lennard-Jones (LJ) pair potential \(u(r) = 4\varepsilon\left[(\sigma/r)^{12} - (\sigma/r)^{6}\right]\), where \(r\) is the scalar distance between two particles and \(\varepsilon\) is the interaction strength. \(u\) is smoothly truncated [14,23] at a cutoff distance \(r_c\) and modified as \(u_1(r) = u(r) - u(r_c) - (r - r_c)du/dr\bigg|_{r=r_c}\). This model exhibits a vapor-liquid transition at a critical temperature \(k_BT_c \approx 0.935\varepsilon\) and a critical density \(\rho_c \approx 0.316\sigma^3\) [24]. The non-equilibrium coarsening dynamics within this model is simulated using MD in the canonical ensemble using the Nose-Hoover thermostat [25]. An initial configuration is prepared at a high temperature \(5T_c\) such that it corresponds to the homogeneous phase and at time \(t = 0\) it is quenched inside the binodal to a temperature \(T = 0.6 T_c\). The subsequent dynamics which involves the formation of droplets and their coalescence is monitored at a time interval of \(10\tau\); \(\tau = \sqrt{m\sigma^2/\varepsilon}\) is the LJ time unit. Periodic boundary conditions [26] are applied along all Cartesian directions. The reduced temperature \(T^*\) and...
system size $L^*$ are defined as $T^* = k_B T/\epsilon$, $L^* = L/\sigma$, $\varrho = N/V$ is the overall density of the fluid where $N$ is the total number of particles and $V = (L a)^3$ is the volume of the cubic simulation box. Unless otherwise mentioned, data correspond to averaging over 6 independent configurations.

**Droplet identification.** – One major hurdle in quantifying the CP dynamics is to track the coalescing droplets accurately. This is done by using the following procedure [23]: i) The local density around each particle is calculated and if this density is higher than a critical value the particle is marked as an element of any of the droplets. ii) Next, depending upon the spatial distance between these marked particles, different droplets are identified. This method provides complete information about the total number of droplets in the system, the radius and volume of each droplet. The centre of mass and average radius of each droplet can next be calculated by assuming their spherical shape. From this one can easily measure $a_L/a_S$ and $R_L/R_S$ for a pair of coalescing droplets.

**Results.** – In fig. 2, we show an exemplary coalescence event between two droplets of different sizes produced by using MD simulations. As the droplets collide, a thin liquid bridge forms. With time the liquid bridge grows and the composite structure relaxes until at $t \approx 13250\tau$ which with time increases in thickness and the elongated composite drop relaxes to form a spherical product drop. Results correspond to $L^* = 100$ and $T^* = 0.67 T^\epsilon$.

Binder-Stauffer’s mechanism of Brownian coagulation and coalescence [22,27]. First of all, with increasing size inequality $R_L/R_S$, the product drop moves much closer towards the bigger parent, i.e., $a_L/a_S$ decreases. This is because increasing size inequality increases the Laplace pressure difference between the parents and hence the merged drop is formed much closer to the larger parent. For equal-size droplets ($R_L/R_S = 1$) the product always forms exactly at the centre of the two parents, i.e., $a_L/a_S = 1$. In the limiting case of $R_L/R_S \rightarrow \infty$ the merged drop should form over the infinite parent, i.e., $a_L/a_S \rightarrow 0$. The dashed lines in fig. 3 stand for the previously reported values of $q = 2.1$ and 5.1 in the literature [10,11]. Clearly, our computational data in fig. 3 for various $L^*$, $\varrho$, and $T^*$ (marked by different symbols) do not follow these early findings. The dotted lines mark the limiting values 2.8 and 3.4 of $q$ which the data corroborate. This yields $q \approx 3.1 \pm 0.3$. No prominent system size dependence is observed in fig. 3.

Of course a broader range of abscissa, in fig. 3, will lead to more accurate quantification. However, it should be noted that at early times during non-equilibrium phase separation the size dispersion of droplets is very low which results in lower values of $R_L/R_S$. At very late times the size dispersion increases. However, for the densities and system sizes chosen in this work the droplet size distribution is not broad enough to give rise to values of $R_L/R_S$ much larger than 2.

Note that from the locations of the centres of the coalescing and the merged droplets one can also calculate the distance $d$ between the center of the droplet appearing due to a coalescence event and the line of centers of two parental droplets: $d^2 = c^2 - a_L^2$; where $a_L = b(a^2 + c^2 - b^2)/(2ac)$ and thus study the time dependence of $d$. This we leave out as future exercise.

In fig. 3, the obtained value of $q$ is different from the previous reports [10] $q = 2.1$ and 5.1 in the literature.
We attribute this difference to the dominant microscopic mechanism of the coalescence event. It was already demonstrated [11] that \( q = 5 \) arises from the surface energy release during a coalescence event. Here, we propose that the CP dynamics in our work is governed mainly by the \textit{centre-of-mass theory}. Assuming that the product drop forms at the centre of mass (c.m.) of the two coalescing parents, one obtains: 

\[
    m_p \mathbf{r}_p = m_S \mathbf{r}_S + m_L \mathbf{r}_L,
\]

where \( \mathbf{r}_S, \mathbf{r}_L, \mathbf{r}_P \) are the position vectors of the c.m. of the smaller, larger parents and the product, respectively, and \( m_S, m_L, m_P \) stand for their masses. Simple vector algebra [29] and geometric arguments lead to 

\[
    (a_L/a_S) \sim (m_L/m_S)^{-1}.
\]

Now, assuming that each droplet has constant spatial density within it and using the relation: 

\[
    m = 4\pi R^3/3,
\]

one obtains the coalescence preference relation 

\[
    (a_L/a_S) \sim (R_L/R_S)^{-3}.
\]

Our MD simulation result \( q \approx 3.1 \), in fig. 3, is in excellent agreement with this scaling argument.

To further verify the applicability of the c.m. theory for CP dynamics in phase-segregating fluids, we investigate various kinetic properties of the coalescence events. In fig. 4(a), we test the law of mass conservation, \( V_P = V_S + V_L \), for the coalescing drops considered in fig. 3. Constant density approximation provides the volume of a droplet \( V \) to be proportional to the total number of particles \( N \) in the droplet. The mass conservation law thus leads to 

\[
    N_P/N_S = 1 + N_L/N_S.
\]

Data in fig. 4(a) nicely corroborates this conservation law. This observation is also supported by the predictions of inelastic “sticky” collisions between coarsening droplets within the framework of Brownian coagulation theory [22].

Next, in fig. 4(b), we show the time evolution of the \( x \)-coordinate, marked by filled symbols, of the centres of the two coalescing droplets considered in fig. 2. The larger and smaller parents are marked by \( L \) and \( S \), respectively. Before \( t \approx 13400 \), the centers of the individual parent droplets approach each other during coalescence and shape relaxation of the composite object takes place. Finally, beyond \( t \approx 13400 \), the shape of the composite structure does not change much and the spherical final product droplet has almost formed. As a result, after \( t = 13400 \) locations of the two coalescing droplets coincide with each other. It should be noted that the smaller parent \( S \) shifts more towards the bigger one \( L \) and the final product is formed closer to the larger parent, as expected for the coalescence preference phenomenon. The \( x \)-coordinate of the c.m. of the parent droplets was calculated, at time \( t \approx 13300 \) when the \( x \)-ordinates of the droplets just touch, as 

\[
    x_{\text{cm}} = (x_1 R_1^3 + x_2 R_2^3)/(R_1^3 + R_2^3)
\]

and was calculated to be \( \approx 52.76 \) (marked by an arrow). The final product indeed forms very close to this location.

Figure 4(a) and fig. 4(b), therefore, convincingly demonstrate that the merged droplet during droplet coalescence, via the Brownian motion and coagulation mechanism [22] in a phase-separating single-component fluid, forms at the centre-of-mass location of the parents, and it leads to a coalescence preference exponent \( q \approx 3 \). This is in strong contradiction with previous studies [10,11] where the product drop does not form at the c.m. location.

Droplet phase separation may involve more complex coalescence events as well. One such example being the coalescence-induced coalescence [30]. A droplet undergoing coalescence changes its shape with time and during this shape relaxation process it may touch another neighboring droplet due to geometrical reasons. This will in turn lead to further coalescence events. This is particularly relevant when the average droplet density in the fluid is very high [30]. We hope that our analysis on rather simple binary coalescence will motivate future studies on complex droplet collisions.

\textbf{Summary.} – In summary, the coalescence preference (CP) dynamics during non-equilibrium phase separation of a single-component fluid has been investigated using molecular dynamics simulations and scaling arguments. Due to the off-critical order parameter of the fluid, phase
separation progresses via coalescence of liquid droplets. To the best of our knowledge, this is the first study on CP during any coarsening process following a sudden temperature quench. Our results reveal that the location of the product drop exhibits strong dependence on the size inequality of the parents. With increasing size inequality the product drop forms much closer to the larger parent. Although this general trend is in accord with previous experiments [10] on microbubble/droplet the CP exponent, which characterizes the “closeness” of preferential location to the larger parent, it is found to be different from previous reports in the literature. Specifically, for the present study $\simeq 3.1$ and earlier studies [10,11] reported $q \simeq 2.1$ and 5.1. Such a discrepancy is attributed to the underlying mechanisms of the coalescence event. If the motion of the merged bubble after the initial bridge formation, in a low-density system, is controlled by the surface energy difference between the parent and the merged bubble it leads to a CP exponent $q \simeq 5.1$. For a densely packed system, this exponent gets modified to $q \simeq 2.0$ because of the presence of multiple neighbouring bubbles which might block the position of the product bubble. On the other hand, in a phase-separating single-component fluid droplet coalescence occurs via the inelastic sticky collision and coagulation and after coalescence the product droplet forms at the centre-of-mass (c.m.) location of the parents. Scaling arguments provide a value of $q = 3$ for this c.m. theory with which the MD simulation data accord well. This is in contrast with droplet coalescences in [11] where the merged product does not form at the c.m. location, but much closer to the larger parent. We hope that our study on droplet coalescence in a phase-separating single-component fluid will promote future investigations on droplet coalescence in binary-liquid mixtures, coalescence of bubbles, which will in turn lead to the understanding of the universality in coalescence preferences effects.

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