Highlights

- Effect of density ratio on the mechanisms of secondary breakup has been studied. A density ratio-Weber number phase plot is presented to depict different breakup mechanisms.

- Reynolds number and viscosity ratio shift the breakup transition Weber number values.

- Instantaneous Weber number is defined to explain the non-breakup of deforming droplets at low density ratios.
Secondary breakup of drops at moderate Weber numbers: 
Effect of Density ratio and Reynolds number

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Abstract

Breakup of liquid drops occurs in several natural and industrial settings. Fully resolved volume of fluid based simulations presented in this study reveal the complete flow physics and droplet dynamics that lead to the breakup of a drop in the moderate Weber number regime. We have investigated the effects of density ratio and Reynolds number on the initial dynamics of drop deformation. A density ratio-Weber number phase plot is presented that indicates the variation in the deformation of the drop at various density ratios and Weber numbers. We show that the initial deformation and dynamics of the droplets at low density ratios is significantly different from that observed at high density ratios. We also study the temporal characteristics of the initial droplet deformation and motion.

1 Introduction

When a drop is accelerated in a high speed gas flow, it deforms due to the aerodynamic forces and eventually fragments into tiny droplets; this is termed as secondary breakup. This phenomenon has been studied over many decades in the interest of its numerous applications, for example, in rainfall, sprays, combustion and chemical industries. Complete understanding of the breakup phenomenon is essential for an accurate determination of the drop size distribution which dictates the surface to volume ratio and hence the efficiency of drying, chemical reaction and combustion. Further, a better understanding of the breakup also helps in developing accurate closure relations for Lagrangian and Eulerian multi-fluid modeling approaches.

Over the years, numerous experimental and numerical studies have been performed to study the secondary breakup of drops. Several articles (Pilch and Erdman (1987); Faeth et al. (1995); Guildenbecher (2009)) have periodically reviewed the advances in this field. The secondary breakup of a drop can be broadly categorized into four modes of deformation and breakup, primarily based on the aerodynamic Weber number and liquid Ohnesorge number: (a) Vibrational mode, where a drop oscillates at its natural frequency and it may (or may not) undergo breakup (Hsiang and Faeth, 1992) and when it breaks it produces fewer daughter drops of the size comparable to that of the parent

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drop (Pilch and Erdman, 1987), (b) Bag mode, where a drop deforms into a flat disc and then is
blown into a thin bag, attached to a toroidal ring, that expands and eventually ruptures, followed by
the breakup of the toroidal ring (Chou and Faeth, 1998; Jain et al., 2015). Bag fragments into a large
number of smaller sized drops and the ring breaks up into a smaller number of larger sized drops.
With an increasing Weber number, some interesting features appear such as bag with a stamen (Pilch
and Erdman, 1987) and bag with multiple lobes (Cao et al., 2007). This phenomenon is thought to
be due to Rayleigh-Taylor (RT) instability (Theofanous et al., 2004; Zhao et al., 2010; Jain et al.,
2015) or a combined RT/aerodynamic drag mechanism (Guildenbecher, 2009). (c) Sheet thinning
mode for higher Weber number, where the ligaments and small daughter drops break off from the
thinning rim of the parent drop until the core of the parent drop reaches a stable state. Earlier,
shear stripping (a viscous phenomenon) was assumed to be the mechanism (A. A. Ranger, 1969),
but later Liu and Reitz (1997) proposed the sheet thinning mechanism, pointing out that it is an
inviscid phenomenon. (d) Catastrophic mode, where a drop breaks up into multiple fragments due to
unstable surface waves at high speeds (Liu and Reitz, 1993). The transition from one breakup mode
to the other occurs very gradually as a function of Weber number ($We$) (as shown later in Figure 9).
Different studies have proposed different transitional values of $We$ (subject to the inaccuracies in
the exact calculation of $We$ and also the presence of impurities that alter the properties of the fluid
used in the experiments; see Table 1) and hence the reliability of the transitional values of $We$ has
remained a moot point. Other parameters that influence the breakup mechanism are the density
ratio and the gas Reynolds number (and the liquid Ohnesorge number).

Several experimental studies have been performed in the last decades to unravel the physics of
secondary breakup of droplets (see Guildenbecher (2009) for an elaborate review). Some of the
studies have been performed at low density ratios (in the range 1 – 10) while most of the studies
have been performed for water-air systems. Simpkins and Bales (1972) and Harper et al. (1972)
presented experimental and theoretical studies, respectively, of secondary breakup of droplets at
high Bond numbers. Harper et al. (1972) showed that Rayleigh-Taylor instability is dominant at
higher Bond numbers after a short time algebraic deformation in time. Experiments of Simpkins
and Bales (1972) corroborated the theoretical findings. Patel and Theofanous (1981) studied the
fragmentation of drops moving at high speed for mercury/water system (density ratio $\sim$ 10). They
showed that the breakup time even at low density ratios correlates well with the time constant
for the growth of unstable Taylor waves in the entire range of Bond numbers. Later, Theofanous
et al. (2004) studied droplet breakup at different static pressures over a wide range of gas densities,
all in the rarefied range. In this study, they noted that the upstream face of the droplet, upon
droplet flattening, becomes immediately susceptible to Rayleigh-Taylor instability. Further, they
showed that for low Weber numbers, Bag formation corresponds to one Rayleigh-Taylor wave on
the ‘disc-shaped’ droplet. Lee and Reitz (2000) performed experiments for a range of gas densities
(corresponding to density ratios 100 – 1000). Using a pressurized chamber the ambient air pressure
was controlled to vary the density ratio from 100 to 1000. They concluded that the Rayleigh number
and also the density ratio have little effect on the drop breakup mechanisms, although the transition
liquid systems and discussed the similarities between the features observed in the two systems. He
noted that the value of the first critical Weber number in liquid-liquid systems is higher than the
gas-liquid systems. He also noted that the breakup features such as bag and bag-stamen that are
observed at low $We$ (around 15 to 40) for gas-liquid systems, do not appear clearly in the liquid-liquid
systems.

Most of the numerical studies have been performed with low density ratios and only a few with
high density ratio. Efforts in numerical studies have only started to pay off recently and most
numerical simulations attempt to study the breakup at low density ratios ($\rho^* < 100$), essentially
Table 1: Transition Weber number (\(We\)) for different breakup regimes.

<table>
<thead>
<tr>
<th>Breakup regime</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vibrational</td>
<td>(We &lt; 12)</td>
<td>(We &lt; 10)</td>
<td>(We &lt; 11)</td>
<td>(We &lt; 13)</td>
<td>(We &lt; 18)</td>
<td>(We &lt; 35)</td>
<td>(We &lt; 35)</td>
</tr>
<tr>
<td>Bag</td>
<td>12 &lt; (We &lt; 50)</td>
<td>10 &lt; (We &lt; 18)</td>
<td>11 &lt; (We &lt; 35)</td>
<td>13 &lt; (We &lt; 35)</td>
<td>13 &lt; (We &lt; 18)</td>
<td>12 &lt; (We &lt; 24)</td>
<td></td>
</tr>
<tr>
<td>Bag-stamen</td>
<td>50 &lt; (We &lt; 100)</td>
<td>18 &lt; (We &lt; 30)</td>
<td>35 &lt; (We &lt; 80) (multimode)</td>
<td>35 &lt; (We &lt; 80) (multimode)</td>
<td>18 &lt; (We &lt; 40)</td>
<td>28 &lt; (We &lt; 41)</td>
<td>45 &lt; (We &lt; 65)</td>
</tr>
<tr>
<td>Plume-shear</td>
<td>63 &lt; (We &lt; 80)</td>
<td>10 &lt; (We &lt; 40)</td>
<td>40 &lt; (We &lt; 80)</td>
<td>40 &lt; (We &lt; 80)</td>
<td>65 &lt; (We &lt; 85)</td>
<td>65 &lt; (We &lt; 85)</td>
<td></td>
</tr>
<tr>
<td>Sheet (Shear)</td>
<td>100 &lt; (We &lt; 350)</td>
<td>(We &gt; 63)</td>
<td>(80 &lt; We &lt; 800)</td>
<td>(85 &lt; We &lt; 120)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>thinning</td>
<td>(We &gt; 350)</td>
<td>(We &gt; 800)</td>
<td>(We &gt; 800)</td>
<td>(We &gt; 800)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>


Due to numerical convergence issues at high density ratios. Nevertheless, these studies find direct applications in high-pressure environment applications as well as in manufacturing of metal pellets by quenching liquid metal droplets. Zaleski et al. (1995) performed one of the earliest numerical studies of the secondary breakup of drops in 2D. They observed a backward-bag at low Weber number (\(We\)) for \(\rho^* = 10\) and reported that their results contradict the general experimental observations (which were mostly done for higher \(\rho^*\) values), where a forward-bag is seen at this \(We\). They suggested that this mismatch is a result of the discrepancy in their initial conditions. Han and Tryggvason (2001) extensively studied the breakup of drops for two \(\rho^*\) values, 1.15 and 10. For \(\rho^* = 10\), they observed a forward-bag at low \(We\) and backward-bag at higher \(We\), and for \(\rho^* = 1.15\) they observed backward-bag for all moderate \(We\). They concluded that the formation of forward-bag is due to the detachment of the wake downstream of the drop and the formation of backward-bag is due to the entrapment of the drop in the vortex ring. On decreasing \(Re\), they also observed that a higher \(We\) is required to obtain the same mode of breakup. Aalburg et al. (2003) reported that the secondary atomization is essentially independent for \(\rho^* > 32\) and that there is no effect of \(Re\) on \(We_{crit}\) beyond \(Re > 100\). Kékesi et al. (2014) studied the breakup of drops for \(\rho^* = 20, 40\) and 80 and reported to have observed new breakup modes such as Bag, Shear, Jellyfish shear, thick rim shear, thick rim bag, rim shear and mixed. The new breakup modes were due to the influence of the viscous effects in their simulations (some of these cases are at \(Oh > 0.1\) and \(Re_g < 100\)). Yang et al. (2016) also studied the effect of \(\rho^*\) on the breakup but at a very high \(We = 225\) value in the regime of catastrophic breakup for \(\rho^* = 10\), 25, 32, 60. On decreasing \(\rho^*\), they observed a lower deformation rate but the range of \(\rho^*\) values chosen was probably too low at such high \(We\) to see any discernible effect of changing \(\rho^*\) on the breakup. Formation of spherical cap and ligaments and the fragmentation of ligaments further into multiple drops were the common features they observed in their study. Recently, 3D simulations were performed for water and air at atmospheric conditions (\(\rho^* \sim 1000\)) by Xiao et al. (2014) where they validated their LES code. In more recent studies, Xiao et al. (2016) and Xiao et al. (2017) have studied the effect of \(Oh\) and Mach number on the breakup of a drop, respectively. We, in our previous work (Jain et al., 2015), have extensively studied the breakup and its characteristics for \(\rho^* = 1000\) using fully resolved 3D simulations.

For the systems with low density ratios (<100) and at moderate Weber numbers (20-80), backward-bag (opening of the bag facing the downstream direction followed by sheet thinning) has been seen as the predominant breakup mode in the numerical simulations (see Kékesi et al. (2014); Khosla et al. (2006)). In the present work, we numerically study the effect of a wide range of density ratios on the drop breakup mechanisms at different aerodynamic Weber numbers. We focus on the effect of flow features in the surrounding medium on the drop deformation. We note that the initial deformation into a bag or bag-stamen essentially drives the subsequent fragmentation.
The paper is organized as follows: Section 2 describes the equations solved, the computational domain and the grid independence study. Results and discussion are presented in Section 3. The summary of the study and important conclusions are discussed in Section 4.

2 Problem description and Formulation

Figure 1 shows the schematic of the computational domain for the axisymmetric simulations performed in this study with the dashed line marking the axis of symmetry. The domain is 10\(d_0\) along the radial direction and 20\(d_0\) along the axial direction, where \(d_0\) is the diameter of the drop. Liquid and gas densities are \(\rho_l\) and \(\rho_g\), respectively, and the ratio \(\rho^* = \rho_l/\rho_g\) is varied from 10 to 1000 by keeping the gas density as unity and varying the liquid density. Viscosity of the liquid and the gas are given by \(\mu_l\) and \(\mu_g\), respectively. Surface tension coefficient at the liquid-gas interface is given by \(\sigma\).

Boundary condition on the left end of the computational domain is that of gas inlet and a uniform velocity of \(U_g\) is prescribed. Outlet flow boundary conditions are imposed at the right end of the computational domain. Slip boundary conditions are applied at the other (side) walls of the domain to minimize the confinement effects and axisymmetric boundary conditions are imposed at the axis of symmetry marked by the dashed line in Figure 1. The drop is accelerated by the high-speed gas flow and its breakup is governed by the following five non-dimensional numbers: aerodynamic Weber number \(We = \rho_g U_g^2 d_0/\sigma\) (ratio of the gas inertial forces to the surface tension forces), liquid Ohnesorge number \(Ob = \mu_l/\sqrt{\rho_l d_0 \sigma}\) (ratio of the viscous capillary time scale, \(\mu_l d_0/\sigma\), to the inertial capillary time scale \(\sqrt{\rho_l d_0 \sigma}\)), gas Reynolds number \(Re = \rho_g U_g d_0/\mu_g\) (ratio of gas inertial forces to the gas viscous forces), viscosity ratio \(M = \mu_l/\mu_g\) (ratio of drop viscosity to the gas viscosity) and the density ratio \(\rho^* = \rho_l/\rho_g\) (ratio of drop density to the gas density). For the 3D simulations presented in the later part of the paper, we have used the computational setup similar to the one discussed in Jain et al. (2015).

A one-fluid formulation is used for the numerical simulations (Mirjalili et al., 2017). The governing equations for the coupled liquid and gas flow simulated in this study are described in the following. Considering both the drop fluid and the surrounding gas to be incompressible, the corresponding continuity equation is given by,

\[ \nabla \cdot \mathbf{u} = 0, \]  

Figure 1: Schematic of the simulation setup (not drawn to scale). Flow direction represents the direction of the gas flow. Drop is shown at the initial time \((t = 0)\).
where \( \mathbf{u} \) is the divergence free velocity field. The governing equations for the momentum conservation are given by the Navier–Stokes equations (Eqn. 2) modified to implicitly account for the surface tension forces and the interfacial boundary conditions of continuity of velocity, and normal and tangential stress balance:

\[
\rho(C) \left( \frac{\partial \mathbf{u}}{\partial t} + \nabla \cdot (\mathbf{u} \mathbf{u}) \right) = -\nabla p + \nabla \cdot (2\mu(C) \mathbf{D}) + \sigma \kappa \mathbf{n} \delta_s. \tag{2}
\]

Here, \( C \) is the volume fraction of liquid that takes a value of zero in the gas phase and one in the liquid phase. The density and viscosity for the one-fluid formulation are expressed as, \( \rho = \rho_l C + \rho_g (1 - C) \) and \( \mu = \mu_l C + \mu_g (1 - C) \), respectively. The deformation rate tensor is given by \( \mathbf{D} = \frac{1}{2}(\nabla \mathbf{u} + (\nabla \mathbf{u})^T) \).

The last term in the equation \( (\sigma \kappa \mathbf{n} \delta_s) \) accounts for the surface tension force \( (\sigma \kappa, \kappa \text{ is the local interface curvature}) \) acting on the interface, expressed as a volumetric force using the surface Dirac delta function \( \delta_s \) and modeled using the continuum surface force approach (Brackbill et al., 1992). The direction of this force is along the local normal \( (\mathbf{n}) \) at the interface. The evolution equation for the interface is given as an advection equation in terms of the volume fraction, \( C \), obtained by applying kinematic boundary condition at the interface:

\[
\frac{\partial C}{\partial t} + \mathbf{u} \cdot \nabla C = 0. \tag{3}
\]

We use a cell-based Octree grid adaptive mesh refinement (AMR) geometric volume of fluid (VOF) algorithm in Gerris (see Popinet, 2003, 2009; Tomar et al., 2010) to solve the above set of equations. Gerris uses a second-order accurate staggered time discretization for velocity, volume-fraction and pressure fields. Balanced-force algorithm by Francois et al. (2006) is used to accurately calculate the surface tension forces and minimize spurious currents. The discretization of the equations (Eqn. 1–4) are described in detail in Popinet (2003) and will be discussed here only briefly.

Discretized Navier–Stokes equations are solved using a projection method. First, an auxiliary velocity field is obtained using the following discretization (Popinet (2003)):

\[
\rho \left( \frac{\mathbf{u}^* - \mathbf{u}_n}{\Delta t} + \nabla \cdot (\mathbf{u} \Delta \mathbf{u} + \frac{1}{2}) \right) = \nabla \cdot (\mu \left( \mathbf{D}_n + \frac{1}{2} \right) + (\sigma \kappa \mathbf{n} \delta_s) \mathbf{n} + \frac{1}{2}). \tag{4}
\]

Void fraction is updated using the following equation with the fluxes computed geometrically (Popinet (2009)):

\[
\frac{C_{n+\frac{1}{2}} - C_{n-\frac{1}{2}}}{\Delta t} + \nabla \cdot (C_n \mathbf{u}_n) = 0. \tag{5}
\]

The pressure Poisson equation:

\[
\nabla \cdot \left( \frac{\nabla p}{\rho(C)} \right) = \frac{\nabla \cdot \mathbf{u}_n}{\Delta t}, \tag{6}
\]

is solved using a geometric multigrid method and the auxiliary velocity field is updated as following,

\[
\mathbf{u}_{n+1} = \mathbf{u}_n - \frac{\Delta t}{\rho_{n+\frac{1}{2}}} \nabla p_{n+\frac{1}{2}}, \tag{7}
\]

to obtain a divergence free velocity field \( (\nabla \cdot \mathbf{u}_{n+1} = 0) \). Adaptive mesh refinement (AMR) is performed using a cost function based on the local vorticity in the field and the gradient of the
void-fraction field, thus using a very fine refinement in the regions of high velocity gradient and at the interface. We use four different grid resolutions for the refinement of the surrounding gas flow (refined based on the local vorticity values in the flow) and the interface region - $d_0/\Delta x_{\text{min}} = 102, 204, 410$ and 820 for our 2D axisymmetric simulations. These resolutions are more than that employed in any of the previous studies in the literature. For example, Han and Tryggvason (2001) used around 100 and (Kékesi et al., 2014) used 32 number of grid points per droplet diameter. Figure 2 shows the drop shapes at different grid refinements and density ratios.

Drop shapes for grid resolutions $d_0/\Delta x_{\text{min}} = 410$ and 820 are very close (in a physically meaningful way) and show clear convergence. Hence, we use the mesh refinement of $d_0/\Delta x_{\text{min}} = 410$ for all the axisymmetric simulations presented in this study. We also perform a few 3D simulations using a mesh refinement of $d_0/\Delta x_{\text{min}} = 412$ in the interface region and $d_0/\Delta x_{\text{min}} = 102$ in the surrounding flow region (grid refinement in the surrounding gas flow region has a negligible effect on the drop shape as long as the $Re_l$ is not too high) to show the validity of our axisymmetric assumption. We found that the grid refinement depends on the liquid Reynolds number of the flow $Re_l = \rho_l U_g d_0/\mu_l$ and for $Re_l$ values of about 10000 and above, a grid refinement of at least $d_0/\Delta x_{\text{min}} \sim 410$ is required to avoid any unphysical surface fragmentation and corrugations at the interface. Thus, in order to maintain high-fidelity of the simulations (and consistency between axisymmetric and 3D simulations), we performed all the simulations with an $Re_l$ that is sufficiently smaller than 10,000.

In order to perform high density ratio simulations, as discussed in (Jain et al., 2015), since a sharp interface (single grid transition region) results in an inexplicable spike in the kinetic energy, we use a thin transition region for better convergence. We use a transition region of two cells for $\rho^* \geq 500$ and one cell for $500 > \rho^* \geq 100$ on either sides of the interface for smoothing the jump in the physical properties. To test the efficacy of the numerical algorithm in capturing high density ratios, mainly the use of a thin smoothing width at the interface, we had presented validation test cases in (Jain et al., 2015) which show good agreement with the corresponding analytical results for a density ratio of 1000. In the following section, we discuss the simulations for a wide range of liquid to gas density ratio to study its effects on the dynamics of drop deformation.
Table 2: Parameters for the different simulations presented in this study.

<table>
<thead>
<tr>
<th>$\rho^*$</th>
<th>$We$</th>
<th>$Re$</th>
<th>$M$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$10, 50, 100, 150, 200, 250, 500, 1000$</td>
<td>$20, 40, 60, 80, 100, 120$</td>
<td>$4000$</td>
<td>$100$</td>
</tr>
<tr>
<td>$1000$</td>
<td>$20$</td>
<td>$141, 500, 1414$</td>
<td>$100$</td>
</tr>
<tr>
<td>$1000$</td>
<td>$20$</td>
<td>$1414, 4000, 6000$</td>
<td>$1000$</td>
</tr>
<tr>
<td>$10$</td>
<td>$20, 40, 60, 80, 100, 120$</td>
<td>$20000$</td>
<td>$100$</td>
</tr>
</tbody>
</table>

3 Results and discussion

In order to investigate the effects of density ratio, $\rho^*$, on the secondary breakup of a drop, we perform a large set of well resolved simulations with different values of $\rho^*$, Reynolds number $Re$ and aerodynamic Weber number, $We$. Table 2 lists the parameter range covered in this study. For each value of $\rho^*$ listed in the table, we vary the Weber number to study the effects of density ratio on the drop breakup regimes namely, bag breakup, bag with stamen, multi-bag breakup and sheet-thinning breakup which are observed experimentally (Guildenbecher, 2009) for the range of $We$ chosen here for the simulations. The liquid Ohnesorge number for all the simulations is $\leq 0.1$ and therefore, the critical Weber number based on the previous studies (Hsiang and Faeth, 1992; Krzeczkowski, 1980) is not expected to vary significantly.

3.1 Validity of axisymmetric assumption

In this section, we show comparisons of 2D axisymmetric and 3D simulations, to validate the 2D axisymmetric assumption that we make in the analysis throughout the article. Since 3D simulations are more expensive, we use a mesh resolution of $d_0/\Delta x_{\text{min}} = 102$ for the simulations presented only in this section for both 2D axisymmetric and 3D simulations (only to show the validity of axisymmetry assumption by making a one-to-one comparison). For more refined and grid converged axisymmetric simulations of the same, refer to Figure 10. First, we present the case with density ratio of $\rho^* = 10$ and Weber number $We = 60$. The drop shapes in Figure 3a and 3b match very well to first order for all times.

The comparison between 2D and 3D calculations for the density ratio of $\rho^* = 50$ and Weber number $We = 60$ are presented next. Here again, the drop shapes in Figures 4a and 4b agree very well for all times. Hence, the validity of the axisymmetric assumption is well justified, at least up to the onset of breakup. After the onset of breakup, a daughter drop that breaks from the rim of the parent drop in the 2D axisymmetric simulation representing a toroidal ligament (a ring) in 3D, and the ring further undergoes breakup into multiple smaller droplets due to capillary instability. Thus, the axisymmetric assumption is not valid after the onset of breakup.

We note here that although the axisymmetric simulations cannot be employed to study the complete fragmentation of the drop, the axisymmetric assumption is valid for initial drop deformation which essentially determines the final breakup mode. For example, for a drop breaking up via the bag-stamen mode, the initial deformation, formation of the bag and the stamen are nearly axisymmetric deformations, however, the rupture of the bag and subsequent breakup of the rim and the stamen are clearly three dimensional features. In the present study, in order to perform a large set of simulations, we have restricted ourselves to axisymmetric simulations.
Figure 3: 2D axisymmetric and 3D simulations showing the evolution of the drop shape in time for $\rho^* = 10$ at $We = 60$. Time $t^*$ values are noted on top for each drop.
Figure 4: 2D axisymmetric and 3D simulations showing the evolution of the drop shape in time for $\rho^* = 50$ at $We = 60$. Time $t^*$ values are noted on top for each drop.
3.2 Effect of density ratio

As discussed earlier in Sec.1, there are conflicting views on the effect of density ratio on the breakup mechanism for a given $We$. Where Aalburg et al. (2003) reported that there is little effect of density ratio (for $\rho^* > 32$) on the breakup mechanism, Jing and Xu (2010) presented simulations for density ratios 10, 100 and 1000 and showed that a change in the density ratio alters the critical Weber numbers for the different breakup regimes. Several simulations have been reported for low density ratios where for even a reasonably lower $We$, backward bag was observed (see, for example, Han and Tryggvason (2001),Khosla et al. (2006)). In this section, we present simulations for a wide range of density ratios and discuss the physical mechanisms that alter the breakup modes.

In what follows, we discuss the effect of density ratio on the deformation and motion of the droplet. The time is non-dimensionalized with the characteristic time scale, $t^* = t/t_c$, where,

$$t_c = \frac{d_0 \sqrt{\rho_l/\rho_g}}{U_g}.$$  \hspace{1cm} (8)

The characteristic time scale is defined by using the diameter of the drop as the characteristic length scale and the velocity scale in the drop obtained by comparing the dynamic pressures in the gas and the drop. Figure 5 shows the time evolution of the drop shape and the displacement for $\rho^* = 10$ and $\rho^* = 1000$ at $We = 20$. The flow of the gas is from left to right. Centroid of the drop of $\rho^* = 10$ moves a distance of $0.79d_0$, for the drop of $\rho^* = 200$ it moves a distance of $0.48d_0$ (not shown in the figure) and for the drop of $\rho^* = 1000$ it moves a distance of $0.34d_0$ in $t^* = 1$. The leeward side of the drop for $\rho^* = 10$ also moves downstream with time, whereas the leeward side of the drop for $\rho^* = 1000$ remains virtually stationary until $t^* \sim 1$, though the centroid is moving in the streamwise direction in both the cases. The significant difference in the motion of the centroid is primarily due to the differences in the velocity of the drop and the rate of momentum transmitted to the leeward side of the drop, which depends on the kinematic viscosity, $\nu$. The value of $\nu$ for $\rho^* = 10$ is 100 times the $\nu$ for $\rho^* = 1000$. We can also observe the formation and motion of the capillary waves emanating from the rim of the drops in both the cases (more evidently for the drop with $\rho^* = 1000$).

Capillary time-scale based on inertia, also called as the Rayleigh time-scale (Rayleigh, 1879), is given by $t_R \sim \sqrt{\rho_l d^3/\sigma}$. This is around 3 times the characteristic time scale $t_c$ of the drop in both the cases, since both $t_c$ and $t_R$ are proportional to $\sqrt{\rho_l}$. Capillary time-scale based on viscous forces is given by $t_M \sim \mu d/\sigma$. This is around 0.16 times $t_c$ for the drop with $\rho^* = 10$ and around 0.016 times $t_c$ for the drop with $\rho^* = 1000$, thus suggesting a lower resistance to the waves by the viscous forces in the case of high density ratio relative to the low density ratio case, since the $\nu$ is smaller for high density ratios. The stretching time-scale (for the rim) is obtained by the scaling $t_s \sim d/u_{rim}$. This is around 0.76 times the $t_c$ for the drop with $\rho^* = 10$ and 1.12 times the $t_c$ for the drop with $\rho^* = 1000$. Comparing these time scales, we expect the capillary reorganization to occur at a rapid rate in the high-density ratio case in comparison with the low density ratio cases. However, when we performed another simulation with a higher viscosity for $\rho^* = 1000$ case (not shown here; this case is similar to the one in Figure 5b but with $\mu_l$ increased by 10 times so that it would have the same capillary time-scale based on viscous forces as the drop in Figure 5a when non-dimensionalized by $t_c$; this case would also have the same Ohnesorge number as the one in Figure 5a), we still observe formation of a forward bag. Therefore, we note that capillary rearrangement timescales alone do not result in the formation of backward or forward bag formation and resolution of this aspect of drop deformation required further attention. Note that our definition of backward bag is the one where the rim of the bag is stretched in the direction of the flow relative to the bag. This is different from the one proposed by Han and Tryggvason (1999) but is consistent with the one used in Jain et al. (2015).
Figure 5: Time evolution of the drop movement for $\rho^* = 10$ and $\rho^* = 1000$ at $We = 20$.

Figure 6: Drop shapes at $t^* = 1$ for $\rho^* = 10, 50, 100, 150, 200, 250, 500$ and 1000 (left to right) and $We = 20$. Arrow denotes the direction of gas flow.
Figure 6 shows the drop shapes at $t^* = 1$ for $We = 20$ and for different values of $\rho^*$. The flow of the gas is from left to right. We note that the drops at high density ratios deform into a flat disc at around $t^* = 1$ from an initially spherical shape, whereas at low density ratios ($\rho^* = 10$), drop progressively deforms into a backward bag without achieving a proper flat disc shape (see Fig.5a). Note that our definition of backward bag is the one where the rim of the bag is stretched in the direction of the flow relative to the bag. This is different from the one proposed by Han and Tryggvason (1999) but is consistent with the one used in Jain et al. (2015). For intermediate density ratios, the extent of bending of the disc progressively decreases with increase in $\rho^*$ and a near flat disc is obtained for $\rho^* = 1000$. We would like to note here that although $t^* = 1$ is same for all the profiles shown in Fig.6, the dimensional time varies as $\sqrt{\rho^*}$. Hence the physical time corresponding to $t^* = 1$ is smaller for low density ratios when compared to higher density ratios. Drop shape for $\rho^* = 10$ is of nearly uniform thickness whereas for intermediate density ratios ($50 < \rho^* < 150$), the drops become thinner near the rim. The thickness at the center of the drop initially decreases and beyond $\rho^* = 150$ it increases slightly and saturates, whereas the diameter of the disc initially increases and beyond $\rho^* = 150$ it undergoes a sudden decrease and saturates (see Fig.6). The variation in the curvature of the drop with increasing $\rho^*$ suggests a decreasing tendency of forming a backward bag.

Figure 7 shows the non-dimensional (with respect to $U_g$) velocities of the drop and the surrounding gas at various locations for the drops with $\rho^* = 10, 150$ and 1000 at $t^* = 1$. As expected (based on the scaling relation $U_l \sim \sqrt{\frac{1}{\rho^*} U_g}$), the axial and radial components of the velocity at the center ($u_{center}$) and at the rim ($u_{rim}$) of the drop decrease with an increase in $\rho^*$. A vortex is formed behind the drop due to the flow separation as shown in the Figure 7. We can see that the velocities of the vortex (strength of the vortex) in the gas flow is increasing with an increase in $\rho^*$ due to the higher relative velocity of the drop ($u_l - U_g$) for higher $\rho^*$ values. Interestingly, the axial component of the velocity at the rim of the drop is significantly higher than that at the center of the drop at lower $\rho^*$. But this difference in velocity at the center and rim $u_{center} - u_{rim}$ is decreasing with increase in $\rho^*$, and for $\rho^* = 1000$, we can see that the velocities are almost the same at the rim and that at the center. Percentage of the difference in the velocity at the rim and that at the center (relative to the velocity at the center of the drop), $(u_{rim} - u_{center})/u_{center}$, is 13.3% for $\rho^* = 10,
Figure 8: Evolution of the drop shape in time for $\rho^* = 10$ at $We=20$, 40 and $\rho^* = 50$ at $We = 20$. Non-dimensionalized time, $t^*$, values are mentioned on top for each drop shape. Arrows show the direction of the gas flow and the dotted lines mark the axis of symmetry. Note that the distance between consecutive droplet profiles plotted here does not represent the actual displacement of the drop.

6.5% for $\rho^* = 150$ and $\sim 1.2\%$ for $\rho^* = 1000$. Hence, there is more stretching of the rim in the direction of the flow for lower $\rho^*$ values. This explains the bending of the drop for lower $\rho^*$ values and the formation of flat disc for higher $\rho^*$ values.

Figure 8 shows the time evolution of the drop shapes for the cases where drops do not breakup. Drop with $\rho^* = 10$ at $We = 20$ deforms into a concave-disc facing downstream and then bends in the opposite direction and finally collapses without breakup, encapsulating a bubble within it. For $We = 40$ and $\rho^* = 10$, it deforms into a backward-bag and again collapses onto itself before it could break. For $\rho^* = 50$ at $We = 20$, the drop deforms into a concave-disc facing downstream and then into the shape of a canopy-top. Subsequently, with further deformation of the drop, the rim tends to pinch-off from the core drop, but before it could pinch-off, the drop relaxes back collapsing onto itself without breakup. This also shows the highly complicated unsteady behavior of evolution of the drop shapes. To understand this behaviour of no-breakup, we calculate the instantaneous Weber number (based on the velocity of the gas relative to the drop velocity) at the onset of breakup using $We_{\text{inst}} = \rho_g (U_g - u_{\text{drop}})^2 d_0/\sigma$. Estimating the centroid velocity of the drop $u_{\text{drop}}$ from the simulations at $We = 20$, we find that the $We_{\text{inst}} = 3.69$ for $\rho^* = 10$, $We_{\text{inst}} = 8.91$ for $\rho^* = 50$ and $We_{\text{inst}} = 11.1$ for $\rho^* = 100$, and $We_{\text{inst}}$ increases further with increase in $\rho^*$ value. Clearly for $\rho^* = 10$ and $\rho^* = 50$, $We_{\text{inst}}$ is below the $We_{\text{crit}} \sim 10 - 12$, implying that the drop would not breakup. Similarly, at $We = 40$, $We_{\text{inst}} = 8.92$ for $\rho^* = 10$ and $We_{\text{inst}} = 18.31$ for $\rho^* = 50$. Here again for $\rho^* = 10$, $We_{\text{inst}}$ is below the $We_{\text{crit}}$ whereas, for $\rho^* = 50$, $We_{\text{inst}} > We_{\text{crit}}$ for an initial aerodynamic Weber number $We = 40$, and thus we observe breakup of the drop. These predictions based on the criterion $We_{\text{inst}} > We_{\text{crit}}$ for the breakup of drop are in good agreement with our numerical results (as also shown in Figure 8). Thus, we can conclude that the breakup of a drop not only depends on the initial $We$ value but also on the instantaneous dynamics of the drop. More importantly, for low density ratio, for the same momentum transfer the relative velocity decreases much faster in comparison to the rates of deformation of the drop, thus, the instantaneous $We$ decreases sharply and vibrational modes, without breakup, are observed.

In order to study the morphology of the drops during the breakup, typical shapes of the drops
Table 3: Typical shapes of the drop at the onset of breakup for $\rho^* = 10 - 1000$ at $We = 20 - 120$, $Re_g = 4000$, $M = 100$ and $Oh = 0.003 - 0.9$. The time $t^*_b$ where the profiles are taken is plotted in the Figure 11a.
at the onset of the breakup have been tabulated in Table 3 for all the conditions listed in Table 2. Again, for the cases presented in Figure 8, no breakup is observed. Comparing the drop shapes for different \( \rho^* \) values (for the same \( We \)) in Table 3 reveals that at \( We = 20 \), a forward-bag (bag opening facing the gas flow) is seen for \( \rho^* = 10 \), transient drop shapes (canopy-top which can also be seen as a shape in between forward-bag and backward-bag) for \( \rho^* = 50 \) and 100, and a forward-bag (bag facing the flow) with stamen for \( \rho^* \geq 150 \). For \( \rho^* = 50 \) and 100, the drop shapes are similar, but for \( \rho^* = 50 \) the rim does not pinch-off from the core drop whereas, for \( \rho^* = 100 \), the rim eventually pinches-off from the drop breaking into a toroidal ring and a smaller drop. There seems to be a progressive change with increase in \( \rho^* \), from canopy shaped drop for \( \rho^* = 50 \) to a drop with a bag and not-so-clear stamen for \( \rho^* = 100 \) and very prominent stamen with a bag for \( \rho^* > 150 \). Interestingly, the stamen is very long for \( \rho^* = 150 \) and it decreases in size with increase in \( \rho^* \). This can be understood by studying the velocities at the tip of the stamen. The non-dimensional velocity at the tip of the stamen (with respect to the centroid velocity of the drop), \( u_{\text{stamen}}/u_{\text{drop}} \), is 0.74 for \( \rho^* = 150 \), 0.85 for \( \rho^* = 250 \) and 0.9047 for \( \rho^* = 1000 \). This clearly implies that there is a higher relative velocity between the stamen and the drop for lower \( \rho^* \) which results in more stretching of the stamen for lower \( \rho^* \) values and hence results in the formation of a longer stamen. This forward-bag with stamen mode of breakup at \( We = 20 \) was observed before in Jain et al. (2015) at \( We = 40 \). This difference in \( We \) may be due to the significantly different values of Oh in Jain et al. (2015) and in the present simulations, though in both cases \( Oh < 0.1 \) was maintained. For example, the \( Oh \) used in Jain et al. (2015) was 0.1 for \( \rho^* = 1000 \) in all the simulations, whereas here we use \( Oh = 0.0035 \) for a similar case of \( \rho^* = 1000 \) at \( We = 20 \). Gas Reynolds number used for \( \rho^* = 1000 \) discussed above is 4000, whereas Jain et al. (2015) performed the simulations at \( Re_g = 1414 \). This effect of \( Oh \) on the drop deformation and breakup will be discussed in detail later.

At \( We = 40 \) and higher, a backward-bag is seen for \( \rho^* = 10 \) (as also observed by Han and Tryggvason (2001) for \( \rho^* = 10 \) at \( Re = 242 \) and \( We \geq 37.4 \)), for \( \rho^* = 50 \) a transient form of sheet-thinning, where the thin rim oscillates like a whiplash (ensuing the motion from the vortex shedding in the surrounding gas flow) and for \( \rho^* = 100 \) and higher, drop deforms into a concave-disc (facing downstream) and eventually breaks up due to sheet-thinning. For \( \rho^* = 200 - 1000 \) at \( We = 40 \) and for \( \rho^* = 200 - 250 \) at \( We = 60 \), we see an interesting “cowboy-hat” shape of the drop. A similar drop shape was observed by Khodaei et al. (2006). For \( \rho^* = 10 \), the length of the rim increases with an increase in \( We \) value, whereas for higher \( \rho^* \) (\( \rho^* = 100 - 1000 \)), the length of the rim decreases with increase in \( We \) and at the bottom-right corner of the table for \( \rho^* = 500 \) and 1000 at \( We = 100 \) and 120, the drops at the onset of breakup are essentially flat discs without any discernible rim. The length and the thickness of the rim is very small and as a result very fine drops are formed during sheet-thinning breakup. Another interesting observation is that the rim is thicker for drops of lower \( \rho^* \). This is possibly due to higher Taylor-Culick velocity \( u_{tc} \) for lower \( \rho^* \), where \( u_{tc} = \sqrt{2\sigma/\rho h} \). Hence, a higher \( u_{tc} \) (retraction velocity) and a higher stretching velocity \( u_s \) at the rim (equal to the values of \( u_{rim} \) in Figure 4), and is of the order of the velocity given by \( \sqrt{T/\rho^* U_g} \) and acts in the opposite direction to \( u_{tc} \) results in the formation of a swollen rim for the drops with lower \( \rho^* \). Consequently, the stretching of the fluid in this swollen rim takes more time resulting in the delayed breakup/pinch-off of the rims of the drops for low \( \rho^* \) values. This is also in agreement with our observed breakup time, \( t^*_{bc} \), for \( \rho^* = 10 \) case as shown later in the Figure 11a.

To summarize the breakup modes presented in Table 3, we draw a phase plot of \( \rho^* \) vs \( We \) shown in Figure 9. Typical shapes for each of the breakup mode is shown beside the plot. We characterize the drop shapes as following, from top to bottom: forward-bag no-breakup, transient no-breakup, transient, forward-bag with stamen, backward-bag, backward-bag with sheet-thinning, whiplash with sheet-thinning and finally the bottom most is sheet-thinning. Hatched region marks
the transition regime indicating transition from bag (forward/backward) to sheet-thinning. Note, the differences between the drop shapes for backward-bag with sheet thinning (for e.g. $\rho^* = 250$ at $We = 40$),whiplash with sheet thinning (for e.g. $\rho^* = 50$ at $We = 60$) and sheet-thinning (for e.g. $\rho^* = 500$ at $We = 120$) are not very evident from the instantaneous shapes of the drop presented in Table 3 at the onset of the breakup. However, the temporal evolution of these drop shapes (see Figure 10) suggest the classification shown in Figure 9. Figure 10a shows the time evolution of the drop for $\rho^* = 10$ and $We = 60$. The whiplash of the rim of the drop for $\rho^* = 10$ is clearly very different from that for $\rho^* = 50$ shown in Figure 10b. For $\rho^* = 50$ and $We = 60$, time evolution in figure 10b shows the whiplash action of the drop rim along with the sheet-thinning at the edges of the drop. Figure 10c, for $\rho^* = 250$ and $We = 40$, shows the formation of a backward bag and subsequent detachment of its rim. Sheet-thinning sets up only after the detachment of the rim. Figure 10d, for $\rho^* = 500$ and $We = 120$, shows the sheet-thinning mode of breakup. To make the differences between these breakup modes and the backward bag even more clearer, the time evolution of the drop for the backward bag case for $\rho^* = 10$ and $We = 60$ has also been included in Figure 10a.

In addition to these differences in the deformation, breakup morphologies and breakup modes, the breakup mechanism is also different for higher and lower $\rho^*$ values. We note here that the final fragmentation is determined by the mode of deformation the droplet undergoes and as will be discussed in the subsequent section, the axisymmetric assumption for drop deformation are valid up to near fragmentation point before rupture of the bag/rim starts. The breakup is due to the Rayleigh-Taylor (RT) instability at higher $\rho^*$ values ($\rho^* \geq 150$) (Zhao et al., 2010; Jain et al., 2015), whereas the breakup is due to the dynamics of the rim at lower $\rho^*$ values and is significantly influenced by the surrounding gas flow (Section 3.5). Hence, drops for roughly $\rho^* \geq 150$ behave similarly at similar values of $We$. This difference in breakup for different $\rho^*$ values (with $Re_g$, $M$ being constant and $Oh < 0.1$) makes “Density Ratio” a crucial parameter in characterizing the secondary breakup of drops.

Figures 11a, 11b and 11c show the drop breakup time $t_b^*$ (defined here as the time when the fragmentation begins), the drop displacement, $x_l/d_0$, and the relative velocity of the centroid of the
Figure 10: Evolution of the drop shape in time for $\rho^* = 10$ at $We = 60$, $\rho^* = 50$ at $We = 60$, $\rho^* = 250$ at $We = 40$ and $\rho^* = 500$ at $We = 120$. Non-dimensionalized time, $t^*$, values are mentioned on top for each drop shape. Arrows show the direction of gas flow and the dotted lines mark the axis of symmetry. Note that the distance between consecutive droplet profiles plotted here does not represent the actual displacement of the drop.

Relative velocity, $u_r = (U_g - u_l)/(U_g)$, respectively, at the onset of breakup for the conditions listed in the first row of Table 3. Clearly, $t_b^*$ and $x_l/d_0$ are quite different for the drops with $\rho^* = 10$ and for the drops with $\rho^* = 50 - 1000$. With an increase in $We$, both $t_b^*$ and $x_l/d_0$ decrease following a power-law given by $t_b^* = 9.5 We^{-0.5}$ and $x_l/d_0 = 17 We^{-0.25}$, respectively. Relative velocity $u_r$ on the other hand has a continuous variation from $\rho^* = 50$ to $\rho^* = 1000$ following a general power-law given by $4(10^{-4} \rho + 0.4) We^{(0.15 - 10^{-4} \rho)}$ with average values increasing from 0.76 to 0.95, though it is significantly different for $\rho^* = 10$ with an average value of 0.36. Zhao et al. (2010) reported an average value of 0.9 for ethanol and water drops combined, which are in good agreement with the simulations presented here (also shown in the figure as a line) and Dai and Faeth (2001) reported 0.87 for water drops. Relative velocity, $u_r$, increases with an increase in $\rho^*$ value indicating that the drop for lower $\rho^*$ would attain higher velocity at the onset of breakup. Here, we note that the drops for $\rho^* = 10$ at $We = 20$ and 40 and for $\rho^* = 50$ at $We = 20$, do not breakup at all. This is in good agreement with the observations of Han and Tryggvason (2001). The values corresponding to these values of $\rho^*$ and $We$ reported in Figure 11 indicate only a point where the drops show a tendency to breakup. This tendency to breakup is based on the criteria that the drop could have pinched-off at the thinnest section attained during the deformation process. However, when the simulations are run for a longer duration, the rim retracts and the breakup does not occur.
We... of breakup. Solid line represents the power law fit.

(b) Distance travelled by the drop up to the onset of breakup. Solid line represents the power law fit.

(c) Relative velocity at the onset of breakup. Solid line represents the results by Zhao et al. (2010). Dotted lines show the general power law fit for $\rho^* = 1000$ and $\rho^* = 200$ case.

Figure 11: Relative velocities, time taken and the distance travelled by the drop at the onset of breakup for the parameters listed in the Table 3.

3.3 Rayleigh-Taylor instability

The role of Rayleigh-Taylor instability in the breakup of a drop in catastrophic regime has been extensively discussed in the last decades (Harper et al., 1972; Simpkins and Bales, 1972; Joseph et al., 1999; Patel and Theofanous, 1981; Lee and Reitz, 2000; Guildenbecher, 2009). Harper et al. (1972) showed that, at high Bond numbers (above $10^5$), Rayleigh-Taylor instability dominates the algebraic aerodynamic deformation and leads to the formation of waves on the windward side of the drops. However, the bag formation is mostly considered as blowing out of a thin liquid sheet due to large stagnant pressure. Nevertheless, formation of stamen and multibag at a higher Weber number suggests the role of instability in the formation of bag with a decrease in the wavelength of...
the instability as the acceleration of the droplet increases. Theofanous et al. (2004) showed that the Rayleigh-Taylor based theoretical predictions of number of waves agreed well with the experimental observations for drops with a wide range of viscosity values. Experimental observations of Zhao et al. (2010) also showed good agreement with the theoretical expression for the number of expected waves (1 for bag and 2 for bag-with-stamen) proposed by Theofanous et al. (2004).

In our numerical simulations, we also observe the formation and growth of RT waves on the windward surface of the drop as shown in Figure 12. Shapes of the drop for $\rho^* = 1000$ at $Re_g = 1414$, $M = 1000$, $We = 20$ are shown for four different times, $t^* = 1.006, 1.028, 1.341$ and $1.565$. It is quite evident from these figures that the amplitude, $A$, of the wave grows with time. The drop eventually deforms into a forward-bag and then breaks. We assume this wave on the surface of the drop as an RT wave (highlighted in the Figure 12 using thin red lines) and calculate the non-dimensional RT wavenumber in the maximum cross-stream direction of the drop at $t^* \sim 1.3$ (breakup initiation time at $We = 20$ (Xiao et al., 2014)) as,

$$ N_{RT} = \frac{D_{max}}{\lambda_{max}} = \frac{0.1754}{0.18} = 0.97 $$

This value of $N_{RT} = 0.97$ lies in the range of $1/\sqrt{3} < N_{RT} < 1.0$, which implies that the drop should deform into a forward-bag (Zhao et al., 2010) and then breakup. This is in very good agreement with what we observe in our simulations. Growth rate of the RT wave obtained from the numerical simulations, $\omega_n$, can be calculated by comparing the amplitude of the wave, $A$ (shown in the Figure 12) at two different time instances; assuming normal mode growth of the waves, $A = A_0 e^{\omega t}$. Considering the surface tension effects to be negligible, theoretical growth rate, $\omega_t$, can be calculated using the following expression,

$$ \omega_t = \sqrt{k \alpha \left( \frac{\rho_l - \rho_g}{\rho_l + \rho_g} \right)} $$

We calculated the growth rate of these RT waves at two time instances (a)$t^*_1 = 1.006$ to $t^*_2 = 1.028$ and (b)$t^*_1 = 1.341$ to $t^*_2 = 1.565$. Note that the acceleration of the drop changes over time due to
the change in the drag force with the change in shape and velocity. Therefore, the corresponding
growth rate of the instability on the instantaneous Bond number \( (Bo = \rho g ad^2/4\sigma) \) remains
unchanged (see the relation in Appendix A). For the case (a), the non-dimensional numerical growth
rate, \( \omega^*_n \), is found to be \( \omega^*_n = 2.31Bo^{3/4}/\sqrt{We} \) and the non-dimensional theoretical growth rate is
found to be \( \omega^*_n = 0.7598Bo^{3/4}/\sqrt{We} \), where the instantaneous Bond number is 7.48, and for the case
(b), \( \omega^*_n = 2.54Bo^{3/4}/\sqrt{We} \), where the instantaneous Bond number is 10.73. Interestingly, for both the
cases, \( \omega^*_n \) is \( \sim 3 \) times that of \( \omega^*_g \). These significantly different, yet consistent growth rate values,
could be due to the end-effects associated with the growth of RT waves on the surface of the drop.

3.4 Flow around the drop and the effect of Reynolds number

In this section, we study the flow field around the drop to identify the effects of the flow structures on
the deformation and breakup of the drops. Although, some attempt has been made to comprehend
this through experimental and numerical observations (see Flock et al. (2012); Strotos et al. (2016)),
a more systematic analysis is required to completely characterize the flow field. Figures 13 and 14
show the flow field around the drop at \( t^* = 1.52 \) for \( \rho^* = 1000, M = 1000 \) and \( We = 20 \) at two
different gas Reynolds numbers, \( Re_g = 4000 \) and 1414, obtained from the 3D and 2D axisymmetric
simulations. We observe the formation of a Hill’s spherical vortex in the wake region for the drop in
axisymmetric simulations (Figures 13a and 14a), whereas in 3D simulations (Figures 13b and 14b),
a three-dimensional vortex shedding is seen in the wake region for both the cases. At very early
stages of the drop deformation we also observed the formation of vortex ring in the 3D simulations
(not shown here). This vortex ring starts to shed at around \( t^* \sim 0.15 \) for the drop with \( \rho^* = 1000 \)
value and subsequently at \( t^* = 1.52 \), we see a three-dimensional eddy formation and stretching
resulting in a highly unsteady complex flow in the wake region of the drop. Though, the formation
of a vortex and its shedding behind the drop need not be similar to the flow past a cylinder, a
qualitative comparison could help us in better understanding the flow. Jeon and Gharib (2004)
performed a comparative study of the the vortex in the wake of the cylinder and the formation of
a vortex ring and reported that the shedding of the vortex starts at the non-dimensional time,
\( t/t_{cv} = 4 \), expressed in terms of the characteristic time \( t_{cv} = d_0/U \), where \( U \) is the average velocity
of the vortex generator (relative velocity of the drop with respect to gas, \( U_g - u_{drop} \) in our case) and
d_0 is its diameter. For the present case, we find that the relative velocity, \( (U_g - u_{drop})/U_g \) is 0.92
from our simulations. Hence the vortex shedding is expected to start at \( t^* \sim 0.14 \), which is in very
good agreement with our observations of \( t^* \sim 0.15 \) in our simulations.

Although, the flow around the drops in 2D is significantly different from that in 3D, the shapes
of the drops are surprisingly the same, and both the 2D and 3D simulations predict the formation
of stamen and a forward bag, for two different Reynolds numbers, as shown in Figure 15. This
indicates that the flow structures in the wake do not significantly affect the drop morphology and
breakup for the drops with higher \( \rho^* \) values (lower \( \rho^* \) values are investigated in later sections).
Flock et al. (2012) also reported a similar conclusion based on the differences in the observations
of the PIV realized instantaneous flow fields, showing alternating vortices in the wake region, and
the ensemble-averaged of the flow fields, showing a symmetric twin vortex pair in the wake region
around the drop.

Another interesting observation from Figures 13-15 is that the drop shapes are different for
different \( Re \) (4000 and 1414) values for the same \( M = 1000, \rho^* = 1000 \) and \( We = 20 \). Figure 15
clearly shows the tendency of the drop to form a stamen at \( Re = 4000 \), whereas there is no sign of
formation of stamen for \( Re = 1414 \). Further, Table 4 lists the drop breakup modes along with the
non-dimensional RT wavenumber, \( N_{RT} \), and Table 5 lists the corresponding breakup shapes of the
Figure 13: Comparison of flow field around the drop for $\rho^* = 1000$ at $Re_g = 4000$, $M = 1000$, $We = 20$ and at $t^* = 1.52$ obtained from a 3D and 2D axisymmetric simulation.
Figure 14: Comparison of flow field around the drop for $\rho^* = 1000$ at $Re_g = 1414$, $M = 1000$, $We = 20$ and at $t^* = 1.52$ obtained from a 3D and 2D axisymmetric simulation.
Figure 15: Comparison of drop shapes for $\rho^* = 1000$ at $Re_g = 1414$ and 4000, $M = 1000$, $We = 20$ and at $t^* = 1.52$ obtained from a 3D and 2D axisymmetric simulation showing a good match between 2D and 3D results.

<table>
<thead>
<tr>
<th>$Re$</th>
<th>$M = 100$</th>
<th>$M = 1000$</th>
</tr>
</thead>
<tbody>
<tr>
<td>4000</td>
<td>Bag-Stamen (1.196)</td>
<td>Bag-Stamen (0.8729)</td>
</tr>
<tr>
<td>1414</td>
<td>Bag-Stamen (1.084)</td>
<td>Bag-Stamen (1.092)</td>
</tr>
<tr>
<td>500</td>
<td>Bag (0.71004)</td>
<td>Bag (0.7014)</td>
</tr>
<tr>
<td>141</td>
<td>Bag (0.4548)</td>
<td>Bag (0.057)</td>
</tr>
</tbody>
</table>

Table 4: Breakup modes at different $Re$ values for a drops with $\rho^* = 1000$ at $We = 20$ and $M = 100$ and 1000. Values in the bracket denotes the $N_{RT}$ values.

The drop for $\rho^* = 1000$ at $We = 20$, $M = 100$ and 1000 and at different $Re$ values. It can be seen that with an increase in $Re$ value for the same viscosity ratio $M$, the drop deformation and hence the breakup mode is changing, effectively altering the breakup transition value of $We$, that is with an increase in $Re$ the breakup transition value of $We$ is decreasing.

Non-dimensional RT wave number, $N_{RT}$, listed in Table 4 also shows an evident increase with increase in $Re$ value, thus reinforcing the argument that the RT-instability is the mechanism of the breakup of drops for high $\rho^*$ values. For different $M$ values, the value of $Re$ for which the the breakup mode changes is also different. Therefore, Ohnesorge number, $Oh$, which varies with $M$ and $Re$ values when everything else is kept constant, is a better parameter to represent this behavior.

In terms of Ohnesorge number, a decrease in $Oh$, shifts the transition $We$ to lower values. We note that, although the values of $Oh$ are less than 0.1 (a value below which it is considered that there are no effects of viscous forces on the breakup mechanism), changing the value of $Oh$ still affects the breakup of drops but not to the extent for the values of $Oh > 0.1$. This possibly explains the discrepancy in breakup transitional value of $We$ observed by different authors in their experimental results (Table 1).
Table 5: Breakup shapes at different Re values for the drops with $\rho^* = 1000$ at We = 20 and $M = 100$ and 1000 along with the time $t^*$ beside them.

<table>
<thead>
<tr>
<th>Re</th>
<th>$M = 100$</th>
<th>$M = 1000$</th>
</tr>
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<tbody>
<tr>
<td>4000</td>
<td>![Drop Image]</td>
<td>![Drop Image]</td>
</tr>
<tr>
<td>1414</td>
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<td>![Drop Image]</td>
</tr>
<tr>
<td>500</td>
<td>![Drop Image]</td>
<td>![Drop Image]</td>
</tr>
<tr>
<td>141</td>
<td>![Drop Image]</td>
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</tbody>
</table>

### 3.5 Rim dynamics

For the drops with high $\rho^*$, flow around the drop has relatively low effect on the drop deformation, morphology and the breakup. Hence the drops deform into a flat disc and further the RT-instability governs the breakup as already explained in Sections 3.3 and 3.4. But for the drops of lower $\rho^*$, the flow field has a greater impact on the drop morphology, deformation and the breakup due to the higher velocities induced in the drops by the surrounding gas flow, as explained in the Section 3.2, and the flow patterns around the rim guide the direction of alignment of the rim and hence we conclude that “rim dynamics” govern the breakup.

Figures 16 and 17 show the 2D axisymmetric and 3D simulations of the drop for $\rho^* = 10$, $M = 100$ at We = 20, $Re_{\rho} = 4000$, $t^* = 2.02$ and at $t^* = 2.53$. Interestingly, turbulent vortex shedding is not seen in the wake of the drop even in the 3D simulations, unlike the drops with high $\rho^*$ values. Instead, undisturbed vortex rings are seen in both 3D and 2D axisymmetric simulations. Since a 2D axisymmetric simulation predicts drop shape as well as the flow around the drop accurately, the initial phase of the deformation of the drop during the secondary breakup for low $\rho^*$ values can indeed be considered axisymmetric.

The primary vortex ring, seen in Figures 16a and 17a, is formed on the leeward surface of the drop due to the separation of an initially attached boundary layer on the surface of the drop. This vortex ring pulls the rim of the drop along, resulting in the formation of a backward-bag. Eventually this vortex ring pinches-off from the boundary layer and moves downstream with respect to the drop and a pair of secondary counter-rotating vortex rings are formed at the rim (Figures 16a and 17a) due to the flow separation behind the rim. This counter-rotating vortex rings induce opposite directional velocity in the rim of the drop and deflects it more towards the upstream direction, eventually turning the drop into a forward-bag.

At later times, this primary vortex ring becomes strongly asymmetrical and eventually sheds
Figure 16: Comparison of flow field around the drop for $\rho^* = 10$ at $Re_g = 4000$, $M = 100$, $We = 20$ and at $t^* = 2.02$ obtained from a 3D and 2D axisymmetric simulation.
Figure 17: Comparison of flow field around the drop for $\rho^* = 10$ at $Re_g = 4000$, $M = 100$, $We = 20$ and at $t^* = 2.53$ obtained from a 3D and 2D axisymmetric simulation.
vortex at the vortex formation time of $t/t_{cv} = 4$ (as discussed earlier in Section 3.4). For example, $(U_g - u_{drop})/u_g = 0.45$ for the case shown in Figure 16 and hence the asymmetric vortex formation is expected to occur at $t^* \sim 3.04$. But according to our observations, based on our simulations, vortex ring never reaches an asymmetrical state because the relative velocity between the drop and the gas, $U_g - u_{drop}$, which feeds into the circulation of the vortex ring reduces substantially with time, and hence the critical circulation required for asymmetry in the vortex rings is never achieved for the drops with low $\rho^*$ values. In contrast, for high density ratios $U_g - u_{drop}$ reduces relatively slowly with time and hence the critical circulation is quickly achieved at very early stages of the deformation of the drop that leads to asymmetric vortex rings and eventually a turbulent wake is observed.

4 Summary and Conclusions

In the present study, we performed fully resolved numerical simulations of a drop in a high speed gas flow to investigate the effect of density ratio and Reynolds number on the secondary breakup of the drops. These simulations were performed for a moderate Weber number range (20-120), where bag breakup, multi-mode and sheet-thinning breakup modes have been observed in experiments.

Previous studies reported conflicting views on the effect of density ratio on the breakup modes and drop morphology (Aalburg et al., 2003; Kékesi et al., 2014; Yang et al., 2016). To resolve these discrepancies, we performed a large set of simulations with different values of $\rho^*$ from 10 to 1000, and $We$ from 20 to 120. Further, we vary $Re$ and $M$ independently to delineate their effects on drop morphology. As discussed in previous sections, in this study we primarily present axisymmetric simulations along with a few 3D simulations to support the axisymmetric approximations. We would like to emphasize here that although the final fragmentation of the drop is a 3D phenomenon, the initial deformation of the drop into the breakup modes, bag, bag-with-stamen and shear-stripping mode is essentially an axisymmetric phenomenon.

In what follows, we present the important conclusions from this study.

1. For high $\rho^*$ values, drops deform into a flat disc, whereas for low $\rho^*$ values the drops do not deform into a flat disc at all, instead they bend towards the downstream direction and for intermediate $\rho^*$ values, there is a gradual variation in the bend.

2. Axial and radial components of the velocities at the center of the drop, $u_{center}$, and at the rim, $u_{rim}$, decrease with an increase in $\rho^*$ values, which also follows from the scaling relation based on momentum transfer $U_l \sim \sqrt{1/\rho^*} U_g$. Further, the difference in axial velocity, $(u_{center} - u_{rim})$, decreases with an increase in the $\rho^*$ values, which explains the higher stretching and bending of the drops for low $\rho^*$ values.

3. Displacement of the drops decreases with increase in $\rho^*$ values. This is due to the differences in the centroid velocity of the drop, $u_{drop}$, and the momentum transferred to the leeward side of the drops, which depends on the kinematic viscosity of the drop, $\nu$.

4. Breakup time, $t_b^*$, and the distance travelled in the streamwise direction, $x_l/d_0$, at the onset of breakup are higher for the drops with $\rho^* = 10$ than for the drops with $\rho^* = 50 - 1000$ and they decrease with an increase in $We$, whereas relative velocity, $u_r = (u_g - u_l)/u_g$ has a continuous variation from $\rho^* = 50 - 1000$ and for high density ratios the values are in good agreement with the experimental observations of Dai and Faeth (2001); Zhao et al. (2010).

5. Drops for $\rho^* = 10$ at $We = 20$ and 40 and for $\rho^* = 50$ at $We = 20$ do not breakup at all, which is also in agreement with the simulations of Han and Tryggvason (2001). We explain
this using the instantaneous Weber number, \( We_{\text{inst}} = \rho g (U_g - u_{\text{drop}})^2 d_0 / \sigma \). The values of \( We_{\text{inst}} \) for these conditions are less than \( We_{\text{crit}} \) at the onset of breakup or the time where there could have been a pinch-off at the thinnest section of the drop, whereas \( We_{\text{inst}} > We_{\text{crit}} \) at the onset of breakup for all other cases where breakup is observed.

6. Interesting drop shapes are observed with varying \( \rho^* \) values keeping \( We \) constant. At \( We = 20 \), a forward-bag is seen at \( \rho^* = 10 \), transient canopy-top shape for \( \rho^* = 50 \) and 100 and forward-bag with stamen for \( \rho^* \geq 150 \). The size of the stamen decreases with increase in \( \rho^* \) value. At \( We = 40 \) and higher, backward-bag is seen for \( \rho^* = 10 \), whiplash with sheet-thinning for \( \rho^* = 50 \) and sheet-thinning for \( \rho^* \geq 100 \). The rim is thicker for low \( \rho^* \) values, which could be due to higher Taylor-Culick velocity for low \( \rho^* \) values.

7. In addition to the differences in deformation, breakup morphology and breakup modes, the breakup mechanism is also different for higher and lower \( \rho^* \) values. At higher \( \rho^* \) values, the formation of bag is due to the Rayleigh-Taylor instability. The non-dimensional RT wavenumber, \( N_{\text{RT}} \) in the cross-stream direction of the drop agrees very well with the range proposed by Zhao et al. (2010). Numerically calculated non-dimensional growth rates, \( \omega^* \), on the other hand are not in direct quantitative agreement with the theoretical estimates for the instability on a planar surface; but they are proportional to the theoretical growth rate, implying that there could be “end-effects” associated with the growth of RT wave.

8. Study of the flow around the drop, in 3D simulations, shows that the vortex ring formed due to flow separation in the wake region of the drop, develops asymmetries and sheds leading to the formation of turbulent wake region and the time taken to form these asymmetries agrees very well with the vortex formation time scales. However, the vortex ring observed in 2D axisymmetric simulations is stable and never develops any asymmetries. Nevertheless, the drop shapes are same in 2D and 3D cases, implying that the flow has only a weak effect on the drop shape for higher \( \rho^* \) values, which is in agreement with the experimental observations of Flock et al. (2012).

9. Increasing gas Reynolds number, \( Re \) alters the breakup and we see that the drop breakup mode transitions from a bag to bag-with-stamen for an increase of \( Re \) from 1414 to 4000 at viscosity ratio \( M = 1000 \). Non-dimensional RT wavenumber, \( N_{\text{RT}} \), also increases with an increase in \( Re \), and the \( N_{\text{RT}} \) values again conform to the range given by Zhao et al. (2010), indicating that the RT-instability is indeed the breakup mechanism at higher \( \rho^* \) values. The effect of \( M \) is that it shifts the value of transition value of \( Re \) (critical \( Re \) across which there is a change in the breakup mode for a given \( We \)). Hence, combining the effect of \( M \) and \( Re \), Ohnesorge number, \( Oh \), is a better parameter to represent this behavior. This explains the discrepancy in breakup transitional values of \( We \) observed by different authors in their experiments (Table 1).

10. For lower \( \rho^* \) values, breakup is governed by the dynamics of the rim. Flow around the drop has a greater impact (in comparison to high \( \rho^* \) drops) on the drop shape, deformation and breakup due to higher velocity induced in the drop by the surrounding gas flow. For low \( \rho^* \), unlike for higher \( \rho^* \) values, the vortex ring formed at the rim never develops asymmetries until the breakup of the drop.

To conclude, the drops for \( \rho^* < 150 \) behave differently from \( \rho^* \geq 150 \) at the same \( We \), making “Density ratio” an important parameter in characterizing the secondary breakup of drops and also in the study of liquid jets in gas crossflow. The present study, describes the differences in the behavior of
the drops for different $\rho^*$ values, such as in the air-water system in atmospheric conditions, where the $\rho^* \sim 1000$, and in high pressure applications where $\rho^*$ can be $\sim 100$, and also in the manufacturing of pellets by quenching molten metal in a pool of cold water where $\rho^* \sim 1 - 10$.

In the present work, we have essentially focused on the drop deformation and breakup modes in the moderate Weber number regime. Based on our results, we believe that a similar systematic study on vibrational mode of breakup as well as high Weber number breakup would reveal interesting effects of the density ratio on the deformation and breakup mechanisms.

**Supplementary material**

The data and videos from the simulations can be accessed from the following link: goo.gl/7qVEKF

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**Declarations of interest**

None.

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**Appendix A: Relation between non-dimensional growth rate and instantaneous Bond number**

Non dimensional growth rate is given by,

$$\omega^* = \omega \frac{d\sqrt{\rho^*}}{U_g}$$  \hspace{1cm} (11)

From equation 10,

$$\omega^* = \sqrt{ka \left( \frac{\rho^* - 1}{\rho^* + 1} \right) \frac{d\sqrt{\rho^*}}{U_g}}$$  \hspace{1cm} (12)

Substituting for wavenumber, $k = \sqrt{\rho o a/3\sigma}$ and rearranging,

$$\omega^* = \frac{1}{3^{\frac{1}{4}}} \frac{At^\frac{1}{2} Bo^\frac{3}{4}}{\sqrt{We}}$$  \hspace{1cm} (13)

Here $At$ is the Atwood number. Hence the $\omega^*$ scales as $Bo^\frac{3}{4}$ for a given $\rho^*$ and $We$. 


References


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