Gas bubble formation from a submerged orifice under constant-flow conditions in a quiescent high-density liquid metal, lead–bismuth eutectic (LBE), at high Reynolds numbers was investigated numerically. The numerical simulation was carried out using a coupled level-set and volume-of-fluid method governed by axisymmetric Navier–Stokes equations. The ratio of liquid density to gas density for the system of interest was about 15,261. The bubble formation regimes varied from quasi-static to inertia-dominated and the different bubbling regimes such as period-1 and period-2 with pairing and coalescence were described. The volume of the detached bubble was evaluated for various Weber numbers, \( \text{We} \), at a given Bond number, \( \text{Bo} \), with Reynolds number \( \text{Re} \). It was found that at high values of the Weber number, the computed detached bubble volumes approached a \( 3/5 \) power law. The different bubbling regimes were identified quantitatively from the time evolution of the growing bubble volume at the orifice. It was shown that the growing volume of two consecutive bubbles in the period-2 bubbling regime was not the same whereas it was the same for the period-1 bubbling regime. The influence of grid resolution on the transition from period-1 to period-2 with pairing and coalescence bubbling regimes was investigated. It was observed that the transition is extremely sensitive to the grid size. The transition of period-1 and period-2 with pairing and coalescence is shown on a Weber–Bond numbers map. The critical value of Weber number signalling the transition from period-1 to period-2 with pairing and coalescence decreases with Bond number as \( \text{We} \), which is shown to be consistent with the scaling arguments. Furthermore, comparisons of the dynamics of bubble formation and bubble coalescence in LBE and water systems are discussed. It was found that in a high Reynolds number bubble formation regime, a difference exists in the transition from period-1 to period-2 with pairing and coalescence between the bubbles formed in water and the bubbles formed in LBE. © 2015 American Institute of Chemical Engineers

Keywords: numerical simulation, bubble formation, liquid metal, bubbling regimes, bubble volume, transition

Introduction

The growth and detachment of bubbles by injection of a gas through a submerged orifice in a quiescent liquid are of great interest for diverse applications in chemical, metallurgical, and biomedical systems. Applications of gas bubble generation in high-density liquid metals include the operation of liquid metal circulation in metallurgical and nuclear power systems. In nuclear power systems, the liquid metal lead–bismuth eutectic (LBE) is preferred as the target system taking into account the high production rate of neutrons, effective self-circulating heat removal, and minimal radiation damage.

Few studies have been reported on the problem of bubble dynamics in liquid metals. Some experimental studies have focused on bubble formation and bubble rising in liquid metals. Experimental studies have been carried out concerning the effect of gas injection through submerged orifices or small holes in a liquid metal, because a liquid metal is not a transparent liquid and there are very few measuring techniques in experimental physics to evaluate the bubble diameters, bubble velocity, and bubble shape compared with transparent liquids, especially water. Considering these difficulties, numerical studies on the dynamics of bubble formation from an orifice in
a quiescent high-density liquid metal were undertaken in this work.

Many investigations of the basic case of bubble formation at a single orifice were reviewed by Kumar and Kuloor,2, Clift et al.,3 Kulkarni and Joshi,4 Tsuge,5 and Sadhal et al.6 Different experimental studies7–9 and theoretical approaches10–18 were applied to study the problem of bubble formation at a single orifice or a single nozzle. In most of the theoretical models17,23,32,33,38 developed by these workers, the effect of the previously formed bubble on the growth of the bubble at the orifice were not taken into account, although Zhang and Shoji,21 Chuang and Goldschmidt,39 and Deshpande et al.40 developed models by taking into account the wake effect of the previous bubble or bubble coalescence. Depending on the operating parameters (gas flow rate, orifice diameter, surface wetting characteristics, and physical properties of the liquid phase), different regimes19–21,23,36,27,41 are discerned as period-1, period-2 with pairing or double coalescence (see below), period-3 (triple coalescence), period-4 (quadruple coalescence), and finally chaotic bubbling regimes.32 Most importantly, Higuera and Medina33 reported that the period-2 regime consists of the pairing regime20,21,23 and the double coalescence regime.20,21,23 The pairing regime is characterized when two already detached bubbles rise as a pair and coalesce far away or near the orifice (detachment–detachment coalescence).44,35 The double coalescence regime is defined when one already detached bubble coalesces with the growing bubble at the orifice (detachment–coalescence–detachment). In this article, the behavior of the pairing regime is henceforward referred to as period-2 with pairing and coalescence bubbling regime.

Conversely, the dynamics of the growth and detachment of drops formed from a capillary tube in air have been extensively studied experimentally and numerically by Zhang and Basaran,43 Wilkies et al.,44 Yildirim et al.,45 Amb ravaneswaran et al.,46–48 Subramani et al.,49 and others. The period-1 dripping regime with or without satellite drop formation,43–46 period-1 to complex dripping47–49 (period doubling bifurcations, chaos, and hysteresis) to period-58,50 and the transitions among different regimes have been discussed in detail.47–51

In addition to the experimental and theoretical investigations, numerical simulations of bubble formation processes have been carried out using the boundary integral method (BIM),23,35,52 volume-of-fluid method,53 level-set method,54 coupled level-set and volume-of-fluid (CLSVOF) method,55–60 and front tracking method.61 Oguz and Prosperetti,32 Higuera and Medina,33 and Gordillo et al.52 studied the bubble formation process from an orifice in a quiescent inviscid liquid at high Reynolds numbers by means of the potential flow approximation. Oguz and Prosperetti32 did not discuss interactions between bubbles at high gas flow rates. Higuera and Medina33 found a critical Weber number (high gas flow rates) for a given Bond number above which pairs of bubbles coalesce either in the vicinity of or far away from the orifice. However, the map of the transition between period-1 and period-2 bubbling regimes with a wide range of Bond and Weber numbers was not clearly shown. Gordillo et al.52 reported numerical results of the characteristics of the minimum neck radius of the bubbles with time at high Reynolds numbers. Gerlach et al.,55 Buwa et al.,56 and Chakraborty et al.37 used the CLSVOF method to evaluate the bubble formation process at a single submerged orifice under low and high gas flow rates. In their work, the effects of the orifice diameter,55 gravity,37 and gas flow rates on the period-1 and period-2 bubbling regimes and their transition characteristics55,56 were reported. To the best of the authors’ knowledge, there has been no numerical study to investigate the dynamics of bubble formation and rising in a high-density liquid metal as one of the fluids. The numerical investigation of bubble generation in a quiescent liquid with a high density ratio of liquid to gas, such as above 10,000, is a challenging task. For the design of efficient two-phase flow systems,55–57 a detailed knowledge of the bubble shapes and different bubbling regimes is of fundamental importance.

The objective of this article is an ab initio numerical analysis of the effects of Weber and Bond numbers on the global characteristics of the bubble formation process at high Reynolds numbers, such as the frequency of the formation and the volume of bubbles created via injection of a gas at constant flow rate through a single orifice in quiescent high-density liquids. The combined level-set and volume-of-fluid (CLSVOF) method57,58,62–64 was used to perform the numerical simulations. First, we validate our computational results with the benchmark scaling laws from very low to high flow-rate regimes of bubble formation. Then, we extend previous studies32,35,56–58 to include time periodic bubbling regimes by means of qualitative and quantitative analyses, the interaction and coalescence behavior of bubbles in period-2 with pairing and coalescence bubbling regime, the transition between period-1 and period-2 with pairing and coalescence bubbling regimes characterized by a map that depends on We and Bo and a suitable scaling analysis for the transition. The analysis of other regimes (period-2 with double coalescence regime, period-3, period-4, and chaotic bubbling regimes) of bubble formation at sufficiently high gas flow rates was not investigated. It was observed that in period-2 with double coalescence regime, the coalesced bubble after departing from the orifice breaks up into toroidal bubbles due to strong liquid jet generation at the rear part of the coalesced bubble.35 The axisymmetric solutions of the formation of toroidal bubbles are not realistic.35 The three-dimensional (3-D) effect of the liquid wake or turbulent eddies stabilizes the shape and weakens the liquid jet65 and no toroidal bubbles are formed. 3-D fluid flow was not taken into account in the present simulations. Therefore, numerical simulations using axisymmetric governing equations to study the problems of period-2 regime with double coalescence to the chaotic regime were not carried out.

In the present work, nitrogen and liquid LBE at atmospheric pressure and 220°C served as reference working fluids. Their physical properties are listed in Table 1. The effect of the nitrogen gas flow rate Q through the orifice in the range 0.5 m L/min \(\leq Q \leq 140\) mL/min and the effect of the orifice radius \(R_o\) in the range 0.72 mm \(\leq R_o \leq 3\) mm were studied. In this study, the five dimensionless parameters Weber number \((We = \rho g Q^2/\pi \sigma R_o^3)\), Bond number \((Bo = \rho g R_o^2/\sigma)\), Reynolds number \((Re = \rho g Q/\pi \rho \rho g R_o)\), density ratio of gas to liquid \((\eta = \rho_g/\rho_l)\), and viscosity ratio of gas to liquid \((\lambda = \mu_g/\mu_l)\) were provided for computations, where \(\sigma\) is the surface tension, \(\rho_l\) is the liquid density, \(\rho_g\) is the gas density, \(\mu_l\) is the liquid viscosity, and \(\mu_g\) is the gas viscosity. Quantitative and qualitative results were obtained by varying one dimensionless parameter while keeping other parameters fixed. We carried out investigations for \(4.31 \times 10^{-4} \leq We \leq 24.81, 0.13 \leq Bo \leq 2.28,\) and \(Re \gg 1\). The other two parameters were \(\eta = 6.55 \times 10^{-5}\) [the ratio of the liquid (LBE) density to the gas density is about 15.261] and \(\lambda = 1.12 \times 10^{-2}\). The orifice plate material was assumed to be a hydrophilic surface. In this
problem, the static contact angle was taken as $\phi_s = 0^\circ$, indicating that the bubble base is pinned to the orifice rim.

**Formulation of the Problem**

**Problem description**

The growth and detachment of gas bubbles at a single submerged orifice immersed in a quiescent liquid are discussed. When the gas flows through the orifice, the bubble volume increases continuously. According to classical Refs. 32 and 36, there are two stages of bubble formation, namely the expansion stage and the collapse stage. In the expansion stage, the bubble mainly grows in the radial direction and the detaching (upward direction) forces are weak compared with the attaching forces (downward direction). In the collapse stage, the bubble mainly grows in the axial direction (upward direction) and the detaching forces dominate over the attaching forces. In this stage, the bubble is attached to the orifice mouth through a neck forming in the vicinity of the orifice. At the end of this stage, the bubble is detached from the orifice and moves upwards through the liquid. In the present work, complete numerical simulation of the processes of bubble formation under constant-flow conditions$^{37,58}$ was performed using a cylindrical coordinate system $(r, z, \theta)$ as shown in Figure 1. The origin of the $(r, z, \theta)$ coordinate system is placed at the center of the orifice rim, where $r$ is the radial coordinate, $z$ is the axial coordinate measured in the opposite direction of gravity $g$, and $\theta$ is the azimuthal coordinate. In this article, the dynamics are assumed to be axisymmetric and the problem is independent of the azimuthal coordinate $\theta$. A gas is injected at a constant flow rate $Q$ through a single orifice of radius $R_o$, at the horizontal hydrophilic bottom wall (wetting surface) submerged in a quiescent Newtonian liquid. We assume the ambient liquid and the gas filling the bubble to be incompressible with isothermal conditions. In the following, we discuss the different forces that act during the growth of the bubble at the orifice in a quiescent liquid.

At very low gas flow rates, the buoyancy (upward direction) and surface tension (downward direction) forces balance each other at the moment of bubble detachment$^{56}$ and the detached bubble volume $V_F$ is determined by

$$2\pi R_o \sigma = V_F (\rho_g - \rho_l) g$$

where $V_F$ is the Fritz volume$^{42,66}$ of a detached bubble which is assumed to be a sphere. This balance gives $V_F = 2\pi R_o \sigma / (\rho_g - \rho_l) g$ or $V_F = V_F/R_o^3 = 2\pi/Bo$ in dimensionless variables. The influence of $\rho_l$ is neglected, since $\rho_l \ll \rho_g$ here. This regime of bubble formation is known as the quasi-static bubble formation regime$^{32,36,37}$ in which the bubble volume is independent of gas flow rate.

An inertia-dominated region of periodic bubbling exists at high gas flow rates.$^{17,18,32,35}$ In this high flow rate region, the bubble is detached when the buoyancy force has to overcome the force due to liquid inertia and the bubble volume can be estimated by an order of magnitude balance

$$\rho_l V_b g \sim \rho_g (Q/V_b^{2/3})^2 V_b^{2/3}$$

which yields $V_b \sim Q^{6/5}/g^{3/5}$ or, in dimensionless terms, $V_b R_o^3 \sim (We/Bo)^{3/5}$. The transition from the quasi-static region to the liquid inertia region can be characterized when the surface tension, buoyancy, and liquid inertia forces are all of the same order

$$\sigma R_o \sim \rho_l V_b g \sim \rho_g (Q/V_b^{2/3})^2 V_b^{2/3}$$

from which the critical gas flow rate$^{32}$ is defined by $Q_{cr} = \pi (16/3)^{1/6} (\sigma R_o / \rho_l)^{3/6}$, which can be written in dimensionless form as $We_{cr} = (16/3)^{1/3} Bo^{2/3}$. It can be found from Eqs. 1–3 that $V_b V_F \sim (We/We_{cr})^{3/5}$, which is independent of Bond number. According to Ref. 32, $V_b V_F$ is constant when $Q \ll Q_{cr}$, $(We \ll We_{cr})$ and increases with $(Q/We_{cr})^{3/5}$ when $Q \gg Q_{cr}$, $(We \gg We_{cr})$.

At the onset of the high-flow rate region, where the inertia of the liquid starts to influence the dynamics of the growing bubble at the orifice, the Reynolds number is of the order of $Re_{cr} = (\rho_l R_o \sigma / \mu_l^2)^{1/2}$ (obtained from Eq. 3). The Reynolds number is $Re_{cr} \sim 900$ for liquid LBE when the orifice radius $R_o = 1 mm$, and higher Reynolds numbers are obtained for higher gas flow rates. This shows that in the present study, the effect of viscous force is very small during the process of bubble formation. However, the influence of liquid viscosity becomes important at the collapse stage of axisymmetric bubble collapse in a quiescent liquid, as reported by Thoroddsen et al.$^{56}$ Gordillo$^{67}$ and many others$^{9,21}$ in the case of pinch-off of liquid drops immersed in a fluid of negligible density when the minimum radius of the neck $R_{min}$ becomes sufficiently small. Gordillo$^{67}$ found that viscous effects would not come into play in liquids with viscosities below 4.2 cP when $R_{min} > 10 \mu m$. In our present computations, it was found that the bubble pinched off when the minimum radius of the neck was $R_{min} = 110 \mu m$, as also suggested elsewhere.$^{60}$ Therefore, a very fine grid and very small time step are needed to capture the final instants prior to pinch-off. However, our computations are unable to resolve it owing to limitations of computational resources. Therefore, in LBE and water, the bubble pinch-off process also can be treated as an inviscid flow.

**Table 1. Physical Properties of the Fluids for a Nitrogen–Liquid LBE System at 220°C**

<table>
<thead>
<tr>
<th>Property</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\rho_l$</td>
<td>10.443 4652 kg/m$^3$</td>
</tr>
<tr>
<td>$\rho_g$</td>
<td>0.648444 kg/m$^3$</td>
</tr>
<tr>
<td>$\mu_l$</td>
<td>$2.28 \times 10^{-3}$ Pa s</td>
</tr>
<tr>
<td>$\mu_g$</td>
<td>$2.55 \times 10^{-3}$ Pa s</td>
</tr>
<tr>
<td>$\sigma$</td>
<td>$404.462 \times 10^{-3}$ N/m</td>
</tr>
</tbody>
</table>

Figure 1. Schematic of bubble formation from an orifice and computational domain.
Governing equations and numerical method

In this study, the two-phase flow system consists of a gas and a liquid which are immiscible and incompressible fluids. It is assumed that the flow in each phase is axisymmetric. The problem is governed by the single set of continuity and Navier–Stokes equations for treating a single fluid continuum. The continuity equation is given in cylindrical axisymmetric coordinates \((r, z)\) as

\[
\frac{1}{r} \frac{\partial(ru)}{\partial r} + \frac{\partial v}{\partial z} = 0
\]  
(4)

where \((u, v)\) are radial and axial components of the velocity field, respectively. The momentum equations are as follows:

\[
\rho(\ddot{z}) \left( \frac{\partial u}{\partial t} + u \frac{\partial u}{\partial r} + v \frac{\partial u}{\partial z} + \frac{\partial p}{\partial r} + S_{rr} \right) + \frac{\partial S_{zr}}{\partial r} = \frac{\partial \tau_{rz}}{\partial z} + S_z
\]

\[
\frac{\rho(\ddot{z}) g_r + \sigma(\phi) \frac{\partial \phi}{\partial r}}{r} \]

\[
\rho(\ddot{z}) \left( \frac{\partial v}{\partial t} + u \frac{\partial v}{\partial r} + v \frac{\partial v}{\partial z} + \frac{\partial p}{\partial z} + S_{zz} \right) + \frac{\partial S_{rz}}{\partial z} = \frac{\partial \tau_{zr}}{\partial z} + S_r
\]

\[\frac{\rho(\ddot{z}) g_z + \sigma(\phi) \frac{\partial \phi}{\partial z}}{r} \]

where \(\rho\) is the pressure, \(g_r(=0)\), and \(g_z(= - g)\) are the radial and axial components of the gravitational acceleration, respectively, and \(S_{rr}, S_{zz},\) and \(S_{zr}\) are the components of Newtonian stress tensor, which are expressed as

\[
S_{rr} = 2 \mu(\ddot{z}) \frac{\partial u}{\partial r}, \quad S_{zz} = 2 \mu(\ddot{z}) \frac{\partial v}{\partial z}, \quad S_{zr} = S_{rz}
\]

\[
\mu(\ddot{z}) \left( \frac{\partial u}{\partial r} + \frac{\partial v}{\partial z} \right)
\]  
(7)

The influence of surface tension \(\sigma\) is incorporated into the momentum equations following the continuum surface force model of Brackbill et al.\(^{72}\) Here, \(\ddot{z}\) is the smoothed void fraction field, which is defined using a Heaviside function,\(^{62}\) \(H(\phi)\), as

\[
\ddot{z} = H_\epsilon(\phi)
\]

\[
\begin{align*}
1 & \quad \phi > \epsilon \\
\frac{1}{2} + \frac{\phi}{2\epsilon} + \frac{1}{2\pi} \sin \left( \frac{\pi \phi}{\epsilon} \right) & \quad |\phi| \leq \epsilon \\
0 & \quad \phi < -\epsilon
\end{align*}
\]  
(8)

where \(\phi\) is the level set function, which is the signed distance function from the interface. \(\phi\) is zero at the interface and has positive values in the liquid region and negative values in the gas region. Here, \(2\epsilon\) is the interface thickness over which the fluid properties are interpolated. The present simulations were performed using \(\epsilon = 1.5\Delta r\), where \(\Delta r\) is the size of the computational cell. The smoothed density \(\rho(\ddot{z})\) and viscosity \(\mu(\ddot{z})\) can be expressed by a Heaviside function \(H_\epsilon(\phi)\) as

\[
\rho(\ddot{z}) = \rho_l \ddot{z} + \rho_g (1 - \ddot{z})
\]

\[\mu(\ddot{z}) = \mu_l \ddot{z} + \mu_g (1 - \ddot{z})
\]  
(9)

where subscripts \(l\) and \(g\) represent liquid and gas, respectively. The density and viscosity of the liquid and gas are assumed to be constant. The local curvature \(\kappa\) at the gas–liquid interface is computed as

\[
\kappa = -\nabla \cdot \hat{n}
\]

(11)

where the unit normal vector \(\hat{n}\) is derived from level set function \(\phi\)

\[
\hat{n} = \frac{\nabla \phi}{|\nabla \phi|}
\]  
(12)

and it is directed into the liquid region. In the CLSVOF method, the advection equations for the volume fraction \(\phi\) and the level set function \(\phi\) are

\[
\frac{\partial \phi}{\partial t} + \frac{1}{r} \frac{\partial (ru \phi)}{\partial r} + \frac{\partial (v \phi)}{\partial z} = 0
\]

\[
\frac{\partial \phi}{\partial t} + \frac{1}{r} \frac{\partial (ru \phi)}{\partial r} + \frac{\partial (v \phi)}{\partial z} = 0
\]  
(13)

The governing equations are nondimensionalized by introducing nondimensional variables using \(r^* = r/\bar{r}, \quad z^* = z/\bar{z}\), \(\kappa^* = \kappa \bar{r}, \quad u^* = \frac{u}{\bar{u}}, \quad v^* = \frac{v}{\bar{v}}, \quad \rho^* = \frac{\rho}{\bar{\rho}}, \quad t^* = \frac{t}{\bar{t}}\)

\[
\rho^*(\ddot{z}) = \frac{\rho(\ddot{z})}{\rho_l}, \quad \mu^*(\ddot{z}) = \frac{\mu(\ddot{z})}{\mu_l}
\]

where \(\bar{r}\) is the characteristic length and average inflow velocity of gas at the orifice inlet \(v_{d,avg}(= Q/\pi \bar{R}_o^2)\) is the characteristic velocity. The resulting governing equations\(^{68}\) are as follows:

Continuity equation:

\[
\nabla \cdot \bar{V} = 0
\]  
(15)

Momentum equations

\[
\rho^*(\ddot{z}) \left( \frac{\partial \bar{V}^+}{\partial t} + \nabla \left( \bar{V} \bar{V}^+ \right) \right)
\]

\[
= -\nabla p^* + \rho^*(\ddot{z}) \left( \frac{B_0}{We} \nabla \left[ \mu^*(\ddot{z}) \left( \nabla \bar{V}^+ + (\nabla \bar{V}^+)^T \right) \right] \right)
\]

\[+ \left( \frac{1}{We} \right) \kappa^*(\phi) \nabla \ddot{z}
\]

(16)

where \(\bar{V} = (u, v)\) is the velocity vector. In Eq. 16, the density and viscosity of the single fluid continuum are computed by

\[
\rho^*(\ddot{z}) = \bar{\rho} \ddot{z} + \bar{\rho}_l (1 - \ddot{z})
\]

\[\mu^*(\ddot{z}) = \bar{\mu} \ddot{z} + \bar{\mu}_l (1 - \ddot{z})
\]  
(17)

Additionally, we shall provide two other parameters, the Froude number \(Fr = \frac{v_{d,avg}^2}{g \bar{R}_o} \left( = \frac{\bar{V}}{\bar{V}_{avg}} \right)\) and the Ohnesorge number \(Oh = \frac{\mu v_{d,avg}}{\rho g \bar{R}_o}\). In this study, the range of the Ohnesorge number is 0.0006 < Oh < 0.0013 and the range of the Reynolds number is 10 < Re < 4000.

The CLSVOF numerical method\(^{57,62}\) for an evolving moving interface was implemented in the present study. In the CLSVOF method, the level set function\(^{75}\) \(\phi\) is used to capture the interface and volume of the fluid fraction\(^{74}\) \(\bar{z}\) is used to conserve mass due to the moving interface. The governing equations are discretized using the finite-difference method on an axisymmetric coordinate and equidistant grid \(\Delta r = \Delta z\) in the radial and axial directions. The MAC algorithm\(^{75}\) is employed to solve the single set of governing equations on a
staggered grid with scalars \((p, \phi, \text{ and } z)\) located at the cell centers and velocity components \((V)\) at the center of the cell faces. The convection and the viscous terms are discretized by a second-order ENO method and central differencing, respectively. In the present work, the time stepping procedure is based on the explicit method to maintain the stability of the solution. During the computations, time steps are chosen to satisfy CFL, capillary, viscous and gravitational time conditions (discussed in Section 2.3 in Ref. 76). The details of the numerical method have been described by Chakraborty et al.76

**Computational domain and boundary conditions**

The computational domain established for axisymmetric numerical simulations of bubble formation is shown in Figure 1. For low and high gas flow rates involved in the present problem, the width and height of the computational domain are taken as \(R = 20R_{b}\) and \(Z = 60R_{b}\), respectively, so that the bubble formation process is unaffected by the side and outflow boundaries, as also discussed elsewhere.58

As boundary conditions, symmetry or slip condition at the left or right boundaries, no-slip condition at the bottom wall and Neumann condition at the outflow boundary are imposed. At the orifice inlet, gas flows upwards with a defined parabolic profile of Poiseuille flow.57,58 At the orifice, the Reynolds number \(Re = \frac{r_o Q}{
u g H_{ref} D_o}\) based on the gas properties is deployed, where \(D_o\) is the orifice diameter. The range of \(Re\) is below of the laminar pipe flow condition (\(Re < 2300\)). To ensure incompressibility in the flow domain, a “reference” pressure \(p_{ref}\) of the liquid is applied at a particular point for all the computations. In these simulations, \(p_{ref} = 0\) is applied as the reference pressure. At the outflow boundary (outlet) the pressure is the “reference” pressure \(p_{ref}\) minus the hydrostatic pressure \(p(r, Z, t) = p_{ref} - \rho(z)gZ = -\rho(z)gZ\). At the bottom wall, Neumann condition of the pressure is applied. The bubble is initially assumed to be a hemisphere of radius equal to the orifice radius. Both the gas phase and the liquid phase are assumed initially to be quiescent and at uniform pressure.

**Grid independence test and validation**

The grid independence test is performed using \(40 \times 120, 80 \times 240, 100 \times 300\), and \(160 \times 480\) mesh grids in a computational domain, as shown in Figure 2 for the conditions \(We = 0.0727, Bo = 0.253\), and \(Re = 242.33\). The period-1 bubbling regime (see below in the Results section) can be observed under these conditions. Figure 2 shows the difference between profiles at the final instant prior to bubble pinch-off. The maximum relative difference in height of the profiles obtained is around 21.92% for \(40 \times 120\) and \(80 \times 240\) mesh grids, around 6.22% for \(80 \times 240\) and \(100 \times 300\) mesh grids and around 1.33% for \(100 \times 300\) and \(160 \times 480\) mesh grids. Figure 2 also reveals that the bubble shapes before detachment of the bubble for \(100 \times 300\) and \(160 \times 480\) mesh grids are almost identical, but very different from those for \(40 \times 120\) and \(80 \times 240\) mesh grids.

In addition to the above analysis, the volume of a detached bubble is calculated using \(40 \times 120, 80 \times 240, 100 \times 300\), and \(160 \times 480\) mesh grids with the same parameters as in Figure 2. Table 2 shows the volumes of detached bubbles for different mesh grids. The difference in detached bubble volume for \(40 \times 120\) and \(80 \times 240\) mesh grids is around 6.43%, for \(80 \times 240\) and \(100 \times 300\) mesh grids it is around 24% and for \(100 \times 300\) and \(160 \times 480\) mesh grids it is around 0.3%.

Therefore, in this study, the computations were performed on \(100 \times 300\) mesh grids to save computing time without affecting the accuracy of the numerical results. Depending on the orifice radius \(R_o\) and volumetric gas flow rate \(Q\) through the orifice, a grid resolution \(\Delta r = \Delta z = 0.2 \times 10^{-3} m\) in both the \(r\) and \(z\) directions was chosen to capture the bubble formation process accurately. The corresponding uniform dimensionless mesh size of \(\Delta r/R_o = \Delta z/R_o = 0.2\) was employed in all simulations in both directions. During our computations, the time step was chosen as \(\Delta t = 10^{-6} s\) to satisfy the Courant–Friedrichs–Lewy (CFL) and capillary time step conditions.76

The computed volume of a dimensional detached bubble for this case is \(V_b = 2.2585 \times 10^{-3} m^3\). The calculated detached bubble volumes from the analytical correlations given by Jamialahmadi et al.22 and Gaddis and Vogelpohl38 are \(V_b = 2.407 \times 10^{-3} m^3\) and \(V_b = 2.520 \times 10^{-3} m^3\), respectively. The comparison shows good agreement with the analytical correlations22,38 resulting in errors of about 6.17% and 10.38% relative to the analytical correlations of Jamialahmadi et al.22 and Gaddis and Vogelpohl,38 respectively. The discrepancy found in the bubble volume in LBE between the predicted result and the studies of Jamialahmadi et al.22 and Gaddis and Vogelpohl38 is due to the spherical bubble assumption in the latter models. The accuracy of the correlation given in Ref. 38 was assessed with the experimental results for highly viscous liquids. The volume of a detached

<table>
<thead>
<tr>
<th>Number of grid meshes</th>
<th>Detached bubble volume ((\times 10^{-3} m^3))</th>
</tr>
</thead>
<tbody>
<tr>
<td>40×120</td>
<td>2.981</td>
</tr>
<tr>
<td>80×240</td>
<td>2.801</td>
</tr>
<tr>
<td>100×300</td>
<td>2.2585</td>
</tr>
<tr>
<td>160×480</td>
<td>2.265</td>
</tr>
</tbody>
</table>

The non-dimensional numbers are \(We = 0.0727, Bo = 0.253, Re = 242.33, \eta = 6.55 \times 10^{-5}\), and \(\lambda = 0.0112\).\footnote{The period-1 bubbling regime (see below in the Results section) can be observed under these conditions. Figure 2 shows the difference between profiles at the final instant prior to bubble pinch-off. The maximum relative difference in height of the profiles obtained is around 21.92% for \(40 \times 120\) and \(80 \times 240\) mesh grids, around 6.22% for \(80 \times 240\) and \(100 \times 300\) mesh grids and around 1.33% for \(100 \times 300\) and \(160 \times 480\) mesh grids. Figure 2 also reveals that the bubble shapes before detachment of the bubble for \(100 \times 300\) and \(160 \times 480\) mesh grids are almost identical, but very different from those for \(40 \times 120\) and \(80 \times 240\) mesh grids. In addition to the above analysis, the volume of a detached bubble is calculated using \(40 \times 120, 80 \times 240, 100 \times 300\), and \(160 \times 480\) mesh grids with the same parameters as in Figure 2. Table 2 shows the volumes of detached bubbles for different mesh grids. The difference in detached bubble volume for \(40 \times 120\) and \(80 \times 240\) mesh grids is around 6.43%, for \(80 \times 240\) and \(100 \times 300\) mesh grids it is around 24% and for \(100 \times 300\) and \(160 \times 480\) mesh grids it is around 0.3%. Therefore, in this study, the computations were performed on \(100 \times 300\) mesh grids to save computing time without affecting the accuracy of the numerical results. Depending on the orifice radius \(R_o\) and volumetric gas flow rate \(Q\) through the orifice, a grid resolution \(\Delta r = \Delta z = 0.2 \times 10^{-3} m\) in both the \(r\) and \(z\) directions was chosen to capture the bubble formation process accurately. The corresponding uniform dimensionless mesh size of \(\Delta r/R_o = \Delta z/R_o = 0.2\) was employed in all simulations in both directions. During our computations, the time step was chosen as \(\Delta t = 10^{-6} s\) to satisfy the Courant–Friedrichs–Lewy (CFL) and capillary time step conditions.76 The computed volume of a dimensional detached bubble for this case is \(V_b = 2.2585 \times 10^{-3} m^3\). The calculated detached bubble volumes from the analytical correlations given by Jamialahmadi et al.22 and Gaddis and Vogelpohl38 are \(V_b = 2.407 \times 10^{-3} m^3\) and \(V_b = 2.520 \times 10^{-3} m^3\), respectively. The comparison shows good agreement with the analytical correlations22,38 resulting in errors of about 6.17% and 10.38% relative to the analytical correlations of Jamialahmadi et al.22 and Gaddis and Vogelpohl,38 respectively. The discrepancy found in the bubble volume in LBE between the predicted result and the studies of Jamialahmadi et al.22 and Gaddis and Vogelpohl38 is due to the spherical bubble assumption in the latter models. The accuracy of the correlation given in Ref. 38 was assessed with the experimental results for highly viscous liquids. The volume of a detached

![Figure 2. Effect of mesh size on bubble shape at departure for \(Q = 1.67 \times 10^{-7} m^3/s\), \(R_o = 1 \times 10^{-3} m\) in a nitrogen–liquid metal LBE system.](image-url)
bubble in water using the same geometric and flow conditions, leading to $Bo = 0.1363$, $We = 0.0391$, and $Re = 53.05$, is $V_b = 3.9 \times 10^{-8} \text{m}^3$, obtained by Gerlach et al.\textsuperscript{55} (see Table 2 in Ref. 55). It is observed that the detached bubble volume in LBE is smaller than that in water for low gas inflow rate conditions when the static contact angle $\omega_s = 0^\circ$. This is primarily due to higher Bond number leading to a higher buoyancy force compared with the surface tension force at the bottom of the growing bubble in LBE than that of water. This effect causes early detachment of the bubble in LBE. However, Irons and Guthrie,\textsuperscript{13} Bai and Thomas,\textsuperscript{14} and Gnyloskurenko and Nakamura\textsuperscript{77} found that the average bubble sizes in liquid metal are larger than those of air bubbles in water. This may be due to the lower Bond numbers used in liquid metals compared with those in water. It was also reported\textsuperscript{13,14,77} that the dynamics of the bubble formation process in a liquid metal were also influenced by the presence of nonwetting or hydrophobic materials with $\omega_s > 90^\circ$ (referred to as poor wettability). This influences the bubble base diameter along the surface material and the final detached bubble volume. Gnyloskurenko et al.\textsuperscript{28} Byakova et al.\textsuperscript{29} Lin et al.\textsuperscript{76} and Corchero et al.\textsuperscript{79} experimentally and Gerlach et al.\textsuperscript{55} numerically found that the size of the detached bubble is controlled by the orifice diameter when $0^\circ \leq \omega_s < 68^\circ$ (referred to as good wettability) and is independent of the contact angle. It was found that when $\omega_s < 68^\circ$ the bubble base coincides with the edge of the orifice during the bubble formation process. In the present work, the effect of the contact angle was not considered. Therefore, studies on the problem of the dynamics of bubble formation on hydrophobic surfaces are beyond the scope of the present work.

Unfortunately, the dynamics of bubble formation for different materials (with different static contact angles $\omega_s$), both hydrophilic and hydrophobic, in liquid metal LBE systems are lacking in the literature. To justify the accuracy of our numerical code, computations were carried out to determine the volume of detached bubbles in argon–liquid aluminum, where the liquid density and surface tension are $\rho_l = 2373 \text{kg/m}^3$ and $\sigma = 860 \times 10^{-5} \text{N/m}$, respectively, when $\omega_s = 0^\circ$. The computed results are compared with the experimental results of Gnyloskurenko and Nakamura\textsuperscript{77} and Gnyloskurenko et al.\textsuperscript{28} where the gas flow rates injected through the orifice of radius $R_o = 0.5 \times 10^{-3} \text{m}$ are $Q = 0.648 \times 10^{-5} \text{m}^3/\text{s}$ for liquid

Figure 3. (a) Time evolution of the volume of four consecutive bubbles (top) and time instant of first bubble growth and breakup (bottom) at the orifice and (b) volume of detached bubbles (top) after bubble breakup and the generation of bubbles (bottom) under the conditions $We = 0.172$, $Bo = 0.1425$, $Re = 325.09$, $\eta = 6.55 \times 10^{-5}$, and $\lambda = 0.0112$. 

AIChe Journal November 2015 Vol. 61, No. 11 Published on behalf of the AIChE DOI 10.1002/aic 4001
aluminum ($Bo = 0.0067$ and $We = 0.933$) and $Q = 0.033 \times 10^{-6} m^3/s$ for water ($Bo = 0.034$ and $We = 0.122$) under low flow rate conditions. The detached volumes obtained from present computations and experiments are $1.81 \times 10^{-3} m^3$ and $1.70 \times 10^{-7} m^3$, respectively (see Ref. 78), with an error of 6.47% in liquid aluminum. The corresponding values in the air–water system are $2.81 \times 10^{-8} m^3$ and $2.84 \times 10^{-7} m^3$, respectively (see Ref. 28), with an error of 1.06%. The discrepancy between our results and the results in Refs. 28 and 77 may be due to the flow rate conditions with time at the orifice in the latter experiments. It is observed that the volume of bubbles is larger in liquid aluminum than in water, which can be attributed to the differences in buoyancy forces in water and liquid aluminum metal systems. Gnyloskurenko et al.28 and Gnyloskurenko and Nakamura77 found that the volume of a detached bubble is independent of $\phi_s$ in the range $0 \leq \phi_s < 68^0$. Therefore, the present results are valid for the wetting or hydrophilic materials with $0 \leq \phi_s < 68^0$.

Results

In this section, we examine bubble generation and bubble dynamics in a high-density liquid metal (liquid LBE). We discuss growing and detached bubble volumes, bubble formation frequency, and the features of the period-1 and period-2 with pairing and coalescence bubbling regimes.

Calculation of detached bubble volume

In this section, the calculation of the dimensional detached bubble volume ($V_b$) is discussed under the conditions of $Q = 1.67 \times 10^{-7} m^3/s$ and $R_o = 0.75 \times 10^{-3} m$, as shown in Figure 3. The bubble volume $V_b$ is defined as

$$V_b = \int \frac{(1-\alpha)dV}{V} \tag{19}$$

where

$$dV = 2\pi r \Delta r \Delta z \tag{20}$$

$\alpha$ is the liquid volume fraction and $V$ is the volume of the computational domain covering growing or detached bubbles. In Figure 3a, the top diagram shows the evolution of the dimensional growing bubble volume at the orifice for four consecutive bubbles with dimensional time and the bottom diagrams show the time sequence of the first growing bubble at the orifice and its detachment. In Figure 3b, the top diagram shows the corresponding detached bubble (rising bubble) volume calculated within a certain height of the computational domain (between $z_1$ and $z_2$) after bubble breakup and the bottom diagrams show the corresponding detached bubbles moving through the ambient liquid. In Figure 3a, the detached bubble volume $V_{b,d}$ is calculated as

$$V_{b,d} = (V_{b,1} - V_{b,2}) \tag{21}$$

where $V_{b,1}$ is the bubble volume just before bubble detachment [where the arrow indicates 3 at the top of Figure 3a]. When the bubble is detached, it leaves another small bubble, with volume $V_{b,2}$ [where the arrow indicates 4 at the top of Figure 3a] attached to the orifice. In Figure 3b, arrows 5–8 indicate the volumes of first rising bubble through distances between $z_1$ and $z_2$ and arrows 9 and 10 denote the volumes of second and third detached bubbles. The results show that the first detached bubble has a smaller volume than those of the second, third, and fourth detached bubbles, which have the same volume. It can be concluded for this case that the bubble formation process reaches a stable state after first bubble detachment for low Weber numbers, that is, for low gas flow rates.
Regimes of low, intermediate and high Weber numbers (gas flow rates)

The dimensionless detached bubble volume $V_b/C_3$ as a function of Weber number or gas flow rate is shown in Figure 4a when $Bo = 0.1425$ and the range of the Weber number is $4.31 \times 10^{-4} \leq We \leq 24.81$. Figure 4a shows that the influence of Weber number (gas flow rate, $Q$) on the volume of detached bubble is significant, and three regimes of the dynamics of bubble formation are observed. For Weber numbers in the range $4.31 \times 10^{-4} \leq We \leq W_{ec1}$, the volume of a detached bubble is constant and independent of $We$, where the computed value of $W_{ec1} = 1.7 \times 10^{-3}$, which is the critical Weber number (gas flow rate) below which the force due to liquid inertia caused by the bubble growth at the orifice exit is neglected and the surface tension force dominates over buoyancy force. The dynamics in this range exhibit a low Weber number region. In this region, the bubble grows quasi-statically (discussed in the section Formulation of the Problem) until the buoyancy force overcomes the surface tension confinement force. In the quasi-static regime, the computationally predicted value of detached bubble volume is $V_b/C_3 = 33.1$. For comparison, the Fritz volume is $V_F/C_3 = 44.09$ (see the section Formulation of the Problem) when $Bo = 0.1425$. It is noteworthy that the ratio of computed detached bubble volume to Fritz volume is $V_b/C_3 = 0.7507$. Our computed volume is smaller than the Fritz volume due to the deviation from an equilibrium spherical shape at its detachment.\textsuperscript{24,32,58} The Bond number based on the size of the bubble at detachment is $\left(\frac{g \ell^2}{V_b/C_3^2}\right) = \frac{Bo \times (V_b/C_3)^{2/3}}{\lambda} = 3$, which is greater than one, and the bubble at detachment is not spherical. When the Weber number exceeds the critical value $We = W_{ec1}$, the inertia of the liquid comes into play during bubble growth at the orifice and

Figure 5. Period-1 bubbling regime for $Bo = 0.1425$ and various values of $We$ and $Re$.

(a) $We = 0.69$, $Re = 530.77$; (b) $We = 2.76$, $Re = 1061.54$; and (c) $We = 6.20$, $Re = 1550$. The others two parameters are $\gamma = 6.55 \times 10^{-2}$ and $\lambda = 0.0112$. 

(a) $W_{ec1} = 1.7 \times 10^{-3}$.
the bubble volume increases with $W_e$. For the range $W_{e,c1} \leq W_e < W_{e,c2} = 6.38$ (see the section Formulation of the Problem for $W_{e,c1}$), an intermediate Weber numbers region is discerned, where the inertia of the liquid adds to the surface tension force and opposes the buoyancy force during the bubble detachment. At $W_e = W_{e,c2}$, the critical Weber number (gas flow rate) above which the opposing force of bubble growth due to liquid inertia can no longer be neglected and the influence of surface tension confinement force decreases. Beyond $W_{e,c2}$, with increasing $W_e$, the bubble volume $V_b$ increases significantly with $W_e$. The results in Figure 4a show that the volumes converge toward the results obtained from the correlation $V_b = 1.378(Q^{6/5}/g^{3/5})$ or $V_b^* = 0.5.4395(W_{ecr})^{1/5}$ given by Davidson and Schuler\textsuperscript{17} for high Weber numbers. The results indicate that at high $W_e$ the bubble volume follows the 6/5 power of the gas flow rate or the 3/5 power of the Weber number. The deviation between our results and the correlation of Davidson and Schuler\textsuperscript{17} is due to the single-stage model assuming the bubble to be growing spherically in the high $W_e$ region reported by Ref. 17. For $W_e \geq 13.95$, the bubbles start to rise in pairs and coalesce far away from the orifice (see below), known as period-2 with pairing and coalescence bubbling regime. The results in Figure 4a show that for $W_e \geq 13.95$ the computed bubble volume is the leading bubble volume of each pair represented by the dashed line. The numerical calculations of bubble volume reveal good qualitative agreement with the analytical correlation given by Jamialahmadi et al\textsuperscript{22}. The present results show a qualitatively similar trend to the results obtained by Oguz and Prosperetti\textsuperscript{32} (as shown in Figure 2 in Ref. 32) and Higuera and Medina\textsuperscript{35} (as shown in Figure 2 in Ref. 35). The numerical results of Higuera and Medina\textsuperscript{35} cover the range of Weber numbers $0.1 \leq W_e \leq 25328.72$ with $B_o = 0.1$ and $1.0$, $\eta = 0.001$ and $\lambda = 0.01$. Those of Oguz and Prosperetti\textsuperscript{32} cover wide ranges of Weber numbers, $0.0017 \leq W_e \leq 72513.601$, and Bond numbers, $0.0014 \leq B_o \leq 0.545$, $\eta = 0.001$ and $\lambda = 0.01$. Further, the comparisons between our computed results and the numerical results of Oguz and Prosperetti\textsuperscript{32} based on the BIM are shown in Figure 4b, where the volume of detached bubbles scaled with the Fritz volume $V_f$ is plotted as a function of $(W_e/W_{e,c2})^{1/2}$ (see the section Formulation of the Problem). It is observed that the present results match the results of Oguz and Prosperetti\textsuperscript{32} reasonably well. As can be seen from Figure 4, the present numerical simulations were not carried out beyond $W_e > 24.81$ or $(W_e/W_{e,c2})^{1/2} \geq 2$, the reason being that the coalesced bubble formed in period-2 with pairing and coalescence breaks up, leading to toroidal bubble formation (not shown here) for the nitrogen–LBE system. The results in Figure 4 depict that the present simulations cover bubble formation from a quasi-static regime in the very low Weber number region to an inertia-dominated regime in the high Weber number region.

We report the results of numerical simulations performed for the investigations of period-1 and period-2 with pairing and coalescence bubbling regimes. Grid sensitivity analysis of both regimes was performed using $\Delta r/R_o = \Delta z/R_o \approx 0.333$, $\Delta r/R_o = \Delta z/R_o = 0.2$, and $\Delta r/R_o = \Delta z/R_o = 0.1$ grid sizes. The dynamics of bubble formation using $\Delta r/R_o = \Delta z/R_o = 0.2$ and $\Delta r/R_o = \Delta z/R_o = 0.1$ grid sizes showed almost no significant difference (not shown here) in the two regimes. Therefore, the simulations were carried out on a $100 \times 300$ mesh grid with a uniform dimensionless mesh size of 0.2 in both directions. Figure 5 shows the period-1 bubbling regime at $B_o = 0.1425$ for (a) $W_e = 0.69$, (b) $W_e = 2.76$, and (c) $W_e = 6.20$. The simulation corresponds to nitrogen bubble formation in a liquid metal LBE and the Morton number ($M = g\mu^3/\rho^2\sigma^4$) of the liquid is $3.84 \times 10^{-13}$. The diameter of the detached bubble becomes larger than the diameter of the orifice owing to continuous injection of a constant air flow rate at the orifice. Figure 5 depicts that for each value of $W_e$, the shape of the rising bubble deviates from spherical and oscillates after detachment. At such a low value of $M$ and a moderate bubble size, the liquid inertia force dominates and shape oscillation can be observed.\textsuperscript{280} Figure 5a indicates that at low $W_e$ (low gas flow rate) the detached bubble rises sufficiently away from the orifice. With further increase in $W_e$ (increasing gas flow rate), the distance between the bubbles decreases, as shown in Figures 5b, c. However, the influence of the wake behind the rising (leading) bubble on the growing bubble (trailing bubble) at the orifice becomes insensitive and therefore no interaction can be observed. Qualitatively, the results in Figure 5 show that a regular train of nearly identical bubbles of equal size is produced from the orifice. In addition to the equal size, the bubbles are seen to form above the orifice at equal intervals of dimensionless time $t_1 = t_2$, where the bubble detachment period $t$ is defined as the time interval between the detachment instant of a bubble and that of the next bubble. Figure 6 shows the time evolution of the
dimensional growing bubble volume at the orifice for $0 \leq t < 0.25$s using the same parameters as in Figure 5. Figure 6 shows that after the first bubble detachment as in (a) and after the second bubble detachment as in (b) and (c), all consecutive detached bubbles have the same bubble volume $V_b$. The computed values of $V_b$ ($V_b^*$) are $2.08 \times 10^{-8}$ $m^3$ (49.30) for the second bubble detachment as in (b) and (c), all consecutive detached bubbles have the same bubble volume $V_b$. The computed values of $V_b$ ($V_b^*$) are $2.08 \times 10^{-8}$ $m^3$ (49.30) for...
We = 0.69, 2.805×10^{-8} m^3 (66.49) for We = 2.76, and 3.44 \times 10^{-8} m^3 (81.54) for We = 6.20. The bubble formation process reaches a stable state after the first bubble detachment at low We (low gas flow rate) as shown in Figure 6a and after the second bubble detachment at moderate We (moderate gas flow rates) as shown in Figures 6b, c. Similar observations were reported by Ohta et al.\textsuperscript{59} It can be expected from Figures 5 and 6 that with increasing We the bubble formation frequency increases. Figure 6 illustrates that for the period-1 bubbling regime, simulations can be carried out for the formation of 2–4 bubbles after discarding the initial transients.

Figures 7 and 8 show time sequence snapshots of bubble formation in the period-2 with pairing and coalescence bubbling regime at Bo = 0.1425 when We = 13.95 and We = 24.81, respectively. When We is gradually increased from 13.95 (see Figure 7), the wake behind the detached (leading) bubble begins to influence the growth of the forming bubble (trailing bubble) attached to the orifice. The liquid drag force exerted on the surface of the forming bubble is of the order $\rho_l(Q/V_{b}^{3/2} - v_w)^2V_{b}^{2/3}$ due to the wake of the leading bubble, where $v_w$ is the average wake velocity acting on the growing bubble in the presence of the leading bubble. Owing to the reduction of the liquid drag force acting on the forming bubble, the forming bubble (trailing bubble) detaches earlier than the leading bubble and the size of the trailing bubble is smaller than that of the leading bubble (see Figure 10). As a result, the leading and trailing bubbles rise as a pair and eventually merge coaxially some distance above the orifice and a compound bubble is formed. In this regime, two distinct bubble formation periods are observed ($t_1 \neq t_2$). It is found that the bubble formation period of the trailing bubble is shorter than that of the leading bubble ($t_2 < t_1$). This regime of bubbling is known as the period-2 with pairing and coalescence bubbling regime.\textsuperscript{20,21,23,35} With further increase in Weber number to We = 24.81 (see Figure 8), pairing of bubbles and coalescence move close to the orifice. The results in Figure 8 indicate that the formation periods of the trailing bubble ($t_2$)

Figure 9. Time instant of (a) the shapes (left) and the corresponding velocity field (right) before and after coalescence of two coaxial (leading and trailing) bubbles, (b) film (gap) thickness and film shape and an enlarged view (right) between the leading and trailing bubbles before coalescence, and (c) after contact of two bubbles, the motions of the minimum radius of the air bridge and the shape of the coalesced bubble.

The nondimensional parameters are Bo = 0.1425 and We = 13.95 (same parameters as used in Figure 7).
forces are greater than the restoring forces due to interfacial values of the Eotvos number greater than one, the buoyancy of leading bubble are more influenced by the leading bubble (formation period Figure 10. Time evolution of the growing bubble volume at the orifice in the period-2 with pairing and coalescence bubbling regime for $Bo = 0.1425$ under the conditions (a) $We = 13.95$ and (b) $We = 24.81$. are more influenced by the leading bubble (formation period of leading bubble $t^*$). This is because at higher Weber number, the detached (leading) bubble becomes larger in volume and the wake behind it becomes stronger. It is interesting that the coalesced bubble does not influence the growth of the next forming bubble at the orifice and the period-2 with pairing and coalescence bubbling regime repeats. In addition, at high $We = 24.81$ a tiny liquid drop inside the detached bubble is observed, as shown in Figure 8; see Figures 4 and 6 in Ref. 35 with numerical and experimental visualizations in the high $We$ region. Further, an unexpected tiny bubble is observed inside the ambient liquid; this may be due to the numerical error. In contrast, the formation of this tiny bubble may be due to the liquid drop formed inside the bubble hitting the upper part of the bubble surface after bubble coalescence. Sometimes it may not appear in the liquid, as can be seen from Figure 8. However, in this work, the formation and motion of tiny bubbles inside the ambient liquid have not been considered in detail numerically. We need more fine grids to capture this kind of tiny bubble dynamics. The same phenomena in air–aqueous glycerol solution were reported by Tufaile and Sartorelli.23 In their experiments, tiny bubbles, known as antibubbles, were produced after the coalescence of two bubbles in the period-2 bubbling regime at high gas flow rates. An antibubble is a shell of air surrounding a drop of liquid inside the liquid. The details of anti-bubble formation are described in Ref. 27.

As noted in Figures 7 and 8, the leading and trailing bubbles rise as a pair and finally coalesce above the orifice in the liquid. This kind of interaction and coalescence between two buoyancy-driven bubbles (leading and trailing bubbles) has already been discussed in the experimental studies of Bhaga and Weber81 and Katz and Meneveau82 and the numerical studies of Chakraborty et al.,76 Yuan and Prosperetti,83 and others. Studies of two coaxial drop coalescence were reported by Manga and Stone84 for the buoyancy-driven motion case. It is found from the results in Figure 10a (see below) that the calculated equivalent spherical bubble diameters of the leading, trailing and coalesced bubbles are $5 \times 10^{-3}$, $4.85 \times 10^{-3}$, and $6.23 \times 10^{-3}$ m, respectively, when $We = 13.95$ and $Bo = 0.1425$ as used in Figure 7. The corresponding values of the Eotvos number are 6.33, 5.96, and 9.83, respectively. At large values of the Eotvos number greater than one, the buoyancy forces are greater than the restoring forces due to interfacial tension and a liquid jet or wake is formed behind each bubble that pushes the bottom surface of each bubble, leading to deformations, as shown in Figure 9a (see the top part, left). However, the shapes of deformed bubbles oscillate and undergo continual deformations because of the weak restoring interfacial tension forces and strong liquid inertia. Similar observations can be also made for the case in Figure 8. The results in Figures 7–9a show that the leading bubble becomes almost flattened (mostly oblate shape) whereas the trailing bubble becomes elongated (prolate shape), as observed and discussed elsewhere.21,23,35,56,76 This prolate shape occurs due to the flow induced by the leading bubble. It can be seen that when the trailing bubble enters to the wake of the leading bubble, the drag force on the trailing bubble decreases rapidly.76,83 As a result, the trailing bubble moves faster than the leading bubble and accelerates, which eventually causes coalescence. After coalescence, the lower surface of the coalesced bubble is accelerated to its top surface and simultaneously a liquid jet is formed behind the merged bubble, as shown in Figure 9a (see both top right and bottom parts). The coalescence occurs by the dynamics of thinning of the liquid film (see Figure 9b), which was investigated by Manga and Stone,84 Chesters and Hofman,85 and Oolman and Blanch.86 It can be seen from Figure 9b that as time progresses, the film thickness decreases and the portions of the bubbles close to each other become flattened.84–86 Once the liquid film between bubbles has substantially thinned and ruptured, the bubbles make contact. We observed from our numerical investigations that the bubbles make contact when the spacing between them is less than 110 μm. At the moment of contact of two bubbles, an air bridge with large curvature is formed. As time progresses, the bridge due to its large curvature expands in the radial direction under the influence of the Laplace pressure driving the coalescence, leading to a larger coalesced bubble with a smaller surface area [Figure 9c]. The shape (oblate/prolate) oscillations and continual deformations of the coalesced bubble can be observed (Figure 9) due to strong liquid inertia and weak interfacial stresses. The detailed analysis of the coalescence between two bubbles or drops when they are allowed to contact quasi-statically and the subsequent interfacial motion has been discussed by Stover et al.,87 Thoroddsen et al.,88 and Basaran.89 The process of the coalescence of two deformable buoyancy-driven bubbles is not discussed in detail here and will be addressed further in our future work.
Figures 10a, b show the time evolution of the dimensional growing bubble volume at the orifice for $0 \leq t < 0.3$ s using the same parameters as in Figures 7 and 8, respectively. It can be seen from Figure 10 that after the second bubble detachment, bubbles of two different volumes $V_b$ are formed successively, exhibiting the period-2 bubbling regime. The bubble formation process reaches a stable state after the second bubble detachment for the case in Figure 10a and after the fourth bubble detachment in Figure 10b. The computed values of the leading bubble volume $V_{bL}$ and the trailing bubble volume $V_{bT}$ when $We = 13.95$ and 24.81 are given in Table 3. It is found that the difference in the volumes of the leading and trailing bubbles $(V_{bL} - V_{bT})$ increases with increasing Weber number. Figure 10 shows that the number of bubbles produced remains almost the same with time $0 \leq t < 0.3$ s, showing that in the high Weber number region the bubble formation frequency remains relatively constant. This regime is denoted the constant frequency bubble formation regime. Figure 10 depicts that for the period-2 with pairing and coalescence bubbling regime the simulation results are consistent after the formation of 2–3 pairs and coalescence of bubbles.

**Transition of bubbling regimes**

Studies of the transition from period-1 to period-2 with pairing and coalescence regimes in a quiescent liquid LBE are very important for keeping bubbles small by avoiding their coalescence, which is a highly undesirable process in many industrial applications. Before discussion of the transition, we need to clarify carefully the grid refinement studies on the transition position from period-1 to period-2 with pairing and coalescence regimes. Bonometti and Magnaudet reported that “the position of the transition from spherical cap to toroidal bubbles is extremely sensitive to grid resolution . . . .” Therefore, grid sensitivity analyses on the transition positions were conducted. We studied the formation of 20 bubbles under the conditions $Bo = 0.1425$, $We = 6.20$ with a dimensionless mesh size of $\Delta r/R_o = \Delta z/R_o \approx 0.333$ (as shown in Figure 11) and $\Delta r/R_o = \Delta z/R_o = 0.2$ [as shown in Figures 5c and 6c]. For grid size $\Delta r/R_o = \Delta z/R_o \approx 0.333$, period-2 with pairing and coalescence can be observed. However, no coalescence process was seen with a grid width of $\Delta r/R_o = \Delta z/R_o = 0.2$. We also carried out the same computation using a grid size of $\Delta r/R_o = \Delta z/R_o = 0.1$ and the period-1 regime can be observed (not shown here). Therefore, we stress that grid refinement studies are needed to evaluate the accuracy of the position of the transition from period-1 to period-2 regimes. Figure 11 also indicates that the motion of the interface is not captured accurately with $\Delta r/R_o = \Delta z/R_o \approx 0.333$. A bubbling regimes map between period-1 and period-2 with pairing and coalescence was constructed using the Bond number and Weber number in the nitrogen–liquid metal LBE system, as shown Figures 12a, b in which $Bo$ was varied as $0.1363 \leq Bo \leq 2.28$. For each value of $Bo$, we determined the critical value of $We$ that signals the transition from period-1 to period-2 with pairing and coalescence. The simulations were conducted using dimensionless grid sizes of 0.333, 0.2, and 0.1 to ensure the grid independence of the transition. Figure 12a indicates that the critical values of Weber number for the dimensionless grid sizes of 0.2 and 0.1 showed almost no significant changes but were very different from that for a dimensionless grid size of 0.333. It is interesting that the results follow a similar trend for three different grid sizes. The results in Figure 12a show that with increasing Bond number, the transition occurs with decreasing Weber number. The transition from period-1 to period-2 with pairing and coalescence regimes can be developed by simple scaling arguments, which have been discussed by Ambravaneswaran et al. and Subramani et al. for the case of transition from dripping to jetting. In the high-flow rate region and with $Oh \ll 1$, the transition occurs when the time scale $t_f = \frac{V_f}{v_{d,avg}}$ is smaller than the time scale for capillary breakup $t_c = \frac{V_f}{v_{d,avg}}$. The time scale $t_f$ is defined based on the capillary length scale (see Refs. 24, 25, 36, and 37) $l_c = \frac{V_f}{v_{d,avg}}$ (taking into account surface tension and gravity) and the average flow velocity of gas $v_{d,avg}$ at the orifice inlet. It is observed that the time scale for flow $t_f = \frac{R_o}{v_{d,avg}}$ is

**Table 3. The Volumes of Leading and Trailing Bubbles for $We = 13.95$ and 24.81 with $Bo = 0.1425$ in a Nitrogen–LBE System: L and T Represent Leading and Trailing Bubbles, Respectively**

<table>
<thead>
<tr>
<th>Volume of detached bubble</th>
<th>Weber number</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$We = 13.95$</td>
</tr>
<tr>
<td>$V_{bL}$</td>
<td>116.023</td>
</tr>
<tr>
<td>$V_{bL} - V_{bT}$</td>
<td>112.69</td>
</tr>
<tr>
<td>$V_{bT}$</td>
<td>3.333</td>
</tr>
</tbody>
</table>
We found that the correlation is shown in Figure 12b, where circle symbols are the computed results in air–water and nitrogen–LBE systems are shown in Figure 12c in terms of Fr and Bo with different grid sizes of 0.333, 0.2, and 0.1. Figure 12c indicates that the bubble pairing and coalescence regimes constructed using dimensionless parameters Fr and Bo for both cases, air–water with η = 0.001 and nitrogen–LBE system with η = 6.55 × 10⁻⁵.

The square symbol is the transition point obtained by Zhang and Shoji in an air–water system, which is at Fr = 258.21 for Bo = 0.1363. The predicted result from our simulation is Fr ≈ 259 for Bo = 0.1363. The transition between period-1 to period-2 with pairing and coalescence is presented using dimensionless grid sizes of 0.333, 0.2, and 0.1.

Kyrriakides et al. and Badam et al. experimentally and Buwa et al. numerically reported the transition of various bubbling regimes using the Bond number (Bo1 = 4gR2/σ) and Froude number (Fr1 = Q2/2πgR3). However, all of their results were based on an air–water system. According to our definition of Froude and Bond numbers, the relations are Fr = 2Fr1 and Bo = 0.25Bo1. Comparative results of the transitions in air–water and nitrogen–LBE systems are shown in Figure 12c in terms of Fr and Bo with different grid sizes of 0.333, 0.2, and 0.1. Figure 12c indicates that the bubble pairing and coalescence processes appear fairly early in LBE compared with those in water. Considering the air–water system, Zhang and Shoji found that for Bo = 0.1363, the period-1 regime transits to the period-2 with pairing and coalescence regime at Fr ≈ 258.21 [square symbols in Figure 12c] and our computed value is Fr ≈ 259. Our predicted result agrees well with the experimental result of Zhang and Shoji. For the same Bond number in the nitrogen–LBE system, we obtained the transition point at Fr = 106.823. Figure 12c illustrates that for the same Fr, the transition is observed in water at higher Bo compared with the case in LBE. Similarly, for same Bo, the transition is observed in LBE at lower Fr compared with the case in water. In addition, It is found from our correlation, We = 1.7Bo⁻¹, which is consistent with the scaling argument. The results in Figure 12b indicate that the critical value of We varies inversely with Bo.

Buwa et al. numerically reported the transition of various bubbling regimes using the Bond number (Bo1 = 4gR2/σ) and Froude number (Fr1 = Q2/2πgR3). However, all of their results were based on an air–water system. According to our definition of Froude and Bond numbers, the relations are Fr = 2Fr1 and Bo = 0.25Bo1. Comparative results of the transitions in air–water and nitrogen–LBE systems are shown in Figure 12c in terms of Fr and Bo with different grid sizes of 0.333, 0.2, and 0.1. Figure 12c indicates that the bubble pairing and coalescence processes appear fairly early in LBE compared with those in water. Considering the air–water system, Zhang and Shoji found that for Bo = 0.1363, the period-1 regime transits to the period-2 with pairing and coalescence regime at Fr ≈ 258.21 [square symbols in Figure 12c] and our computed value is Fr ≈ 259. Our predicted result agrees well with the experimental result of Zhang and Shoji. For the same Bond number in the nitrogen–LBE system, we obtained the transition point at Fr = 106.823. Figure 12c illustrates that for the same Fr, the transition is observed in water at higher Bo compared with the case in LBE. Similarly, for same Bo, the transition is observed in LBE at lower Fr compared with the case in water. In addition, It is found from our correlation, We = 1.7Bo⁻¹, that at Bo = 0.1 and 1 the transition occurs when We¹/² = 4.123 and 1.304, respectively, in the nitrogen–LBE system. It is observed that the Bond number (Bo < 0.1363) becomes relatively insensitive to the value of the Weber number when We > 14 [We¹/² > 3.742; see Figure 12a]. The reason is that the period-1 regime directly transits to the period-2 with double coalescence regime. Conversely, Higuera and Medina found from their numerical simulations that for the same values of Bo, the transition occurs when We¹/² = 15.924 and 6.369, respectively, in the air–water system. The substantial difference from the value of We in the transition of Higuera and Medina may be due to the effect of the density of the ambient liquid and the wake.
effects of the detached (leading) bubble on the forming (trailing) bubble at the orifice. The liquid inertia exerted an upward force on the surface of the forming bubble of the order of $\rho_l v_w^2$ due to the averaged wake velocity, $v_w$, of the previous (leading) detached bubble. The stronger liquid inertia $\rho_l v_w^2$ brings about early detachment of the trailing bubble, resulting in a faster interaction and coalescence process between leading and trailing bubbles in LBE than in water.

**Detached rising bubbles in a high-density liquid**

From low to high gas flow rates, the calculated rising Reynolds number based on the bubble velocity and equivalent spherical detached bubble diameter in a liquid metal LBE is in the range $2000 < Re < 10,000$ and the Eotvos number is in the range $2 < Eo < 7$. In this range of Reynolds numbers, the motion of the detached bubbles may deviate from a vertical line and rise in a zigzag or spiral motion and the flow is nonaxisymmetric.\(^{90-94}\) The path instability (zigzag or spiral) of a rising millimetric air bubble with high Reynolds numbers in water was reported by Magnaudet and coworkers\(^{90-92}\) using 3-D simulations, and established that the primary cause of the onset of path instability is the wake instability in which the flow is 3-D (nonaxisymmetric) and time dependent. It was observed that the path of a rising bubble is rectilinear when the wake is axisymmetric. Magnaudet and Mougin\(^{92}\) found by direct numerical simulation that when the bubble aspect ratio is $\chi < 2.21$, the flow is axisymmetric and the wake is stable whatever the Reynolds number. The bubble rises rectilinearly. Beyond $\chi = 2.21$, the 3-D flow influences the dynamics of the rising bubble and the bubble undergoes zigzag and spiral motion within a finite range of Reynolds number. They found that the flow recovers its axisymmetry again above this finite range of Reynolds number. They also reported that when $\chi = 2.5$ and $Re > 2700$, the flow returns to its axisymmetric form. Magnaudet and coworkers studied the loss of flow axisymmetric flow based on the assumption of a fixed spheroidal bubble shape. However, Cano-Lozano et al.\(^{93}\) investigated the transition from straight to zigzag motion of a fixed real shape of the bubble in terms of the dimensionless parameters that govern the flow dynamics of the rising bubble. In their studies, the shape of the bubble was not considered fixed initially. The real shape of the rising bubble was obtained by means of axisymmetric computations and 3-D simulations were performed using the shape of the bubble and its terminal velocity provided by axisymmetric computations. The results obtained from their studies are more realistic. Furthermore, the stable (considered rectilinear motion) and unstable (considered zigzag or spiral motion) regions of a single bubble rising in quiescent liquids with very low values of Morton number in the range of $3.768 \times 10^{-4} \leq M \leq 7.710 \times 10^{-12}$ were discussed in detail by Cano-Lozano et al.\(^{94}\) and references therein. Cano-Lozano et al.\(^{94}\) using Volume of Fluid technique presented extensive numerical results for buoyancy-driven bubble motion in the stable and close to the unstable regions. The accuracy of their numerical results in terms of terminal bubble shape and terminal velocity was assessed by comparison with results of previous investigations available in the literature. The fundamental understanding of the path instability of a single bubble rising in liquids with high Reynolds number at very low Morton number is still not well established. However, a limitation of the present work is the assumption of axisymmetric fluid flows and bubble shapes.

**Conclusions**

We have carried out numerical simulations of the dynamics of bubble growth, generation and rising in stagnant liquids with very low Morton number, $M = O(10^{-11}-10^{-13})$, and moderately high Reynolds number, $Re \gg 1$, using a CLSVOF method in an axisymmetric computational domain. We examined the dynamics of bubble formation for a wide range of dimensionless parameters, $4.31 \times 10^{-3} \leq We \leq 24.81$ and $0.13 \leq Bo \leq 2.28$, while keeping constant values of $\eta = 6.55 \times 10^{-3}$ and $\lambda = 1.12 \times 10^{-2}$. The computations and subsequent results with respect to detached bubble volume showed good agreement with the analytical models.\(^{22,38}\) It was found that the present simulations cover bubble formation from the quasi-static regime in the very low Weber number region to an inertia-dominated regime in the high Weber number region (see Figure 4). The accuracy of the computed results was assessed by comparison with scaling laws proposed in the literature.\(^{17,32}\) For two regimes of bubble formation: the low flow rate quasi-static regime and high flow rate dynamic regime. In this article, all the computed results are presented after careful grid refinement studies.

The present work was focused on the study of the influence of Weber number (gas flow rate) for a given Bond number (orifice diameter) on the growing bubble volume at the orifice, bubble formation frequency, detached bubble volume and the shape of the bubble. The period-1 and period-2 with pairing and coalescence bubbling regimes were discussed. At high Weber numbers, the volume (Figure 10) and departing period (Figures 7 and 8) of the leading bubbles are larger than those of the trailing bubbles, and this repeats in an alternating fashion exhibiting the period-2 bubbling regime. We also observed that during the period-1 bubbling regime, the bubble formation process reaches the steady state after the second bubble detachment whereas in the period-2 regime the steady-state bubble formation process is reached after 2–3 pairs and coalescence.

The grid refinement tests on the transition point from period-1 to period-2 with pairing and coalescence were investigated. Based on these tests, we can highlight again that the grid resolution is extremely important for finding the critical values for the transition of different bubbling regimes. In addition, the transition map shows that the system always exhibits the period-2 with pairing and coalescence bubbling regime at $We \approx 1$ when the Bond number is sufficiently high, $Bo > 2$.

It is interesting that the dynamics of bubble formation in liquid metal are identical with those of bubble formation in water when the Bond number, $Bo$, Weber number, $We$, and static contact angle, $\phi_s$, are same in both cases. Nevertheless, we found that, even in the high Reynolds number bubble formation region, the possibility of interaction and coalescence in liquid LBE is much faster than that of water for the same value of $Bo$.

**Acknowledgments**

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**Notation**

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
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<tbody>
<tr>
<td>$D_o$</td>
<td>orifice diameter, m</td>
</tr>
<tr>
<td>$g$</td>
<td>gravitational acceleration vector, m/s$^2$</td>
</tr>
<tr>
<td>$H_\sigma$</td>
<td>Heaviside function</td>
</tr>
<tr>
<td>$n$</td>
<td>normal at the interface</td>
</tr>
<tr>
<td>$p$</td>
<td>pressure, N/m$^2$</td>
</tr>
</tbody>
</table>
\[ Q = \text{volumetric flow rate, m}^3/\text{s} \]
\[ R = \text{computational domain size, m} \]
\[ r = \text{radial component of the axisymmetric coordinate, m} \]
\[ S = \text{Newtonian stress tensor, N/m}^2 \]
\[ t = \text{time, s} \]
\[ u = \text{velocity component in r direction, m/s} \]
\[ V = \text{growing bubble volume at the orifice, m}^3 \]
\[ V_b = \text{detached bubble volume, m}^3 \]
\[ v = \text{velocity component in z direction, m/s} \]
\[ \tilde{V} = \text{velocity vector} \]
\[ \nu_{avg} = \text{gas mean velocity at the inlet of the orifice} \]
\[ \nu_{AVG} = \text{Vector of the axisymmetric coordinate, m} \]

Greek letters
\[ \alpha = \text{void fraction of liquid in two-phase cell} \]
\[ \tilde{\alpha} = \text{smoothed void fraction field} \]
\[ \iota = \text{interface numerical thickness, m} \]
\[ \eta = \text{density ratio} \]
\[ \kappa = \text{mean curvature of the interface, m}^{-1} \]
\[ \lambda = \text{viscosity ratio} \]
\[ \mu = \text{viscosity of fluid, Pa s} \]
\[ \rho = \text{density of fluid, kg/m}^3 \]
\[ \sigma = \text{surface tension coefficient, N/m} \]
\[ \tau = \text{bubble detachment period} \]
\[ \phi = \text{level set function} \]

Subscripts
\[ \text{avg} = \text{average value} \]
\[ \text{cr} = \text{critical} \]
\[ d = \text{disperse phase (gas)} \]
\[ g = \text{gas} \]
\[ L = \text{leading bubble} \]
\[ l = \text{liquid} \]
\[ N = \text{number of detached bubbles} \]
\[ o = \text{orifice} \]
\[ T = \text{trailing bubble} \]

Superscripts
\[ * = \text{Dimensionless quantity} \]
\[ \rightarrow = \text{vector} \]
\[ \wedge = \text{unit vector} \]

Dimensionless numbers
\[ Bo = \text{Bond number} \]
\[ Eo = \text{Eotvos number} \]
\[ M = \text{Morton number} \]
\[ Re = \text{Reynolds number} \]
\[ We = \text{Weber number} \]

Literature Cited