Numerical modelling of ultrasonic waves in a bubbly Newtonian liquid using a high-order acoustic cavitation model

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Abstract

To address difficulties in treating large volumes of liquid metal with ultrasound, a fundamental study of acoustic cavitation in liquid aluminium, expressed in an experimentally validated numerical model, is presented in this paper. To improve the understanding of the cavitation process, a non-linear acoustic model is validated against reference water pressure measurements from acoustic waves produced by an immersed horn. A high-order method is used to discretize the wave equation in both space and time. These discretized equations are coupled to the Rayleigh-Plesset equation using two different time scales to couple the bubble and flow scales, resulting in a stable, fast, and reasonably accurate method for the prediction of acoustic pressures in cavitating liquids. This method is then applied to the
context of treatment of liquid aluminium, where it predicts that the most intense cavitation activity is localised below the vibrating horn and estimates the acoustic decay below the sonotrode with reasonable qualitative agreement with experimental data.

Keywords: Acoustic cavitation, Numerical acoustics, Ultrasonic wave propagation, Ultrasonic melt processing, Light metal alloys

1. Introduction

Significant improvements in the quality and properties of metallic materials are observed when treating them near their liquidus temperature [1-3]: the beneficial effects of the treatment include the degassing of dissolved gases, improved wetting, activating inclusions by cleaning the solid-liquid interface, enhancing nucleation, and refining the structure of the solidified metal [1, 4]. These improvements are primarily attributed to acoustic cavitation [5]; the term “cavitation” here follows the definition of Neppiras [6] and is restricted to cases involving the formation, expansion, pulsation, and collapse of existing cavities and bubble nuclei. However, treating large volumes of liquid metal, as is required by industrial processes such as continuous casting, is still problematic: the process is time-consuming and volume-limited so it can currently be applied only to a fixed volume of melt in a crucible. To circumvent these difficulties and facilitate the transfer of this promising technology to industry, a fundamental study of melt cavitation treatment is required [7].

Nastac [8] used the ‘full cavitation model’ [9] developed for hydrodynamic cavitation to model solidification structure evolution in an alloy in the presence of ultrasonic stirring, while computing the acoustic field analytically from the Helmholtz reduced wave equation. An improved version of this model, based on the Keller-Miksis equation [10] and including a turbulent source term arising from the collapse of cavitating bubbles, has been proposed by the authors [11, 12] to model a moving liquid metal volume in a launder. However, the use of a homogeneous cavitation model, e.g. the ‘full cavitation model’, for acoustic cavitation is questionable. Also, the acoustic solver used in [11, 12] was second order in space and
prone to numerical diffusion; hence a higher-order model [13] is desired to improve the accuracy of the acoustic field prediction. The presence of bubbles significantly alters the ultrasonic wave propagation in the melt, and this influence must be accurately quantified to understand the effect of the acoustic field on a volume of treated metal. In this endeavour, we are proposing a macroscopic cavitation model coupled with a high-order acoustic solver, using reference experiments in water [14] for validation.

A plethora of empirical observations of acoustic cavitation in water is available in the literature. For example, observations of streamers and acoustic Lichtenberg figures have been recorded [15]. The conical bubble structure below the radiating surface of the sonotrode has been observed and studied by many authors [14, 16-19]. The tendency of bubbles to form clusters after collapsing has been observed with a high-speed camera [20]. Pressure measurements with calibrated hydrophones in water under ultrasonic treatment are also available [14, 21].

Alongside this empirical evidence, there exists a series of models attempting to explain and reproduce the bubble cloud behaviour numerically. One class of such models is based on a set of non-linear equations proposed by van Wijngaarden [22] to model wave propagation in a bubbly liquid. Caflisch et al. [23] re-derived this set of non-linear differential equations from the microscopic motion equations of a large number of bubbles. Commander and Prosperetti wrote an extensive review of pressure wave propagation models in bubbly liquids [24] following the insight given by Caflisch et al. [23]. A simplified version of the Caflisch equations has recently been used by Louisnard [25, 26] to model bubble structures below a sonotrode in water. Other recent advances include the work of Dahnke et al. [27] who modelled the acoustic pressure field in sonochemical reactors with an inhomogeneous density distribution. Vanhille et al. applied and extended a model consisting of a coupled linear non-dissipative wave and volume variation equations [28] based on the Rayleigh-Plesset equation [29] to model the nonlinear propagation of ultrasonic waves in water-air
bubble mixtures [30-33]. Servant et al. [34-36] considered the Bjernkes forces [37] in their model of mono- and dual-frequency sono-reactors.

Tudela et al. completed a more recent review [38] of the state of the research and outlined the need to model the nonlinear nature of the problem. They highlighted that the Caflisch-type equations have certain drawbacks for the simulation of the effect of bubbles on strong acoustic fields due to: the non-linear nature of the problem, the limits of using assumptions on bubble sizes and distribution, the assumption of low bubble volume fractions in the derivation of the model, and the applicability only in cases where bubble resonance plays a negligible role [38]. Moreover, this class of models requires extremely small time steps for acoustic pressures higher than the Blake threshold, making it unattractive for the design of experiment simulations that seek optimum parameters to enhance cavitation activity. Despite these drawbacks, resolving the complex coupling between the void fraction and acoustic pressure field is necessary and this class of numerical models is therefore unavoidable in acoustic cavitation modelling.

In this paper, a high-order acoustic model coupled with a cavitation model is presented, followed by validation against acoustic pressures measured in water [14] and then applied to the treatment of aluminium in a crucible.

2. Theory

From the conservation of mass and momentum, and using $\frac{\partial p}{\partial \rho} \equiv c^2$, the governing equations for sound propagation in a moving fluid are the Navier-Stokes equations in perturbation form:

$$\frac{\partial p}{\partial t} + v_j \frac{\partial p}{\partial x_j} + \rho c^2 \frac{\partial v_i}{\partial x_j} = S \tag{1}$$

$$\frac{\partial v_i}{\partial t} + v_j \frac{\partial v_i}{\partial x_j} + \frac{1}{\rho} \frac{\partial p}{\partial x_i} = \frac{\partial \tau_{ij}}{\partial x_j} + F_i \tag{2}$$
\( p \) denotes pressure and \( v_j \) are the velocities. \( \rho \) is the liquid density. \( S \) contains mass sources, such as due to vibrating solid surfaces and growing (or collapsing) bubbles. In the model presented in this paper, an additional source term, \( \rho c^2 \partial \phi / \partial t \), is added to the source term \( S \) of equation (1) to account for the acoustic pressure waves induced by the collapse of bubbles, and conversely the sink of acoustic pressure during the creation of bubbles [30].

The forcing terms \( F_i \) are usually set to zero for most practical acoustics problems [13] and contain acoustic velocity sources due to a vibrating surface. Ignoring dissipation due to viscosity \( (\partial \tau_{ij} / \partial x_j \text{ term}) \), the convection terms \( v_j \frac{\partial p}{\partial x_j} \) and \( v_j \frac{\partial v_i}{\partial x_j} \), and considering a constant speed of sound, equations (1) and (2) reduce to the standard, linear Helmholtz equation. However, these assumptions are not accurate for modelling acoustic cavitation.

In this implementation, the viscosity and convection terms are retained: this makes equations (1) and (2) fully coupled and non-linear, unlike the linearized cases in [13, 39]. The effect of the flow on pressure predictions is modelled by including the convection terms. The speed of sound in the liquid is given by \( c = \sqrt{K / \rho} \). However, variations of density and bulk modulus in the bubbly liquid lead to numerical instability due to the discontinuity in derivatives along the saturation curve that separates single phase and two-phase domains [40]. These numerical instabilities can be avoided by treating the speed of sound as a constant, thereby restricting the accuracy of the method to void fractions smaller than 1% [23]. This assumption is applicable to liquid metals where the bubbles originate from dissolved hydrogen.

The gas volume fraction \( \phi = \frac{4}{3} \pi n_0 R^3 \), where \( n_0 \) is the number of stationary bubbles of radius \( R \) per unit volume, is calculated from the solution to the Rayleigh-Plesset equation [29] which governs the dynamics of a bubble in the presence of a strong acoustic field:

\[
R \ddot{R} + \dot{R}^2 = \frac{p_s}{\rho},
\]

where \( p_s \) is given by
\[ p_s = p_b(t) + p_v - \frac{2\sigma}{R} - \frac{4\mu \dot{R}}{R} - p(t), \]  

(4)

and \( p_b \) is the pressure inside the bubble. \( p_v \) is the vapour pressure in the bubble. \( \sigma \) is the surface tension between the gas and liquid interface. \( \mu \) is the dynamic viscosity of the liquid.

The use of the Rayleigh-Plesset equation to derive the gas volume fraction assumes the following:

1. The internal pressure of the bubbles is homogeneous, since the inertia of the gas is negligible.
2. The bubbles remain roughly spherical. Due to the large value of surface tension of an interface of hydrogen with liquid aluminium, bubbles observed during melt cavitation are small, in the region of 10-100 \( \mu \)m in radius [41].

3. Modelling

3.1. Wave equations discretization

The discretization method of Djambazov et al. [13] is used to solve equations (1) and (2). The computational meshes are fully staggered, as described in [13] and illustrated in Figure 1. Fully staggering pressures and velocities allows the formulation of a fully explicit, stable, second order accurate scheme [42]. The accuracy can then be extended to higher orders by allowing the scheme to become implicit provided it retains a strong diagonal dominance to ensure fast convergence [39]. The computational domain is divided into regular cells, with the scalar quantities pressure and bubble volume fraction stored in cell centres, and velocity components stored at cell faces in the middle of each time step. In this formulation, curves are represented by a castellated mesh, i.e. with a bitmap grid-like structure. The DRP (dispersion-relation-preserving) scheme [43] is used with its differentiation and temporal integration steps for the convection integrals only.
3.1.1. Spatial derivatives

The terms $\rho c^2 \frac{\partial v_j}{\partial x_j}$ and $\frac{1}{\rho} \frac{\partial p}{\partial x_i}$ are evaluated with 6-point stencils for the spatial derivatives. At solid boundaries, values are mirrored to provide the missing points, and the free surface is modelled as a layer of cells with fixed acoustic pressure $p = 0$ Pa. For radiative boundary conditions, the derivatives are extrapolated from previous time step values [13].

In all other cells, the first derivative of a function $f$ with respect to $x$ is expressed as

$$\frac{\partial f}{\partial x} = \frac{1}{\Delta x} \sum_{j=-2}^{3} a_j f \left[ x + \left( j - \frac{1}{2} \right) \Delta x \right] \quad (5)$$

The coefficients $a_j$ have been optimized by Djambazov [13] to make the scheme exact to the fourth order in space. These coefficients are provided in the appendix.

Since the fluids considered here are Newtonian, the second derivatives of velocity are required to determine $\partial \tau_{ij}/\partial x_j$. Since viscosity is added as a source term in $F_i$, the viscous forces are computed at each iteration: the second derivatives of velocity $v_i$ along $x_j$ are computed as

$$\frac{\partial^2 v_j}{\partial x_i^2} = \frac{1}{\Delta x} \left( v_{j-1} + v_{j+1} - 2v_j \right) \quad (6)$$
3.1.2. Temporal integration

The terms $\rho c^2 \frac{\partial v_j}{\partial x_j}$ and $\frac{1}{\rho} \frac{\partial p}{\partial x_i}$ are computed in the middle of each time step and are then used to update pressure and velocities at the end of their time steps, using the following approximate integration

$$\int_{\frac{2}{3t}}^{\frac{2}{3t}} f(t) dt = \Delta t \sum_{m=0}^{3} b_m f(-m\Delta t)$$ (7)

with the coefficients $b_m$ chosen to make the scheme third order accurate in time [13]. These coefficients are provided in the appendix.

3.2. Adaptive time stepping for bubble dynamics equation

The Rayleigh-Plesset equation (3) is solved using the Runge-Kutta method with an adaptive time step $h < \Delta t$ evaluated as follows [44]:

1. The Jacobian matrix $[J]$ is calculated as

$$[J] = \begin{bmatrix} \frac{\partial \dot{R}}{\partial R} & \frac{\partial \dot{R}}{\partial \dot{R}} \\ \frac{\partial \dot{R}}{\partial R} & \frac{\partial \dot{R}}{\partial \dot{R}} \end{bmatrix}$$ (8)

2. The eigenvalues $\lambda_1$ and $\lambda_2$ of the amplification matrix $[A] = 1 + h[J]$ are evaluated. The maximum deviation from 1.0 is calculated as $\delta = \max(|\lambda_1 - 1|, |\lambda_2 - 1|)$.

3. The time step $h$ is resized so that $\delta$ becomes close to a target value denoted by $\delta_{\text{max}}$. For stability, $\delta_{\text{max}}$ is set to a small value (0.001) so that the maximum Lyapunov exponent is 1.0 in practice.

Depending on the stage of the bubble cycle, $h$ can vary between $10^{-20}$ s to $10^{-8}$ s.

3.3. Iterative procedure

Starting from an initial guess for the solved variables, the coupled equations (1), (2), and (3) are solved in each time step as follows:
1. The acoustic pressure is solved from equation (1).

2. The Rayleigh-Plesset equation (3) is solved in a separate time scale according to the procedure outlined in Section 3.2. The initial bubble radius and interface velocity in each cell are taken from the previous time step and their values updated with smaller adaptive time increments until the flow time step value is reached. The gas volume fraction is then calculated from the new radii.

3. The velocity components of the pressure perturbation are solved from equation (2) at the end of the time step.

4. The solution is advanced to the next time step and procedures 1 to 3 are repeated until the last time step is reached.

3.4. Material properties

Table 1: Material properties for water and aluminium [1, 45, 46]. Surface tension with air interface for water. Hydrogen interface for aluminium.

<table>
<thead>
<tr>
<th>Material</th>
<th>Water</th>
<th>Aluminium</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature (°C)</td>
<td>20</td>
<td>700</td>
</tr>
<tr>
<td>Density $\rho_l$ (kg m$^{-3}$)</td>
<td>1000</td>
<td>2375</td>
</tr>
<tr>
<td>Dynamic viscosity $\mu$ (10$^{-3}$ Pa s)</td>
<td>1.004</td>
<td>1.0</td>
</tr>
<tr>
<td>Kinematic viscosity $\nu$ (10$^{-6}$ m$^2$ s$^{-1}$)</td>
<td>1.0</td>
<td>0.42</td>
</tr>
<tr>
<td>Speed of sound $c$ (m s$^{-1}$)</td>
<td>1482</td>
<td>4600</td>
</tr>
<tr>
<td>Surface tension $\sigma$ (N m$^{-1}$)</td>
<td>0.079</td>
<td>0.860</td>
</tr>
<tr>
<td>Vapour pressure $p_v$ (kPa)</td>
<td>2.2</td>
<td>negligible</td>
</tr>
<tr>
<td>Bulk modulus $K$ (GPa)</td>
<td>2.15</td>
<td>41.2</td>
</tr>
</tbody>
</table>
The material properties used in the numerical simulations are listed in Table 1. The gas phase is assumed to be adiabatic in each case and therefore $\kappa = \gamma = 1.4$.

3.5. Initial and boundary conditions

The liquid is initially unperturbed (constant hydrostatic pressure and all velocity components set to 0) and contains $n_0$ bubbles of radius $R_0$ per unit volume. In liquid metals, an initial number of nuclei is always assumed since cavitation is attributed to both the hydrogen-containing inclusions and the dissolved hydrogen that is released from aluminium when the local pressure decreases [1]. The vapour pressure of aluminium at its melting point is $0.000012$ Pa [47] and therefore vapour bubbles are unlikely to form in the liquid bulk [48]. Based on the numerical values of acoustic simulations from the literature [30, 31], the number of bubbles per unit volume (bubble density) is $n_0 = 1 \times 10^{11}$ m$^{-3}$ and the initial radius used in water was in the range $R_0 = [1, 10]$ μm. This corresponds to an average distance of 22 radii between bubbles in the extreme case $R_0 = 10$ μm. This separation is long enough to prevent the motion of bubbles due to the effect of secondary Bjerknes forces [49]. For aluminium, $n_0 = 1 \times 10^{11}$ m$^{-3}$ and $R_0 = 1$ μm.

A sinusoidal pressure signal is indirectly prescribed below the sonotrode (see Figure 2) by specifying the acoustic velocity at the sonotrode surface. The acoustic velocity amplitude is calculated from the displacement amplitude $A$ of the sonotrode as

$$v = 2\pi f A$$  \hspace{1cm} (9)$$

The upper boundary is a free surface from which a $180^\circ$ phase shift occurs upon reflection of the acoustic wave: this is approximated by setting $p = 0$ Pa in the top row of computational cells (representing the atmosphere above the interface). All other boundaries, including the sonotrode walls, are fully reflective to sound and are modelled using the mirroring technique from [13]. Radiative boundary conditions are used to approximate absorbent boundaries. The derivatives in transparent cells at the edges of the domain are updated using a second order interpolation on a 3-point stencil [39].
3.6. Geometry and mesh

The geometry of a water vessel is shown in Figure 2 and corresponds to the setup from Campos-Pozuelo et al. [14]. The liquid depth is 18 cm. The radiating surface of the sonotrode, vibrating at 20 kHz, is 1 cm below the free surface. The sonotrode radius is 3.5 cm. The left and right boundaries are fully reflective to sound, as are the sonotrode walls. The bottom boundary consisting of an absorbent material is modelled as a transparent boundary. The top boundary is a free surface. The velocity of the horn is provided in [14] and pressure measurements 4 cm below the sonotrode axis are used for validation.

Figure 2: Geometry of water vessel in the experimental setup from [14]. The origin (black dot) is taken as the point of intersection between the liquid free surface and the axis of the sonotrode. The hydrophone position (clear dot) is 4 cm below the sonotrode surface.

Three mesh densities were used in the simulation and the bubble dynamics at the monitoring point, corresponding to the hydrophone location in [14], were found to be independent of the grid size $\Delta x$. Results are presented in a medium coarse mesh with grid size $\Delta x$ of 2.0 mm. The same grid size is used in all coordinate directions.
The solution is computed on a 2D Cartesian mesh to make the problem tractable with converged meshes. This restriction makes only qualitative comparisons with experiments possible. Full 3D computations are planned in future works after a massively parallel implementation of the acoustic model is completed.

The Courant number is given by $Cou = \Delta t \cdot (c + \max(\vec{v})) / \Delta x$. For numerical stability, the time step and grid size in each case are chosen such that the Courant number is always less than 0.2. Below a Courant number of 0.2, the computed pressures at the validation point are identical: all results are therefore presented with a Courant number of 0.1, corresponding to time steps of the order of $\mu$s.

Another simulation is then run for the case of liquid aluminium in a crucible as depicted in Figure 3, corresponding to the setup available at Brunel University London [50]. The crucible walls are fully reflective and a 180° shift occurs upon reflection from the free surface. The liquid height is 17.5 cm, the radius of the cylindrical base is 6 cm corresponding to a charge of 5.2 kg of commercially pure aluminium at 700 °C. The transducer operates at 17.7 kHz and 3.5 kW input power, the sonotrode tip (20 mm in diameter) is immersed 20 mm below the free surface. The displacement of the horn is calculated from the operating power [51].
4. Results and discussion

This section describes the comparison of predicted pressures in water with experimental data and the profile of the predicted bubble cloud below the horn. Following the comparison with water, the aluminium sonication case is presented with qualitative comparison with experimental data.

4.1. Water

Simulations using the high-order acoustic model are compared with experimental pressure measurements from [14]. Figure 4 shows the predicted and measured pressure evolution at the hydrophone position 4 cm below the sonotrode.
Figure 4: Comparison between numerical results and pressure measurements 4 cm below the sonotrode.

The maximum predicted pressure of 570 kPa is not significantly far from the maximum recorded pressure of 610 kPa. A pressure wave is also emitted at each bubble collapse. Both the peak pressures and the negative pressures are of the same order of magnitude for both numerical predictions and measured values. An exact realization of the experimental data is not possible, since the exact operating conditions (including position of initial nuclei, roughness of surface of vessel, precision in the location of the sonotrode within the mesh resolution …) cannot be possibly determined: this is also why cavitation pressure measurements appear chaotic. Nevertheless, the broad features of the cavitation dynamics, namely the peak pressures and intervals, are correctly predicted.
Figure 5: Comparison between experimental spectrum and numerical prediction.

Figure 5 shows the comparison between the numerical and experimental spectra. There is a general good agreement with the predicted frequencies. The strong subharmonic peaks are due to the influence of the bubble cloud below the sonotrode that grows and collapse at a rate of roughly $\frac{1}{4}$ of the forcing frequency, consistent with the experimental supercavitation observation from [52].

The decay of pressure amplitude with distance from the source is reasonably predicted, as shown in Figure 6. The values for pressure amplitude are obtained by applying a low-pass Butterworth filter to the predicted numerical pressures in the computational cells located at axis of the sonotrode.
Figure 6: Comparison between measured pressures and numerical prediction at cavitating conditions. The pressure amplitude from the numerical simulation is obtained by filtering the numerical pressures using a Butterworth filter.

Figure 7: Development of the bubble cloud below the sonotrode in water. Light grey contours represent volume fractions of 0.1 % and darker contours represent volume fractions of 0.5 %.
Figure 7 shows the evolution of the bubble cloud structure below the sonotrode in water. The cavity first grows (50 μs) and then collapses from the outer rim to form a toroidal shape (79 μs). This is a less dramatic effect of the “acoustic supercavitation” effect reported by [52], due to the lower energy densities involved in this experiment. The large conical bubble structure, that is routinely observed in water experiments [53], is established after a period of initial transient although the structure does not keep a perfectly stable shape (as shown from 174 μs).

A time averaged volume fraction in the last cycle of the simulation is shown in Figure 8. The numerical method does not predict the correct size of the conical structure, with the cone occupying the whole sonotrode surface instead of 2/3 of the area as observed experimentally [18]. With a qualitative 2D comparison, it is not possible to infer the source of this discrepancy since 2D images of the conical bubble structure cover the whole plane below the sonotrode.

Figure 8: Time averaged bubble structure below the sonotrode.
4.1.1. Sensitivity of results to bubble density and initial radius

A parametric study of the effect of the choice of initial radii $R_0$ and bubble density $n_0$ on the pressure predictions has been conducted with Dakota [54] with a Courant number of 0.02 for all cases. The multidimensional study was performed with 3 partitions for each variable and the bounds were $1 \mu m < R_0 < 10 \mu m$ (a factor of 10 lower than bubble size observed during cavitation) and $1 \times 10^{11} \text{ m}^{-3} < n_0 < 1 \times 10^{12} \text{ m}^{-3}$, values commonly used in the literature for water [30-33]. The results of the sensitivity analysis are given in Table 2. The pressure predictions are insensitive to variations of bubble density at low initial volumes and low initial radii. However, large initial radii lead to variations in predictions and should be avoided in numerical simulations: these low values are consistent with the small sizes of stable hydrogen bubbles in the melt before cavitation [1].

Table 2: Multidimensional parametric study

<table>
<thead>
<tr>
<th>$R_0$ (μm)</th>
<th>$n_0$ (m$^{-3}$)</th>
<th>Initial volume (m$^3$)</th>
<th>Maximum pressure (MPa)</th>
<th>Minimum pressure (MPa)</th>
</tr>
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<tbody>
<tr>
<td>1</td>
<td>1.00E+11</td>
<td>4.19E-07</td>
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<tr>
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<td>4.00E+11</td>
<td>1.68E-06</td>
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</tr>
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<td>4.19E-03</td>
<td>0.10</td>
<td>-0.04</td>
</tr>
</tbody>
</table>
4.2. Aluminium

Figure 9: Development of the bubble cloud below the sonotrode in aluminium. Light grey contours represent volume fractions of 0.1 % and darker contours represent volume fractions of 0.5 %.

Results for the aluminium sonication case are presented in this section, using a Courant number of 0.1, \( n_0 = 1 \times 10^{11} \text{ m}^{-3} \), and \( R_0 = 1 \mu \text{m} \). Figure 9 shows the evolution of the bubble cloud structure below the horn. After an initial transient period (< 100 \( \mu \text{s} \)), a stable cone-like structure is developed below the sonotrode: this is the region of intense cavitation activity where nucleation sites are generated during melt treatment.

Figure 10 shows the predicted pressure at 5 selected points in the computational domain: 3 points along the axis of the sonotrode at 2 cm, 6 cm and 10 cm below the sonotrode surface, and 2 off-axis positions all 2 cm below the sonotrode surface – at \( x = 3 \text{ cm} \) (corresponding to the midpoint between the axis and the crucible) and \( x = 6 \text{ cm} \) (at the edge of the crucible). The acoustic pressures predicted decrease dramatically with distance away from the sonotrode and away from the axis, with the point 2 cm below the sonotrode axis registering the highest pressure of 9.5 MPa. Maximum values at the selected points are listed in Table 3: Maximum predicted pressures at selected probe positions in 2 ms to 3 ms range. This pressure dependence on distance from the source is in agreement with experiments.
reported in [55, 56]. A zoom on the minimum acoustic pressures are shown in Figure 11, with minimum pressures confined to 2 bars.

Figure 10: Acoustic pressure predictions at selected points in aluminium domain.
Figure 11: Zoom on the minimum acoustic pressures predicted in the aluminium crucible.

Table 3: Maximum predicted pressures at selected probe positions in 2 ms to 3 ms range.

<table>
<thead>
<tr>
<th>Position</th>
<th>Distance from sonotrode (cm)</th>
<th>Maximum pressure (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>x = 0 cm, y = -2 cm</td>
<td>2.0</td>
<td>9.5</td>
</tr>
<tr>
<td>x = 0 cm, y = -6 cm</td>
<td>6.0</td>
<td>1.4</td>
</tr>
<tr>
<td>x = 0 cm, y = -10 cm</td>
<td>10.0</td>
<td>0.9</td>
</tr>
<tr>
<td>x = 3 cm, y = -2 cm</td>
<td>3.6</td>
<td>2.7</td>
</tr>
<tr>
<td>x = 6 cm, y = -2 cm</td>
<td>6.3</td>
<td>1.6</td>
</tr>
</tbody>
</table>
Figure 12: Variation of maximum pressure with distance from sonotrode and comparison with cavitation intensity from [56].

Figure 12 shows the variation of the maximum pressure with distance from the sonotrode surface. The pressure-distance relationship obeys a power law with an $R^2$ value of 0.95. The decay of pressure with distance is of the order of 1.45 per metre. This pressure dependence on distance is in agreement with qualitative experimental observations with a high-temperature probe [55, 56] plotted on a separate scale, with a decay of the order of 1.28 per metre. The experimental values are quoted in mV since the maximum pressures cannot be recovered from the experimental data. This large decay is expected as the efficiency in acoustic radiation is proportional to the ratio of horn radius to wavelength. The large wavelength in aluminium and the comparatively small sonotrode makes the pressure decrease with distance pronounced.
5. Conclusions

This paper presents a high-order numerical model for predicting acoustic pressures in bubbly liquids subject to acoustic cavitation. This model is computationally attractive for simulations of ultrasonic melt treatment since it is stable for Courant numbers of 0.2 and couples two separate time scales: a bubble dynamics time scale (in the range of $10^{-20}$ s to $10^{-8}$ s) and the flow time scale (or the order of 1 μs). The model has been validated by a water experiment presented in the literature and predicted the reported cavitation characteristics with reasonable agreement between pressure magnitudes, spectra, and decay with distance from the source. When applied to the case of sonication of liquid aluminium, the model predicts a reasonable dependence of acoustic pressure with distance from the ultrasonic source.

Appendix

Spatial discretization coefficients

\[ \Delta_a = 0.047569386 \]
\[ a_1 = \frac{9}{8} + \Delta_a \]
\[ a_2 = -\frac{1}{2} \left( 1 + \frac{1}{12} + \Delta_a \right) \]
\[ a_3 = \frac{1}{10} \Delta_a \]

Temporal discretization coefficients

\[ \Delta_b = 0.08307437 \]
\[ b_0 = 1 + \Delta_b \]
\[ b_1 = \frac{1}{24} - 3 \Delta_b \]
\[ b_2 = 3\Delta_b - \frac{1}{12} \]

\[ b_3 = \frac{1}{24} - \Delta_b \]

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**References**


