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2 A coupled level set and volume of fluid method with a
3 re-initialisation step suitable for unstructured meshes

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8 **Abstract**

9 This paper presents a coupling method of the level set and volume of fluid
10 methods based on a simple local-gradient based re-initialisation approach that
11 evaluates the distance function depending on the computational cell location. If
12 a cell belongs to the interface, the signed distance is updated based on a search
13 in the neighbouring cells and an interpolation procedure is applied depending on
14 the local curvature or the sign of the level set function following [D. Hartmann,
15 M. Meinke, W. Schröder, Differential equation based constrained reinitialisa-
16 tion method for level set methods, J. Comput. Phys. 227 (2008) 6821-6845].
17 The search algorithm does not distinguish between the upwind and downwind
18 directions and hence it is able to be used for cells with an arbitrary number of
19 faces increasing the robustness of the method. The coupling with the volume of
20 fluid method is achieved by mapping the volume fraction field which is advected
21 from the isoface evolution at a subgrid level. Consequently, the coupling with
22 the level set approach is utilised without solving the level set equation. This
23 coupled method provides better accuracy than the volume of fluid method alone
24 and is capable of capturing sharp interfaces in all the classical numerical tests
25 that are presented here.

26 *Keywords:* level set, volume of fluid, re-initialisation

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27 **1. Introduction**

28 *1.1. Scope*

29 In implicit methods for calculating the interface between two fluids flowing
30 in a fixed mesh, the interface is captured using a scalar field advected in space.
31 The scalar field (marker) has to be intrinsically connected to the absence or
32 presence of the liquid phase. These methods are easily extended into three-
33 dimensions but might require fine meshes to resolve the interface. The same
34 limitation holds for front-tracking methods. Here, we are interested in inter-
35 faces for multiphase flows such as bubbles, droplets, and jets for liquid/liquid
36 and liquid/gas interactions. The most commonly cited implicit methods are the
37 volume of fluid method (VOF [1–11] and the level set (LS) methods [12–20].
38 The level set formulation is utilised by transporting a continuous function, as
39 in the VOF method. The level set method has been developed for an accurate
40 representation of complex interface and boundaries for a wide range of appli-
41 cations including among others the areas of shape optimisation [21], computer
42 graphics [22], medical imaging [23], grid generation [24], seismology [25], and
43 superconductors [26]. For fluid interfaces, and particularly in the two-phase
44 flows considered in this paper, the interface of the fluid is defined by the zero-
45 level of a signed distance function and the level set method provides an accurate
46 representation of the curvature of the interface. One common characteristic of
47 this method and VOF is that the user does not need to modify the method re-
48 gardless of the complexity of the geometry since both VOF and level set adjust
49 naturally to any topological changes. One of the main differences between the
50 two methods, is the transition from one fluid to the other, which in the level set
51 method occurs gradually rather than as in the volume of fluid approach where
52 the interface exists in a one-cell layer in between the two fluids.

53 Despite its efficiency in calculating the interface, the level set method has
54 the shortcoming that mass conservation is not guaranteed. This barrier can
55 be overcome by coupling the method with the volume of fluid approach which
56 is conservative, with the level set being highly accurate. This idea was im-

57 plemented first by Bourlioux [27] and Sussman and Puckett [28] giving a new
58 method, the coupled-level-set-volume-of-fluid (CLSVOF) approach. Use of the
59 CLSVOF showed that advecting both the volume fraction and distance func-
60 tions can conserve mass increasing the accuracy of VOF, and providing the basis
61 for different variations of the level set method which have been used in chemical
62 process, aerospace and automotive industries.

63 The coupling of these two approaches does, however pose challenges for the
64 interface reconstruction and the re-initialisation procedures that have to be ad-
65 dressed to successfully simulate fluid flows in the case of three-dimensions or non-
66 orthogonal meshes. In [29] a piecewise-linear interface construction/calculation
67 (PLIC) method is described for advecting the interface, with the level set func-
68 tion used to calculate the curvature. The volume of fluid in a computational cell
69 defines a plane, which is constructed by the intersection points with the cell.
70 The signed distance function is taken as the minimum distance from a finite
71 volume centre to an interface-plane that is defined by a stencil of cells. This
72 VOF-PLIC approach was developed for unstructured meshes in both two and
73 three-dimensions. A similar approach was employed in [30] where the LS-VOF
74 coupling evaluated the level set function from the minimum distance from an
75 arbitrary cell centroid to the zero-level. In addition, no special re-initialisation
76 process was employed, following the geometric operation proposed in [31] (the
77 so-called coupled volume of fluid and level set, a.k.a. VOSET, method) to cal-
78 culate the level set function near the interface. The VOSET method can be
79 applied to accurately compute the curvature and smooth discontinuous physi-
80 cal quantities near the interface for both structured and unstructured meshes.
81 A different LS-VOF coupling suitable for overlapping and moving structured
82 grids was proposed in [32] using a PLIC method for the advection of the volume
83 of fluid approach. The interface was advected using a hybrid split, Eulerian
84 implicit-Lagrangian explicit interface advection scheme which provided good
85 results for the classical test of a deforming three-dimensional sphere. In [33]
86 the idea of flux polygon reconstruction using vertex velocities was employed to
87 evaluate the VOF function. The computed volume fraction was then corrected

88 by a flux corrector estimated using the face velocities. The level set function
89 was advected by a high order total variation diminishing (TVD) scheme and
90 then re-initialised in a narrow band around the interface with a geometric pro-
91 cedure. In [34] the idea of the area of fluid was employed for advecting the
92 volume fraction developing an iterative clipping and capping algorithm for the
93 coupling of the level set and volume of fluid methods. Both the LS and VOF
94 functions are advected by solving a transport equation for each one of them: the
95 volume of fluid is advected employing an interface compression scheme whereas
96 the LS function uses a van Leer TVD scheme. Despite its efficiency in calculat-
97 ing the interface, the LS method has the shortcoming that mass conservation
98 is not guaranteed. This barrier can be overcome by coupling the method with
99 VOF approach which is conservative, and the LS which is highly accurate (see
100 [27] and [28]). In [35] a conservative LS method was developed, which has been
101 demonstrated to conserve mass. This has been the basis of different variations of
102 the LS method which have been used in multiphase flows [34, 36–38]. Coupling
103 the volume of fluid with level set it is possible to combine the benefits of both
104 methods providing an improvement in capturing of the sharp interface with a
105 reasonable accuracy for mass conservation. The ultimate purpose of the correct
106 advection of the level set is the accurate calculation of curvature and mixture
107 properties, in line with the one-fluid approach.

108 *1.2. Objectives*

109 This paper presents a novel coupled LS and VOF method capable of simu-
110 lating the interface of two fluids, of different properties. The first part of the
111 method is the re-initialisation step of the signed distance function. All tradi-
112 tional level set methods face the problem of finding the proper values of the
113 signed distance function, ψ , which satisfy the Eikonal equation, $|\nabla\psi| = 1$. This
114 is usually done by solving the level set equation with a high order approach in
115 time and space to minimise the error, and re-initialising the distance function
116 to avoid the displacement of its initial value ψ_0 [39]. In this paper a partial
117 differential equation re-initialisation method is presented based on the works of

118 Russo and Smereka [40] and Hartmann et al. [41] which allows the simple and
119 efficient calculation of the distance function across the interface. The presented
120 formulation is second-order in space and constructed for computational cells
121 of arbitrary shape, and is tested for both structured and unstructured meshes.
122 The initial value of the distance function ψ_0 in the coupled volume of fluid and
123 level set methods is derived by advecting the volume fraction with either an
124 algebraic or a geometric method. The VOF method for the research presented
125 here, considers the motion of an isoface in a computational cell and advecting
126 it, using the isoAdvector method proposed in [42] and implemented in the open
127 source CFD code OpenFOAM [43]. The isoface is properly advected within a
128 time step, estimating the volume transport across a face before moving on to
129 the next time-step solution. The complete volume fraction advection algorithm
130 is described in detail in the following sections. The coupling of the LS and VOF
131 methods is developed here within OpenFOAM and is done without the need
132 to solve the LS function equation. The approach maps the volume fraction to
133 ψ_0 directly from the VOF step, and then corrects the signed distance function.
134 In order to preserve its distance function character, the level set function is
135 re-initialised in two parts. First, the distance function is calculated for the cells
136 at the interface and is mapped to the level set function. In the second part the
137 re-initialisation equation is solved for the cells adjacent to the cells at the in-
138 terface [39, 41]. Comparisons of the VOF and the coupled LS-VOF for classical
139 numerical tests reveal that the LS step improves the accuracy of solution and
140 boosts the ability of the method to capture sharp interfaces.

141 **2. Motivation and methodology**

142 *2.1. Level set method*

143 The interface which separates the two fluids is represented by the level set
144 function $\psi(\mathbf{x}, t)$. Depending on whether a given point (\mathbf{x}, t) with a normal
145 distance to the interface d , exists in one fluid or the other, $\psi(\mathbf{x}, t)$ is defined as

146 $\psi(\mathbf{x}, t) = +d$, or $\psi(\mathbf{x}, t) = -d$, respectively. The interface Γ is then defined as
 147 the set of points that belong to the zero-level, as follows

$$\Gamma = \{\mathbf{x} | \psi(\mathbf{x}, t) = 0\} \quad (1)$$

148 The level set function is then a distance function that is defined wherever an
 149 interface exists. The distance function can be advected using

$$\frac{\partial \psi}{\partial t} + \mathbf{u} \cdot \nabla \psi = 0 \quad (2)$$

150 where \mathbf{u} is the velocity field. The above equation can be solved using any high
 151 order scheme for hyperbolic systems of the ENO (essentially non-oscillatory)
 152 schemes family or the Runge-Kutta method [44]. A similar advection equation is
 153 used for the marker function in volume of fluid methods. Adding extra algebraic
 154 terms to the right-hand side of this advection term, compresses the interface,
 155 leading to conservative forms of both methods [35, 45].

156 2.2. Re-initialisation step

157 Although the distance function is advected well for $\psi = 0$, it tends to fail to
 158 remain an actual distance function when solving Eq. (2) because of the very
 159 small or large values the magnitude of the gradients $|\nabla \psi|$ might attain on
 160 either side of the interface compromising accuracy [46]. Consequently, a re-
 161 initialisation step is required for the ψ -equation [39]. This is an extra step to
 162 straighten the distorted shape of the function ψ , which might be caused by the
 163 numerical solution of the convection equation or by the complicated fluid veloc-
 164 ity fields. With the re-initialisation step the LS function and the shape of the
 165 interface can be preserved as much as possible throughout the simulation. This
 166 is achieved by solving the following Eikonal equation [15]

$$\frac{\partial \psi_d}{\partial \tau} = \text{sgn}(\psi)(1 - |\nabla \psi_d|) \quad (3)$$

167 where $\text{sgn}()$ is the sign function. The new corrected distance function, ψ_d , is
 168 calculated iteratively knowing ψ which is used as an initial guess in the re-
 169 initialisation process, $\psi_d(t = 0) = \psi$. In Eq. (3), τ is a fictitious time-step

170 which can be associated to the grid resolution. Previous studies suggest vari-
 171 ous values for τ (see [36, 41]) and $\tau = 0.1\Delta x$ is considered in this study. The
 172 re-initialisation equation can be solved in steady state and ideally converges
 173 to $|\nabla\psi_d| = 1$. Previous numerical works have addressed the re-initialisation
 174 problem providing algorithms for either structured [40, 41, 47] or unstructured
 175 computational meshes [48]. In this study, we choose to incorporate and employ
 176 for unstructured meshes the algorithm of [41], which is a partial differential equa-
 177 tion based localised method, which imposes the zero-displacement constraint on
 178 the zero LS. If d_P is the desired signed distance function at the interface for cell
 179 P , then d_P is the value of ψ that has to be used to properly advect the LS func-
 180 tion. An initial guess, \tilde{d}_P of the distance is required during the re-initialisation
 181 step. In [40] a central difference scheme was suggested based upon the knowl-
 182 edge of the upwind or downwind cells of cell P . However, in [41] a modified
 183 expression for \tilde{d}_P was utilised by imposing additional conditions that depend
 184 on how the LS changes along the three co-ordinate directions x, y and z . In
 185 this present study, the distance function is calculated first for the cells at the
 186 interface and is then used to update the level set function. This is done using
 187 first order expressions as proposed in [40]. The Eikonal equation for ψ is then
 188 solved in a narrow band of cells adjacent to the cells that belong to the interface
 189 [41]. The algorithm starts by searching for all the cells at the interface Γ of the
 190 two fluids and performing the re-initialisation step. The algorithm is as follows:
 191 Step 1: For all cells P at the interface Γ , the signed distance function \tilde{d}_P is esti-
 192 mated first using the values of the level set function before the re-initialisation,
 193 denoted for cell P by $\tilde{\psi}_P$. It can be written that

$$\tilde{d}_P = \tilde{\psi}_P / \left[\sum_{\zeta} (\partial_{\zeta} \tilde{\psi}_{\zeta})^2 \right]^{1/2} \quad (4)$$

194 ,where $\partial_{\zeta} \tilde{\psi}_{\zeta}$ is the discrete derivative in the ζ direction [47]. The maximum
 195 between the central and upwind differences can be used for calculating the
 196 derivatives in this expression [40]. In the present study, upwind differences
 197 are considered, employing some of the neighbouring cells. First, a search is

198 performed for all the faces f of the cell P and if a neighbouring cell N that shares
 199 f with P also belongs in Γ , the discrete derivative of $\tilde{\psi}_P$, $\partial_f \tilde{\psi}_P$, is calculated
 200 as:

$$\partial_f \tilde{\psi}_P = \frac{\tilde{\psi}_P - \tilde{\psi}_N}{x_P - x_N} \quad (5)$$

201 Let the number of all these Γ -neighbours be $N_{P,\Gamma}$. Then we use $N_{P,\Gamma}$ discrete
 202 derivatives. In the case of both upwind ($\tilde{\psi}_{i-1}$) and downwind ($\tilde{\psi}_{i+1}$) cells be-
 203 longing to the set Γ , then both differences are considered in the expression for
 204 \tilde{d}_P . Since, $2[(\partial_{i-1} \tilde{\psi}_{i-1})^2 + (\partial_{i+1} \tilde{\psi}_{i+1})^2] \geq [(\tilde{\psi}_{i+1} - \tilde{\psi}_{i-1})/\Delta x]^2$, and given the
 205 fact that we cannot have more than $\lfloor N_{P,\Gamma}/2 \rfloor$ pairs of such central differences,
 206 the sum of all these central differences for cell P in x, y, z (the right part of
 207 the above inequality) will be no greater than the quantity in the denominator
 208 of the signed distance function \tilde{d}_P in Eq.(4), considering all the $N_{P,\Gamma}$ faces e.g.
 209 $N_{P,\Gamma} \left[\sum_f (\partial_f \tilde{\psi}_f)^2 \right]^{1/2}$. For the tests presented here, we use this expression for
 210 calculating \tilde{d}_P . We use these $N_{P,\Gamma}$ derivatives to calculate the target value of
 211 the LS which can be written as:

$$\tilde{d}_P = \frac{\tilde{\psi}_P}{(N_{P,\Gamma} \sum_{k=1}^{N_{P,\Gamma}} (\partial_{f_k} \tilde{\psi}_{f_k})^2)^{1/2}} \quad (6)$$

212 Note that the summation in Eq.(6) is performed through the cell faces and thus
 213 $N_{P,\Gamma}$ is less than or equal to the number of faces of the cell P . The above
 214 expression for \tilde{d}_P is bounded by the special case of a three-dimensional struc-
 215 tured mesh introduced in [41] for Cartesian meshes with pre-defined directions,
 216 x, y, z . The upwind discretisation was employed in [41, 47] and was proven capa-
 217 ble of avoiding any oscillations at the interface that could be caused by central
 218 differences.

219 Step 2: The following step is performed for all the cells in Γ that have
 220 negative curvature κ (calculated from the interface normal \mathbf{n} , $\kappa = \nabla \cdot \mathbf{n}$) or
 221 satisfy the condition $\kappa = 0$ and $\tilde{\psi} \leq 0$, (following [41]). A search is considered
 222 for all the neighbouring cells that share the same face f_k with cell P that have
 223 an opposite sign for ψ_P (for instance, the neighbours N_1, N_2 in Fig.1). Let

224 the number of these cells be M_P , then, the signed distance function and LS
 225 functions $d_P, \tilde{\psi}_P$ for the cell-centre P are interpolated with the same second-
 226 order operator as in [41] as:

$$\begin{aligned}
 (d)_P &= \frac{1}{M_P} \sum_{k=1}^{M_P} d_{f_k} \\
 (\tilde{\psi})_P &= \frac{1}{M_P} \sum_{k=1}^{M_P} \tilde{\psi}_{f_k}
 \end{aligned}
 \tag{7}$$

229 Assuming that the ratio of the LS function with its interpolated value re-
 230 mains the same as the ratio of the signed distance value with its interpolated
 231 value, the position of the zero-level is fixed (a constraint imposed in [41]). The
 232 target value of the LS function is then calculated as [41, 47]

$$d_P = \tilde{\psi}_P \frac{\sum_{k=1}^{M_P} d_{f_k}}{\sum_{k=1}^{M_P} \tilde{\psi}_{f_k}}
 \tag{8}$$

234 where the summation in Eq.(8) is performed over all the M_k cells that are
 235 neighbours of the cell P , at a face f_k with a corresponding d_{f_k} , such that
 236 $\psi_P \psi_{f_k} \leq 0$ (ψ_{f_k} is calculated at the cell centre of the neighbouring cell).

237 Step 3: The LS function for cell P is updated at the interface according to
 238 $\psi_P = d_P$, as suggested by [41].

239 Step 4: The re-initialisation equation is solved for the rest of the cells ad-
 240 jacent to the interface Γ but with a marker function value equal to 0 or 1 (see
 241 Fig.2). Eq. (3) is then solved in steady state:

$$\psi^{n+1} = \psi^n - \Delta \tau S(\tilde{\psi})(|\nabla \psi| - 1)
 \tag{9}$$

242 where $S(\tilde{\psi}) = \tilde{\psi} / \sqrt{\tilde{\psi}^2 + |\nabla \tilde{\psi}|^2 \Delta x^2}$ is a modification of the sign function and
 243 Δx is the cell size. The gradient magnitude is $|\nabla \psi| \cong H_G(D_\zeta^- \psi^n, D_\zeta^+ \psi^n)$ is the
 244 Godunov-Hamiltonian of the LS function from the previous iteration through
 245 all faces of the particular cell. Here, the terms $D_\zeta^- \psi^n, D_\zeta^+ \psi^n$ are the first or-
 246 der approximations of the gradient of ψ along the ζ -direction, $\zeta = x, y$ or z

247 depending on whether the upwind "-" or the downwind cell is considered "+"

$$248 \quad D_{\zeta}^{-} \psi^n = \frac{\psi_i^n - \psi_{i-1}^n}{\Delta \zeta}, \quad D_{\zeta}^{+} \psi^n = \frac{\psi_{i+1}^n - \psi_i^n}{\Delta \zeta} \quad (10)$$

249

250 The need to use first order terms arises from the large gradients across the inter-
 251 face that require an accurate and stable method of calculating H_G . The above
 252 formulation is the basis for various fast marching algorithms with structured
 253 meshes, e.g. [15, 28, 41]. Here, the normal gradient of the level set function ∇_f^{\perp}
 254 is calculated for all the faces f based on the orientation of the normal at the
 255 face. In general

$$\nabla_f^{\perp} = \alpha_{corr}(\psi_P - \psi_n)/|\Delta \zeta| \hat{\zeta} + (\hat{\zeta} - \alpha_{corr} \Delta \zeta) \nabla(\psi)_f \quad (11)$$

256 where α_{corr} is the inverse cosine of the angle between the cell centres and the
 257 normal face, and $\nabla(\psi)_f$ is the linearly interpolated gradient at the face f . This
 258 expression is used for meshes that non-orthogonality is high, and for orthogonal
 259 meshes the second part of the right-hand-side is omitted. Following [34] in the
 260 case of unstructured meshes, one possible approach is to select the upwind cell
 261 from all face-neighbouring cells of the central cell P as the one whose centre
 262 position is closest to the line through the centre and downwind nodes. If the
 263 position vector is $\Delta \zeta$ and the unit vector is denoted as $\hat{\zeta}$, then if $\psi < 0$ and
 264 $\Delta \zeta \cdot \hat{\zeta} < 0$ or $\psi > 0$ and $\Delta \zeta \cdot \hat{\zeta} > 0$ we have:

$$a_{\zeta} = \min \left(\nabla_f^{\perp} \psi \cdot \hat{\zeta} \right) \quad (12)$$

265 If $\psi < 0$ and $\Delta \zeta \cdot \hat{\zeta} > 0$ or $\psi > 0$ and $\Delta \zeta \cdot \hat{\zeta} < 0$ then:

$$a_{\zeta} = \max \left(\nabla_f^{\perp} \psi \cdot \hat{\zeta} \right) \quad (13)$$

266 The Hamiltonian-Godunov term is simply calculated as

$$H_G = \sqrt{\max(a_x^2) + \max(a_y^2) + \max(a_z^2)} \quad (14)$$

267 For the test cases presented here, no significant difference was observed using
 268 an explicit calculation of the gradient of ψ in Eq. (9). This can be performed for

269 unstructured meshes for a cell P with volume $V^{(P)}$, faces f and surface vector
 270 \mathbf{S}_f using:

$$\nabla\psi_d^{(P)} = \frac{1}{V^{(P)}} \sum_f \psi_f \mathbf{S}_f \quad (15)$$

271 The steady solutions of Eq. (9) are distance functions. Furthermore, since
 272 $\text{sgn}(0) = 0$, then $\psi_d(\mathbf{x}, \tau)$ has the same zero-level as ψ .

273 The fictitious time step for the steady state iterations $\Delta\tau$, has to be chosen
 274 so that an accurate value of the LS function is obtained within a reasonable
 275 number of iterations. $\Delta\tau$ can be a fraction of or equal to the mesh size, Δx
 276 [34, 40]. For the simulations performed in this study, $\Delta\tau = 0.1(\Delta x \Delta y \Delta z)^{\frac{1}{3}}$ has
 277 been chosen.

278 A fixed small number of iterations is needed in practice to guarantee the
 279 distance function property near the interface. Following [46], the number of
 280 iterations depends on the thickness of the interface $2\epsilon\Delta x$, with ϵ being the layer
 281 of cells across which the re-initialisation step takes place (the correction is kept
 282 local). The iteration process can stop after 2ϵ time-steps, and a value of $\epsilon = 1.5$
 283 is chosen here so that the interface is spread over a thickness of three cells
 284 (see also Step 4). Finally, for both formulations, Eq.(3) and Eq.(9), the initial
 285 value $\psi_0(\mathbf{x}) = \psi(\mathbf{x}, 0)$ can be taken from the volume fraction α assuming the
 286 interface position is at the iso-surface contour $\alpha=0.5$, (as in [49]) and is written
 287 as a function of the cell size [36].

288 *2.3. Utilising the one-fluid approach*

289 Having calculated the LS function, the mixture properties such as density
 290 and viscosity can be evaluated with the one-fluid approach. The two fluids are
 291 treated as one fluid with properties that change across the interface [7]. To
 292 achieve numerical robustness, a smeared Heaviside function, H , is used [35]
 293 defined as

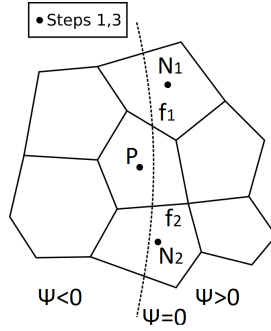


Figure 1: Steps 1-4 performed for a computational cell P in a non-uniform two-dimensional mesh.

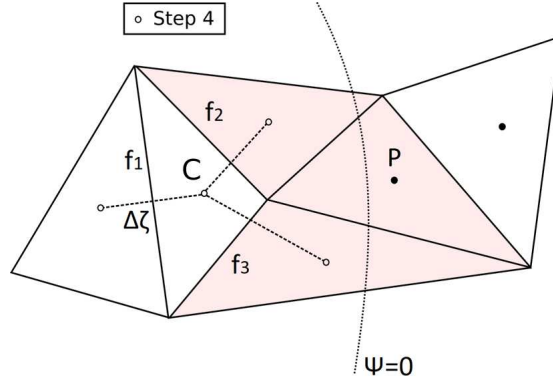


Figure 2: Step 4 performed for a computational cell C that does not belong to Γ .

$$H(\psi) = \begin{cases} 0 & \text{if } \psi < -\epsilon \\ \frac{1}{2} \left[1 + \frac{\psi}{\epsilon} + \frac{1}{\pi} \sin\left(\frac{\pi\psi}{\epsilon}\right) \right] & \text{if } |\psi| \leq \epsilon \\ 1 & \text{if } \psi > \epsilon \end{cases} \quad (16)$$

294 The pseudo-fluid properties can be then calculated as

$$\rho = \rho_1 H + \rho_2 (1 - H) \quad (17)$$

$$\mu = \mu_1 H + \mu_2 (1 - H) \quad (18)$$

295 The surface tension force acting on the interface is calculated as [36, 50]

$$F_\sigma = \sigma \kappa \delta(\psi) \nabla \psi \quad (19)$$

296 Here $\delta()$ is the Dirac function which is employed for limiting the effect of the
 297 surface tension force in the vicinity of the interface, a region with thickness ϵ

$$\delta(\psi) = \begin{cases} 0 & \text{if } |\psi| > \epsilon \\ \frac{1}{2\epsilon} \left[1 + \cos\left(\frac{\pi\psi}{\epsilon}\right) \right] & \text{if } |\psi| \leq \epsilon \end{cases} \quad (20)$$

298 The curvature and the gradient can be discretised (since ψ is a continuous
 299 function) and the surface tension can be calculated at the cell faces using

$$F_\sigma = (\sigma \kappa)_f \delta_f(\psi) \nabla_f^\perp \psi \quad (21)$$

300 2.4. Numerical formulation of coupling Volume of fluid and level set methods

301 The initial value ψ_0 used in Eq. 3 is the starting point for the re-initialisation
 302 iterations and the link with the VOF method. A straightforward approach
 303 without solving the LS equation [36, 49] is to use

$$\psi_0 = (2\alpha - 1) \tilde{\Delta}x \quad (22)$$

304 In the above initial value the $\alpha - 0.5$ iso-surface is used as a starting point for
 305 the signed distance function. The percentage of the interface thickness, $\tilde{\Delta}x$
 306 here equal to $0.8\Delta x$, is introduced for numerical robustness and gives a ψ_0
 307 value within $(-\epsilon, \epsilon)$ for the cells belonging to Γ . The volume fraction α can be
 308 advected with various methods [46, 51–53]. The advection of the volume fraction
 309 depends on the normal to the interface, usually performed employing values in
 310 the neighbouring cells by selecting the orientation of the interface. The coupling
 311 methods used with the VOF approach in [34, 36] use a transport equation for
 312 the volume fraction with the MULES limiter. Even with the addition of an
 313 extra term for compressing the interface, $\alpha(1 - \alpha)U_r$, where U_r is the relative
 314 velocity between the two fluids [45], the interface might still diffuse as previously

315 shown in numerical tests in [30, 42], which can be limited, coupling this scheme
 316 with a level set method [34].

317 The approach in [42] is used to advect the volume and the surface submerged
 318 in one fluid inside each cell at the interface. The idea is that after every time-
 319 step, the iso-surface inside a computational cell (at Γ) splits the cell C with
 320 volume V_C into two different parts: one occupied by fluid 1 with volume frac-
 321 tion α and the other one filled with fluid 2 with volume fraction $(1 - \alpha)$. Let
 322 X_1, X_2, \dots, X_k be the nodes of a cell at the interface, as in Fig.3. The sub-grid
 323 face defined from the line connecting all the intersection points x_m of the iso-
 324 surface and the cell edges, the isoface, is assumed to be advected with a velocity
 325 equal to the velocity of the previous time step and is calculated by solving the
 326 momentum equation. The points x_m at a cell edge $(X_k X_l)$ can be evaluated
 327 from

$$328 \quad x_m = X_k + \lambda_\alpha(X_l - X_k) \quad (23)$$

329 where λ_α is a weight function defined by interpolating at each node X_i the vol-
 330 ume fraction of the surrounding cells. Hence, $\lambda_\alpha = (\alpha - \alpha_k)/(\alpha_l - \alpha_k)$, where
 331 α_k, α_l are the corresponding values at the nodes X_k, X_l . With this linear in-
 332 terpolation the isoface will split the cell into one subcell of fluid 1 with volume
 333 $\Delta V_C(\alpha)$ and another one with volume $V_C - \Delta V_C$ of fluid 2. The isoAdvect
 334 method of Roenby et al. [42] is then performed to find the optimum isovalue
 335 α^* such that $\alpha = \Delta V_C(\alpha^*)/V_C$. A proper value of α^* will cut the cell C into
 336 two subcells with volumetric proportions calculated from the previous guess of
 337 the volume fraction α_C leading to a more accurate reconstruction of the inter-
 338 face than would be obtained using the 0.5-isovalue. The procedure resembles
 339 the re-initialisation step in the correction of the LS method. Using a third
 340 degree polynomial $p_\alpha(x)$ for interpolating the isovalue, we have the following

341 constraints:

$$\begin{aligned}
 & p_\alpha(x_k) = f_k \\
 & p_\alpha\left(x_k + \frac{x_k - x_l}{3}\right) = \frac{f_k}{3} \\
 & p_\alpha\left(x_k + \frac{2(x_k - x_l)}{3}\right) = \frac{2f_k}{3} \\
 & p_\alpha(x_l) = f_l
 \end{aligned}
 \tag{24}$$

343
 344 Finding the polynomial that passes through these points, can then be used to
 345 calculate the isovalue and the isoface which are now defined since both the
 346 vertices and its area, E_m , are known.

347 Next step, is the time evolution of the isoface within a cell. The isoface will
 348 have a centre x_c and velocity \mathbf{u}_c with a normal vector \mathbf{n}_c , as in Fig.3. The
 349 isoface will then pass all the vertices X_k at time t_k which is calculated in [42]
 350 from

$$t_k = t + \mathbf{dx}_i \cdot \frac{\mathbf{n}_c}{\mathbf{u}_c} \tag{25}$$

351 where the vector \mathbf{dx}_k connects the isoface centre with the vertices, $\mathbf{dx}_k =$
 352 $X_k - x_c$. Knowing the time t_k allows calculation of the face-interface line during
 353 the solution time step δt . The submerged area can be integrated using all the
 354 time integrals $[t_k, t_{k+1}]$ such that $t < t_k, t_{k+1} < t + \delta t$. If there are N_{sub} such
 355 time steps then:

$$\int_t^{t+\delta t} E_m d\tau' = \sum_{k=1}^{N_{sub}} \int_{t_k}^{t_{k+1}} E_m d\tau' \tag{26}$$

356 The volume of the fluid with the isoface E_m is easily calculated, since:

$$\Delta V_f^n = \frac{\phi_f}{|\mathbf{S}_f|} \sum_{k=1}^{N_{sub}} \int_{t_k}^{t_{k+1}} E_m d\tau' \tag{27}$$

357 using the volumetric fluxes and the face area at face f , ϕ_f and $|\mathbf{S}_f|$. The volume
 358 fraction is then updated explicitly using the transported volume at each face as

$$\alpha^{t+\delta t} = \alpha^t - \frac{1}{V_C} \sum_{f=1}^{N_f} \Delta V_f^n \tag{28}$$

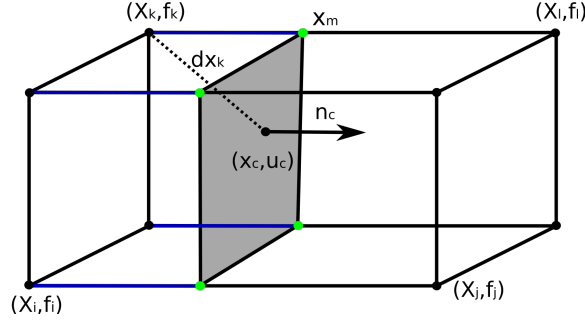


Figure 3: Isoface advection step inside a computational cell. Isoface nodes within the interval $[t_k, t_{k+1}]$ (green points), isoface (shaded face) and face-interface lines (blue).

359 The coupling algorithm of this isoface-level-set-volume-of-fluid (ILSVOF)
 360 method can be described in the following main steps. The numerical fields are
 361 initialised together with the LS function. The dynamic pressure is used to avoid
 362 any sudden changes in the pressure at the boundaries for hydrostatic problems.
 363 The time loop starts by correcting the interface and the volume fraction at the
 364 boundaries. The volume fraction is advected, and corrected, and new values of
 365 α are assigned at the boundaries. The new LS function ψ_0 is calculated using
 366 the results of the advection equation. Next, ψ is re-initialised applying Steps
 367 1-4 described for the re-initialisation procedure, to obtain the signed distance
 368 function, and the interface at the boundaries is corrected. The new interface
 369 curvature is calculated. The mixture properties and fluxes are updated using the
 370 LS function. For instance, for density $\rho = \rho_1 H + \rho_2(1 - H)$ is used, where $H()$
 371 is a Heavyside function of ψ which is used instead of ψ for numerical purposes
 372 in LS methods, [35]. The Navier-Stokes equations are solved for velocity and
 373 pressure using the pressure implicit with splitting of operators (PISO) method.
 374 The process starts again, by first correcting the interface and the volume fraction
 375 at the boundaries and then, following the described steps after that.

376 **3. Numerical tests and discussions**

377 The performance of the proposed numerical methodology is tested using
378 simple test problems involving two fluids of different densities and viscosities.
379 The problems, in both two and three-dimensions, include comparisons between
380 the presented method and other numerical works with or without the level set
381 implementation. Different indicators are used for monitoring the method’s per-
382 formance in terms of shape conservation and boundedness. The numerical tests
383 presented here concern both structured and unstructured meshes and different
384 mesh resolutions are examined for the presented test cases.

385 *3.1. Two-dimensional rotating disc*

386 The rotating disc has been proposed in [54] and considers a disc that under-
387 goes a significant interface deformation and is used here to evaluate the abil-
388 ity of the presented methodology to transport under-resolved interface struc-
389 tures [6, 9, 33, 55]. The computational domain is a unit square which con-
390 tains a disc with radius R placed in the domain so that the disc centre is at
391 $(x, y) = (0.5, 0.75)$. The rest of the domain is filled with a fluid of lower viscosity
392 than the one of the disc. The velocity field is given by

$$u(x, y, t) = -\sin^2(\pi x)\sin(2\pi y)\cos\left(\frac{\pi t}{T}\right) \quad (29)$$

$$v(x, y, t) = \sin(2\pi x)\sin^2(\pi y)\cos\left(\frac{\pi t}{T}\right) \quad (30)$$

393 The velocity field changes in time and space and causes the disc to rotate so that
394 the initially circular disc is stretched with time (the flow lasts for one period T).
395 The resulting filament will then stretch until $t = T/2$ at which point the velocity
396 field reduces to its minimum and becomes zero, according to Eq. (29). For the
397 simulations here, a period of $T = 8$ was used with a total simulation time of $8s$.
398 The physical parameters considered are summarised in Table 1. Two different
399 types of mesh have been used, quadrilateral and triangular, as the ones shown
400 in Fig. 4 each using a coarse, medium and a fine meshes respectively. The

401 results for the new method are shown in Tables 2 and 3. A comparison of
 402 the ILSVOF methodology described here, with the case of using only the VOF
 403 method [42] without the level set, is also included. Previous studies in [42]
 404 have shown that the volume of fluid method of [42] provides higher accuracy for
 405 interface advection than the volume of fluid approach of [45] based on, hence
 406 only the error in shape preservation reported in [42] is included for comparisons
 407 in Tables 2 and 3. Some results obtained using only [45] approach for two and
 408 three-dimensional cases are presented for qualitative comparisons in Fig. 5. The
 409 error

$$E_\alpha = \frac{\sum_i V_i |\alpha_i - \alpha_{exact}|}{\sum_i V_i \alpha_{exact}} \quad (31)$$

410 is used as a quantitative measure of the shape preservation. Here, the exact
 411 solution is defined as the initial position of the rotating disk and E_α is calculated
 412 over all cells of the domain. The extent to which the solution stays within the
 413 range of physical values is also tested, considering the minimum and maximum of
 414 the liquid volume fraction of the rotating disc, $\min(\alpha)$ and $\max(\alpha)$ respectively,
 415 also calculated over all cells of the domain. One should expect ideally to have
 $0 \leq \alpha \leq 1$ for all cells.

Table 1: Physical properties for rotating disc simulations.

	Physical and numerical parameters
Heavy fluid density	1110 kg/m^3
Light fluid density	806 kg/m^3
Heavy fluid kinematic viscosity	1.017e-06 m^2/s
Light fluid kinematic viscosity	2.35e-06 m^2/s
Length of square domain	1 m
Period T	8 s
CFL number	0.3

416

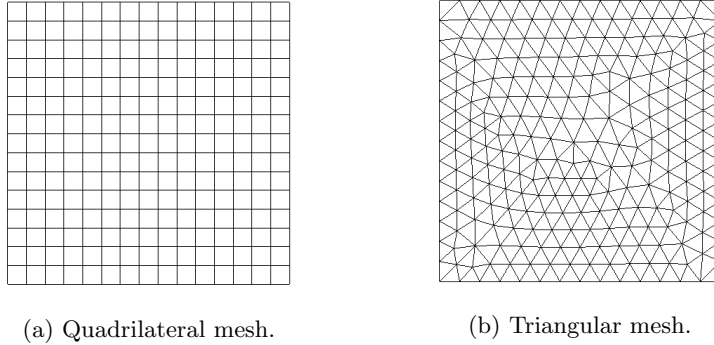


Figure 4: Meshes used to discretise the domain used (coarse mesh study).

417 As indicated in the results of Fig. 5, increasing the mesh resolution, increases
 418 the sharpness of both methods with E_α decreasing. The mesh resolution is
 419 a important parameter that influences the ability of the interface capturing
 420 method to resolve the ligament stretching. The sharp tail at the end of the
 421 deforming ligament fails to be resolved at the subgrid scale. In all cases, at
 422 the time of maximum stretching, $t = T/2$, the rotating spiral thickness becomes
 423 equivalent to the local cell size and fragmentation starts to be noticeable (Fig. 5).
 424 The observed volume sharpness error was generally of the same order as that
 425 obtained from the other approaches, or smaller. The same behaviour is observed
 426 in boundedness were the marker function, α , stays above the minimum value of
 427 0 and below or equal to 1, contrary to the solution obtained without the level
 428 set step. The ILSVOF method maintains similar trends for the error measures
 429 for both structured and unstructured meshes (Table 3). A comparison between
 430 the presented ILSVOF and the VOF methods is shown for $t = T/2$ in Fig. 6 for
 431 the case of the medium size quadrilateral meshes using the OpenFOAM VOF
 432 methods of [42] and [45] which employs the multidimensional universal limiter
 433 with explicit solution (MULES) scheme [56]. As the rotating vortex thickness
 434 becomes progressively smaller, the droplets pinch off and the local interface
 435 curvature becomes of the same order as the mesh size. The decreasing interface
 436 curvature causes the isoface, used here to advect α , to become less accurate for

437 the resulting droplet motion. The coupling with level set shows that it is possible
438 to have less diffusion at the tip for the rotating filament for both quadrilateral
439 and triangular meshes (Fig. 7). The gradients appearing during the deformation
440 of the vortex might have an effect on shape preservation, via Eq. (6), even though
441 the vortex reverses after $t = T/2$ back to its initial position, but these do not
442 seem to reduce the solution accuracy, at least for the re-initialisation steps used
443 here which usually varied from 2 to 5. The ILSVOF retains the vortex shape
444 better than the other methods at the maximum stretching position. Using
445 the interface compression scheme MULES, sharpness can also be maintained
446 at $t = T/2$, but this interface compression might cause the rotating spiral to
447 break-up, fragmenting the vortex in the interval $[T/2, T]$ which is more evident
448 at $t = T$ in Fig. 6 and Fig. 7. The increased error in the averaged volume
449 fraction for this approach is also reported in previous studies in [34], and is
450 observed here in both structured and unstructured methods.

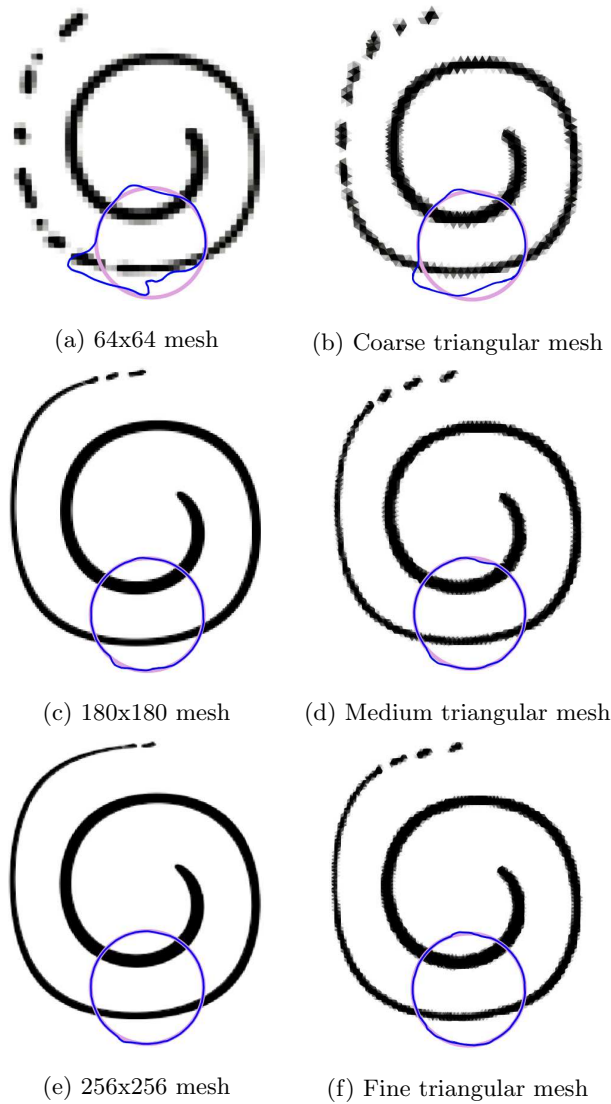


Figure 5: Two-dimensional rotating disc test for level set-VOF method at $t = T/2$. The initial (light purple line) at $t = 0$ and final position of the zero-level set iso-surface (blue line) at $t = T$ are indicated.

451 Comparisons with different grid resolutions ($32^2, 64^2, 128^2$) are shown in Ta-
 452 ble 4 for the values of the calculated $L_1(\alpha)$ error norm for the volume fraction.
 453 Results for $L_1(\alpha)$ from other numerical works are included for comparisons and

Table 2: Comparison of the methods using quadrilateral meshes for the two-dimensional rotating disc case.

Mesh resolution	ILSVOF method			VOF method		
	E_a	$\min(\alpha)$	$\max(\alpha)$	E_a	$\min(\alpha)$	$\max(\alpha)$
64^2	-1.03×10^{-7}	0.0	1.0	3.30×10^{-7}	0.0	1.0
180^2	-1.58×10^{-8}	4.60×10^{-9}	1.0	1.2×10^{-8}	-2.88×10^{-8}	$1.0-1.18 \times 10^{-8}$
256^2	-9.61×10^{-9}	-7.16×10^{-7}	$1.0-1.5 \times 10^{-9}$	-2.48×10^{-8}	-3.34×10^{-5}	$1.0-3.64 \times 10^{-8}$

Table 3: Comparisons of the methods using triangular meshes for the two-dimensional rotating disc case.

Mesh resolution	ILSVOF method			VOF method		
	E_a	$\min(\alpha)$	$\max(\alpha)$	E_a	$\min(\alpha)$	$\max(\alpha)$
17521	-2.77×10^{-7}	0.0	1.0	7.26×10^{-7}	-1.14×10^{-8}	$1.0-9.61 \times 10^{-10}$
79877	3.91×10^{-8}	0.0	1.0	-4.34×10^{-7}	0	1.0
108151	2.11×10^{-8}	0.0	1.0	-1.68×10^{-8}	-9.2×10^{-8}	$1.0-4.78 \times 10^{-8}$

454 the domain and physical properties for the rotating disc test are set as those
455 in [57]. The method presented here, overall demonstrated good accuracy for
456 the two-dimensional rotating vortex case. Compared to other volume of fluid
457 methods that use PLIC such that in [6] the error in $L_1(\alpha)$ is lower for all the
458 meshes tested here. Compared to the tangent of hyperbola for INterface cap-
459 turing, (THINC) scheme and its variations [57–59], the results here are similar
460 or lower. The $L_1(\alpha)$ error is close for the coarse mesh compared to the level
461 set method in [60], but had lower values for the medium and fine meshes. The
462 mass conservation error through time is shown in Fig. 8. The method showed
463 generally reasonable mass conservation error for the different meshes that were
464 used, as shown in Table 4.

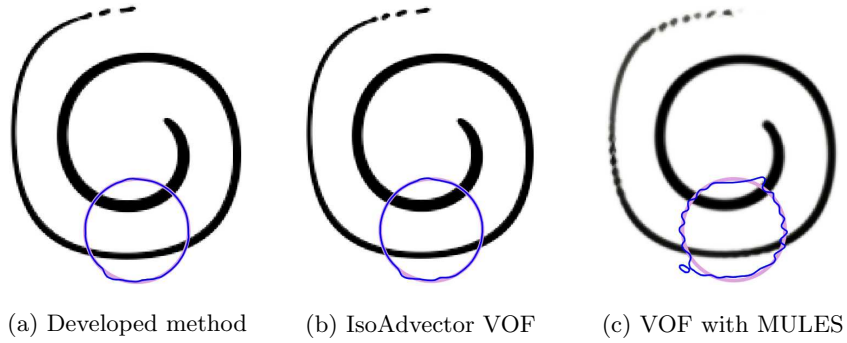


Figure 6: Two-dimensional rotating disc test for the medium quadrilateral mesh at $t = T/2$. The initial (light purple line) at $t = 0$ and final position of the zero-level set iso-surface (blue line) at $t = T$ are indicated.

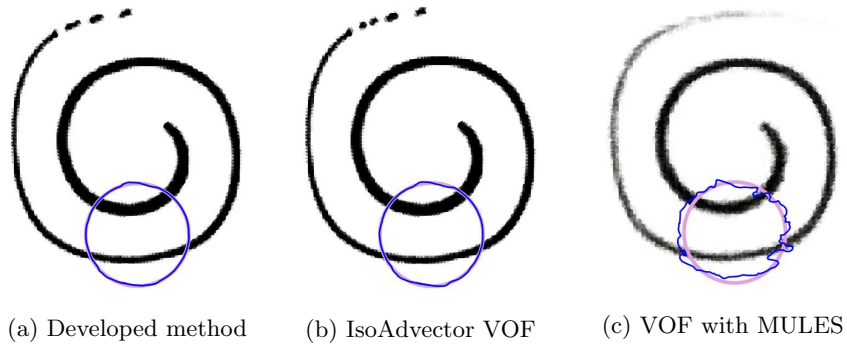


Figure 7: Two-dimensional rotating disc test for the fine triangular mesh at $t = T/2$. Initial (light purple line) at $t = 0$ and final position of the zero-level set isosurface (blue line) at $t = T$ are indicated.

Table 4: Comparisons of the methods using quadrilateral meshes for the two-dimensional rotating disc case. The first order norm $L_1(\alpha)$ is calculated for the three different meshes.

Authors	32^2	64^2	128^2
RiderKothe/Puckett [61]	4.78×10^{-2}	6.96×10^{-3}	1.44×10^{-3}
THINC/WLIC [57]	4.16×10^{-2}	1.61×10^{-2}	3.56×10^{-3}
Markers-VOF [62]	7.41×10^{-3}	2.78×10^{-3}	4.78×10^{-4}
DS-CLSMOF [60]	2.92×10^{-2}	5.51×10^{-3}	1.37×10^{-3}
PLIC [6]	2.53×10^{-2}	2.78×10^{-3}	4.8×10^{-4}
THINC/QQ [58]	6.70×10^{-2}	1.52×10^{-2}	3.06×10^{-3}
THINC/SW scheme [59]	3.90×10^{-2}	1.52×10^{-2}	3.96×10^{-3}
ISLSVOF method	4.19×10^{-2}	1.43×10^{-3}	8.36×10^{-4}

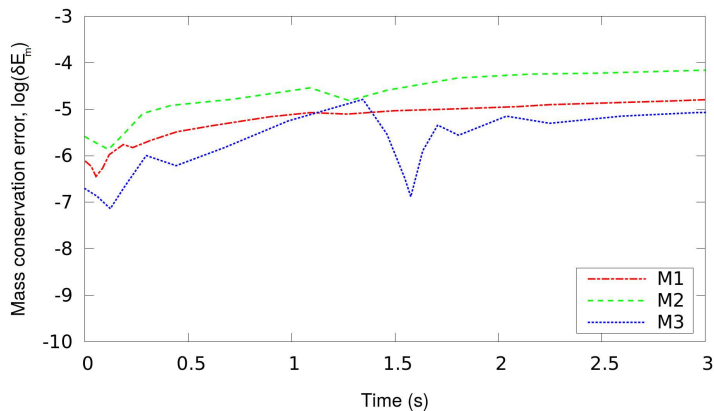


Figure 8: Mass conservation error for the two-dimensional rotating disc case for the three levels of refinement. Here, the meshes M_1 , M_2 and M_3 had 32^2 , 64^2 and 128^2 cells respectively.

465 *3.2. Three-dimensional rotating sphere in a non-uniform velocity field*

466 The next test is the case of a three-dimensional rotating sphere of [54] and
 467 is used to assess the capability of the methodology for capturing the interface
 468 distortion in three dimensions. [34, 36, 42, 63]. In this test, a sphere with radius
 469 $R = 0.15m$ is placed inside a box $[0, 1]^3$ with its centre at $(0.35, 0.35, 0.35)$. The

470 velocity field is defined as

$$\begin{aligned}
 u(x, y, z, t) &= 2\sin^2(\pi x)\sin(2\pi y)\sin(2\pi z)\cos\left(\frac{\pi t}{T}\right) \\
 v(x, y, z, t) &= -\sin(2\pi x)\sin^2(\pi y)\sin(2\pi z)\cos\left(\frac{\pi t}{T}\right) \\
 w(x, y, z, t) &= -\sin(2\pi x)\sin(2\pi y)\sin^2(\pi z)\cos\left(\frac{\pi t}{T}\right)
 \end{aligned}
 \tag{32}$$

473 The period is $T = 3s$, and the density and viscosity of both fluids in the test,
 474 are the same as in the previous section (see Table 1). The sphere is rotating
 475 within the non-uniform velocity field which causes the sphere to deform through
 476 time during $[0, T/2]$. At $t = T/2$ the flow reverses due to the sign change of the
 477 cosine parameter during $[T/2, T]$ causing the deformed sphere to return back
 478 to its original position at $t = T$. The ILSVOF method was first tested for
 479 three different meshes (with $40^2, 64^2, 100^2$ elements) to assess the error in shape
 480 preservation and the boundedness of the marker function. The results given in
 481 Table 5 compare the method with and without the level set step using structured
 482 meshes. Overall, the error E_α decreases with the level set implementation. A
 483 slight increase in E_α is observed for the fine mesh with respect to the medium
 484 mesh, although this is relatively insignificant. Similar trends were observed by
 485 previous authors using LS and VOF method coupling [34]. A more detailed
 486 comparison is given in Fig. 9 at the maximum deformation time, $t = T/2$ for
 487 different grid resolutions ($40^2, 64^2, 100^2, 128^2$ elements). The deforming sphere
 488 appears to be thicker in the case of the ILSVOF method which provides more
 489 detail for the deforming sphere for the different levels of mesh refinement. The
 490 sharp sphere end is also thickened in the ILSVOF case, and the thickening
 491 appears to be more evident for the finer meshes. The time evolution of the
 492 deforming sphere inside the non-uniform flow is shown for different times within
 493 $[0, T]$ in Fig. 10 for a 128^3 mesh. Numerical results revealed as before, relatively
 494 large gradients that the LS function experiences in $[0, T/2]$ which are maintained
 495 and are not reversed in the interval $[T/2, T]$ giving a perturbed profile at the final
 496 position of the sphere. The sphere interface is distorted in all cases as shown in
 497 Fig 11 at $t = T$, with or without level set or interface compression. In general,

498 the ILSVOF method shows better representation of the surface. The most
 499 significant surface distortion is observed in the case of the interface compression
 500 where the compression of the interface decreases the solution accuracy [34]. The
 501 three-dimensional sphere in the same non-uniform flow was also simulated using
 502 unstructured meshes and the results are shown in Table 6. The values of E_α
 503 are higher for the medium and fine meshes compared with those in Table 5 for
 504 structured meshes. The sheet thickness appears to be smaller than the average
 505 edge length even for the finest tetrahedral mesh (~ 0.008), causing the solution
 506 accuracy to drop in the case of unstructured meshes. In addition, the impact of
 507 the steep gradients introduced in the LS method that originate from the initial
 508 value of ψ_0 in Eq. 22 being inserted in Eq. 3 are more evident in the case of
 509 tetrahedral meshes. The L_1 error for the volume fraction for different mesh
 510 resolutions ($32^3, 64^3, 128^3$) for the three-dimensional case of the rotating sphere
 511 are shown in Table 7. Following [56, 59] the error L_1 is calculated for all cells i
 512 and is defined as

$$L_1(\alpha) = \sum_i (\alpha_i - \alpha_{exact}) V_i \quad (33)$$

513 The results are shown alongside with the L_1 error obtained with other volume of
 514 fluid methods. The accuracy of the results here, remained lower than the volume
 515 of fluid with the THINC/SW scheme which uses no geometrical reconstruction
 516 [59] and was more accurate compared to the results obtained in [56] with the
 517 MULES limiter with interFoam. In all cases the L_1 error was very close to the
 518 results from the PLIC VOF method in [64].

Table 5: Comparison of the methods using quadrilateral meshes for the three-dimensional rotating sphere case.

Grid	ILSVOF method			VOF method		
	E_a	$\min(\alpha)$	$\max(\alpha)$	E_a	$\min(\alpha)$	$\max(\alpha)$
40^3	5.43×10^{-7}	-4.05×10^{-17}	1.0	5.84×10^{-7}	-1.87×10^{-16}	1.0
64^3	-9.85×10^{-8}	-1.48×10^{-16}	1.0	-1.04×10^{-7}	-1.53×10^{-9}	$1.0-5.93 \times 10^{-11}$
100^3	1.79×10^{-7}	0.0	1.0	3.22×10^{-7}	0.0	1.0

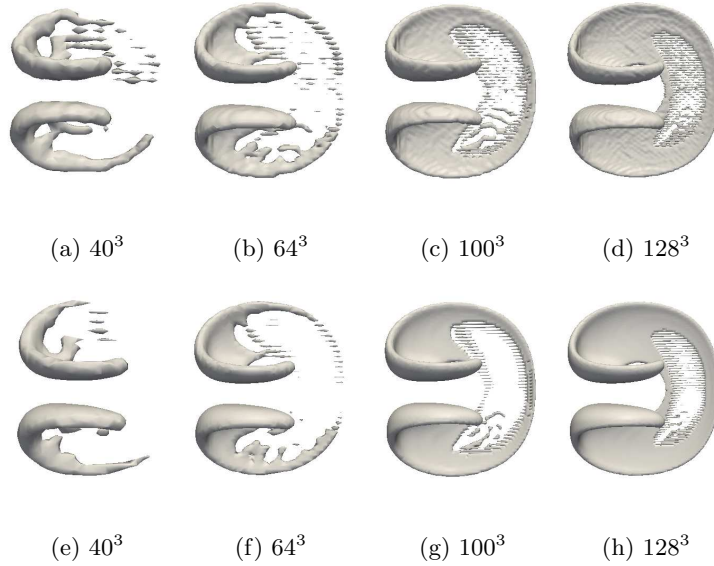


Figure 9: Three-dimensional rotating sphere in a non-uniform flow test for various levels of mesh. The 0.5-iso-surface obtained with the ILSVOF method (top) and without the level set step (bottom) are shown at the maximum deformation time, $t = T/2$.

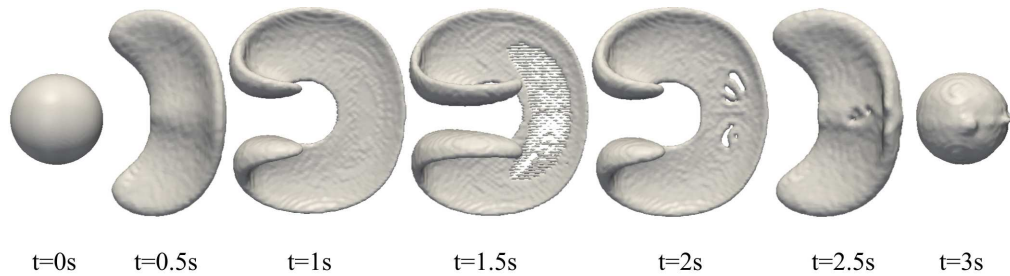


Figure 10: Three-dimensional rotating sphere in a non-uniform flow during time evolution in $[0, T]$ for a fine hexahedral mesh of 128^3 hexahedra.



(a) Developed method (b) Isoface method without (c) Interface compression
 ILSVOF. level set. scheme, MULES.

Figure 11: Three-dimensional rotating sphere in a non-uniform flow test using the 128^3 hexahedral mesh. Initial transparent blue surface at $t = 0$ and final position of the zero-level set iso-surface solid grey iso-surface at $t = T$.

519 *3.3. Three-dimensional dam break case without obstacle*

520 The dam break problem is studied next which consists of a simple three-
 521 dimensional rectangular geometry wherein a liquid column is initially held still
 522 by a dam. When the dam is suddenly removed, the liquid column collapses. The
 523 tank containing the liquid column which collapses in this case is a rectangular

Table 6: Comparison of the methods using tetrahedral meshes for the three-dimensional rotating sphere case.

Mesh resolution	ILSVOF method			VOF method		
	E_a	$\min(\alpha)$	$\max(\alpha)$	E_a	$\min(\alpha)$	$\max(\alpha)$
163208	-6.42×10^{-8}	0	1.0	2.68×10^{-8}	0.0	1.0
322676	-8.7×10^{-3}	0.0	1.0	-3.78×10^{-3}	0.0	1.0
1083126	-2.3×10^{-4}	0.0	1.0	-7.8×10^{-3}	0.0	1.0

Table 7: $L_1(\alpha)$ error norm for different meshes for the three-dimensional rotating sphere case and comparison with other numerical methods.

Authors	32^3	64^3	128^3
RK-3D using PLIC [64]	7.85×10^{-3}	2.75×10^{-3}	7.41×10^{-4}
THINC/SW scheme [59]	8.39×10^{-3}	3.47×10^{-3}	1.08×10^{-3}
interFoam [56]	9.95×10^{-3}	4.78×10^{-3}	2.03×10^{-3}
ISLSVOF method	8.89×10^{-3}	2.96×10^{-3}	8.06×10^{-4}

524 domain with dimensions $4a \times 2.4a \times a$. For a more convenient comparison,
 525 the fluid in the liquid column is assumed to be water and the rest of the tank
 526 is filled with air. Both fluids are assumed to be initially still, and the physical
 527 properties of the three-dimensional case are shown in Table 8. Initially the liquid
 528 column has dimensions $a \times 2a \times a$. The vertical acceleration due to gravity is
 529 taken to be 9.81 ms^{-2} . The velocity before removing the dam, is zero for the
 530 liquid column and the air, and the pressure is set to be the hydrostatic pressure.
 531 Free slip boundary conditions are imposed for all the boundaries of the domain
 532 (assuming zero normal velocity and zero tangential traction) except for the
 533 open top boundary (where the tangential velocity and normal traction are zero).
 534 The displacement of the interface between water and air is tracked in order to
 535 characterise the performance of the developed method using three different grids
 536 ($40 \times 10 \times 20$, $80 \times 20 \times 40$ and $160 \times 40 \times 80$). The results are compared with
 537 the experimental data available in [65] and previous numerical studies using
 538 the LS method from [66] which is a conservative level set method based on the
 539 finite element approach which employs the volume fraction for correcting the
 540 distance function. The results for the position of the water-air interface along
 541 the horizontal (x-axis) and the vertical (y-axis) directions are shown in Fig. 12.
 542 The results for the position are normalised with the length parameter a and are
 543 plotted against the non-dimensional time. The predictions for the leading-edge
 544 position are in good agreement with the experimental data for the examined
 545 time interval. The accuracy in the predictions for the horizontal direction is
 546 closer initially and reasonably close to the experiment during the simulation
 547 until the leading edge reaches the wall, $x/a = 4$. The flow slows down as a
 548 result of wall friction as reported in the experimental work and, as a result, the
 549 calculated interface is expected to differ from the experiment, although this is
 550 less than five to ten percent here for the fine and coarse meshes respectively. The
 551 interface in [66] reaches the position $x/a = 4$ faster than the present simulations
 552 here. In the method presented here, the liquid front propagates slower than in
 553 [66] although the results in [66] for a coarse mesh (not shown here) had similar
 554 trends as for the results obtained here for the different grids. The coarse grid

555 results in the present study are close to the liquid front results reported in [65],
 556 with the results obtained with the other two meshes also being reasonably close
 557 to the experiment. The time evolution of the interface is shown in Figs.13. The
 558 shape of the interface remains almost flat for the considered time steps of the
 559 experiment, and in Fig.13(a-c). Once the collapsing liquid column reaches the
 560 wall, the water rises upwards forming a layer on the right wall. During the time
 561 interval $[0, 0.26s]$ the vertical position of the interface decreases as expected, and
 562 both the present method and the results of Kees et al.[66] are seen to be very
 563 close in Fig. 12b, having the same rate of change in the liquid column height
 564 for the different grid resolutions. The fluid percent mass loss evolution for the
 565 dam break case is shown in Fig. 14. The mass loss approaches zero for the fine
 566 mesh and remains less than 0.01 percent for the coarse mesh.

Table 8: Physical properties for the dam break case.

	Physical and numerical parameters
Water density	1000 kg/m^3
Air density	1.0 kg/m^3
Water kinematic viscosity	1.0e-06 m^2/s
Air kinematic viscosity	1.0e-04 m^2/s
Length a	0.146 m
CFL number	0.5

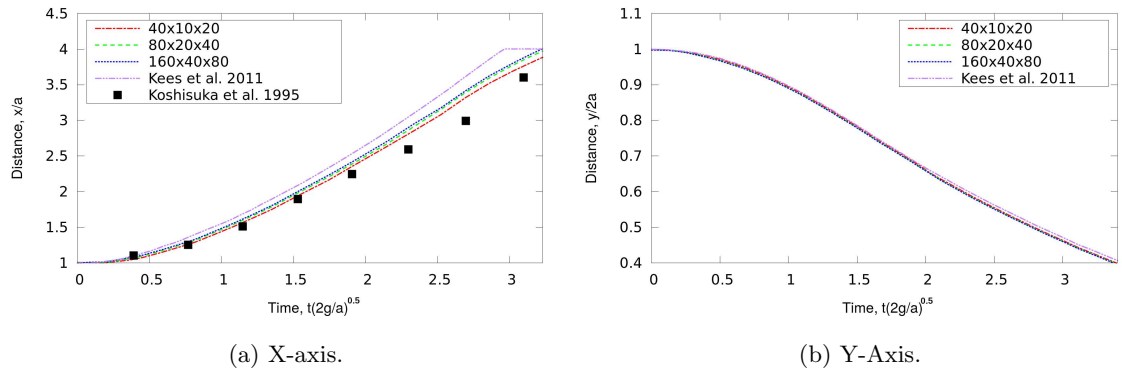


Figure 12: Water-air interface position along the x-axis and y-axis for the three-dimensional dam break case without obstacle.

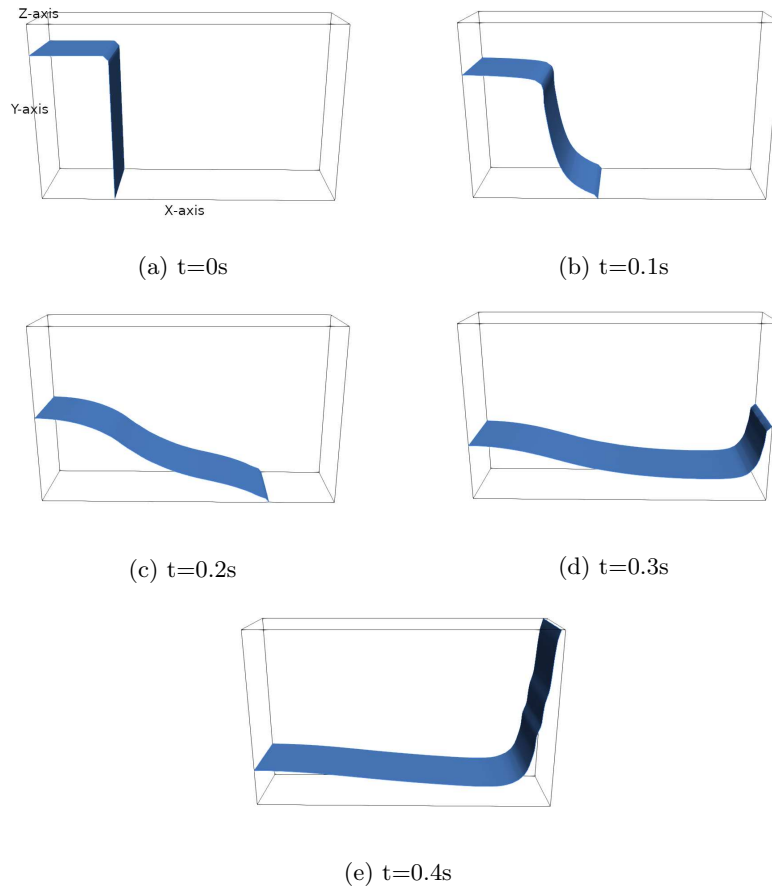


Figure 13: 0.5-isosurface snapshots at different times for the three-dimensional dam break case without obstacle. The liquid column starts to collapse at $t=0s$ and moves towards the right wall until it impinges and rises up forming a layer that keeps moving upwards until it returns back to the tank due to gravity.

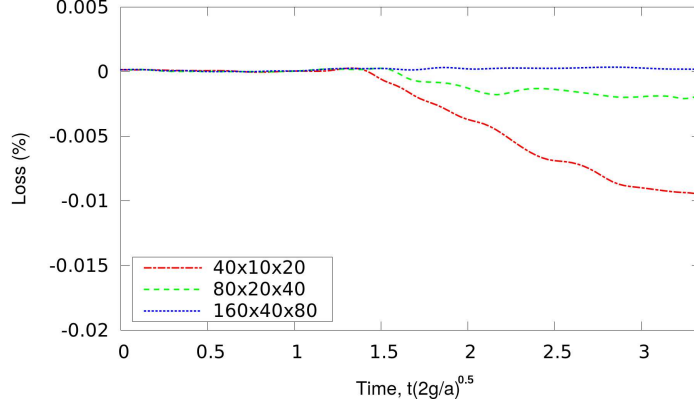


Figure 14: Percent mass loss for the dam break case through time for different mesh resolutions.

567 3.4. Static drop

568 In this test case we are interested in the verification of the methodology
569 for the stationary Laplace solution for a droplet inside a closed domain and
570 assessing the spurious currents [29, 67]. Neglecting any gravitational effects
571 and any external forces the interface between the drop and the ambient fluid is
572 expected to remain at rest. The surface tension force ($\sigma\kappa$) is balanced by the
573 pressure force at the interface according to Laplace's law: $\Delta p_{exact} = \sigma\kappa_{exact}$,
574 where the exact interface curvature is $\kappa = 2/d$ and σ is the surface tension. For
575 a constant pressure outside the droplet p_0 and zero velocity, the pressure inside
576 remains constant and equal to $p_0 + 2\sigma/d$. Due to spurious currents the calculated
577 pressure will differ. A $2d \times 2d$ domain was used for the present numerical tests
578 with $d = 0.5cm$ and the density ratio between the droplet and the surroundings
579 was 10^4 . The viscosities inside and outside were equal to 1 and the surface
580 tension was $1kg \cdot s^{-2}$. For the two-dimensional tests, three meshes of triangular
581 elements were used (with grid size $\Delta x = 1/25, 1/50, 1/100$ as in [29]). In order
582 to evaluate the parasitic currents in the domain, the L_1 error norm is calculated
583 as

$$L_1(\mathbf{u}) = \frac{1}{N_c} \sum_i (\mathbf{u}\mathbf{u})^{1/2} \frac{\mu}{\sigma} \quad (34)$$

584 where the summation is done over the entire domain as in [29] and [68]. The
 585 pressure ranges from p_{out} (pressure outside the droplet) to the inside pressure
 586 (p_{in}). The pressure error used to evaluate the pressure jump denoted by $E(\Delta p)$,
 587 is

$$E(\Delta p) = \frac{|p_{in} - p_{out} - 2\sigma/d|}{2\sigma/d} \quad (35)$$

588 The calculated values for L_1 and $E(\Delta p)$ for the three meshes are shown in
 589 Table 9. The results are also compared to the ones obtained using the VOF
 590 only without using level set and the CLSVOF in [29] using PLIC. The parasitic
 591 currents obtained here are close to those in [29] and smaller than the VOF ap-
 592 proach. In all cases the error L_1 decreases when increasing the mesh resolution.
 593 Similar behaviour is observed for the $E(\Delta p)$ which remained smaller than the
 594 calculated error values for pressure jump reported in [29] for the different grid
 595 resolutions. Fig.15 shows the pressure jump for the three different meshes com-
 596 pared to the exact solution (normalised with the maximum pressure difference
 597 Δp_0). Overall the calculated pressure is close enough to the exact value.

598 Another values that are also used for the static drop test are the L_1 error
 599 norm for pressure, $L_1(p)$ and the maximum velocity after one and fifty time-
 600 steps $U_{max,1}, U_{max,50}$. The drop density for the test is set to $1000kgm^{-3}$ with
 601 a density ratio with the ambient gas phase equal to 1000. The rest of the
 602 properties for the drop and the outside are set as in [68]. For this test, the
 603 drop has a radius $R = 2cm$ and is placed at the centre of a $6cm \times 6cm$ square
 604 domain. Three different grid resolutions were used for this case, with grid size
 605 Δx such that $R/\Delta x = 10, 20, 40$ and the results are shown in Table 10. The
 606 order for the $L_1(p)$ error remains at the order of 10^{-4} or below and the accuracy
 607 of the presented method was close to the CLSVOF works in [68] and [55]. The
 608 unphysical velocity fields that occur in the areas where pressure changes are
 609 monitored using the maximum velocity U_{max} . In this study U_{max} was of order

Table 9: Comparisons of the methods using unstructured meshes for the two-dimensional static drop case.

Mesh resolution	ILSVOF method		VOF method		CLSVOF using PLIC [29]	
	$L_1(\mathbf{u})$	$E(\Delta p)$	$L_1(\mathbf{u})$	$E(\Delta p)$	$L_1(\mathbf{u})$	$E(\Delta p)$
1/25	1.24×10^{-4}	0.011	5.84×10^{-3}	0.026	1.3×10^{-4}	0.02433
1/50	9.81×10^{-6}	0.0046	3.76×10^{-5}	0.0078	3.19×10^{-5}	0.00651
1/100	7.46×10^{-6}	0.0017	6.61×10^{-5}	0.0049	8.82×10^{-6}	0.00215

610 of 10^{-8} for the coarser mesh and was higher for the finer meshes, at the order of
 611 10^{-7} as in [67]. Similar patterns for higher parasitic currents while increasing
 612 the mesh resolution were also reported before in [68], [55] and [67] and has been
 613 reported for different VOF methods which employ the continuous surface force
 614 model [50].

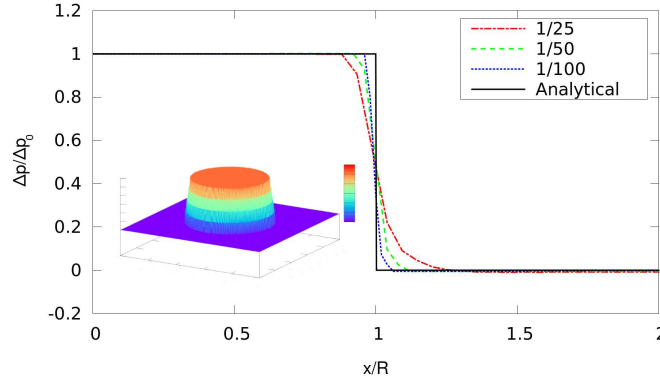


Figure 15: Distribution of the pressure for the static drop test case. Three different mesh resolutions are used to capture the pressure jump across the interface. The pressure is normalised with the exact value Δp_0 and the distance with the droplet radius R . The distribution of the pressure for the finest mesh is also shown.

Table 10: Convergence study for the calculation of pressure, pressure error and velocity for the static drop test. Three different mesh resolutions are used and the density ratio was 1000 on three different grids. Results are compared with different numerical studies.

Authors	$R/\Delta x$	$L_1(p)$	$ u_{max,1} $	$ u_{max,50} $
Gerland et al. 2006 [68]	10	4.81×10^{-3}	7.82×10^{-8}	3.91×10^{-6}
	20	9.48×10^{-4}	1.70×10^{-7}	8.53×10^{-6}
	40	7.04×10^{-5}	4.34×10^{-7}	2.17×10^{-5}
Ningegowda et al. 2014 [55]	10	1.14×10^{-2}	1.12×10^{-6}	5.11×10^{-5}
	20	7.53×10^{-3}	5.88×10^{-6}	3.10×10^{-4}
	40	2.92×10^{-3}	1.30×10^{-5}	9.43×10^{-4}
Jarauta et al. 2018 [67]	10	1.25×10^{-4}	6.08×10^{-9}	4.09×10^{-7}
	20	3.12×10^{-4}	4.38×10^{-8}	1.22×10^{-6}
	40	7.85×10^{-5}	4.26×10^{-7}	3.55×10^{-6}
ISLSVOF method	10	7.36×10^{-4}	2.36×10^{-8}	8.89×10^{-6}
	20	1.66×10^{-4}	5.41×10^{-8}	3.46×10^{-6}
	40	8.21×10^{-5}	7.22×10^{-7}	1.17×10^{-6}

615 *3.5. Rising bubble*

616 The final test case is the rising bubble test proposed in [69]. A circular
 617 bubble is initially placed in a column filled with fluid of higher density than
 618 the density of bubble. Due to the buoyancy force, the bubble rises and deforms
 619 while moving towards the top of the column. The bubble diameter is initially
 620 $d = 0.5$ units and is centred at $(0.5, 0.5)$ in a rectangular domain with dimensions
 621 $2d \times 4d$ as in Fig.16. At the bottom and the top of the column a no-slip boundary
 622 condition is applied with a free-stream boundary condition at the vertical walls
 623 of the domain. The velocity is set to zero in the domain, and inside the bubble
 624 the pressure is constant. The physical properties for the bubble and the heavier
 625 surrounding fluid are listed in Table 11. Different triangular meshes were used
 626 with their resolution varying as: $d/40, d/80, d/160$. The benchmark quantities
 627 used by [69] are the centre mass (y_c), the rise velocity of the bubble (v_c) and its
 628 circularity or sphericity in three dimensions (ζ). These are defined as

$$\begin{aligned}
 y_c &= \frac{\int_{\Omega_b} \mathbf{x} dV}{\int_{\Omega_b} dV} \\
 v_c &= \frac{\int_{\Omega_b} \mathbf{u} dV}{\int_{\Omega_b} dV} \\
 \zeta &= \frac{\pi d}{\Pi}
 \end{aligned}
 \tag{36}$$

631 where Ω_b is the region occupied by the bubble and Π is the perimeter of
 632 the bubble. Fig.17 shows the benchmark quantities through time for the dif-
 633 ferent meshes. Results obtained in the present study are close to the values
 634 obtained in [69] and [70] for x_c , v_c and ζ for the different mesh resolutions. For
 635 the present conditions with $Re = 35$ and $Eo = 10$, where $Eo = gd^2\Delta\rho/\sigma$ is
 636 the Eötvös number, the surface tension force is significant which prevents the
 637 bubble disintegrating. The bubble deforms ($t = 1$) and changes shape from cir-
 638 cular to ellipsoid (see also [70]) reaching its terminal velocity at $t = 2$ which is
 639 approximately 90 percent of the maximum bubble velocity. The change in rise
 640 velocity is in good agreement with the velocity obtained in [70]. At the change
 641 in circularity at $t = 1.9$ where the surface tension effect on the bubble shape is

642 more evident, and is captured with all three computational meshes in this study
643 and is also in close agreement with the calculated ζ in [69]. Similar patterns
644 for the calculated benchmark quantities were also observed in other numerical
645 works [29, 33, 70]. The results for the relative error norms for y_c , ζ and v_c are
646 shown in Table 12. The calculated errors are in close agreement with the results
647 reported in [70] for the three benchmark quantities.

648 For the three-dimensional version of rising bubble test case, a three-dimensional
649 bubble is placed in a cylinder with height $8d$ and diameter $8d$. The bubble is
650 placed at a distance $1.5d$ from the bottom of the cylinder. The density and
651 viscosity ratio between the bubble and the surrounding fluid were set to 100.
652 Three hexahedral meshes with different resolution were used for this study,
653 with a grid size $\Delta x = d/15, 20, 40$. The errors E_{Re} and E_ζ for the calculated
654 Reynolds number Re and sphericity ζ are used to assess the accuracy of the
655 method, where $E_{Re} = (Re - Re_{exact})/Re_{exact}$, $E_\zeta = (\zeta - \zeta_{exact})/\zeta_{exact}$ (where
656 Re_{exact} and ζ_{exact} are the exact values for Re and ζ respectively). The results
657 shown in Table 13 are in good agreement for the three meshes compared to
658 the results from the reference case in [29]. In Fig.18 the mass conservation
659 error is shown through time. The mass conservation error is calculated with
660 respect to the volume fraction at $t = 0$, $\alpha(0)$ and is defined following [29] as
661 $\delta M = |\alpha - \alpha(0)|/\alpha(0)$. The error for the different meshes remained of the order
662 of 10^{-5} or below showing reasonable accuracy for mass conservation. A similar
663 order for δM is reported in the CLSVOF work in [29].

Table 11: Physical properties for the two-dimensional rising bubble test case.

	Physical and numerical parameters
Heavy fluid density	1000 kg/m^3
Bubble density	100 kg/m^3
Heavy fluid viscosity	10 $kg/m \cdot s$
Bubble fluid viscosity	1 $kg/m \cdot s$

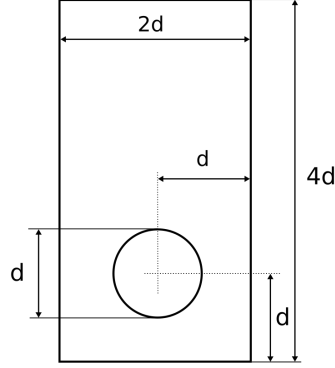


Figure 16: Domain for the two-dimensional bubble rise test case. The diameter of the bubble is initially $d = 0.5$.

Table 12: Relative norm error for the centre of mass, circularity and velocity of the bubble in the two-dimensional rising bubble case. Three different structured meshes are used, and the results are compared with the level set method of [70].

Mesh size	ILSVOF method			LS method of [70]		
	Centre of mass	Circularity	Velocity	Centre of mass	Circularity	Velocity
1/40	1.79×10^{-3}	1.19×10^{-3}	1.06×10^{-2}	2.65×10^{-3}	1.0×10^{-3}	1.19×10^{-2}
1/80	8.91×10^{-4}	3.18×10^{-4}	1.81×10^{-3}	9.64×10^{-4}	3.01×10^{-4}	2.9×10^{-3}
1/160	2.41×10^{-4}	6.37×10^{-5}	6.21×10^{-4}	2.62×10^{-4}	8.83×10^{-5}	7.73×10^{-4}

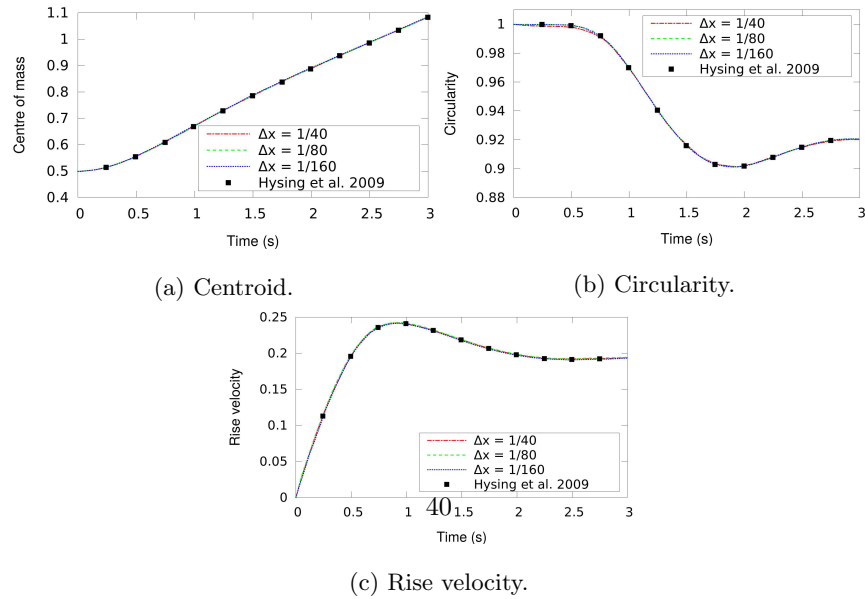


Figure 17: Evolution of rising bubble benchmark quantities through time. The results are compared with the case in [70].

Table 13: Error for Re and sphericity of the bubble in the three-dimensional rising bubble case. Three different structured meshes are used and the results are compared with the CLSVOF method of [29].

Mesh size	ILSVOF method		CLSVOF method of [29]	
	E_{Re}	E_{ζ}	E_{Re}	E_{ζ}
$d/15$	0.00363	0.0126	0.00341	0.0118
$d/20$	0.00314	0.0063	0.00339	0.0074
$d/40$	0.00271	0.0036	-	-

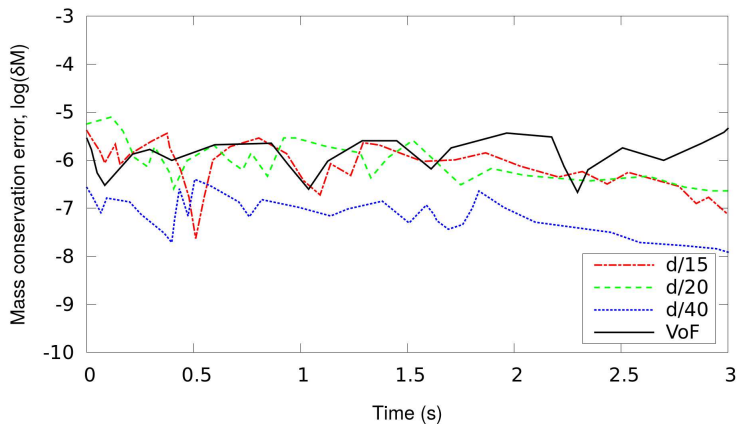


Figure 18: Mass conservation error for three-dimensional rising bubble case for the three levels of refinement.

664 4. Conclusions

665 A novel method for simulating the flow of two immiscible fluids tracking
666 their interface is presented coupling the level set and volume of fluid methods.
667 The new ILSVOF method involves a novel re-initialisation methodology which
668 is described in detail. ILSVOF is simple and can be readily implemented for
669 any type of polyhedral unstructured mesh. A smooth calculation of the gra-
670 dient of the LS function is utilised considering the neighbouring cells via an

671 interpolation at the cell-faces. Using these cell-face calculations for each inter-
672 face cell, it is easy to overcome the limitation of having to arbitrarily define the
673 upwind and downwind cells. The initial value for the re-distancing algorithm is
674 obtained via the advection of the isoface within a time step instead of by using
675 an algebraic method. Overall, the method provides better accuracy compared
676 to the VOF method in most of the numerical tests considered and has been
677 demonstrated to give an accurate representation of the interface in both two
678 and three-dimensional test cases. The mapping of the volume fraction to the
679 distance function is extremely important for the re-initialisation procedure, and
680 alternative ways of doing this, such as by employing a special advection step
681 within each solution time step, should be further investigated. Further com-
682 parisons with high order level set approaches could also be used to improve the
683 level set advection step.

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