ASSESSMENT OF NUMERICAL TECHNIQUES FOR MICRO-DROP FLUID MECHANICS

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Abstract. Are currently available CFD techniques able to model the motion of a submilimiter-scale liquid drop falling in another (quiescent) liquid? In this work we define a very simple two-dimensional test: An initially circular drop of water of diameter 0.1 mm is placed at the center of a square box (size 0.6 mm) filled with oil (relative density 0.8). The goal is to compute the fall of the drop towards the bottom of the box, which would take several seconds based on typical terminal velocities found in the problem. It seems easy, doesn’t it? We have tested several CFD techniques on this problem: FOAM (two available VOF-based methods within FOAM’s platform), GERRIS (free software from Popinet’s group), level-set finite elements (interface capturing) and finally ALE finite elements (interface tracking). The aim is to illustrate on the serious limitations this methodologies have to accurately and efficiently track the dropet’s trajectory.
1 INTRODUCTION

The study of sedimentation of water droplets in oil is of great interest for the petroleum industry (Frising et al., 2006). Crude oil is seldom produced alone. In majority of cases the oil is produced along with water. Due to the presence of active agents in the oil, a water-in-oil emulsion is produced, i.e., an emulsion with oil being the continuous phase and water the dispersed one. Several factors can influence the formation of such emulsions, such as the flow through the porous medium or the turbulence mixing at pipes and valves (Barnea and Mizrahi, 1975). After extraction the two phases are separated which is usually accomplished by using the so called gravity settlers where two important phenomena are observed: droplet coalescence and droplet sedimentation. Both are interesting and challenging phenomena to deal with from a computational standpoint. The former because it involves merging of interfaces and therefore requires robust numerical techniques to handle topological changes of the fluid phases. The latter, which is the main interest here, although simple at a first glance, also presents challenges because it requires tracking of the droplet trajectory for several seconds until it settles down at the bottom of the separator. The diameter of droplets of interest for these applications are in the submilimiter scale with values near 0.1mm. Under these conditions the flow is highly dominated by viscous and surface tension effects and droplets remain spherical along their sedimentation.

State-of-the-art CFD techniques for two-phase flows dominated by surface tension in this operating range suffer from severe time step restrictions (Galusinski and Vigneaux, 2008) and/or from the so called parasitic currents (Scardovelli and Zaleski, 1999) if coarse meshes or low order approximations are used, in which case simulation results may end up being totally meaningless. Refining the time step and the mesh render numerical simulations for the whole history of sedimentation, even for a single droplet, extremely costly. The aim of this article is precisely to illustrate on this point by using several numerical techniques which include, various VOF solvers (Weller et al., 1998; Popinet, 2009; Márquez Damián, 2013), a finite element level set method as proposed in Ausas (2010) and an Arbitrary Lagrangian Eulerian method as proposed in Montefuscolo (2012). The simulation results show the behavior and robustness of the different methodologies to predict the evolution of a single submilimiter water droplet in oil.

By means of outline, after this introduction, the equations and non-dimensional parameters governing the sedimentation of droplets are presented. Next, the different numerical methodologies considered to discretize the problem are suscintly recalled and numerical results from each one are shown to illustrate on their limitations for different time step and mesh resolutions. Finally, some conclusions are drawn.

2 GOVERNING EQUATIONS

All the numerical methodologies used in this work to assess the sedimentation of the droplet implement the resolution of the mathematical problem describe in this section. We consider two-phase immiscible Newtonian flows. The fluid domain $\Omega$ is divided into a “plus” ($+$) region and a “minus” ($-$) region by a closed (smooth) interface $\Gamma$ as shown in figure 1, such that

$$\Omega = \Omega^+(t) \cup \Gamma(t) \cup \Omega^-(t).$$

(1)

The density $\rho$ and viscosity $\mu$ are assumed homogeneous and constant within each region, namely

$$(\rho(x, t), \mu(x, t)) = \begin{cases} (\rho^+, \mu^+) & \text{if } x \in \Omega^+(t) \\ (\rho^-, \mu^-) & \text{if } x \in \Omega^-(t) \end{cases}$$

(2)
On each region we need to solve the dynamical equilibrium equations, i.e.,

\[ \rho \left( \partial_t u + u \cdot \nabla u \right) - \nabla \cdot \tau + \nabla p = -\rho g \hat{e}_2 \quad \text{in} \quad \Omega^+ (t) \cup \Omega^- (t) \tag{3} \]

where \( u \) is the velocity field, \( p \) is the pressure field, \(-g \hat{e}_2\) is the gravitational force and \( \tau = \mu (\nabla u + \nabla^T u) \) is the stress tensor. The incompressibility restriction reads

\[ \nabla \cdot u = 0 \quad \text{in} \quad \Omega^+ (t) \cup \Omega^- (t) \tag{4} \]

In this work we also consider the following initial and boundary conditions

\[
\begin{align*}
\{ u(x, t = 0) &= 0 \quad x \in \Omega^+ (t) \cup \Omega^- (t) \\
u(x, t) &= 0 \quad x \in \partial \Omega \tag{5}
\end{align*}
\]

**Interface conditions**

Important for this problem are the jump conditions at the interface \( \Gamma \). For capillary interfaces, the jump conditions for velocity and stresses on the interface read

\[
\begin{align*}
\left[ u \right] &= 0 \quad \text{on} \ \Gamma \\
\left[ \tau \cdot \hat{n} \right] &= \sigma \kappa \hat{n} - \nabla_\Gamma \sigma \quad \text{on} \ \Gamma \tag{6}
\end{align*}
\]

In which \([\cdot]\) represents the jump of any quantity at \( \Gamma \), \( \sigma \) is the surface tension coefficient, \( \kappa \) is the mean curvature of \( \Gamma \) and \( \hat{n} \) is the normal vector to \( \Gamma \) pointing towards \( \Omega^+ \). The first condition implies that incompressibility is satisfied at the interface. The second condition incorporates the dynamical effects at \( \Gamma \). Here, \( \sigma \) is assumed constant and uniform along \( \Gamma \) for the sake of simplicity and therefore we neglect Marangoni effects coming from its surface gradient \( \nabla_\Gamma \sigma \). These effects are generally due to chemical or thermal gradients. The variables that are expected to be discontinuous at \( \Gamma \) are thus the pressure field and the velocity gradient. For more details see Buscaglia and Ausas (2012).
The interface conditions appear naturally in a variational formulation of the problem which is well suited for the finite element methods used later on. An alternative equivalent approach useful for discretization in other frameworks, is to consider the surface tension force as a singular force at the interface, i.e.,

\[
F_\sigma = \delta_\Gamma \kappa \hat{n} \quad \text{on} \quad \Gamma
\]

(7)

This is the approach followed in the VOF formulation presented in the following section. The Dirac distribution \( \delta_\Gamma \) can be further regularized for discretization purposes leading to different formulations in turn.

2.1 Sedimentation velocity

In this work our aim is to assess different numerical methodologies to simulate the sedimentation of a single water droplet in oil as shown in figure 1. For the sake of simplicity, we consider a two dimensional (planar) setting and therefore study the sedimentation of “large cylinders”. The behavior of droplets in different flow regimes is studied in the classical work of Clift et al. (2005). Here it is reasonable to assume that surface tension effects dominate the problem and the droplet remains circular along its evolution. Since the Reynolds number is also expected to be small, a simple estimation based on Stokes’ flow gives the following expression for the sedimentation velocity

\[
U_s = \sqrt{\frac{gd\pi \rho_w - \rho_o}{C_d \rho_o}},
\]

(8)

where \( \rho_w \) and \( \rho_o \) are the densities of water and oil respectively, \( d \) is the droplet’s diameter, and \( C_d \) is the drag coefficient which for low Reynolds number is roughly proportional to \( \sim 1/\text{Re} \), yielding (see Perry (1950))

\[
U_s = \frac{(\rho_w - \rho_o) g d^2}{18 \mu_o},
\]

(9)

where \( \mu_o \) is the oil viscosity (the continuous phase). Different estimations can be made for the Drag coefficient as proposed in Finn (1953)

\[
C_d = \frac{8 \pi}{\left[ \frac{1}{2} - 0.577 - \ln(\text{Re}/8) \right] \text{Re}},
\]

(10)

but an iterative procedure is then needed to compute \( U_s \). The velocity \( U_s \) is used to effectively compute the non-dimensional parameters in the next subsection and confirm the assumptions already made.

2.2 Non-dimensional groups

Dimensional analysis on the governing equations leads to five non-dimensional parameters for this problem: the Reynolds number, the Froude Number, the Weber number and the ratio between densities \( \rho_o/\rho_w \) and viscosities \( \mu_o/\mu_w \). However, some additional numbers are usually used in the literature to characterize these flows. The parameters of interest here are the Reynolds number, the capillary number \( \text{Ca} \), the Eötvos number \( \text{Eo} \) and the Laplace number \( \text{La} \). They can be all obtained as combinations of the formers and are defined as

\[
\text{Re} = \frac{\rho U d}{\mu}, \quad \text{Ca} = \frac{\mu U}{\sigma}, \quad \text{Eo} = \frac{\Delta \rho g d^2}{\sigma}, \quad \text{La} = \frac{\sigma \rho d}{\mu^2}
\]

(11)
The characteristic velocity is chosen as the terminal or sedimentation velocity \( U_s \) previously introduced. The physical parameters adopted for the numerical simulations are summarized in Table 1. These values correspond to water droplets in oil. Notice the large viscosity ratio \( \mu_o/\mu_w \) of 300 considered. Using the aforementioned rough estimation for the drag coefficient the sedimentation velocity results in

\[
U_s \approx 3.7 \times 10^{-6} \text{ m/s},
\]

finally yielding

\[
\text{Re} = 1.16 \times 10^{-6}, \quad \text{Ca} = 6.54 \times 10^{-5}
\]

\[
\text{Eo} = 10^{-3}, \quad \text{La} = 1.78 \times 10^{-2}
\]

Under these conditions, the flow is clearly in the Stokes regime and dominated by surface tension and viscosity.

Table 1: Physical parameters adopted for the sedimentation of a water droplet in oil.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Symbol</th>
<th>Value [SI]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Droplet diameter</td>
<td>( d )</td>
<td>0.0001 m</td>
</tr>
<tr>
<td>Domain size</td>
<td>( L )</td>
<td>0.0006 m</td>
</tr>
<tr>
<td>Gravity acceleration</td>
<td>( g )</td>
<td>10 m/s(^2)</td>
</tr>
</tbody>
</table>

**Dispersed phase (Water Droplet):**

| Density                    | \( \rho_w \) | 1000 kg/m\(^3\) |
| Viscosity                  | \( \mu_w \) | 0.001 Pa s     |

**Continuous phase (Oil):**

| Density                    | \( \rho_o \) | 800 kg/m\(^3\) |
| Viscosity                  | \( \mu_o \) | 0.3 Pa s       |

**Interface:**

| Surface tension coefficient| \( \sigma \) | 0.02 kg/s\(^2\) |

3 NUMERICAL METHODS

3.1 Parasitic currents

Parasitic currents are a consequence of discretization errors in the numerical modeling of capillary flows. They have been extensively discussed in the literature in various numerical settings. References on the subject can be consulted e.g. in Popinet (2009) and Scardovelli and Zaleski (1999). Some numerical techniques, specially those based on an implicit representation of the interface non-conforming with the edges/faces of the underlying mesh where the flow problem is being solved, are particularly affected by them. Examples of these techniques are the well known level set and VOF methodologies used later on in this work. Several details of the discretization affect the strength and form of these parasitic currents. Although, a complete review of this topic is out of the scope of the article, we may mention:

- mesh quality and resolution (Deshpande et al., 2012; Lafaurie et al., 1993; Harvie et al., 2006);
- size of time step (Galusinski and Vigneaux, 2008);
• treatment of discontinuous variables close to \( \Gamma \), such as the pressure and the velocity gradient (Ausas et al., 2012; Idelsohn et al., 2010);

• accuracy in the discrete representation of \( \Gamma \) and computation of normal and curvature fields (Gross and Reusken, 2007; Cummins et al., 2005; Marchandise et al., 2007),

The behavior of different numerical methods with respect to these factors can vary a lot. As an example, in VOF methods the magnitude of spurious velocities can even increase with mesh refinement as reported in Deshpande et al. (2012). Other classes of methods, as those based on an explicit Lagrangian representation of the interface, such as the Arbitrary Lagrangian Eulerian (ALE) method may exhibit better behaviors as we show later on.

In this work we assess three different classes of methods to solve the sedimentation of the droplet. In the first place, the VOF method, based on a finite volume discretization of the problem. In the second place, an equal order finite element level set method in the form proposed by Ausas (2010). Finally, a finite element ALE method proposed by Montefuscolo (2012). Here, without entering into all the details, we succinctly recall the main ingredients involved on each of them. Prior to this we briefly discuss the time step restrictions that are expected in these problems.

3.2 Time step restrictions

We consider here numerical schemes in which velocity and pressure are solved with a fixed interface geometry and then geometry is updated with the last computed velocity. These are called staggered schemes for which a time step restriction is expected. According to Galusinski and Vigneaux (2008) the time step \( \Delta t \) needs to satisfy the following limiting criterion for stability

\[
\Delta t \leq \Delta t_{\text{lim}} \approx \frac{1}{2} \left\{ C_2 \frac{h \mu}{\sigma} + \sqrt{(C_2 \frac{h \mu}{\sigma})^2 + 4 C_1 \frac{\rho h^3}{\sigma}} \right\},
\]

where \( h \) is the element size and \( C_1 \) and \( C_2 \) are constants depending on the specific method but not on the mesh size and the physical parameters of the problem. For instance, in Deshpande et al. (2012) the authors assess the behavior of interFoam solver (Weller et al., 1998) and experimentally find values of 0.01 and 10 for \( C_1 \) and \( C_2 \) respectively. One would expect that a reasonable value for the mesh size \( h \) in our problem be of the order of \( d/10 \), meaning that a single droplet in a 3D simulation is resolved with 1000 discretization points. This would be an affordable quantity if the aim is at simulating for instance 1000 droplets. The numerical values for viscosity and density considered in table 1 yield \( \Delta t_{\text{lim}} \approx 10^{-5} \) s (at worst). According to the terminal velocity previously computed of \( 3.7 \times 10^{-6} \) m/s, the droplet requires \( \sim 80 \) s to settle down at the bottom of the computational domain, which means that 8 million time steps are needed to complete the simulation! Of course, this restriction can be improved for some of the formulations by using a semi-implicit scheme as proposed by Bänsch (2001) or an implicit scheme as done in Buscaglia and Ausas (2012); Ausas et al. (2009) to deal with the surface tension force.

Now, we describe the main features of the numerical schemes to be tested. Again, we emphasize that a staggered scheme is used in which \( u \) and \( p \) are first solved with a fixed interface, and then \( \Gamma \) is transported with the last computed \( u \).
VOF - Finite volume methods

In the Volume of Fluid (VOF) method initially proposed by Hirt and Nichols (1981) the incompressible Navier-Stokes equations (3) are solved by using a density and viscosity defined by

\[
\mu = \alpha \mu^- + (1 - \alpha) \mu^+ , \\
\rho = \alpha \rho^- + (1 - \alpha) \rho^+ ,
\]

(15)

\( \alpha \) being the volume fraction of the primary phase, i.e., it is a scalar field that assumes values equal to 1 at those cells completely in \( \Omega^- \), 0 at those cells completely in \( \Omega^+ \) and varies between 0 and 1 in those cells on the interface. The advection of \( \alpha \) is governed by the following equation in conservative form

\[
\partial_t \alpha + \nabla \cdot (\alpha \mathbf{u}) + \nabla \cdot [\alpha (1 - \alpha) \mathbf{v}_r] = 0 ,
\]

(16)

where \( \mathbf{v}_r \) is the relative velocity between the two phases. Theoretically, this velocity should be zero, however, in some implementations it is retained to add some compressibility at the interface, which actually turns the problem non-linear. In this work several formulations or implementations are tested, specifically, the InterFoam solver of Weller et al. (1998), a derived solver called interDyMFoam, both in the free software suite OpenFOAM® and the also publicly available Gerris software proposed by Popinet (2003). Some ingredients common to all the aforementioned VOF implementations that were adopted in this work are:

- Discretization of the computational domain into finite volume quadrilateral cells.
- Cell-centered finite volume method.
- Derivation of the PISO method for pressure-velocity coupling.
- Geometric properties and surface tension force computed based on the volume fraction without regularizations.
- Multidimensional Universal Limiter with Explicit Solution (used in FOAM implementations, see Deshpande et al. (2012); Márquez Damián (2013)) and TVD scheme in Gerris to solve the advection equation.

Regarding the computation of the surface tension force mentioned in the itemized list above, at each cell, the normal vector and curvature are computed based on \( \alpha \) as (Brackbill et al., 1992)

\[
\hat{n} = \frac{\nabla \alpha}{\| \nabla \alpha \|} , \quad \kappa = - \nabla \cdot \left( \frac{\nabla \alpha}{\| \nabla \alpha \|} \right),
\]

(17)

yielding the following form for the surface tension force

\[
\mathbf{F}_\sigma = \sigma \kappa \hat{n}
\]

(18)

Level set - Finite element method

In the level set method (Osher and Sethian, 1988) the interface is defined as the zero set of a scalar function \( \phi(x, t) \), i.e.,

\[
\Gamma = \{ x \in \Omega, \phi(x) = 0 \},
\]

(19)

this function should satisfy the condition

\[
\nabla \phi(x) \neq 0 \quad x \in \Gamma ,
\]

(20)
so as the normal to \( \Gamma \) is well defined and given by

\[
\mathbf{n}(x) = \frac{\nabla \phi(x)}{\| \nabla \phi(x) \|} \quad x \in \Gamma
\] (21)

The level set function and implicitly \( \Gamma \), are transported by the velocity field \( u \) that follows after solving the Navier-Stokes equations (3). This transport is done by means of solving the hyperbolic advection equation

\[
\partial_t \phi + u \cdot \nabla \phi = 0 \quad \text{in } \Omega, t > 0
\] (22)

where the initial condition \( \phi(x, 0) = \phi_0(x) \) is defined according to the initial position of the interface. Also, appropriate Dirichlet boundary conditions are needed at the inflow boundaries if present. A thorough description of the complete formulation and details adopted here can be found elsewhere (see Ausas (2010)). The finite element method is adopted for spatial discretization, whose main ingredients are summarized below.

- Discretization of the computational domain into triangles.
- Equal order formulation to solve the Navier-Stokes equations (3) using linear elements for velocity and pressure and algebraic subgrid scale method for stabilization (see e.g. (Codina, 2001; Hughes et al., 1986)).
- SUPG formulation to solve the transport equation (22) with \( P_1 \) elements for the level set function.
- First order temporal accuracy.
- Modified pressure space proposed in Ausas et al. (2010) to capture discontinuities in the pressure field.
- Laplace-Beltrami treatment of the surface tension force.

Using standard \( P_1 \) elements for the level set function implies that the reconstructed interface is elementwise planar. In the Laplace-Beltrami approach, the curvature computation is avoided by means of integrating by parts along \( \Gamma \) (Buscaglia and Ausas, 2012).

**ALE method**

In an ALE formulation Hughes et al. (1981); Baiges et al. (2010); Rabier and Medale (2003) the interface between the two fluids is represented by the edges (in 2D) or faces (in 3D) of the mesh in which the flow problem is being solved. For the particular problem we have at hand, this formulation is expected to be accurate and robust to follow the falling droplet. In the ALE method the finite element partition moves with time. Each nodal position is a function of time that is denoted by \( x^j(t) \). The nodal velocity can be defined as

\[
\mathbf{v}^j(t) = \frac{d x^j(t)}{dt}, \quad j \text{ running over the nodes of the finite element partition}
\] (23)

index \( j \) running over the nodes of the finite element partition. The velocity field is approximated as

\[
\mathbf{u}_h = \sum_j N^j(x, t) \mathbf{u}^j(t), \quad N^j(\cdot, t) \mathbf{e}_n \in W_h(t)
\] (24)
where $e_\alpha$, $\alpha = 1, \ldots, d$ is the canonical basis, and $u^j(t)$ are the nodal velocity values. In (24) $W_h(t) \subset (H^1(\Omega(t)))^d$ is a finite dimensional space that depends parametrically on time. A key point in the derivation of the ALE formulation is the following identity
\[
\partial_t u_h = \sum_j N^j \frac{du^j}{dt} - (v_h \cdot \nabla) u_h
\] (25)
which inserted into a classical spatial discretization of the variational form of (3) yields a differential algebraic equation system to which any temporal discretization scheme can be applied. The formulation adopted here is presented with details in Montefuscolo (2012). The main ingredients of the implementation are
- Discretization of the computational domain into triangles.
- Equal order formulation to solve the Navier-Stokes equations (3) using linear elements for velocity and pressure and algebraic subgrid scale method for stabilization.
- Discontinuous elements for the pressure field at the interface.
- Linear elements for geometry interpolation.
- First order temporal accuracy.
- Elasticity-like problem for the mesh velocity equation.
- Laplace-Beltrami treatment of the surface tension force.

As noticed, many features of the approximation are shared with the finite element level set methodology presented above.

4 NUMERICAL RESULTS

In this section we present results from each numerical methodology to illustrate on their limitations. The numerical values adopted and the corresponding non-dimensional parameters are the ones given in section 2.2. Similar numerical studies using Eulerian and Lagrangian formulations are presented for instance in Bertaki et al. (2010) and Bäumler et al. (2011), however, in none of these articles the authors address the submilliter scale as done here.

Before passing to the results, some comments are in order. For both finite element based formulations, namely, the level set and the ALE methods, simulations are carried out for different grid resolutions. The central region of the computational domain is refined so as to have mesh resolutions $h$ equal to $d/10$, $d/20$ and $d/40$, while a coarser mesh is generated far away from the region of interest. For the VOF simulations, uniform and locally refined meshes only near the interface are used. This allows us to save some computational time and assess the behavior of the different formulations with respect to grid resolution. Simulations runs until one second of physical time when possible. Although this is a short time compared to the total one needed for complete sedimentation, and the steady state velocity may not be even reached for some of the simulations, we have found this sufficient to characterize the behavior of the numerical schemes. Finally, we also report on the center of mass position and velocity computed as
\[
r_{cm}(t) = \frac{1}{\text{meas}(\Omega^+(t))} \int_{\Omega^+(t)} x \, dx, \quad u_{cm}(t) = \frac{1}{\text{meas}(\Omega^+(t))} \int_{\Omega^+(t)} u(x, t) \, dx
\] (26)
remberming the value previously estimated for the sedimentation velocity is $0.0037 \text{ mm/s}$. 

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4.1 VOF results

In this section we present results corresponding to different VOF formulations: interFoam, interDyMFoam and Gerris. Other formulations also tested as iInterFoam showed very similar results or did not succeed in completing a significant part of the total simulation time and therefore are not shown here for the sake of brevity. The finite volume meshes used in the simulations are shown in figure 2. For interFoam two meshes are considered: a regular one with a cell size $\Delta x = d/10$ and a adaptively locally refined one with up to three levels of refinement obtained by uniform cell splitting near the droplet’s interface. In Gerris code the grid is also adaptively refined as the simulation proceeds similarly to interDyMFoam. The initial grid is always consisting of an uniform regular grid with $\Delta x = d/10$. The time step $\Delta t$ used in simulations are $10^{-6}s$ for the uniform mesh used by interFoam and $10^{-7}s$ for the refined grids used by interDyMFoam and Gerris. For the finer meshes, simulations are computationally very demanding and we were able to report results only for the different times indicated. The velocity fields are shown in figure 3. As observed the strenght of parasitic currents is totally polluting the results for interFoam and interDyMFoam. The droplet remains essentially at its initial position during most part of the simulation and for this reason the plot is not shown. Further mesh and time step refinements do not solve the problem. For Gerris, on the other hand, results start to improve with the level of refinement considered. This is due to the improved treatment for the curvature field by means of the so called height functions used in this implementation. The droplet did not attain the terminal velocity but the flow pattern is clearly symmetric as expected. These results already show serious limitations of some of the VOF implementations, even when very fine meshes are used.

Using Gerris software, with the adaptively refined grid and time step, the complete sedimentation can be simulated in approximately 67 days on a 4-core desktop computer with i7 processor at 2.9 GHz.

![Figure 2: Finite volume meshes used for the VOF simulations.](image)

4.2 Level set results

The finite element meshes built for the level set simulations are displayed in figure 4. As previously mentioned they are finer in the region of interest, being this more than sufficient to simulate 1 s of time. The time step $\Delta t$ used in simulations is $10^{-4}s$ for the first mesh, $5 \times 10^{-5}s$ for the second one and $2.5 \times 10^{-5}s$ for finest one. Remember the modified pressure space is being used to capture the discontinuous behavior of $p$ at the interface, however, this is not enough to avoid the nocive effect of the parasitic currents as shown in figure 5 where the velocity...
field at $t = 1$ is plotted. The maximum of the colour scale in these figures is dictated by the maximum parasitic current which sometimes is limited to few elements, hindering the fact that most of the droplet phase is moving at the same velocity. This is particularly the case for the third refinement level and to some extent for the second refinement. Figure 6 shows the droplet’s vertical position and velocity as a function of time for the three meshes. Results start to be acceptable for the third refinement in which the size of the parasitic currents is sufficiently reduced being their effect limited to a very small region and/or for small periods of time. For the sake of clarity, in these plots only a fraction of the total number of simulated points is shown. The poorest results are obtained for the first refinement (red dots) for which the strenght of the spurious velocities completely pollutes the droplet’s trajectory.

For the finest mesh (consisting of 14406 triangular elements), using a time step $\Delta t = 2.5 \times 10^{-5}$, the complete sedimentation can be simulated in approximately 15 days on a 4-core desktop computer with i7 processor at 2.9 GHz.

4.3 ALE results

The finite element meshes built to compute the flow with the ALE formulation are shown in figure 7. A finer region in the central part of the computational domain is also considered, although a smoother transition from fine to coarse elements is used. The droplet interface is coincident with the element edges as can be notice by zooming in at these plots. The corre-

sponding velocity fields are shown in figure 8. The time step $\Delta t$ used in simulations is $10^{-4}$s for the first mesh and $10^{-5}$s for the rest of the meshes. The superior behavior of this formulation
Figure 5: Velocity field (in mm/s) obtained with the level set formulation on the different meshes at $t = 1$.

Figure 6: Droplet’s center of mass position (left) and velocity (right) as a function of time obtained with the level set formulation for the different meshes.

as compared to previous ones is quite evident. This is also seen by plotting the evolution of the center of mass position and velocity as a function of time (see figure 9). As observed in figure 9, the droplet does not attain the terminal value of the sedimentation velocity at the end of the simulation, and the time needed to do so seems to be quite dependent on mesh resolution. The main limitations observed are related to the size of the time step needed in principle for stability if the staggered scheme is used, although this can be partially relaxed.

For the coarsest mesh (consisting of 1516 triangular elements) for which results are already reasonably accurate, using a time step $\Delta t = 10^{-4}$, the complete sedimentation can be simulated in approximately 30 days on a 4-core desktop computer with i7 processor at 2.9 GHz.

Figure 7: Finite element meshes used for the ALE simulations.
5 CONCLUSIONS

We have shown the behavior of three different classes of numerical methods for two-phase capillary flows to assess a relevant problem for the oil industry related to the sedimentation of submilimiter water droplets in oil. It has been shown that there exist serious limitations of such methods to efficiently solve the problem: millions of very short time steps are needed to follow the whole history of sedimentation when staggered schemes are used and/or very fine meshes becomes necessary to reduce the size and extent of parasitic currents. If the final goal is the simulation of real systems involving thousands of 3D droplets, these are extremely limiting factors. For the VOF method, the only implementation that seems to be sufficiently accurate for this problem is the one used in the publicly available software Gerris. In some of the implementations tested, including some commercial ones, the simulation even reached the final time, or the droplet’s behavior was totally polluted by the parasitic currents with disastrous consequences. For the level set method, the linear interpolation used for interface representation becomes a very limiting factor to as regards accuracy. Results start to be reasonably accurate for moderately refined meshes. In this case, as well as for the ALE approach, adding implicitness to the formulation is relatively easy and certainly mitigates the time step restriction issue as has been verified although not reported here for the sake of brevity. As expected, the ALE formulation outperformed the rest of the methodologies with respect to accuracy for a given mesh resolution.
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