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QEEFoam: a Quasi-Eulerian-Eulerian model for polydisperse turbulent gas-liquid flows. Implementation in OpenFOAM, verification and validation.

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Abstract

In this paper we present a new multiphase computational model for polydisperse turbulent gas-liquid flows. In this model the gas phase is transported by a single convection equation and the effect of turbulent dispersion is addressed by including a diffusion term. In order to close the system of equations, the gas phase velocity is calculated by employing the slip velocity concept or by solving an ordinary differential equation. This procedure shares similarity with the Euler-Lagrange (E-L) method, in which the gas velocity is updated by bubble Lagrangian tracking, and with the Euler-Euler (E-E) method, and for this reason it is called the Quasi-Eulerian-Eulerian (Q-E-E) method. In order to account for polydispersity one single transport equation is added to describe the effects of bubble breakage and coalescence on the bubble size distribution. The novel Q-E-E method was implemented in the open-source code OpenFOAM-7 and was used to simulate turbulent gas-liquid flows with three different geometries operating under different conditions. The predictions for the dynamical vortex structures, local phase fraction, global gas holdup, bubble size distribution and vertical/horizontal liquid velocities were verified against the solution provided by the E-L solver, or against published experimental data. Good agreement was found and extremely small computational costs were detected.

Keywords: Computational fluid dynamics, Bubbly flow, Eulerian-Eulerian model, OpenFOAM, Bubble column, Bubble size distribution

1. Introduction

Bubbly flows are omnipresent in engineering and are found in many applications, ranging from the chemical to the nuclear industry. For bubble-driven flows, the bubble movement is the main source of momentum for the flow field; these flows are often characterized by low liquid velocities and relatively high superficial gas velocities. The bubble-driven flow in ducts and columns with a small aspect ratio is usually complicated. Recirculating flow is quite common even at relatively low gas flow rates [30], which can speed up mass and heat transfer. If the gas superficial velocity is high, it leads to intense interaction between bubbles and bubble breakage and coalescence can be dominating. In vertical bubbly flows with a high aspect ratio, the small bubbles migrate to the wall resulting in a steeper velocity gradient in the near-wall region. On the opposite, the large bubbles migrate towards the center [71, 15]. The velocity profile resembles that of single phase flows. In all the cases, there

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is a strong coupling between the gas bubbles spatial distribution, the liquid velocity, the momentum exchange between gas and liquid and the resulting flow structure is quite complex.

Besides experimental investigations, bubbly flows can be also analyzed by using computational models, which cannot be solved analytically except in very special cases. For process design and analysis, computational fluid dynamics (CFD) has become an important and indispensable tool. Specifically, the simulation methods for gas-liquid flows in CFD can be divided mainly into three categories. In the first category one finds multiphase direct numerical simulation methods, based on the simple idea of directly tracking the interface between the involved phases. In the case of bubbly flows this implies tracking the evolution of the gas-liquid interface around each bubble. This is an extremely accurate method, but characterized by very high computational costs. These methods will not be discussed here, since they are usually restricted to very low Reynolds numbers and are suitable for dilute gas-liquid flows. Readers interested in the details are referred to the specialized literature [55].

In the Eulerian-Lagrangian (E-L) method [10, 29] the continuous phase is processed via the Eulerian framework, while the individual bubble motion is simulated through the solution of the corresponding Newton equation of motion. Bubbles trajectories are calculated and by considering fixed control volumes the coupling with the continuous phase can be finalized. These discrete bubbles can be coarse-grained into larger bubble clusters to speed up the simulation and the bubble cluster is usually called parcels [64]. The E-L method features no numerical diffusion as the most important merit.

In the Eulerian-Eulerian (E-E) method [5, 51, 61], both phases are evolved in the Eulerian framework, and a different set of Navier-Stokes equations can be constructed for each phase. The coupling between these phases is achieved by the so-called momentum interfacial exchange term appearing in the momentum equations. In quadrature-based moments methods (QBMM) [46], the dispersed phase is described by the generalized population balance equation while the continuous phase is controlled by the traditional Eulerian equations. Depending on the closure of the disperse phase velocity, this method can be easily extended to poly-disperse systems and bubble coalescence and breakage can be included. This method was investigated extensively in our previous works [38, 40, 39, 36]. Another promising method is the mixture model [45, 26] and one of its most popular implementation is based on the so-called algebraic slip model [65]. In the mixture model, instead of describing each phase by using its own continuity and momentum balance equation, only one continuity equation and one momentum balance equation are solved for the mixture of the two phases. All of these models except the E-L method do not keep track of the individual bubbles but represent an averaged description in which the space domain may contain a fraction of both the continuous and the disperse phases. Readers are suggested to other works for more information on the derivation and application of these mathematical models [25, 43, 68].

However, none of these methods is satisfactory. The governing equations of the E-E method are quite complex, which implies that complicated numerical techniques are needed to handle the phase segregation problem and the phase coupling [53, 37]. Handling the numerical diffusion in the E-E method is also non-trivial, since high-order spatial discretization schemes usually generate boundedness problems, resulting in turn in catastrophic divergence. The realizability of the moments in the E-QBMM cannot be easily ensured either and dozens of moment equations are needed to be solved for a three-dimensional test case. Meanwhile, fine meshes cannot be used in the traditional E-L method since the bubble size should be smaller than the cell size in order to calculate the momentum transfer between phases accurately [11]. Statistical noise cannot be avoided either. For a given desired accuracy, this greatly increases the computational cost and a large number of numerical bubbles are required to eliminate statistical noise. For the mixture model, the boundary condition of the mixture velocity needs to be carefully treated and it is mainly used for gas-solid settling problems

with very small slip velocity [1, 48].

Is it possible to have a computational model that simplifies the governing equations of the multi-phase flow beyond these existing methods? The answer to this question seems to be yes. Sokolichin et al. [62] developed a simplified gas-liquid model from the two-phase E-E method. The spirit of this method is that, if one sums up the momentum equation of the gas and the liquid and neglects the terms involving the gas density ($\rho_g \ll \rho_l$), a liquid momentum equation without phase fraction is constructed. In this model, the governing equations fully correspond with the mass and momentum balance equations and the equation of state of a single-phase flow. Meanwhile, a phase fraction depended acceleration term exists in the momentum equation. This term is used to describe a force which is directed upwards and is proportional to the local gas holdup. Therefore, the gas-liquid flow can be interpreted as a one-phase flow with an additional buoyancy source term in the momentum balance. However, this method is restricted to a very low local phase fractions. Ferry and Balachandar proposed a so-called fast or equilibrium Eulerian-Eulerian method for small particle response times τ [18, 19]. They express the particle velocity as an expansion in τ to remove the particle momentum partial differential equation. This equation is first-order accuracy with respect to τ but and higher-order equations can also be derived. It was further extended to account near-wall approximation, which can accurately capture the high shear region behavior [20]. The very same approach was also used in other works to simulate particle-laden flows for small Stokes flows [60, 59]. Only the concentration field needs to be solved for particles of certain size range. Another possible similar simplification originates from the E-L method. If one replaces the Lagrangian particles equation with a gas phase convection equation under the Eulerian framework, the time consuming Lagrangian tracking is not necessary and the Lagrangian noise can be avoided. However, the equation system is not closed since the gas phase velocity is unknown. Therefore, the gas phase velocity needs to be modelled and a closed set of equation is finally obtained and can be solved.

The methods described so far are generally all solved assuming a monodisperse population of bubbles with constant and given size. However real bubbly flows are characterized by bubble breakage and coalescence, leading to polydispersity. To account for these effects the bubble size distribution must be calculated by solving the population balance equation (PBE). Different methods were proposed to solve the PBE, such as the class method (CM) [32], the quadrature method of moments (QMOM) [47] and they were employed to reconstruct number density functions in many works [23, 70, 67, 57, 12, 21, 22, 8, 31]. However, these methods require large computational resources [38, 36] and a cheaper alternative could be the so-called one primary and one secondary particle method (OPOSPM) [17, 2, 3], which conserves only the total gas bubble number density and the total gas bubble volume density (i.e. volume fraction). In this method two transport equations for the total number density and the total volume fraction of the gas bubbles are solved together with a one-quadrature point for closing the unclosed integrals in the PBE. This method can be further simplified to a one-equation model for the solution of the PBE based on the mathematically consistent d_{30} instead of the classical mean droplet diameter. Yet another computationally cheap alternative is the interfacial area transport equation (IATE) method [69, 27] which considers the gas-liquid specific surface area as one of the most important parameters and is based on the solution of the corresponding transport equation (for the interfacial area). The effect of bubble coalescence, due to the random turbulent collisions between bubbles, and of the wake entrainment process, due to the relative motions of the bubbles, can be accounted for. As far as bubble breakup is concerned instead the impact of turbulent eddies is considered. A similar method was also developed in [33, 34].

In this work, we present a new fast multiphase computational model for turbulent gas-liquid bubbly flows accounting for polydispersity. The unknown gas convection velocity is modelled by 1) employing the slip velocity method or 2) solving an ordinary differential equation. This equation resembles the E-L method, but the actual Lagrangian particle tracking is removed to achieve compu-

tational efficiency. It also resembles the Eulerian quadrature-based moments methods (E-QBMM), in which the velocity is modelled by the velocity polynomial assumption [46]. For these reasons we call this computational model the Quasi-Eulerian-Eulerian (Q-E-E) method. The first ‘‘Eulerian’’ denotes that the gas bubbles are convected by a single transport equation under the Eulerian framework, the second ‘‘Eulerian’’ denotes that the liquid phase continuity and momentum balance equations are employed. An additional transport equation was included to address bubble coalescence and breakage. The fluid phase equations and the additional transport equation for bubble coalescence and breakage are coupled by the drag force. We then validate the model predictions by comparison with predictions and experiments obtained in two classical gas-liquid bubble columns and one relatively new airlift column reported in the literature. To the author’s best knowledge this method, although quite straightforward but was never published before nor implemented nor validated. It is eventually worth mentioning that the model has been implemented in OpenFOAM and is available for download and use.

The remainder of the document is as follows: in Section 2, the governing equations and numerical discretization of the Q-E-E method are discussed. The solver development and implementation are also described. In Section 3 and 4, the Q-E-E method solver is validated against three test cases. Finally, conclusions are drawn in Section 5.

2. Governing equations

The multiphase system is composed by different phases. It is common to assume the liquid phase as continuous phase and model it under the Eulerian framework [4]. In this manner, without taking account for mass transfer, the mass conservation equation for the incompressible liquid phase (continuous phase) is described by the following continuity equation:

$$\frac{\partial \alpha_c}{\partial t} + \nabla \cdot (\alpha_c \mathbf{U}_c) = 0 \quad (1)$$

where α_c is the phase fraction of the liquid phase, and \mathbf{U}_c is the velocity. The momentum conservation of the liquid phase can be described by the volume-averaged incompressible momentum equation:

$$\rho_c \frac{\partial (\alpha_c \mathbf{U}_c)}{\partial t} + \rho_c \nabla \cdot (\alpha_c \mathbf{U}_c \otimes \mathbf{U}_c) = -\alpha_c \nabla p + \alpha_c \nabla \cdot \boldsymbol{\tau}_c + \alpha_c \mathbf{g} - \mathbf{M}, \quad (2)$$

where p is the average pressure, ρ_c is the density, \mathbf{g} is the gravity acceleration vector, and \mathbf{M} is the momentum interface exchange term to account for the momentum exchange between the gas phase and liquid phase, which will be discussed in the next section; $\boldsymbol{\tau}_c$ represents the stress tensor, which can be calculated by:

$$\boldsymbol{\tau}_c = \mu_c [\nabla \mathbf{U}_c + (\nabla \mathbf{U}_c)^T] - \frac{2}{3} \mu_c (\nabla \cdot \mathbf{U}_c) \mathbf{I} - \frac{2}{3} \rho k \mathbf{I}. \quad (3)$$

where k is the turbulent kinetic energy. Equation (1) and (2) are employed in the E-L, E-E and E-QBMM to predict the liquid phase flow field.

For the gas phase, it is quite common to address it as a disperse phase. The dispersed phase can be modelled with different procedures. In the Lagrangian framework, the position and velocity of the bubbles can be computed by solving the force balance over the individual bubbles:

$$\frac{d\mathbf{X}}{dt} = \mathbf{U}_d, \quad (4)$$

$$m_d \frac{d\mathbf{U}_d}{dt} = \mathbf{F}, \quad (5)$$

where \mathbf{X} is the position vector, \mathbf{U}_d is the bubble velocity, m_d is the bubble mass, and \mathbf{F} is the force exerted on discrete bubbles, which requires a certain degree of modelling. Here we assume the bubble mass is constant and that there is no net momentum exchange due to mass transfer. Among the many relevant fluid-bubble interaction forces, the most important is produced by the stresses of the continuous phase on the bubble surface, resulting in the drag and buoyancy forces. The balance of these forces determines the terminal bubble velocity. The bubble phase fraction (α_d) and liquid phase fraction (α_c) can be updated by using the following equations:

$$\alpha_d = \frac{\sum V_d}{V_{\text{cell}}}, \alpha_c = 1 - \alpha_d, \quad (6)$$

where V_d is the single bubble volume in a cell and V_{cell} is the cell volume.

The gas phase can be also modelled in the Eulerian framework resulting in equations similar to Eq. (1) and Eq. (2):

$$\frac{\partial \alpha_d}{\partial t} + \nabla \cdot (\alpha_d \mathbf{U}_d) = 0 \quad (7)$$

where α_d is the phase fraction of the gas phase, and \mathbf{U}_d is the average gas bubble velocity. The momentum conservation can be described by the volume-averaged incompressible momentum balance equation:

$$\rho_d \frac{\partial(\alpha_d \mathbf{U}_d)}{\partial t} + \rho_d \nabla \cdot (\alpha_d \mathbf{U}_d \otimes \mathbf{U}_d) = -\alpha_d \nabla p + \nabla \cdot \tau_d + \alpha_d \mathbf{g} + \mathbf{M}, \quad (8)$$

where ρ_d is the density and τ_d represents the stress tensor.

On the other hand, one can also use the generalized population balance equation for the gas phase, which can be written as follows [46]:

$$\frac{\partial f}{\partial t} + \nabla_{\mathbf{x}} \cdot (\mathbf{U}_d f) + \nabla_{\mathbf{U}_d} \cdot (\mathbf{A} f) = \mathcal{S}, \quad (9)$$

where f is the number density function for the gas phase phase, characterizing the number of bubbles per unit volume with a certain velocity and size, \mathbf{A} is the acceleration term, which is given by:

$$\mathbf{A} = -\frac{1}{\rho_d} \nabla p + \mathbf{g} + \frac{1}{\rho_d} \mathbf{M}. \quad (10)$$

The solution of Eq. (9) is usually achieved in terms the moments of the number density function and the corresponding closure problem is usually overcome by using a quadrature approximation.

Equation (4) and Eq. (5) can be solved via a meshless method by updating the Lagrangian bubble location without any numerical problems. Equations (1) - (6) constitute the set of equations employed in any E-L method. In the E-L method, numerical problems arise and need special treatments. For example, the phase fraction is a strictly bounded value ($0 \leq \alpha_d < 1$). It should be smaller than one otherwise a vacuum state of the liquid phase can occur in certain cells. This also explains the reason why the bubble size should be smaller than the cell size and why a two-dimensional E-L simulation is not strictly applicable. Meanwhile, the bubble location tracking algorithm should be highly efficient otherwise the whole algorithm would be quite slow. It is suggested to employ the face penetrating method to locate the bubble location, but it sometimes fails and the bubble disappears if the bubble moves just through the cell vertices instead of the cell faces [14]. Equations (1), (2), (7) and (8) constitute the equations typically employed in the E-E method. The resulting transport

equations are quite complex and difficult to solve. Depending on the source terms, the mathematical characteristic also changes and the numerical scheme should respect it. Equations (1), (2) and (??) constitute the equations employed in the E-QBMM. This equation system is not closed and sub-models are necessary. Moreover, it is not trivial to maintain the realizability of the moments if a high-order scheme is used [58, 35, 54].

2.1. Gas phase velocity formulation

The governing equations employed in the Q-E-E method can be derived from the existing method using justifiable approximations. It can be seen from Eq. (1) and Eq. (2) that once the phase fraction α_d is calculated, they can be used to solve for the continuous phase velocity \mathbf{U}_c and pressure p . In the E-E method, α_d is convected by Eq. (7), in which \mathbf{U}_d is unknown. In the E-QBMM, the unknown \mathbf{U}_d is calculated from the moments, this method was also called the velocity-polynomial-approximation (VPA) method. On the other hand, considering a stagnant liquid environment ($\mathbf{U}_c = 0$), it was proven that the balance between the buoyant force and the drag force can be achieved. Eventually, the bubble will ascent at a nearly constant velocity, namely the so-called terminal velocity. The force balance equation can be written as follows:

$$\rho_d V_d \mathbf{g} \left(1 - \frac{\rho_c}{\rho_d} \right) = \frac{3}{4} C_D \frac{\rho_c}{d} |\mathbf{U}_d| \mathbf{U}_d V_d, \quad (11)$$

where V_d is the volume of single bubble, d is the bubble diameter, and C_D is the drag coefficient which requires a certain degree of modelling and different models were proposed. For example, in the Schiller and Naumann model, it can be calculated as follows:

$$C_D = \begin{cases} 24 (1 + 0.15 \text{Re}^{0.687}) / \text{Re}, & \text{Re} \leq 1000, \\ 0.44, & \text{Re} > 1000, \end{cases} \quad (12)$$

where Re is the Reynolds number defined by:

$$\text{Re} = \frac{\rho_c d |\mathbf{U}_c - \mathbf{U}_d|}{\mu_c}, \quad (13)$$

where μ_c is the viscosity of the liquid phase. Substituting Eq. (12) into Eq. (11), the bubble terminal velocity can be calculated. Simulations showed that the bubble terminal velocity and the drag coefficient are strongly dependent on the bubble diameter under turbulent conditions [62, 66]. For air bubbles in clean water with size ranging from 3 mm to 5 mm, the bubble terminal velocity is approximately constant and equal to 0.23 m/s. If the Reynolds number is larger than 2000, C_D ranges from approximately 0.3 to 3 as measured in experiments. Therefore, it is safe to calculate the bubble velocity from the following equation:

$$\mathbf{U}_d = \mathbf{U}_c + \mathbf{U}_s, \quad (14)$$

where \mathbf{U}_s is the slip velocity. This slip velocity assumption was also used in the mixture/drift-flux model. A similar approximation was also employed in other works and the justification was proven by several authors [4, 19]. In this work, we adopt a similar approach and we assume a constant slip velocity due to the reason that the bubble terminal velocity typically ranges only from 0.2 m/s to 0.25 m/s. A similar procedure was also adopted in other works [63]. In this manner, the gas phase velocity can be calculated and the capability of this method is investigated in this work. This is the first procedure investigated in this work.

Besides Eq. (14), an iterative procedure can be used to obtain the gas phase velocity, which is more consistent with the theory. In the field of E-L and E-QBMM, the gas phase velocity can be updated with the following ordinary differential equation (ODE):

$$\frac{d\mathbf{U}_d}{dt} = \mathbf{g} - \frac{1}{\rho_d} \nabla p_c + \frac{1}{\tau} (\mathbf{U}_c - \mathbf{U}_d), \quad (15)$$

where τ is the relaxation time. Substituting the Bernoulli equation:

$$\nabla p_c = \rho_c \mathbf{g} \quad (16)$$

into Eq. (15) leads to:

$$\frac{d\mathbf{U}_d}{dt} = \mathbf{g} \left(1 - \frac{\rho_c}{\rho_d} \right) + \frac{1}{\tau} (\mathbf{U}_c - \mathbf{U}_d). \quad (17)$$

When the Euler-implicit time scheme is employed to solve Eq. (17), the velocities for the next time step can be written as follows:

$$\mathbf{U}_d^{t+\Delta t} = \frac{\mathbf{U}_d^t + \Delta t (S_p \mathbf{U}_c + S_u)}{1.0 + S_p \Delta t}, \quad (18)$$

where

$$S_u = \mathbf{g} \left(\frac{\rho_d - \rho_c}{\rho_d} \right), S_p = \frac{3}{4} \frac{\mu_c}{\rho_d d^2} C_D \text{Re}. \quad (19)$$

Since Eq. (19) is dependent on Re , which is further dependent on \mathbf{U}_d , an iterative procedure should be adopted. In practise, Eq. (19) can be solved 5 - 10 times to obtain a convergent gas phase velocity. Since no differential equation is evolved, the iterative procedure is quite fast and can be easily implemented. In this manner, the gas phase velocity can be obtained without assigning the value of the slip velocity. It should be noted here that other momentum interfacial exchange terms (e.g., lift force) can be included in S_u as reported in Eq. (19) and the solving procedure is straightforward. This is the second procedure investigated in this work.

Another problem that arises is how to model the bubble path dispersion. It is well known that the width of the rising bubble swarm increases with the height of the column. Thus, radial mixing takes place in the gas phase. But this effect cannot be recognized because only a convective term exists in the gas phase transported equation. Under turbulent flow conditions, Eq. (7) should be averaged and the following equation can be obtained:

$$\frac{\partial \alpha_d}{\partial t} + \nabla \cdot (\alpha_d \mathbf{U}_d) + \nabla \cdot (\overline{\alpha'_d \mathbf{U}'_d}) = 0, \quad (20)$$

where $\overline{\alpha'_d \mathbf{U}'_d}$ needs to be modelled. The last term on the left-hand side of Eq. (20) allowed us to have a turbulent diffusion term in the phase transported equation. In this work, we adopted the approach extensively employed in the field of slurry simulations [13, 49]. In this model, the turbulent dispersion term was modelled by the following assumption:

$$\nabla \cdot (\overline{\alpha'_d \mathbf{U}'_d}) = -\nabla \cdot \left(\frac{\nu_t}{\sigma} \nabla \alpha \right), \quad (21)$$

where ν_t is the turbulent kinematic viscosity of the liquid phase, determined in turn by turbulence model, and σ is the turbulent Schmidt number for gas bubble phase fraction. The turbulent Schmidt number may be interpreted as the ratio of turbulent momentum transport to the turbulent transport

of the mass associated to the gas phase. However, the numerical value of it is not well established in the literature because no single constant value can be used. In the proposed model, we simply set the turbulent Schmidt number for volume fractions equal to 1.0, since this value results in the best overall agreement with the experimental data. The turbulent kinematic viscosity of the liquid can be calculated from the turbulence model. In this work, the standard $k - \varepsilon$ model was employed, which can be written as follows:

$$\frac{\partial \varepsilon}{\partial t} + \nabla \cdot (\mathbf{U}\varepsilon) - \nabla \cdot (D_\varepsilon \nabla \varepsilon) = C_1 G \frac{\varepsilon}{k} - \frac{2}{3} C_1 \varepsilon \nabla \cdot \mathbf{U} - C_2 \frac{\varepsilon}{k} \varepsilon, \quad (22)$$

$$\frac{\partial k}{\partial t} + \nabla \cdot (\mathbf{U}k) - \nabla \cdot (D_k \nabla k) = G - \frac{2}{3} \nabla \cdot \mathbf{U}k - \varepsilon, \quad (23)$$

where

$$G = \nu_t \left(\nabla \mathbf{U} + \nabla \mathbf{U}^T - \frac{2}{3} (\nabla \cdot \mathbf{U}) \mathbf{I} \right) : \nabla \mathbf{U}. \quad (24)$$

Once the k and ε equation was solved, the turbulent viscosity can be calculated by

$$\nu_t = C_\mu \frac{k^2}{\varepsilon}. \quad (25)$$

2.2. Polydispersity and bubble size distribution

To account for bubble breakage and coalescence, another model is needed. The QMOM or EQ-MOM was implemented in our previous works but they are not suitable for this work due to the large demand of computational resources [42, 41, 39]. Therefore, we adopted a simpler procedure as reported in [33, 34]. In this method, the total bubble number density is calculated as follows:

$$n = \frac{\alpha_d}{\pi/6d^3}. \quad (26)$$

From the values of n at each point in the computational domain, the local average bubble size d can be calculated. An additional transport equation is solved for n , accounting for the transport of bubbles (by convection and turbulent diffusion) and changes in bubble size (and therefore bubble number) by breakage and coalescence according to [33, 34]:

$$\frac{\partial n}{\partial t} + \nabla \cdot (\mathbf{U}_d n) - \nabla \cdot (\nu_t \nabla n) = S_{\text{break}} - S_{\text{coal}}, \quad (27)$$

where S_{break} is the source term to account for bubble breakage and S_{coal} is the sink term to account for bubble coalescence. The rate of bubble break-up is assumed to be related to the collision rate with turbulent eddies of similar size, since larger eddies merely transport the bubbles and smaller eddies have insufficient energy. The collision rate with eddies depends on the turbulent velocity, for which the bubble size is assumed to lie within the inertial sub-range of turbulence. Additionally, break-up is conditional on exceeding a critical value of bubble Weber number, We_c , representing the ratio of turbulent shear forces to the restoring surface tension forces. Following these considerations, the expression for the break-up is assumed to be: [33, 34, 69]

$$S_{\text{break}} = C_{\text{br}} \frac{U_t}{1.4d} \sqrt{1.0 - \frac{-We_c}{We}} \exp\left(\frac{-We_c}{We}\right), \quad We > We_c, \quad (28)$$

where C_{br} is user-defined breakage coefficient, and U_t is taken to be the velocity of eddies in the inertial sub-range of the turbulent eddy spectrum, which may be written as:

$$U_t = 1.4(\varepsilon d)^{1/3}. \quad (29)$$

The bubble Weber number defines the tendency for bubbles to break up or remain stable, which can be written as:

$$We = \frac{\rho_c U_t d}{\sigma}, \quad (30)$$

where σ is the surface tension between the gas and liquid phases. The bubble coalescence can be modelled by random collision process or wake induced collision process. In this work, we consider random collision, though the wake induced collision can be included straightforwardly. In the theory of random collision, the bubble coalescence sink term can be written as [33, 34, 69]

$$S_{coal} = n^2 C_{turb} U_t \frac{d^2}{\alpha_{max}^{1/3} \alpha_d^{1/3}} \left(1 - \exp \left(-C_{turb2} \frac{\alpha_{max}^{1/3} \alpha_d^{1/3}}{\alpha_{max}^{1/3} \alpha_d^{1/3} - \alpha_{max}^{1/3} \alpha_d^{1/3}} \right) \right), \quad (31)$$

where C_{turb} , C_{turb2} are user-defined coalescence coefficients, and α_{max} is the maximum packing limit for the bubbles. Eq. (20), Eq. (14), Eq. (1), Eq. (2) together with the equations of the turbulent model Eq. (22), Eq. (23) and the bubble diameter equation Eq. (27) form the governing equations of the Q-E-E method, which is suitable for polydisperse turbulent bubbly flows. It can be solved in the same way as in the single phase flow by some iterative procedure under the finite volume method, which will be discussed in the next section.

2.3. Numerical discretization

In this section, we summarise the iterative solving procedure based on the PISO algorithm [28]. The solving procedure of the Q-E-E method resembles the procedure of the E-QBMM and the E-L method. The spirit of PISO algorithm is to use the momentum to obtain a non-divergence free velocity field, then use the continuity equation to construct a pressure Poisson equation to update the velocity field to ensure divergence free. The pressure equation is suggested to be solved 2 - 3 times within a time step. Special numerical techniques are necessary to prevent oscillations, especially when there are gradient terms and large body forces in the source term of the momentum equation.

In order to obtain a smooth solution, the pressure term (gradient term), gravity term and others are neglected at this step and the discretized form of Eq. (2) can be written as follows:

$$a_P \mathbf{U}_P + \sum a_N \mathbf{U}_N = S_P, \quad (32)$$

where a_P is the matrix diagonal coefficients, a_N is the matrix non-diagonal coefficients, \mathbf{U}_P is the unknown velocity defined at the cell center, and \mathbf{U}_N is the known velocity for the cell neighbours. The subscript $_d$ was neglected for brevity. Eq. (32) constitutes a matrix system which can be solved by any sparse linear system to obtain the predicted velocity. However, in practise, it is not solved and this step can be safely replaced by a one step Jacobian point iterative procedure. The solution of this one step iterative procedure produces the predicted velocity:

$$\mathbf{HbyA}_P = \frac{-\sum a_N \mathbf{U}_N + S_P}{a_P}. \quad (33)$$

Due to the crude approximation, \mathbf{HbyA} is meaningless and needs to be corrected. The convergent velocity and pressure should obey the continuity equation. The discretized continuity equation can

be written as

$$\frac{(\alpha_{c,P}^{t+\Delta t} - \alpha_{c,P}^t)\Delta V_P}{\Delta t} + \sum (\alpha_{c,f} \mathbf{U}_f^{t+\Delta t} \cdot \mathbf{S}_f) = 0, \quad (34)$$

where Δt is the time step, and ΔV_P is the cell volume. Owing to the Rhie and Chow interpolation, the cell face velocity can be expressed by [56]

$$\mathbf{U}_f^{t+\Delta t} = \mathbf{HbyA}_f - \frac{1}{a_P} \alpha_{c,f} (\nabla p)_f + \alpha_{c,f} \frac{1}{a_P} \mathbf{g} - \frac{1}{a_P} \mathbf{M}_f. \quad (35)$$

Substituting Eq. (35) into Eq. (34) leads to

$$\sum \frac{\alpha_{c,f}^2}{a_P} (\nabla p)_f = \frac{(\alpha_{c,P}^{t+\Delta t} - \alpha_{c,P}^t)\Delta V_P}{\Delta t} + \sum \left(\alpha_{c,f} \left(\mathbf{HbyA}_f + \frac{\alpha_{c,f}}{a_P} \mathbf{g} - \frac{1}{a_P} \mathbf{M}_f \right) \cdot \mathbf{S}_f \right). \quad (36)$$

The continuous form of Eq. (36) is the so-called pressure Poisson equation and its solution gives the pressure of the continuous phase. It should be noted here that the term associated with \mathbf{g} is the gravity/buoyancy term and the term associated with \mathbf{M} is the momentum exchange interfacial term. In some works, it was claimed that the effect of the buoyancy term is larger than that of the momentum exchange term. In this work, the effect of the buoyancy term was evaluated. In summary, the solving procedure consists of the following steps:

1. From the initial field, calculate the momentum exchange term and construct the velocity matrix system as reported in Eq. (32); Calculate the predicted velocity as reported in Eq. (33) by the one point Jacobian method;
2. Solve the pressure Poisson equation as reported in Eq. (36) to update pressure.
3. Reconstruct the velocity from the pressure as reported in Eq. (35). Repeat step 2 and 3 two or three times and a convergent velocity/pressure field of the continuous phase can be obtained.
4. Update the dispersed phase velocity by using Eq. (14) (i.e. constant slip or procedure 1) or Eq. (18) (i.e. ODE or procedure 2) and calculated the phase transported equation as reported in Eq. (20). Update the continuous phase fraction by $\alpha_c = 1 - \alpha_d$.
5. Correct the turbulence equations and bubble size equation and move to the next time step.

The implementation of the algorithm mentioned previously is straightforward and it is briefly discussed here. The velocity matrix system can be constructed by:

```
fvVectorMatrix UcEqn
(
    fvm::ddt(alphaC, Uc) + fvm::div(alphaPhiC, Uc)
  - fvm::Sp(fvc::ddt(alphaC) + fvc::div(alphaPhiC), Uc)
  + continuousPhaseTurbulence->divDevRhoReff(Uc)
  ==
  - fvm::Sp(dragSp, Uc)
);
```

in which the drag force was discretized semi-implicitly. The predicted velocity \mathbf{HbyA} can be solved by

```
HbyA = UEqn.H()/UEqn.A();
```

The face velocity of the continuous phase as reported in Eq. (35) can be implemented by

```
Kd = 0.75*alphad*muc/sqr(diameter)*CdRe;
```

```

preDispersed = Kd*Ud;
surfaceScalarField phicForces
(
    fvc::flux(rAUc*preDispersed/rhoc)
    - (
        fvc::interpolate(rAUc*(Su)) & mesh.Sf()
    )
    + alphacf*rAUcf*(g & mesh.Sf())
);
surfaceScalarField phiHbyA
(
    "phiHbyA",
    (
        fvc::flux(HbyA)
        + alphacf*rAUcf*fvc::ddtCorr(Uc, phic)
        + phicForces
    )
);

```

where $CdRe$ is $C_D Re$. Afterwards, the pressure Poisson equation can be implemented by

```

fvScalarMatrix pEqn
(
    fvm::laplacian(sqr(alphacf)*rAUcf, p)
    ==
    fvc::ddt(alphac) + fvc::div(alphacf*phiHbyA)
);

```

and the velocity can be reconstructed by

```

Uc = HbyA
    + rAUc*fvc::reconstruct((phicForces - pEqn.flux())/alphacf)/rAUcf);

```

All the governing equations are implemented in the open-source CFD code OpenFOAM and wrapped into a new solver called QEEFoam. In this work, the code and test cases are open-sourced. Readers are referred to the supplementary files for download.

3. Test cases and numerical details

The predicted results obtained with the QEEFoam are compared with the measurements available in the literature both for qualitative and quantitative comparisons. The first test case (Test A) simulated in this work is a partially aerated rectangle bubble column investigated by Becker et al. [6]. The second test case (Test B) is a rectangle bubble column investigated by Díaz et al. [16]. The third test case (Test C) is an airlift bubble column investigated by Mandalahalli et al. [44]. The first two cases are characterised by a simple geometry with a complicated flow structure, which are suitable for numerical investigation. Depending on the experimental conditions, the flow regimes resulting from time/volume-averaged procedures can differ considerably from the unsteady flow structures. The fully developed vortex flow has been proved to be a very useful tool to evaluate the hydrodynamic properties of bubble columns. The last test case is rather new and the geometry is quite different from the previous two. It could be a good test case for further validation.

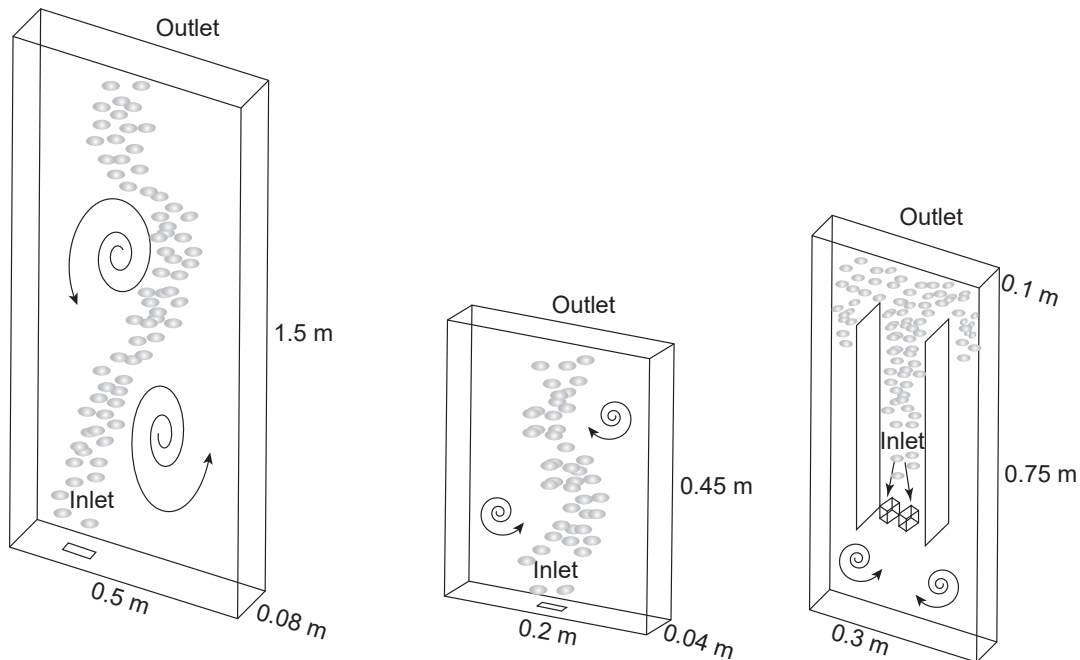


Figure 1: Bubble column geometry investigated experimentally by Becker et al. (left) [6], Díaz et al. (middle) [16]. Airlift column geometry investigated experimentally by Mandalahalli et al. (right) [44].

The column was filled with tap water (i.e. continuous liquid phase) at room temperature and atmospheric pressure, while the air was fed through an aluminium sparger. A sketch of the setup is shown in Fig. 1. A fully orthogonal hexahedral grid measuring 52 (height) \times 32 (width) \times 18 (depth) cells for test A and 40 (height) \times 21 (width) \times 9 (depth) cells for test B were generated by `blockMesh` utility, as we confirmed that these non-uniform hexahedral grids are enough to capture the plume oscillations in our previous investigations. A similar mesh resolution was also reported in other works [16, 37, 52, 50]. The sparger inlet was generated by `topoSet` and `createPatch` utilities, through which gas enters the domain was modelled as a rectangle with an area equal to 0.001197 m² and 0.00021 m², respectively. This simplification has been proven to be efficient for meshing, and previous works have shown that the results are not sensitive to the inlet shape [16, 9]. For test case C, a non-uniform hexahedral grid was employed. The diameter of bubbles in test A, B and C is 3 mm, 5.05 mm and 2.2 mm, respectively. Different gas superficial velocities were used and will be reported in the following sections. The buoyancy force, drag force and the turbulent dispersion force were considered in all the test cases in the Q-E-E simulations. Second order with a limiter convection scheme was used for all variables. In our test cases, the unbounded problem of the phase fraction and bubble number density was not shown.

Some results were also compared with that predicted by the OpenFOAM E-L solver `DPMFoam`. In the E-L simulations, rebounded wall boundary and “escape” boundary were given for the bubbles. The bubble number injected per second to the computational domain was calculated from the superficial gas velocity as

$$n = \frac{\mathbf{U}_s \mathbf{S}_{\text{outlet}}}{V_{\text{bubble}}}, \quad (37)$$

where n the bubble number injected per second, $\mathbf{S}_{\text{outlet}}$ is the surface vector, \mathbf{U}_s is the gas superficial velocity, and V_{bubble} is the bubble volume. Sizes of the bubble were assumed to be uniform. Bubble

collision is not considered and therefore a two-way coupling was assumed. In all the simulations, the liquid phase was assumed to be stagnant at the beginning. The non-slip velocity boundary condition was given for the liquid velocity. For the liquid pressure, a zero gradient with the hydrodynamic correction was applied. The standard turbulent wall function was applied for turbulent variables.

As previously mentioned two algorithms are tested for the calculation of the gas bubble velocity:

1. by assuming a constant gas-liquid slip velocity and by using Eq. (14),
2. by updating the gas bubble velocity with the solution of the ODE discretized in Eq. (18).

4. Results and discussions

4.1. Test case A

In this test case, the gas was injected on the left side of the bottom. One can certainly expect that vortexes may exist due to the non-symmetric injection. It was also shown in previous works that the dynamic liquid circulation is quite complex and can only be predicted by three-dimensional simulations [50, 63]. Therefore, our algorithm and implementation should capture such quasi-periodic transient liquid circulations. The predicted and measured horizontal liquid velocities for test case A at different positions with slip velocity method and ordinary differential equation method were reported in Fig. 2 and table 1. It can be seen that the quasi-periodic transient liquid circulations were successfully captured. The horizontal liquid velocity at point A is slightly larger than that predicted for point B since point A locate at the left side, which is in good agreement with that as measured in the literature [6, 63]. The effect of the turbulent dispersion term on the period of bubble plum oscillation (POP) is negligible (within 10 %) as it is seen in table 1. Meanwhile, it can be found that the user-defined slip velocity has a substantial effect on the POP. A larger slip velocity corresponds to large relative velocity and the coupling between the gas and liquid phase is stronger, which implies a small relaxation time and smaller POP. It was also shown that a vertical slip of 0.2 m/s predicted the best results compared with the experimental data. Such slip velocity value agrees well with the experimental founding as mentioned in section 2. On the other hand, the ODE method predicts quite good POP as well, since the gas phase velocity was predicted by solving the ODE system and the user-defined slip velocity is not necessary.

	With dispersion term	Without dispersion term
Slip vel. method 0.15 m/s	54 s	53 s
Slip vel. method 0.2 m/s	40 s	37.5 s
Slip vel. method 0.25 m/s	35 s	30 s
ODE method	39 s	37 s

Table 1: Comparison of predicted and experimentally measured POP for the test case A with/without turbulent dispersion term. Different vertical slip velocity was used. Measured POP: 41 s.

The prediction of the oscillating phase fraction, velocity, and streamlines coloured by pressure at different time instants ($\Delta t = 2$ s) for the test case A by the slip velocity method are reported in Fig. 3. Results predicted by the ODE method was not shown for brevity since the difference is negligible. They show a surprisingly good agreement with experiments. Several liquid circulation cells can be observed in the column, typically no more than 4 circulation zones can be found. The vortex at the top tends to increase its size. When the upper vortex reaches a critical size, it starts to move downwards. The size of the largest vortex is as large as half of the column. Once these large vortexes reached the bottom, they are smeared due to the wall and new vortexes are generated at the top.

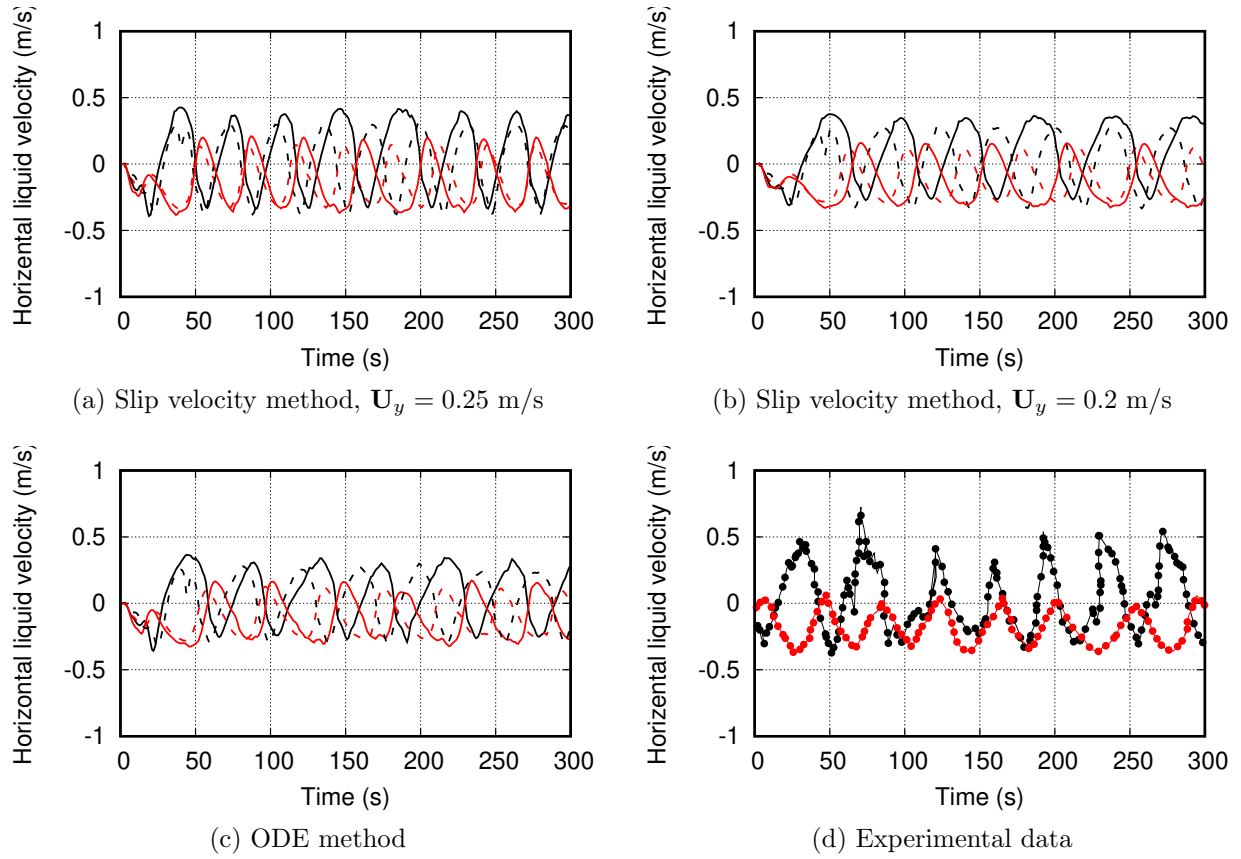


Figure 2: Calculated horizontal liquid velocities for test case A at different positions by slip velocity method and ODE method. Black line: (0.035 0.9 0.04), red line: (0.45 1.05 0.04). Solid line: with turbulent dispersion term. Dashed line: without turbulent dispersion term.

At this point, the "cycle" is repeated again. Meanwhile, the center of the vortex has a relatively low pressure, which is consistent with the vortex theory. It can be seen that such a dynamic vortex movement can be surprisingly captured by our code. On the other hand, the transient phase fraction shows similar oscillating plumes. After the gas was injected from the bottom, it was dispersed by the turbulent dispersion term and moved forward due to buoyancy. The time-averaged vertical liquid velocity by the slip velocity method and ODE method at different positions were reported in Fig. 4. The predicted results by the ODE method seem to be better, whereas those predicted by the slip velocity method (0.25 m/s) were overshoot and those predicted by the slip velocity method of 0.15 m/s were undershot. On the other hand, when the slip velocity method was 0.2 m/s. Good agreements with experimental results [6, 63] were observed. This test case shows that the flow field can be successfully predicted by the slip velocity method and the ODE method, where the results may be influenced by the user-defined slip velocity and a fitting procedure may be necessary for the first one. All these results show that our method is capable of predicting the turbulent bubbly plume.

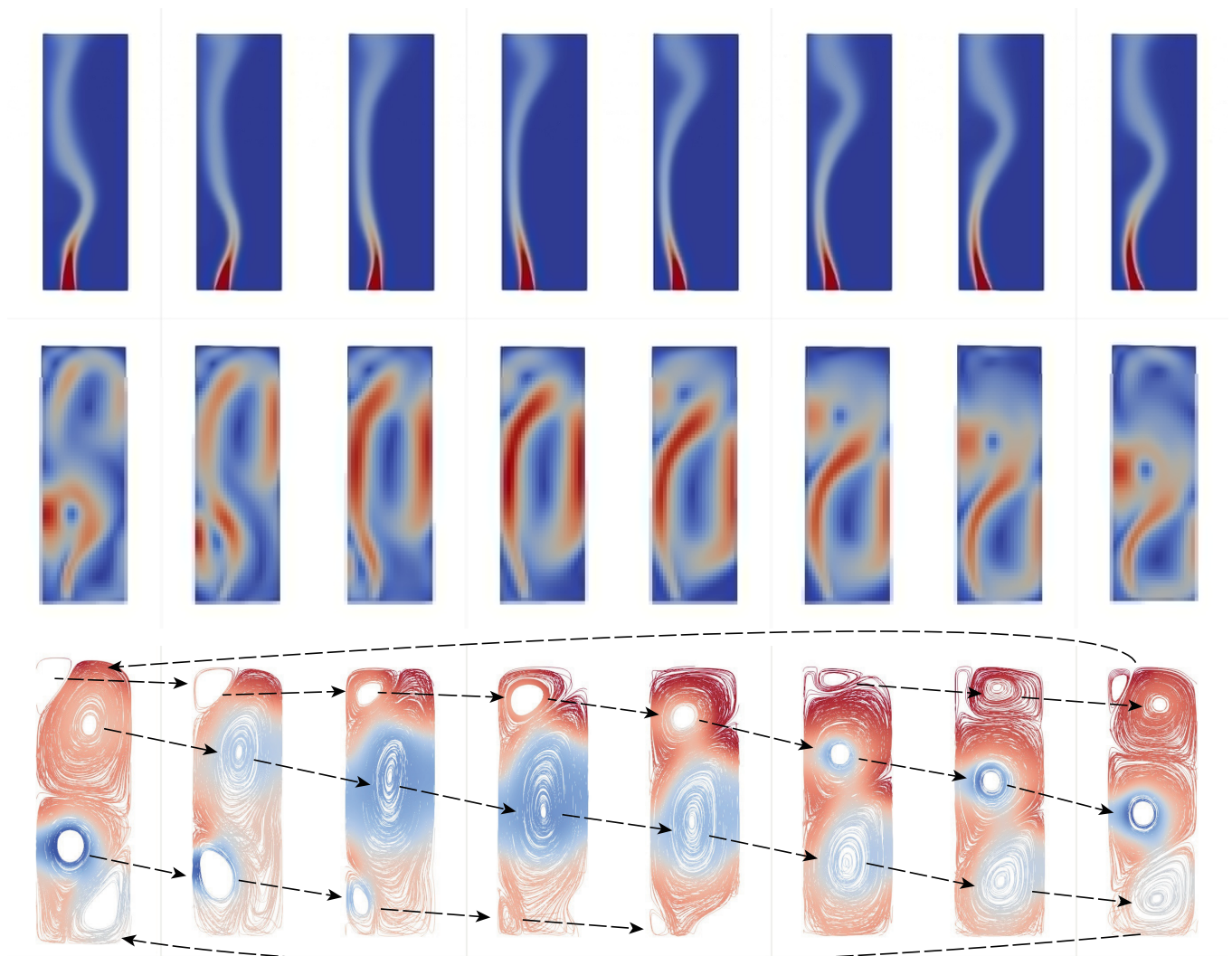


Figure 3: Prediction of the oscillating phase fraction (top, [0:0.03]), velocity (middle, [0:0.4]), and streamlines (bottom, colored by pressure, [-0.07:0.04]) at different time instants ($\Delta t = 2$ s) for the test case A.

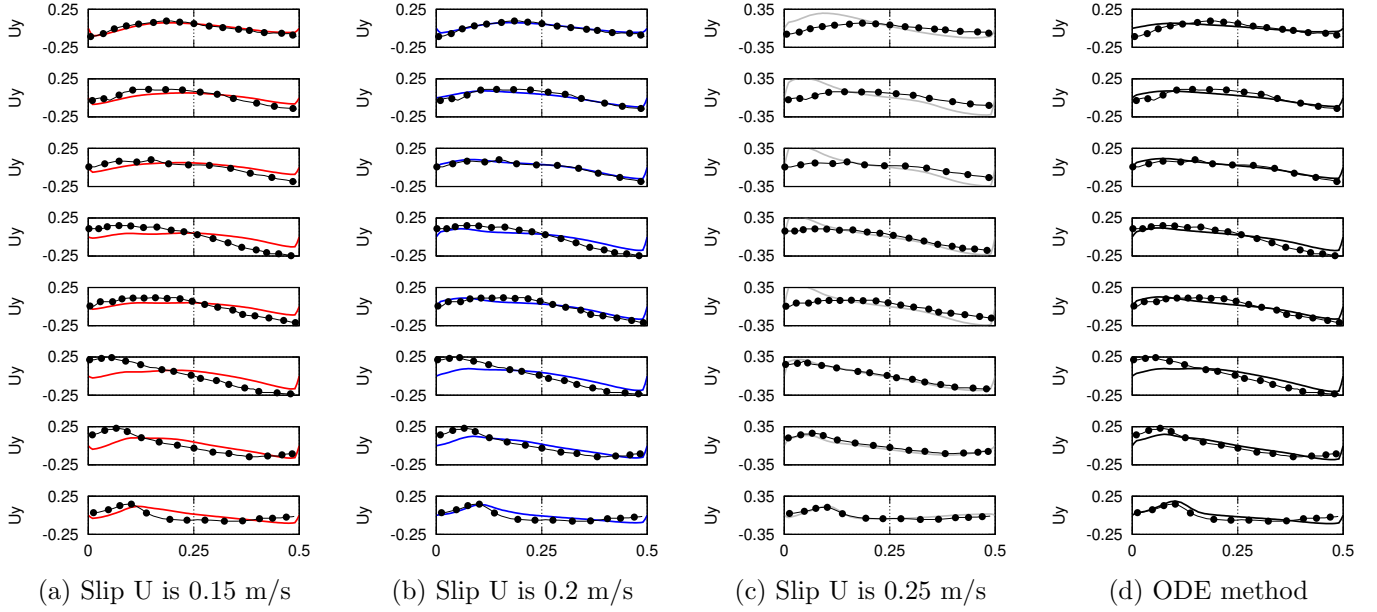


Figure 4: Comparison of the time-averaged upward liquid velocity with experimental data (points) at different heights on the mid-depth plane (from bottom to top: $y/H = 0.11, 0.22, 0.35, 0.49, 0.6, 0.7, 0.78, 0.89$).

4.2. Test case B

In this test case, the gas was injected in the middle of the bottom, which implies the geometry is quite symmetric. Nonetheless, it was also shown in previous work that the dynamic liquid circulation is quite complex, as also confirmed by experiments [16]. In the experiments, different gas superficial velocities were employed. The predicted horizontal liquid velocity at different positions was reported in Fig. 5. The drag model developed by Grace et al. was employed [24]. Since the slip velocity method was investigated in the previous case, in this test case, the magnitude of the slip velocity was assumed to be 0.25 m/s. Specifically, the predicted values of the gas POP were reported in table 2. It can be seen that the results predicted by the Q-E-E method agree well with the simulated results reported in the literature [16] and our previous works [37, 42, 9] in which the E-E and E-QBMM were used to simulate bubbly flows. Specially, the POP predicted by the slip velocity method is comparable with that predicted by the ODE method, which implies that a slip velocity of 0.25 m/s is suitable for this test case. It should be noted here that a 0.2 m/s slip velocity seems to be better for the previous case. This slip velocity acts as a tuning parameter in our method and a fitting procedure is required.

The predicted gas POP when the superficial velocity was assumed as 0.012 m/s with and without the gravity term exerted on the continuous phase is reported in Fig. 6. It can be seen that the predicted gas POP with or without the gravity term differs within 5%. Meanwhile, it was found that the momentum interfacial exchange term's effect is very important, which implies that the coupling of the gas and liquid phases by the momentum transfer is quite important. Similar predictions were also obtained with the E-L simulations solved by DPMFoam, which are not shown here for brevity. Such finding may contradict with others where the phase coupling was seen as more important than the momentum coupling. The prediction of the oscillating phase fraction and time-averaged phase fraction with and without turbulent dispersion term were reported in Fig. 7. It is obvious that the predicted gas phase fraction was quite sharp if the turbulent dispersion term was neglected in the transported equation, which is consistent with the theory. With the turbulent dispersion force, those

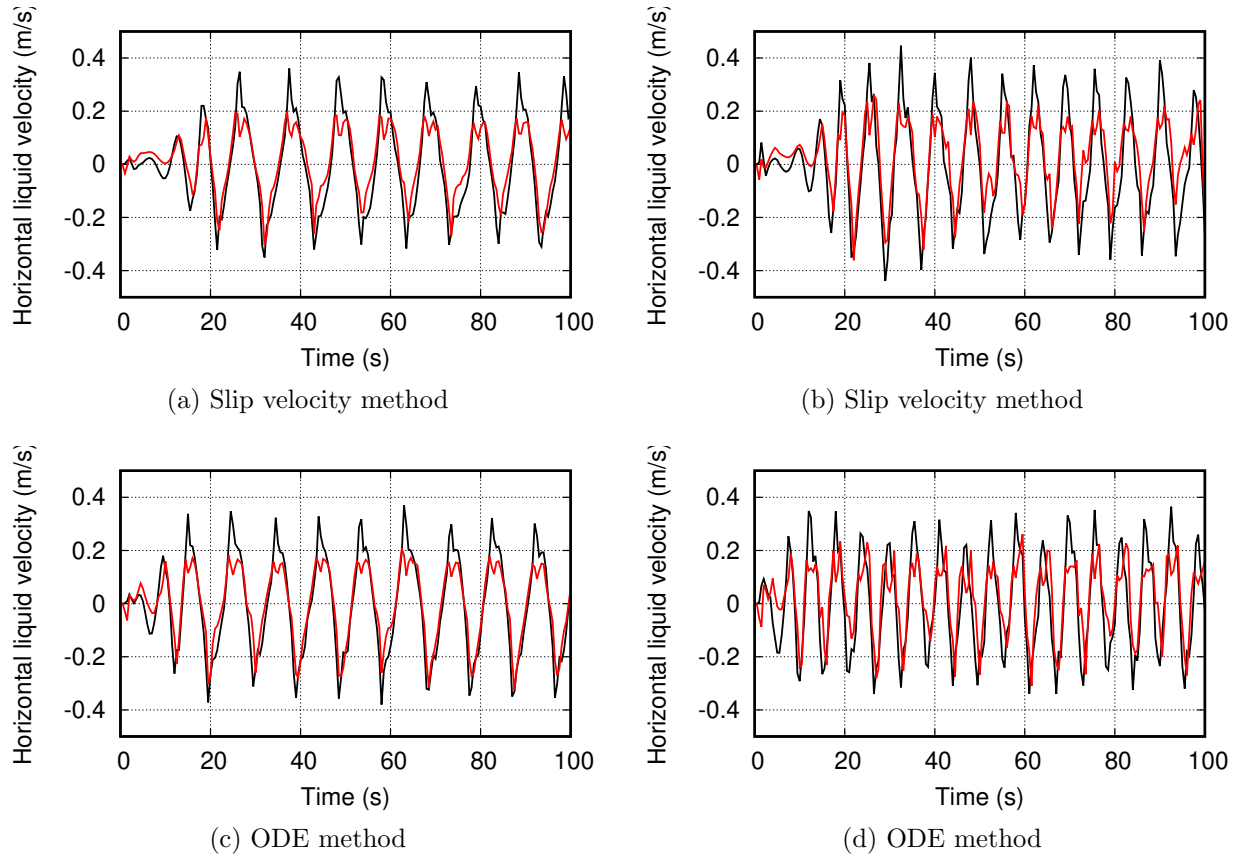


Figure 5: Calculated horizontal liquid velocity for test case B operated at different superficial gas velocities (left: 0.0024 m/s, right: 0.0071 m/s) at different positions (black line: $x = 0.1$ m, $y = 0.225$ m, $z = 0.02$ m; red line: $x = 0.05$ m, $y = 0.225$ m, $z = 0.02$ m) by slip velocity method ($|\mathbf{U}_s| = 0.25$ m/s) and ODE method.

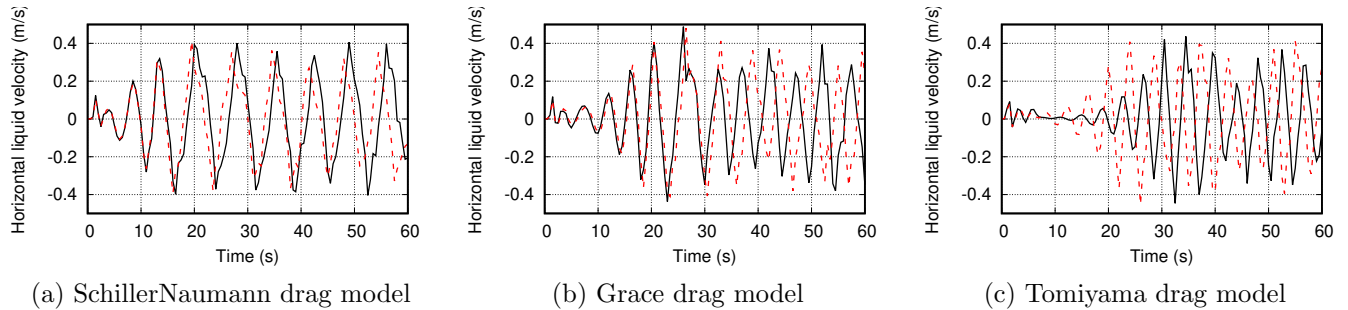


Figure 6: Calculated horizontal liquid velocity for test case B operated with (black solid line) or without (red dashed line) gravity term by slip velocity method. Different drag models were employed. Position: $x = 0.1$ m, $y = 0.225$ m, $z = 0.02$ m. Superficial gas velocities is 0.012 m/s.

bubbles also move deeper into the column. If one compares the predicted transient phase fraction with the experiments, it is also obvious that the predicted results with turbulent dispersion term are more accurate.

Gas superficial velocity	Experiments	Predictions by slip vel.	Predictions by ODE
0.0024 m/s	11.37 s	11.5 s	11.4 s
0.0071 m/s	5.69 s	6.3 s	5.45 s
0.012 m/s	4.3 s	4.9 s	4.1 s

Table 2: Comparison of predicted and experimentally measured POP for the test case B with turbulent dispersion term by the slip velocity method and ODE method.

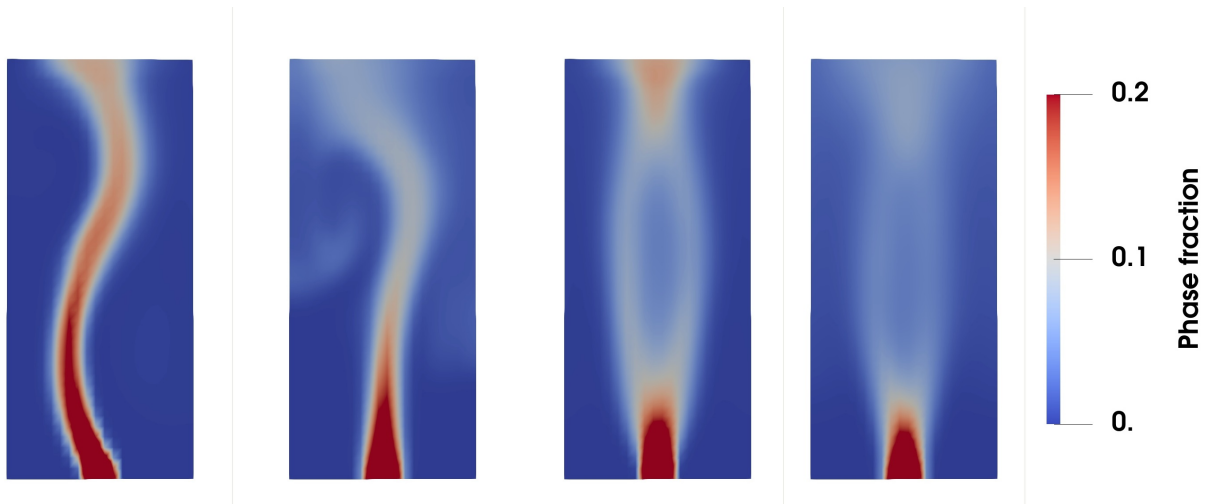


Figure 7: Prediction of the oscillating phase fraction and time-averaged phase fraction with and without turbulent dispersion term by the ODE method.

The time-averaged gas hold-up and vertical liquid velocity at different positions with and without turbulent dispersion term are reported in Fig. 8. Here the gas superficial velocity is 0.012 m/s. Different drag coefficients were employed and only the results predicted by the slip velocity method are reported. Predictions obtained by the E-L method are also used as a comparison. It can be seen that the phase fraction predicted without the turbulent dispersion term shows an over-estimated double peak, which can be also seen in Fig. 7. This double peak comes from the periodic oscillating flow fields. On the other hand, results predicted with the turbulent dispersion term by the Q-E-E method agree well with those predicted by the E-L method. The drag model also has an effect on the predictions but the effect can be neglected. The predicted vertical liquid velocities by the Q-E-E method also agree well with that predicted by the E-L method. It can be observed that the resulting time-averaged flow pattern consists of a non-uniform velocity distribution presenting an upward flow in the column center and a downward flow along the column walls, a liquid circulation mode. This type of flow differs considerably from the instantaneous flow pattern previously described. The calculated global gas hold-up by the Q-E-E method at different superficial gas velocities is reported in Fig. 9. It can be seen that the predictions agree well with the experiments. All these results reveal that the dynamic vortex flow fields in a symmetric geometry can be predicted by the Q-E-E method and the results are surprisingly good. The prediction of the bubble size distribution of test case B with different gas superficial velocities was reported in Fig. 10. It can be seen that the

bubble's diameter is larger at the center of the column than those at sides. Contour plots of the diameter show that, with the increase of the gas superficial velocity, smaller bubbles tend to stay in the liquid recirculation path, whereas bigger bubbles are concentrated into the central plume zone where coalescence and breakup occur with faster rates due to turbulence. Such results were also observed in other works where a quadrature-based moments method was used [7].

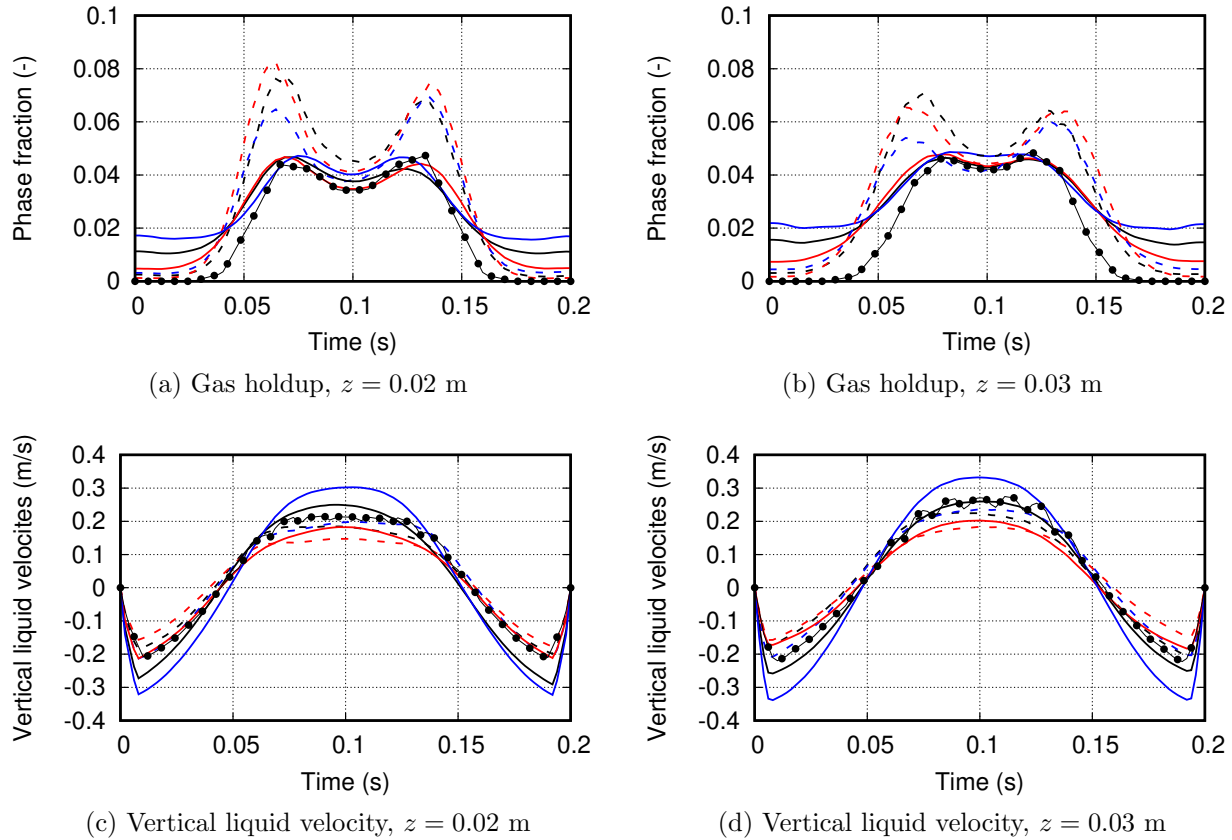


Figure 8: Time-averaged gas hold-up and vertical liquid velocity at different positions for test Case B. Gas superficial velocity is 0.012 m/s. Different drag coefficients were employed. Solid line: with turbulent dispersion term. Dashed line: without turbulent dispersion term by the slip velocity model. Black: Grace drag model, red: SchillerNaumann drag model, blue: Tomiyama drag model. Black line-points: data simulated by E-L method with SchillerNaumann drag model.

4.3. Test case C

The feature of test case C is that it is a new test case and the geometry is quite complex. The Q-E-E method should be able to predict accurate results no matter what the geometry it is. The time-averaged gas hold-up at different positions with different gas superficial velocities are reported in Fig. 11. It is seen that the profile of the time-averaged local gas fraction above the sparger plate ($z = 0.25$ m) shows an expected M-shape profile, which was also observed in the experiments. The peaks correspond to the gas fraction at the centers of the spargers. With increasing superficial velocity, there is an increase in the overall number of bubbles and thus an increased gas fraction. The turbulent dispersion effect was negligible, since the bubbles were just injected into the column and the flow was not fully developed. For the middle section ($z = 0.35$ m), the effect of the turbulent dispersion cannot be omitted and it can be seen that the phase fraction can be only predicted accurately when the turbulent dispersion was included, otherwise a over-estimated double peak was

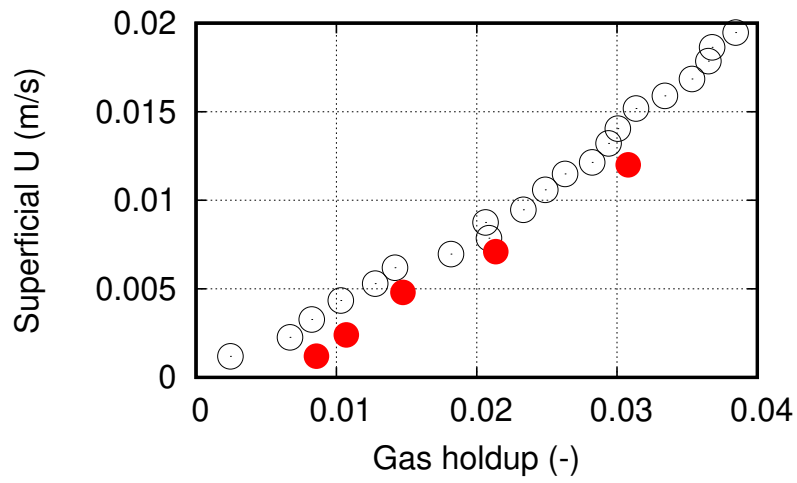


Figure 9: Comparison between experimental (black circles) and calculated global gas hold-up (red circles) at different superficial gas velocities for test case B slip velocity method.

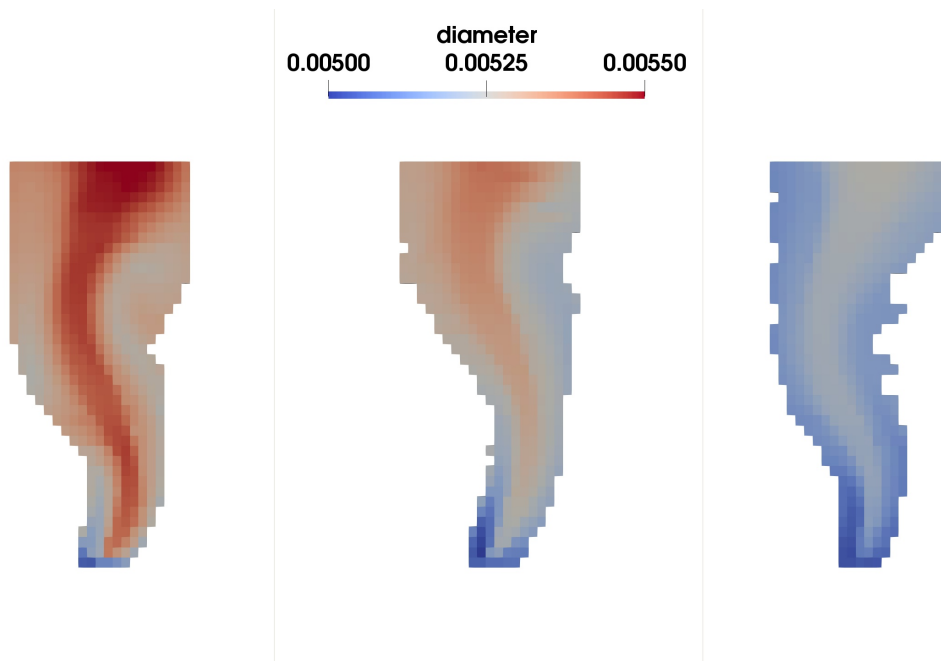


Figure 10: Prediction of the bubble size distribution of test case B with different gas superficial velocity (left: 0.012 m/s, middle: 0.0071 m/s, right: 0.0024 m/s).

predicted. The predicted phase fraction at the upper riser section was underestimated ($z = 0.58$ m). Improvements can be obtained by selecting other drag models or by fitting the bubble breakage and coalescence parameters. Meanwhile, the polydisperse predictions showed a significant difference with monodisperse results for high gas superficial velocity (0.0032 m/s). The difference of the results between monodisperse and polydisperse predictions for small gas superficial velocity is negligible. This can be explained by the fact that bubble breakage and coalescence is not frequent for these operating conditions. On the other hand, the difference of the results between monodisperse and polydisperse predictions for large gas superficial velocity is notable. This was also observed in test case B since the bubble coalescence and breakage are dominant for flows operated under larger superficial gas velocities.

5. Conclusions

In this work, a fast Eulerian-based approach which is suitable for polydisperse turbulent gas-liquid flow was proposed and implemented in open-source CFD code OpenFOAM-7 as `QEEFoam`. This method solves only one set of Navier-Stokes equations for the liquid phase. The gas phase was transported under the Eulerian framework by a convection equation. Turbulent effects can be included by adding a turbulent dispersion term in the gas phase equation.

The unknown gas phase velocity is calculated by:

1. assuming a constant slip velocity between the gas bubble and the surrounding liquid phase
2. solving an ODE for the gas bubble velocity which accounts for the interfacial forces acting on the bubble

A user-defined slip velocity is needed for the first method, while no empirical parameters are necessary for the second one. The bubble size is determined by including a transport equation for the total bubble number density. Bubble breakage and coalescence are modelled as source and sink terms. The whole equation system can be solved by any iterative solution procedure (e.g., PISO algorithm). Our algorithm and implementation are validated against three test cases characterized by different geometries and by different gas superficial velocities. The uncertainty associated with the constant slip velocity assumption and the effect of turbulent dispersion are assessed. The importance of the drag momentum exchange term is also assessed.

Predictions for the local/global gas phase fraction, period of oscillation the gas bubble plume and horizontal/vertical transient/time-averaged liquid velocity are compared with experimental data or simulation results predicted by the E-L method. Our results show that the methodology is capable of adequately describing the very complex dynamic flow fields in the bubble columns. Good agreements between the predictions with experimental data can be obtained. The slip velocity method can predict comparable results with that predicted by the ODE system if a suitable velocity is chosen, with slightly smaller computational costs. Compared with the existing CFD-PBE coupling algorithm (e.g., QMOM or CM), `QEEFoam`, which solves for the total bubble number density transport equation, is stable and fast. In conclusion, our algorithm seems a very promising and fast strategy for the simulation of industrial polydisperse turbulent gas-liquid flow.

Appendix

In order to compare the computational efficiency, the running times of `reactingTwoPhaseEulerFoam` and `QEEFoam` were reported in Fig. 12 ($\Delta t = 0.01$ s). All the simulations were launched by one CPU processor (Intel i7-5820K).

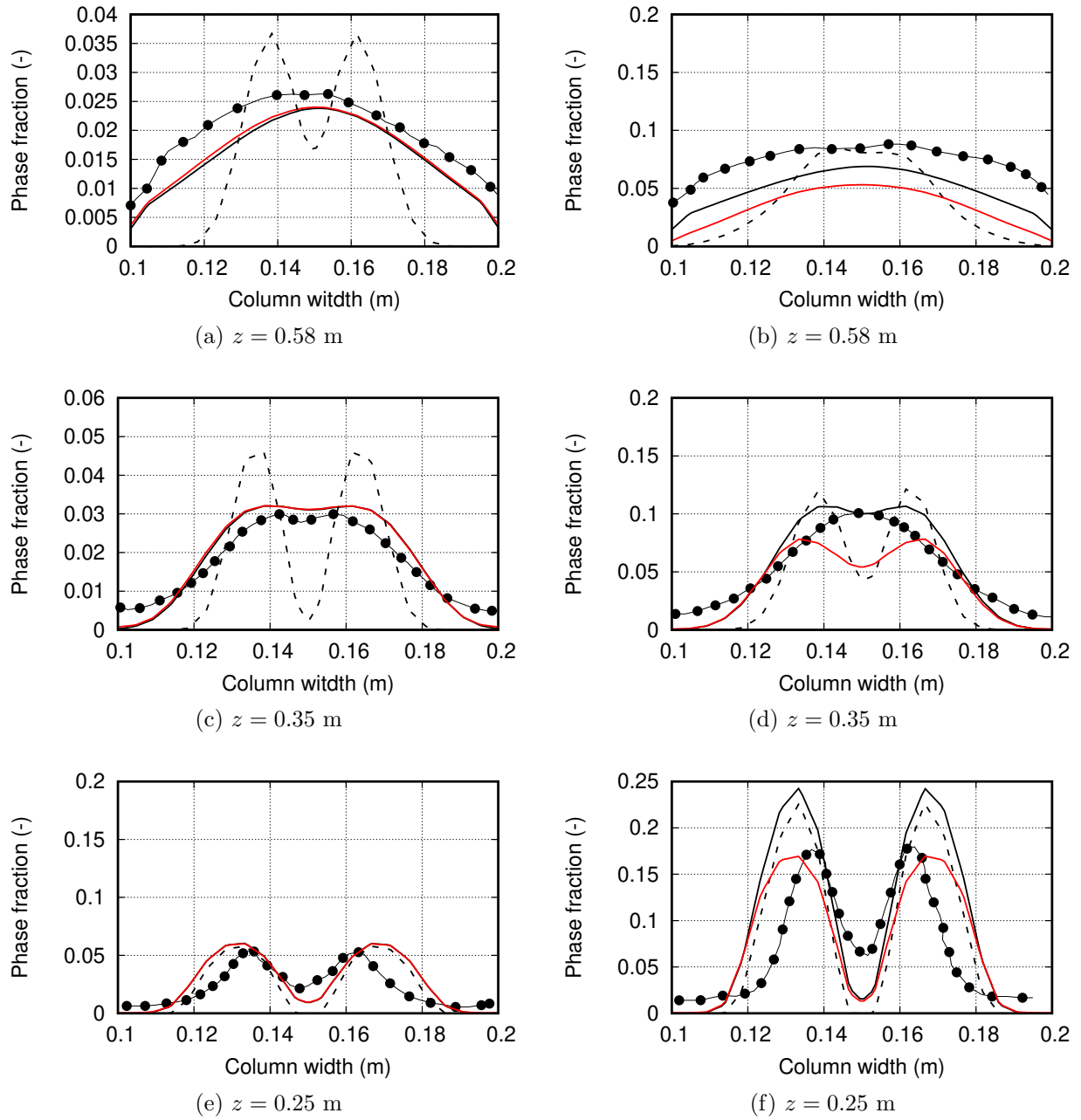


Figure 11: Time-averaged gas hold-up different positions by the ODE method for test case C. Gas superficial velocity is 0.008 m/s (left) and 0.032 m/s (right). Solid line: with turbulent dispersion term. Dashed line: without turbulent dispersion term. Line-points: experimental data. Black: monodisperse. Red: polydisperse.

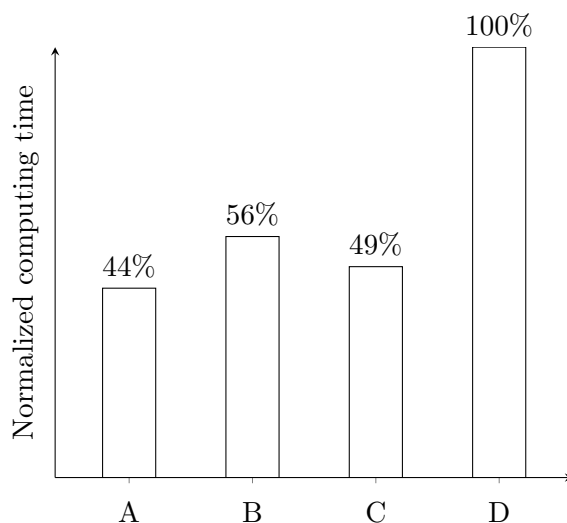


Figure 12: Comparison of the computing efficiency between `QEEFoam` and `reactingTwoPhaseEulerFoam` for test case B. A: monodisperse `QEEFoam` (slip vel. method, 327 s). B: monodisperse `QEEFoam` (ODE method, 413 s). C: polydisperse `QEEFoam` (slip vel. method, 362 s). D: monodisperse `reactingTwoPhaseEulerFoam` (732 s).

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