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A novel methodology for simulating vibrated fluidized beds using two-fluid models

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HIGHLIGHTS

- A novel strategy for the Eulerian simulation of vibrated fluidized beds is proposed.
- Vibration is incorporated through body forces in the transformed governing equations.
- Results compare well with DEM simulations and experiments for vertically vibrated beds.
- Phenomena like the bubble growth with the vibration strength are simulated.

GRAPHICAL ABSTRACT

Keywords:
Fluidization
Two-fluid model
Vibrated fluidized bed
Vibration strength
Bubble
Body force

ABSTRACT

The present work considers the use of the two fluid (Euler-Euler) CFD approach for the continuum description of vibrated fluidized beds as a less computationally demanding alternative to the discrete description given by Lagrangian Eulerian methods such as DEM. In particular, a novel simulation strategy consisting of solving the two fluid model equations in a coordinate reference system that moves with the vibrating walls of a gas-solid fluidized bed is proposed. By this way, vibration is transformed into simple alternating acceleration terms that are introduced through body forces in both the gas and the particle phase equations. The results of a series of two fluid model simulations compare well with discrete particle simulations as well as with experimental data reported for beds containing Geldart group B particles. In general, the results of a series of two fluid model simulations show similar trends to those seen in discrete particle simulations as well as in experimental data reported for beds containing Geldart group B particles. Exception of that is the velocity of bubbles, for which the two fluid simulations compare less satisfactorily with the available experimental data. The two fluid model simulations are also able to reproduce expected phenomena like the bubble growth with the vibration amplitude and the dependence of the pressure drop fluctuation on the vibration strength. In view of these promising results, the proposed two fluid model formulation opens the possibility of increasing the scale of the vibrated fluidized beds currently simulated.

1. Introduction

Fluidization is a process for powder handling widely used in industry owing to the favorable gas-solid and solid-solid contact efficiencies it provides. Nevertheless, gas fluidization alone has a limited ability to reduce particle agglomeration and gas channeling. The vibrated fluidized bed (VFB) is a gas-solid fluidization technology that introduces an external vibration force to the conventional fluidized bed (FB). Vibration prevents the solids from channeling and seriously agglomerating by supplying the bed with the energy required to break the bubble preferential paths and overcome the interparticle forces. At present, mechanical vibration has already been used in the process of relatively large (>1 mm) as well as fine micron scale particles [1-6] and very small nanoparticles in the range 1-100 nm [7,8], and it has been proved to be an effective means in the fluidization of cohesive particles [3,4,9], drying of granular material [10] and agglomeration control [8].
The dynamics of particle and gas phases in vibrated fluidized beds has been traditionally simulated employing the Lagrangian Eulerian approach (or discrete element method, DEM) that calculates the individual trajectory of each particle in the bed together with the solution of the continuum gas flow field. This allows the description of all the resulting particle-particle and particle-wall collisions [21,22]. Tatemoto et al. [23] and Xiang et al. [24] used the Lagrangian Eulerian method to simulate the motion of Geldart group B particles in a vibrated two dimensional bed that was fluidized with air. The thickness of the simulated bed was equal to the particle diameter and 1500 particles were considered. Changes in void fraction and air pressure fluctuations with time obtained from the simulation were analyzed and the distribution of particles in the bed was compared to experimental results [23]. Xiang et al. [24] also studied the distribution of velocity and concentration of particles at different amplitudes and frequencies of vibration, providing valuable qualitative information on the VFB behavior. Nevertheless, as stated in [24], quantitative evaluation of the data was difficult since the number of particles in the calculation was too small. The Lagrangian Eulerian approach has been also employed in the simulation of three dimensional (3 D) VFB. A 3 D simulation of a cylindrical (2.41 mm in diameter) vibrated fluidized bed filled with 14,135 cohesive fine particles was presented by Limtrakul et al. [25], where the effects of particle type, amplitude and frequency of vertical vibration, as well as the superficial gas velocity on the particle movement in the cylindrical bed were analyzed.

According to the literature reviewed, the use of discrete models for simulating VFB is restricted to beds of reduced dimensions, i.e. reduced number of particles, with the additional drawback of being highly computationally intensive [25]. Thus, a different computational strategy would be desirable to numerically simulate larger beds while achieving computing times similar to the ones required for the simulation of conventional FB. In this regard, simulation of small and medium scale conventional gas fluidized beds is commonly undertaken by means of two fluid computational fluid dynamic (CFD) models, also known as Eulerian Eulerian two fluid models, which are primarily based on the representation of the gas phase and the particulate phase as two interpenetrating continua [22,26]. Two fluid models provide information about the macroscopic hydrodynamics (i.e. velocity and volume fraction) of the two phases, including the bubble formation and motion in two and three dimensions [27,29].

The present work considers the use of the continuum description of VFB as a less computationally demanding alternative to the discrete description. In particular, a novel simulation strategy consisting on solving the two fluid model CFD equations in a co-ordinate reference system that moves with the vibrating walls of the bed is proposed here. By this way vibration is transformed into simple alternating acceleration terms that are introduced through body forces in both the gas and the solids phases. To assess the capabilities of two fluid models in describing realistically vibrating fluidized beds, the results of a series of two fluid model simulations are analyzed and compared with previously reported Lagrangian Eulerian simulations and experiments of sinusoidally
vibrated beds filled with Geldart group B particles and fluidized with air in bubble regime.

2. Theory

2.1. Two fluid model equations

The description of gas fluidized beds as two interpenetrating continua is the essence of two fluid models [22,26]. One continuum refers to the gas phase in the bed and the other to the particle phase (also known as particulate or solids phase). This allows the Eulerian description of both phases without resorting to the individual tracking of each particle in the bed. Two fluid models make use of the general equations of conservation of mass and momentum for the mass in the gas (g) and particle (p) phases. Table 1 lists these equations expressed in absolute coordinates and standard vector notation [22]. From the definition of volume fraction it is implicit that \( x_g + x_p = 1 \). In the momentum equation, the particle phase is treated as a fluid with effective transport properties [26]. Commonly, the conservation equations of mass and momentum are solved together with the differential equation for the granular temperature, which is based on the kinetic theory of granular flows and provides information about the level of random fluctuation of particle velocity due to collisions [26]. The granular temperature is required for the closure expressions of the drag coefficient \( K_{gp} \), the solid viscosities \( \mu_p \) and \( \lambda_p \), and the effective particle pressure \( p_p \). Also functions of \( \Theta \) are the diffusion coefficient of granular temperature \( k_{\Theta} \) and the collision dissipation energy \( \gamma_{\Theta} \). In the present work, the closure equation of Gidaspow et al. [30] has been chosen for calculating the drag coefficient \( K_{gp} \) owing to its robustness at the beginning of the simulation sequence. Previous studies indicate that differences between this and other well-known closure equations are not large in medium size two dimensional beds [28]. Table 2 summarizes these and other closure models selected for the present study.

2.2. Transformed two fluid model equations

As commented in Section 1, two fluid models have been traditionally employed for simulating non vibrating beds. However, as these models are based on universal laws of physics such as the mass conservation and the Newton’s second law, nothing precludes their use in situations in which the walls vibrate. This is true provided the vibration has a frequency sufficiently smaller than the characteristic collision frequency of particles in the bed, that is, there is separation of scales between collision frequencies and wall vibration frequencies. At this regard, characteristic collisions times smaller than \( 10^{-4} \) s (i.e. frequencies greater to \( 10^4 \) Hz) have been reported for conventional FB of Geldart B particles beyond the onset of bubbling [33]. In fact, bubble rise in conventional FB pushes upwards and downwards the particles within a region close to each bubble, the size of this region being of order of the bubble diameter [34]. The fluctuation induced by bubbles on the particle velocity has a characteristic frequency that can be roughly estimated dividing the bubble velocity by the size of the perturbed area \( f_b = 0.7 \sqrt{gD_b/D_h} = 2/\sqrt{D_b} \). Hence, for bubble diameters of order 10 cm, the fluctuation induced by bubbles on the particle velocity in conventional FB has a characteristic frequency of order 10 Hz, which is the same order of the vibration frequency considered in the present work. Therefore, for the vibrated bed regime studied here, it is expected the range of applicability of two fluid model equations to include vibration without the need of modifying or adding new closure models.

In order to simulate vibration using two fluid Eulerian models, a direct imposition of vibration by means of oscillatory displacement of the boundary condition at the walls of the bed has disadvantages related to the practical implementation of moving boundaries and the consistency of the mass and momentum balances in the computational cells at inlet flow boundaries (i.e. the bed distributor). In the present work, a novel strategy is presented to get rid of the disadvantages of the direct imposition of wall vibration. In particular, instead of using the traditional absolute system of reference for the spatial coordinates \( x = (x, y, z) \) in the governing equations, Eqs. (T1.1) (T1.5) in Table 1, a new relative system of reference \( x' = (x', y', z') \) that moves attached to the bed vessel (i.e. bed walls) can be employed. Note that it is assumed that the walls are non deformable so that they move as a rigid solid. In this moving system of reference \( x' \) there is no apparent movement of the walls and vibration information is incorporated in the governing equations through coordinate transformation factors. Thus, the computational mesh to be used for the solution of the governing equations is static in a moving system of reference, which greatly simplifies the numerical solution of the equations. Conceptually, the results should be equivalent to those obtained from the solution of the governing equations in the absolute coordinate system using a mesh that moves with the bed walls. Fig. 1 sketches the procedure of coordinate transformation to a relative system of reference which moves with the bed vessel.

Without loss of generality, the movement of the bed walls due to vibration in an absolute system of reference will be expressed here as a sinusoidal displacement in any of the three spatial directions, but other displacement functions can be used:

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Governing equations in absolute coordinates.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Conservation of mass</strong></td>
<td>( \frac{\partial}{\partial t} (x_g \rho_g v_g) + \nabla \cdot (x_g \rho_g v_g v_g) = 0 ) (T1.1)</td>
</tr>
<tr>
<td><strong>Conservation of momentum</strong></td>
<td>( \frac{\partial}{\partial t} (x_g \rho_g v_g) + \nabla \cdot (x_g \rho_g v_g v_g) = \nabla p_g + \nabla \cdot (x_g \tau_{gp}) + K_{gp}(v_g - v_p) + \tau_{p}\mathbf{g} ) (T1.2)</td>
</tr>
<tr>
<td><strong>Conservation of momentum</strong></td>
<td>( \frac{\partial}{\partial t} (x_p \rho_p v_g) + \nabla \cdot (x_p \rho_p v_p v_g) = \nabla p_p + \nabla \cdot (x_p \tau_{gp}) + K_{gp}(v_g - v_p) + \tau_{p}\mathbf{g} ) (T1.3)</td>
</tr>
<tr>
<td><strong>Balance of granular temperature</strong></td>
<td>( \frac{3}{2} \frac{\partial}{\partial t} (x_p \rho_p \Theta) + \nabla \cdot (x_p \rho_p \Theta \mathbf{v}<em>p) = (p_p \mathbf{J} + T_p) \cdot \nabla \mathbf{v}<em>p + \nabla \cdot (k</em>{\Theta} \nabla \Theta) + \gamma</em>{\Theta} \frac{3}{2} K_{gp} \Theta ) (T1.4)</td>
</tr>
</tbody>
</table>
\[ \begin{align*}
\delta(t) &= \{a_x \sin(2\pi f_x t), a_y \sin(2\pi f_y t), a_z \sin(2\pi f_z t)\} \\
&= \{a_x, a_y, a_z\} \sin(2\pi f \cdot t) \\
\end{align*} \]

where \( t \) is the time, the trio \( a_x, a_y, \) and \( a_z \) represents the vibration amplitude in the \( x, y \) and \( z \) direction, respectively, and \( f_x, f_y, \) and \( f_z \) are their corresponding vibration frequencies. Since the moving system of reference is attached to the bed walls the following transformation of coordinates is going to be used:

\[ \begin{align*}
\dot{x} &= x' \dot{t} \\
\dot{y} &= y' \dot{t} \\
\dot{z} &= z' \dot{t}
\end{align*} \]

Thus, the gas or particle phase velocities expressed in the moving system of reference are

\[ \begin{align*}
\mathbf{v'} &= \mathbf{v} + \nabla \delta/
\end{align*} \]

Applying the chain rule it is easy to demonstrate that

\[ \begin{align*}
\frac{\partial}{\partial t} &= \frac{\partial}{\partial t'} \frac{\partial}{\partial x} \\
\frac{\partial}{\partial x'} &= \frac{\partial}{\partial x} \\
\frac{\partial}{\partial \delta} &= \frac{\partial}{\partial \delta'} \nabla' \nabla
\end{align*} \]

\[ \text{Table 2} \]
Summary of closure models.

\[ \begin{align*}
A - \text{Coefficient of drag between gas and particles} \ [30] \\
K_{sp} &= 3 \frac{1}{4} \frac{\mu_p \rho_p}{\rho_g} \left( \frac{\rho_g}{\rho_p} \right)^{2/3} \text{ for } x_g > 0.8 \\
K_{sp} &= 150 \frac{\mu_p}{2 \rho_p^2} + 1.75 \frac{\rho_p}{\rho_g} \left( \frac{\rho_g}{\rho_p} \right)^{2/3} \text{ for } x_g \leq 0.8 \\
\text{where the drag coefficient is defined as:} \\
C_D &= \frac{24}{\sqrt{\rho_p \rho_g}} \left[ 1 + 0.15 \left( \frac{\rho_p}{\rho_g} \right)^{3/2} \right] \\
\text{With } \rho_p &= \rho_p \left( \frac{\mu_p}{\rho_p} \right)^{1/2} \\
B - \text{Particle phase pressure} \ [31] \\
\rho_p &= 2\rho_p \mu_p \theta^2 + 2\rho_p \left( 1 + \mu_p \right) \frac{\theta^2}{\rho_g} \frac{\partial \sigma_{pp}}{\partial \theta} \\
\text{where the radial distribution function is:} \\
\sigma_{pp}(r) &= \left( \frac{\rho_g}{\rho_p} \right)^{1/3} \left( \frac{\rho_p}{\rho_g} \right)^{1/2} \\
C - \text{Gas phase stress tensor} \\
\tau_x &= 2\mu_p \mu_v \left( \nabla v_x + \nabla v_x' \right) + 2\left( \frac{\mu_p}{\rho_p} \right) \nabla \cdot \mathbf{v} \\
\text{With } \mu_v &= 0 \\
D - \text{Particle phase stress tensor} \\
\tau_p &= 2\rho_p \mu_p \theta \left( \nabla v_x + \nabla v_x' \right) + 2\left( \frac{\mu_p}{\rho_p} \right) \nabla \cdot \mathbf{v} \\
\text{where particle phase bulk viscosity is} \ [31]: \\
\mu_p &= \frac{4}{\pi} \frac{\rho_p \mu_p \mu_v}{\rho_g} \left( \frac{\theta}{\rho_g} \right)^{1/2} \\
\text{and particle phase shear viscosity is:} \\
\mu_p &= \mu_{p, b} + \mu_{p, sh} \\
\text{which is composed of a kinetic viscosity} \ [30]: \\
\mu_{p, b} &= \frac{10 \rho_p \theta^2 \mu_v}{27 \theta_g \left( 1 + \rho_p \sigma_{pp} \right)} \left( 1 + \frac{4}{\sigma_{pp}} \frac{\rho_g}{\rho_p} \right)^{1/2} \\
\text{a collisional viscosity} \ [30]: \\
\mu_{p, col} &= \frac{4}{\pi} \frac{\rho_p \mu_p \mu_v}{\rho_g} \left( \frac{\theta}{\rho_g} \right)^{1/2} \\
\text{and a frictional viscosity} \ [32]: \\
\mu_{p, fr} &= \frac{\rho_p \sin \theta}{\sqrt{2} \rho_p^2} \\
\text{where } \theta \text{ is the angle of internal friction, and } \mu_{p, sh} \text{ is the second invariant of the deviatory stress tensor.} \\
E - \text{Diffusion coefficient of granular temperature} \ [30] \\
\kappa_g &= \frac{150 \rho_p \theta^2 \mu_v}{27 \theta_g \left( 1 + \rho_p \sigma_{pp} \right)} \left( 1 + \frac{6 \rho_p}{\theta^2 \sigma_{pp} \left( 1 + \rho_p \sigma_{pp} \right)} \right)^{1/2} \\
\text{F - Collisional dissipation of energy} \ [31] \\
\gamma_{a, b} &= \frac{12}{\rho_p} \frac{\sigma_{pp}}{\sqrt{\pi}} \rho_p \frac{\theta^2}{\rho_g} \left( \frac{\rho_g}{\rho_p} \right)^{3/2} \\
\end{align*} \]
where $\nabla' = \{\partial / \partial x', \partial / \partial y', \partial / \partial z'\}$ is the gradient operator in the moving system of reference. Note also that $\partial\delta/\partial t = \partial\delta'/\partial t$. Introducing the transformed time derivative and gradient in Eq. (T1.1),

$$\frac{\partial}{\partial t} (\zeta_0 \rho_k) \frac{\partial \delta}{\partial t} \nabla' (\zeta_0 \rho_k p) + \nabla' \cdot (\zeta_0 \rho_k v \rho_k) = 0$$

and using Eq. (3) together with the fact that $\partial\delta/\partial t = \nabla' (\zeta_0 \rho_k) = \nabla' (\zeta_0 \rho_k \delta / \partial t)$ since $\delta$ is only a function of $t$, the equation of mass conservation for the gas phase after transforming coordinates is shown in Eq. (T3.1) in Table 3, which is analogous to the equation in the absolute system of coordinates. This result should be expected since the conservation of mass is a property independent of the system of reference. A similar result is obtained for the conservation of mass in transformed coordinates for the particle phase, Eq. (T3.2) in Table 3.

The same transformation of coordinates can be done for the momentum equations:

$$\frac{\partial}{\partial t} (\zeta_0 \rho_k \zeta_0 v) \frac{\partial \delta}{\partial t} \nabla' (\zeta_0 \rho_k \zeta_0 v) + \nabla' \cdot (\zeta_0 \rho_k v \rho_k v) = \zeta_0 \nabla' p + \nabla' \cdot (\zeta_0 \rho_k v \rho_k v) + K_{gp} (v \rho_k v)$$(T3.2)

$$\frac{\partial}{\partial t} (\zeta_0 \rho_k \zeta_0 v) \frac{\partial \delta}{\partial t} \nabla' (\zeta_0 \rho_k \zeta_0 v) + \nabla' \cdot (\zeta_0 \rho_k v \rho_k v) = \zeta_0 \nabla' p + \nabla' \cdot (\zeta_0 \rho_k v \rho_k v) + K_{gp} (v \rho_k v)$$(T3.3)

Again, incorporating Eq. (3) in Eqs. (6) and (7) and realizing that $\nabla' (\partial\delta'/\partial t)$ = 0 and

$$\nabla' \cdot \dot{\zeta_0} v' + \alpha \frac{\partial \zeta_0 v'}{\partial t} + \frac{\partial \zeta_0 v}{\partial t}$$

The following transformed momentum equations are obtained:

$$\frac{\partial}{\partial t} (\zeta_0 \rho_k \zeta_0 v) \frac{\partial \delta}{\partial t} \nabla' (\zeta_0 \rho_k \zeta_0 v) + \nabla' \cdot (\zeta_0 \rho_k v \rho_k v) = \zeta_0 \nabla' p + \nabla' \cdot (\zeta_0 \rho_k v \rho_k v) + K_{gp} (v \rho_k v)$$(T3.4)

$$\frac{\partial}{\partial t} (\zeta_0 \rho_k \zeta_0 v) \frac{\partial \delta}{\partial t} \nabla' (\zeta_0 \rho_k \zeta_0 v) + \nabla' \cdot (\zeta_0 \rho_k v \rho_k v) = \zeta_0 \nabla' p + \nabla' \cdot (\zeta_0 \rho_k v \rho_k v) + K_{gp} (v \rho_k v)$$(T3.5)

where $\zeta_0$ = $\mu_0 (\nabla' v + \nabla' v') + (1 - 2 \mu_0) (\nabla' v')$ is the tensor of shear stresses for the velocities of the gas ($k=q$) and particle ($k=p$) phases in the moving coordinates.

The transformed momentum conservation expressions, Eqs. (9) and (10), can be further simplified noticing that $\frac{\partial}{\partial t} \nabla' (\zeta_0 \rho_k \zeta_0 v)$ = $\nabla' \left( \frac{\partial}{\partial t} (\zeta_0 \rho_k \zeta_0 v) \right) + \frac{\partial}{\partial x} \left( \frac{\partial}{\partial t} (\zeta_0 \rho_k \zeta_0 v) \right) = \nabla' (\partial (\zeta_0 \rho_k \zeta_0 v) / \partial t) + \nabla' (\partial (\zeta_0 \rho_k \zeta_0 v) / \partial \rho_k)$.

Besides, $\frac{\partial}{\partial t} \nabla' (\zeta_0 \rho_k \zeta_0 v)$ = 0 owing to the conservation of mass, Eqs. (T3.1) and (T3.2). This leads to the final form of the transformed equations for the momentum conservation included in Table 3, Eqs. (T3.3) and (T3.4).

Regarding the granular temperature $\theta$, it accounts for the quadratic difference between the individual velocity of particles and the bulk velocity $v_e$ of the particle phase [26]. By definition this velocity difference is stochastic, which implies that changing from a static to a moving system of reference modifies the bulk velocity as well as the individual velocity of particles, but does not alter the difference between individual and bulk velocities. This can be seen in Table 3.

### Table 3

<table>
<thead>
<tr>
<th>Equation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>(T3.1)</td>
<td>Conservation of mass</td>
</tr>
<tr>
<td>(T3.2)</td>
<td>Conservation of momentum</td>
</tr>
<tr>
<td>(T3.3)</td>
<td>Conservation of momentum</td>
</tr>
<tr>
<td>(T3.4)</td>
<td>Balance of granular temperature</td>
</tr>
</tbody>
</table>

*Time $t$, coordinates $x'$ and velocity $v'$ in a moving system of reference are given by Eqs. (2)-(4) and (9) respectively.*
noticing that both the individual and bulk velocities of particles in a moving system of reference are modified by the same apparent velocity, $\frac{\partial \mathbf{v}}{\partial t}$. Thus, this apparent velocity vanishes when calculating the difference between the individual and bulk particle velocities. Therefore, the granular temperatures observed from the moving and the static systems of reference are equal, i.e. $\Theta' = \Theta$, which means that, for a given bubbling bed, the observed granular temperature is independent of the system of reference. As for the case of the mass conservation equations, transformation of coordinates and velocities using Eqs. 3, 4a, 4b, 5 leave unchanged the structure of the governing equation for the granular temperature, Eq. (T3.5) in Table 3.

All the deductions above have been done for two phases (i.e. gas and particles), but the same procedure can be equally carried out in beds containing a mixture of gas, liquid and solid phases or multiple particle phases representing differentiated particle size distributions.

2.3. Acceleration and equivalent mass forces

In the previous Section 2.2 the governing equations for the gas and particulate phases expressed in a coordinate system that moves with the walls of the vibrated bed have been deduced. As noticed previously, the resulting equations for the mass conservation and granular temperature balance, Eqs. (T3.1), (T3.2) and (T3.5), are similar to the equations in a static system of reference, Eqs. (T1.1), (T1.2) and (T1.5). In contrast, the equations of momentum conservation in a moving system of reference, Eqs. (T3.3) and (T3.4), differ from the original ones, Eqs. (T1.3) and (T1.4), in that they posses an extra term in their right hand side related to the acceleration of the system of reference, $\phi \frac{\partial \mathbf{v}}{\partial t}$. In fact this acceleration term can be viewed as an apparent volumetric force, acting over both the gas and the particulate phases, that reflects the counter reaction to the acceleration of the system of reference. Therefore,

$$ f_{m,eq} = g \frac{\partial \delta}{\partial t} $$

is the equivalent vector of mass force that should be implemented in the model, instead of the gravity vector $g$, to simulate a vibrating bed with two fluid models. For a sinusoidal vibration, Eq. (1), the equivalent vector of mass force is

$$ f_{m,eq} = \{g_x + 4\pi^2 f_x^2 a_x \sin(2\pi f_x t), g_y + 4\pi^2 f_y^2 a_y \sin(2\pi f_y t), g_z + 4\pi^2 f_z^2 a_z \sin(2\pi f_z t)\} $$

Obviously, after solving the system of transformed equations, Eqs. (T3.1) (T3.5) in Table 3, the absolute position $x$ of the phases, the velocity $\mathbf{v}$, and the shear stresses $\tau$, can be recuperated by undoing the transformation given by Eqs. (2) and (3).

2.4. Effect of the transformation on the boundary conditions

It is to be noticed that in the transformed coordinates, given by the moving system of reference, all the vibration effects in the model are concentrated in Eqs. (T3.3) and (T3.4), whereas the apparent movement of the walls and other boundaries is null. That means that the boundary conditions to be employed by the transformed equations in Table 3, are similar to the standard conditions used in a non vibrating bed. Implicit in this reasoning is the assumption that the plenum and the distributor vibrate attached to the rest of the bed vessel. Thus, the gas superficial velocity in the moving system of reference is steady and only depends on the gas flow rate entering the plenum as in a non vibrating bed.
lected as the base case of the slim bed, with vibration frequency equal to 15 Hz and amplitude 1.5 mm. Similarly, case HA 2 (with vibration frequency 15 Hz and amplitude 15 mm) was utilized as the base case for the simulation of the high amplitude vibrated bed. For these base cases, as well as for the non vibrated cases, there are three different meshes used to analyze the simulation sensitivity on the mesh number of nodes.

### 3.2.2. Boundary condition parameters

A time independent and spatially uniform superficial velocity condition, $U_g$ in Table 4, was imposed at the bottom of both bed configurations for the gas phase, see Figs. 2b and 3b. Null velocity and concentration were set for the particle phase in order to avoid it to cross the distributor. Pressure outlet conditions (denoted as PO in Figs. 2b and 3b) were selected for the upper boundary of

![Fig. 2. Configuration-1 consisting in a vibrated slim bed: (a) domain dimensions and computational mesh used for the two-fluid simulation; (b) boundary conditions in relative coordinates and vertical positions for data sampling.](image)

![Fig. 3. Configuration-2 consisting in a fluidized bed with high-amplitude vibration: (a) domain dimensions and computational mesh used for the two-fluid simulation; (b) boundary conditions in relative coordinates and data sampling regions.](image)

### 3.2.3. Solution procedure

The commercial code ANSYS FLUENT v12.0 was used to solve the transformed two fluid model differential equations of continuity and momentum, Eqs. (T3.1)–(T3.4) for the gas and particle phases together with the granular temperature balance equation, Eq. (T3.5). This code resorts to a finite volume formulation with an implicit time advance algorithm to solve these transient governing equations, the coupling between the gas and particle phases being done by a modified SIMPLE algorithm (Phase coupled SIMPLE), which has given satisfactory results in the prediction of the pressure and bubble behavior of conventional fluidized beds [17]. Since the transformed governing equations to be solved are expressed in a moving system of coordinates, vibration of the bed vessel was incorporated in the simulation through the equivalent vector of mass force, Eq. (12), which was programmed in a separated C++ code and interpreted by ANSYS FLUENT as a user defined function. A second order accuracy scheme was selected for the discretization of the advective transport terms as well as for the time advance algorithm.

The restitution coefficient for the collisions between particles was $e_{pp} = 0.9$ for all the simulations. For the thin bed (configuration

### Table 4

Summary of simulation parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Configuration-1: vibrating slim bed</th>
<th>Configuration-2: high-amplitude vibrating bed</th>
</tr>
</thead>
<tbody>
<tr>
<td>$W$ (mm)</td>
<td>25</td>
<td>300</td>
</tr>
<tr>
<td>$H$ (mm)</td>
<td>250</td>
<td>600</td>
</tr>
<tr>
<td>$h_0$ (mm)</td>
<td>55</td>
<td>205</td>
</tr>
<tr>
<td>$d_p$ (μm)</td>
<td>1000</td>
<td>198</td>
</tr>
<tr>
<td>$\rho_p$ (kg/m$^3$)</td>
<td>1550</td>
<td>2520</td>
</tr>
<tr>
<td>$\mu_p$ (kg/m$^3$)</td>
<td>1.167</td>
<td>1.167</td>
</tr>
<tr>
<td>$\mu_g$ (Pa s)</td>
<td>1.22e−5</td>
<td>1.22e−5</td>
</tr>
<tr>
<td>$U_g$ (m/s)</td>
<td>0.5</td>
<td>0.0907</td>
</tr>
<tr>
<td>$f$ (Hz)</td>
<td>3–10</td>
<td>15–30</td>
</tr>
<tr>
<td>$A$ (mm)</td>
<td>1–2</td>
<td>5–15</td>
</tr>
<tr>
<td>$e_{pp}$ (–)</td>
<td>0.9</td>
<td>0.9</td>
</tr>
<tr>
<td>$e_{pw}$ (–)</td>
<td>0.9</td>
<td>0.2</td>
</tr>
<tr>
<td>$z_{pr}$ (–)</td>
<td>0.6</td>
<td>0.61</td>
</tr>
<tr>
<td>$x_{max}$ (–)</td>
<td>0.605</td>
<td>0.63</td>
</tr>
<tr>
<td>$\phi$ (–)</td>
<td>0.6</td>
<td>0.6</td>
</tr>
<tr>
<td>$\Delta t$ (s)</td>
<td>2.5e−4</td>
<td>1e–4</td>
</tr>
<tr>
<td>$N_i$ (iterations)</td>
<td>40</td>
<td>50</td>
</tr>
</tbody>
</table>
tion 1) the maximum packet limit was assumed to be $\alpha_{pm\text{max}} = 0.605$, which is calculated from a hexagonal arrangement of spheres occupying a bed of thickness equal to their diameter. For the high amplitude vibrating bed (configuration 2), since the bed thickness is several times the particle diameter, the standard value $\alpha_{pm\text{max}} = 0.63$ was chosen to represent the maximum packing limit. In all the simulations the value chosen for the friction packing limit $\alpha_{fr}$ was slightly smaller than $\alpha_{pm\text{max}}$. It was found that differences between these two packing limits did not affect markedly the simulation results, though convergence of the solution seemed to proceed smoothly if $\alpha_{fr}$ was inferior to $\alpha_{pm\text{max}}$.

The simulations commenced with air entering at the distributor with the bed at rest. The time step of the simulations was fixed to values of order $10^{-5}$ s, that is, two orders of magnitude smaller than the characteristic vibration time period. The number of iterations per time step, $N_e$, was 40 and 50 (see Table 4). This ensured that the residuals of the numerically solved equations remained below $10^{-3}$. To obtain statistically steady state results, a total simulated time of more than 25 s for the slim bed configuration and 12 s for the high amplitude bed configuration were computed. The first 4 s of simulations were not used in the results in order to eliminate the transient effects during the fluidization start up.

4. Results

4.1. Mesh sensitivity analysis

Three different meshes were studied for each vibrating bed configuration in order to assess the sensitivity of the simulation on the spatial resolution of the computational mesh. The mesh sensitivity analysis was focused on the non vibrating case as well as the base cases for the vibrating beds. For the slim bed (configuration 1), cases SB 1A and SB 2A in Table 5 were solved on a mesh having cells of size $\Delta s = 1.78$ mm in both $x$ (horizontal) and $z$ (vertical) directions in the region occupied by the bed particles. In cases SB 1B and SB 2B the mesh is refined by halving the cells size in $x$ and $z$ directions so that $\Delta s = 0.892$ mm. Finally, after refining again the mesh in the two spatial directions, the cell size was reduced to $\Delta s = 0.446$ mm for cases SB 1C and SB 2C. Analogously, for the high amplitude vibrated fluidized bed, the cases HA 1A and HA 2A in Table 6 have a computational mesh composed of square cells whose size is $\Delta s = 10$ mm in the region containing particles. Refining in horizontal and vertical directions, the cell size is $\Delta s = 5$ mm for the cases HA 1B and HA 2B, and $\Delta s = 2.5$ mm for the cases HA 1C and HA 2C.

Fig. 4 depicts the mean and standard deviation of the gas pressure at the distributor. To enhance visualization, both variables have been normalized dividing by the value obtained from the mesh of smallest cell size (i.e. meshes for SB 1C and SB 2C in Table 5 and for HA 1C and HA 2C in Table 6). The gas pressure was calculated performing the arithmetic mean of the local values of the gas static pressure at the distributor. Pressure is indicative of the bed dynamics and is affected by the interaction between the emulsion phase and the bubbles located anywhere in the bed. On view of Fig. 4a, it is clear that passing from the meshes SB 1B and SB 2B to the meshes SB 1C and SB 2C produces little difference in the results at the expense of increasing the computational cost, which was roughly proportional to the number of nodes in the mesh. Therefore, the computational mesh for the cases SB 1B and SB 2B was selected to carry out the remaining simulations for the slim bed (i.e. cases SB 3 to SB 7 in 5). For similar reasons, see Fig. 4b, the mesh used in cases HA 1B and HA 2B was the one chosen to simulate the rest of cases in Table 6 for the high amplitude vibrated bed.

4.2. Comparison with DEM results

In this section the two fluid model methodology described in Section 3 is employed to simulate a slim bed of small scale, i.e. configuration 1 in Table 4, previously characterized in the literature by Lagrangian Eulerian (DEM) simulations [23,24].

4.2.1. Time evolution of particle volume fraction and pressure drop

The evolution of the simulated particle volume fraction and gas pressure drop along time is first presented to elucidate whether the general dynamics of small vibrating beds can be satisfactorily predicted with two fluid models. The results presented here have been constructed by transversally averaging the simulated volume fraction and pressure along a horizontal line, covering the whole bed width, at a given distance from the distributor, $z$. Apart from the distributor, $z = 0$, three different distances have been selected to perform the horizontal averaging, which are $z_1$, $z_2$ and $z_3$ in Fig. 2b. From here on this averaging operation will be denoted as $x$ average.

Fig. 5a shows at different positions from the distributor the time variation of the $x$ average particle volume fraction taken from the simulation without vibration (case SB 1B in Table 5). The same simulated variables are depicted in Fig. 5b for the fluidized bed with vertical vibration at frequency $f_z = 15$ Hz with amplitude $A = 1.5$ mm (case SB 2B). In the case without vibration, Fig. 5a, near the bottom of the bed ($z = 1.56$ mm), there is no appreciable change of particle volume fraction with time. The changes in volume fraction values, at higher positions in the bed, are due to the presence of voids (i.e. bubbles). In the case of the vibrated bed, Fig. 5b, a similar result to the one obtained with a DEM simulation strategy by Tatemoto et al. [23] is found: void fraction oscillations (1 $\alpha_{fr}$) appear close to the distributor ($z = 1.563$ mm) at the same frequency of the imposed vibration, i.e. 15 Hz. Moreover, at this lower position, the $x$ averaged particle volume fraction oscillates along time around a mean volume fraction value that is lower than

<table>
<thead>
<tr>
<th>Cases</th>
<th>Mesh nodes (nodes in horizontal direction)</th>
<th>Vibration frequency $f$ (Hz)</th>
<th>Vibration amplitude $A$ (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SB-1A</td>
<td>1279 (14)</td>
<td>No vibration</td>
<td></td>
</tr>
<tr>
<td>SB-1B</td>
<td>3701 (28)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SB-1C</td>
<td>12,314 (56)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SB-2A</td>
<td>1279 (14)</td>
<td>15</td>
<td>1.5</td>
</tr>
<tr>
<td>SB-2B</td>
<td>3701 (28)</td>
<td>15</td>
<td>1.5</td>
</tr>
<tr>
<td>SB-2C</td>
<td>12,314 (56)</td>
<td>15</td>
<td>1.5</td>
</tr>
<tr>
<td>SB-3</td>
<td>3701 (28)</td>
<td>5</td>
<td>1.5</td>
</tr>
<tr>
<td>SB-4</td>
<td>3701 (28)</td>
<td>20</td>
<td>1.5</td>
</tr>
<tr>
<td>SB-5</td>
<td>3701 (28)</td>
<td>30</td>
<td>1.5</td>
</tr>
<tr>
<td>SB-6</td>
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<td>15</td>
<td>1</td>
</tr>
<tr>
<td>SB-7</td>
<td>3701 (28)</td>
<td>15</td>
<td>2</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Cases</th>
<th>Mesh nodes (nodes in horizontal direction)</th>
<th>Vibration frequency $f$ (Hz)</th>
<th>Vibration amplitude $A$ (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HA-1A</td>
<td>2511 (30)</td>
<td>No vibration</td>
<td></td>
</tr>
<tr>
<td>HA-1B</td>
<td>9211 (60)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HA-1C</td>
<td>36,421 (120)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HA-2A</td>
<td>2511 (30)</td>
<td>15</td>
<td>15</td>
</tr>
<tr>
<td>HA-2B</td>
<td>9211 (60)</td>
<td>15</td>
<td>15</td>
</tr>
<tr>
<td>HA-2C</td>
<td>36,421 (120)</td>
<td>15</td>
<td>15</td>
</tr>
<tr>
<td>HA-3</td>
<td>9211 (60)</td>
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<td>HA-4</td>
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<tr>
<td>HA-5</td>
<td>9211 (60)</td>
<td>20</td>
<td>5</td>
</tr>
<tr>
<td>HA-6</td>
<td>9211 (60)</td>
<td>30</td>
<td>5</td>
</tr>
</tbody>
</table>
for the case without vibration. This behavior was also reported in Tatemoto et al. [23] and Xiang et al. [24], who showed that a void gap appears close to the distributor at high frequencies that lowers the mean volume fraction. At higher positions in the bed, \( z = 26.6 \) mm and 51.6 mm in Fig. 5b, the time evolution of the \( x \)-averaged volume fraction does not closely follow the imposed vibration and have a pattern that resembles more the solid volume fraction of the bed without vibration, \( z = 26.6 \) mm and 51.6 mm in Fig. 5a. Thus, the presence of bubbles is probably the dominant phenomenon affecting the particle volume fraction variation at these higher positions in the bed.

Simulated pressure signals are presented in Fig. 6 for the two fluid model cases without vibration (SB 1B) with vibration at 15 Hz and displacement 1.5 mm (SB 2B). The pressure signal was obtained by \( x \)-averaging the air relative pressure close to the distributor in the simulated bed. Here, the air relative pressure is the air pressure minus the air pressure at the top of the freeboard. Thus, the pressure signal can be interpreted as the pressure drop that suffers the air when passing through the whole bed. As for the particle volume fraction signals, a higher number of pressure oscillations are found in the case with vibration compared to the non vibrated case. This two fluid simulation outcome is in harmony with the results obtained with DEM simulations by Xiang et al. [24]. The vertical vibration of the bed creates an oscillatory bulk motion of the bed, which directly affects the pressure at the

![Fig. 4](image)

**Fig. 4.** Normalized mean and standard deviation of the gas pressure drop as a function of the number of nodes in the computational mesh used for the two-fluid simulations: (a) slim bed; (b) high-amplitude vibrating bed.

![Fig. 5](image)

**Fig. 5.** Time evolution of the \( x \)-averaged particle volume fraction at different heights over the distributor: two-fluid simulation results for the fluidized slim bed without vibration (a), and with vibration at \( f = 15 \) Hz and \( A = 1.5 \) mm (b).

![Fig. 6](image)

**Fig. 6.** Instantaneous and mean gas pressure drop in the fluidized slim bed. The results were obtained from the two-fluid simulations without vibration and with vibration at \( f = 15 \) Hz and \( A = 1.5 \) mm.
According to Fig. 6, the main frequency of the pressure oscillation in the simulations is similar to the vibration frequency imposed to the bed (15 Hz). On the contrary, as shown in Fig. 6, the frequency of the simulated pressure drop for the bed without vibration coincides with the natural frequency of the bed estimated with $f_n = g/h_0^{1/2}/\rho = 4.3$ Hz [36]. This natural frequency is still present in the vibrated fluidized bed since the amplitude of the 15 Hz pressure drop oscillations in Fig. 6 are modulated with a frequency equal to $f_n$.

### 4.2.2. Effect of vibration frequency and amplitude

The values obtained for the simulated mean and the standard deviation of the pressure drop at different vertical vibration frequencies, are represented in Fig. 7a. For that purpose, cases SB1B, SB2B, SB3, SB4 and SB5 in Table 5 have been used in the present analysis. Increasing the vibration frequency entails higher values of the standard deviation of pressure fluctuations, while their mean is almost constant, that is, independent of vibration. This trend is in good agreement with the results obtained in the DEM simulation by Xiang et al. [24].

Fig. 7. Mean value and standard deviation of the air pressure drop in the fluidized slim bed versus vibration frequency for $A = 1.5$ mm (a) and vibration amplitude for $f = 15$ Hz (b).

In Fig. 8, the normalized pressure fluctuations and distributor displacement over time are shown for two different vibration frequencies. Normalization was done by subtraction of the mean value and division by the standard deviation of each signal. As commented previously, the bed vibration frequency is recovered in the simulated pressure fluctuation curve, but with a certain delay time. Following Xiang et al. [24], the delay time is defined as the difference between the time at the maximum value of pressure and the time of maximum displacement in the distributor (see Fig. 8).

Fig. 8. Time evolution of the normalized distributor displacement and gas pressure drop taken from the two-fluid simulations of the fluidized slim bed with vibration at $f = 5$ Hz (a) and $f = 15$ Hz (b). In both cases $A = 1.5$ mm.

The delay time is characterized in Fig. 9. According to the two fluid simulation results, the delay time is highly dependent on the vibration frequency, as shown in Fig. 9a. A change in the vibration frequency modifies the vibration time period ($1/f$) which alters the available time for the particles to reproduce the bed vessel vibration. In contrast, the vibration time period remains constant in Fig. 9b and, as the two fluid simulations reveal, the effect of the vibration amplitude on the delay time is not as sharp. This tendency is in fine agreement with the DEM results reported in Xiang et al. [24]. Though quantitative differences appear between the
two fluid simulation and DEM results in Fig. 9b, the obtained values for the delay time are of the same order of magnitude, which is a remarkable finding considering that the fluid like behavior of the particle phase (an assumption implicit in two fluid models) may not be fully reached if the number of particles is small as in the case of this slim bed configuration.

4.3. Comparison with experiments

In the previous section it has been seen that the two fluid model methodology presented here can provide comparable results to the Lagrangian Eulerian simulation of a fluidized bed of small scale under reduced vibration amplitudes. In order to extend this assessment of capabilities of two fluid models in simulating vibrated beds, the next results are focused on a larger bed with high amplitude vibration (configuration 2 in Table 6). The results from the two fluid model simulation are compared with the experimental findings shown in Zhou et al. [19] and Mawatari et al. [20].

4.3.1. Effect of vibration on the bubble behavior

Fig. 10 shows four snapshots of the instantaneous solids volume fraction distribution obtained with two fluid model simulations at different vibration intensities characterized by the vibration strength parameter $A_z$.

$$A_z = \frac{4\pi^2Af^2}{|g|}$$

The values of $A_z$ in Fig. 10 correspond to a fixed vibration frequency of 15 Hz while the vibration amplitude takes the values of 0, 5, 10, and 15 mm (cases HA 1B, HA 2B, HA 3 and HA 4 in Table 6). The presence of bubbles in all cases shown in Fig. 10 demonstrates the bubbling condition of the bed without and with vibration. Apparently, an increase of the vibration amplitude promotes the presence of larger bubbles in the bed. This effect is in good agreement with the experimental photographs obtained in [19,20]. The rationale of this effect is that an increase of the vibration strength creates higher acceleration forces suffered by the bed during the sinusoidal displacement of the vessel. In other words, the vertical vibration strength is more intense with larger vibration amplitudes $A$. As observed by Mawatari et al. [20], the increasing of the vibration strength reduces the minimum fluidization velocity and, as a consequence, increases excess gas velocities, so the expected visible flow in form of bubbles grows. Besides, it has been reported that increasing of the bed agitation amplitude, produced when increasing $A$, can favor the coalescence of bubbles [19], which is a way of increasing the size of the bubbles. The effect of vibration frequency on the local mean diameter of bubbles is illustrated in Fig. 11a for a vibration amplitude equal to 5 mm (cases HA 1B and HA 4 to 6 in Table 6). In the figure, simu
Bubbles are detected in the simulation from snapshots of solids volume fraction exported every $5 \times 10^{-3}$ s of computed time. A threshold value of particle volume fraction equal to $\varphi_{p,th} = 0.3$ was used to detect the bubble contour. Information concerning the selection of $\varphi_{p,th}$ and the calculation of the bubble diameter and velocity from the bubble contours can be found elsewhere [28]. As in Zhou et al. [19], bubbles are captured when their centroids pass through any of the two sampling regions marked in Fig. 3b. In each sampling region, the mean diameter is calculated performing the arithmetic mean of the diameters of all the captured bubbles. These two sampling regions are relatively thin (2 cm) and span all the bed width; one region ranges from $z = 9$ cm to 11 cm, and the other region from $z = 15$ cm to 17 cm. As Fig. 11a indicates, the mean bubble diameter obtained with the two fluid simulations and the experiments are reasonably similar. The simulations show that the bubble size increases with the vibration frequency, though this increment is not evident in the case of the experimental data. Regarding the vibration amplitude, Fig. 11b shows that the mean bubble diameter obtained from the two fluid simulations (cases HA 1B, HA 2B, HA 3 and HA 4 in Table 6) increases with the vibration amplitude. As Fig. 11b demonstrates, this result is in harmony with the experimental data reported in [19]. Also in good agreement with the experimental evidence is the fact that the mean bubble diameter seems to be more sensitive to the vibration amplitude than to the vibration frequency (i.e. doubling the vibration amplitude alters the mean bubble diameter more than doubling the vibration frequency). Besides, both the simulated and experimentally acquired bubbles have slightly larger diameters when increasing the distance of the sampling region to the distributor, $z$. The mean vertical velocity of bubbles obtained from the two fluid model simulations as a function of the vibration frequency and amplitude is presented in Fig. 12. The results have been taken from the same simulation cases utilized in Fig. 11. In addition, experimental bubble velocity data from [19] have been included for comparison purposes. In Fig. 12 the mean velocity depicted is relative to the bed distributor movement. It was found that expressing the results in an absolute system of reference did not change significantly the mean velocity because the instantaneous differences between the relative and absolute velocity are compensated after calculating the mean over the time.

Both the two fluid simulation and the experiments yield mean bubble velocities of the same order or magnitude. In some cases (see vibration with $A = 5$ mm and $f = 15$ Hz) the value of the mean bubble velocity provided by the two fluid simulation is virtually
equal to the experimental data. According to Fig. 12a, changes in the vibration frequency do not produce a clear tendency on the change of the mean vertical velocity estimated with the two fluid simulation results at \( z = 9.11 \) cm, which is in agreement with the experimental results. Unexpectedly, the major discrepancies between the simulation and the experimental data appear for the bed without vibration (\( f = 0 \) Hz). The effect of the vibration amplitude on the mean vertical velocity of bubbles is illustrated in Fig. 12b. Again, the bubble velocities obtained from the simulation and from the experiments by Zhou et al. [19] have the same order of magnitude. Bubbles in the experiments have a mean velocity that slightly increases and then decreases with the vibration amplitude, though this dependency is weak. In contrast, the tendency revealed by the two fluid simulation is that the bubble velocity monotonically decreases with the vibration amplitude. However, it is worth mentioning that the estimation of the bubble velocity in simulations and experiments is an operation sensitive to the way the displacement of the bubble centroid is calculated. Changes in shape of an irregular bubble may produce an apparent displacement that is difficult to separate from what would be expected to be the ascending velocity of the bulk volume of the bubble. In fact, bubbles in VFB can be larger and with more irregular boundaries than in conventional FB. In experimental vibrating beds, the estimation of the bubble velocity has the additional burden, not encountered in the simulation, of the matching between the camera and the bed vessel vibration, which can penalize the resolution of the bubble boundaries in the particle images. These factors could contribute to the differences encountered between the two fluid simulations and the experimental results concerning the mean velocity of bubbles. Nevertheless, the ultimate causes of the differences between the present two fluid simulations and the experiments by Zhou et al. [19] are unknown and further experimental data and computational studies are required to give an answer.

### 4.3.2. Effect of vibration on the pressure drop

Accordingly to the previous results, since the main dynamics of the bed is promoted by the presence of bubbles, and they become larger in size while varying the vibration strength, an analogous result to that obtained for the slim bed (configuration 1 in Section 4.2.2) is expected for the high amplitude vibrated bed (configuration 2). In particular, as the two fluid simulations presented in Fig. 13a indicate, the time averaged value of the pressure drop in the bed remains practically unchanged with the vibration frequency whereas the standard deviation of the pressure drop increases more than proportionally, i.e. in an approximately quadratic way. This trend is not so clear in the slim bed (configuration 1) where particles occupy a narrow volume affected by the presence of walls. When the vibration amplitude is augmented, Fig. 13b, the standard deviation also increases. However the standard deviation of the pressure drop in the bed grows linearly. The different growth rate of the standard deviation with frequency and amplitude can be explained resorting to the vibration strength, which is naturally incorporated through the equivalent mass force used in the proposed methodology for the two fluid simulation. In particular, after introducing Eq. (13) into Eq. (12) for the case of downward gravity acceleration, i.e. \( g_z = 9.81 \) m/s², it is easy to see that:

\[
f_{\text{m,eq}} = \{0, 0, g_z, [1, \sin(2\pi f t)]\}
\]

According to Eq. (14), an increase of the vibration strength augments the fluctuating intensity of the equivalent mass force \( f_{\text{m,eq}} \). Therefore, given that the vibration strength is proportional to the amplitude and the square of the frequency, Eq. (13), it seems evident that the vibration strength is the parameter that actually controls the magnitude of the oscillations observed for the gas pressure drop in a vibrated fluidized bed. This is confirmed in Fig. 14, in which all the results for the pressure standard deviation of Fig. 13a and b tend to collapse on the same curve as a function of the vertical vibration strength \( \Lambda_z \).
5. Summary and conclusions

A novel methodology to simulate vibrating beds using Eulerian Eulerian two fluid models was presented in this work. The method is based on expressing the governing equations in a moving system of reference for which vibration is incorporated through equivalent body forces. Thus, the method is relatively simple to implement. It eliminates the need for moving boundaries to cause vibration and the computational cost is similar than that for two fluid models applied to conventional non vibrating beds. According to the ob tained results, the dynamics of a slim vibrating fluidized bed sim ulated with the proposed two fluid model strategy was in reasonable agreement with the results of previous Lagrangian Eulerian simulations (DEM) present in the literature. This agree ment is remarkable considering that the small number of particles contained in the studied slim bed may be insufficient to fully reach the fluid like behavior of the particle phase that is assumed in two fluid models. Comparison with reported experimental studies of a vibrating quasi two dimensional bed seems to indicate that two fluid models are capable of predicting realistically the bubble behavior in vibrated fluidized beds even if the vibration is of large amplitude. Nevertheless, some discrepancies between the simulation results and the experimental measurements concerning the mean vertical velocity of bubbles would require further confir mation with additional experimental data.

The potential applications of the two fluid methodology pre sented here are significant. As two fluid models are able to deal with beds having larger amounts of particles than Lagrangian (DEM) models, the two fluid model methodology here described opens the way of dramatically increasing the scale of the vibrated fluidized beds currently simulated. Furthermore, nanofluidization processes that are mechanically assisted by vibration to reduce fluidized beds currently simulated. Furthermore, nanofluidization opens the way of dramatically increasing the scale of the vibrated (DEM) models, the two fluid model methodology here described presents here are significant. As two fluid models are able to deal with beds having larger amounts of particles than Lagrangian (DEM) models, the two fluid model methodology here described opens the way of dramatically increasing the scale of the vibrated fluidized beds currently simulated. Furthermore, nanofluidization processes that are mechanically assisted by vibration to reduce fluidized beds currently simulated. Furthermore, nanofluidization opens the way of dramatically increasing the scale of the vibrated fluidized beds currently simulated. Furthermore, nanofluidization opens the way of dramatically increasing the scale of the vibrated fluidized beds currently simulated. Furthermore, nanofluidization opens the way of dramatically increasing the scale of the vibrated fluidized beds currently simulated. Furthermore, nanofluidization opens the way of dramatically increasing the scale of the vibrated fluidized beds currently simulated. Furthermore, nanofluidization opens the way of dramatically increasing the scale of the vibrated fluidized beds currently simulated. Furthermore, nanofluidization opens the way of dramatically increasing the scale of the vibrated fluidized beds currently simulated. 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Furthermore, nanofluidization opens the way of dramatically increasing the scale of the vibrated fluidized beds currently simulated. Furthermore, nanofluidization opens the way of dramatically increasing the scale of the vibrated fluidized beds currently simulated. Furthermore, nanofluidization opens the way of dramatically increasing the scale of the vibrated fluidized beds currently simulated. Furthermore, nanofluidization opens the way of dramatically increasing the scale of the vibrated fluidized beds currently simulated. Furthermore, nanofluidization opens the way of dramatically increasing the scale of the vibrated fluidized beds currently simulated. Furthermore, nanofluidization opens the way of dramatically increasing the scale of the...