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Gas-liquid flow modeling in columns equipped with structured packing

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Abstract

This paper deals with the modeling of gas-liquid flow in distillation columns equipped with structured packing. The devices are seen as bi-structured porous media and a macro-scale model is proposed taking into account this specific geometry. In this model, the two liquid films, one-per-sheet, are treated separately and are allowed to exchange matter at the vicinity of the contact points between corrugated sheets. The model emphasizes mechanisms that lead to the liquid radial dispersion effects: a main part comes from the geometry itself, another part is due to the capillary effects. A particular attention is paid to model these phenomena from a macro-scale point of view. Finally, the simulation results are confronted to tomography imaging within a lab-scale column and show a qualitative good agreement of the liquid distribution.

Introduction

Structured packings play a large role in chemical engineering processes involving gas-liquid separation such as air distillation or CO₂ absorption columns. They are made of an assembly of vertically aligned corrugated sheets in which two adjacent sheets present symmetrical properties (i.e., ridges

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and furrows are symmetrically inclined by a given angle from one sheet to another, see Figure 1). One of their main advantages lies in their ability to allow a large surface exchange between gas and liquid all the while offering low pressure drops, which made their success in the early 1990’s when they progressively outpaced the plate distillation columns industry. In fact, the pressure drop through structured packing is only one-fifth to one-tenth that of a trayed column.¹

In order to design enhanced packings, it is of great importance to understand all the physical phenomena involved in the process and to be able to predict pressure loss, mass transfer efficiency and liquid spreading. Indeed, it is now well-established that a maldistribution of the liquid will heavily impact the efficiency of mixture separation.² Spiegel and Meier³ and Olujic et al.⁴ have provided a complete road map for improving the distillation processes over the next decades. One of the most important points they mentioned in their papers is the numerical modeling of gas-liquid flows. As a matter of fact, flow processes can be very complex and their modeling requires a large spectrum of knowledge in the fluid mechanics field. In a general way, the columns are operated in a counter-current flow mode: a thin gravity liquid film is sheared by the turbulent flow of a gas phase. All of these phenomena can be well captured by Computational Fluid Dynamic tools, which become more and more accurate and robust as they improve. For example, Sun et al.⁵ have recently simulated a 3D two-phase flow within a representative elementary volume of a structured packing using a Volume-Of-Fluid (VOF) method. However, since a huge number of grid cells is required, these kinds of simulations are restricted to very small domains and cannot therefore be conceivably used for simulating flows within the entire columns representing the complete packing geometry.

To do so, the henceforth well-established strategy consists in a multi-scale approach such as the one proposed by Raynal and Royon-Lebeaud,⁶ in which structured packings are apprehended as porous media with high porosity defining two scales of description: a small, pore-scale one and a macro-scale one assimilated to the packing scale. At the smaller scale, flows and other transfer phenomena are simulated using the exact geometry of the packing. Basically, to a cell of the mesh grid corresponds either a fluid or the solid structure and the two-phase flow can be computed using a VOF method involving gas turbulent modeling. On the other hand, at the macro-scale, the flow is governed by averaged equations. In this case, several continua are superimposed and a cell
of the grid contains both fluids and solids in the average sense. Information regarding the exact
topology of the packing and the interfacial phenomena between the gas and the liquid or between
the fluid and the solid structure are embedded in effective parameters such as permeability tensors
and exchange coefficients, which can be evaluated by taking into account the periodicity of the
pore-scale simulations.

In dry operating conditions, the flow at the larger scale is commonly modeled using a Darcy-
Forchheimer law.7 The effective parameters that appear in this law, also called Ergun’s coefficients,8
are evaluated from either lab-scale and industrial-scale measurements which relate pressure drops
to mass flow rates in the columns9–12 or using turbulent pore-scale simulations.13–21 Although this
modeling at the macro-scale of the turbulent single gas flow in columns equipped with structured
packings provides good predictions, there is still no consensus regarding the macro-scale model that
should be used for simulating gas-liquid flows in such devices. Indeed, due to this peculiar structured
geometry, the two-phase flow modeling from a macroscopic point of view remains a challenging
problem that is yet to be overcome. In particular, the macroscopic phenomena that lead to the
spreading of a liquid point source at the top of a packing are still misunderstood, and the classical
two-phase flow models in porous media, i.e. generalized Darcy’s laws, fail to properly capture the
liquid distribution within the column. This latter discussion was made possible by improvements in
tomography imaging,22–26 which provides a mapping of the liquid distribution within the packing
and is therefore an interesting tool for confronting simulation results with experimental ones.

By analogy with two-phase flow modeling in a fixed-bed, macroscopic models have emerged. For
instance, Iliuta et al.27 proposed a one-dimensional Eulerian two-fluid model with additional source
terms that represent the flow resistance due to the interaction of the fluids with the solid structure
and the mutual friction between liquid and gas. Although the overall pressure loss and the liquid
hold-up are in good agreement with the experimental results, this model does not reproduce the
liquid radial spreading. To simulate this phenomenon, chemical engineering authors28–32 used what
they called a "dispersion" model which consists in an advection-diffusion equation solved in a vertical
cylinder. Fourati et al.26 experimentally showed that the spread factor coefficient that appears in
this equation does not depend on the flow conditions. However, although these dispersion models are
in chemical engineering, they have no sufficient theoretical justifications. Some authors to model forces that lead to the mechanical dispersion by adding dispersive terms in tum equations. The origin of this additional term is confusing and can be interpreted ways: Lappalainen et al.\textsuperscript{33} assume that it comes from capillary pressure phenomena et al.\textsuperscript{34} suggest that it is a higher order term, associated to the local convection results from the upscaling procedure from the pore-scale to the larger scale.

the original macro-scale approaches developed to simulate this liquid radial dispersion columns, we can note the proposal of Aroonwilas et al.\textsuperscript{35,36} more recently used by Sun et modeling, the liquid distribution simulation relies on a discrete method. They consider the nodes represent the points of contact between two adjacent corrugated sheets.
t, the authors assume that the liquid film can flow in four different directions (two per represent either a corrugated angle or an angle due to the gravity effects. Mass flow h direction are then evaluated using a probability law. Sun et al.\textsuperscript{5} evaluate this stream VOF pore-scale simulations. The original work of Trifonov et al.\textsuperscript{37} is also based on

in mind that liquid flows in such devices with two preferential directions, Mahr and have found it convenient to split the liquid film into two separate phases flowing along t corrugated sheets. These phases are not (except perhaps at very low saturation) independent since adjacent sheets are in contact and the wetting liquid can flow from the other. This latter point is confirmed in the experimental study done by Alekseenko et al.\textsuperscript{40} who measured by fiber-optic sensors the thickness of the liquid film in the space confined by two adjacent corrugated sheets. They have shown that, in the vicinity of the contact points between corrugated sheets, menisci form is at the origin of local liquid flow redistribution over the corrugated surfaces. Based on Mewes et al.’s early work\textsuperscript{41} who investigated anisotropic porous structures, Mahr and Mewes\textsuperscript{38} have built a macroscale model from the integration over a control volume of a pore-scale two-phase problem where the liquid flow was split in two distinct phases. Their resulting model is a three-phase Eulerian one, consisting of a gas phase and two liquid phases between which mass transfer processes occur. They forced the preferential direction of liquid films
due to sheet orientations by using anisotropic permeability tensors whose values differ for each liquid phase.

From our point of view, the idea of separating a liquid film into two fictitious phases that exchange matter at the vicinity of the contact points between corrugated sheets is a major breakthrough in the modeling of the liquid distribution in the columns equipped with structured packings. This additional degree of freedom has a strong potential to catch the liquid radial spreading. In the present paper, we present an alternative model based on a similar idea. While Mahr and Mewes have heuristically considered the mass transfer term as a difference of saturation between the two fictitious phases, we introduce a different modeling that relies on a preliminary theoretical work done by Soulaine et al. In this work, authors qualified structured packings as bi-structured porous media and proposed macroscopic equations based on the volume averaging method to simulate fully saturated flows in such materials. They showed that the mass transfer between these two fictitious phases can be expressed as a difference between the two averaged pressure fields. We will discuss in the present study how this result can be extended to gas-liquid flow in structured packings.

The paper is organized as follows. In Section 2, we present the mathematical model developed to simulate gas-liquid flow in structured packings. A particular attention is paid to the treatment of mass exchange between liquid films. In Section 3, we present a simple model to evaluate multiphase permeability tensors from an analogy with inclined plane. Section 4 deals with the numerical procedure used to solve this problem. Finally, in Section 5 we numerically solve the gas-liquid flow through a Mellapak 250.X and compare our results with tomography images obtained by Fourati et al.

**Mathematical model**

The mathematical model introduced in this section is a heuristical extension to the case of gas-liquid flow of the two-pressure model proposed by Soulaine et al., which simulates flows in fully saturated bi-structured porous media. In the present model, a turbulent gas phase denoted $\gamma$ is flowing at counter-current of a liquid phase. Following the old saying *divide ut regnes*, this latter phase is split into two separate gravity films denoted $\beta_1$ and $\beta_2$ respectively with the same physical
fluid properties (density $\rho_\beta$, viscosity $\mu_\beta$, ...). Each film flows though the solid structure with two different preferential directions, namely one direction per corrugated sheet. Moreover, they can exchange matter at the vicinity of contact points between two adjacent corrugated sheets. The set of macro-scale equations that makes up the model combines three continuity equations and three multi-phase Darcy’s laws and is directly based on the Darcy generalized model originally proposed by Muskat\textsuperscript{44} for two-phase immiscible flows in porous media.

When studying multi-phase flows in porous media, it is handy to introduce the notion of saturation $S_i$. This variable corresponds to the filling rate of the pore-space by the $i$-phase. It is bounded by 0 and 1, and the sum of all saturation fields satisfies the obvious relationship,

$$S_\gamma + S_{\beta_1} + S_{\beta_2} = 1. \quad (1)$$

The whole liquid volume fraction is simply $S_\beta = S_{\beta_1} + S_{\beta_2}$. At packing-scale and assuming isothermal and incompressible flows, with no mass transfer at the gas-liquid interface and no chemical reactions, the saturation profiles are governed by the following macro-scale mass balance equations,

$$\varepsilon \frac{\partial S_\gamma}{\partial t} + \nabla \cdot \mathbf{U}_\gamma = 0, \quad (2)$$

$$\varepsilon \frac{\partial S_{\beta_1}}{\partial t} + \nabla \cdot \mathbf{U}_{\beta_1} = \dot{m}, \quad (3)$$

$$\varepsilon \frac{\partial S_{\beta_2}}{\partial t} + \nabla \cdot \mathbf{U}_{\beta_2} = -\dot{m}. \quad (4)$$

In these equations, $\mathbf{U}_i$ stands for the superficial velocity vector of the $i$-phase, $\varepsilon$ is the porosity of the packing and $\dot{m}$ models the amount of liquid in the $\beta_2$-phase which is transferred to the $\beta_1$-phase at the corrugated sheets contact points. We will discuss further ahead the expression of this mass exchange term.

In the generalized Darcy’s model, the superficial velocity of each phase is estimated using a phase Darcy’s law, which means that the superficial velocities and the pressure gradients are related by
an apparent permeability tensor:

\[ U_{\gamma} = -\frac{K_{\gamma}}{\mu_{\gamma}} \left( \nabla P_{\gamma} - \rho_{\gamma} g \right), \]  
\[ U_{\beta_1} = -\frac{K_{\beta_1}}{\mu_{\beta}} \left( \nabla P_{\beta_1} - \rho_{\beta} g \right), \]  
\[ U_{\beta_2} = -\frac{K_{\beta_2}}{\mu_{\beta}} \left( \nabla P_{\beta_2} - \rho_{\beta} g \right). \]  

In these equations, \( \mu_{\gamma} \) and \( \mu_{\beta} \), \( \rho_{\gamma} \) and \( \rho_{\beta} \) are respectively the viscosity and the density of the gas and liquid phases, \( g \) represents the gravity acceleration, \( K_i \) and \( P_i \) are the multi-phase permeability tensors and the averaged pressure fields of the \( i \)-phase respectively. This modeling suggests that the pore space where the gas phase flows is reduced by the presence of the liquid phase near the solid packing structure, and vice versa. Consequently, the multi-phase permeability tensors strongly depend on the liquid saturation profiles. Moreover, to account for inertia and turbulence effects, these tensors can also depend on the velocity of each phase.\(^7,8,45\) This point will be discussed in the next section.

The present momentum equations could be complexified to account for the viscous coupling due to the gas-liquid shear-stress\(^{46,47}\) and the coupling effects between the two fictitious liquid films.\(^{39,42}\) This effect can be of importance near the flooding point when hydrodynamics is mostly determined by the shear stresses on the gas/liquid interface. However, as observed in the numerical results of the last section, these simplifications emphasize the roles played by the packing geometry and by the situation in the vicinity of the contact points between corrugated sheets to explain the macroscopic phenomena that lead to the spreading of a liquid point source on top of a packing. In our model, as shown in the next section, \( K_{\beta_1} \) and \( K_{\beta_2} \) are tensors with non-zero off-diagonal coefficients. Both have different values that express different preferential directions due to the inclined angle of a corrugated sheet. If the shear stress effect participates to the liquid radial dispersion, it is not at leading order.

Besides this main liquid spreading effect due to the packing geometry, one can notice in Eqs (2)-(4) that at the opposite of most models proposed in the literature to simulate flows in gas-liquid contactors,\(^{27,39,41,48}\) we do not assume that the averaged pressure field in the liquid phases is equal
to the gas one. This results in an additional dispersion effect, as pointed out by Lappalainen et al.,\textsuperscript{33} caused by the capillary effects. Actually, these phenomena are well-known by multi-phase porous media investigators who introduce a macro-scale capillary pressure $p_c$ defined as the difference between both gas and liquid averaged pressure fields and function, at least, of saturation,\textsuperscript{49–51}

$$p_c(S_\beta) = P_\gamma - P_\beta.$$  

(8)

Wettability effects are generally included in the relative permeability and capillary pressure relationships for the generalized Darcy’s law model, which is based on a quasi-static view of the interface locations. In the case for structured packings, this latter assumption is not necessarily acceptable, and it is possible that modifications of the equation structure itself would be necessary. In general, this is still an open question. In the case of structured packings with film flows, i.e., low liquid saturation, we may have various situations. In cryogenic conditions, wettability is stronger and we believe that the potential dewetting of the solid surface is small, while this is probably likely to occur more rapidly for hydrocarbon compounds, for example. In this paper, we will skip that potential difficulty for the time being.

Since our model involves two liquid phases, we can, without any assumptions, introduce the following regional capillary pressures,

$$p_c(S_\beta_1, S_\beta_2) = P_\gamma - P_\beta_i \text{ with } i = 1, 2,$$  

(9)

and since by definition of the intrinsic averages, $P_\beta = \frac{S_\beta_1}{S_{\beta_1} + S_{\beta_2}} P_{\beta_1} + \frac{S_\beta_2}{S_{\beta_1} + S_{\beta_2}} P_{\beta_2}$ we may write the trivial relation,

$$p_c(S_1 + S_2) = \frac{S_\beta_1}{S_{\beta_1} + S_{\beta_2}} p_{c_1}(S_\beta_1, S_\beta_2) + \frac{S_\beta_2}{S_{\beta_1} + S_{\beta_2}} p_{c_2}(S_\beta_1, S_\beta_2).$$  

(10)

Actually, at the macro-scale, the capillary pressure translates the local curvilinear gas-liquid interface dynamic. The effect of capillary dispersion is therefore consistent with Alekseenko et al.\textsuperscript{40} experiments that showed the presence of a meniscus at the sheet’s contact points. They also showed
that, at this very spot, the liquid is redistributed over the corrugated surface. This means that some amount of liquid can be transferred from one sheet to its adjacent one, and inversely. In our model, this quantity is described by $\dot{m}$. When a fluid is split in two separate fictitious phases for up-scaling purposes, Soulaine et al.\textsuperscript{42} demonstrated that, for the one-phase flow case, the mass exchange rate is evaluated in terms of an averaged pressure fields difference

$$\dot{m} = \frac{h}{\mu_\beta} (P_{\beta_1} - P_{\beta_2}),$$ \hspace{1cm} (11)

where $h$ is a mass exchange coefficient. We adopt such a formulation for the multiphase flow case since it ensures that the sign of $\dot{m}$ will always follow the liquid stream-wise, something which is not granted by a mass exchange expressed as a linear function of the difference in both liquid volume fractions as proposed by Mahr and Mewes.\textsuperscript{39} Indeed, one may imagine geometries where the pressure difference and the saturation difference have opposite signs. Finally, introducing Eq (8) into Eq (11), the mass exchange equation reads

$$\dot{m} = \frac{h}{\mu_\beta} (p_{c_2}(S_{\beta_1}, S_{\beta_2}) - p_{c_1}(S_{\beta_1}, S_{\beta_2})), \hspace{1cm} (12)$$

and is therefore directly linked to the capillary effects in the vicinity of the contact points between corrugated sheets.

**Flow characterization at large scale**

The mathematical model introduced in the previous section involves several effective parameters as the multi-phase permeability tensors that relates the superficial velocities to the pressure gradients, or the mass exchange coefficient between the two fictitious liquid films. These parameters embed information related to both the packing geometry and flow configuration (gas-liquid and liquid-liquid interfaces). Therefore, they directly depend on the pore-scale physics and they can be evaluated from pore-scale simulations over a representative elementary volume of the packing. If this multi-scale analysis is now well-established for the evaluation of the gas permeability tensor,\textsuperscript{13–21} the prediction
of $K_{\beta_1}$ and $K_{\beta_2}$ from local simulations is more complicated since it requires the knowledge of the exact location of the gas-liquid interface. However, according to some assumptions, this local two-phase flow problem can be highly simplified and has some helpful analytical solutions.

Considering a corrugated sheet inclined by an angle $\theta$ from the vertical axis, a liquid film will flow downward according to an effective corrugation angle $\theta^*$ that varies between the corrugation angle $\theta$ and the gravity vector orientation. In order to characterize the liquid film, it is then common to consider the flow over a plane inclined by $\theta^*$ (see Figure 2). Actually, $\theta^*$ is an effective parameter that includes the 3D features of the corrugations. It may be evaluated from an analytical formula (see the proposal in Shilkin et al.\textsuperscript{52} as an example) or from a pore-scale simulation of films flowing over the corrugated surface. Moreover, most of the industrial packings are made of perforated metal sheets. These perforations lead to deviations of the liquid stream-wise and therefore impact the value of $\theta^*$. The dependency of $\theta^*$ to the sheet holes could be caught by film simulations over the real topology of the pierced corrugated surface.

Besides this geometrical conceptualization, further simplifications are made regarding the two-phase flow motion itself. First, we assume a free surface liquid film and neglect shear stress effects at the gas-liquid boundary. Then, we neglect inertial effects, which means that the film’s thickness is constant along the inclined plane. With such a configuration, the liquid flow is governed by a simple analytical solution\textsuperscript{53} in the coordinate system related to the inclined plane. The averaging of this solution leads to a linear relationship between the superficial velocity and the gradient of the averaged pressure field. Actually, this relationship looks like a multi-phase Darcy’s law with a relative permeability that depends on the $\beta_1$-liquid saturation. With simple linear algebra, this relation can be expressed in the coordinate system related to the packing, we have

$$
K_{\beta_1} = \frac{K_0}{2} S_{\beta_1}^{3} \begin{pmatrix}
\sin^2(\theta^*) & \cos(\theta^*)\sin(\theta^*) & 0 \\
\cos(\theta^*)\sin(\theta^*) & \cos^2(\theta^*) & 0 \\
0 & 0 & 0
\end{pmatrix}.
\tag{13}
$$

The complete derivation that leads to this formula can be found in Soulaine’s work.\textsuperscript{45} Similarly,
from the solution of a liquid film over a plane inclined by $\theta^*$ we get,

$$K_{\beta_2} = \frac{K_0}{2} S_{\beta_2}^3 \begin{pmatrix} \sin^2(\theta^*) & -\cos(\theta^*)\sin(\theta^*) & 0 \\ -\cos(\theta^*)\sin(\theta^*) & \cos^2(\theta^*) & 0 \\ 0 & 0 & 0 \end{pmatrix}. \quad (14)$$

In these two multi-phase permeability tensors, $K_0$ represents the permeability value if two adjacent corrugated sheets were put together with $\theta = 0^\circ$. We can notice the presence of off-diagonal terms in the $xOy$ plane of the sheet. Moreover, all the zero values in the $z$ direction mean that there is no liquid flow in this direction. In fact, some amount of liquid can flow in this direction by crossing the sheet perforations, but this effect is probably very small. The permeability tensors of the packing rotated by $90^\circ$ are merely obtained by switching the coefficients of subscript $x$ with the ones of subscript $z$. In Eqs (13)-(14), we recognize a relative permeability in terms of the liquid saturation flowing over each corrugated surface to the power of three ($S_{\beta_1}^3$). This relative permeability, reminiscent of the proposal of Brooks and Corey, is intrinsic to the assumption leading to the laminar free surface flow over an inclined plane. A deeper investigation of the saturation dependency could be performed from pore-scale simulations on a more realistic topology of the corrugated sheet. This could also provides some information regarding the effect of inertia and enlighten a potential dependency to the averaged liquid film velocity.

Concerning the vapor permeability tensor, the coefficients could be obtained from dry gas turbulent pore-scale simulations over a representative elementary volume of the packing or from lab-scale measurements (see the review in the section of this paper). In the spirit of a Forchheimer’s law, the permeability tensor should depend on the Reynolds number to account for the inertia and turbulence effects. The resulting tensor is weighted by a relative permeability to account for the reduction of the pore space where the gas flows, due to the presence of the liquid films at the sheet walls. In the absence of accurate two-phase flow simulations over the real structure and to remain consistent with the liquid permeability tensors, we postulate a Brook and Corey weighting function. Finally,
we have
\[
K_\gamma = \frac{K_0}{1 + \alpha Re_\gamma} S^3_{\gamma} \begin{pmatrix}
\sin^2(\theta) & 0 & 0 \\
0 & \cos^2(\theta) & 0 \\
0 & 0 & 0
\end{pmatrix},
\] (15)

where \(\alpha\) is a model parameter that denotes the inertia/turbulence effects in the gas phase.

The mathematical model we have introduced in the previous section involves two capillary pressures, \(p_{c1}\) and \(p_{c2}\), that should satisfy the relation Eq (10). Actually, without deeper theoretical investigations, we have no further informations regarding the saturation dependency of \(p_{ci}\). To simplify the problem, we postulate, in agreement with Eq (10), that,
\[
p_{ci}(S_{\beta_1}, S_{\beta_2}) = p_c(S_{\beta_i}) \text{ with } i = 1, 2.
\] (16)

The problem is therefore reduced to the evaluation of the capillary pressure \(p_c\) that could be measured experimentally using classical methods. Here, we are going to use the well-known Brook and Corey correlation:\(^{50}\)
\[
p_c(S_{\beta_i}) = p_{c0} (S_{\beta_i})^{\frac{1}{\lambda}}
\] (17)

where \(p_{c0}\) is the entry capillary pressure and \(\lambda\) is the pore-size distribution index. This coefficient has a small value for heterogeneous pore size distribution and a large value when the pore size distribution is regular. We will fix its value at \(\lambda = 2.\)\(^{57}\) Hence, only \(p_{c0}\) remains a model parameter.

The exchange coefficient \(h\) is the latest effective parameter that appears in our model. It quantifies the amount of liquid that flows from a sheet to its closest neighbor at a sheet contact point. \textit{Stricto sensu}, it should depend on the saturation profiles and on the liquid Reynolds number if the film flow involves inertia effects. In the absence of accurate pore-scale simulations, we will consider that \(h\) is only weighted by the liquid saturation,
\[
h = (S_{\beta_1} + S_{\beta_2}) h^*,
\] (18)

where \(h^*\) is a constant dimensionless parameter. Further investigations will be needed to clarify
Numerical implementation

In this section, we focus on the implementation of the mathematical problem introduced in Section 2. We chose to code it using the finite volume CFD library OpenFOAM®. From a numerical point of view, this problem can be seen as a three-phase flow problem in which a gas phase and two liquid phases are flowing through an anisotropic porous medium. Since the sum of the 3 saturations is equal to unity and since the liquid pressures can be related to merely the gas pressure from the capillary pressure concept, the problem can be reduced to the resolution of a gas pressure \( P_\gamma \) and two liquid saturation fields \( S_{\beta_1} \) and \( S_{\beta_2} \). The numerical treatment of this kind of problem, well-known by the petroleum industry, has received a lot of attention in the literature and several books offer comprehensive reviews of the resolution methods available. Among the different methods available, we retained the IMPES algorithm (IMplicit Pressure, Explicit Saturation). The main idea of this widely used method is to solve the pressure and saturation equations in a segregated manner, this choice being compatible with the OpenFOAM® algorithmic structure. At each time step, the pressure equation is solved implicitly whereas the saturation equations are solved explicitly. This method has the multiple advantages of being quite simple to implement and requiring less computer memory compared with other methods such as a simultaneous solution method or the Euler-Euler models with porous source terms. Moreover, as it will be further discussed, an efficient time-step management leads to stable and very fast computations. The 3-phase solver is developed on an existing two-phase IMPES solver \textit{impesFoam} that has been developed in a previous work. The following paragraphs introduce the main parts of the numerical model solving multi-phase flows in structured packings.

The first step of the IMPES method consists in the prediction of the gas pressure field within the computational domain. This stage requires a partial differential equation that governs \( P \equiv P_\gamma \).
To form this latter, the three velocities are expressed as

\[
U_\gamma = -M_\gamma \nabla P + L_\gamma \cdot g, \tag{19}
\]

\[
U_{\beta_1} = -M_{\beta_1} \nabla P + L_{\beta_1} \cdot g + M_{\beta_1} \nabla P_{c_1}, \tag{20}
\]

\[
U_{\beta_2} = -M_{\beta_2} \nabla P + L_{\beta_2} \cdot g + M_{\beta_2} \nabla P_{c_2}, \tag{21}
\]

where \(M_i\) represents the mobility tensor of the \(i\)-phase, and the tensor \(L_i\), which has a time dimension, deals with the contribution of gravitational effects. It must be noted that more complex models involving viscous resistance terms between phases for example could be written using this generic formulation. Therefore, that same solver could be used for further, and possibly more complex, investigations of this problem.

Defining the total velocity \(U\) as the sum of the phase-velocities, we have

\[
U = -M \cdot \nabla P + L \cdot g + M_{\beta_1} \nabla P_{c_1} + M_{\beta_2} \nabla P_{c_2}, \tag{22}
\]

where

\[
M = \sum_i M_i \quad \text{and} \quad L = \sum_i L_i \quad \text{with} \quad i = \gamma, \beta_1, \beta_2, \tag{23}
\]

since the sum of the continuity equations Eqs (2) to (4) give a divergence-free velocity \((\nabla \cdot U = 0)\), we can therefore formulate the following pressure equation:

\[
\nabla \cdot ( -M \nabla P ) + \nabla \cdot ( L \cdot g + M_{\beta_1} \nabla P_{c_1} + M_{\beta_2} \nabla P_{c_2} ) = 0. \tag{24}
\]

This equation is solved implicitly using the \(M_i\) and \(L\) tensors and the capillary pressures computed with the latest values of the liquid saturation profiles \((S_{\beta_i}^{n-1})\). The velocity fields are then deduced from Eqs (19)-(21) using the new computed gas pressure field \(P^n\). Then, the liquid mass balance equations Eq (3) and Eq (4) are solved explicitly one after the other. Finally, the gas saturation can be merely deduced using the values of \(S_{\beta_1}^n\) and \(S_{\beta_2}^n\) from Eq (1). At the end of this stage, the multi-phase permeability tensors and the capillary pressures are updated with the new values of the saturation profiles.
Because of the hyperbolic feature of these conservation laws and the strong non-linearities that exists in the model (relative permeabilities and capillary pressure correlations), solving this set of equations can lead to high instabilities. To ensure stability of the numerical simulations, the flux terms have been discretized with a first order upwind scheme.

Because of their explicit treatment, the computation of both saturation equations requires small time steps limited by a CFL condition. On the contrary, the implicit treatment of the pressure equation is time-consuming. To significantly reduce the CPU time, we embrace the smart time-step management method proposed by Chen et al. who noticed that the pressure field relaxation was much longer than the evolution of the saturation. Hence, in their method, the pressure equation is no longer solved at each time step. It results in a very stable and fast algorithm that could allow the use of this solver in an industrial background.

Results and discussion

Now that we have introduced the conceptual, mathematical and numerical basis of our gas-liquid flow model in structured packing, we are going to compare the simulation results with the liquid distribution experimentally obtained by Fourati et al. In their experiment, they have mapped liquid distribution by means of gamma-ray tomography at different positions downstream a source point injection. Their column was 40 cm in diameter and 1.5 m in height and was filled with six 22 cm high layers of structured packings Mellapak 250.X manufactured by Sulzer Chemtech (see Figure 3). The packing geometry is characterized by a large porosity $\varepsilon \approx 0.98$ and an inclination angle of the flow channels with the vertical direction of $\theta \approx 30^\circ$. The packing elements were alternatively rotated around the axis of the column by 90° relative to each other. The column was operated in the counter-current flow mode and was fed at the top by a liquid (water) point source (the inlet was approximatively a 50 mm circular section). As for the gas (air), it was injected at the bottom of the column. Local liquid hold-up profiles were mapped for several cross-sections ($Z_1$, $Z_2$ and $Z_3$ in Figure 3).

We have tried to reproduce these experimental results with the model we have developed. However, all the exact protocol of Fourati et al. has not been respected and we have adopted some
assumptions. First, we do not consider gas injection in the column to focus on the radial liquid dispersion only. This is consistent with the conclusions of Fourati et al.\textsuperscript{26} who have operated the gas-liquid flow at counter-current for a wide range of mass flow rates, and found that the radial dispersion coefficient does not vary significantly with the flow conditions. Hence, this assumption seems suitable at leading order. In addition, we have supposed that the corrugated sheets were not perforated. In fact, these perforations should modify the value of $\theta^*$ and the inclined plane analogy is no longer true. From the Mellapak 250.X geometrical features and using the formula proposed by Shilkin and Kenig,\textsuperscript{65} we obtain that $\theta^* \approx 20^\circ$. According to our definition, $K_0$ is the permeability of the packing if two adjacent sheets were assembled with corrugations aligned with the vertical axis. Hence, using a hydraulic radius this permeability can be approximated by $K_0 = \frac{R^2}{8} = \frac{H^2b^2}{4H^2b^2}$ where $H = 1.2$ cm and $b = 2.4$ cm are the dimensions of the corrugation (see Figure 1). We obtain $K_0 \approx 10^{-5}$ m$^2$. Since we do not account for gas turbulence in these simulations, $\alpha \equiv 0$. Our model requires additional informations, especially those regarding the capillary pressure coefficients $p_{c0}$ and $h^*$ in the mass exchange function ($\dot{m}$). An adjustment from experiments or pore-scale simulations is beyond the scope of this paper. Instead, we have performed a sensitivity analysis to these parameters and compared the results with the saturation profiles in the cross-section $Z_2$ that belongs to the first pack. Therefore, we have simulated the flow in the first layer for different values of $p_{c0}$ and $h^*$. The computational domain is a cylinder whose height is 22 cm and the diameter is 40 cm, meshed with 72000 hexaedea (see Figure 4).

The liquid injection is carried out at the center of the highest cross-section as depicted in Figure 4 with $U_{\beta_1} = U_{\beta_2} = 0.015$ m/s. The physical properties of air and water in the simulation are respectively $\rho_\gamma = 1.17$ kg/m$^3$, $\mu_\gamma = 1.85 \times 10^{-5}$ kg/m/s and $\rho_\beta = 998$ kg/m$^3$, $\mu_\beta = 10^{-3}$ kg/m/s. Once the steady-state is reached, in the order of a few seconds, we have qualitatively compared the liquid distribution at the cross-section $Z_2$. In this plane, located at the bottom of the pack, Fourati et al.\textsuperscript{26} have shown that the liquid hold-up, initially circular, has spread in the direction of the corrugated sheets and that the liquid saturation was more important at the center of the cross-section (see Figure 5a). We have tried different sets of values for $p_{c0}$ and $h^*$. In the absence of deeper experimental investigations like quantitative measurements of the capillary effects at the
vicinity of the contact points, it is very difficult to decide which set of values is the most realistic. Nevertheless, this sensitive analysis helps to understand how the model behaves according to the weight of the liquid mass exchange function. In Figure 5 we have plotted the simulation results we obtained for \( p_{c0} = 0.1 \text{ kg/m/s}^2 \) and \( h^* \) varying in the range 0 to 10. For \( h^* = 0 \) (see Figure 5b), one notices two separate liquid distributions. In this case, there is no liquid transfer from one sheet to another (\( \dot{m} = 0 \)) and a film remains on the same sheet during all the process. For larger values up to \( h^* = 3 \) (see Figure 5c,d,e), the simulation results are in keeping with the experimental hold-up mapping. However, there are two peaks on both sides of the packing center which does not appear in Figure 5a. The value \( h^* = 4 \) is in better agreement with the tomography results and is the one we will chose for the following simulations. For the highest values of the mass exchange coefficient, the radial liquid spreading is very weak, and the saturation pattern at the bottom of the packing is similar to the injection one. In this case, \( i.e., \) for \( h^* \) high enough we are in a situation of local equilibrium and we recover the dynamic of the classical two-phase flow in porous media model. This sensitivity analysis illustrates that, besides the role played by the structured packing geometry, the radial dispersion phenomenon is also governed by capillary effects at the vicinity of the contact points between two adjacent corrugated sheets.

Once the coefficients related to the liquid mass exchange are determined, we have simulated the flow within the three first packing layers. The first and the third layers are characterized by the permeability tensors \( K_{\beta_1}, K_{\beta_2} \) and \( K_\gamma \) defined by Eqs (13), (14) and (15). For the second pack which is rotated around the axis of the column by 90° relative to the former layer, its permeability tensors are merely obtained switching the coefficients with index \( x \) by those indexed \( z \) in these equations. In this work, we simulate the gas-liquid flow in the column one structured packing at a time. An alternative solution could be to simulate the gas-liquid flow within the whole column by defining two different porous media with different permeability tensors according to their rotation angle around the vertical axis. The first method combines the advantages to reduce CPU time and to directly use the solver as detailed in the previous section. Once the steady-state is reached in the first pack, we get the outlet boundary values of the velocity fields and apply them at the inlet of the next pack following,
\[ U_{\beta_1}^{n+1} \cdot n_{inlet} = U_{\beta_2}^{n+1} \cdot n_{inlet} = \frac{U_{\beta_1}^n \cdot n_{outlet} + U_{\beta_2}^n \cdot n_{outlet}}{2}, \]  

(25)

where \( U_{\beta_1}^{n+1} \cdot n_{inlet} \) and \( U_{\beta_2}^{n+1} \cdot n_{inlet} \) are the liquid velocity boundary values at the inlet of a pack while \( U_{\beta_1}^n \cdot n_{outlet} \) and \( U_{\beta_2}^n \cdot n_{outlet} \) denote the boundary values at the outlet of the previous pack. This relation means that there is no additional redistribution of the liquid flow rate across the section at packing elements junctions. Actually, an accumulation of liquid in this zone due to capillary effects that generates a liquid film attached to the extremities of the packing metal sheets has been experimentally reported.\(^{66-68}\) The liquid saturation at junctions between packing units is twice more important than in the interior of the packing unit. If this phenomena should create additional pressure drop, Fourati et al.\(^{26}\) have experimentally shown that it does not strongly participate to liquid redistribution. The hereby model does not account for this accumulation phenomena. However, it could be easily enhanced in this direction considering an additional porous medium layer with different physical properties in between two successive packs. An alternative model would introduce a specific boundary condition instead of Eq (25). Such a condition for the split model discussed in this paper has not been developed yet and is beyond the scope of this paper.

The liquid saturation profile is plotted in Figure 6 according to a slice in the \( xOy \) plane (on the left) and another in the \( zOy \) plane (on the right), the \( y \) axis being the vertical axis, \( xOy \) corresponding to the orientation of the corrugated sheet of the first and third packing elements and, as a matter of fact, \( zOy \) those of the second pack. We notice that the liquid only spreads according to the \( x \) direction within the first and the third packs while the radial dispersion only occurs into the \( z \) direction in the second packing unit. In Figure 7, we have compared the simulation results with the liquid hold-up maps in the cross-section of the column obtained by tomography imaging.\(^{26}\) Simulations are qualitatively comparable to the experimental results: as expected we get a liquid spreading following a preferential direction (\( x \) or \( z \) according to the structured packing orientation) that leads to a liquid distribution almost homogeneous after several packing transitions. However, the order of magnitude of the liquid saturation is twice lower than the experimental values and we can notice that, in the third pack (Figure 7, \( Z_3 \) plane), the liquid spreading is not as diffusive as the experimental results. We have identified several points that can explain this result. First,
the effective corrugation angle estimation we use does not account for the sheet perforations. A more accurate value of $\theta^*$ could be obtained from pore-scale simulations of the liquid film flow over a realistic topology of the corrugated sheets. Furthermore, strong assumptions (laminar flow, constant film thickness...) have been made to derive the simple liquid film model we used to evaluate the multiphase permeability tensors. Indeed, regarding this latter point, Raynal et al. have shown that the liquid thickness evaluated from the Nusselt analytical solution may be underestimated up to twice the expected value, even more. Third, the coefficients of the transfer function ($\dot{m}$) have been approximatively adjusted and further investigations deserve to be carried out. Finally, as it has already been mentioned, the condition we used at the junction between two successive packing elements is a simplified view of the reality.

**Conclusion**

We have developed a mathematical model based on a multi-scale analysis to simulate gas-liquid flow through distillation columns equipped with structured packing. In this modeling, the packing is seen as a bi-structured porous medium considering each group of oriented sheets as a separate structure. The two liquid films, one-per-sheet, are then seen as two overlapping continua where the interaction between each other is modeled through a transfer function.

This modeling allows one to understand the mechanisms that lead to the liquid spreading. A major part of this spreading is directly induced by the criss-crossing structured geometry of the packing that forces the liquid films to flow following two different preferential directions. Combined to these geometric features, the capillary effects play an important role since they govern the transfer of some amount of liquid from a corrugated sheet to another sheet at the vicinity of the contact points. The simulations performed with this model have been successfully compared to laboratory-scale measurements. In particular, the model can properly catch the liquid distribution within the column.

However, the numerical results are mainly qualitative and show the model potential. Indeed, the effective properties like relative permeabilities, the effective corrugation angle and the capillary pressures have been evaluated from simple analytical solutions. Further investigations regarding their
saturation-dependency from either accurate pore-scale simulations or experimental measurements could make the model becomes quantitative as well.

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