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Numerical modeling of local capillary effects in porous media as a pressure discontinuity acting on the interface of a transient bi-fluid flow

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Abstract

Transient flows through porous media can be controlled by local capillary forces. In an attempt to ease the representation of these complex multiscale flows, this article presents a new numerical approach to account for these local forces, viewed as a global pressure discontinuity acting in bi-fluid flows through smeared-out porous media. A finite element discretization of the Darcy's equations is considered and a pressure enriched space is locally introduced at the fluid interface in order to capture the pressure discontinuity. Then, a Variational Multiscale Stabilization (VMS) method is selected to take into account the subgrid effects on the finite element solution and hence ensure the consistency of the finite element formulation. The fluid front is represented by a level set function, convected with the fluid velocity thanks to a finite element scheme stabilized with a Streamline-Upwind/Petrov-Galerkin (SUPG) method. Both convergence and implementation are first validated with the Method of Manufactured Solution (MMS) and the model shows a good convergence. Second, a comparison with experimental measurements in the case of capillary wicking of water into carbon reinforcements shows a very good correlation between experimental and numerical results.

Keywords: Capillary stress, Darcy's equations, stabilized finite element

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method, discontinuous pressure, capillary wicking

1 1. Introduction

Capillary effects define the ability of a liquid to maintain contact with a solid. They are localized and play a key role in the description of liquid flows in porous media. Their best visualization is the spontaneous wicking mechanisms, where the liquid flows without the assistance of any external force. Such phenomena are related in one hand to the surface tension between the liquid and the surrounding media such as air, and on the other hand, to the liquid-solid and air-solid surface energies.

During flows in a porous medium, micro- and macro-voids may develop 9 following the competition between viscous and capillary effects. This work 10 focuses on the capillary effects occurring in porous media such as fibrous re-11 inforcements during infusion process, especially Liquid Resin Infusion (LRI) 12 process. Indeed, capillary effects are traditionally neglected in the flow simu-13 lation during high pressure composite manufacturing processes. Meanwhile, 14 experimental studies have shown that the capillary stress resulting from the 15 interaction of carbon reinforcements and liquid, such as water or epoxy resin, 16 can reach a value of 0.3 - 0.4 bar in quasi-UD fabrics [62, 63], at the fibre 17 scale [76] or in carbon woven fabrics [5]. This value represents approxima-18 tively one third of the 1 bar driving force available in the LRI process, which 19 is too significant to be neglected and even permits to manufacture compos-20 ite parts that could not be without its contribution. Consequently, the aim 21 of this paper is to introduce these local capillary effects, in order to assess 22 their influence on the filling stage scenarios at the scale of composite parts 23 [46, 70].24

Indeed, following the multi-scale nature of high performance composites, 25 the study can be conducted at three different scales as shown in many stud-26 ies [15, 22, 36, 60]: at the fiber or microscopic scale ($\sim 10^{-6}$ m), at the 27 tow or mesoscopic scale (~ 10^{-3} m) and at the process or macroscopic scale 28 $(\sim 10^{-1} {\rm m})$. At microscopic scale, numerical modeling of the capillary rise 29 is well-documented [11, 13, 52, 71]. Capillary effects are accounted for into 30 Navier-Stokes or Stokes equations by the mean of the surface tensions be-31 tween the three phases (solid-liquid-air) [4, 20, 24]. At mesoscopic scale, 32 capillary effects are taken into account through the micro-diffusion within 33 fiber tows. Generally, unsaturated flow models are adopted to describe this 34

problem. A sink term which depends on the capillary number is incorporated in the governing equations to model the capillary effects [65, 67, 72], but standing only for isotropic representations. In many research areas such as ground water infiltration or oil recovery, the capillary pressure is related to the saturation according to different analytical parameterizations such as the ones proposed by Van Genuchten, Stauffer, Kalaydjian, Hassanizadeh, Gray and Bareblatt [38, 47, 50].

This approach has been extended to composite manufacturing fields in 42 order to simulate the filling stage and to assess void formation [17, 54, 59]. 43 Capillary effects are usually represented through the introduction a single 44 capillary pressure, although capillary effects do come from local mechanisms 45 related to both orthotropic micro(meso)-structure architecture and surface 46 tensions. Dedicated studies concentrate on these local phenomena [24] which 47 are, for the moment, out of reach in tractable models at the structure scale. 48 An alternative way of introducing these 3D effects at an upper scale is to 49 consider their effect as a capillary stress tensor (3D representation) acting 50 on the fluid-gas interface in a slug-flow approach, *i.e.* no saturation zone is 51 considered at this scale. Accounting properly for the 3D pressure disconti-52 nuity in a numerical approach will allow to complete full models of infusion 53 processes at the structure scale [15] including coupling with the wet/dry pre-54 form mechanical response, as well as to model dual-scale flows at the tow 55 scale [75] provided an equivalent homogeneous medium can be used to rep-56 resent populations of fibres. Notwithstanding any local effects related either 57 to velocity or pressure fields which are of utmost interest to represent lo-58 cal physical changes, such as for shear-rate dependent fluids, void creations 59 issues, or fluid pressure acting in wet preforms for coupling issues [22, 15]. 60

In this work, an innovative macroscopic approach is adopted. The cap-61 illary action is described by a capillary stress tensor acting on the liquid-air 62 interface. The subsequent jump of the pressure field at the flow front is taken 63 into account numerically in the weak formulation of Darcy's equations. Those 64 equations, established in a velocity-pressure mixed form, are solved using a 65 Finite Element Method (FEM). Both velocity and pressure are approximated 66 by continuous and linear fields. According to the Brezzi - Babuška theory, 67 such an approximation is not stable. This issue is overcome by stabilizing the 68 Finite Element (FE) formulation thanks to the Variational MultiScale (VMS) 60 framework introduced by Hughes [39, 40] and extensively used and studied 70 by Badia & Codina to stabilize Stokes', Darcy's and Maxwell's equations in 71 a unified setting [7, 8, 9, 10]. However, a special attention is mandatory to 72

⁷³ accurately capture the pressure discontinuity across the flow front, which is⁷⁴ described here with a level-set method [57, 58, 69].

The literature provides several techniques to capture this local phenomenon. 75 For instance, the Extended-Finite Element Method (X-FEM) [21, 32, 45] con-76 sists in enriching the pressure approximation space by discontinuous func-77 tions. This enrichment, which is not localized in the mesh elements, pro-78 vides additional degrees of freedom, resulting in some computational issues 79 when the discontinuity is moving (the mesh, and consequently the global 80 "stiffness" matrix, need to be updated). A discontinuous Galerkin formu-81 lation [1, 12, 48, 56] can be another way of dealing with singular forces at 82 the interface. As the continuity between elements is weakly imposed, it al-83 lows the solution to be discontinuous: each element has its own degrees of 84 freedom and is connected to its neighboring by numerical fluxes. However, 85 in our simulations, the interface does not necessarily correspond to edges of 86 elements, but can cut these elements. In this work, the jump of pressure 87 field is captured using the technique developed by Ausas *et al.* [6], which 88 consists in a local enrichment of the pressure space by discontinuous func-89 tions. Unlike the X-FEM approach, the corresponding additional degrees of 90 freedom are local to an interface element, and can therefore be eliminated at 91 the elementary level before the final assembly. 92

The rest of this article is divided into four parts. Section 2 focuses on 93 the mathematical description of the fluid flow problem, and the finite ele-94 ment strategy implemented. Section 3 describes the level-set method used 95 to capture the fluid front, *i.e.* the interface across which pressure is discon-96 tinuous. An error analysis is given in Section 4 to assess the accuracy of the 97 numerical developments. Finally, Section 5 compares simulation and exper-98 imental results for water capillary wicking in carbon reinforcements. Also, 99 a 3D simulation of the resin flow through an orthotropic stiffener is carried 100 out. 101

¹⁰² 2. Fluid flow problem

¹⁰³ 2.1. Physical and mathematical description

Let Ω be a region of \mathbb{R}^d (with d = 2, 3 the spatial dimension) bounded by $\partial \Omega$ (see Fig. 1). Ω represents a porous medium, the fibrous preform in our context, considered as an equivalent homogeneous orthotropic medium characterized by a porosity ϕ and a saturated permeability K independent on the fluid. The permeability is a measure indicating the capacity of the material to allow fluids to pass through it. In a realistic description, the fibrous reinforcement is anisotropic and \boldsymbol{K} is a symmetric tensor. The domain Ω is filled with two immiscible, Newtonian and incompressible fluids: a liquid of viscosity μ_l , occupying the subdomain Ω_l and a surrounding medium (for instance, the air) of viscosity $\mu_a \ll \mu_l$, occupying Ω_a . Hence: $\Omega = \Omega_l \cup \Omega_a$. The interface is denoted $\Gamma_{l/a}$: $\Gamma_{l/a} = \partial \Omega_l \cap \partial \Omega_a$.

The domain boundary $\partial \Omega$ is divided into two types of boundary: $\partial \Omega_D$ and $\partial \Omega_N$ such that $\partial \Omega = \partial \Omega_D \cup \partial \Omega_N$ and $\partial \Omega_D \cap \partial \Omega_N = \emptyset$, where respectively Dirichlet and Neumann boundary conditions are prescribed.



Figure 1: 2D-description of the domain Ω .

¹¹⁸ Darcy's equations [28] express the flow velocity \boldsymbol{v} at the scale of the ¹¹⁹ homogeneous equivalent medium, *i.e.* the superficial fluid velocity, with ¹²⁰ respect to the pressure gradient ∇p and two parameters: the fluid viscosity ¹²¹ and the permeability (Eq. 1). The mass conservation is introduced through ¹²² the divergence of the velocity \boldsymbol{v} (Eq. 2). Hence, the governing equations are

$$\mu \boldsymbol{K}^{-1} \boldsymbol{v} + \nabla p = \boldsymbol{f}, \quad \mu = \begin{cases} \mu_l & \text{in } \Omega_l \\ \mu_a & \text{in } \Omega_a \end{cases}$$
(1)

$$\nabla \cdot \boldsymbol{v} = h \tag{2}$$

with f the external forces and h a source/sink term, equal to zero when the fluids are assumed to be incompressible. Assuming, for a while, that the axes of the orthonormal coordinate system $\{x, y, z\}$ coincide with the normals to the three symmetry planes of the orthotropic material, the permeability ¹²⁷ tensor writes in this eigen-system:

$$\boldsymbol{K} = \boldsymbol{K}_{LOC} = \begin{pmatrix} K_x & 0 & 0\\ 0 & K_y & 0\\ 0 & 0 & K_z \end{pmatrix}$$
(3)

with $K_{\{x,y,z\}}$ the values of the permeability in the x-, y- and z-directions, the index $_{LOC}$ refers to the local or material coordinate system.

The capillary effects are described at the macroscopic scale by a capillary stress tensor, σ_{cap} , having the same eigen-directions as the permeability tensor, since they correspond to the symmetries of the orthotropic porous medium. Hence,

$$\boldsymbol{\sigma}_{cap} = \boldsymbol{\sigma}_{cap}^{LOC} = \begin{pmatrix} \sigma_{cap}^{x} & 0 & 0\\ 0 & \sigma_{cap}^{y} & 0\\ 0 & 0 & \sigma_{cap}^{z} \end{pmatrix}$$
(4)

with $\sigma_{cap}^{\{x,y,z\}}$ the components of the equivalent capillary stress at the interface $\Gamma_{l/a}$ in the x-, y- and z-directions. In general situations, described in section 5.3, the eigen-directions of the previous tensors vary from point to point, and consequently, do not match with the axes of the global coordinate system. In this case, tensor K is expressed in the global system by: K = $QK_{LOC}Q^{T}$, where Q is the orthogonal tensor expressing the passage from the local to the global bases. Similarly, $\sigma_{cap} = Q\sigma_{cap}^{LOC}Q^{T}$.

¹⁴¹ Capillary effects give rise to a jump of pressure across $\Gamma_{l/a}$. This jump, ¹⁴² denoted [p], is expressed as

$$[p] = \boldsymbol{n} \cdot \boldsymbol{\sigma}_{cap} \cdot \boldsymbol{n} \quad \text{on} \quad \Gamma_{l/a} \tag{5}$$

¹⁴³ where n is the normal vector to the interface.

Finally, system (Eq. 1), (Eq. 2), completed by (Eq. 5), is closed by prescribing a normal velocity v_0 on $\partial \Omega_D$ and a pressure p_0 on $\partial \Omega_N$:

$$\boldsymbol{v} \cdot \boldsymbol{n} = v_0 \qquad \text{in } \partial \Omega_D \qquad (6)$$

$$p = p_0 \qquad \text{in } \partial \Omega_N \tag{7}$$

$$[\boldsymbol{v} \cdot \boldsymbol{n}] = 0 \qquad \text{on } \Gamma_{l/a} \tag{8}$$

where the last condition (Eq. 8), the continuity of the normal velocity, expresses the mass conservation across the interface $\Gamma_{l/a}$.

148 2.2. Weak formulation

In order to solve the previous Darcy's system with a finite element method, 149 the weak formulation of these equations has first to be established. Two ap-150 proaches exist to express the weak formulation. First, the Darcy's problem 151 can be formulated in pressure only, and the velocity post-calculated apart. 152 However, mass conservation issues can appear when considering a jump of 153 material properties such as a jump of permeability [29]. Second, and this 154 is the strategy adopted here, a full velocity/pressure mixed weak formula-155 tion can be formulated, ensuring the mass conservation. Moreover, what is 156 called the dual formulation of Darcy's equations [8, 34] is chosen, in order to 157 naturally enforce the pressure discontinuity. 158

The dual variational formulation is obtained by multiplying the strong equations (Eq. 1) and (Eq. 2) respectively by any admissible and smooth enough velocity test function \boldsymbol{w} and pressure test function q, and then by integrating by part the term $\boldsymbol{w} \cdot \nabla p$. The natural enforcement of the capillary stress results from this integration by parts:

$$\begin{split} < \boldsymbol{w}, \nabla p >_{\Omega} &= < \boldsymbol{w}, \nabla p >_{\Omega_l} + < \boldsymbol{w}, \nabla p >_{\Omega_a} \\ &= - < \nabla \cdot \boldsymbol{w}, p >_{\Omega} + < \boldsymbol{w} \cdot \boldsymbol{n}, [p] >_{\Gamma_{l/a}} + < \boldsymbol{w} \cdot \boldsymbol{n}, p >_{\partial \Omega_N} \\ &= - < \nabla \cdot \boldsymbol{w}, p >_{\Omega} + < \boldsymbol{w} \cdot \boldsymbol{n}, \boldsymbol{n} \cdot \boldsymbol{\sigma}_{cap} \cdot \boldsymbol{n} >_{\Gamma_{l/a}} + < \boldsymbol{w} \cdot \boldsymbol{n}, p_0 >_{\partial \Omega_N} \end{split}$$

where, for a bounded region R, the bilinear form $\langle .,. \rangle_R$ denotes the $L^2(R)^n$ inner-product (n = 1 if a and b are scalars, n = d if they are vectors): $\langle a, b \rangle_R = \int_R a \cdot b \, dR$, for $a \text{ and } b \text{ in } L^2(R)^n$ the classical Lebesgues functional space,

$$L^{2}(R) = \{q: R \to \mathbb{R} \mid \int_{R} q^{2} dR < \infty\}$$

In order to complete the functional setting associated with the weak Darcy's equations, the Sobolev space $H(\nabla \cdot, \Omega)$ is also introduced:

$$H(\nabla \cdot, \Omega) = \{ \boldsymbol{u} \in L^2(\Omega)^d \mid \nabla \cdot \boldsymbol{u} \in L^2(\Omega) \}$$

Finally, the dual formulation of the mixed Darcy system (Eq. 1) - (Eq. 2) - (Eq. 5) - (Eq. 6) reads: Find $(\boldsymbol{v}, p) \in H(\nabla \cdot, \Omega) \times L^2(\Omega)$, with $\boldsymbol{v} \cdot \boldsymbol{n} = v_0$ on $\partial \Omega_D$, such that

$$< \mu \mathbf{K}^{-1} \mathbf{v}, \mathbf{w} >_{\Omega} - < \nabla \cdot \mathbf{w}, p >_{\Omega} = < \mathbf{f}, \mathbf{w} >_{\Omega} + < \mathbf{w} \cdot \mathbf{n}, p_0 >_{\partial \Omega_N} + < \mathbf{w} \cdot \mathbf{n}, \mathbf{n} \cdot \boldsymbol{\sigma}_{cap} \cdot \mathbf{n} >_{\Gamma_{l/a}}$$
(9)
$$< \nabla \cdot \mathbf{v}, q >_{\Omega} = < h, q >_{\Omega}$$
(10)

¹⁷³ $\forall (\boldsymbol{w}, q) \in H(\nabla \cdot, \Omega) \times L^2(\Omega), \text{ with } \boldsymbol{w} \cdot \boldsymbol{n} = 0 \text{ on } \partial\Omega_D, \text{ and } \mu = \mu_i \text{ in } \Omega_i.$

174 2.3. Stabilized FE formulation

The computational domain Ω is discretized by using a mesh made up of 175 triangles in 2D or tetrahedrons in 3D. Let Ω_h be this discretized domain. The 176 velocity \boldsymbol{v} and the pressure p are approximated by \boldsymbol{v}_h and p_h , which are both 177 continuous and piecewise linear functions (P1/P1 approximation). However, 178 such an approximation is not stable [8, 3, 29] according to Ladysenskaya-179 Brezzi-Babuška theory. In this work, this difficulty is overcome by using a 180 Variational Multi-Scale (VMS) technique [8, 40] consisting in adding some 181 stabilization terms to the Galerkin formulation. More precisely, the velocity 182 and pressure functional spaces, $\mathcal{V} \equiv H(\nabla \cdot, \Omega)$ and $\mathcal{P} \equiv L^2(\Omega)$ are split as 183

$$\mathcal{V} = \mathcal{V}_h \otimes \mathcal{V}' \quad ext{ and } \quad \mathcal{P} = \mathcal{P}_h \otimes \mathcal{P}'$$

where \mathcal{V}_h and \mathcal{P}_h are the velocity and pressure finite element spaces and \mathcal{V}' and \mathcal{P}' are the so-called subgrid or unresolvable scale spaces of velocity and pressure. Following this approach, the solution (\boldsymbol{v}, p) of the variational problem (Eq. 9)-(Eq. 10), as well as the test functions (\boldsymbol{w}, q) are divided as

$$oldsymbol{v} = oldsymbol{v}_h + oldsymbol{v}', \quad p = p_h + p'$$

 $oldsymbol{w} = oldsymbol{w}_h + oldsymbol{w}', \quad q = q_h + q'$

Subsequently, the variational problem is broken down into a problem at 188 the resolvable scale, the finite element problem, and a subgrid scale problem, 189 which cannot be explicitly solved. Consequently, the strategy of VMS meth-190 ods consists in approximating the effects of the subgrid scale onto the finite 191 element scale, leading to additional terms in the finite element formulation. 192 In this work, the Algebraic SubGrid Scale (ASGS) technique is used, a sub-193 type of VMS method developed by Badia and Codina in [8, 9, 25, 37]. The 194 subgrid terms are expressed as a function of the finite element residual such 195 that, on a mesh element e196

$$\boldsymbol{v}'_{|e} \approx -\boldsymbol{\tau_u}^e (\mu \boldsymbol{K}^{-1} \boldsymbol{v_h} + \nabla p_h - \boldsymbol{f})_{|e}$$
 (11)

$$p'_{|e} \approx -\tau_p^e (\nabla \cdot \boldsymbol{v_h} - h)_{|e}$$
 (12)

where τ_u^e and τ_p^e are stabilization parameters (Eq. 13) on the element *e*. They depend on the mesh size h_e , the geometry (through L_0 , a characteristic length of the domain Ω), the fluid viscosity, the porous medium permeability and the stabilization coefficients c_u and c_p (in this work, $c_u = c_p = 1$). As we are using P1/P1 approximation, these two parameters are expressed as [2, 8]:

$$\boldsymbol{\tau_u}^e = \frac{h_e \boldsymbol{K}_{|e}}{c_u L_0 \mu_{|e}}, \quad \boldsymbol{\tau_p}^e = \frac{\mu_{|e} c_p L_0 h_e}{K_{m|e}}$$
(13)

with K_m^e an equivalent permeability, chosen as [15]

$$K_m^e = \frac{1}{d} trace(\boldsymbol{K}_{|e})$$

Including the subgrid scale effects, the discrete FE system reads: Find $(\boldsymbol{v}_{\boldsymbol{h}}, p_{h}) \in \mathcal{V}_{h} \times \mathcal{P}_{h}$, with $\boldsymbol{v}_{\boldsymbol{h}} \cdot \boldsymbol{n} = v_{0}$ on $\partial \Omega_{hD}$, such that

$$<\mu \mathbf{K}^{-1} \mathbf{v}_{h}, \mathbf{w}_{h} >_{\Omega_{h}} - <\nabla \cdot \mathbf{w}_{h}, p_{h} >_{\Omega_{h}} - <\nabla \cdot \mathbf{v}_{h}, q_{h} >_{\Omega_{h}} + \sum_{e} \tau_{p}^{e} <\nabla \cdot \mathbf{v}_{h}, \nabla \cdot \mathbf{w}_{h} >_{e} + \sum_{e} \tau_{u}^{e} < -\mu \mathbf{K}^{-1} \mathbf{v}_{h} - \nabla p_{h}, \mu \mathbf{K}^{-1} \mathbf{w}_{h} + \nabla q_{h} >_{e} = <\mathbf{w}_{h} \cdot \mathbf{n}, \mathbf{n} \cdot \boldsymbol{\sigma}_{cap} \cdot \mathbf{n} >_{\Gamma_{hl/a}} + <\mathbf{w}_{h} \cdot \mathbf{n}, p_{0} >_{\partial\Omega_{hN}} + <\mathbf{f}, \mathbf{w}_{h} >_{\Omega_{h}} + _{\Omega_{h}}$$
⁽¹⁴⁾
$$+ \sum_{e} \tau_{u}^{e} <\mathbf{f}, \mu \mathbf{K}^{-1} \mathbf{w}_{h} + \nabla q_{h} >_{e} + \sum_{e} \tau_{p}^{e} _{e}$$

²⁰⁴ $\forall (\boldsymbol{w}_h, q_h) \in \mathcal{V}_h \times \mathcal{P}_h$ with $\boldsymbol{w}_h \cdot \boldsymbol{n} = 0$ on $\partial \Omega_{hD}$. In this formulation, \sum_{e} ²⁰⁵ stands for the summation over all the mesh elements e.

Note that when the porous medium is assumed to be isotropic, previous formulation (Eq. 14) can be slightly simplified, since the permeability K, the stabilization parameter τ_u and the capillary stress become scalar.

209 2.4. Pressure discontinuity

The capillary stress generates a pressure discontinuity at the liquid/air 210 interface. Moreover, for two different liquids the jump of viscosity across 211 this same interface leads to a discontinuity of the pressure gradient. In a FE 212 framework, these two kinds of discontinuities represent a numerical difficulty 213 to be dealt with. A first approach found in the literature consists in circum-214 venting the discontinuity by considering a smooth transition area around 215 the interface [16]. The performance of this method depends strongly on the 216 smoothing function, on the transition region thickness, and consequently on 217 the local mesh size [27]. Since no mesh adaptation strategy is used in this 218 work, the liquid-air interface will be identified by a continuous set of seg-219 ments (2D) or triangles (3D) crossing the mesh elements [15, 61] and built 220 locally thanks to the level-set front-capturing method described in Section 3. 221

This approach allows to integrate the capillary term into (Eq. 14) directly on a segment or triangle, using one integration point if the capillary stress is piecewise constant. Moreover, additional integration points are considered in the elements *e* crossed by the interface, in order to evaluate accurately the term $\langle \mu K^{-1} v_h, w_h \rangle_e$. Thus, in the 2D configuration (Fig. 2), assuming that both viscosities of the liquid and air are constant, 3 integrations points are used in each sub-element deriving from the element split.

However, such a split is not sufficient to ensure the accurate capture of 229 the pressure and pressure gradient discontinuities. Especially, continuous 230 and piecewise linear approximation of the pressure, piecewise linear approx-231 imation of the interface, give rise to the parasitic current phenomenon (even 232 if the curvature is not involved in the equations), which consists in spu-233 rious oscillations of the velocity, possibly deteriorating the interface [33]. 234 Here again, several options are available in the literature to reduce these 235 oscillations [23, 30, 55]. In particular, an enrichment of the pressure space 236 [6, 20, 26, 42, 43, 44] can be set up, locally in the elements crossed by the 237 fluid front. This work considers the pressure enrichment developed by R. 238 Ausas et al. [6]. Originally introduced to deal with discontinuities involved 239 in Navier-Stokes equations, this technique is applied here to Darcy's equa-240 tions. This consists in adding, in the elements crossed by the interface, the 241 two discontinuous shape functions M_1 and M_2 described in Fig. 2 and derived 242 243 as following:

$$M_1(\boldsymbol{x}) = (1 - S(\boldsymbol{x}))\chi^l(\boldsymbol{x})$$
(15)

$$M_2(\boldsymbol{x}) = S(\boldsymbol{x})\chi^a(\boldsymbol{x}) \tag{16}$$

244 with

$$S(\boldsymbol{x}) = \sum_{J \in \mathcal{J}^a} N_J(\boldsymbol{x}) \tag{17}$$

where N_J is the usual linear shape function associated with node J, χ^l is equal to 1 in the liquid region, to 0 elsewhere, and $\chi^a = 1 - \chi^l$. The set \mathcal{J}^a corresponds to the element nodes being in Ω_a .

In such elements, the pressure field p_h is expressed as

$$p_h(\boldsymbol{x}) = \sum_J P_J N_J(\boldsymbol{x}) + C_1 M_1(\boldsymbol{x}) + C_2 M_2(\boldsymbol{x})$$
(18)

where P_J are the degree of freedom associated with the element vertices J, while C_1 and C_2 are those associated with the discontinuous shape functions



Figure 2: 2D local pressure enrichment and surface reconstruction (inspired of [6]).

 M_1 and M_2 . However, since C_1 and C_2 are defined per element, they can be eliminated by static condensation, at the elementary level, prior to the final assembly. Therefore, the main advantage of combining the interface reconstruction and local pressure enrichment is that the discontinuity generated by the capillary stress is treated without increasing the number of final degrees of freedom and affecting the computation time.

²⁵⁷ 3. Fluid front capturing: level-set method

The moving flow front is captured by a level-set method [57, 69], consisting in choosing a continuous function, the so-called level-set function, $\psi(\boldsymbol{x},t): \Omega \times \mathbb{R}^+ \to \mathbb{R}$, negative in Ω_l and positive in Ω_a . Consequently, the interface $\Gamma_{l/a}$ is implicitly described as the zero-isovalue of function ψ :

$$\Gamma_{l/a}(t) = \{ \boldsymbol{x} \in \Omega \mid \psi(\boldsymbol{x}, t) = 0 \}$$
(19)

where t denotes the time variable. Note that the gradient of ψ allows the computation of the normal vector normal at the interface.

Assuming the flow velocity \boldsymbol{v} , defined both in Ω_l and Ω_a , known at each instant $t \in [0, T]$ (T is the final time of the simulation), the level-set function is then convected according to the hyperbolic equation (Eq. 20):

$$\frac{\partial \psi}{\partial t} + \boldsymbol{v} \cdot \nabla \psi = 0 \quad \forall (\boldsymbol{x}, t) \in \Omega \times [0, T]$$
(20)

$$\psi(\boldsymbol{x}, t=0) = \psi_0 \quad \forall \boldsymbol{x} \in \Omega \tag{21}$$

$$\psi(\boldsymbol{x},t) = g(\boldsymbol{x},t) \quad \forall (\boldsymbol{x},t) \in \partial \Omega^{-} \times [0,T]$$
(22)

where $g(\boldsymbol{x},t)$ (Eq. 22) corresponds to the value of ψ to be imposed on the incoming boundary $\partial \Omega^-$ (Fig. 1)

$$\partial \Omega^{-}(t) = \{ \boldsymbol{x} \in \partial \Omega \mid \boldsymbol{v}(\boldsymbol{x}, t) \cdot \boldsymbol{n} < 0 \},\$$

while (Eq. 21) states for the initial condition at t = 0.

270 3.1. SUPG formulation

Transport equation (Eq. 20) is solved by a FE technique, using the same 271 mesh as for Darcy's equations. The variational formulation is first obtained 272 by multiplying (Eq. 20) by any admissible and smooth enough test function 273 and integrating the product over Ω . The time interval is discretized by a set 274 of points $0 = t_0 < t_1 < \cdots < t_n < t_{n+1} < \cdots < t_N$, and a finite difference 275 scheme in time, the implicit Crank-Nicholson scheme, is then applied to 276 (Eq. 20). It results that at each instant t_n , $\psi(\cdot, t_n)$ is approximated by 277 $\psi_h(\cdot, t_n)$ a continuous piecewise linear function. However, the usual Galerkin 278 approach is known to be not stable for hyperbolic equations. This issue can 279 be avoided by considering a Streamline Upwind Petrov-Galerkin (SUPG) 280 method, introduced by Hughes in [19], and consisting in taking the test 281 functions in a space different of the shape functions. More precisely, shape 282 functions are still the nodal functions N_J already introduced, but the test 283 functions, denoted N_J^* , are now defined on a mesh element e, as 284

$$N_J^* = N_J + \tau^e \boldsymbol{v} \cdot \boldsymbol{\nabla} N_J$$

where the stabilization parameter τ^e is chosen as

$$\tau^e = \frac{1}{2} \frac{h_e}{v_e}$$

with h_e the size of element e and v_e the norm of the average velocity in e.

This modification of the test functions adds, in a consistent way, an upwind artificial diffusion term stabilizing the FE formulation, at least as long as the convective term remains under control.

290 3.2. Filtered level-set

For the level-set procedure described below to be effective, the level-set function ψ has to be initialized with a specific expression. Let $d_0(\boldsymbol{x})$ denote the signed distance function from point \boldsymbol{x} to the initial liquid-air interface $\Gamma_{la}(0)$. The initial expression of the level-set function, involved in the initial condition (Eq. 21) is then chosen as

$$\psi_0(\boldsymbol{x}) = \varepsilon \tanh\left(\frac{d_0(\boldsymbol{x})}{\varepsilon}\right)$$
 (23)

where ε can be viewed as the thickness of the interface. In practice: $\varepsilon = 3h_e$.

Outside a narrow band around the interface, ψ_0 quickly tends towards 297 the constant values $\pm \varepsilon$. Therefore, condition (Eq. 22) to be prescribed on 298 the inflow boundary can easily be enforced. Additionally, within this tiny 299 band close to the interface, ψ_0 is equal, in the first order, to the distance func-300 tion d_0 . A distance function have, by definition, a unit gradient: $\|\nabla d_0\| = 1$. 301 This property ensures the "control" of the convection term in transport equa-302 tion (Eq. 20) and thus the efficiency of the SUPG stabilization. However, 303 the initial "tanh-like" shape (Eq. 23) is not preserved under the transport 304 of ψ with the Darcy's velocity field \boldsymbol{v} . That is why, as this velocity varies 305 abruptly (but continuously) through the liquid-air interface, steep gradients 306 of level-set function will develop in its vicinity, and the SUPG stabilization 307 will fail. This problem is avoided by periodically reinitializing the level-set 308 function: the zero-isovalue is preserved, while the tanh property is applied 309 elsewhere. Based on the relation $d \tanh(x)/dx = 1 - \tanh^2(x)$, function ψ is 310 of the form (Eq. 23) if 311

$$\|\nabla\psi\| = \left|1 - \left(\frac{\psi}{\varepsilon}\right)^2\right| \tag{24}$$

At a given time t_n , the reinitialization step consists in solving iteratively the Hamilton-Jacobi equation

$$\frac{\partial \tilde{\psi}}{\partial \tau} + sgn(\tilde{\psi}) \left(\left\| \nabla \tilde{\psi} \right\| - \left| 1 - \left(\frac{\tilde{\psi}}{\varepsilon} \right)^2 \right| \right) = 0$$
 (25)

$$\tilde{\psi}(x,\tau=0) = \psi(x,t_n) \tag{26}$$

³¹⁴ until reaching the steady state, *i.e.* $\partial \tilde{\psi} / \partial \tau = 0$, corresponding consequently ³¹⁵ to the property (Eq. 24). This state gives the reinitialized level-set func-³¹⁶ tion. In practice, only a few increments (3 in our simulations) are necessary ³¹⁷ to recover the unit gradient property in the narrow band around the inter-³¹⁸ face. In (Eq. 25), τ is a time-like variable, and sgn is the regularized sign ³¹⁹ function [57]

$$sgn(\psi) = \frac{\psi}{\sqrt{\psi^2 + \|\nabla\psi\|^2 h_e^2}}$$
(27)

Note that, classically, Hamilton-Jacobi equation (Eq. 25) can be considered as a transport equation with a right-hand-side, and is then solved in the same way as the level-set convection equation (Eq. 20). The reinitialization velocity is equal to $sgn(\psi) \frac{\nabla \tilde{\psi}}{\|\nabla \tilde{\psi}\|}$, while the non-linear terms are explicitly evaluated at the previous iteration.

325 3.3. Time-stepping strategy

The time-stepping strategy consists, for a given time increment, in solving Darcy's equations, then updating the flow front position by solving the levelset transport equation using the Darcy's velocity, and moving on to the next time increment. To sum up, the algorithm coupling Darcy's and level-set problems is as following:

Algorithm 1 Staggered algorithm for Darcy's and level-set problems
Require: $\psi(\boldsymbol{x}, t = 0) = \psi_0$ the initial value for the level set function
while $0 < t^{n+1} < T$ do
1- Fluid problem:
Find $(\boldsymbol{v}_h, p_h) \in \mathcal{V}_h \times \mathcal{P}_h$ by solving Darcy's equations (Eq. 14)
2- Flow front problem:
Find ψ_h by solving the level-set equations
3- Reinitialization problem:
Repeat 3 times: Solving Hamilton-Jacobi's equations (Eq. 25)
end while

331 4. Convergence analysis

The FE model presented in the previous section has been implemented 332 in the FE software Z-set [68]. The efficiency of the implementation, as well 333 as the accuracy of the approach, are evaluated by an error analysis based 334 on the Method of Manufactured Solutions (MMS) [64]. This consists in 335 selecting velocity and pressure fields that satisfy Darcy's equations (Eq. 1)-336 (Eq. 2) and calculating the corresponding right-hand-side terms that are then 337 prescribed in the FE problem. Performance of the implementation measures 338 the capability of reproducing the initial fields. 339

The 2D-computational domain is the unit square $\Omega = [0, 1] \times [0, 1]$. The analytical pressure field is defined as

$$p(x,y) = \begin{cases} \sin(2\pi x)\sin(2\pi y) & \text{for } y < \frac{1}{2} \\ \sin(2\pi x)\sin(2\pi y) + \sigma_{cap} & \text{for } y > \frac{1}{2} \end{cases}$$
(28)

342 with σ_{cap} the scalar value of the capillary stress in the isotropic case. 343

Replacing (Eq. 28) inside Darcy's equation (Eq. 1) gives the components of the velocity, v_x and v_y

$$v_x = \frac{K}{\mu} 2\pi \cos(2\pi x) \sin(2\pi y)$$

$$v_y = \frac{K}{\mu} 2\pi \sin(2\pi x) \cos(2\pi y)$$
(29)

These fields (Eq. 28) and (Eq. 29) satisfy Darcy's system (Eq. 1)-(Eq. 347 2) with the term h taken as

$$h = \nabla \cdot \boldsymbol{v} = \frac{K}{\mu} 8\pi^2 \sin(2\pi x) \sin(2\pi y) \tag{30}$$

Only one type of boundary condition is considered here, the Dirichlet one (Eq. 6): $\boldsymbol{v} \cdot \boldsymbol{n} = v_0 = 0$ on $\partial \Omega$. Thus, the so-called compatibility condition is fulfilled, that is

$$\int_{\Omega} \nabla \cdot \boldsymbol{v} \, d\Omega = \int_{\partial \Omega} \boldsymbol{v} \cdot \boldsymbol{n} \, d\Gamma = \int_{\partial \Omega} v_0 \, d\Gamma = 0$$

In all the simulations shown in this section, the pressure jump across the 351 line $\{y = \frac{1}{2}\}$ is equal to 1, $[p] = \sigma_{cap} = 1$, while the ratio K/μ is also unit 352 (isotropic case). Pressure and velocity obtained by the FE strategy described 353 before are plotted in a 3D-representation in Fig. 3 using an unstructured mesh 354 of element size $h_e = 0.0125$. The pressure discontinuity is well-captured, 355 without apparent oscillations of pressure. This is qualitatively confirmed 356 in Fig. 4, where the computed pressure is satisfactorily compared to the 357 analytical one along two lines, $\{x = \frac{1}{4}\}$ and $\{x = \frac{1}{2}\}$. 358

Next, a quantitative analysis of the error made on velocity and pressure is performed by considering 4 structured meshes of size, respectively, $h_e = 1/20$, 1/40, 1/80 and 1/160. On each of these meshes, pressure error is calculated with the usual L^2 -norm denoted $\|\cdot\|_{L^2}$, while velocity error is estimated both in L^2 -norm and in $H(\nabla \cdot)$ -norm denoted $\|\cdot\|_{H(\nabla \cdot)}$ (Eq. 31):

$$\|u\|_{L^{2}} = \left(\int_{\Omega} u^{2} d\Omega\right)^{\frac{1}{2}}, \quad \|u\|_{H(\nabla \cdot)} = \left(\|u\|_{L^{2}}^{2} + \|\nabla \cdot u\|_{L^{2}}^{2}\right)^{\frac{1}{2}}$$
(31)



Figure 3: 3D-plot of pressure and velocity fields (Eq. 28)-(Eq. 29), obtained by the FE solution.

The theoretical convergence rates, without discontinuity of pressure, are 364 2 in L²-norm both for the pressure and velocity, and 1 in the $H(\nabla \cdot)$ -norm 365 for the velocity [3, 8, 53]. Therefore, three different cases are proposed here: 366 a continuous case, corresponding to $\sigma_{cap} = 0$, in order to assess the Darcy's 367 solver in a classical situation and have a reference situation; two discontinuous 368 cases with $\sigma_{cap} = 1$ as mentioned above, but one without pressure enrichment 369 of Section 2.4, and one with this technique. Results are summarized in 370 Tables 1-2-3 and Fig. 5. 371

Mesh	h_e	$ p - p_h _{L^2}$	$\ oldsymbol{v}-oldsymbol{v}_{oldsymbol{h}}\ _{L^2}$	$\ oldsymbol{v}-oldsymbol{v}_{oldsymbol{h}}\ _{H(abla\cdot)}$
20×20	0.05	0.0213803	0.0839136	6.76775
40×40	0.025	0.00389416	0.0188836	3.3258
80×80	0.0125	0.000714466	0.00416498	1.61289
160×160	$6.25 \cdot 10^{-3}$	0.000148175	0.00100372	0.812385

Table 1: Error in the L^2 -norm for the pressure, and both the L^2 -norm and $H(\nabla \cdot)$ -norm for the velocity. Case with a continuous pressure.

We observe that without pressure discontinuity, the convergence rate obtained is in agreement with the optimal one, since the rate is slightly higher



Figure 4: Comparison between analytical pressure field (Eq. 28) (continuous line) and results of simulation (dots), along the lines $x = \frac{1}{4}$ and $x = \frac{1}{2}$.

Mesh	h_e	$ p - p_h _{L^2}$	$\ oldsymbol{v}-oldsymbol{v}_{oldsymbol{h}}\ _{L^2}$	$\ oldsymbol{v}-oldsymbol{v}_{oldsymbol{h}}\ _{H(abla\cdot)}$
20×20	0.05	0.0889837	0.109918	7.2842
40×40	0.025	0.0475362	0.0203123	3.33272
80×80	0.0125	0.0367567	0.00888546	2.1614
160×160	$6.25 \cdot 10^{-3}$	0.0231142	0.00189264	0.906225

Table 2: Error in the L^2 -norm for the pressure, and both the L^2 -norm and $H(\nabla \cdot)$ -norm for the velocity. Case with pressure discontinuity and no pressure enrichment.

Mesh	h_e	$ p - p_h _{L^2}$	$\ oldsymbol{v}-oldsymbol{v}_{oldsymbol{h}}\ _{L^2}$	$\ oldsymbol{v}-oldsymbol{v}_{oldsymbol{h}}\ _{H(abla\cdot)}$
20×20	0.05	0.0615517	0.0933167	6.94098
40×40	0.025	0.0211956	0.0247864	3.63211
80×80	0.0125	0.00571188	0.00801094	1.9443
160×160	$6.25 \cdot 10^{-3}$	0.00237119	0.00323276	1.55267

Table 3: Error in the L^2 -norm for the pressure, and both the L^2 -norm and $H(\nabla \cdot)$ -norm for the velocity. Case with pressure discontinuity and local pressure enrichment.

than 2 for both pressure and velocity in L^2 -norm, and equal to 1 for the velocity in $H(\nabla \cdot)$ -norm. As expected, the two cases with pressure discontinuity let show lower convergence rates compared to the continuous case, especially for the pressure. Without enrichment strategy, a sub-optimal convergence rate





Figure 5: Error analysis: continuous case (a), discontinuous case without pressure enrichment (b) and discontinuous case with local pressure enrichment (c).

of 0.62 is obtained for pressure, corresponding to approximatively 1/3 of the 378 theoretical order predicted for the continuous case. However, when enriching 379 locally the pressure space, this rate is greatly improved, since jumping up to 380 80% of this same theoretical order, with the value of 1.6. Note that, with this 381 same enrichment technique, but considered in the context of Navier-Stokes 382 equations, Ausas and co-authors [6] obtained a pressure convergence rate 383 equal to 75% of the one predicted with a continuous pressure. Hence, we can 384 conclude that the numerical approach presented below allow us to describe 385 with accuracy the pressure discontinuity when solving the Darcy's equations 386 with a capillary stress applied on the moving flow front. 387

5. Numerical applications 388

This section assesses and demonstrates the performance of our numerical 389 model in realistic contexts. First, numerical simulations of flows through 390 porous media with a very low permeability are carried out, and the interest 391 of local pressure enrichment is highlighted. Next, capillary wicking simula-392 tions are performed and the results are compared with experimental studies. 393 Finally, a first approach of flows in 3D orthotropic materials is provided 394 through the simulation of the filling stage of a T-stiffener during a LRI pro-395 cess. It is also used to demonstrate that simply shifting the pressure at a 396 boundary condition by the capillary stress may hold for UD cases, but is not 397 satisfactory in terms of filling scenario and results for general 3D cases. 398

5.1. Ascending capillary flow and realistic parameters 399

The numerical strategy is assessed by simulating a unidirectional flow 400 in a porous medium with realistic properties, in terms of permeabilities, 401 viscosities and capillary stresses, in the context of composite materials man-402 ufacturing. Thus, the computational domain Ω is a square of 1 meter side. 403 The isotropic permeability K is equal to $3.0 \times 10^{-13} \text{m}^2$, while the isotropic 404 capillary stress, applied on the interface $\Gamma_{l/a} \equiv \{y = h = 1/2\}$, is of 32×10^3 405 Pa. Viscosities are $\mu_l = 10^{-3} Pa.s$ and $\mu_a = 10^{-5} Pa.s$ This pressure is the 406 only driving force, since boundary conditions on both planes $\{y = 0\}$ and 407 $\{y = 1\}$ are set to the atmospheric pressure. The remaining boundaries are 408 considered as impervious walls, thus the $\boldsymbol{v} \cdot \boldsymbol{n} = 0$ condition is applied on the 409 vertical edges of the domain, $\{x = 0\}$ and $\{x = 1\}$. All numerical values of 410 material properties and boundary conditions are sum up in Fig. 6. 411

In the case of a unidirectional flow, the analytical solution of Darcy's 412 equations is quite simple to determine. Indeed, the pressure is piecewise 413 linear, while the velocity is constant. With the notations introduced in Fig. 6, 414 the pressure and velocity fields can be written as 415

$$p(x,y) = \mu_l \frac{p_1 - p_0 - \sigma_{cap}}{h\mu_l + (1-h)\mu_a} y + p_0 \quad \text{in } \Omega_l$$

$$p(x,y) = \mu_a \frac{1}{h\mu_l + (1-h)\mu_a} (y-1) + p_1 \text{ in } \Omega_a$$

$$v_x(x,y) = 0 \qquad \text{in } \Omega$$

$$v_y(x,y) = -K \frac{p_1 - p_0 - \sigma_{cap}}{h\mu_l + (1-h)\mu_l} \qquad \text{in } \Omega$$
(32)

$$v_y(x,y) = -K \frac{p_1 - p_0 - \sigma_{cap}}{h\mu_l + (1-h)\mu_a}$$
 in Ω

$$p(y = 1) = p_1 = 10^5 \text{ Pa} \quad (1, 1)$$

$$\Omega_a, K = 3.0 \times 10^{-13} \text{m}^2$$

$$\mu_a = 10^{-5} \text{ Pa.s}$$

$$\sigma_{cap} = 32 \times 10^3 Pa$$

$$y = \frac{1}{2}$$

$$\Omega_l, K = 3.0 \times 10^{-13} \text{m}^2$$

$$\mu_l = 10^{-3} \text{ Pa.s}$$

$$(0, 0) \quad x \quad p(y = 0) = p_0 = 10^5 \text{ Pa}$$

Figure 6: Material properties and boundary conditions used in the numerical simulation of unidirectional flow.

The velocity and pressure fields computed with a structured mesh of size 416 $h_e = \frac{1}{50}$ (4800 triangular elements corresponding to 2499 nodes) and an 417 interface crossing the elements, are given in Fig. 7 and 8. Two cases are 418 considered: without and with the pressure enrichment introduced in section 419 2.4. In the first case, the pressure jump is not well-captured at the interface 420 (Fig. 7(b)), resulting in a spurious velocity around this interface (Fig. 7(a)). 421 On the contrary, the discontinuity of the pressure field is accurately computed 422 with the enrichment (Fig. 8(b)) leading to a uniform velocity field as expected 423 by Equation (Eq. 32) (Fig. 8(a)). Numerical and analytical values of the 424 velocity are identical, and equal to $1.901 \times 10^{-5} \text{ m.s}^{-1}$ in norm. This also 425 proves the accuracy of the pressure description. To complete this analysis, 426 analytical and numerical pressures have been plotted on the line $\{x = \frac{1}{2}\}$ 427 in Fig. 9, for different structured meshes. Again, no pressure oscillation is 428 observed. 429



Figure 7: Velocity and pressure fields obtained without pressure enrichment



Figure 8: Velocity and pressure fields obtained with pressure enrichment



Figure 9: Comparison between analytical and numerical pressures, plotted along the line $\{x = \frac{1}{2}\}$, for different mesh sizes h_e .

430 5.2. Capillary wicking

In this section, simulation of wicking in carbon reinforcements is confronted to experimental data [62, 63]. The only driving force is thus due to capillary effects.

434 5.2.1. Experimental approach

Pucci *et al.* [62] proposed an experimental procedure to determine the scalar capillary stress σ_{cap} in the three main directions of a unidirectional (UD) carbon fabric (Fig. 11). On the one hand, for a given direction, the mass of water in the fabric m(t) is recorded over time using a tensiometer. Wicking is commonly described by a modified Washburn equation [74] for porous media relating mass and time

$$m^{2}(t) = \left[\frac{(c\bar{r})\phi^{2}(\pi R^{2})^{2}}{2}\right]\frac{\rho_{l}^{2}\gamma\cos\theta_{a}}{\mu_{l}}t$$
(33)

where c is a constant accounting for the tortuous path of liquid in the equivalent capillary tube arrangement of mean radius \bar{r} . ϕ is the porosity and R the inner radius of the cylindrical sample holder. The first term in square brackets finally represents a geometric factor of the porous medium. ρ_l and μ_l are, respectively, the liquid (water) density and its viscosity. θ_a is the apparent mean advancing contact angle during the capillary rise and γ_l the liquid surface tension.

⁴⁴⁸ On the other hand, from Darcy's equation applied to a unidirectional ⁴⁴⁹ flow (Eq. 32) following the assumption of spontaneous impregnation under ⁴⁵⁰ the effect of capillary stress σ_{cap} (Fig. 10) the square of the water height $h^2(t)$ ⁴⁵¹ (see Fig. 11) can be expressed as a function of time

$$h^2(t) = \frac{2K\sigma_{cap}}{\mu_l\phi}t\tag{34}$$

This expression is easily obtained from the last equation of the analytical model (Eq. 32), considering that $v_y = \phi \frac{dh}{dt}$, $p_1 = p_0$, $\mu_a = 0$ and integrating it with respect to time.

Taking into account the cylindrical shape of radius R, the mass gain can be related to the height by

$$m^{2}(t) = h^{2}(t)\phi^{2}\rho_{l}^{2}(\pi R^{2})^{2}$$
(35)

⁴⁵⁷ Considering the equivalence between Eq. 33 and Eq. 35, it is then possible ⁴⁵⁸ to describe capillary stress σ_{cap} for a given permeability K [62].



Figure 10: Capillary wicking in a cylindric quasi-UD carbon reinforcement according to Darcy law for the three principal directions of the medium.

459 5.2.2. Numerical simulations and results

Three 2D numerical simulations of wicking of water into a cylindrical 460 quasi-UD carbon reinforcement have been carried out. For each simulation, 461 the fabric is oriented in one of the directions x, y or z. Note that consequently, 462 each of these simulations is reduced to an isotropic case. The computational 463 domain, the $2R \times H$ rectangle described in Fig. 11, is discretized with a 464 fixed mesh of 2,352 triangular elements and 1,250 nodes. The boundary 465 conditions prescribed for this simulation are a zero normal velocity on the 466 vertical sides and a pressure of 1 bar on the two other sides. 467

The definition of the orthotropic permeability tensor, orthotropic capillary stress tensor applied on the flow front and identified from experience, are given in Table 4, as well as the water and "air" viscosities, the water density, the porosity, and the dimensions 2R and H of the computational domain.

In order to have realistic simulations, the porosity has to be taken into account in Darcy's equations. This is achieved by substituting $v\phi$ for v Darcy's equations. From the position of the water height h(t) obtained by simulation, the corresponding water weight is calculated by Eq. 35 and compared (Fig. 12), with experimental data and the analytical expression given by Eq.

Permeabilities [49]	(m^2)
K_x	$3 \cdot 10^{-11}$
K_y	$1.5 \cdot 10^{-11}$
K_z	$3 \cdot 10^{-13}$
Capillary stress [62]	(kPa)
σ^x_{cap}	1.15 ± 0.30
σ^{y}_{cap}	0.51 ± 0.14
σ_{cap}^{z}	32.10 ± 11.60
Others	
μ_{water}	10^{-3} Pa.s
$ ho_{water}$	$10^3 { m kg.m^{-3}}$
μ_{air}	$10^{-5} { m Pa.s}$
ϕ	0.40
2R	$12 \mathrm{mm}$
Н	20 mm

Table 4: Capillary wicking parameters.



Figure 11: Geometrical parameters and boundary conditions of the capillary wicking.

478 34-35. It can be shown that numerical simulations and analytical expres-479 sion give comparable results. Therefore, the numerical simulations reflect 480 the experimental wicking. It confirms that the numerical model is correct 481 since wicking in each main directions were already correctly described by 482 the analytical model. However, the key point is that the proposed numerical 483 methodology simulates wicking, a transient phenomenon, here in an isotropic 484 context, but with realistic values of parameters involved in the model.



Figure 12: Comparison of the time dependent numerical and experimental weight in the x (a), y (b) and z (c) directions.

Parameters	Values	Units
K_t	10^{-14}	m^2
K_p	10^{-12}	m^2
σ^t_{cap}	0.3	bar
σ^p_{cap}	0.01	bar
μ_{resin}	0.1	Pa.s
μ_{air}	10^{-5}	Pa.s
ϕ	40%	
Δt	100	\mathbf{S}

Table 5: Inputs - LRI simulation.

485 5.3. Full 3D-simulations of LRI process filling stage

This section investigates the 3D-simulation of the filling stage of a Liquid 486 Resin Infusion (LRI) process [15, 22]. More precisely, the objective is to 487 evaluate the influence of capillary effects on the resin impregnation. The part 488 to be infused is the aeronautic-like stiffener shown in Fig. 13. Furthermore, 489 the preform is assumed to have an additional symmetry: two eigen-values 490 of the permeability tensor, as well as two eigen-values of the capillary stress 491 tensor are equal. Hence, Table 5 gives the values of the in-plane permeability 492 K_p , which is a hundred times larger than the transverse permeability K_t . As 493 capillary forces are more significant in less permeable media, the capillary 494 stress σ_{cap}^{t} in the transverse direction is higher than the value in the plan 495 σ^p_{cap} . These values, completed by the resin viscosity, the air viscosity, the 496 porosity and the time step Δt are also provided in Table 5. Figure 13 shows 497 the corresponding materials eigen-directions y^t and x^p on a cutting plane. 498 That is the transverse direction and the plane orthogonal to this direction in 499 three different areas, allowing to compute numerical values of permeabilities 500 and capillary stresses at each integration point of finite elements. Moreover, 501 the boundary conditions both in velocity and pressure are given. The resin 502 flow front is initialized as the plane $\{y = 0.5cm\}$. The flow is driven by 503 the difference of pressure between the "inlet" (plane $\{y = 0\}$) and the "top" 504 (plane $\{y = 12cm\}$) boundaries, equal to 10^5 Pa, and additionally by the 505 capillary stress on $\Gamma_{l/a}$ when this is taken into account. 506

Regarding the solution for 1D cases, one may question about a simple way of accounting for capillary effects by modifying the overall pressure gradient. Although an extension to 3D is not straightforward, in order to assess



Figure 13: Geometrical dimensions and boundary conditions of the stiffener.

also such a basic approach, 3 simulations for the T-stiffener were considered: 510 one carried out without capillary effects, one accounting for capillary effects 511 by modifying the overall pressure gradient through changing the boundary 512 condition in the plane y = 0: $\boldsymbol{n} \cdot \boldsymbol{\sigma} \cdot \boldsymbol{n} = 10^5 Pa \rightarrow \boldsymbol{n} \cdot \boldsymbol{\sigma} \cdot \boldsymbol{n} = 10^5 Pa + \sigma_{cap}^t$ 513 Modified BC-, and finally integrating orthotropic capillary stresses with the 514 proposed method - Discontinuous Pressure-. Figure 14 compares the flow 515 front position during infusion for the corresponding three simulations. As 516 expected, the part is filled more quickly when the capillary effects are taken 517 into account: 1h58 min with the Modified BC approach using the highest 518 capillary stress σ_{cap}^t , 2h 10min with our discontinuous pressure numerical 519 model and 3 hours without any capillary effects. One can verify that cap-520 illary effects will help the filling of the preform. Besides, the pressure and 521 fluid front kinetics resulting from the approaches integrating these effects 522 differ largely. 523

More precisely, the pressure computed from the 3 methods are presented in 524 Figure 15, for locations along a vertical line $\{0.01; y; 0.2\}$ as sketched in Fig-525 ure 14b. One can verify that the pressure profiles obtained without capillary 526 effects and with the Discontinuous Pressure method are quite close, showing 527 the ability of the latter method to capture properly the pressure field out of 528 the interface region (pressure gradient) while integrating locally the capillary 520 effects. As for the Modified BC, the obtained pressure profile is largely modi-530 fied. Also, it can be noticed that the Discontinuous Pressure approach yields 531



Figure 14: Numerical results - position of the fluid front during the filling of an orthotropic stiffener at different times

a pressure jump whose magnitude is a combination of both capillary stresses
in the transverse and plane directions. This method is intrinsically able to
account for the capillary stress orthotropy, and the corresponding flow front
follows the preform principal directions.

It can be concluded that with the Modified BC method, first the or-536 thotropic character of the capillary effects will not be accounted for by the 537 simple overall gradient correction, and second the pressure field will not be 538 discontinuous, opposite to the physics of two-phase flows. Consequently, a 539 finer analysis is not possible with this approach, and especially it will no 540 longer hold for a more exhaustive modelling approach relying on velocity 541 and fluid pressure fields. Conversely, the proposed approach with discontin-542 uous pressure will yield relevant pressure and velocity distributions which can 543 then be incorporated in more exhaustive approaches of the filling stage in-544 cluding solid-fluid mechanics couplings through the fluid pressure. Of course, 545



Figure 15: Comparison of the pressure fields with the *Modified BC* - *Discontinuous Pressure* methods and without capillary effects at t=3000s along a vertical line $\{0.01; y; 0.2\}$ plotted in Fig. 14b.

further experimental studies are required to validate and calibrate the numerical model for industrial-like structures. However, these results show that
capillary effects may have huge impact on the filling strategy for the out-ofautoclave processes targeted here.

550 6. Conclusion

In our macroscale configuration, the capillary action is represented by 551 capillary stresses, acting at the liquid/air interface by the mean of the nor-552 mal vector. This stress is weakly enforced in Darcy's equations as an input 553 parameter of the model and generates a pressure jump at the interface. These 554 equations are discretized using a stabilized mixed FE method, linear in both 555 velocity and pressure. The spurious velocities due to the pressure discontinu-556 ity are limited by using a local pressure enrichment technique. The numerical 557 model gives the expected convergence rates, both for velocity and pressure. 558 Besides, the 2D isotropic simulations of a capillary wicking of water inside 559 a carbon fabric show a good correlation between the numerical results and 560 the experimental data, as well as with the analytical model of Washburn's 561 equation. Finally, a full 3D and orthotropic case has been investigated: the 562 filling of an aeronautic part with a LRI process. This simulation enlightens 563 the influence of the capillary effects on the progress of the filling stage, and 564 demonstrates that the overall response is of highest importance, but also the 565

proper representation of the pressure discontinuity is mandatory for velocity
and pressure fields predictions to be used for solid-fluid mechanics couplings
for instance [15]. Further experimental studies are now required in order to
confirm this scenario.

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