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# An adaptive strategy for discontinuous Galerkin simulations of Richards' equation: Application to multi-materials dam wetting



Advance

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#### ABSTRACT

Numerical solution of Richards' equation remains challenging to get robust, accurate and cost-effective results, particularly for moving sharp wetting fronts. An adaptive strategy for both space and time is proposed to deal with 2D sharp wetting fronts associated with varying and possibly vanishing diffusivity caused by nonlinearity, heterogeneity and anisotropy. Adaptive time stepping makes nonlinear convergence reliable and backward difference formula provides high-order time scheme. Adaptive mesh refinement tracks wetting fronts with an *a posteriori* error indicator. The novelty of this paper consists in using this technique in combination with a weighted discontinuous Galerkin framework to better approximate steep wetting fronts by a discontinuity. The potential of the overall approach is shown through various examples including analytical and laboratory benchmarks and simulation of full-scale multi-materials dam wetting experiment.

# 1. Introduction

Predicting variably-saturated flows in porous media is a major issue for many fields in science and engineering. For example, such flows arise in soil physics, hydrogeology, environment, agriculture or oil industry for problems like subsurface contaminant transport, petroleum reservoir, water resources, surface ponding, etc.

In the present work, flows in variably-saturated porous media are described by Richards' equation. What makes Richards' equation very attractive is that it models the porous medium as a whole part including both saturated and unsaturated zones. Richards' equation is a nonlinear parabolic equation which can degenerate into an elliptic equation under complete saturation condition. Richards' equation has been extensively used for numerical simulations by the hydrogeology community, see e.g. (Thoms et al., 2006; Szymkiewicz, 2013). Despite this research effort, many numerical challenges remain for solving Richards' equation with some simulations being still unreliable and/or expensive (Farthing and Ogden, 2017; Zha et al., 2019). This numerical complexity prevented a more general use of Richards-based model for a number of applications. Indeed, the solution of Richards' equation involves sharp wetting fronts which evolve both in space and time and are difficult to resolve. Besides, the simulation must treat simultaneously unsaturated/saturated regions of parabolic/elliptic natures, several porous media of different hydraulic properties and possibly fast-changing boundary conditions.

These aspects can also lead to steep gradients acting like discontinuities. Considering the set of nonlinear functions for hydraulic properties, it is seldom possible to foresee the behaviour of Richards' equation and getting numerical solutions can be hard to achieve. Especially, numerical schemes often fail to converge or need fine discretization, making computation costly. Many numerical methods are dealing with Richards' equation to treat wetting fronts and heterogeneous soils in various contexts. Each of them shows advantages and drawbacks and that is the reason why research is still undergoing to push back the limits for the use of Richards' equation. For instance, to handle layered soils, some recent publications include a Kirchhoff integral transform-based method (Suk and Park, 2019), a transversal method of lines (Berardi et al., 2020), a domain decomposition technique with L-scheme linearization (Seus et al., 2018) and even upscaled models of Richards' equation for fractured porous media (Kumar et al., 2020).

In this paper, a discontinuous Galerkin (DG) method is chosen to solve Richards' equation. DG methods are based on a variational formulation in an element-wise fashion, sharing advantages both with finite elements and finite volumes methods. In particular, they are locally conservative which is crucial in fluid dynamics (Rivière, 2008). Moreover, the nature of DG formulation enables to work on non-conforming mesh and to change locally the degree of polynomial approximation. This is an important benefit since adaptive mesh refinement (AMR), the so-called  $\mathfrak{h}$ -adaptation, and high-order accuracy, the so-called *p*-adaptation, be-

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Received 19 October 2020; Received in revised form 23 February 2021; Accepted 19 March 2021 Available online 24 March 2021 0309-1708/© 2021 Elsevier Ltd. All rights reserved. come possible (Dolejší and Feistauer, 2015), promoting a growing use of DG methods for transport phenomena in porous media such as twophase flow problems (Klieber and Rivière, 2006; Bastian, 2014). However, DG methods remain sparsely applied to Richards' equation. In 2007, Li et al. solved Richards' equation in 1D with a local discontinuous Galerkin method (Li et al., 2007b; 2007a). Sochala discretized Richards' equation through a SIPG mixed DG formulation in 2008 (Sochala, 2008; Sochala et al., 2009). Recently, in 2019, Dolejší et al. (2019) proposed a space-time DG method for solving Richards' equation.

To reach robustness and accuracy, an adaptive strategy is developed in this paper. Richards' equation is known to be a stiff differential equation with difficult convergence (Paniconi and Putti, 1994; Lehmann and Ackerer, 1998; List and Radu, 2016) so the time discretization and the nonlinear solver have to be addressed carefully. Adaptive mesh refinement will be used to capture moving wetting fronts thanks to a posteriori estimation. h-Adaptation is employed for two-phase flow in porous media (Klieber and Rivière, 2006) but also for Richards' equation (Miller et al., 2006; Li et al., 2007a). Making the most of DG methods flexibility, adaptive mesh refinement is combined with a weighted discontinuous Galerkin (WDG) framework which allows discontinuity in the solution according to the nonlinear diffusivity. Following earlier works, this approach was formulated by Ern, Di Pietro and other collaborators (Ern et al., 2008; Di Pietro et al., 2008) and by Proft and Rivière under the name improved and adapted discontinuous Galerkin methods (Proft and Rivière, 2006; 2009). Application of such strategy for Richards' equation, combining adaptive mesh refinement, a posteriori estimation and WDG framework, is the main novelty of the present study.

The Richards' equation framework is first recalled in Section 2. In Section 3, Richards' equation discretization is presented through a DG framework and then solved by a nonlinear iterative process aiming to robustness. Section 4 is dedicated to the adaptive strategy proposed to improve solving of Richards' equation. Several numerical experiments are discussed in Section 5. In particular, the modelling strategy is tested on a challenging benchmark case of full-scale wetting of a multi-materials dam. Finally, Section 6 is devoted to some concluding remarks.

# 2. Model problem

#### 2.1. Richards' equation

Richards' equation is a classical nonlinear parabolic equation used to describe flow in unsaturated/saturated zones of an aquifer. Derivation of Richards' equation is described for instance in Szymkiewicz (2013). Variables in play are  $\theta$  the water content [],  $\psi$  the pressure head [L] and  $\mathbb{K}$  the hydraulic conductivity tensor [L  $\cdot$  T<sup>-1</sup>]. The mixed formulation is selected in the present study owing to its versatility (Farthing and Ogden, 2017):

$$\partial_t \theta(\psi) - \nabla \cdot (\mathbb{K}(\psi)\nabla(\psi + z)) = 0. \tag{1}$$

Alternative pressure-based or saturation-based formulations have been discarded because they are either non-conservative or undefined for complete saturation and heterogeneous soils (Celia et al., 1990). The mixed formulation can be rewritten in the hydraulic head form which is more common in hydrology:

$$\partial_t \theta(h-z) - \nabla \cdot (\mathbb{K}(h-z)\nabla h) = 0, \tag{2}$$

where  $h = \psi + z$  is the hydraulic head [L]. One can also add a source/sink term Q [T<sup>-1</sup>] to Richards' equation to model various processes, like bacteria colony or plant roots uptake, or to couple Richards' equation with free surface flow.

Solving Eq. (1) requires two constitutive laws: one for hydraulic conductivity and one for water content. Several models have been proposed, depending on the hydraulic properties of the porous medium.

The hydraulic conductivity  $\mathbb{K}$  is generally supposed to react to saturation identically for each space direction. This leads to write:

$$\mathbb{K}(\psi) = \mathbb{K}_{s} K_{r}(\psi), \tag{3}$$

where  $\mathbb{K}_{s}$  the intrinsic or saturated hydraulic conductivity tensor  $[L \cdot T^{-1}]$  and  $K_{r}$  the relative hydraulic conductivity [].

For practical purposes, the water content is often described in terms of effective saturation  $S_{\rm e}$  []:

$$S_{\rm e}(\psi) = \frac{\theta(\psi) - \theta_{\rm r}}{\theta_{\rm s} - \theta_{\rm r}},\tag{4}$$

where  $\theta_s$  denotes the saturated water content [] and  $\theta_r$  the residual water content [], corresponding to the maximal and minimal saturations, respectively.

The hydraulic properties present two different behaviours depending on whether the porous media is saturated ( $\psi \ge 0$ ) or not ( $\psi < 0$ ):

$$S_{\rm e}(\psi) = \begin{cases} 1 & \text{if } \psi \ge 0, \\ S_{\rm e} & \text{otherwise,} \end{cases} \quad \text{and} \quad K_{\rm r}(\psi) = \begin{cases} 1 & \text{if } \psi \ge 0, \\ K_{\rm r} & \text{otherwise.} \end{cases}$$
(5)

 $S_{\rm e}$  and  $K_{\rm r}$  are monotonic increasing functions of pressure head  $\psi$  in the unsaturated zone. The water table corresponds to  $\psi = 0$  by definition and is considered belonging to the saturated zone. The capillary fringe is the layer above water table where water is raised due to capillary actions. It belongs to the unsaturated zone and there is no standard definition for its upper limit. Throughout this paper, several constitutive laws will be used to model hydraulic properties in the unsaturated zone. They are compiled in Table 1.

It is worth noting that:

- under complete saturation, hydraulic properties become constant and Richards' equation degenerates into an elliptic equation characterised by fast diffusion;
- under almost complete unsaturation, hydraulic properties get very near-zero values which stop diffusion and may be inconvenient numerically;
- for particular set of parameters, when  $\psi \to 0^-$ , constitutive laws may exhibit very steep gradients.

These constitutive laws are mainly responsible for the numerical challenges of Richards' equation, such as nonlinearities, degenaracies and instabilities, which are often observed in the presence of material heterogeneities or dynamic boundary conditions like seepage boundary condition. Special treatment may be done to regularize the constitutive relations (Jger and Kačur, 1995; Pop, 2002; Radu et al., 2006). For instance, Dolejší et al. (2019) modify slightly the functions to avoid some types of degeneracy mentioned above and improve the convergence properties. It can cause difficulties for numerical analysis but, in practice, we find that the convergence of our solving algorithm described in Section 3 can be ensured choosing carefully the parameters of the computations. As a result, no modification on the constitutive relations is done in this study.

### 2.2. Seepage boundary condition

The seepage boundary condition is specific to subsurface model (Scudeler et al., 2017). This condition is used to model the interface between a porous medium and the atmosphere. If the porous medium is saturated and an outflow occurs, then water pours out at atmospheric pressure:  $\psi = 0 \iff h = z$ . Otherwise, the interface acts as an impervious boundary and there is no flux. This condition mimics an outflow condition. Fig. 1 depicts the situation. The treatment of the seepage boundary condition is difficult because the length of seepage face is unknown a priori: it depends on the sought solution.

There are many ways to express and realize the seepage boundary condition. A simple and common approach is to refer to its base definition by considering a switch between a Dirichlet  $\psi_D$  and a Neumann  $q_N$  boundary condition (Cooley, 1983; Beaugendre et al., 2006; Šimůnek et al., 2009; Scudeler et al., 2017):

$$\begin{cases} h = z & \text{if } h \ge z \text{ and } -\mathbb{K}(h-z)\nabla h \cdot \boldsymbol{n} > 0, \\ -\mathbb{K}(h-z)\nabla h \cdot \boldsymbol{n} = 0 & \text{otherwise.} \end{cases}$$
(6)

#### Table 1

Hydraulic relations used in this paper.

Name	Expression	Parameters
Gardner–Irmay relations (1958) (Irmay, 1954; Gardner, 1958)	$S_{\rm e} = e^{\frac{\alpha \psi}{m}} K_{\rm r} = e^{\alpha \psi}$	a: pore-size distribution [-] m: tortuosity [-]
Vachaud's relations (1971) (Vachaud and Thony, 1971)	$S_{\rm e} = \frac{C}{C +  \psi ^{\rm D}} K_{\rm r} = \frac{A}{A +  \psi ^{B}}$	A, C: empirical shape parameters $[L^{B,D}]$ B, D: empirical shape parameters []
Van Genuchten-Mualem relations (1980) (Mualem, 1976; van Genuchten, 1980)	$S_{\rm e} = (1 + (\alpha  \psi )^n)^{-m} K_{\rm r} = S_{\rm e}^l (1 - (1 - S_{\rm e}^{\frac{1}{m}})^m)^2$	$l = \begin{cases} 0.5 & \text{for Mualem (1976)} \\ 1 & \text{for Burdine (1953)} \end{cases} : \text{pore connectivity []} \\ a: \text{ parameter linked to air entry pressure inverse [L-1]} \\ n > 1: \text{ pore-size distribution []} \\ m = 1 - \frac{1}{2}: \text{ pore-size distribution []} \end{cases}$



Fig. 1. Seepage modelling by the boundary condition.

The realization of such condition should be done inside the nonlinear iterative process according to the previous solution guess at a local level. However, it is not always the case because of the involved numerical methods and discretization schemes. Then, the exit point is searched iteratively until a convergence criterion is reached according to different techniques.

Alternatively, the seepage boundary condition may be interpreted as a nonlinear Robin boundary condition:

$$\mathbb{1}_{S}(h)(h-z) - (1 - \mathbb{1}_{S}(h))\mathbb{K}(h-z)\nabla h \cdot n = 0,$$
(7)

where the seepage indicator function is:

$$\mathbb{1}_{S}: \Gamma_{S} \to \{0, 1\} \tag{8}$$

$$h \mapsto \begin{cases} 1 & \text{if } h \ge z \text{ and } -\mathbb{K}(h-z)\nabla h \cdot \boldsymbol{n} > 0, \\ 0 & \text{otherwise.} \end{cases}$$
(9)

This compact formulation will be used here for the discretization to incorporate directly the seepage boundary condition into the weak formulation, keeping in mind that it encompasses a mix of a Dirichlet and a Neumann boundary conditions whose actual realization is done such as in Eq. (6).

# 3. Numerical methods

#### 3.1. Discontinuous Galerkin discretization

More careful developments about discontinuous Galerkin (DG) methods can be found in Rivière (2008) or Dolejší and Feistauer (2015). Let  $d \in \{1, 2, 3\}$  be the dimension. The porous medium is represented by the computational domain  $\Omega \subset \mathbb{R}^d$  of boundary  $\partial\Omega$  and the final time is  $T \in \mathbb{R}^*_+$ . The boundary  $\partial\Omega$  is subdivided into three mutually disjoint boundaries,  $\partial\Omega = \Gamma_D \cup \Gamma_N \cup \Gamma_S$ , corresponding to the Dirichlet, Neumann and seepage boundary conditions respectively. The following problem is considered:

find 
$$h(\mathbf{x}, t) : \Omega \times (0, T) \longrightarrow \mathbb{R}$$
 such that

$$\begin{cases} \partial_{t}\theta(h-z) - \nabla \cdot (\mathbb{K}(h-z)\nabla h) &= 0, \quad \text{in } \Omega \times (0,T), \\ h &= h_{0}, \quad \text{in } \Omega \times \{0\}, \\ h &= h_{D}, \quad \text{on } \Gamma_{D} \times (0,T), \\ -\mathbb{K}(h-z)\nabla h \cdot \boldsymbol{n} &= q_{N}, \quad \text{on } \Gamma_{N} \times (0,T), \\ \mathbb{1}_{S}(h)(h-z) - (1-\mathbb{1}_{S}(h))\mathbb{K}(h-z)\nabla h \cdot \boldsymbol{n} &= 0, \quad \text{on } \Gamma_{S} \times (0,T). \end{cases}$$
(10)

The time duration (0,T) is subdivided into N time intervals such that  $0 = t^0 < t^1 < \ldots < t^N = T$ . Let  $n \in \mathbb{N}, 0 < n \le N$ : if the time interval  $T^n = [t^{n-1}, t^n]$  is considered, the corresponding time step is  $\tau^n = t^n - t^{n-1}$ .

 $\Omega$  is subdivided into  $N_E$  mutually disjoint polygonal elements E forming a mesh. The mesh at the time partition  $T^n$  is denoted by  $\mathcal{E}_{\mathfrak{h}}^n = \{E^n\}_{E \subset \Omega}$ . So,  $\overline{\Omega} = \bigcup_{E \in \mathcal{E}_{\mathfrak{h}}^n} E$ . Moreover, the boundary of one element

 $E \in \mathcal{E}_{\mathfrak{h}}^{n}$  is  $\partial E$  while its diameter  $\mathfrak{h}_{E}$  is defined by:

$$\mathfrak{h}_E := \sup_{\mathbf{x}, \mathbf{y} \in E} |\mathbf{x} - \mathbf{y}|. \tag{11}$$

For every mesh  $\mathcal{E}_{h}^{n}$ , the space step is set as:

$$h^{n} := \max_{E \in \mathcal{E}_{\kappa}^{n}} (\mathfrak{h}_{E}).$$

$$(12)$$

The set of all open faces of all elements  $E \in \mathcal{E}_{\mathfrak{h}}^{n}$  is denoted by  $\mathcal{F}_{\mathfrak{h}}^{n}$ . Furthermore, these definitions stand:

$$\mathcal{F}_{\mathfrak{h},\mathrm{D}}^{n} := \bigcup_{F \in \Gamma_{\mathrm{D}}} F, \quad \mathcal{F}_{\mathfrak{h},\mathrm{N}}^{n} := \bigcup_{F \in \Gamma_{\mathrm{N}}} F, \quad \mathcal{F}_{\mathfrak{h},\mathrm{S}}^{n} := \bigcup_{F \in \Gamma_{\mathrm{S}}} F$$
(13)

$$\mathcal{F}_{\mathfrak{h},\mathrm{B}}^{n} := \mathcal{F}_{\mathfrak{h},\mathrm{D}}^{n} \cup \mathcal{F}_{\mathfrak{h},\mathrm{N}}^{n} \cup \mathcal{F}_{\mathfrak{h},\mathrm{S}}^{n}, \quad \mathcal{F}_{\mathfrak{h},\mathrm{I}}^{n} := \mathcal{F}_{\mathfrak{h}}^{n} \setminus \mathcal{F}_{\mathfrak{h},\mathrm{B}}^{n}.$$
(14)

For each face  $F \in \mathcal{F}_{\mathfrak{h}}^n$ , there is a unit normal vector  $\mathbf{n}_F$ . For  $F \in \mathcal{F}_{\mathfrak{h},1}^n$ , its orientation is arbitrary but kept fixed and, for  $F \in \mathcal{F}_{\mathfrak{h},B}^n$ , it is oriented outward.

Let two neighbouring elements  $E_1$  and  $E_r$  sharing one face  $F \in \mathcal{F}_{\mathfrak{h},\mathbf{I}}^n$ . There are two traces of a function u along F denoted by  $u_1$  and  $u_r$ :

$$\mathbf{x} \in F, u_{1}(\mathbf{x}) := \lim_{\varepsilon \to 0^{+}} u(\mathbf{x} + \varepsilon \mathbf{n}_{F}), \qquad \mathbf{x} \in F, u_{r}(\mathbf{x}) := \lim_{\varepsilon \to 0^{-}} u(\mathbf{x} + \varepsilon \mathbf{n}_{F}).$$
(15)

Formally, the jump and the mean of the function *u* across a face  $F \in \mathcal{F}_{\mathfrak{h}, \mathfrak{l}}^n$  are defined respectively by:

$$\llbracket u \rrbracket := u_{1} - u_{r}, \qquad \llbracket u \rrbracket := \frac{1}{2}(u_{1} + u_{r}).$$
<sup>(16)</sup>

In the case that  $F \in \mathcal{F}_{\mathfrak{h},B}^n$ ,  $\llbracket u \rrbracket = \llbracket u \rrbracket = u_1$  where  $u_1$  is the trace of the function *u* from the element which holds *F*. Throughout this paper, the quantity  $\mathfrak{h}_F$  denotes the length of  $F \in \mathcal{F}_{\mathfrak{h}}^n$  and  $p_F$  the mean of polynomial degree of neighbouring elements:

$$\mathfrak{h}_F := |F|, \qquad p_F := \sqrt{\{\!\!\!\!\ p^2\}\!\!\!\!\}}. \tag{17}$$

In the following derivation of the DG variational formulation, several interior penalty methods are considered. Their numerical properties are slightly different. Further information are available in Rivière (2008);

**Dolejší and Feistauer (2015)** for example. They use discontinuous approximations over the mesh  $\mathcal{E}_{\mathfrak{h}}^n$  defining a so-called broken Sobolev space  $H^s(\mathcal{E}_{\mathfrak{h}}^n)$  with s > 3/2. The solution is sought in a subspace of this broken Sobolev space, taken to be:

$$S_p(\mathcal{E}^n_{\mathfrak{h}}) := \{ v \in L^2(\Omega) : v \Big|_E \in \mathbb{P}_p(E), \forall E \in \mathcal{E}^n_{\mathfrak{h}} \},$$
(18)

where  $\mathbb{P}_p(E)$  denotes the space of polynomial functions on *E* of degree less than  $p \in \mathbb{N}$ .

Eq. (10) is multiplied by a test function  $v \in S_p(\mathcal{E}_{\mathfrak{h}}^n)$  and then integrated on each element  $E \in \mathcal{E}_{\mathfrak{h}}^n$ . Green's theorem is used element-wise before summing over all elements in  $\mathcal{E}_{\mathfrak{h}}^n$ . The Neumann boundary condition has appeared naturally within the formulation and has been substituted. So it stands for the homogeneous Neumann part of the seepage boundary condition.

On one hand, knowing  $\forall u, v \in S_p(\mathcal{E}_{\mathfrak{h}}^n), [\![uv]\!] = [\![u]\!] [\![v]\!] + [\![v]\!] [\![u]\!]$  and assuming  $[\![\mathbb{K}(h-z)\nabla h \cdot n_F]\!] = 0$  because this quantity is smooth enough, product jump is decomposed and simplified where it is possible.

On the other hand, two penalty terms are considered. One mimics the solution continuity by constraining the interior solution jump to be zero, and one enforces the Dirichlet boundary condition, including the remaining part of the seepage boundary condition. If the solution is a continuous function satisfying the Dirichlet boundary conditions, the two penalty terms are vanishing so they can be added to the formulation:

$$J_{\mathbf{I}}(h,v) = \sum_{F \in \mathcal{F}_{\mathfrak{h}}^{\mathbf{I}}} \int_{F} \rho_{F}^{\mathbf{I}} \llbracket h \rrbracket \llbracket v \rrbracket \, \mathrm{d}F, \qquad J_{\mathbf{D}}(h,v) = \sum_{F \in \mathcal{F}_{\mathfrak{h}}^{\mathbf{D}}} \int_{F} \rho_{F}^{\mathbf{D}}(h-h_{\mathbf{D}})v \, \mathrm{d}F.$$
(19)

The interior and Dirichlet penalty weights are set as:

$$\varrho_F^{\rm I} = \frac{\sigma_F^{\rm I} \gamma_F}{\mu_F} \quad \text{and} \quad \varrho_F^{\rm D} = \frac{\sigma_F^{\rm D} \gamma_F}{\mu_F}.$$
(20)

where the face measure is defined as  $\mu_F = \frac{\mathfrak{h}_F}{p_F^2}$  and where  $\sigma_F^{\mathrm{I}}$ ,  $\sigma_F^{\mathrm{D}}$  and  $\gamma_F$  are suitably chosen positive constants.  $\sigma_F^{\mathrm{I}}$  and  $\sigma_F^{\mathrm{D}}$  are user-defined parameters which must be above a threshold value to provide coercivity and guarantee stability for some of the DG methods (Epshteyn and Rivière, 2007).  $\gamma$  is a diffusion penalty coefficient set to one generally whereas some studies consider other values, see e.g. Schötzau and Zhu (2009).

A first weak formulation rises but it is not is not symmetric between the unknown *h* and the test function *v*. Making the formulation symmetrical can be useful to preserve the natural symmetry in the discrete diffusion operator, to use appropriate solvers or to enhance numerical properties of the formulation. This can be done by adding the corresponding symmetric term which vanishes because  $[\![h]\!] = 0$  if *h* is a sufficiently smooth function.

Finally, the space semidiscretization reads:

find 
$$h \in S_p(\mathcal{E}_{\mathfrak{f}}^n)$$
 such that  $\forall v \in S_p(\mathcal{E}_{\mathfrak{f}}^n)$ ,  
 $m_{\mathfrak{f},n}(\partial_t \theta(h-z), v) + a_{\mathfrak{f},n}(h, v) = l_{\mathfrak{f},n}(v)$ , (21)

where the DG bilinear forms, m and a, and linear form l are:

$$m_{\mathfrak{h},n}(\partial_t \theta(h-z), v) = \sum_{E \in \mathcal{E}_{\mathfrak{h}}} \int_E \partial_t \theta(h-z) v \, \mathrm{d}E,$$
(22)

$$\begin{split} a_{\mathfrak{h},n}(h,v) &= \sum_{E \in \mathcal{E}_{\mathfrak{h}}} \int_{E} \mathbb{K}(h-z) \nabla h \cdot \nabla v \, \mathrm{d}E \\ &- \sum_{F \in \mathcal{F}_{\mathfrak{h}}^{\mathrm{I}}} \int_{F} \left[ \mathbb{K}(h-z) \nabla h \cdot \boldsymbol{n}_{F} \right] \left[ v \right] \, \mathrm{d}F \\ &- \sum_{F \in \mathcal{F}_{\mathfrak{h}}^{\mathrm{D}}} \int_{F} \mathbb{K}(h-z) \nabla h \cdot \boldsymbol{n}_{F} v \, \mathrm{d}F + \sum_{F \in \mathcal{F}_{\mathfrak{h}}^{\mathrm{D}}} \int_{F} \rho_{F}^{\mathrm{D}} h v \, \mathrm{d}F \end{split}$$

$$-\sum_{F \in \mathcal{F}_{\mathfrak{h}}^{S}} \int_{F} \mathbb{1}_{S}(h) \mathbb{K}(h-z) \nabla h \cdot \boldsymbol{n}_{F} v \, \mathrm{d}F$$

$$+ \sum_{F \in \mathcal{F}_{\mathfrak{h}}^{I}} \int_{F} \rho_{F}^{I} \llbracket h \rrbracket \llbracket v \rrbracket \, \mathrm{d}F + \sum_{F \in \mathcal{F}_{\mathfrak{h}}^{S}} \int_{F} \rho_{F}^{D} \mathbb{1}_{S}(h) h v \, \mathrm{d}F$$

$$- \Theta \sum_{F \in \mathcal{F}_{\mathfrak{h}}^{I}} \int_{F} \llbracket \mathbb{K}(h-z) \nabla v \cdot \boldsymbol{n}_{F} \rrbracket \llbracket h \rrbracket \, \mathrm{d}F$$

$$- \Theta \sum_{F \in \mathcal{F}_{\mathfrak{h}}^{D}} \int_{F} \mathbb{K}(h-z) \nabla v \cdot \boldsymbol{n}_{F} h \, \mathrm{d}F$$

$$- \Theta \sum_{F \in \mathcal{F}_{\mathfrak{h}}^{S}} \int_{F} \mathbb{1}_{S}(h) \mathbb{K}(h-z) \nabla v \cdot \boldsymbol{n}_{F} h \, \mathrm{d}F, \qquad (23)$$

$$l_{\mathfrak{h},n}(v) = \sum_{F \in \mathcal{F}_{\mathfrak{h}}^{\mathrm{D}}} \int_{F} \sigma_{F}^{\mathrm{D}} h_{\mathrm{D}} v \, \mathrm{d}F + \sum_{F \in \mathcal{F}_{\mathfrak{h}}^{\mathrm{S}}} \int_{F} \sigma_{F}^{\mathrm{D}} \mathbb{1}_{\mathrm{S}}(h) z v \, \mathrm{d}F - \sum_{F \in \mathcal{F}_{\mathfrak{h}}^{\mathrm{N}}} \int_{F} q_{\mathrm{N}} v \, \mathrm{d}F$$
$$-\Theta \sum_{F \in \mathcal{F}_{\mathfrak{h}}^{\mathrm{S}}} \int_{F} \mathbb{K}(h-z) \nabla v \cdot \boldsymbol{n}_{F} h_{\mathrm{D}} \, \mathrm{d}F$$
$$-\Theta \sum_{F \in \mathcal{F}_{\mathfrak{h}}^{\mathrm{S}}} \int_{F} \mathbb{1}_{\mathrm{S}}(h) \mathbb{K}(h-z) \nabla v \cdot \boldsymbol{n}_{F} z \, \mathrm{d}F.$$
(24)

 $\Theta = \{-1, 0, 1\}$  is a constant leading to different symmetric versions of the DG formulation listed in Table 2.

The seepage boundary condition is treated numerically thanks to the indicator function  $\mathbb{1}_S$  which alternates dynamically between the Dirichlet part when  $\mathbb{1}_S = 1$  and the Neumann part for  $\mathbb{1}_S = 0$ , directly in the usual weak framework of the DG methods. This treatment can model many physical situations, like multiple seepage faces simultaneously, because it does not require any assumption about the mesh or seepage face (Scudeler et al., 2017).

This treatment was also done by Dolejší et al. (2019) where they pointed out this is a great benefit compared to conforming FE methods which need to modify the FE space or use numerical tricks, resulting in difficult implementation. However, the indicator function  $1_S$  is not differentiable which causes difficulties for the numerical analysis. A regularization of this function can be done (Schweizer, 2007; Pop and Schweizer, 2011). For example, Dolejší et al. (2019) smoothed the indicator function  $1_F$  to avoid a brutal switch between the values 0 and 1. In this work, no regularization is performed because numerical experiments do not show major convergence solver problems. Besides, DG methods handle already discontinuities and weak penalty naturally. This makes the numerical scheme even more simple.

# 3.2. Time discretization

To get the fully discrete DG formulation, the time derivative has now to be discretized. Such a procedure which takes care of space discretization before time discretization is called the method of lines. It is used commonly by others studies for Richards' equation, see e.g. (Miller et al., 2006; Li et al., 2007b; Farthing and Ogden, 2017).

Backward differentiation formula (BDF) are implicit numerical integration methods especially used for stiff differential equations thanks to their wide region of stability. These linear multistep methods use qalready computed solutions to produce a method of order  $q \le 6$ . Analysis can be found in Sli and Mayers (2003); Dolejší and Vlasák (2008). The 1-step BDF is the backward Euler scheme. For this study, BDF are interesting because they provide high-order accuracy in time which is suitable if one wants to reach high-order in space without loss of gain. The following notation will be used for any function  $u \in L^2(0, T; \Omega)$ :

 $\forall n \in \mathbb{N}_+, u^n := u(\mathbf{x}, t_n). \tag{25}$ 

Table 2
Different types of DG methods.

		Symmetrization		
		$\Theta = -1$	$\Theta = 1$	
Penalization	$ \begin{aligned} \forall F \in \mathcal{F}_{\mathfrak{h}}, \sigma_{F}^{\mathrm{I}} = \sigma_{F}^{\mathrm{D}} = 0 \\ \forall F \in \mathcal{F}_{\mathfrak{h}}, \sigma_{F}^{\mathrm{I}} \neq 0, \sigma_{F}^{\mathrm{D}} \neq 0 \end{aligned} $	OBB method NIPG	– IIPG	global element method SIPG

NIPG, non-symmetric interior penalty Galerkin; SIPG, symmetric interior penalty Galerkin; IIPG, incomplete interior penalty Galerkin; OBB method, Oden–Baumann–Babuška method.

**Table 3**BDF coefficients  $\alpha_{a,k}$  for all methods.

q	1 2	1 <u>3</u>	-1 -2	<u>1</u>				
	3	$\frac{1}{6}$	-3	$\frac{2}{3}{2}$	$-\frac{1}{3}$			
	4	$\frac{23}{12}$	-4	3	$-\frac{4}{3}_{10}$	4 5	1	
	5 6	60 49	-5 -6	<u>15</u>	$\frac{3}{20}$	4 15	5	1
$\alpha_{s,k}$		<b>0</b>	1	<b>2</b> <sup>2</sup>	3	4	5	6
	k							

The DG formulation in Eq. (21) is integrated over  $T^n$  and the time integral is approximated by the BDF methods of 0 < q < 7 steps:

find a sequence of  $(h^n)_{n \in \mathbb{N}_+} \in S_p(\mathcal{E}^n_{\mathfrak{h}})$  such that

$$\begin{cases} h^{0} = h_{0}, \\ \forall v \in S_{p}(\mathcal{E}_{\mathfrak{h}}^{n}), \quad m_{\mathfrak{h},n}(\sum_{k=0}^{q} \frac{a_{q,k}}{\tau^{n}} \theta(h^{n+1-k} - z), v) + a_{\mathfrak{h},n}(h^{n+1}, v) = l_{\mathfrak{h},n}(v, t^{n+1}), \end{cases}$$
(26)

where BDF coefficients are given in Table 3.

However, stability of BDF methods has some restrictions. Only BDF methods of order 1 and 2 are A-stable according to the second Dahlquist barrier for implicit linear multistep methods as defined and proved in Dahlquist (1963). BDF methods of order 3 to 6 are conditionally stable or A( $\alpha$ )-stable where  $\alpha$  decreases with the order, see e.g. Hairer and Wanner (1996); Ascher and Petzold (1998). Then, time stepsizes have to be small enough to make the method stable. BDF methods of order q > 6 are unconditionally unstable so they cannot be used.

BDF methods presented here are based on fixed time steps so initialization and a change of time stepsize has to be treated carefully. For a scheme with q-step BDF, the first time step is subdivided into as many times as necessary to compute the sub-time steps with an increasing sequence of lower-order BDF methods. The remaining next macrotime steps are still subdivided while the number of computed solution, stored gradually, is not sufficient to go on with the *q*-step BDF method. Working with BDF methods of variable time stepsize is possible using divided differences-based formula but with some restrictions on time step variations to keep stability. This is done for Richards' equation in Clément (2021) following the method of Hay et al. (2015). For this study, the backward Euler scheme will be mainly used because it provides good stability properties and remains simple to work with adaptive time stepping. For Tracy's benchmark, the time step is kept unchanged to deal with high-order BDF methods which are necessary to decrease time error regarding space error.

This study uses the mixed form of Richards' equation (physically mass conservative) and DG methods (numerically local mass conservative). Then, mass conservation should be very good which is checked, globally at machine precision and locally with small discrepancies, in Clément (2021).

# 3.3. Linearization

Richards' equation is a nonlinear equation usually solved by an iterative procedure such as fixed-point iteration (also called Picard's) or Newton–Raphson method whose choice is determining for computa-

tion time performances and convergence. Lots of derived methods exist: modified Newton-Raphson, mixed Picard-Newton, quasi-Newton, L-scheme, etc. Studies have been carried out to compare these methods, and others, for solving Richards' equation for various problems (Paniconi and Putti, 1994; Lehmann and Ackerer, 1998; List and Radu, 2016). They emphasize that fixed-point iteration and even Newton-Raphson scheme are very sensitive and do not converge systematically according to Richards' equation's formulations, initial and boundary conditions and because of the nonlinearities of constitutive laws. In particular, the nonlinear iterative solver can oscillate between two solutions. There are still recent and significant works on this topic like Anderson acceleration (Lott et al., 2012), nested Newton's type algorithm (Casulli and Zanolli, 2010) or modified L-schemes (Mitra and Pop, 2019). For this study, a damped Newton-Raphson method and a fixed-point method were implemented. Let m be the number of the nonlinear iteration. The residual of Eq. (26) is expressed as:

$$r_{\mathfrak{h},n}(h^{n+1},v) := m_{\mathfrak{h},n}(\sum_{k=0}^{q} \frac{\alpha_{s,k}}{\tau^{n}} \theta(h^{n+1-k}-z), v) + a_{\mathfrak{h},n}(h^{n+1},v) - l_{\mathfrak{h},n}(v;t^{n+1}).$$
(27)

The difference between two successive iterations is written  $\delta_h^{n+1,m} := h^{n+1,m+1} - h^{n+1,m}$  so the iterative procedure reads:

$$\begin{cases} \frac{d\mathbf{r}_{\mathfrak{h},n}(h^{n+1,m},\upsilon)}{dh^{n+1,m}}\delta_{h}^{n+1,m} = -r_{\mathfrak{h},n}(h^{n+1,m},\upsilon),\\ h^{n+1,m+1} = h^{n+1,m} + \delta_{h}^{n+1,m}. \end{cases}$$
(28)

Underlying ideas are already described in Lehmann and Ackerer (1998) and have just been adapted here to the DG system. Roughly speaking, a fixed-point method is obtained from the Newton–Raphson method by avoiding the first derivative terms inside  $\frac{dr_{h,n}(h^{n+1,m},v)}{dh^{n+1,m}}$ . The damped Newton–Raphson method relaxes the increment  $\delta_h^{n+1,m}$  while the new residual is greater than the previous one. Such a procedure is described in Dolejší et al. (2019). One important choice for nonlinear iterative process is the stopping criterion which is set for this study as:

$$\frac{\|r_{\mathfrak{h},n}(h^{n+1,m},v)\|_{L^{2}(\Omega)}}{\|a_{\mathfrak{h},n}(h^{n+1},v)\|_{L^{2}(\Omega)}} < \varepsilon_{1} \quad \text{and} \quad \frac{\|\delta_{h}^{n+1,m}\|_{L^{2}(\Omega)}}{\|h^{n+1,m}\|_{L^{2}(\Omega)}} < \varepsilon_{2}, \tag{29}$$

where  $\varepsilon_1, \varepsilon_2$  are a user-defined tolerances. These two criteria are relative in order to be independent from the characteristic quantities of the problem. The first criterion is residual and the second one is incremental. The numerical tests of Section 5 were carried out with the fixed-point method because it was more robust concerning initialization.

For each nonlinear iteration, the linear system has to be constructed and solved. We use an in-house direct solver based on LU decomposition and working with skyline matrix storage.

# 3.4. Adaptive time stepping

Time adaptation is motivated by the convergence of the nonlinear solver. On one hand, transient simulations have difficulties to converge if the time step is too large but, on the other hand, shorter time steps mean more time steps and so, a longer computational time. That is the reason why time adaptation is very attractive and common for Richards' equation. Different strategies can be used to adjust the time step (Farthing and Ogden, 2017), either heuristic and mainly based on convergence performance of the nonlinear solver or rational and based on error control. The latter ones are generally more efficient but heuristic methods remains a relevant approach due to their simplicity.

For this study, the time step is adjusted heuristically according to the previous number of iterations  $N_{\rm it}$  from the nonlinear solver such as (Bergamaschi and Putti, 1999; Thoms et al., 2006). Indeed, the time stepsize has a direct effect on the convergence of the solver. The simulations begin with a time step  $\tau^0$ . Then, the future time steps are calculated accordingly to the following rule: each time step is kept if convergence at the previous time iteration is achieved between  $m_{it}$  and  $W_{it}$  nonlinear iterations, it is increased by an amplification factor  $\lambda_{amp}$  if the convergence requires fewer than  $m_{it}$  nonlinear iterations and it is decreased by a reduction factor  $\lambda_{\rm red}$  if the convergence requires more than  $M_{\rm it}$ nonlinear iterations. In the case where the convergence is not fulfilled because of a solver failure (poor initial guess, bad condition number) or because the nonlinear iterations go over a prescribed maximum bound  $W_{\rm it}$ , the time step is started again (back-stepping) using a stepsize reduced by  $\lambda_{red}$ . The time stepping scheme to choose the next time step  $\tau^{n+1}$  is calculated from the previous one  $\tau^n$  can be summarised with:

$$\tau^{n+1} = \begin{cases} \lambda_{amp} \tau^n & \text{if } N_{it} \le m_{it}, \\ \tau^n & \text{if } m_{it} < N_{it} \le M_{it}, \\ \lambda_{red} \tau^n & \text{if } M_{it} < N_{it} \le W_{it}, \end{cases}$$
(30)  
$$\tau^n = \lambda_{red} \tau^n & \text{if } W_{it} < N_{it}$$

or if the solver has failed (time step is started again).

The factors  $0 < \lambda_{red} < 1$  and  $1 < \lambda_{amp}$  as well as the threshold values  $0 < m_{it} \le M_{it} < W_{it}$  are prescribed by the user and found empirically. Default values are  $\lambda_{amp} = 2$ ,  $\lambda_{red} = 0.5$ ,  $m_{it} = 3$ ,  $M_{it} = 7$  and  $W_{it} = 10$ . A minimum time step can be defined to avoid excessive small time steps.

With this approach, the nonlinear solver is more robust because the time step is adjusted until success of convergence independently on  $\tau^0$ . Thanks to the amplification/reduction coefficients, the time step is adjusted smoothly. Nevertheless, the method depends on fixed empirical parameters and does not provide an optimal time step. The resulting loss/gain in computational time is difficult to assess in regards to balance between nonlinear iterations number and time steps length.

# 4. Adaptive strategy

#### 4.1. Adaptive mesh refinement

The Adaptive Mesh Refinement techniques (AMR) are now widely used and have since proven their efficiency, whether on 2D or 3D mesh, structured or unstructured mesh, conforming or not conforming mesh, with domain decomposition or not, see e.g. (Min and Gibou, 2007; Losasso et al., 2004; Fuster et al., 2009; Coupez and Hachem, 2013).

In this work, applications with complex geometries are aimed, or even later couplings of models on hybrid mesh. This is why, as in a previous work (Altazin et al., 2016), a non conforming unstructured mesh is used. Therefore, although computing time may be lost compared to fully structured Cartesian codes, Block-Based Adaptive Mesh Refinement strategy is adopted. In accordance with the simulations presented in this paper, only the bidimensional case is presented. The strategy adopted is then as follows and illustrated in the Fig. 2.

The unstructured mesh is composed of quadrilateral and/or triangular elements, where each element, as in Altazin et al. (2016), defines a *root element* or *block*, see Fig. 2(a). Then, a mesh refinement level is defined for each block, initially by the user and thereafter in accordance with the chosen mesh refinement criterion, see Fig. 2(b). Coarsening and refinement are decided according to threshold values  $0 < \beta_c < \beta_r$ . If  $\eta$  denotes the block-level error estimate, three situations are encountered:

For β<sub>c</sub> < η < β<sub>r</sub>, the block remains unchanged and so for the related elements;



Fig. 2. AMR Strategy: (a) Block mesh; (b) Level of mesh refinement; (c) mesh generation; (d) Morton numbering.



Fig. 3. Mesh refinement for quadrangle and triangle using quadtree.

- For β<sub>r</sub> < η, the block is refined so its elements are split into four isotropic subelements;
- For η < β<sub>c</sub>, the block is coarsened so four neighbouring subelements are merged into one element.

Limitations of such selection are described in Pons and Ersoy (2019) as well as a procedure to overcome them. The mesh refinement level is adapted according to the rule that the ratio of mesh refinement should not exceed 2 between two neighbouring blocks. As illustrated in Ersoy et al. (2013), this constraint allows smooth transitions between refined and unrefined regions. The mesh is then built using a quadtree graph (available for quadrangle and triangle) in order to define easily and precisely the neighbouring elements of each face. This procedure is pursued until the desired level of mesh refinement is reached as shown in Fig. 2(c). In the case of evolutionary problems, this implies frequent remeshing to follow the phenomenon studied. This is why we prefer to widen the stencil with a coarse root mesh, namely blocks, and therefore to use the whole quadtree graph, in order to remesh less often. It is illustrated for quadrangles and triangles in Fig. 3. Finally, a space filing curve using Morton numbering (Z-order) is built in order to number easily the degrees of freedom line in Fig. 2(d).

During a simulation, if a mesh refinement or coarsening process takes place, a new evaluation of the unknown field must be carried out. The prolongation (refinement) and restriction (coarsening) process are facilitated by the fact that the quadtree graph connect explicitly the "mother" element to its four "daughter' elements. To do that, one may use local interpolation/extrapolation. This is not straightforward in practice because it requires to identify the number of points and their position on the element to construct a right-determined system giving a fairly close solution. In the present study, the projection is made by solving a local DG problem. This technique gives good degrees of freedom by performing calculations at quadrature points. The element-wise weak formulation corresponding to  $h_{E_{new}} = h_{E_{old}}$  reads:

$$\int_{E_{\text{new}}} h_{E_{\text{new}}} v \, \mathrm{d}E_{\text{new}} = \int_{E_{\text{new}}} h_{E_{\text{old}}} v \, \mathrm{d}E_{\text{old}}.$$
(31)

Then, mass matrix can be used to project the solution on the new mesh. This method is also used to enforce the initial condition  $h = h_0$ .

#### 4.2. A posteriori error estimation

An *a posteriori* error indicator is employed to show how a simple quantity linked to problem physics is able to provide information about refinement. This error indicator is inspired from Miller et al. (2006) where they use changes in effective saturation. Here, the error indicator is based directly on changes in the solution that is to say the gradient of hydraulic head *h*. This allows to work both for the saturated and unsaturated regions. So, it reads:

$$C_E = \frac{1}{\max_{E \in \mathcal{E}_k} (\frac{1}{|E|} \|\nabla h\|_{L^2(E)})} \frac{1}{|E|} \|\nabla h\|_{L^2(E)}.$$
(32)

The error indicator is normalized so that  $0 \le C_E \le 1$ .

Another a posteriori error indicator is introduced but it is based on error estimation. Indeed, Verfürth developed a posteriori estimationbased error indicators based on residual and derived thanks to suitable norms in the context of finite elements methods for many equations (Verfürth, 2013), in particular for nonlinear parabolic problems such as Richards' equation. For this kind of problem, Verfürth used some variants to prove reliability as well as global and local efficiency for implicit Runge-Kutta schemes with some restrictions (Verfürth, 1998) and more standard time discretization in Verfürth (2003); Verfrth (2004). Later on, this type of residual-based derivation was adapted by Melenk and Wohlmuth to  $h_p$ -adaptive finite elements methods for elliptic problems (Melenk and Wohlmuth, 2001). Afterwards, Houston et al. (2007) as well as Schötzau and Zhu (2009); Zhu and Schotzau (2010) developed residual-based a posteriori error estimation for hp-adaptive interior penalty DG methods applied to elliptic problems and convectiondiffusion equations respectively.

Greatly inspired by these works and supported by an heuristic analysis, the following *a posteriori* estimation-based error indicator is used:

$$(\eta_E^n)^2 = (\eta_{E,R}^n)^2 + (\eta_{E,F}^n)^2 + (\eta_{E,J}^n)^2,$$
(33)

where  $\eta_{E,R}^n$ ,  $\eta_{E,F}^n$  and  $\eta_{E,J}^n$  are respectively the element residual, the face residual and the solution jump sub-estimates.  $\rho_F^I$  and  $\rho_F^D$  were defined in Eq. (20). The quantities  $\lambda_m(\mathbb{K})$  and  $\lambda_M(\mathbb{K})$  stand respectively for the lowest and largest eigenvalue of  $\mathbb{K}$  on *E*. For a face *F* of normal *n*, the minimum of normal component from neighbouring  $\mathbb{K}$  is chosen:  $\kappa_m :=$ min $(\kappa_l, \kappa_r)$  where  $\kappa = \mathbf{n}^T \cdot \mathbb{K}(u_h^{n+1}) \cdot \mathbf{n}$ . So sub-estimates write:

$$(\eta_{E,\mathsf{R}}^n)^2 = \frac{\mathfrak{h}_E^2}{p_E^2 \lambda_{\mathsf{m}}(\mathbb{K})} \| \frac{\theta(u_{\mathfrak{h}}^{n+1}) - \theta(u_{\mathfrak{h}}^n)}{\tau^n} - \nabla \cdot (\mathbb{K}(u_{\mathfrak{h}}^{n+1}) \nabla u_{\mathfrak{h}}^{n+1}) \|_{L^2(E)}^2, \tag{34}$$

$$\begin{aligned} (\eta_{E,F}^{n})^{2} &= \sum_{F \in \partial E \cap \mathcal{F}_{\mathfrak{h}}^{1}} \frac{\mathfrak{h}_{F}}{2p_{F}\kappa_{\mathfrak{m}}} \| \left[ \mathbb{K}(u_{\mathfrak{h}}^{n+1}) \nabla u_{\mathfrak{h}}^{n+1} \cdot \boldsymbol{n} \right] \|_{L^{2}(F)}^{2} \\ &+ \sum_{F \in \partial E \cap \mathcal{F}_{\mathfrak{h}}^{N}} \frac{\mathfrak{h}_{F}}{p_{F}\kappa_{\mathfrak{l}}} \| q_{\mathfrak{N}} - \mathbb{K}(u_{\mathfrak{h}}^{n+1}) \nabla u_{\mathfrak{h}}^{n+1} \cdot \boldsymbol{n} \|_{L^{2}(F)}^{2}, \end{aligned}$$
(35)

$$\begin{aligned} (\eta_{E,\mathbf{J}}^{n})^{2} &= \sum_{F \in \partial E \cap \mathcal{F}_{\mathfrak{h}}^{1}} \frac{1}{2} (\rho_{F}^{1} + \frac{\mathfrak{h}_{F}}{p_{F}\kappa_{m}}) \| [\![u_{\mathfrak{h}}^{n+1}]\!] \|_{L^{2}(F)}^{2} \\ &+ \sum_{F \in \partial E \cap \mathcal{F}_{\mathfrak{h}}^{D}} (\rho_{F}^{D} + \frac{\mathfrak{h}_{F}}{p_{F}\kappa_{1}}) \| u_{D} - u_{\mathfrak{h}}^{n+1} \|_{L^{2}(F)}^{2}. \end{aligned}$$
(36)

Here, the time discretization is based on the implicit Euler scheme but the *a posteriori* estimation-based error indicator can be determined for other time schemes such as in Verfürth (1998). Global estimates are obtained with:

$$(\eta^n)^2 = \sum_{E \in \mathcal{E}_h^n} (\eta_E^n)^2.$$
 (37)

Unlike the aforementioned papers, there is no rigorous mathematical proof for this residual-based energy norm a posteriori error estimation. Extension to the case of nonlinear and time-dependant parabolic equation solved by hp-adaptive DG methods remains an open problem. However, some remarks can be drawn. Firstly, the numerical treatment of the time dependence and of the nonlinearity resorts to consider a sequence of linear and steady problems. Secondly, as noted by Verfürth (2013), the element residual term is related to the residual of the numerical solution with respect to the strong form of the equation which may be viewed as the error from the nonlinear process. The face residual term is related to the boundary operator which is associated with the strong and weak forms of the differential equation. It reflects, on one hand, that the numerical solution gradient - the flux - is discontinuous and on the other hand, that Neumann boundary conditions may not be satisfied. The solution jump term is related to the penalization which are associated with the weak form of the equation. It reflects that the numerical solution is discontinuous at interior faces in the DG framework and Dirichlet boundary conditions may be slightly violated according to the penalty coefficients. Moreover, the second term and third parts of the estimation-based error indicators measure how valid the hypothesis on the seek solution are to derive the DG formulation.

#### 4.3. Weighted discontinuous Galerkin framework

In the context of convection-diffusion equations or coupling of parabolic/hyperbolic domains, the solution may hold spurious oscillations at internal layers for vanishing or varying diffusion (Ern and Proft, 2006). This situation is typical of porous media problems where there are material heterogeneity and degenerate hydraulic properties. One interpretation is the flow stays continuous but the solution mimics a discontinuity. Then, the numerical scheme is unable to recognize the sharp internal layer leading to numerical instabilities.

Even though this phenomenon may be resolved by mesh refinement, the corresponding computational cost is substantial to maintain a physically acceptable solution. Slope limiters are another possibility to control these undershoots/overshoots but further computational developments are needed to cope with the geometry of elements in high-dimension (Sochala, 2008; Rivière, 2008). A satisfactory approach would be to design a DG method that can handle internal layers in an automated fashion. That is the reason why the weighted discontinuous Galerkin (WDG) framework was introduced (Ern et al., 2008; Rivière, 2008). These methods work thanks to two key ingredients which can be seen as a way to incorporate into the definition of the scheme some partial knowledge of the solution. The first one is the use of weighted averages instead of the standard arithmetic average inside the discontinuous formulation. Then, the amount of diffusivity flux for each side of a face is controlled. The second ingredient is to relax the face penalization used for continuity constraint by a coefficient depending on the harmonic mean of the diffusivity of the neighbouring elements. Such a penalty strategy turns out to tune automatically the amount of local penalty to regulate the degree of smoothness of the approximate solution. Analysis and details can be found in Ern et al. (2008); Rivière (2008) and related.

Internal layers depend on the spectral structure of the diffusivity. The choice is to take its normal component for face evaluation (Ern et al., 2008; Burman and Zunino, 2006). Weights across a face are positive numbers such that  $\omega_l + \omega_r := 1$ . The weighted average and the conjugate weighted average are respectively defined as:

$$\llbracket u \rrbracket_{\omega} := \omega_{\mathbf{l}} u_{\mathbf{l}} + \omega_{\mathbf{r}} u_{\mathbf{r}}, \qquad \llbracket u \rrbracket_{\overline{\omega}} := \omega_{\mathbf{r}} u_{\mathbf{l}} + \omega_{\mathbf{l}} u_{\mathbf{r}}. \tag{38}$$

Then, the relation  $\forall u, v \in S_p(\mathcal{E}^n_{\mathfrak{h}}), [\![uv]\!] = [\![u]\!]_{\varpi}[\![v]\!] + [\![u]\!] [\![v]\!]_{\overline{\omega}}$  is replacing Eq. (3.1) in the weak formulation derivation. The weighted diffusion

penalty coefficient is taken to be the harmonic mean of the neighbouring normal hydraulic conductivities across the face (Ern et al., 2008):

$$\gamma_F = \frac{2\kappa_1 \kappa_r}{\kappa_1 + \kappa_r}.$$
(39)

Standard DG methods assume  $\omega_l = \omega_r = \frac{1}{2}$ , which reduces the weighted average to the arithmetic average, and  $\gamma_F = 1$ . The WDG method chooses other definitions. Following definitions in Ern et al. (2008), let the weights be:

$$\begin{cases} \omega_{l} = \frac{\kappa_{r}}{\kappa_{l} + \kappa_{r}}, & \omega_{r} = \frac{\kappa_{l}}{\kappa_{l} + \kappa_{r}} & \text{if } \kappa_{l} + \kappa_{r} \neq 0, \\ \omega_{l} = \omega_{r} = \frac{1}{2} & \text{otherwise.} \end{cases}$$
(40)

With this definition choice for weights,  $\gamma_F$  results to be equivalent to the weighted average of diffusivities:

$$\gamma_F = \frac{2\kappa_l \kappa_r}{\kappa_l + \kappa_r} = \{\![\kappa]\!]_{\omega}. \tag{41}$$

WDG methods assume that diffusivity discontinuities fit the mesh (Ern et al., 2008). This is generally the case for heterogeneous media but it is not true for nonlinear diffusivity such as in Richards' equation. Then, sharp internal layers may occur inside an element so the weighted framework is not expected to work in this situation. Nevertheless, a suitable adaptive mesh refinement can be used as a capturing technique. This idea is one of the main novelties proposed in this paper. The weighted framework and mesh adaptation are working dynamically in synergy: the former changes the smoothness nature of the numerical solution while the latter tracks the internal layer linked to nonlinearity thanks to refinement driven by the *a posteriori* estimation-based error indicator Eq. (33).

## 5. Numerical results

Simulations are performed to highlight some numerical behaviour of Richards' equation and the ability of the numerical methods to treat them. The IIPG method is employed in every simulation thereafter.

#### 5.1. Polmann's test-case

This test-case considers a soil from New Mexico whose hydraulic properties are provided by Polmann et al. (1988). A 1D vertical sample of this soil subject to downward infiltration was simulated by Celia et al. (1990), Manzini and Ferraris (2004) as well as Sochala (2008). This test-case can be challenging because, on one hand, the prescribed pressure head on both sides of the 100 cm soil column has a difference of 925 cm resulting in steep solicitation, and, on the other hand, hydraulic conductivity shows strong variations under the set of values taken by pressure head. The Polmann's test-case employs Van Genuchten–Mualem relations with  $K_s = 9.22 \times 10^{-3}$  cm s<sup>-1</sup>,  $\theta_s = 0.368$ ,  $\theta_r = 0.102, \ \alpha = 3,35 \times 10^{-2} \text{ cm}^{-1}, \ n = 2, \ m = 0.5 \text{ and } l = 0.5.$  The computational domain  $\Omega$  is a rectangle  $(0, 20) \times (0, 100)$  cm. The test-case is solved for pressure head  $\psi$  during s (48 h) with a constant time step  $\tau = 120$  s and the initial condition is  $\psi_0 = -1000$  cm. Throughout the examples, implicit Euler scheme, p = 1,  $\sigma_{\rm I} = \sigma_{\rm B} = 100$  are used. Computation is done for two meshes: M100 is a coarse mesh of 100 elements and M1000 is a refined mesh of 1000 elements.

For the mesh M100, nonlinear iterations have difficulties to reach the requirements of stopping criteria and the solution holds spurious oscillations, in particular an undershoot ahead of the sharp wetting front as showed in Fig. 4. This behaviour was already noticed by Celia et al. (1990) and Sochala (2008). The latter decides to eliminate this undershoot by adding a slope limiter which causes a small late compared to the non-limited solution. Better results are obtained with the mesh M1000, the oscillations in the solution vanish, see Fig. 5. The M1000-results agree with those from Celia et al. (1990) as well as from Manzini and Ferraris (2004). This shows that a good discretization is necessary for Richards' equation to get quality solution. Such requirement can be fulfilled at optimal cost by a suitable adaptive local refinement. The error indicator from Eq. (32) is employed with refinement threshold values  $\beta_c = \beta_r = 50$ . Adaptation is performed every 5 time steps, starting with the M100 mesh. In Fig. 5, results show that refinement is able to capture the wetting front dynamically and eliminate the undershoot. The average number of elements is 211. More frequent adaptations allow to follow perfectly the front while less frequent adaptations are less costly.

In addition, the WDG method from Section 4.3 is tried without any adaptation. The mesh M100 is used. Results in Fig. 5 show that the undershoot is still present but reduced to one single element. It is relevant to point out that the solution holds jumps where the wetting front is steep. They prevent oscillations to propagate and make solution catch the front steepness. The remaining undershoot depends on the wetting front localization in relation with the mesh geometry. This is due to the nonlinear nature of diffusivity and that is why it is important to associate mesh adaptation.

#### 5.2. Tracy's benchmark

This test-case has an analytical solution given by Tracy (2006) for 2D and 3D problems which can be used as a benchmark for Richards' equation (Tracy, 2007; 2010). Tracy's benchmark is particularly relevant because it is transient, has a simple parameter to vary nonlinearities for Richards' equation and holds differentiated steep regions. It can serve to check the nonlinear solver robustness, convergence properties and adaptive mesh refinement. Then, it is very convenient to assess the solution quality and compare codes: Solin and Kuraz (2011) and Dolejší et al. (2019) used it to evaluate the performance of their respective method.

Tracy's benchmark employs Gardner–Irmay relations with  $K_s = 1.1 \text{ m d}^{-1}$ ,  $\theta_s = 0.5$ ,  $\theta_r = 0$ ,  $\alpha = 0.1 \text{ m}^{-1}$  and m = 1. It is solved for pressure head  $\psi$ . The residual pressure head  $\psi_r$  is a parameter. The computational domain  $\Omega$  is a square  $(0, a) \times (0, b)$  m. Here, a = b = 1 m and  $\psi_r = -10$  m. A specific Dirichlet boundary condition is prescribed on the top edge with the function:

$$\psi_{\text{top}} = \frac{1}{\alpha} \log(e^{\alpha \psi_{\text{r}}} + (1 - e^{\alpha \psi_{\text{r}}}) \sin(\frac{\pi x}{a})). \tag{42}$$

While for other edges, a constant Dirichlet boundary condition is imposed:  $\psi = \psi_r$ . The initial condition is  $\psi_0 = \psi_r$ . Tracy gives the 2D exact solution  $\psi_{ex}$  in Tracy (2006). Fig. 6 shows the solution obtained for a fixed grid with  $\tau = 10^{-4}$  day and  $T = 10^{-2}$  day.

The treatment of initial condition is not straightforward. First, because the initial condition does not respect the top boundary condition: they are inconsistent. This issue can be troublesome for discretizations like finite elements methods (Solin and Kuraz, 2011) but it is natural for DG methods where boundary conditions are usually enforced by weak penalization (Dolejší et al., 2019). Second, because the top boundary condition condition is particularly stiff compared to the initial state. Therefore, the very first moments are the most interesting to simulate in order to address the time-boundary inconsistency and the early stiffness. Nonlinearities are controlled by the parameter  $\alpha$  in Eq. (42). More  $\alpha$  is large, more the relative hydraulic conductivity drops quickly to near zero, and so, nonlinearities are increased for Richards' equation (Tracy, 2007). Moreover, errors are linked to the top corners since the top boundary condition induces a steep slope at these corners (Tracy, 2007; 2010). As Tracy stated, these features make this problem a good option to investigate adaptive mesh refinement.

The classic  $L^2$ -norm and, following Rivière (2008), Schötzau and Zhu (2009); Zhu and Schotzau (2010) or Dolejší et al. (2016), the energy norm, also called the DG intrinsic norm, are introduced for a scalar function  $u : \mathbb{R}^d \longrightarrow \mathbb{R}$ :

$$\|u\|_{L^{2}(E)}^{2} := \int_{E} u^{2} dE \qquad \|\|u\|_{\mathcal{E}(E)}^{2} := \|u\|_{\mathcal{R}(E)}^{2} + \sum_{F \in \partial E} \|u\|_{\mathcal{J}(F)}^{2}, \tag{43}$$



Fig. 4. Pressure head along the vertical for Polmann's test-case with the mesh M100.



**Fig. 5.** Comparison of pressure head at t = 24 h for Polmann's test-case: IIPG on mesh with 100 elements (M100), IIPG on mesh with 1000 elements (M1000), IIPG with adaptive mesh refinement (AMR), WDG with 100 elements (WDG).



Fig. 6. Hydraulic head for Tracy's benchmark with p = 1 and 6-step BDF.



(a) Errors in the  $L^2$ -norm

(b) Errors in the energy norm

Fig. 7. Numerical convergence for Tracy's benchmark. P1, P2, P3 and P4 stands respectively for linear, quadratic, cubic and quartic approximations.

where the residual seminorm and the jump seminorm are respectively:

$$\|u\|_{\mathcal{R}(E)}^{2} := \int_{E} \mathbb{K}(u) \nabla u \cdot \nabla u \, \mathrm{d}F$$
(44)

$$\|u\|_{\mathcal{J}(F)}^2 := \int_F \rho_F [\![u]\!]^2 \,\mathrm{d}F.$$
(45)

Errors between numerical and exact solution are computed for a time step such that:

$$\|e\|_{X} = \|\psi_{ex} - \psi\|_{X} \quad \text{with } X = \{L^{2}, \mathcal{E}, R, J\}.$$
(46)

To measure how effective the estimation-based error indicator is, the effectivity indices for the estimation-based error indicator (33), the element and face residuals (34) and (35), and the solution jump (36) will be used:

$$I_{\mathcal{E}}^{\text{eff}} := \frac{\eta_{E}^{n}}{\|\|e\|\|_{\mathcal{E}(E)}}, \qquad I_{R}^{\text{eff}} := \frac{((\eta_{E,R}^{n})^{2} + (\eta_{E,F}^{n})^{2})^{\frac{1}{2}}}{\|e\|_{R(E)}}, \qquad I_{J}^{\text{eff}} := \frac{\eta_{E,J}^{n}}{\|e\|_{J(F)}}.$$

$$(47)$$

Due to the lack of rigorous derivation, the effectivity index is not expected to represent properly the true error. The estimator efficiency is rather appreciated to set up a mesh refinement strategy.

First of all, several simulations are performed to show convergence properties. Because of symmetry, one half vertical plan of the domain is considered to speed up the simulation. Duration is set to  $T = 10^{-4}$ day to focus on small times. The computations are carried out with the 6-step BDF and time steps are set to  $\tau = 10^{-6}$  s. This is needed to reduce time discretization errors expecting that they will be negligible compared to space discretization errors in order to observe space convergence. Indeed, according to theoretical error estimates (Dolejší and Feistauer, 2015), error in  $L^2$ -norm and  $H^1$ -seminorm behave such that:

$$\|e\|_{L^2} \approx c_{\mathfrak{h}} h^p + c_\tau \tau^q, \tag{48}$$

$$\|e\|_{\mathcal{E}} \approx c_h' h^{p+1} + c_r' \tau^q,\tag{49}$$

where  $c_{\mathfrak{h}}, c'_{\mathfrak{h}}, c_{\tau}$  and  $c'_{\tau}$  are constants independent of *h* and  $\tau$ . Numerical experiments involve four meshes of 50, 200, 400 and 3200 quadrilateral elements. Order approximation varies from one to four. Fig. 7 shows the convergence of the error  $e_{\mathfrak{h}}$  for the  $L^2$ - and energy norms with respect to the number of degrees of freedom. It is interesting to note that space error are so small for the most dense mesh with p = 4 that time error saturates the convergence.

In Fig. 8, effectivity indices are plotted. The error measured with the jump seminorm is independent from the exact solution and so is equivalent to the jump indicator (36). Then, the scaling factor of the jump indicator was chosen to be the penalty weight in order to be equal to jump seminorm of the error. As a consequence,  $I_{\rm I}^{\rm eff}$  controls only the numerical representation of exact solution (interpolation error) and must be equal to one as observed in Fig. 8e. On the contrary,  $I_{\varepsilon}^{eff}$  and  $I_{R}^{eff}$  are not representative of the true error in term of magnitude like observed on Fig. 8a and c. They are blowing up, at least for P1 and P2. It is probably due to a wrong choice for the scaling factors. The factors come from from Schötzau and Zhu (2009); Zhu and Schotzau (2010) who considered another problem. They were kept because they scale properly the sub-indicators to drive mesh adaptation as a capturing technique like showed later on. Moreover, the scaling factors always overestimate the true error which is good for mesh refinement monitoring. True a posteriori error analysis for the nonlinear problem of Richards' equation is known to be difficult and remains completely open.



**Fig. 8.** Effectivity indices for different indicators (left) and convergence diagrams for the three parts of the indicator  $\eta_E^n$  (right). P1, P2, P3 and P4 stands respectively for linear, quadratic, cubic and quartic approximations.

The behaviour of the three contributions with respect to the number degrees of freedom is shown in Fig. 8 by varying the order of approximation and the mesh size. Since they do not represent properly the true error, observations are difficult to relate with the true error. However, some remarks can be drawn. Firstly, these diagrams show the convergence up to a certain extent. Secondly, one can see that the volume contribution – the residual indicator from Eq. (34) – behaves quite differently compared to the surface contributions – the flux and jump indicators from Eqs. (34) and (35). The convergence is globally maximised with *p*-refinement for the surface contributions while  $\mathfrak{h}$ -refinement seems to improve convergence better for the volume contribution. This observation should be assessed in more details but could drive the  $\mathfrak{h}p$ -decision making.

Error estimation is evaluated in terms of ability to drive the mesh adaptation. The error in  $L^2$ -norm is above the error measured with en-

ergy norm as stated in Fig. 9. It is worth to notice that the estimationbased error indicator

Error estimation is evaluated in terms of ability to drive the mesh adaptation. The distribution of  $\|e\|_{L^2(E)}$  with no mesh adaptation is given in Fig. 9 for reference to localize the zones of interest. It is above the error measured with energy norm. In Fig. 10, it is worth to notice that the estimation-based error indicator  $\eta_E^n$  is able to give the shape (but not the magnitude) of the true error measured with the energy norm, particularly for steady state, even though it was not expected because there is no theoretical proof. Besides, during transient state, the estimation-based error indicator spots the error in the top corner where Tracy stated there were problems for this test-case, see Fig. 9 at  $t = 10^{-4}$  d. The gradient-based error indicator  $C_E$  was used successfully in a previous paper (Clément et al., 2020) to capture the moving wetting front on a simple 1D infiltration case. Yet, as shown in Figs. 11 and



Fig. 9. True error distribution in the  $L^2$ -norm on Tracy's benchmark.

12, it performs badly for the Tracy's benchmark where steep gradients arise on a large part of the domain. The refinement thresholds are difficult to tune which leads to over-refinement for large regions at the top. They were set up to  $\beta_c = \beta_r = 0.5$  in order to give  $||e||_{L^2(\Omega)} \approx 0.167$  at  $t = 10^{-2}$  d. Then, the computation is very costly since it takes around 227 min. For the estimation-based error indicator, the ability to drive

the mesh adaptation is more convincing.  $\beta_c = \beta_r = 0.85$  in order to target  $\|e\|_{L^2(\Omega)} = 0.137$  at  $t = 10^{-2}$  d. The mesh is adapted dynamically without inducing over-refinement. Elements are refined where it is needed because adaptation occurs locally while the error is controlled. Then, the cost is reasonable with a computation around 35 min.

# 5.3. Wetting of the La Verne dam

#### 5.3.1. Setting

The model is here used to simulate the full-scale wetting of a multimaterials dam: the La Verne dam. La Verne dam was constructed in 1991 to supply Gulf of Saint–Tropez (south of France) with drinkable water. La Verne dam is 42 m high and peaks at 90 m above sea level. Its width ranges from –102.3 m to 110.5 m. The reservoir is on the left in Fig. 13. The genuine La Verne dam inner section shows a complicated heterogeneous material patterns, partially described in Fleureau and Fry (1991); Bonelli et al. (1993). The dam is an earth-filled embankment dam. A clay core allows for impermeability while outer zones are semi-permeable alluvial shells. The dam is protected by rip-rap upstream. Filters and drains of very permeable gravel-like materials secure the dam inner saturation. The dam foundation lies on an impervious rock stratum thanks to con-





Fig. 10. Comparison between distributions of true error in energy norm with the indicator  $\eta_{E}^{n}$  on Tracy's benchmark.



(a) Mesh and error distribution in  $L^2$ -norm for  $C_E$ -driven AMR(b) Mesh and error distribution in  $L^2$ -norm for  $\eta_E^n$ -driven AMR





(a) Mesh and error distribution in energy norm for  $C_E$ -driven(b) Mesh and error distribution in energy norm for  $\eta_E^n$ -driven AMR AMR

Fig. 12. Comparison with energy norm between  $C_E$ -based adaptation (left) and  $\eta_E^n$ -based adaptation on Tracy's benchmark.



Fig. 13. Geometry, materials and boundary conditions of La Verne dam for the numerical case. S1, S2, S3 are numerical sensors corresponding to the experimental sensors while S4 and S5 are additional purely numerical sensors.

crete injections. A fine loam layer covers the downstream dam part. The filling of La Verne dam was controlled with instruments (Fleureau and Fry, 1991) during 40 days, which provides a unique and challenging dataset to test the model. In particular, the reservoir height evolution is available as well as three groundwater hydraulic head measurements from the core, see Figs. 13 and 15. The three sensors are at z = 55 m and their x-coordinates are -10.5 m, -3 m and 5 m.

For the simulation, the reservoir height is used as a forced boundary condition in the form of a Dirichlet boundary condition monitored by a function based on experimental data (see Fig. 14). Since comparison focusses on the core of the dam, it is assumed that outer high permeable and/or fine layers materials do not have a significant impact on inner groundwater flows and saturation. The downstream thin loam layer, the filter-isolated toe drain and the lower submerged rip-rap are therefore not represented by the simulation and assimilated to the adjacent mate-

rials (see Table 4 for description of numerical dam structure). A Dirichlet boundary condition for hydraulic head is prescribed at the downstream boundary. The upper rip-rap and the cofferdam are simulated as such to assess the method robustness owing to their direct contact with the upstream dynamic boundary condition (forcing) and their hydraulic properties are steep. Rock/concrete foundations are supposed to be perfectly impervious which means a zero-valued Neumann boundary condition can be prescribed. Finally, seepage boundary condition are prescribed everywhere else because outflow can drain from the exposed upstream and downstream shell slopes. The actual simulated configuration of La Verne dam is sketched in Fig. 13. Five different materials are simulated (Table 4), with hydraulic properties provided by Bonelli et al. (1993). The dam wetting simulation involves each tool presented in this paper for the adaptive strategy: the WDG method in combination with the AMR technique driven by the estimation-based error indicator. Mesh



Fig. 14. Time evolution of hydraulic head h, mesh and water table (white line) during La Verne dam wetting.

Table 4Materials of the numerical La Verne dam.

Materials	Components	Constitutive laws	$\theta_{\rm s}$	$\theta_{\rm r}$	$\mathbb{K}_{\mathrm{s}}(1,1)(m\cdot d^{-1})$	$\mathbb{K}_{s}(2, 2) (m \cdot d^{-1})$	Specific parameters
m1	Core Cofferdam	Van Genuchten–Mualem	0.23	0	$2.592\times10^{-3}$	$8.64\times10^{-4}$	$\alpha = 0.08  m^{-1}, n = 1.2, l = 0.5$
m2	Upstream shell Downstream shell	Van Genuchten-Mualem	0.25	0	5.184	1.728	$\alpha = 0.01  m^{-1}, n = 2.1, l = 0.5$
m3	Outer upstream shell	Vachaud	0.22	0	5.184	1.728	$A = 2.99 \times 10^{-4} \text{ m}^5$ , $B = 5 C = 6.34 \times 10^{-2} \text{ m}^{2.9}$ , $D = 2.9$
m4	Protection rip-rap	Vachaud	0.27	0	17.28	8.64	$A = 2.99 \times 10^{-4} \text{ m}^5$ , $B = 5 C = 6.34 \times 10^{-2} \text{ m}^{2.9}$ , $D = 2.9$
m5	Filters Drains	Vachaud	0.32	0	17.28	8.64	$A = 2.99 \times 10^{-4} \text{ m}^5$ , $B = 5 C = 6.34 \times 10^{-2} \text{ m}^{2.9}$ , $D = 2.9$

adaptation is done every five time steps. Fixed-point iteration is used together with adaptive time stepping. Order of approximation is quadratic for space and one for time (implicit Euler) in order to have a robust and cost-effective computation. Penalty parameters are  $\sigma_{\rm I} = \sigma_{\rm B} = 100$ .

The simulation of La Verne dam case is a very challenging benchmark for numerical models, combining heterogeneous materials, steep constitutive laws and dynamic boundary conditions. However, despite of its interest, the La Verne dam benchmark remains only partly documented by in-situ instruments, precluding a comprehensive quantitative confrontation between observations and model. In particular, the hydraulic properties, which have a drastic effect on flow dynamics, remain approximative due to the absence of direct in-situ characterization. In addition, note that no initial in-situ values for water table elevation and saturation are available inside the dam, which can also strongly affect the subsequent evolution. For the present simulation, the initial water table is imposed at 67 m to fit the initial values of experimental hydraulic head. Finally, Bonelli et al. pointed out that hydromechanical coupling should be considered to be fully consistent with the real case, see Fleureau and Fry (1991).

#### 5.3.2. Results

Fig. 14 depicts hydraulic head (colour contours), water table (white line) and adaptive mesh for both simulations every 10 days. Time evolution of simulated hydraulic head and water content are displayed in Fig. 15 at the position of the three experimental sensors (S1 to S3) and for two additional relevant points called numerical sensors (S4 and S5). It is recalled that initial reservoir free surface and water table elevation are at 57 and 67 m, respectively. Note that the water table is not displayed in Fig. 14 at some points due to insufficient resolution and oscillations w.r.t the front sharpness, which are detailed in the Discussion section. The numerical model is observed to provide a good overall description of the dam wetting process. Fig. 14 shows that the propagation of the wetting wave inside the dam is strongly dependent on the inner materials properties. The first eleven days are characterized by an ad-



**Fig. 15.** Time evolution of inner dam hydraulic parameters. a) Hydraulic head at sensors S1 (short-dashed lines), S2 (dash-dotted lines) and S3 (long-dashed lines) from the present simulation (red), the experimental data (light blue) and the previous results of Bonelli et al. (1993). The reservoir height is depicted in dark solid line. b) Hydraulic head at numerical sensors S4 (orange) and S5 (yellow). c) Water content at S1 to S5 sensors (note that S1, S2 and S3 show similar values). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)



(a) Distribution of water content at t = 40 days





Fig. 16. Heterogeneity and nonlinear varying diffusivity highlighted respectively by water content and Péclet number for La Verne dam simulation with the reference simulation.

justement phase during which the reservoir surface remains below the initial water table elevation. Hydraulic head and water content at numerical sensors S4 and S5 show a very constrasting behaviour (Fig. 15b and c). This reveals that, even if both m2- and m3-materials show the same hydraulic conductivity at saturation, the difference in constitutive laws induces a radically different dynamics of the capillary fringe. At S4, the head remains nearly constant during the adjustement phase while the water content is very low. The m3-material is here nearly desaturated, well above a thin capillary fringe. By contrast, S5 is within the large capillary fringe associated to the m2-material, providing a much higher water content. The hydraulic head shows first a small decay, probably due to the drain proximity, before starting to rise in response to the reservoir filling. Around day 14, both S4 and S5 sensors show a simultaneous regular rising, slightly lagged from the reservoir level. A strong jump is observed at S4 (Fig. 15c), indicating the crossing of a sharp capillary fringe. While rising, the water table remains nearly horizontal in the upstream m2- and m3-materials zone but a sharp front is observed to develop across the left inner drain (m5-material), see Fig. 14. Small numerical oscillations in hydraulic head are observed in Fig. 15 but do not affect the global dynamics. Further downstream and lower within the dam, the hydraulic head simulated at S1, S2 and S3 shows the slow wetting wave propagation, with attenuation and damping (Fig. 15b). At S1, S2, S3, the water content remains constant during the whole simulation, in accordance to the sensors position below water table.

When compared to hydraulic head experimental data (see Fig. 15a), the overall time evolution and order of magnitude are well represented. Some discrepancies are observed, in particular the late increase of head which is experimentally observed on each sensor around 30 days, which tends to be underestimated by the model. However, recalling the lack of experimental control evoked before, more detailed comparison should be made with caution. For the sake of comparison, the previous numerical results presented by Bonelli et al. (1993) are recalled in Fig. 15 in green lines. While the overall time evolution is rather similar between models, a better agreement with experimental data is obtained with the present simulation.

#### 5.4. Discussion

The proposed modelling strategy for Richard's equation showed very satisfactory performance in reproducing analytical and laboratory test cases and in simulating complex full-scale experiment. In this latter case, the model is able to capture automatically wetting fronts which are moving dynamically thanks to AMR. In particular, the fronts associated with heterogeneity are well resolved by the WDG technique. This is illustrated by water content distribution in Fig. 16a. Moreover, the simulation is robust since it is able to compute completely without user intervention and handle many features: degeneracy for saturated/unsaturated zones, dynamic forcing boundary condition and seepage. However, the simulation struggles to capture wetting fronts associated with nonlinear varying diffusivity resulting in severe overshoots/undershoots for the transition between saturated and unsaturated zones in very permeable materials. Further insight on this behaviour can be discussed using the Péclet number framework. Richards' equation is an elliptic-parabolic equation predominantly diffusive, it shares properties with convectiondiffusion equation and can behave like them. For example, sharp wetting fronts and internal layers can be reinterpreted within this context. The Péclet number represents the ratio of the rate of advection to the rate of diffusion. This dimensionless number is useful to analyse locally the nature of the flow regime and determine whether advection or diffusion is dominant. For this study, the Péclet number is extracted from the saturation-based formulation of Richards' equation which is similar to a convection-diffusion equation with the water content  $\theta$  as single unknown. This formulation is undefined in saturated zones because hydraulic diffusivity D tending to infinity but this zone is not of interest for the Péclet number since Richards' equation is purely diffusive. Moreover, it is assumed that  $\theta \neq 0$ . This assumption is not restrictive because if water content drops to zero, nothing happens and, in practice, there is often a residual water content. The total flux can be separated into a

diffusive flux defined by hydraulic diffusivity and driven by capillarity, and an advective flux defined by hydraulic conductivity/water content and driven by gravity. The saturation-based formulation can be rewritten as:

$$\partial_t \theta - \nabla \cdot (D(\theta) \nabla \theta + \frac{\mathbb{K}(\theta) \nabla z}{\theta} \theta) = 0,$$
(50)

with the hydraulic diffusivity  $D(\theta) = \mathbb{K}(\theta) \frac{d\psi(\theta)}{d\theta} [L^2 \cdot T^{-1}].$ 

The Péclet number can therefore be defined as:

$$Pe = \frac{\mathbb{K}(\theta)L}{D(\theta)\theta} = \frac{C(\theta)L}{\theta},$$
(51)

with the capillary capacity  $C = \frac{d\theta(\psi)}{d\psi}$  [L<sup>-1</sup>] and *L* the characteristic linear dimension assimilated to an element size  $\mathfrak{h}_E$ . High Péclet numbers are clearly reached in the sharp-gradient areas in Fig. 16b, with m1-, m2- and m3-materials (i.e. drains, rip-rap and upstream shell) showing relatively high Péclet numbers, from 1 to more than 5. Consequently, water table is not straight anymore and a delay in elevation is observed upstream, see Fig. 14.

In order to overcome these limitations, an additional ad-hoc configuration, named augmented simulation, has been implemented. This one does not use WDG framework and other difference lies in the mesh and tolerances for nonlinear convergence criterion. Finer discretization is allowed in permeable material that is to say the m1-, m2- and m3materials. The mesh is also refined around water table and both for the gradient-based and estimation-based error indicators. Instead of the reference simulation, 4-order BDF is employed and hydraulic properties are relaxed with a numerical minimal value to avoid complete degeneracy. The *augmented* computation allowed to eliminate spurious oscillations from the solution thanks to a more refined mesh, which explains why WDG method is not needed. As the mesh is extremely refined, the augmented simulation takes approximately 13.5 times longer to compute (42 h 37 min) than the reference simulation (3 h 11 min). One core Intel(R) Xeon(R) CPU 5-2630 v3 with 2.40 GHz was used. Additional information and results can be found in Clément (2021).

The simulation of La Verne dam filling shows many difficult points which lead to two unwanted effects. The first one is the development of non-physical oscillations around steep wetting fronts associated with nonlinear varying diffusivity which affect solution accuracy. The other one is nonlinear convergence possible only with very small time steps so that the calculation duration is high. These challenges illustrate numerical issues of Richards' equation for heterogeneous media with highvarying diffusivity. The improvements offered by the *augmented* simulation support the idea that a series of heuristic tools (threshold values, maximum refinement level, refinement frequency, adaptive time steps) should be investigated to improve accuracy while maintaining cost-effectiveness.

#### 6. Conclusion

In this work, a new strategy has been formulated for the solution of Richards' equation on the basis of discontinuous Galerkin method and adaptive mesh refinement. The approximation can reach high-order both in space and time. Issues outlined in the introduction have been tackled and further discussed throughout the paper so the following concluding remarks can be drawn:

- The seepage boundary condition is directly incorporated into the DG weak formulation which makes its treatment natural and simple.
- The adaptive time stepping allows the nonlinear iterative solver to converge giving robustness.
- Mesh adaptation is employed to monitor spatial errors of the Richards' equation by capturing the moving wetting fronts thanks to an *a posteriori* error indicator.
- The adaptive mesh refinement is based on a block structure to surround large regions where wetting fronts move aiming to save computational time by avoiding systematic refinement. It also prepares a future parallelization.

 Oscillations in sharp wetting fronts of Richards' equation have been interpreted within the context of convection-diffusion equation. Wetting fronts are considered as internal layers due to the nonlinear varying, and possibly vanishing, diffusivity as well as material heterogeneity and anisotropy. A local Péclet number is exhibited to highlight troublesome regions. A weighted discontinuous Galerkin method is used to allow for jumps in the solution which better approximate the sharp internal layers. Mesh adaptation and the weighted framework work in synergy to capture and resolve sharp wetting fronts through the proposed *a posteriori* error indicator.

The strategy leads to mass conservative, efficient and robust solution of Richards' equation. However, the method holds heuristic parameters so that questions remain open regarding the numerical analysis, the  $\mathfrak{h}_p$ decision making, the nonlinear convergence or the error balance. All these aspects have great potential of improvements and will be investigated in future studies to optimize the solving of Richards' equation.

# **Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

# CRediT authorship contribution statement

Jean-Baptiste Clément: Conceptualization, Software, Visualization, Writing - original draft. Frédéric Golay: Software, Writing - review & editing. Mehmet Ersoy: Methodology, Writing - review & editing. Damien Sous: Supervision, Writing - review & editing.

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