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Adaptive dynamic multilevel simulation of fractured geothermal reservoirs



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ABSTRACT

An algebraic dynamic multilevel (ADM) method for fully-coupled simulation of flow and heat transport in heterogeneous fractured geothermal reservoirs is presented. Fractures are modeled explicitly using the projection-based embedded discrete method (pEDFM), which accurately represents fractures with generic conductivity values, from barriers to highlyconductive manifolds. A fully implicit scheme is used to obtain the coupled discrete system including mass and energy balance equations with two main unknowns (i.e., pressure and temperature) at fine-scale level. The ADM method is then developed to map the finescale discrete system to a dynamic multilevel coarse grid, independently for matrix and fractures. To obtain the ADM map, multilevel multiscale coarse grids are constructed for matrix as well as for each fracture at all coarsening levels. On this hierarchical nested grids, multilevel multiscale basis functions (for flow and heat) are solved locally at the beginning of the simulation. They are used during the ADM simulation to allow for accurate multilevel systems in presence of parameter heterogeneity. The resolution of ADM simulations is defined dynamically based on the solution gradient (i.e. front tracking technique) using a user-defined threshold. The ADM mapping occurs algebraically using the so-called ADM prolongation and restriction operators, for all unknowns. A variety of 2D and 3D fractured test cases with homogeneous and heterogeneous permeability maps are studied. It is shown that ADM is able to model the coupled mass-heat transport accurately by employing only a fraction of fine-scale grid cells. Therefore, it promises an efficient approach for simulation of large and real-field scale fractured geothermal reservoirs. All software developments of this paper is publicly available at https://gitlab. com/DARSim2simulator.

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1. Introduction

Geothermal energy resources are becoming more important in the transition towards renewable energy. Due to low carbon footprints [1–4] and high potential of sustainability, the demand for geothermal energy has been rising and is expected to increase further within the next decades. Accurate scalable modeling of fluid and heat transport in geothermal reservoirs

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is of paramount importance in order to fulfil both scientific and societal expectations on successful field development plans [5,6].

While maintaining field scalability, simulation models should accurately capture high contrasts in both mass and heat transport properties inside the porous media. Moreover, a large number of reservoirs contain complex geological networks of fractures and faults with a broad range of conductivities. Such extreme contrasts have significant impact on flow and heat patterns. Therefore, high fidelity representation of the physical phenomena within the heterogeneous fractured reservoirs is crucial [7]. However, while achieving this task, the computational science community faces a variety of complex challenges, as the models demand high-resolution grids to be imposed on the entire reservoir (in orders of kilometres) [8]. Non-linear behavior of the system due to strong mass-heat coupling results in poor stability and convergence. In case of multi-phase flow (e.g., high-enthalpy systems) these issues become more severe [9]. The geo-mechanical processes (i.e., elastic and plastic deformation) [10–12], reactive transport (e.g., geo-chemical interaction between the substances) [13–15] and compositional changes are other notable challenges that the system might face. Expectedly, the presence of fractures and faults [16–18] with wide heterogeneity contrasts compared to the rest of the formation significantly increases the computational complexity. These challenges introduce high demands for developing advanced simulation methods that are able to provide efficiency (i.e., applicable to real field-scale problems) while maintaining accuracy at the desired level. To address these challenges, various advanced simulation schemes have been developed.

Discrete Fracture Models (DFM) aim to represent fractures as conforming lower dimensional domains by discretizing them at the matrix cell interfaces [19–21]. On the other hand, the Embedded Discrete Fracture Model (EDFM) makes use of non-conforming grids to discretize fractures as lower dimensional objects fully independent from the matrix grid structure [22–26]. Conservative flux transfer terms are introduced between overlapping matrix and fracture elements. EDFM is applicable to provide acceptable solutions for highly conductive fractures. Even for high-conductive fractures, it leads to very small, yet unphysical, flux leakage. More importantly, EDFM cannot accurately represent flow barriers (e.g., sealing fractures and impermeable faults). In order to resolve this limitation, projection-based EDFM (pEDFM) was introduced [27] where consistent connectivities between matrix and fractures were developed. Therefore, pEDFM can be applied to fractured porous media with generic range of conductivity contrast between fracture and the hosting rock. Even with pEDFM independent grid, the size of the reservoir and density of the fractures make any fine-scale simulation approach impractical.

Multiscale Finite-Volume (MsFV) [28–33] and Multiscale Finite-Element (MsFE) [34–36] methods have been devised to accomplish efficient simulations in presence of heterogeneity, by solving coarse-scale systems, while avoiding upscaling, yet preserving fine-scale heterogeneities [37,38]. Dynamic Local Grid Refinement (DLGR) [39–47] is another method, which employs fine-scale resolution in parts of the domain where needed (i.e., moving fronts). Algebraic dynamic multilevel (ADM) method has been developed to address the multilevel multiscale challenges on equations with elliptic, parabolic and hyperbolic systems of PDEs [48]. By extending the grid refinement strategy and taking the advantages of multilevel multiscale basis functions, the ADM method is able to algebraically map the fine-scale discrete systems (either fully implicit or sequential) to a dynamic multilevel system for all unknowns. Once the system is solved at ADM resolution, it is prolonged back to fine-scale resolution. In case of fully implicit (FIM) formulation, as the system is solved at ADM resolution for all unknowns, reconstruction of conservative flux field is not required. This method has been further developed for multiphase flow in heterogeneous formations with compositional and capillary effects [49], and fractured reservoirs using EDFM scheme [52,50]. Recent developments also include multiscale finite element method for flow-heat transfer modeling using EDFM approach [37,38]. However, such method has not yet been developed for coupled mass-heat transport in fractured porous media with pEDFM method.

In this article, an Algebraic Dynamic Multilevel (ADM) method for simulation of non-isothermal single-phase flow in fractured porous media is presented. Due to the complex non-linearity of the system (strong mass-heat coupling), two sets of equations (mass and energy conservation) are solved using the FIM approach. In the energy balance equation, conduction terms are described for both fluid and rock. Projection-based EDFM (pEDFM) is employed in order to explicitly represent fractures and to provide independent gridding of matrix and fractures. Here, the applicability of the pEDFM implementation [27] has been extended to a fully generic 3D geometry where it allows for including fractures (or flow barriers) with any orientation on the Cartesian coordinate system. This is crucial for practical applications. While maintaining the geometrical flexibility of EDFM, the matrix-matrix and fracture-matrix connectivities are adjusted to account for projection of fracture plates on the matrix transmissibilities. This allows for consistent modeling of fractures with all conductivity (from high to low) compared with that of the matrix. The fine-scale discrete system including both matrix and fracture network is obtained for two main unknowns, namely, pressure and temperature. We note that local thermal equilibrium is assumed, meaning that the temperatures in fluid and rock are considered to be equal. This assumption can be recognized as safe for the majority (or perhaps all) of the geo-science applications [51] due to large contact area between the liquid and solid phases. Once the fine-scale system is discretized using pEDFM, it is mapped into a dynamic multilevel resolution grid (i.e., ADM grid). The ADM grid is constructed based on a hierarchically nested multilevel multiscale grid resolution, where locally computed multiscale basis functions are employed at each coarsening level [34,28,30]. These basis functions are computed only at the beginning of the simulation as a pre-processing step. This provides higher computational efficiency, while capturing the fine-scale heterogeneous properties accurately.

In order to construct the multilevel multiscale operators for the geothermal fractured system, different types of basis functions are computed for pressure and temperature unknowns. For pressure, as a global variable, multiscale basis functions are computed. However, temperature resembles a local variable for which constant basis functions are used. This

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distinction in computation of basis functions is preserved on a hierarchical pattern for all the coarsening levels across the domain. To capture the effect of permeable and impermeable fractures, all basis functions are fully coupled, meaning that extra basis functions are computed for fractures as well. The construction of all basis functions are done fully algebraically. In this process, the multilevel prolongation operator is assembled so that the partition of unity holds [53,54]. The multilevel finite-volume restriction operator ensures mass conservation inside the domain. At each time-step, the ADM map is obtained by using a front tracking technique to detect the region where the solution gradient is higher than a user-defined threshold. These sub-domains of sharp gradient are resolved at fine-scale level. Moreover, regions near the source terms are always kept at fine-scale resolution to maintain accuracy. The rest of the domain is resolved at hierarchically nested coarser resolution. This ADM map is able to adapt itself to the moving front dynamically and efficiently, while capturing the physics of interest in both flow and heat transport with an accuracy set by the user-defined threshold. The Newton linearization scheme is employed to find nonlinear solutions iteratively. Once the ADM solution is found, it is prolonged back to the fine scale using prolongation operators.

Various homogeneous and heterogeneous test cases are considered and their numerical results are presented to demonstrate the accuracy and applicability of the devised method. Furthermore, its sensitivity towards fractures with different ranges of conductivity is studied.

This article is organized as follows. The governing equations and FIM approach are described in Section 1. The fine-scale discretization using pEDFM is covered in Section 2. The ADM method as the simulation strategy for fractured geothermal reservoirs is presented in Section 3. The test cases and their numerical results are shown in Section 4. The paper is concluded in Section 5.

2. Fine-scale equations and solution strategy

2.1. Governing equations

In this section, the two main governing equations are described. Here, the temperatures of fluid and rock are assumed to be identical.

2.1.1. Mass balance

Mass balance equation for non-isothermal single-phase fluid flow in porous media with n_{frac} discrete embedded fractures reads

$$\frac{\partial}{\partial t} \left(\phi^m \rho_f \right) - \nabla \cdot \left(\rho_f \frac{1}{\mu_f} \mathbf{K}^m \cdot \nabla p^m \right) = \rho_f q^{mw} + \sum_{i=1}^{n_{\text{frac}}} \rho_f^* \mathcal{Q}^{mf_i}, \quad \text{on} \quad \Omega_m \subseteq \mathfrak{R}^n, \tag{1}$$

for the rock matrix (m) and

$$\frac{\partial}{\partial t} \left(\phi^{f_i} \rho_f \right) - \nabla \cdot \left(\rho_f \frac{1}{\mu_f} \mathbf{K}^{f_i} \cdot \nabla p^{f_i} \right) = \rho_f q^{f_i w} + \rho_f^* \mathcal{Q}^{f_i m} + \sum_{j=1}^{n_{\text{frac}}} \left(\rho_f^* \mathcal{Q}^{f_i f_j} \right)_{j \neq i},$$
on $\Omega_{f_i} \subset \mathfrak{R}^{n-1} \quad \forall i \in \{1, ..., n_{\text{frac}}\}, \quad (2)$

for the lower dimensional fracture (f_i) . Here, the main unknown is the pressure p. Moreover, ϕ and \mathbf{K} are the porosity and permeability of each medium. \mathbf{K} is a tensor to account for a generic anisotropic case. Superscripts m, f_i and w correspond to matrix, fracture i and well, respectively. Subscripts f and r denote fluid and rock. ρ and μ are the density and viscosity of the fluid respectively. In addition, q^{mw} and q^{f_iw} are the source terms (i.e., wells) on matrix m and fracture f_i . Moreover, \mathcal{Q}^{mf_i} and \mathcal{Q}^{f_im} are the flux exchange between matrix m and overlapping fracture f_i corresponding to the grid cells where overlap occurs. $\mathcal{Q}^{f_if_j}$ is the flux exchange from j-th fracture to the i-th fracture on the intersecting elements. This means that the mentioned flux exchange terms are non-zero only where matrix-fracture overlap or fracture-fracture intersection exists. Due to mass conservation, $\iiint_V \mathcal{Q}^{mf_i} dV = - \iint_{A_{f_i}} \mathcal{Q}^{f_im} dA$, and $\iint_{A_{f_i}} \mathcal{Q}^{f_if_j} dA = - \iint_{A_{f_j}} \mathcal{Q}^{f_jf_i} dA$ hold.

The Peaceman well model is used to obtain the well source terms for matrix

$$q^{mw} = \frac{PI \cdot \Lambda^* \cdot (p^w - p^m)}{\Lambda V} \tag{3}$$

and fractures

$$q^{f_i w} = \frac{PI \cdot \Lambda^* \cdot (p^w - p^{f_i})}{\Delta A}.$$
(4)

Here, *PI* is the well productivity index and Λ^* is the effective mobility ($\Lambda = K/\mu$) between the well and the penetrated grid cell in the medium. ΔV and ΔA are the control volume and control area used in the discrete system for matrix *m* and

fracture f_i respectively. The flux exchange terms Q^{mf_i} , Q^{f_im} (matrix-fracture connectivities) and $Q^{f_if_j}$ (fracture-fracture connectivities) are written as:

$$\mathcal{Q}^{mf_i} = C I^{mf_i} \cdot \Lambda^* \cdot (p^{f_i} - p^m) \tag{5}$$

$$\mathcal{Q}^{mf_i} = CI^{f_i m} \cdot \Lambda^* \cdot (p^m - p^{f_i}) \tag{6}$$

$$\mathcal{Q}^{f_i f_j} = \mathcal{C} I^{f_i f_j} \cdot \Lambda^* \cdot (p^{f_j} - p^{f_i}), \tag{7}$$

where *C1* denotes the connectivity index between each two non-neighboring elements. As an example, the connectivity index between *i*-th matrix element and *j*-th fracture element is calculated as $CI_{ij} = \frac{A_{ij}}{\langle d \rangle_{ij}}$, where A_{ij} is the area fraction of fracture cell *j* overlapping with matrix cell *i* and $\langle d \rangle_{ij}$ is the average distance between these cells.

2.1.2. Energy balance

Assuming local equilibrium, energy balance on the entire domain reads

$$\frac{\partial}{\partial t} \left((\rho \, U)_{eff}^{m} \right) - \nabla \cdot \left(\rho_{f} H_{f}^{m} \frac{1}{\mu_{f}} \mathbf{K}^{m} \cdot \nabla p^{m} \right) - \nabla \cdot \left(\mathbf{\Lambda}_{eff}^{m} \cdot \nabla T^{m} \right) = \\ = \rho_{f} H_{f}^{m} q^{mw} + \sum_{i=1}^{n_{\text{frac}}} \rho_{f}^{*} H_{f}^{*} \mathcal{Q}^{mf_{i}} + \sum_{i=1}^{n_{\text{frac}}} \mathcal{R}^{mf_{i}}, \quad \text{on} \quad \Omega_{m} \subseteq \mathfrak{R}^{n}, \quad (8)$$

in the rock matrix (m) and

$$\frac{\partial}{\partial t} \left((\rho U)_{eff}^{f_i} \right) - \nabla \cdot \left(\rho_f H_f^{f_i} \frac{1}{\mu_f} \mathbf{K}^{f_i} \cdot \nabla p^{f_i} \right) - \nabla \cdot \left(\mathbf{\Lambda}_{eff}^{f_i} \cdot \nabla T^{f_i} \right) = \\
= \rho_f H_f^{f_i} q^{f_i w} + \rho_f^* H_f^* \mathcal{Q}^{f_i m} + \sum_{j=1}^{n_{\text{frac}}} \left(\rho_f^* H_f^* \mathcal{Q}^{f_i f_j} \right)_{j \neq i} + \mathcal{R}^{f_i m} + \sum_{j=1}^{n_{\text{frac}}} \left(\mathcal{R}^{f_i f_j} \right)_{j \neq i}, \\
\text{on} \quad \Omega_{f_i} \subseteq \mathfrak{R}^{n-1} \quad \forall i \in \{1, ..., n_{\text{frac}}\}, \quad (9)$$

in the lower dimensional fracture (f_i). Here, beside pressure p, the second main unknown is T as the temperature in both fluid and solid rock. (ρU)_{eff} is the effective property defined as

$$(\rho U)_{eff} = \phi \rho_f U_f + (1 - \phi) \rho_r U_r,$$
(10)

where U_f and U_r denote the specific internal energy in fluid and rock respectively. Distinctly, $(\rho U)_{eff}^m = \phi^m \rho_f U_f + (1 - \phi^m)\rho_r U_r$ and $(\rho U)_{eff}^{f_i} = \phi^{f_i}\rho_f U_f + (1 - \phi^{f_i})\rho_r U_r$. Moreover, H_f is the specific fluid enthalpy. The three mentioned terms can be expressed as non-linear functions of pressure and temperature. Λ_{eff} is the effective thermal conductivity of the saturated rock defined as

$$\Lambda_{eff} = \phi \Lambda_f + (1 - \phi) \Lambda_r. \tag{11}$$

Here, Λ_f and Λ_r are the thermal conductivities in fluid and rock, respectively. The subscripts f and r indicate fluid and solid rock. Note that, $\Lambda_{eff}^m = \phi^m \Lambda_f + (1 - \phi^m) \Lambda_r$ and $\Lambda_{eff}^{f_i} = \phi^{f_i} \Lambda_f + (1 - \phi^{f_i}) \Lambda_r$. Lastly, \mathcal{R}^{mf_i} and $\mathcal{R}^{f_i m}$ are the conductive heat flux exchange between matrix m and the overlapping fracture f_i . $\mathcal{R}^{f_i f_j}$ denotes the conductive heat flux exchange from j-th fracture to the i-th fracture where the intersection occurs. Similar to mass flux exchange, the conductive flux exchange terms are non-zero only for the existing matrix-fracture overlaps or fracture-fracture intersections. $\iiint \mathcal{R}^{mf_i} dV =$

$$-\iint_{A_{f_i}} \mathcal{R}^{f_i m} dA, \text{ and } \iint_{A_{f_i}} \mathcal{R}^{f_i f_j} dA = -\iint_{A_{f_j}} \mathcal{Q}^{f_j f_i} dA \text{ hold as well to honor the conservation of energy.}$$

To obtain the conductive heat flux exchanges, i.e., \mathcal{R}^{mf_i} , \mathcal{R}^{f_im} (matrix-fracture connectivities) and $\mathcal{R}^{f_if_j}$ (fracture-fracture connectivities), the embedded discrete scheme is used, i.e.,

$$\mathcal{R}^{mf_i} = C I^{mf_i} \cdot \mathbf{\Lambda}^*_{eff} \cdot (T^{f_i} - T^m)$$
(12)

$$\mathcal{R}^{mf_i} = CI^{f_i m} \cdot \mathbf{\Lambda}^*_{eff} \cdot (T^m - T^{f_i}) \tag{13}$$

$$\mathcal{R}^{f_i f_j} = C I^{f_i f_j} \cdot \Lambda_{eff}^* \cdot (T^{f_j} - T^{f_i}), \tag{14}$$

where Λ_{eff}^* is obtained as harmonically-averaged property between the two non-neighboring elements. *C1* is identical to the connectivity index used in Eq. (5).



Fig. 1. The independent matrix 3D grid and the fractures 2D grids are shown. Note that each domain has its own grid and that any fracture orientation can be considered.

2.1.3. Correlations

The following correlations are used to compute certain fluid and rock properties.

Fluid viscosity: Viscosity as a function of temperature is calculated via the following correlation [55],

$$\mu_f(T) = 2.414 \times 10^{-5} \times 10^{\frac{247.8}{T-140}}.$$

Fluid density: The density of fluid is a function of pressure and temperature, defined as [51]

$$\rho_f(P, T) = \rho_{fs}(T) [1 + c_w(T) (P - P_s)],$$

where $P_S = 1$ bar. $c_w(T)$ and $\rho_{fs}(T)$ are computed from the following empirical correlations [8,56]

$$c_w(T) = (0.0839 T^2 + 652.73 T - 203714) \times 10^{-12}$$

$$\rho_{fs}(T) = -0.0032 T^2 + 1.7508 T + 757.5.$$

Fluid Entalphy: Fluid enthalpy is obtained via [51]

$$H_f(P,T) = u_{ws} + C_{pf}(T-T_s) + \frac{P}{\rho_f},$$

where $u_{ws} = 420000 \text{ J/kg}$.

2.2. Fine-scale discrete system using pEDFM

The coupled system of non-linear equations described above, i.e., Eqs. (1)-(9) with two main unknowns (i.e., p, T) is discretized spatially using two-point-flux-approximation (TPFA) finite-volume scheme and temporally using the backward (implicit) Euler scheme [57]. The discretization results in set of structured grids for a three-dimensional (3D) porous medium with 2D fracture planes included that are generated independently. A visual example of the gridding structure is shown in Fig. 1. The obtained non-linear FIM system is solved using the Newton-Raphson iterative method [57].

2.2.1. Mass and heat fluxes

The mass flux exchange between each two neighboring control volumes i and j (inside one medium) using TPFA scheme can be written as

$$F_{ij} = \frac{\rho_f^*}{\mu^*} T_{ij} (p_i - p_j),$$
(15)

where $T_{ij} = \frac{A_{ij}}{d_{ij}} K_{ij}^H$ denotes the transmissibility between grid cells *i* and *j*. A_{ij} is the interface area between the two grid cells, d_{ij} is the distance between their cell centers and K_{ij}^H is the harmonic average of the permeabilities set at the interface of grid cells *i* and *j*. The properties with superscript * are obtained using the upwind scheme. Note that density ρ and viscosity μ are functions of pressure and temperature.

The mass flux between each two non-neighboring grid cells i and j (either matrix-fracture or fracture-fracture connectivities) is obtained via EDFM formulation. This flux between a matrix (denoted as m) cell i and a fracture (denoted as f) cell j is defined as



Fig. 2. Figure on the left: An illustration of a 2D fracture plate overlapping a matrix cell. In this example, the overlapping section forms an irregular hexagon. Figure on the right: A visualization of two intersecting fracture plates (left) and discrete elements of different intersecting fractures (right), with the intersection lines shown in red.

$$\mathcal{F}_{ij}^{mf} = -\mathcal{F}_{ij}^{fm} = -\frac{\rho_f^*}{\mu^*} T_{ij}^{mf} (p_i^m - p_j^f), \tag{16}$$

where the transmissibility $T_{ij}^{mf} = K_{ij}^H C I_{ij}$. K_{ij}^H is the harmonically averaged permeability between the overlapping matrix and fracture elements, and $C I_{ij}$ is the connectivity index between them. The EDFM models the matrix-fracture connectivity index as $CI_{ij} = \frac{A_{ij}^{mf}}{\langle d \rangle_{ij}}$ where A_{ij}^{mf} is the area fraction of fracture cell *j* overlapping with matrix cell *i* (see Fig. 2, left) and $\langle d \rangle_{ij}$ is the average distance between these cells [24].

The mass flux exchange between two intersecting fracture elements i (on fracture f) and j (on fracture g) is given as

$$\mathcal{F}_{ij}^{fg} = -\mathcal{F}_{ij}^{gf} = -\frac{\rho_{f}^{*}}{\mu^{*}} T_{ij}^{fg} (p_{i}^{f} - p_{j}^{g}), \tag{17}$$

where T_{ij}^{fg} is the transmissibility between the two cells obtained via a lower dimensional connectivity index formulation. Mind that the intersection of 2D fracture plates is a line and for 1D fracture line-segments it is a point. Fig. 2 (right) shows a visualization example of intersection between two non-neighboring 2D fracture elements. The intersection forms a line segment I_{ij} (shown in red color) with the average distances from the intersection segment of $\langle d \rangle_{II_{ii}}^f \neq \langle d \rangle_{II_{ii}}^g$. Thus, T_{ii}^{fg} is obtained using an harmonic-average formulation, i.e.,

$$T_{ij}^{fg} = K_{ij}^{H} \frac{CI_{il_{ij}}^{f} \times CI_{jl_{ij}}^{g}}{CI_{il_{ij}}^{f} + CI_{jl_{ij}}^{g}},$$
(18)

where, e.g., $CI_{II_{ij}}^{f}$ represents the connectivity index between the 2D fracture element *i* and the 1D intersection line segment I_{ii} , as shown in Fig. 2.

The convective heat flux exchange between the neighboring control volumes *i* and *j* reads

$$\check{F}_{ij} = \frac{\rho_f^* H_f^*}{\mu^*} T_{ij} (p_i - p_j).$$
⁽¹⁹⁾

Here, H_f^* is the enthalpy of fluid determined at the interface between grid cells *i* and *j*. One can conclude $\check{F}_{ij} = H_f^* F_{ij}$. Similarly, the convective heat flux exchange between non-neighboring elements can be obtained via multiplication of their mass flux exchange (\mathcal{F}_{ij}) by the effective enthalpy (H_f^*) determined at the intersection of two overlapping elements. Namely, $\check{\mathcal{F}}_{ij} = H_f^* \, \mathcal{F}_{ij}$.

The conductive heat flux exchange between each two neighboring cells i and j is written as

$$G_{ij} = \mathbb{T}_{cond,ij}(T_i - T_j), \tag{20}$$

where, $\mathbb{T}_{cond,ij} = \frac{A_{ij}}{d_{ij}} (\Lambda_{eff})_{ij}^{H}$ is the transmissibility between grid cells *i* and *j*. A_{ij} , d_{ij} and $(D_{eff})_{ij}^{H}$ are the interface area, the distance from the cell centers and the harmonic average of the effective conductivity set at the interface between grid cells *i* and *j* respectively.

The conductive heat flux exchange between non-neighboring elements i and j is obtained as

$$\mathcal{G}_{ij}^{mf} = -\mathcal{G}_{ij}^{fm} = -\mathbb{T}_{cond,ij}^{mf}(T_i^m - T_j^f),\tag{21}$$

for matrix-fracture connectivities and

$$\mathcal{G}_{ij}^{fg} = -\mathcal{G}_{ij}^{fg} = -\mathbb{T}_{cond,ij}^{fg} (T_i^f - T_j^g), \tag{22}$$

for fracture-fracture connectivities

Here, $\mathbb{T}_{cond,ij} = (\Lambda_{eff})_{ij}^H Cl_{ij}$, where $(\Lambda_{eff})_{ij}^H$ is the harmonic average of the effective conductivity between the overlapping matrix and fracture elements. A similar procedure as in Eqs (18) will be used to obtain the fracture-fracture connectivity indices.

At each time-step the fine-scale discrete mass balance equation reads

$$\frac{\left(\phi^{m}\rho_{f}\right)_{i}^{n+1} - \left(\phi^{m}\rho_{f}\right)_{i}^{n}}{\Delta t} + \sum_{j=1}^{N_{n}}F_{ij} + \sum_{k=1}^{n_{\text{frac}}} \left(\sum_{j=1}^{N_{f_{k}}}\mathcal{F}_{ij}^{mf_{k}}\right) = \left(\rho_{f}q^{mw}\right)_{i}, \quad \forall i \in \{1, ..., N_{m}\}$$
(23)

for element i in matrix m and

$$\frac{\left(\phi^{f_h}\rho_f\right)_i^{n+1} - \left(\phi^{f_h}\rho_f\right)_i^n}{\Delta t} + \sum_{j=1}^{N_n} F_{ij} + \sum_{j=1}^{N_m} \mathcal{F}_{ij}^{f_hm} + \sum_{k=1}^{n_{\text{frac}}} \left(\sum_{j=1}^{N_{f_k}} \mathcal{F}_{ij}^{f_hf_k}\right) = \left(\rho_f q^{f_hw}\right)_i, \quad \forall i \in \{1, ..., N_{f_h}\}$$
(24)

for element *i* in fracture f_h . N_m and N_{f_k} denote the number of elements in matrix *m* and fracture f_k respectively. N_n is the number of neighboring grid cells.

Similarly, the fine-scale discrete form of the energy balance equation at each time-step is written as:

$$\frac{\left(\left(\rho \ U\right)_{eff}^{m}\right)_{i}^{n+1} - \left(\left(\rho \ U\right)_{eff}^{m}\right)_{i}^{n}}{\Delta t} + \sum_{j=1}^{N_{n}} \check{F}_{ij} + \sum_{k=1}^{n_{\text{frac}}} \left(\sum_{j=1}^{N_{f_{k}}} \check{F}_{ij}^{mf_{k}}\right) + \\
+ \sum_{j=1}^{N_{n}} \check{G}_{ij} + \sum_{k=1}^{n_{\text{frac}}} \left(\sum_{j=1}^{N_{f_{k}}} \check{G}_{ij}^{mf_{k}}\right) = \left(\rho_{f} H_{f}^{m} q^{mw}\right)_{i}, \quad \forall i \in \{1, ..., N_{m}\} \quad (25)$$

for element i in matrix m and

$$\frac{\left(\left(\rho \ U\right)_{eff}^{f_h}\right)_i^{n+1} - \left(\left(\rho \ U\right)_{eff}^{f_h}\right)_i^n}{\Delta t} + \sum_{j=1}^{N_n} \check{F}_{ij} + \sum_{j=1}^{N_m} \check{F}_{ij}^{f_hm} + \sum_{k=1}^{n_{\text{frac}}} \left(\sum_{j=1}^{N_{f_k}} \check{F}_{ij}^{f_hf_k}\right) + \\
+ \sum_{j=1}^{N_n} \check{G}_{ij} + \sum_{j=1}^{N_m} \check{G}_{ij}^{f_hm} + \sum_{k=1}^{n_{\text{frac}}} \left(\sum_{j=1}^{N_{f_k}} \check{G}_{ij}^{f_hf_k}\right) = \left(\rho_f H_f^{f_h} q^{f_hw}\right)_i, \quad \forall i \in \{1, ..., N_{f_h}\} \quad (26)$$

for element *i* in fracture f_h . The residual form of equations (23)-(26) can be written as subtracting the left-hand side from the right-hand side in each equation. The residual of the mass balance equation for each element *i* in matrix *m* (i.e., $(r_{MB}^m)_i^{n+1}$) reads

$$(r_{MB}^{m})_{i}^{n+1} = \left(\rho_{f}q^{mw}\right)_{i} - \frac{\left(\phi^{m}\rho_{f}\right)_{i}^{n+1} - \left(\phi^{m}\rho_{f}\right)_{i}^{n}}{\Delta t} - \sum_{j=1}^{N_{n}}F_{ij} - \sum_{j=1}^{N_{m}}\mathcal{F}_{jj}^{f_{h}m} - \sum_{k=1}^{n_{frac}}\left(\sum_{j=1}^{N_{fk}}\mathcal{F}_{ij}^{f_{h}f_{k}}\right) = 0, \quad \forall i \in \{1, ..., N_{m}\}.$$
(27)

The residuals for the other three equations (i.e., the mass balance in fractures $(r_{MB}^{f_h})_i^{n+1}$, the energy balance in matrix $(r_{EB}^m)_i^{n+1}$ and the energy balance in fractures $(r_{EB}^{f_h})_i^{n+1}$) is obtained similar to Eq. (27). The full residual vector at time-step n is obtained by a vertical concatenation of the residuals per medium. Namely,

$$r^{n} = [(r_{MB}^{m})^{n}, (r_{MB}^{f_{1}})^{n}, (r_{MB}^{f_{2}})^{n}, ..., (r_{EB}^{f_{n_{\text{frac}}}})^{n}, (r_{MB}^{m})^{n}, (r_{EB}^{f_{1}})^{n}, (r_{EB}^{f_{2}})^{n}, ..., (r_{EB}^{f_{n_{\text{frac}}}})^{n}]^{T}.$$
(28)

As the residual at time-step n + 1 (r^{n+1}) is a non-linear function of pressure and temperature unknowns at time-step n + 1 (i.e., p^{n+1} and T^{n+1}), the Newton-Raphson iteration method is applied to solve the system at each time-step. In other words,

$$r^{\nu+1} = r^{\nu} + \frac{\partial r}{\partial p} \bigg|^{\nu} \delta p^{\nu+1} + \frac{\partial r}{\partial T} \bigg|^{\nu} \delta T^{\nu+1} = 0.$$
⁽²⁹⁾

Here, the superscript ν denotes the iteration index. At each Newton iteration, the linearized system of equation is written as $\mathbf{J}^{\nu} \delta x^{\nu+1} = -r^{\nu}$. In this system, \mathbf{J}^{ν} is the Jacobian matrix including all derivatives. Each block $J_{e_{\alpha}}$ contains the derivatives of equation e with respect to unknown α , i.e., $J_{e_{\alpha}} = \partial r_e / \partial \alpha$ (containing the list of, $\frac{\partial r_{MB}}{\partial p} |^{\nu}$, $\frac{\partial r_{BB}}{\partial T} |^{\nu}$, and $\frac{\partial r_{EB}}{\partial T} |^{\nu}$) and $\delta x^{\nu+1}$ is the full vector of updates to the unknowns, precisely, $\delta x^{\nu+1} = [\delta p, \delta T]^T$. Clearly, $\delta p = [\delta p^m, \delta p^{f_1}, ..., \delta p^{f_{n_{\text{frac}}}}]^T$ and $\delta T = [\delta T^m, \delta T^{f_1}, ..., \delta T^{f_{n_{\text{frac}}}}]^T$. Subscripts *MB* and *EB* refer to equations of mass balance, and energy balance. The full linear system can be illustrated as:

$$\underbrace{\begin{pmatrix} \begin{pmatrix} J_{MB_{p}}^{mm} & J_{MB_{p}}^{mf} \\ J_{MB_{p}}^{fm} & J_{MB_{p}}^{ff} \end{pmatrix}}_{\begin{pmatrix} J_{MB_{T}}^{fm} & J_{MB_{T}}^{ff} \\ J_{MB_{T}}^{fm} & J_{MB_{T}}^{ff} \end{pmatrix}^{\nu}}_{\begin{pmatrix} J_{BB_{T}}^{mm} & J_{MB_{T}}^{ff} \\ J_{BB_{p}}^{fm} & J_{EB_{p}}^{ff} \end{pmatrix}}_{\begin{pmatrix} J_{EB_{p}}^{mm} & J_{EB_{T}}^{ff} \\ J_{EB_{p}}^{fm} & J_{EB_{p}}^{ff} \end{pmatrix}} \underbrace{\begin{pmatrix} J_{BB_{T}}^{mm} & J_{BB_{T}}^{mf} \\ J_{EB_{T}}^{fm} & J_{EB_{T}}^{ff} \end{pmatrix}^{\nu}}_{J_{0}^{fm}} \underbrace{J_{EB_{T}}^{mf} & J_{EB_{T}}^{ff} \end{pmatrix}^{\nu}}_{\delta x_{0}^{\nu+1}} = -\underbrace{\begin{pmatrix} r_{MB}^{m} \\ s_{MB}^{r} \\ r_{BB}^{r} \\ r_{EB}^{r} \\ r_{0}^{\nu} \\ r_{0}^{\nu}$$

In order to achieve a converged solution, the following conditions have to be assured:

$$\frac{||r_{(MB)}||_2}{||r_{(MB)}^0||_2} < \epsilon_{(MB)} \wedge \frac{||r_{(EB)}||_2}{||r_{(EB)}^0||_2} < \epsilon_{(EB)} \wedge \frac{||\delta p||_2}{||p||_2} < \epsilon_{(p)} \wedge \frac{||\delta T||_2}{||T||_2} < \epsilon_{(T)}$$
(31)

Here, each threshold (ϵ_x) is a user-defined tolerance set initially as input at the beginning of the simulation.

To improve the convergence, at the beginning of each time-step, the mass balance and energy balance equations are solved as decoupled systems in a sequential manner. In other words, first, decoupled linear system of mass balance equation (32) is solved and a vector of pressure updates (i.e., δp) is obtained. After updating the pressure, all dependent properties and derivatives are re-calculated. Similarly, in the next step, the decoupled linear system of energy balance equation (33) is solved and a vector of temperature updates is acquired. Consequently, the temperature unknowns are updated and all dependent parameters and derivatives are recalculated. After these steps, the fully coupled linear system (30) is solved iteratively.

$$\underbrace{\begin{pmatrix} J_{MB_{p}}^{mm} & J_{MB_{p}}^{mf} \\ J_{MB_{p}}^{fm} & J_{MB_{p}}^{ff} \end{pmatrix}^{\nu}}_{\mathbf{J}_{BB_{p}}^{mp}} \underbrace{\begin{pmatrix} \delta p^{m} \\ \delta p^{f} \end{pmatrix}^{\nu+1}}_{\mathbf{J}_{BB_{p}}^{mm}} = -\underbrace{\begin{pmatrix} r_{MB}^{m} \\ r_{MB}^{f} \end{pmatrix}^{\nu}}_{r_{MB_{\nu}}^{r_{MB_{\nu}}}} \\ \underbrace{\begin{pmatrix} J_{EB_{T}}^{mm} & J_{EB_{T}}^{mf} \end{pmatrix}^{\nu}}_{\mathbf{J}_{EB_{T}}^{fm}} \underbrace{\begin{pmatrix} \delta T^{m} \\ \delta T^{f} \end{pmatrix}^{\nu+1}}_{\mathbf{J}_{EB_{T}}^{fm}} = -\underbrace{\begin{pmatrix} r_{EB}^{m} \\ r_{EB} \end{pmatrix}^{\nu}}_{r_{EB_{\nu}}^{r_{EB}}} \end{aligned}$$
(32)

The extension to pEDFM is explained in section 2.2.2.

Obtaining the solution of this linearized system of equations is computationally challenging due to the strong coupling of mass and heat transfer, especially for real field-scale applications. In section 3 the simulation framework of this work is presented with which resolving the computational challenge is aimed using the ADM method.

2.2.2. pEDFM implementation

To correct for EDFM limitation on fractures with generic conductivity due to parallel transmissibilities, the matrix-matrix, fracture-matrix and fracture-fracture connectivities are modified in the overlapping regions. The mentioned modifications eliminate the parallel transmissibilities, such that pEDFM is applicable to any conductivity contrast between the matrix and fractures. Initially, all connectivities between two neighboring matrix cells that are disconnected due to the overlapping fractures are detected. Due to geometrical algorithm devised during the development of this method, a continuous projection path (visible in Fig. 3 as solid lines in light blue color) is automatically obtained on the interfaces as such it disconnects the neighboring connections letting the flux occur only on one route (i.e., through matrix-fracture-matrix).

Let fracture element f overlap matrix grid cell i (as shown in Fig. 3) with an area fraction A_{if} . A set of projections is defined on the interface between the overlapped matrix grid cell i and its neighboring grid cells (in orange) that are affected by the crossing (i.e., j and k). Please note that in the 3D dimensional case, there will be three projections. For each dimension (i.e., x, y and z) the projection area fractions are obtained via:



Fig. 3. pEDFM illustration for a 2D matrix with structured grid and a 1D overlapping fracture. The overlapped matrix cells are highlighted in yellow color. With classical EDFM method, these cells are connected to each overlapping fracture element via the connectivity index in the matrix-fracture flux exchange. pEDFM introduces extra non-neighboring connections between fracture elements and matrix cells highlighted in orange.

$$A_{if\perp x_e} = A_{if} \times \cos(\gamma), \quad x_e \in \{x, y, z\},\tag{34}$$

where γ is the angle between the fracture element and the interface connecting the matrix grid cell *i* and the neighboring grid cell in the corresponding dimension (shown in Fig. 3 on the zoomed-in section and highlighted in red color as $A_{if \perp x}$ and $A_{if \perp y}$). New transmissibilities are defined to connect the fracture element *f* to each non-neighboring matrix grid cells (i.e., *j* and *k* in the 2D example shown in Fig. 3):

$$T_{i_ef} = \frac{A_{if \perp x_e}}{\langle d \rangle_{i_ef}} \Lambda_{i_ef}, \quad x_e \in \{x, y, z\},$$
(35)

where, $\langle d \rangle_{i_e f}$ is the average distance between the fracture element f and matrix grid cell i_e . $\Lambda_{i_e f}$ is the effective fluid mobility between these cells. As a result of the new transmissibilities, the connectivity between the matrix grid cell i and its corresponding neighboring cells is modified:

$$T_{ii_e} = \frac{A_{ii_e} - A_{if \perp x_e}}{\Delta x_e} \Lambda_{i_e f}, \quad \mathfrak{X} \in \{x, y, z\}.$$
(36)

For simplicity of the implementation, the modified transmissibilities are obtained by multiplication of coefficient α as a fraction of the projected cross-section area, and the cross-section area of the corresponding interface. Please note that for all overlapping fracture elements (except for the boundaries of the fractures), the projection will cover the whole interface. Therefore, α is 1.0 for most of the cases, resulting in zero transmissibility between the matrix grid cells (i.e., $T_{ii_e} = 0$), thus removing the parallel transmissibilities [27].

3. ADM method for flow and heat transfer

3.1. Solution strategy

At the beginning of every Newton iteration, the fine-scale fully-implicit linearized system of equations from Eq. (30) is used to algebraically assemble a reduced linear system on a dynamic multilevel grid resolution which is defined at the beginning of each time-step. At the first stage (i.e., obtaining the dynamic multilevel grid), sets of N_m^l and $N_{f_i}^l$ hierarchically nested grids are imposed on matrix *m* and fracture f_i . Please note that *l* refers to the coarsening level at which the corresponding Cartesian grid is specified. Here, the fine-scale level is referred to as l = 0, meaning N_m^0 and $N_{f_i}^0$ grid cells are defined at fine-scale level for matrix and fracture respectively. It is assumed that the fine-scale resolution is a sufficiently accurate representation of the domain to capture the physical phenomena. The coarsening ratio (i.e., the ratio between the resolutions of level *l* and l - 1 is written as

$$\gamma_m^l = \frac{N_m^{l-1}}{N_m^l} \quad \text{and} \quad \gamma_{f_i}^l = \frac{N_{f_i}^{l-1}}{N_{f_i}^l}$$
(37)

for matrix *m* and fracture f_i . The coarsening ratios γ_l are constant on the hierarchical set of coarsening levels *l*, but both properties are defined independently for matrix and each fracture, providing practical flexibility. Combination of grid cells at different resolutions results in construction of the ADM grid. To map from the fine-scale resolution to this dynamic multilevel resolution, ADM employs a series of restriction (**R**) and prolongation (**P**) operators, which are constructed and computed only at the beginning of the simulation (see the offline stage of Fig. 4). The ADM system is written as



Fig. 4. Schematic description of ADM strategy.

$$\underbrace{\hat{\mathbf{R}}_{l}^{l-1}\dots\hat{\mathbf{R}}_{1}^{0}\mathbf{J}_{0}\hat{\mathbf{P}}_{0}^{1}\dots\hat{\mathbf{P}}_{l-1}^{l}}_{\mathbf{J}_{ADM}}\delta\hat{\mathbf{x}}_{ADM} = -\underbrace{\hat{\mathbf{R}}_{l}^{l-1}\dots\hat{\mathbf{R}}_{1}^{0}r_{0}}_{r_{ADM}}.$$
(38)

In this system, $\hat{\mathbf{R}}_{l}^{l-1}$ is the so-called restriction operator which maps the sections of the solution vector ($\delta \hat{x}_{ADM}$) at level l-1 to level l. Additionally, $\hat{\mathbf{P}}_{l-1}^{l}$ is the prolongation operator mapping parts of the solution vector from level l to level l-1 [48,50]. After solving the ADM system, the solution acquired at ADM level ($\delta \hat{x}_{ADM}$) is prolonged back to fine-scale resolution to obtain an approximated solution (i.e., $\delta x'_{0}$), namely

$$\delta x_0 \approx \delta x'_0 = \hat{\mathbf{P}}_0^1 \dots \hat{\mathbf{P}}_{l-1}^l \, \delta x_{ADM}. \tag{39}$$

Note that δx_0 represents the reference fine-scale solution. The ADM Restriction ($\hat{\mathbf{R}}_l^{l-1}$) and prolongation ($\hat{\mathbf{P}}_{l-1}^l$) operators are obtained from the static multilevel multiscale restriction (\mathbf{R}_l^{l-1}) and prolongation (\mathbf{P}_{l-1}^l) operators respectively. The static prolongation operator \mathbf{P}_{l-1}^l is constructed by assembling the locally computed basis functions (see section 3.2) at level *l* and is written as

$$\mathbf{P}_{l-1}^{l} = \begin{pmatrix} [(P_{p})_{l-1}^{l}]_{mm} & [(P_{p})_{l-1}^{l}]_{mf} & 0 & 0\\ [(P_{p})_{l-1}^{l}]_{fm} & [(P_{p})_{l-1}^{l}]_{ff} & 0 & 0\\ 0 & 0 & [(P_{T})_{l-1}^{l}]_{mm} & [(P_{T})_{l-1}^{l}]_{mf} \\ 0 & 0 & [(P_{T})_{l-1}^{l}]_{fm} & [(P_{T})_{l-1}^{l}]_{ff} \end{pmatrix}_{N_{l-1} \times N_{l}}$$
(40)

Note that \mathbf{P}_{l-1}^{l} consists of two main diagonal blocks for the two unknowns (i.e., pressure *p* and temperature *T*). Within each main block, the off-diagonal terms refer to the coupling terms between matrix and fractures. The static restriction operator \mathbf{R}_{l}^{l-1} reads

$$\mathbf{R}_{l}^{l-1} = \begin{pmatrix} [R_{l}^{l-1}]_{m} & 0 & 0 & 0\\ 0 & [R_{l}^{l-1}]_{f} & 0 & 0\\ 0 & 0 & [R_{l}^{l-1}]_{m} & 0\\ 0 & 0 & 0 & [R_{l}^{l-1}]_{f} \end{pmatrix}_{N_{l} \times N_{l-1}}.$$
(41)

To assure mass conservation, a finite-volume restriction operator is used. Therefore, \mathbf{R}_{l}^{l-1} elements contain either 1 or 0 where

$$R_l^{l-1}(u, v) = \begin{cases} 1 & \text{if cell } v \text{ is inside coarser cell } u, \\ 0 & \text{otherwise.} \end{cases}$$
(42)

For the test cases that are presented in this article, different interpolators are employed for each of the variables. The prolongation operator for the pressure (i.e., $(P_p)_{l-1}^l$) uses multilevel multiscale basis functions as interpolator (see section 3.2), whereas the temperature prolongation operator contains constant basis functions, namely $(P_T)_{l-1}^l = [R_l^{l-1}]^T$ where superscript *T* refers to transpose operator. Therefore, the matrix-fracture coupling blocks of prolongation operator related to temperature (i.e., $[(P_T)_{l-1}^l]_{mf}$ and $[(P_T)_{l-1}^l]_{fm}$) are zero.

3.2. Multilevel multiscale basis functions

The pressure prolongation operator $(P_p)_{l-1}^l$ used in this work employs multiscale basis functions at each coarsening level. It contains diagonal sub-blocks $[(P_p)_{l-1}^l]_{mm}$, $[(P_p)_{l-1}^l]_{ff}$ and off-diagonal sub-blocks $[(P_p)_{l-1}^l]_{mf}$ and $[(P_p)_{l-1}^l]_{fm}$. The ADM method allows for different matrix-fracture coupling strategies for calculation of basis functions. In case of decoupled basis functions, where the fractures and matrix do not influence the local solutions of each other, the off-diagonal sub-blocks consist of only zero elements. However, by choosing the fully-coupled approach, each medium will influence the local solution (i.e., the basis functions) of the overlapping domain, hence the off-diagonal blocks contain non-zero elements as well [54,50].

The prolongation operator is constructed fully algebraically [54] and it includes the basis functions at all coarsening levels. To compute the basis functions, first the fine-scale pressure system of equations is assembled. Consequently, the coarse grids are imposed on the fine-scale computational grid cells for matrix and fractures. The approach to construct the hierarchically nested primal and dual coarse grids is flexible in this work. One option is to impose the coarse grids by determining identical primal coarse grids at each coarsening level, namely each primal coarse grid at level *l* has identical size of $\gamma^l = \gamma_x^l \times \gamma_y^l \times \gamma_z^l$. In this case the coarse nodes are not located on the corners or edges of the global domain. Fig. 5 shows an example of the primal and coarse grids of this option for a 2D domain of 75 × 75 grid including 3 fractures. The other option is to include the coarse nodes on the corners and edges of the global domain, resulting in primal coarse grids with different sizes. Fig. 6 illustrates the coarse grid construction of this approach. Please note that the level at which each of the media (either matrix or fractures) is coarsened can be defined independently.

Once the coarse grids are constructed, the basis functions are calculated algebraically for each coarsening level. The multiscale procedure converts the two-point flux approximation (TPFA) connectivities of the fine-scale system (i.e., A^0) to multi-point flux approximation (MPFA) scheme at coarse-scale systems for each coarsening level l (i.e., A^l). Therefore, prior to construction of coarse-scale system at level l, the system at level l - 1 is reduced to the TPFA scheme, namely

$$[A^{l-1}]_{TPFA} = TPFA_Operator([A^{l-1}]_{MPFA}).$$
(43)



Fig. 5. Coarse grid structure with identical primal coarse grids at each coarsening level for a 2D 75 × 75 matrix and 3 fractures. The coarsening ratio in matrix is $\gamma_{m,x}^{l} = 5$ and $\gamma_{m,y}^{l} = 5$ in x and y direction respectively. The number of coarse grid at level l = 2 is $N_{m,x}^{l} = \frac{75}{(\gamma_{m,x}^{l})^{2}} = 3$ in x direction and $N_{m,y}^{l} = \frac{75}{(\gamma_{m,x}^{l})^{2}} = 3$ in y direction.



Fig. 6. Coarse grid structure including primal coarse nodes on the corners and edges for a 2D 76 × 76 matrix and 3 fractures. The coarsening ratio in matrix is $\gamma_{m,x}^l = 5$ and $\gamma_{m,y}^l = 5$. The number of coarse grid at level l = 2 is $N_{m,x}^l = \frac{76-1}{(\gamma_{m,x}^l)^2} + 1 = 4$ and $N_{m,y}^l = \frac{76-1}{(\gamma_{m,y}^l)^2} + 1 = 4$ in x and y direction respectively.

Here, the *TPFA_Operator* removes the MPFA components from the entries of the coarse-scale system matrix. This procedure is performed independently on each medium [50]. Fig. 7 shows coupled basis functions for a matrix coarse node (left) and two fracture coarse nodes (right).

3.3. Selection of the grid resolution

The fine-scale grid resolution is mapped into the ADM grid resolution using the ADM operators. This dynamic grid resolution is obtained by a user-defined grid selection criterion as a front tracking technique. At each time-step n, the ADM grid is selected based on the solution at time-step n - 1 (explicit scheme) using a threshold set as the input parameter for the grid selection criterion. The condition for which the criterion is considered, compares the difference of the values of a main unknown (in this case, the temperature) between neighboring cells and the grid cells in a coarse node (spatial gradient of the unknown) [48]. Let Ω_l^I and Ω_l^J be the set of two neighboring coarse grid cells I and J at coarsening level l. The indices of fine-scale grid cells belonging to the coarse grid cells Ω_l^I and Ω_l^J are indicated by i and j respectively. The temperature difference $\Delta \overline{T}_{IJ}$ is obtained as

$$\Delta \bar{T}_{IJ} = max(|T_i - T_j|) \quad \forall i \in \Omega_I^I \text{ and } \forall j \in \Omega_I^J.$$

$$\tag{44}$$



(a) A matrix basis function

(b) Two basis functions of a fracture

Fig. 7. Example of coupled basis functions for a 2D homogeneous domain. The figure on the left shows the basis function of a matrix coarse node and the figure on the right illustrates the basis functions of two coarse nodes on a fracture.



Fig. 8. Example of an ADM grid for a 3D domain with one fracture plate at a specific time-step during the simulation. Two coarsening levels are employed for the matrix and one coarsening level is used in the fracture. Here, the moving front of the fluid is captured in fine-scale resolution.

The coarse grid block *I* is refined from coarsening level *l* to resolution l - 1 if the condition

$$\Delta \bar{T}_{IN} > \text{tol} \tag{45}$$

holds. Here, N refers to all coarse grid cells neighboring coarse grid cell I (at coarsening level l). Fig. 8 shows an example of a 3D domain including one fracture with two coarsening levels over matrix and one coarsening level over fracture. Please note the independency of coarsening strategy and grid selection procedure for matrix and fracture. It can be realized that the moving front of the flooding fluid will be captured with higher resolution as it corresponds to sharper gradients. Moreover, the grid cells around the wells are always kept at fine-scale resolution to guarantee accurate capture of source term fluxes.

4. Numerical results

Numerical results are presented in this section. First, the pEDFM model is validated. Thereafter, ADM results are compared against fine-scale simulations, used as a reference.

4.1. Test case 1: validation of pEDFM

This test case demonstrates the validation of pEDFM implementation. For this purpose, a 2D homogeneous domain of 3 m × 3 m is considered (see Fig. 9). Two fractures intersect in the middle of the domain forming a cross-shape. Both fractures are 1.5 [m] long with aperture of 8×10^{-3} [m]. In terms of fractures conductivity, two scenarios are taken into account. In the first scenario, fractures are $a_f = 10^8$ times more permeable than the rock matrix ($K_m = 10^{-14}$ m² and



Fig. 9. Visualization of test case 1 geometry, a 2D domain with a cross-shaped fracture network at the center.

Input parameters	of test case	1 for fluid	and rock	properties.

Table 1

Property	Value	
Rock thermal conductivity (Λ_r)	4 [W/mK]	
Fluid thermal conductivity (Λ_f)	negligible	
Rock density (ρ_r)	2750 [kg/m ³]	
Fluid specific heat (C_{p_f})	4200 [J/kgK]	
Rock specific heat (C_{p_r})	790 [J/kgK]	
Matrix porosity (ϕ)	0.2	
Matrix permeability	$10^{-14} [m^2]$	
High-perm fractures permeability	10 ⁻⁶ [m ²]	
Low-perm fractures permeability	10 ⁻²² [m ²]	
Fractures aperture	$8 \times 10^{-3} \ [m]$	

 $K_f = 10^{-6} \text{ m}^2$) and in the second scenario, fractures are 10^8 times less permeable than the rock matrix ($K_m = 10^{-14} \text{ m}^2$ and $K_f = 10^{-22} \text{ m}^2$). The initial pressure and temperature of the reservoir are $p_0 = 1.5 \times 10^7$ [pa] and $T_0 = 400$ [K] respectively. Cold water with pressure $p_{inj} = 2 \times 10^7$ [pa] and temperature $T_{inj} = 400$ [K] is injected from the left boundary and hot water is produced from the right boundary with pressure $p_{prod} = 10^7$ [pa]. No-flow boundary conditions applies to top and bottom boundaries. Table 1 shows the input parameters of this test case.

To obtain the reference solution (denoted as DNS), a 375 × 375 grid resolution is imposed on the domain. This allows for fully resolving the flow inside the fracture. In this case fractures are defined as channels along the middle row and column of the discretized domain. EDFM and pEDFM simulations are run with four different matrix gridding resolution of 75 × 75, 45 × 45 and 25 × 25. At all runs, the fractures aperture is identical and set to be $a_f = 8 \times 10^{-3}$ [m] which is approximately equal to the matrix grid cell size of DNS discretization. The size of grid cells inside the fractures have similar dimensions as matrix grid cells. In cases with highly conductive fractures, the simulation is run for 6 [hours] and in the cases with low-permeability fractures, the simulation is run for 12 [hours]. The results are provided at 50 isochronal intervals.

In the first scenario, fractures are considered to be highly conductive (see Table 1 for more details). The results of the simulations as plots of pressure and temperature solutions are shown in the Fig. 10 and 11.

The plots in Fig. 12 present the pressure and temperature errors as functions of the simulation time. The error of variable x (i.e., pressure or temperature), denoted as e_x , is calculated via

$$e_{x} = \frac{||x_{\text{DNS}} - x_{\text{EDFM/pEDFM}}||_{2}}{||x_{\text{DNS}}||_{2}}.$$
(46)

At the center of the domain where the fractures intersect, a difference in the temperature distribution between the DNS, EDFM and pEDFM methods can be observed. Please note that the difference arises from the discretization approaches. In DNS method, fractures are actually channels with aperture of one grid cell. As a result, those matrix grid cells are flooded with the cold fluid. This is not the case for EDFM and pEDFM.

In the second scenario, both fractures are set to be 10^8 times less permeable than the matrix rock. Figs. 13 and 14 show the simulation results of this scenario as pressure and temperature solutions respectively.

Fig. 15 shows the pressure and temperature errors versus the simulation time for the second scenario. As the EDFM is incapable of capturing the low permeable fractures and the results are not accurate and representative, the errors are only shown for pEDFM.



Fig. 10. Test case 1: High-permeable fractures: comparison of pressure solution between the fully resolved DNS (single plot at left column) using 375×375 grid cells, pEDFM results (top row) and EDFM results (bottom row). Both pEDFM and EDFM are run with different grid resolutions (left to right: 75×75 , 45×45 and 25×25) at 25th time interval (i.e., t = 3 hours).



Fig. 11. Test case 1: High-permeable fractures: comparison of temperature solution between the fully resolved DNS, pEDFM and EDFM. Similar to previous figure, the single plot at left column shows DNS solution, top row shows the pEDFM results and bottom row shows the EDFM results.



Fig. 12. Test case 1: High-permeable fractures: Pressure and Temperature errors of pEDFM and EDFM simulations with respect to the reference solution (DNS).

4.2. Test case 2: ADM with a 2D homogeneous fractured reservoir

In this test case, the accuracy of the ADM method for mass-heat transport in a homogeneous domain containing fractures with mixed conductivities is studied. For this purpose, a 2D fractured 100 m × 100 m domain with 30 fractures is considered. The length of each fracture is different but the size of their aperture is identical and set to $a_f = 5 \cdot 10^{-3}$ m. A 136 × 136



Fig. 13. Test case 1: Low-permeable fractures: comparison of pressure solution between the fully resolved DNS (single plot at left column) using 375×375 grid cells, pEDFM results (top row) and EDFM results (bottom row). Both pEDFM and EDFM are run with different grid resolutions (left to right: 75×75 , 45×45 and 25×25) at 25th time interval (i.e., t = 3 hours).



Fig. 14. Test case 1: Low-permeable fractures: comparison of temperature solution between the fully resolved DNS, pEDFM and EDFM. Similar to previous figure, the single plot at left column shows DNS solution, top row shows the pEDFM results and bottom row shows the EDFM results.



Fig. 15. Test case 1: Low-permeable fractures: Pressure and Temperature errors of pEDFM compared to reference solution (DNS).

grid is imposed on the rock matrix and the fracture network consists of 315 grid cells (in total 18811 cells). The matrix permeability is $K_m = 10^{-14}$ m² and the permeability of the fracture network has the range of $K_{f_{min}} = 10^{-20}$ m² and $K_{f_{max}} = 10^{-7}$ m². Two injection wells are located at the bottom left and top left corners with injection pressure of $p_{inj} = 2 \cdot 10^7$ Pa. Additionally, there are two production wells at the bottom right and the top right corners with pressure of $p_{prod} = 1 \cdot 10^7$ Pa. Table 2 demonstrates the input parameters of this test case.

The coarsening level for matrix is $l_m = 2$ and for fractures is $l_f = 1$. Over the entire computational grids, the coarsening ratio is $\gamma = 3 \times 3$. The ADM simulations are run for three different thresholds of coarsening criterion, namely,

 Table 2

 Input parameters of test case 2 for fluid and rock properties.

Property	Value	
Rock thermal conductivity (Λ_r)	4 [W/mK]	
Fluid thermal conductivity (Λ_f)	negligible	
Rock density (ρ_r)	2750 [kg/m ³]	
Fluid specific heat (C_{p_f})	4200 [J/kgK]	
Rock specific heat (C_{p_r})	790 [J/kgK]	
Matrix porosity (ϕ)	0.2	
Matrix permeability	$10^{-14} [m^2]$	
fractures permeability (min)	$10^{-20} [m^2]$	
fractures permeability (max)	$10^{-7} [m^2]$	
Fractures aperture	5×10^{-3} [m]	



Fig. 16. Test case 2: pressure plots. Fig. 16a illustrates the fracture network of this test case. The figure on top right is the fine-scale solution (16b), and the figures on the bottom row show the ADM solution with thresholds of $\Delta T = 5$ (16c), $\Delta T = 10$ (16d) and $\Delta T = 20$ (16e) respectively.

 $\Delta T = 5,10,20$ [°K] and the ADM results are compared to result of fine-scale simulation as reference solution. The ADM error for pressure and temperature is obtained via

$$e_x = \frac{||x_{\rm FS} - x_{\rm ADM}||_2}{||x_{\rm FS}||_2} \tag{47}$$

where, *x* is either pressure or temperature and the subscripts ADM and FS denote ADM and fine-scale, respectively. The simulation is run for 1000 [days] and the results are printed at 100 isochronal intervals.

Figs. 16 and 17 show the pressure and temperature solution of fine-scale and ADM simulations. The errors (e) and the percentage of active grid cells (AGC) are mentioned under the ADM plots.

Fig. 18 provides more details of the error analysis for test case 2. On the top row pressure and temperature errors (Fig. 18a and 18b) versus simulation time-steps are shown. Fig. 18d plots the percentage of active grid cells used during the simulation. Lastly, averaged errors and average active grid cells are plotted over the entire simulation for all three thresholds.

Please note that the regions around the wells are always kept at fine-scale resolution and as a result (depending on the size of the domain) a noticeable portion of active grid cells are imposed. In this test case, approximately 5 percent of total grid cells are kept at fine-scale due to near-well refinement.

Please note that in this test case, intersection between permeable fractures and impermeable fractures occur. By default, the impermeable fractures are considered as "non-sealing" flow barriers, meaning that the crossing of the highly permeable



Fig. 17. Test case 2: temperature plots. Fig. 17b on top right shows the fine-scale solution. The ADM solutions are shown on the figures of bottom row with thresholds of $\Delta T = 5$ (17c), $\Delta T = 10$ (17d) and $\Delta T = 20$ (17e).



Fig. 18. Test case 2: error analysis. Fig. 18a and 18b show ADM error of pressure and temperature over simulation time-steps. The errors are calculated based on Eq. (47). Fig. 18c shows the percentage of active grid cells used during simulation versus the time-steps. Lastly, Fig. 18d demonstrates the averaged values of the mentioned parameters over the entire simulation time for three thresholds.



Fig. 19. Test case 3: Pressure plots for the 2D heterogeneous test case with 30 highly conductive fractures. Shown are the permeability map 19a, fine-scale solution 19b, ADM with thresholds $\Delta T = 5$ 19c, $\Delta T = 10$ 19d, $\Delta T = 20$ 19e and $\Delta T = 40$ 19f respectively.

fractures cause flow leakage. This effect can be seen in the Fig. 16 where the pressure gradient across the permeable fracture stays negligible on both sides of the impermeable fracture.

4.3. Test case 3: ADM with a 2D heterogeneous fractured reservoir

This test case, assesses the ADM performance on a 2D heterogeneous and fractured domain. Similar to the previous test case, a 2D domain with dimensions of 100 m × 100 m is considered. A network of 30 fractures with identical geometry as in test case 2 exists. The rock matrix is discretized into 136×136 fine-scale grid cells and 291 grids are imposed on fracture network (in total 18516 grid cells). The permeability of the matrix ranges from $K_{m_{min}} = 1.2 \times 10^{-15}$ m² to $K_{m_{max}} = 1.2 \times 10^{-12}$ m². All fractures are highly conductive with permeability of $K_f = 10^{-8}$ m². The well pattern used in this test case is identical to test case 2. Moreover, the same coarsening strategy is used in this test case. The input parameters of Table 2 holds for this test case as well, except for permeability values of matrix and fractures. The ADM simulations are run for four different thresholds ($\Delta T = 5$, 10, 20, 40 [°K]) their results are compared to the reference solution (fine-scale). The simulation is done for 200 [days] and the results are extracted at 100 isochronal intervals. The pressure and temperature solutions of fine-scale and ADM simulations are displayed on Fig. 19 and 20 respectively. Similar to the previous test case, the errors (*e*) and the percentage of active grid cells (*AGC*) are indicated under the ADM plots.

Fig. 21 shows the error analysis for test case 3. Pressure and temperature errors versus simulation time-steps are illustrated on the top row (Fig. 21a and 21b). Fig. 21c shows the percentage of active grid cells during the simulation. Average errors and active grid cells over the entire simulation time can be seen for all four ADM thresholds on Fig. 21d.

4.4. Test case 4: ADM with a 3D homogeneous fractured reservoir

In this test case, a 3D 100 m × 100 m × 40 m containing 15 lower dimensional fractures with different geometrical properties is considered. An 81 × 81 × 27 Cartesian grid is imposed on rock matrix and fracture network is discretized in to 1377 grid cells (summation of 178524 grid cells). The rock matrix has permeability of $K_m = 10^{-14}$ m². Fracture network consists of both highly conductive fractures with permeability of $K_f = 10^{-6}$ m² and flow barriers with permeability of $K_f = 10^{-22}$ m². Fig. 23a shows the structure of fracture network for this 3D test case. Two injection wells exist on the bottom left and top left boundaries with pressure of $p_{inj} = 5 \cdot 10^7$ Pa. Similarly, two production wells are located at the bottom right and top right boundaries with pressure of $p_{prod} = 1 \cdot 10^7$ Pa. All wells are vertical and perforate the entire thickness of the reservoir. Simulation is run for 200 days and the results are printed at 100 isochronal intervals. Two coarsening levels with coarsening ratio of $\gamma_m = 3 \times 3 \times 3$ is considered for matrix and one coarsening level with coarsening ratio of $\gamma_m = 3 \times 3 \times 3$ is considered for matrix and one coarsening level with coarsening ratio of $\gamma_f = 3 \times 3$ for fractures. Fig. 22 illustrates the ADM grid at fine-scale

Simulations are run for fine-scale and ADM with four different thresholds ($\Delta T = 10, 20, 40, 80$ [°K]). The results of ADM are compared to the reference solution (fine-scale). The temperature solutions are shown in Fig. 23.

Fig. 24 provides the error analysis of this test case.



Fig. 20. Test case 3: Temperature plots for the 2D heterogeneous test case with 30 high conductive fractures. Shown in these plots are the permeability map (20a), fine-scale solution (20b), ADM with thresholds $\Delta T = 5$ (20c), $\Delta T = 10$ (20d), $\Delta T = 20$ (20e) and $\Delta T = 40$ (20f) respectively.



Fig. 21. Test case 3: error analysis. Fig. 21a and 21b show ADM error of pressure and temperature over simulation time-steps. The errors are calculated based on Eq. (47). Fig. 21c shows the percentage of active grid cells used during simulation versus the time-steps. Lastly, Fig. 21d demonstrates the averaged values of the mentioned parameters over the entire simulation time for three thresholds.



Fig. 22. Test case 4: Illustration of the ADM grid at time interval 10 (t = 20 [days]) for $\Delta T = 40$ [°K]. Note that the grid cells near the wells are always kept at fine-scale resolution.

5. Conclusions

An algebraic dynamic multilevel (ADM) method for fully-implicit simulation of single-phase thermal flow in fractured heterogeneous porous media using pEDFM is presented. First, the pEDFM model [27] is extended to account for 3D fracture geometries. The fine-scale fully implicit system resulting from geothermal-pEDFM model is then mapped into a dynamic multilevel (i.e., ADM) grid, defined for matrix and fractures independently. Such map is obtained using sequences of ADM prolongation and restriction operators which are assembled from the static multilevel multiscale operators. The static operators are calculated only at the beginning of the simulation after the construction of the coarse grids for both matrix and fractures on all coarsening levels. The prolongation operator is constructed based on locally computed basis functions. Important to note is that the matrix-fracture coupling is taken into account for multiscale basis functions. The ADM grid selection strategy with front tracking technique employs the fine-scale grid cells only where and when needed.

Fine-scale simulation for a 2D geothermal system allows to investigate the pEDFM and EDFM accuracy, both compared to the direct numerical solution (DNS). Both highly conductive and flow-barrier fractures are considered. It is shown that in the presence of highly conductive fractures, pEDFM and EDFM performed accurate, though EDFM still allows for small unphysical leakage. However, EDFM fails to capture the low conductive fractures, while pEDFM shows a good representation of DNS with satisfactory accuracy.

Moreover, numerical results are obtained for 2D and 3D geothermal reservoirs to investigate ADM performance. The ADM results on these test cases are compared to those obtained from fine-scale simulations. The sensitivity of ADM to different grid coarsening criteria is also studied. It is observed that ADM is capable of providing accurate simulations by employing only a fraction of the fine-scale grid cells in the sub-domains where it is needed. Due to the rarefaction of the temperature distribution (highly diffused temperature at the front), more fine-scale grid cells are used at the temperature front depending on the threshold. Additionally, ADM provides an algebraic framework which brings a scalable simulation method for thermal fluid flows. One can expect that by increasing the size of the domain, the percentage of active grid cells used during simulation reduces. Therefore, ADM introduces a promising simulation strategy for real-field geothermal reservoir simulations.

All software developments of this work is made available open source at https://gitlab.com/DARSim2simulator.

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.

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(a) Reservoir geometry and fracture network

(b) Fine-scale



(c) $\Delta T = 10$ $err = 1.6 \times 10^{-3}, AGC = 58\%$



(d) $\Delta T = 20$ err = 2.9 × 10⁻³, AGC = 54%



Fig. 23. Test case 4: temperature plots. Fig. 23b on top right shows the fine-scale solution. The ADM solutions are presented on the figures of middle and bottom row with thresholds of $\Delta T = 10$ (23c), $\Delta T = 20$ (23d), $\Delta T = 40$ (23e) and $\Delta T = 80$ (23f). For sake of visualization, only the grid cells with temperature values of 300 [°K] < ΔT < 350 [°K] illustrated. On these plots, the ADM grid belonging to fine-scale resolution are shown with gray color, but the ADM grid of coarser levels are hidden for better visualization. The temperature distribution corresponds to simulation time interval 25 (t = 50 [days]). The figure on the top left (23a) shows the fracture network including the discrete fracture plates.

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(c) Active grid cells

(d) Average values

Fig. 24. Test case 4: error analysis. Fig. 24a and 24b present ADM error of pressure and temperature versus simulation time-steps. The errors are calculated using Eq. (47). Fig. 24c shows the percentage of active grid cells versus simulation time-steps. Finally, Fig. 24d demonstrates the averaged values of the mentioned parameters over the entire simulation time for four thresholds.

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