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Parallel Framework for Complex Reservoir Simulation with Advanced Discretization and Linearization Schemes

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Abstract

The continuous progress of reservoir monitoring technology provides encouraging capacities to reduce uncertainties in the subsurface characterization and to mitigate risks in field development applying the reservoir simulation approach. However, it is always challenging to take full advantage of the observation data, since an accurate representation of strong heterogeneities requires a high-resolution grid. Most of the discretization methods cannot handle full tensor permeability, and high nonlinearity introduced by complex physical process drastically reduces simulation efficiency. In this work, we develop an advanced parallel framework for reservoir simulation with the implementation of state of the art discretization and linearization methods. We apply the multipoint flux approximation (MPFA) method to handle the full tensor permeability in unstructured grids. To keep the fidelity of the geological model and improve computational efficiency, we use massively parallel computations via Message Passing Interface (MPI). Complex subsurface physics is described by mass-based formulations making the framework flexible for general-purpose reservoir simulation. However, the representation of phase behavior introduces additional workload when compared with the phase-based formulations in the traditional approach. Here, we apply the Operator-Based Linearization (OBL) approach which not only overcomes this drawback but also turns it to an advantage. In this method, the conservation equations are described in an operator form. By constructing a library of tabulated operators, the repeated work spent on complex phase behavior and property evaluation can be significantly reduced. We benchmark the parallel framework with analytical solutions under singlephase flow and multiphase flow. The results demonstrate that the parallel framework provides accurate simulation results for structured and unstructured grids. We validate that MPFA implemented in our parallel framework converges to real solutions when the permeability is a full tensor. Besides, several realistic cases have been rigorously tested confirming high computational capacity, efficiency, and accuracy of the advanced massively parallel framework for general-purpose reservoir simulation. With the implementation of MPFA and OBL approaches, the parallel framework is fully equipped for the simulation of problems with full tensor permeability, high-heterogeneities, and complex physical processes.

Introduction

With the development of reservoir monitoring technology, more valuable data has been observed during the exploration and development stages. Assisted with reservoir simulation, this data can help to reduce uncertainties in the subsurface characterization and mitigate risks in field development. However, it is always challenging to integrate detailed information into reservoir simulation. For example, limited by the computational efficiency, a highly heterogeneous geological model can not be applied for reservoir simulation of complex physical processes (e.g. compositional simulation) directly.

The most popular approach to reducing the number of degrees of freedom is a coarsening of geological models which increases numerical errors inevitably. However, another challenge may be introduced after the upscaling procedure due to the appearance of full tensor permeability. The most commonly used two-point flux approximation (TPFA) method can not provide convergent solutions for reservoirs with full permeability tensor [Abushaikha et al., 2017; Hjeij and Abushaikha 2019a and b]. Also, the solutions using the TPFA method are not always numerically convergent for general unstructured grid cases [Abushaikha et al., 2020]. Besides, the complex physics of multiphase flow makes it hard for the development of a reservoir simulator and enhances the nonlinearity of the system which can drastically reduce simulation efficiency. Thus, an accurate and computationally efficient reservoir simulator is still strongly required.

To overcome the limitations introduced by TPFA and to have a better understanding of the flow behavior in the subsurface domain, several advanced numerical methods have been developed in reservoir simulation [Aavatsmark et al., 1998; Aavatsmark, 2002; Abushaikha et al., 2015; Abushaikha et al., 2018; Hjeij et al. 2019a; Hjeij et al. 2019b; Zhang et al., 2019a; Zhang et al., 2019b]. Among them, the O-method of multipoint flux approximation (MPFA-O) has been proved to be an efficient and convergent method. The flow behavior in strong heterogeneities [Abushaikha et al., 2008; Li et al., 2014; Wang et al., 2019; Li et al., 2018a; Wu et al., 2018] and the effect of phase behavior on flow response [Voskov et al. 2008; Voskov et al. 2009; Li et al., 2016; Liu et al., 2018; Li et al., 2018b] have also been widely discussed.

In this work, we introduce an advanced parallel framework for reservoir simulation. We apply the MPFA-O method on a general unstructured grid and compare results with the TPFA approach. We demonstrate that our framework can provide convergent solutions for the cases with full tensor permeability and unstructured grid. For high-fidelity geological models, we use massively parallel computations via Message Passing Interface (MPI) which can improve the simulation efficiency significantly. To make the framework more general, we apply the mass-based formulations which unify the single-phase, dead oil, black oil, and the compositional models.

For the solution of the nonlinear governing equations, we apply the state of art linearization method named as operator-based linearization (OBL) [Voskov, 2017; Khait et al., 2017; Khait et al., 2018a; Khait et al., 2018b]. With the application of OBL, the phase behavior computation workload can be drastically reduced. Also, it simplifies the programming of Jacobian assembly which in turn improves the flexibility of the simulation framework. We benchmark the numerical solutions with analytical solutions under single-phase flow and multiphase flow conditions. The results demonstrate that the parallel framework developed in this work is capable to provide accurate and efficient solutions for reservoir simulation. The simulations of a highly heterogeneous geo-model on different numbers of processors demonstrate the strong scalability of the parallel framework.

Parallel Framework for Reservoir Simulation

In this section, we describe the governing equations, discretization method, linearization method, and the implementation of massively parallel computation.

Modelling approach

Assuming that there are n_c components and n_p phases in an isothermal system, the transport equations can be written as below.

$$\frac{\partial}{\partial t} \left(\phi \sum_{j=1}^{n_p} x_{cj} \rho_j s_j \right) + \operatorname{div} \sum_{j=1}^{n_p} x_{cj} \rho_j \mathbf{u}_j + \sum_{j=1}^{n_p} x_{cj} \rho_j \widetilde{q}_j = 0, c = 1, \dots, n_c.$$
(1)

Here, ϕ is the reservoir porosity; *t* is the time; subscript *c* is the index for mass components; subscript *j* is the index for phases; x_{cj} is the mole fraction of component *c* in phase *j*; ρ is the phase molar density; *s* is the saturation; \tilde{q} is the phase rate per unit volume. \mathbf{u}_j is the Darcy velocity of phase *j*:

$$\mathbf{u}_{j} = -\left(\mathbf{K}\frac{k_{rj}}{\mu_{j}}\nabla P\right), \quad j = 1, \dots, n_{p}$$

$$\tag{2}$$

where **K** is the permeability tensor; k_{rj} is the relative permeability of phase *j*; μ is the viscosity; *P* is the pressure.

There are two kinds of formulations, including mass-based and phase-based formulations, used for solving the system of governing equations. In this study, to unify the formulations of the single-phase, dead oil, black oil, and the compositional models, we apply the mass-based formulations of which the unknowns are pressure and overall compositions $z_c = \sum_{j} x_{cj} \rho_j s_j / \sum_{j} \rho_j s_j$.

Multipoint flux approximation method

As a common discretization method, the TPFA method is widely used in many reservoir simulators. However, it is limited to provide an accurate solution for the simulations which apply unstructured grid or full tensor permeability. To handle these two situations, by using functions related to geometry calculation in INMOST (Terekhov et al., 2019), we apply the MPFA-O method for discretization. As shown in Figure 1, there are six control volumes in black bold lines and two intersection volumes in red lines. x_1 , x_2 , x_3 , x_4 , x_5 , and x_6 are the centers of cell 1, cell 2, cell 3, cell 4, cell 5, and cell 6; x_1^- , x_2^- , x_3^- , x_4^- , x_5^- , x_6^- and $x_7^$ are the midpoints of edges. The fluxes through half-edges including x_1x_7 , x_3x_7 , $x_7x_2^-$ and $x_7x_4^-$ are computed through intersection volume $x_1x_2x_4x_3$, and the fluxes through half-edges including x_2x_8 , x_6x_8 , $x_8x_5^-$ and $x_8x_7^-$ are computed through intersection volume $x_3x_4x_6x_5$.



Figure 1—The schematic of control volumes and intersection volumes

As shown in Figure 2, $v_1^{(1)}$ is the normal on the connection line between x_1 and x_3 , the superscript denotes the index of cell 1; \mathbf{n}_1 , \mathbf{n}_2 , \mathbf{n}_3 , and \mathbf{n}_4 are the normal vectors on the half-edges. In the sub-region of the intersection volume, falls inside cell 1, the gradient of potential can be written as:

grad
$$P = \frac{1}{2F_1} \left[\mathbf{v}_1^{(1)} \left(\overline{P}_1 - P_1 \right) + \mathbf{v}_2^{(1)} \left(\overline{P}_3 - P_1 \right) \right],$$
 (3)



b) Half-edge normals of control volumes

a) Half-edge normals of intersection volume

Figure 2—Normal vectors in an interaction volume

where F_1 is the area of the triangle $x_1x_1\overline{x_3}$.

The flux through half-edge is represented by $f_i^{(k)}$ where *i* is the index of the edge and *k* is the index of the cell. Take the cell 1 for example, the fluxes through half-edges can be written as:

$$\begin{bmatrix} f_1^{(1)} \\ f_3^{(1)} \end{bmatrix} = -\begin{bmatrix} \Gamma_1 \mathbf{n}_1^T \\ \Gamma_3 \mathbf{n}_3^T \end{bmatrix} \mathbf{K}_1 \operatorname{grad} P = -\frac{1}{2F_1} \begin{bmatrix} \Gamma_1 \mathbf{n}_1^T \\ \Gamma_3 \mathbf{n}_3^T \end{bmatrix} \mathbf{K}_1 \begin{bmatrix} \mathbf{v}_1^{(1)} & \mathbf{v}_2^{(1)} \end{bmatrix}_{-\frac{1}{P_3 - P_1}}^{-\frac{1}{P_3 - P_1}}$$
(4)

where \mathbf{K}_1 is the permeability of cell 1 in full tensor format; Γ is the length of half-edge.

By defining G_1

$$G_{1} = \frac{1}{2F_{1}} \begin{bmatrix} \Gamma_{1} \mathbf{n}_{1}^{T} \\ \Gamma_{3} \mathbf{n}_{3}^{T} \end{bmatrix} \mathbf{K}_{1} \begin{bmatrix} \mathbf{v}_{1}^{(1)} & \mathbf{v}_{2}^{(1)} \end{bmatrix} = \frac{1}{2F_{1}} \begin{bmatrix} \Gamma_{1} \mathbf{n}_{1}^{T} \mathbf{K}_{1} \mathbf{v}_{1}^{(1)} & \Gamma_{1} \mathbf{n}_{1}^{T} \mathbf{K}_{1} \mathbf{v}_{2}^{(1)} \\ \Gamma_{3} \mathbf{n}_{3}^{T} \mathbf{K}_{1} \mathbf{v}_{1}^{(1)} & \Gamma_{3} \mathbf{n}_{3}^{T} \mathbf{K}_{1} \mathbf{v}_{2}^{(1)} \end{bmatrix},$$
(5)

equation (4) can be written as

$$\begin{bmatrix} f_1^{(1)} \\ f_3^{(1)} \end{bmatrix} = -G_1 \begin{bmatrix} -P_1 \\ -P_1 \\ -P_3 - P_1 \end{bmatrix}.$$
(6)

By using the same approach, we can obtain matrixes G_2 , G_3 , and G_4 . The fluxes through half-edges in each cell can be written as:

$$\begin{bmatrix} f_1^{(1)} \\ f_3^{(1)} \end{bmatrix} = -G_1 \begin{bmatrix} -P_1 \\ -P_1 \\ -P_3 - P_1 \end{bmatrix} \begin{bmatrix} f_1^{(2)} \\ f_4^{(2)} \end{bmatrix} = -G_2 \begin{bmatrix} P_2 - P_1 \\ -P_4 - P_2 \end{bmatrix}$$

$$\begin{bmatrix} f_2^{(3)} \\ -P_3 \\ -P_3 \end{bmatrix} = -G_3 \begin{bmatrix} -P_2 - P_3 \\ -P_3 \\ -P_3 \end{bmatrix} \begin{bmatrix} f_2^{(4)} \\ -P_4 \\ -P_4 \end{bmatrix} = -G_4 \begin{bmatrix} P_4 - P_2 \\ -P_4 \\ -P_4 \end{bmatrix}$$

$$(7)$$

By assuming that the flux on the interface of two neighboring cells is continuous, equation (7) can be rewritten as:

$$f_{1} = -g_{1,1}^{(1)} (P_{1} - P_{1}) - g_{1,2}^{(1)} (P_{3} - P_{1}) = g_{1,1}^{(2)} (P_{1} - P_{2}) - g_{1,2}^{(2)} (P_{4} - P_{2})$$

$$f_{2} = g_{1,1}^{(4)} (\overline{P}_{2} - P_{4}) + g_{1,2}^{(4)} (\overline{P}_{4} - P_{4}) = -g_{1,1}^{(3)} (\overline{P}_{2} - P_{3}) + g_{1,2}^{(3)} (\overline{P}_{3} - P_{3})$$

$$f_{3} = -g_{2,1}^{(3)} (\overline{P}_{2} - P_{3}) + g_{2,2}^{(3)} (\overline{P}_{3} - P_{3}) = -g_{2,1}^{(1)} (\overline{P}_{1} - P_{1}) - g_{2,2}^{(1)} (\overline{P}_{3} - P_{1})$$

$$f_{4} = g_{2,1}^{(2)} (\overline{P}_{1} - P_{2}) - g_{2,2}^{(2)} (\overline{P}_{4} - P_{2}) = g_{2,1}^{(4)} (\overline{P}_{2} - P_{4}) + g_{2,2}^{(4)} (\overline{P}_{4} - P_{4})$$
(8)

By defining $\mathbf{f} = [f_1, f_2, f_3, f_4]^T$, $\mathbf{P} = [P_1, P_2, P_3, P_4]^T$, $\mathbf{P} = [P_1, P_2, P_3, P_4]^T$, we can obtain two equations:

$$\mathbf{f} = \mathbf{C}\mathbf{P} + \mathbf{F}\mathbf{P}.\tag{9}$$

$$\mathbf{AP} = \mathbf{BP}.\tag{10}$$

By combining equations (9) and (10), we can obtain:

$$\mathbf{f} = \mathbf{T}\mathbf{P}, \quad \mathbf{T} = \mathbf{C}\mathbf{A}^{-1}\mathbf{B} + \mathbf{F}. \tag{11}$$

As **T** is computed based on intersection volume, it can only be used to compute the fluxes through halfedges. To simplify the Jacobian assembly, we integrate the **T** of intersection volumes as connection-based transmissibility vector. For example, in Figure 1, we get a vector $\mathbf{T}_{integ} = [T_1 T_2 T_3 T_4 T_5 T_6]$ for the connection between cell 3 and cell 4. With the vector of pressure $\mathbf{P} = [P_1 P_2 P_3 P_4 P_5 P_6]^T$, we get the flux through edge x_7x_8 by $flux_{x,x_8} = \mathbf{T}_{integ} \cdot \mathbf{P}$. Thus, by applying the MPFA-O method to discretize the mesh and applying backward Euler approximation to discretize in time, the governing equations can be transformed as:

$$\left(V\phi\sum_{j=1}^{n_p} x_{cj}\rho_j s_j\right)^{n+1} - \left(V\phi\sum_{j=1}^{n_p} x_{cj}\rho_j s_j\right)^n - \varDelta t\sum_l \left(\sum_{j=1}^{n_p} x_{cj}^l \rho_j^l \lambda_j^l \mathbf{T}_{inneg}^l \mathbf{P}\right) + \varDelta t\sum_{j=1}^{n_p} x_{cj}\rho_j q_j = 0, c = 1, \dots, n_c.$$
(12)

Here *V* is the volume of a control volume; *l* denotes the edges/faces of a control volume; $\lambda_j^l = (k_{rj} / \mu_j)^l$ is the mobility of phase *j* over the interface *l* by upstream weighting, $q_j = q_j V$ is the source of phase *j*; *n*+1 is the current time step; *n* is the previous time step.

Operator-based linearization method

As an unconditionally stable method, the implicit method has been implemented in many reservoir simulators. However, it is always challenging to construct the Jacobian matrix, especially when the phase behavior is complex. In general, the main challenges are introduced by properties and their derivatives. During a simulation run, properties may be computed multiple times for the same state of the system (similar values of nonlinear unknowns). The repeated work reduces the simulation efficiency a lot.

There are three ways to determine the derivatives including numerical approach, straightforward handdifferentiation approach, and automatic differentiation techniques. The numerical approach is quite flexible but often fails to offer a robust solution and can be expensive for multicomponent systems [Pruess et al., 1999; Pruest, 2004]. The straightforward hand-differentiation approach is the most accurate strategy and is implemented in many commercial simulators. However, the ensemble of the Jacobian becomes quite complex which usually reduces the flexibility to add/change governing physics in the simulation framework [Cao, 2002]. The automatic differentiation techniques [Voskov 2012; Zaydullin et al. 2014; Garipov et al. 2016; Garipov et al. 2018] are proved to provide robust solutions and help to keep the flexibility of a reservoir simulator. However, the automatic differentiation usually introduces an overhead and limits the efficiency of a reservoir simulator.

In this work, we apply the state-of-the-art operator-based linearization (OBL) method proposed by Voskov (2017). By using the OBL, equations (12) are transformed in an operator format:

$$r_{c}\left(\boldsymbol{\xi},\boldsymbol{\omega},\mathbf{u}\right) = V\left(\boldsymbol{\xi}\right)\phi_{0}\left(\boldsymbol{\xi}\right)\left[\alpha_{c}(\boldsymbol{\omega})-\alpha_{c}(\boldsymbol{\omega}_{n})\right] - \sum_{i} \Delta t \mathbf{T}_{imteg}^{l} \mathbf{P}\boldsymbol{\beta}_{c}^{l}\left(\boldsymbol{\omega}\right) + \theta_{c}\left(\boldsymbol{\xi},\boldsymbol{\omega},\mathbf{u}\right), c = 1, \dots, n_{c},$$
(13)

where, the operators are defined as:

$$\alpha_c \! \left(\omega \right) = \left[1 + c_r \! \left(P - P_{ref} \right) \right] \! \sum_{j=1}^{n_p} \! x_{cj} \rho_j s_j, \tag{14}$$

$$\beta_c \! \left(\omega \right) = \sum_{j=1}^{n_p} \! x_{cj}^l \rho_j^l \lambda_j^l, \tag{15}$$

$$\theta_c \left(\boldsymbol{\xi}, \boldsymbol{\omega}, \mathbf{u} \right) = \Delta t \sum_{j=1}^{n_p} x_{cj} \rho_j q_j \left(\boldsymbol{\xi}, \boldsymbol{\omega}, \mathbf{u} \right).$$
(16)

Here, P_{ref} is the reference pressure for the porosity ϕ_0 . From equations (14) to (16), α_c and β_c are only dependent on the phase and rock properties and independent of spatially distributed properties. The term θ_c can also be separated into space-dependent and state-dependent operators, but considering the complexity of the well conditions, we do not show it here. Therefore, the most severe nonlinearity and complexity related to phase behavior and property calculation are introduced by α_c , β_c and their derivatives.

In the OBL, the parameter space of the nonlinear unknowns of α_c and β_c are discretized using a uniform grid. Once a status falls inside a hypercube with certain coordinates, we compute the operators on the vertices of this hypercube. Then, the operators and their derivatives inside the hypercube can be determined by interpolation. To save the workload spent on operator computation, the operators on each node are only computed single time and stored. The robustness and reliability of the OBL approach have already been proofed in multiple numerical studies (Voskov, 2017; Khait and Voskov, 2017; 2018a, b). In addition, the utilization of the OBL approach simplifies the programming of the linearization stage which makes the simulator more general and flexible.

Implementation of massively parallel computation

As an efficient and promising approach to improve computational performance, the massively parallel computation is chosen as a base technology for our reservoir simulation framework. To utilize that, we should first assign workload to the processors on cluster nodes evenly. As schematically shown in Figure 3a, a geo-model with 8×8 grids is divided into four sub-domains through reservoir domain decomposition. Then we determine shared cells in blue (Figure 3b) and ghost cells in red (Figure 3c) in each sub-domain. By assigning the grids in each sub-domain to a unique processor, the assignment of the workload is completed. Next, during a simulation run, to integrate the workload of each sub-domain, we apply the Message Passing Interface (MPI) to exchange the information among different processors by sharing the variables of shared cells with corresponding ghost cells in neighboring processors.



Figure 3—The grids for parallel simulation with MPFA-O

Numerical results

In this section, we first benchmark the numerical solutions with analytical solutions in single-phase flow to validate the accuracy of the parallel framework. Second, to demonstrate the performance of the OBL method, we perform a benchmark in two-phase flow. In the end, we test the parallel framework with field cases on structured and unstructured grids.

Benchmark of single-phase flow

Here, we will benchmark the numerical solutions with analytical solutions under a single-phase transient flow state. We apply two kinds of grids to meshing a homogeneous geo-model with a cubic size of $1 \times 1 \times 1$ m. First, we mesh the domain with four resolutions based on hexahedra shown in Table 1. Second, by splitting the hexahedra grid into six tetrahedrons grids, we generate an unstructured grid. The rock and fluid are incompressible; the reservoir is anisotropic with a full tensor permeability shown below:

$$\mathbf{K} = \begin{bmatrix} 10 & 0.5 & 0.0 \\ 0.5 & 10 & 0.5 \\ 0.0 & 0.5 & 10 \end{bmatrix}.$$
(17)

Table 1—Grid dimension and grid size

Grid dimensions	8×8×8	16×16×16	32×32×32	64×64×64
Grid size/m	0.125	0.0625	0.03125	0.015625

The transport equation can be written as:

$$\nabla \cdot (\vec{\mathbf{u}}) = f, \tag{18}$$

where f is a force term constraint by an analytical solution:

$$P_a = 10 + \sin(\pi x) \sin(\pi y) \sin(\pi z) e^{-t}.$$
(19)

By defining a function shown in equation (20), we can investigate the accuracy of the numerical solutions.

$$L_{2-imteg} = \int_{t=0}^{T} L_2 dt, \quad L_2 = \sqrt{\sum_{i=1}^{N} V_i (P_{a-i} - P_{n-i})}.$$
 (20)

Where V_i is the volume of the *i*th grid; P_{a-i} is the analytical pressure solution of the *i*th block; P_{n-i} is the numerical pressure solution of the *i*th block; N is the number of blocks.

Through a detailed sensitivity analysis, we find that the optimum values of simulation time and timestep are equal to 5 days and 0.1 days. Prolongation of simulation time or chopping of timestep will barely change the $L_{2-integ}$. Finally, we compare the numerical solutions of TPFA and MPFA-O methods using four grid resolutions. From Figure 4, we can see that with the mesh refinement, the solutions of the MPFA-O method converge to a real solution while the TPFA method fails to converge. It demonstrates that our simulator is capable to provide accurate solutions for reservoir simulation in unstructured and full tensor permeability domains.



Figure 4—L_{2-integ} analysis of the MPFA-O and TPFA methods

Benchmark of multiphase flow

In this section, we will benchmark numerical solutions of two-phase flow with analytical solutions of the Buckley-Leverett equation. Here, the reservoir size is equal to $10 \times 1 \times 1$ m; the rock and fluid are incompressible; the permeability is 1 mD; the porosity is 0.3; the reservoir is saturated with oil; the viscosities of oil and water are equal to 2 and 1 cP respectively; the relative permeabilities are defined as $k_{ro} = s_o^2$ and $k_{rw} = s_w^2$; the resolution of the parameter space of each nonlinear known is set as 64. We inject water from the left side and produce oil from the right side with a constant rate equaling 0.001 m³/day and simulation time equaling 1000 days. We apply hexahedra to meshing the geo-model with grid dimensions equaling to $1024 \times 5 \times 5$.

The numerical solutions for pressure and water saturation are shown in Figure 5 and Figure 6. From Figure 6, we can observe a shock of the water saturation. And numerical solution approaches to the real solution which is obtained by solving the Buckley-Leverett equation. By changing the parameters of rock and fluid in the model, we can investigate the effect of the properties on flow behavior which helps us further understand the flow underground.



Figure 5—Pressure distribution of the numerical solution



Figure 6—Water saturation of the numerical and analytical solutions

Next, we perform simulations with different grid resolutions including $64 \times 5 \times 5$, $128 \times 5 \times 5$, $256 \times 5 \times 5$, and $512 \times 5 \times 5$. By defining a function shown in equation (21), we can investigate the effect of grid size on simulation accuracy. As shown in Figure 7, with the mesh refinement, the water saturation approaches to the real solution. It demonstrates that the OBL method is capable to provide accurate and convergent solutions for multiphase flow.



Figure 7—The effect of grid size on the numerical solution

$$L_2 = \sqrt{\sum_{i=1}^{N} V_i (S_{w-a-i} - S_{w-n-i})}.$$
(21)

Where S_{w-a-i} is the analytical water saturation solution of the i^{th} block; S_{w-n-i} is the numerical water saturation solution of the i^{th} block.

Field cases

Since we are still working on a high-performance linear solver [Nardean et al., 2019], the solver applied in this study is just a common one that constrains the simulation efficiency. Thus, in this section, we test a three-dimensional problem based on the top 20 layers of SPE10 reservoir on structured and unstructured grids, which can be taken as a reference for initial testing between the two discretization schemes, and also investigate the scalability of the new parallel simulator.

Case 1. The permeability is shown in Figure 8. The rock and fluid are incompressible; the initial pressure is 300 bar; the porosity is homogeneous and equals 0.25; the initial water saturation is 0.2; the viscosities of oil and water are equal to 2 and 1 cP respectively; the relative permeabilities are

defined as $k_{rw} = [(S_w - S_{wc})/(1 - S_{wc} - S_{or})]^2$, $k_{ro} = [(S_o - S_{or})/(1 - S_{wc} - S_{or})]^2$, $S_{wc} = 0.2$, $S_{or} = 0.2$; the resolution of the parameter space of each nonlinear known is set as 64; two wells are imposed diagonally with one injector and one producer; the bottom hole pressure of injector is 400 bar; the bottom hole pressure of producer is 200 bar; the simulation time is 5000 days.



Figure 8—Permeability of the top 20 layers of SPE10 reservoir

We test the MPFA-O and TPFA methods for the strong heterogeneous model which is meshinged by hexahedra. As shown in Figure 9, the water saturation distribution is quite complex due to the strong heterogeneities underground. The MPFA-O and TPFA methods provide the same solution since we apply a structured grid and diagonal tensor permeability.



Figure 9—Water saturation distribution after 5000 days (8 cores)

Case 2. Here, we bend the geo-model of the previous case and obtain a new model shown in Figure 10. The parameters applied are the same as the last case. From Figure 11a, we can see that the bent strong heterogeneous geo-model introduces a complex flow response and water breakthrough time is 1200 days. By observing the error between MPFA-O and TPFA in Figure 11b, we find that the water breakthrough time is delayed by using the TPFA.



Running the simulation up to 5000 days, we obtain a much more complex water saturation distribution in Figure 12. By performing this simulation on 8, 16, 32, and 64 cores, we can investigate the scalability of the new parallel simulator. As shown in Figure 13, the speedup versus the number of processors demonstrates the strong scalability of this parallel framework for reservoir simulation.



Figure 12—Water saturation distribution of the MPFA-O after 5000 days (8 cores)



Figure 13—Scalability of the new parallel framework for reservoir simulation

Conclusions

In this work, an advanced parallel framework for reservoir simulation has been developed. To make the framework flexible for complex geological models and capable to handle full tensor permeability, the MPFA-O method is applied on a general unstructured mesh.

As a general-purpose reservoir simulator, the governing equations are represented by mass-based formulations which could unify different flow models. To overcome the Jacobian assemble hussle and improve the performance of complex phase behavior evaluation, we apply state-of-the-art Operator-Based Linearization (OBL) approach. In the OBL, the governing equations are transformed into an operator form. The state-dependent operators are uniformly discretized in the parameter space of the problem. By computing the values of operators in vertices of hypercubes, we construct a tabulated representation of physics. During a simulation run, once the status falls inside a hypercube, we determine the values of operators and their derivatives by multi-linear interpolation. This way, the phase behavior and property evaluation at each node is reduced to one time only. Besides, the programming complexity or Jacobian assemble is drastically simplified.

In order to simulate results at geological scale, we use massively parallel computations via Message Passing Interface (MPI) to improve the computational efficiency. We benchmark the numerical solutions with analytical solutions of single-phase flow which demonstrates that the parallel framework is capable to provide accurate and convergent solutions for reservoir simulations. Furthermore, to validate the modelling capabilities of multiphase flow, we compare the numerical solutions with analytical solutions obtained by solving the Buckley-Leverett equation. The results demonstrate that our advanced parallel framework for reservoir simulation is capable to handle multiphase flow problems. By using a strong heterogeneous real field geo-model, we run simulations at different numbers of processors. The results demonstrate the strong scalability of the new parallel framework for reservoir simulation.

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