Large-scale Reservoir Simulations on IBM Blue Gene/Q

Hui Liu, Kun Wang, and Zhangxin Chen

University of Calgary 2500 University Dr NW, Calgary, AB, Canada, T2N 1N4 {hui,j.liu, wang30, zhachen}@ucalgary.ca

Abstract. This paper presents our work on simulation of large-scale reservoir models on IBM Blue Gene/Q and studying the scalability of our parallel reservoir simulators. An in-house black oil simulator has been implemented. It uses MPI for communication and is capable of simulating reservoir models with hundreds of millions of grid cells. Benchmarks show that our parallel simulator are thousands of times faster than sequential simulators that designed for workstations and personal computers, and the simulator has excellent scalability.

Keywords: Large-scale reservoir, simulation, Blue Gene/Q, parallel computing

1 Introduction

Nowadays, large-scale reservoir simulations are becoming more and more popular in the oil and gas industry in order to simulate complex geological models. However, when a model is large enough, a simulator may take days or even weeks to finish one run using regular workstations and personal computers. This problem can also be observed in black oil, compositional and thermal simulations. Efficient computational methods and fast reservoir simulators should be investigated.

Reservoir simulations have been studied for decades and various models and methods have been developed. Coats studied black oil, compositional and thermal models, and he also investigated numerical methods, linear solver, preconditioner, grid effects and stability issues in his publications [1,2,3,8,17,11]. Kaarstad et al. [6] implemented a parallel two-dimensional two-phase oil-water simulator, which could solve problems with millions of grid cells. Rutledge et al. [4] implemented a compositional simulator for parallel computers using the IMPES (implicit pressure-explicit saturation) method. Shiralkar et al. [5] developed a portable parallel production qualified simulator, which could run on a variety of parallel systems. Killough et al. [7] studied locally refined grids in their parallel simulator. Dogru et al. [9] developed a parallel black oil simulator, which was highly efficient and was capable of simulating models with up to one billion cells. Zhang et al. developed a scalable general-purpose platform to support adaptive finite element and adaptive finite volume methods, which was also applied to

2 Large-scale reservoir simulations on IBM Blue Gene/Q

reservoir simulations using Discontineous Galerkin method [10,12,24]. For many reservoir simulations, most of the running time is spent on the solution of linear systems. We know that the most important is to develop efficient preconditioners. Many preconditioners have been proposed, such as constrained pressure residual (CPR) methods [13,14], multi-stage methods [15], multiple level preconditioners [23] and fast auxiliary space preconditioners (FASP) [16,18]. Chen et al. studied parallel reservoir simulations and developed a family of CPR-like preconditioners for black oil simulations and compositional simulations, including CPR-FP, CPR-FPF and CPR-FFPF methods [21].

A black oil simulator has been developed based on our in-house parallel platform. The black oil model has three mass conservation equations for three components (water, gas and oil). The system is fully coupled nonlinear system, which is solved by inexact Newton-Raphson methods, and structured grids and finite difference methods are applied. The performance of the black oil simulator is studied on IBM Blue Gene/Q system using large-scale reservoir models for standard black oil model and two-phase oil-water model. Numerical experiments show that our simulator is scalable and it is capable of simulating models with hundreds of millions of grid cells.

2 Reservoir Simulation Models

The black oil model and its simplified model, two-phase oil-water model, are briefly introduced here.

2.1 Black Oil Model

The black oil model has three phases (water, oil and gas), and three components. The model assumes that there is no mass transfer between water phase and the other two phases, and gas component can exist in gas and oil phases. Oil component is also assumed that it can exist in oil phase only. The reservoir is isothermal and no energy change is considered.

The Darcy's law is applied for black oil model, which establishes a relationship between volumetric flow rates of three components and their pressure changes in a reservoir, which is described as:

$$Q = -\frac{\kappa A \Delta p}{\mu L},\tag{1}$$

where κ is the absolute permeability of rock, A is a cross-section area, Δp is the pressure difference, μ is viscosity of fluid, and L is the length of a porous medium. In three-dimensional space, the differential form of Darcy's law is:

$$q = \frac{Q}{A} = -\frac{\kappa}{\mu} \nabla p. \tag{2}$$

By combining Darcy's law, black oil model has the following mass conservation equations for each component:

$$\begin{cases} \frac{\partial}{\partial t}(\phi s_o \rho_o^o) &= \nabla \cdot \left(\frac{KK_{ro}}{\mu_o} \rho_o^o \nabla \Phi_o\right) + q_o, \\ \frac{\partial}{\partial t}(\phi s_w \rho_w) &= \nabla \cdot \left(\frac{KK_{rw}}{\mu_w} \rho_w \nabla \Phi_w\right) + q_w, \\ \frac{\partial(\phi \rho_o^g s_o + \phi \rho_g s_g)}{\partial t} &= \nabla \cdot \left(\frac{KK_{ro}}{\mu_o} \rho_o^g \nabla \Phi_o\right) + \nabla \cdot \left(\frac{KK_{rg}}{\mu_g} \rho_g \nabla \Phi_g\right) + q_o^g + q_g, \end{cases}$$
(3)

where, for phase α ($\alpha = o, w, g$), Φ_{α} is its potential, ϕ and K are porosity and permeability of a resevoir, and s_{α} , μ_{α} , p_{α} , ρ_{α} , $K_{r\alpha}$ and q_{α} are its saturation, phase viscosity, phase pressure, density, relative permeability and production (injection) rate, respectively. ρ_o^o and ρ_o^g are density of the oil component in the oil phase and the density of the solution gas in the oil phase, respectively. They have the following relations:

$$\begin{cases} \Phi_{\alpha} = p_{\alpha} + \rho_{\alpha}gz, \\ S_o + S_w + S_g = 1, \\ p_w = p_o - p_{cow}(S_w), \\ p_g = p_o + p_{cog}(S_g), \end{cases}$$
(4)

where z is reservoir depth, p_{cow} is capillary pressure between water phase and oil phase, p_{α} is pressure of phase α , and p_{cog} is capillary between gas phase and oil phase.

The properties of fluids and rock are functions of pressure and saturation. The pressures of water and gas phases are functions of oil phase pressure and saturation; see equation (4). The density of water is a function of its pressure:

$$\rho_w = \rho_w(p_w) = \rho_w(p_o, s_w),$$

and the density of the oil phase is a function of its phase pressure and the bubble point pressure:

$$\rho_o^o = \rho_o^o(p_o, p_b),$$

where p_b is bubble point pressure. The bubble point pressure is the pressure at which infinitesimal gas appears. The water viscosity μ_w is assumed to be a constant. The oil phase viscosity is a function of its pressure p_o and the bubble point pressure p_b :

$$\mu_o = \mu_o(p_o, p_b).$$

The relative permeabilities K_{rw} , K_{ro} and K_{rg} are functions of water and gas saturations S_w and S_g :

$$\begin{cases} K_{rw} = K_{rw}(S_w), \\ K_{rg} = K_{rg}(S_g), \\ K_{ro} = K_{ro}(S_w, S_g), \end{cases}$$

4 Large-scale reservoir simulations on IBM Blue Gene/Q

where K_{ro} is calculated using the Stone II formula.

For real simulations, the relative permeabilities are given by tables or analytic formulas. Other properties, such as density, viscosity and capillary pressure, have analytic formulas and they can be calculated by table input too. With proper boundary conditions and initial conditions, a close system is given. Here no flow boundary condition is assumed.

2.2 Two-Phase Flow Model

This model is a simplified model of the standard black oil model, which assumes that the reservoir has two phases, oil and water, and they are immiscible. The model is similar to black oil model, which is written as [19]:

$$\begin{cases} \frac{\partial}{\partial t}(\phi s_o \rho_o) &= \nabla \cdot \left(\frac{KK_{ro}}{\mu_o} \rho_o \nabla \Phi_o\right) + q_o \\ \frac{\partial}{\partial t}(\phi s_w \rho_w) &= \nabla \cdot \left(\frac{KK_{rw}}{\mu_w} \rho_w \nabla \Phi_w\right) + q_w. \end{cases}$$
(5)

2.3 Well Modeling

Different well constraints can be set for each active well. One commonly-used method is a sink-source model. For each perforation block m, its well rate (production or injection) $q_{\alpha,m}$ is calculated by:

$$q_{\alpha,m} = W_i \frac{\rho_{\alpha} K_{r\alpha}}{\mu_{\alpha}} (p_h - p_{\alpha} - \rho_{\alpha} \wp(z_h - z)), \tag{6}$$

where p_h is bottom hole pressure of a well, W_i is its well index, z_h is reference depth of bottom hole pressure, z is depth of the perforated grid block m, and p_{α} is phase pressure of the perforated grid block, such as oil, gas and water. W_i can be calculated by several different models. In our simulator, a Peaceman model [25] is chosen.

Many operation constraints and their combinations may be applied to each well at different time stages, such as a fixed bottom hole pressure constraint, a fixed oil rate constraint, a fixed water rate constraint, a fixed liquid rate constraint and a fixed gas rate constraint. When the fixed bottom hole pressure condition is applied to some well, its bottom hole pressure, p_h , is known and its well rate $q_{\alpha,m}$ is known if we have phase pressure of the perforated block. The constraint equation for the well is

$$p_h = c, \tag{7}$$

where c is a constant set by the user input. No known exists for this constraint.

When a fixed rate constraint is applied to a well, its bottom hole pressure is an unknown. For the fixed water rate constraint, the equation is

$$\sum_{m} q_{w,m} = q_w,\tag{8}$$

where q_w is constant. For the fixed oil rate constraint, its equation is

$$\sum_{m} q_{o,m} = q_o, \tag{9}$$

where q_o is constant and known. A well may be applied different constraints at different time period. A schedule can be set by input, in which users can set operation changes for each well.

2.4 Numerical Methods

In this paper, conservative finite difference schemes are employed to discretize these models. The inexact Newton method is employed to solve the nonlinear equations. The time term is discretized by the backward Euler difference scheme. If we let f^n represent the value of a function f at any time step n, then its derivative at time step (n + 1) is approximated by

$$\left(\frac{\partial f}{\partial t}\right)^{n+1} = \frac{f^{n+1} - f^n}{\Delta t}.$$
(10)

The space terms are discretized by cell-centered finite difference method [19]. Here if we assume d is a space direction and A is the area of the corresponding face of a grid cell, the transmissibility term $T_{\alpha,d}$ can be written as:

$$T_{\alpha,d} = \frac{KK_{r\alpha}}{\mu_{\alpha}}\rho_{\alpha}\frac{A}{\Delta d}.$$
(11)

Inexact Newton Method The nonlinear system can be represented by

$$F(x) = 0, (12)$$

where x is unknown vector, including oil phase pressure, water saturation and well bottom hole pressure. For black oil model, a gas saturation (or bubble point pressure) is also included. After linearization, a linear system, Ax = b, is obtained in each Newton iteration, where A is Jacobian matrix, x is unknown to be determined, and b is right-hand side. The standard Newton method solves the linear system accurately. However, it is computationally expensive and it is not necessary sometimes. In our implementation, the inexact Newton method is applied, whose algorithm is described in Algorithm 1.

The only difference between standard Newton method and inexact Newton method is how to choose θ_l . Usually the parameter, θ_l , for standard Newton method is fixed and small, such as 10^{-5} . The parameter, θ_l , for inexact Newton method is automatically adjusted. Three different choices are listed as follows

6 Large-scale reservoir simulations on IBM Blue Gene/Q

Algorithm 1 The inexact Newton Method

- 1: Give an initial guess x^0 and stopping criterion ϵ , let l = 0, and assemble right-hand side b.
- 2: while $||b|| \ge \epsilon$ do
- 3: Assemble the Jacobian matrix A.
- 4: Find θ_l and δx such that

$$\|b - A\delta x\| \le \theta_l \|b\|, \tag{13}$$

5: Let l = l + 1 and $x = x + \delta x$.

7: x is the solution of the nonlinear system.

[24]:

$$\theta_{l} = \begin{cases} \frac{\|b^{l} - r^{l-1}\|}{\|b^{l-1}\|}, \\ \frac{\|b^{l}\| - \|r^{l-1}\|}{\|b^{l-1}\|}, \\ \gamma \left(\frac{\|b^{l}\|}{\|b^{l-1}\|}\right)^{\beta}, \end{cases}$$
(14)

where r^{l} and b^{l} are residual and right-hand side of *l*-th iteration, respectively. The residual is defined as,

$$r^l = b^l - A\delta x. \tag{15}$$

Linear Solver If a proper matrix ordering (numbering of unknowns) is applied, the matrix A derived from each Newton iteration can be written as

$$A = \begin{pmatrix} A_{pp} & A_{ps} & A_{pw} \\ A_{sp} & A_{ss} & A_{sw} \\ A_{wp} & A_{ws} & A_{ww} \end{pmatrix},$$
(16)

where A_{pp} is the matrix corresponding to oil phase pressure unknowns, A_{ss} is the matrix corresponding to other unknowns in each grid cell, such as water saturation, gas saturation and bubble point pressure, and A_{ww} is the matrix coefficients corresponding to well bottom hole pressure unknowns, and other matrices are coupled items.

The matrix A is hard to solve in large-scale reservoir simulations. Many multi-stage preconditioners have been developed to overcome this problem, such as CPR, FASP, CPR-FP and CPR-FPF methods. The key idea is to solve a sub-problem (A_{pp}) using algebraic multi-grid methods (AMG). In this paper, the CPR-FPF method developed by Chen et al. [21] is applied. Matrix decoupling techniques are also employed, such as ABF decoupling and Quasi-IMPES decoupling.

3 Numerical Experiments

An Blue Gene/Q from IBM is employed to run reservoir simulations. The system, Wat2Q, is located in the IBM Thomas J. Watson Research Center. Each node has 32 computer cards (64-bit PowerPC A2 processor), which has 17 cores. One of them is for the operation system and the other 16 cores for computation. The system has 32,768 CPU cores for computation. The performance of each core is really low compared with Intel processors. However, the system has strong network relative to CPU performance, and the system is scalable.

3.1 Oil-water Model

The SPE10 model is described on a regular Cartesian grid, whose dimensions are $1,200 \times 2,200 \times 170$ (ft) [22]. The model has $60 \times 220 \times 85$ cells $(1.122 \times 10^6$ cells). It has one injection well and four production wells. The original model is designed for two-phase oil-water model and it has around 2.244 millions of unknowns.

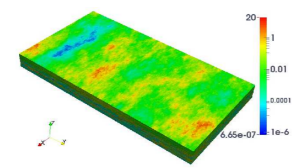


Fig. 1. Permeability in X Direction of the SPE10 benchmark

The model is highly heterogeneous. Its permeability is ranged from 6.65e-7 Darcy to 20 Darcy, and the x-direction permeability, K_x , is shown in Figure 1. Its porosity shown in Figure 2, ranges from 0 to 0.5. Data sets for porosity and permeability can be downloaded from SPE10's official website. The relative permeability of water phase is calculated by

$$K_{rw}(s_w) = \frac{(s_w - s_{wc})^2}{(1 - s_{wc} - s_{or})^2},$$
(17)

and the relative permeability of oil phase is calculated by

$$K_{ro}(s_w) = \frac{(1 - s_{or} - s_w)^2}{(1 - s_{wc} - s_{or})^2},$$
(18)

where $s_{wc} = s_{or} = 0.2$. Capillary pressure is ignored.

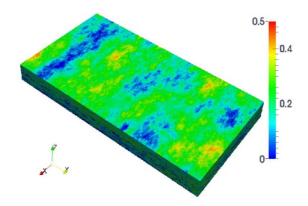


Fig. 2. Porosity of the SPE10 benchmark

3.2 Numerical Examples

Example 1. The original SPE10 project is simulated. The termination tolerance for inexact Newton method is 10^{-2} and its maximal Newton iterations are 20. The linear solver BiCGSTAB is applied and its maximal inner iterations are 50. The Quasi-IMPES decoupling strategy is used. Simulation period is 2,000 days and maximal time step is 100 days. Summaries of numerical results are shown in Table 1 [21], and its scalability is shown by Figure 3.

Table	1.	Summaries	of	Example	1
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# Pro	$\cos \# \operatorname{Ste}$	eps # New	ton # Solve	er # Avg.	solver Time (s)	Avg. time (s)
8	50	298	7189	24.1	27525.6	92.3
16	50	297	7408	24.9	13791.8	46.4
32	51	322	7467	23.1	7044.1	21.8
64	50	294	7609	25.8	3445.8	11.7

Table 1 presents results for time steps, Newton iterations, total linear iterations, average linear iterations per Newton iteration, overall running time and average running time per Newton iteration. From this table, we can see each case has similar time steps and Newton iterations. The linear solver and preconditioner are robust, where each Newton iteration terminate in around 25 linear iterations. The table and Figure 3 show our simulator has excellent scalability.

Example 2. This example tests a refined SPE10 case, and each grid cell is refined into 27 grid cells. It has around 30 millions of grid cells and around 60 millions of unknowns. The stopping criterion for the inexact Newton method is 1e-3 and

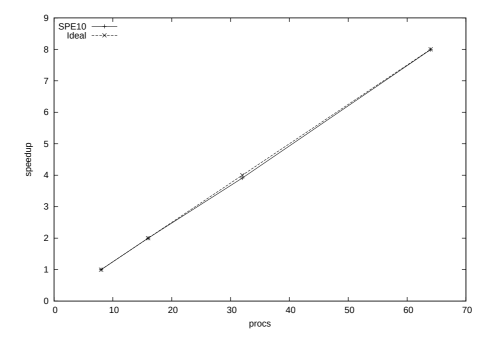


Fig. 3. Scalability of preconditioners, Example 1 $\,$

its maximal Newton iterations are 20. The BiCGSTAB solver is applied and its maximal iterations are 100. Potential reordering and Quasi-IMPES decoupling strategy are applied. The simulation period is 10 days. Up to 128 computer cards are used. The numerical summaries are shown in Table 2, and its speedup (scalability) is shown in Figure 4.

Table 2. Numerical summaries of Example	2	2
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# Proc	cs # Step	s # New	ton $\#$ Solve	er # Avg.	solver Time (s)	Avg. time (s)
128	40(1)	295	2470	8.3	43591.8	147.7
256	39	269	2386	8.8	20478.4	76.1
512	40	260	2664	10.2	10709.8	41.1
1024	39	259	2665	10.2	5578.7	21.5

The numerical summaries in Table 2 show the inexact Newton method is robust, where around 40 time steps and around 260 Newton iterations are used

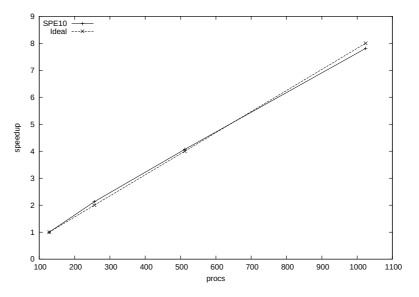


Fig. 4. Scalability of Example 2

for each simulation with different MPI tasks except the case with 128 MPI tasks due to one time step cut that contributes 20 Newton iterations. The linear solver BiCGSTAB and the preconditioner show good convergence, where the average number of linear iterations for each nonlinear iteration is between 8 and 11. The results mean our linear solver and preconditioner are effective and robust. The overall running time and average time for each Newton iteration show our simulator has excellent scalability, which is almost ideal. The scalability is also demonstrated by Figure 4.

Example 3. This example also tests a refined SPE10 case, where each grid cell is refined into 125 grid cells. It has around 140 millions of grid cells and around 280 millions of unknowns. The stopping criterion for inexact Newton method is 1e-2 and its maximal Newton iterations are 20. The BiCGSTAB solver is applied and its maximal iterations are 100. The Quasi-IMPES decoupling strategy is applied. The simulation period is 10 days. The numerical summaries are shown in Table 3, and the speedup (scalability) curve is shown in Figure 5.

This case is difficult. However, results from Table 3 show our nonlinear and linear methods are robust. Each Newton iteration terminate in less than 10 linear iterations. Running time and Figure 5 show our simulator has good scalability. For the case with 512 MPI tasks, it has super-linear scalability.

Example 4. This case simulates a refined SPE1 model, which has 100 millions of grid cells. The termination tolerance for inexact Newton method is 10^{-2} and maximal Newton iterations is 15. The BICGSTAB linear solver is chosen and its maximal iterations is 20. ABF decoupling strategy is enabled. The simulation

Table 3. Numerical summaries of Example 3

# Pro	cs # St	eps # New	ton # Solve	er # Av	g. solver Time (s) Avg. time (s)
256	57	328	2942	8.9	168619.6 514.0
512	60	328	2236	6.8	72232.4 220.2
1024	62	341	3194	9.3	43206.5 126.7
2048	59	327	3123	9.5	22588.8 69.0

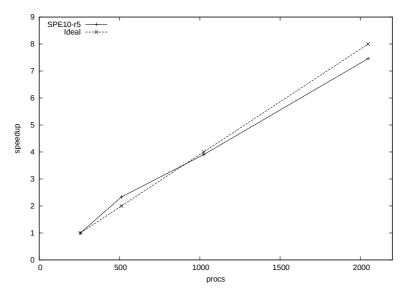


Fig. 5. Scalability of Example 3

period is 10. Summaries of numerical results are shown in Table 4 and scalability curve is shown by Figure 6.

# Proc	cs # Step	s # Newto	n # Solv	rer # Avg.	solver Time (s)	Avg. time (s)
512	27(1)	140	586	4.1	11827.9	84.4
1024	27	129	377	2.9	5328.4	41.3
2048	26	122	362	2.9	2708.5	22.2
4096	27	129	394	3.0	1474.2	11.4

Table 4. Summaries of Example 4

The original SPE1 project is a small one with 300 grid cells $(10 \times 10 \times 3)$ and the project is refined to 100 millions of grid cells $(1000 \times 1000 \times 100)$. It has a homogeneous geological model. Table 4 presents numerical summaries for nonlinear method, linear solver and performance. All four simulations use around

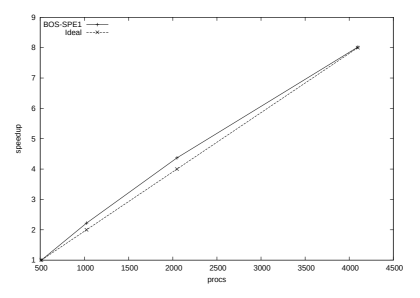


Fig. 6. Scalability (speedup) of Example 4

27 time steps. The simulation with 512 MPI tasks uses 140 Newton iterations, which is more than other cases due to one time step cut. The linear solver and preconditioner are robust, which can solve a linear system in a few iterations. Again, running time and Figure 6 show our simulator has excellent scalability.

4 Conclusion

Parallel reservoir simulations are studied in the paper, which are based on our in-house parallel platform. The platform provides grids, data, linear solvers and preconditioners for reservoir simulators. A black oil model is implemented. Numerical experiments show that our simulator has excellent scalability and simulations can be sped up thousands of times faster. The paper also demonstrates that parallel computing techniques are powerful tools for large-scale reservoir simulations and IBM Blue Gene/Q system is scalable.

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