EXTENDING THE PETREL MODEL BUILDER FOR EDUCATIONAL AND RESEARCH PURPOSES

A Thesis

by

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MASTER OF SCIENCE

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Reservoir Simulation is a very powerful tool used in the Oil and Gas industry to perform and provide various functions including but not limited to predicting reservoir performance, conduct sensitivity analysis to quantify uncertainty, production optimization and overall reservoir management. Compared to explored reservoirs in the past, current day reservoirs are more complex in extent and structure. As a result, reservoir simulators and algorithms used to represent dynamic systems of flow in porous media have invariably got just as complex. In order to provide the best solutions for analyzing reservoir performance, there is a need to continuously develop reservoir simulators and reservoir simulation algorithms that best represent the performance of the reservoir without compromising efficiency and accuracy.

There exists several commercial reservoir simulation packages in the market that have been proven to be extremely resourceful with functionality that covers a wide range of interests in reservoir simulation yet there is the constant need to provide better and more efficient methods and algorithms to study and manage our reservoirs. This thesis aims at bridging the gap in the framework for developing these algorithms. To this end, this project has both an educational and research component. Educational because it leads to a strong understanding of the topic of reservoir simulation for students which can be daunting especially for those who require a more direct experience to fully comprehend the subject matter. It is research focused because it will serve as the foundation for developing a framework for integrating custom built external simulators.
and algorithms with the workflow of the model builder of our reservoir simulation package of choice i.e. Petrel with the Ocean programming environment in a seamless manner for simulating large scale multi-physics problems of flow in highly heterogeneous flow of porous media.

Of particular interest are the areas of model order reduction and production optimization. In-house algorithms are being developed for these areas of interest and with the completion of this project. We hope to have developed a framework whereby we can take our algorithms specifically developed for areas of interest and add them to the workflow of the Petrel Model Builder.

Currently, we have taken one of our in-house simulators i.e. a two dimensional, oil-water five-spot water flood pattern as a starting point and have been able to integrate it successfully into the “Define Simulation Case” process of Petrel as an additional choice for simulation by an end user. In the future, we will expand this simulator with updates to improve its performance, efficiency and extend its capabilities to incorporate areas of research interest.
DEDICATION

To my family and friends
ACKNOWLEDGEMENTS

I want to take this opportunity to thank everyone that was instrumental directly or indirectly in helping me accomplish my research objectives. Of note is my Committee Chair, Dr. Eduardo Gildin and other committee members, Dr. Mike King and Dr. Yuefeng Sun.

I would also like to extend my gratitude to Schlumberger for giving me access to the relevant software and tools to complete this project on schedule and to the Society of Petroleum Engineering, Dallas Section for the grant awarded to me.

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CHAPTER I

INTRODUCTION AND LITERATURE REVIEW

Introduction

A Reservoir Simulator is a computer generated model that aims to represent the full geological extent and structure of a reservoir. The reservoir simulator gives the reservoir engineer the ability to study and analyze the performance of the reservoir under various operating conditions in order to adopt an efficient and maximum profit-driven development strategy for producing hydrocarbons from the reservoir.

As part of this effort to develop an appropriate management strategy, the reservoir engineer works with an interdisciplinary team made up of geologists, geophysicists, production engineers, drilling engineers and other functions to develop a reservoir management plan. Developing a reservoir management plan is a long process that first begins with adopting a strategy. According to Hoang et al. (1992), “recognizing the specific need and setting a realistic and achievable purpose are the first steps in reservoir management.” During this stage, the nature of the reservoir from its characterization i.e. geology, rock and fluid properties, fluid flow, recovery mechanisms to be implemented, drilling and well completion and past production performance are established.

Following reservoir characterization, is a study of the “total environment” to better understand the co-operate goals, culture, political stability, economic climate, social issues including safety and environmental regulations. The next stage is developing operational strategies for depleting the reservoir to recover hydrocarbons.
Data acquisition and analysis is conducted to further understand the nature of the reservoir and with the information gathered, we are able to feed inputs to geological models and numerical simulators to calibrate against historic production data and conduct prediction runs for future reservoir performance under various operating conditions. There are many stages in the reservoir management process including but not limited to strategy implementation, evaluation and monitoring but in all cases, accurate reservoir simulation is of central importance due to its connection to all phases of the management plan. In this project, we focus on the reservoir simulation stage of reservoir management.

To conduct reservoir performance prediction, simulation studies and analysis, many commercial reservoir simulation packages are available in the market. Reservoir simulation technology has gone through lots of changes from times of simplified models to give estimates of reservoir performance to reservoir simulators that are capable of capturing in great detail the structure and dynamic nature of a reservoir and provide very accurate results. Reservoir simulators are developed based on a set of partial differential equations that govern the flow of hydrocarbons in highly heterogeneous porous media. Current commercial reservoir simulators have the ability to simulate single phase and multiphase flow behavior in porous media and represent complex geology with complex geometry systems in an increasing order of billions of grid blocks in size. Although very efficient numerical solution techniques have been implemented in these industrial strength software packages, the large scale nature of many reservoir models still pose difficulty in getting results in a timely manner. This difficulty is evident especially when
one couples optimization algorithms in which several runs of forward modeling are required to complete the specific task. It is with this realization of increased system complexity that many model reduction algorithms have been developed in recent years Antoulas (2005), Gildin et al. (2006), Gildin and Lopez (2011). In this case, reduced order models can represent such large and complex reservoirs with both lower complexity and less computationally demanding models that will capture subsurface behavior and physical attributes without compromising efficiency and accuracy.

As mentioned by Hoang et al. (1992), during the reservoir characterization stage of reservoir management, very important information concerning the nature of the fluid(s) flowing, the nature of the rock and other important information are established. Reservoir simulators are designed with the capability to model the variation in structure of porous media to give a good representation of the reservoir in order to calibrate the model against past production and use as a base to conduct predictions of reservoir performance. Among these simulation packages, one can single out Petrel (Petrel Reservoir Engineering), the Pre and Post Processor of Eclipse (ECLIPSE Reservoir Engineering), which are both owned by Schlumberger. Petrel gives a user access to three types of simulators namely a black oil simulator i.e. Eclipse 100, a compositional simulator i.e. Eclipse 300 and FrontSim, the streamline simulator. As part of building simulation models, the user of Petrel has the ability to create a “simulation case” using processes in the process pane that span every domain of Petrel from Reservoir Engineering, Production Engineering, Geophysics and Geology to develop a simulation case.
The Model builder of Petrel has a lot of functionality to prepare simulation cases for reservoir simulation but as implied earlier, reservoir simulation is continually growing and so functionality that captures these developments in reservoir simulation have to be incorporated into the workflow process of developing models in model builders. New efficient algorithms to enhance performance and provide more accurate solutions are developed frequently based on the research in academia, industry and other third parties. A major part of this project is to extend the functionality of Petrel by using the Schlumberger accompanying application program interface, Ocean-in-Petrel which enables a software developer to program new processes and efficient algorithms that provide unique solutions and add them to the Petrel workflow for creating models and cases for reservoir simulation.

The objective of this project is to develop a framework for incorporating in-house developed algorithms that provide our unique solutions into the workflow of Petrel in the form of custom external reservoir simulators among the choices of simulators for simulation of reservoir simulation cases. Currently, under the supervision of my advisor and a Schlumberger Software Development team, I have achieved this goal by using my two-phase, two-dimensional oil-water 5-Spot water flood pattern simulator as a basis and implemented the equivalent external custom simulator. The outputs of the custom simulator are visualization of pressures, water and oil saturation maps at any given time along with the production rates of oil and water.

As the product of this project will be used in a classroom environment, it is our belief that this project will have immense educational value for students as it will
explore various aspects of reservoir simulation and show how models under various operating conditions perform by accepting input data and making a development strategy and finally defining a simulation case to be simulated in a timely manner. We believe that this project will serve as the foundation for a series of external custom simulators (plug-ins) in the general area of closed-loop reservoir management, model reduction and production optimization.

**Motivation**

In many scientific development and practical applications, one is faced with the task of simulating and controlling complex dynamic systems. In reservoir simulation, the key challenges are to represent oil and gas reservoirs both as realistically and accurately as possible in order to provide scenarios of prediction under various operating conditions to select an optimal development strategy for reservoir management.

A major challenge of reservoir simulation is the accurate geological representation of large scale models by numerical simulators. These simulators are mathematical models derived from first principle, conservation laws and profound knowledge of material physics as its foundation and discretized sets of partial differential equations. In order to represent these large scale reservoirs, highly accurate and detailed description induced dynamic systems of large dimensions in the state space as millions of grid blocks are often necessary in the discretization process. Direct numerical simulation of the associated large scale models leads to unmanageable large demands in computational effort often requiring massive parallel computation, complex hardware and trade-off between accuracy and complexity. To avoid the challenges of
direct simulation, proposed methods such as Upscaling as mentioned by Christie (1996), Chen et al. (2005) which is a tool used to develop geo-statistical reservoir descriptions has become increasingly popular to represent fine scale descriptions of reservoir porosity, permeability and other flow functions with coarser scale models more suitable for simulation grids.

As mentioned, model order reduction can be used to obtain fast simulation models Christie (1996), Chen et al. (2005). The Framework created in this project can be easily extended to incorporate the reduced order model when available into the Petrel software package. In this case, when these algorithms are fully developed, plug-ins that contain reduced models will be added to the workflow of the Petrel Model Builder and be made available possibly for commercial use.

**Literature Review**

Petrel is the pre and post processor of Eclipse a reservoir simulation interface that is well known in the Oil and Gas Industry. With Petrel, a user is able to develop reservoir simulation cases that aid in studying and analyzing the performance of reservoirs under various operating conditions to facilitate final decision making for reservoir management. I began my literature review by bearing in mind the sole objective of this project which is to develop a framework and foundation for extending the reservoir model builder in Petrel for reservoir simulation of defined simulation cases. Suffice to say, this project is more software development centric than it is about making considerable improvement in a particular area of research. The end product will document our way of extending the functionality of Petrel by implementing an external
custom simulator with an in-house developed two-phase, two-dimensional oil-water 5-spot water flood pattern.

For all intent and purposes, this is a literature review but in actual sense, this is the process of how I came to understand and learn how to extend the functionality of Petrel. It began by undergoing the training conducted by Schlumberger on Ocean-In-Petrel Software Development Kit, which is the application program interface through which a software developer can work in any domain of reservoir modeling such as Production Engineering, Reservoir Engineering, Geology etc. and apply their creativity and develop extensions or plug-ins to implement algorithms to add to the workflow of Petrel for model building.

In the Ocean store “www.ocean.slb.com”, which is the one stop shop to purchase plug-ins that have been developed by software developers, you will find custom solutions in different areas of interest spanning every domain of reservoir model building and analysis. This brings me back to the motivation for this project. To have a good idea of where we are going, we need to have an equally good idea of the past, what has been achieved and the areas where attention is most needed. Reservoir simulation tools come in various forms from simple mathematical models to complicated computer generated simulators for predicting reservoir behavior (Blackwell and Richardson 1971). In this project, we are more interested in computer generated simulators. After the inception of computer generated reservoir simulators, the main issues encountered involved but weren’t limited to simulation speed, computational demand, accuracy and general efficiency (Watts et al. 1997).
As computing speed and power increased, so did the ability to model subsurface geological structures with large numbers of grid blocks, complex geometry requiring complex gridding schemes. The capability of reservoir simulators has gone from being able to handle single-phase, one-dimensional models to multiphase three-dimensional models. According to Watts et al. (1997), “many of the advances in computational methods made the technical transition possible.” Some of these advances include Aronofsky and Jenkins radial gas model, Alternating-direction implicit procedure, IMPES computational method, Upstream weighting, Implicit method, Geostatistics, Upscaling, Voronoi gridding etc.

To the benefit of end users, all the afore-mentioned computational advances are included in most reservoir simulating packages today to simulate multiphase fluid models of varying dimensions, realistic geometries, well conning, compositional simulation, simulation of miscible fluid models, fractured reservoirs, integration with non-reservoir computations etc.

As computing power increased, simulation of reservoirs that were once thought impossible became more feasible. Among such reservoirs are those that require large number of grid blocks of the order of billions of grid blocks with complex geometry that impose heavy computational demands on our simulators. In the industry, to solve these large scale problems, solutions such as Upscaling have been adopted where a fine grid with properties of porosity, permeability and flow functions are represented with a system of smaller grids that maintains the structure and implicit physical nature of the
reservoir with the goal of reducing computational cost, demand and provide solutions that doesn’t compromise accuracy and efficiency.

Another proposed solution is model order reduction which is a counterpart of upscaling which also aims to represent the finer grid model with a coarser grid model with the main distinguishing factor being the algorithms implemented to reduce to a system of smaller states. Model order reduction and production optimization is a key area of interest and like other areas in reservoir simulation that have undergone breakthroughs over the years, our research team is focusing on making strides in these areas. It is the intention of this thesis, that with the framework for extending the functionality of Petrel established, already developed plug-ins such as the custom simulator for our two-phase 5-spot water flooding pattern will be used in the class room environment to educate students on the topic of reservoir simulation and provide a basis to develop other plug-ins to implement in the future, custom simulators for our model order reduction and production optimization algorithms.

**Scope of Study**

Reservoir simulation is an area in the oil and gas industry that plays an extremely important role in the decision making process concerning how reservoirs are appraised, production potential evaluated and how implementation of development plans are executed to extract hydrocarbons for maximum profit. In an earlier section of my introduction, I talked briefly about the development of reservoir simulators and how present day simulators are capable of handling multiphase flow behavior and representing complex geological patterns that are a direct consequence of gradual
technical advances as noted by Watts (1997). In continuation of these developments, research is on-going to solve identified problems and provide unique efficient solutions. Among these areas of study, model order reduction and production optimization has been identified as a potential area for breakthrough.

The scope of this project is limited to establishing the framework for extending the functionality of Petrel, the pre and post processor of Eclipse by using Ocean-in-Petrel, an application program interface to add our algorithms as processes in the model building workflow or create an external custom simulator that implements our algorithms. We have achieved this objective with the implementation of an in-house simulator that was initially written in Matlab (Matlab R2012b), a highly technical programming language very suitable for matrix and numerical manipulation and then re-programmed in Ocean (Ocean 2012) which is the primary programming language for extension of Petrel functionality.

The contents of this thesis, is broken into three distinct parts. The first part explains the governing equations involved in describing the system of flow behavior in our oil-water 5-spot water flood system and how they are applied to this particular system to obtain responses such as pressures, saturations and production rates at different points in time. The second part is the implementation of the same reservoir simulation scenario using Petrel to explain how it is developed step by step from creating a simple grid, creating rock, saturation functions, fluid models, initial conditions other flow functions, well placement, development strategy and finally defining a simulation case and submitting it for simulation. The third part describes the extension
of Petrel by using Ocean-in-Petrel to develop an external custom simulator to implement and visualize results of our water flood case. Figure 1 below shows a simplified flow process of the implemented external custom simulator plug-in which works by first defining a set of inputs setup in Petrel including grid sizes and dimensions, fluid and rock parameters, well information and well location etc. that are fed to the external custom simulator for simulation. The results of the simulation are sent back into Petrel workspace for visualization.

**Figure 1: Process Flow for Custom Simulator Plug-in**

The results of this project will first and foremost serve as an educational tool to be used in a classroom environment teaching students important aspects of reservoir simulation and secondly, it will establish the framework for implementing plug-ins developed in the research areas of model order reduction and production optimization.
CHAPTER II
MODEL CONSTRUCTION

The phenomenon of simulating the behavior of a reservoir is bore by first understanding the mechanics, properties and dependencies of fluids and the governing equations that determine the flow of these fluids i.e. hydrocarbons in porous media. These governing equations are partial differential equations that are discretized in both space and time by using discretizing schemes such as finite differences, finite elements etc. In this project, the partial differential equations are discretized using finite differences.

Our reservoir is a two phase and two dimensional oil-water system that is been implemented with a typical shoebox model that has no flow boundaries on all sides of the reservoir. The reservoir is constructed with an equal number of grid blocks in both the x and y directions with one layer in the z direction and initialized with an initial pressure, water saturation and corresponding fluid and rock properties in every grid block. The flow of fluids in the reservoir is dictated by applying the theory of conservation of mass, flow equations and equations of state to a control volume with the firm assumption that the net mass rate of the control volume is equal to zero. The manner in which flow into grid blocks is determined is done is by using a 5-point stencil i.e. two dimensional problem that takes into consideration flow from the four surrounding grid blocks namely north, south, east and west into the “center” grid block as shown in Figure 2 below.
In the following section, I will present the governing partial differential flow equations utilized in my reservoir beginning with its differential form and end up with its discretized form.

**Partial Differential Flow Equations**

The reservoir system described in this problem resembles that of a black oil system which places each flowing fluid into one of three categories namely oil, water or gas. In reservoir simulation, flow of multiphase problems is treated simultaneously with certain assumptions been made. For instance, in a black oil system, oil and water are immiscible; hence there is no transfer of mass between both fluids and in the presence of gas, gas is soluble only in oil and in that case, we will have both free gas and dissolved gas i.e. solution gas. In this project, we restrict our work to an oil-water fluid system.

Flow of hydrocarbons in porous media is based on three fundamental principles namely conservation of mass, application of flow equations and application of an
equation of state. In this chapter, I will highlight the main parts in the discretization process of our flow equations. For full derivation of the partial differential equations, refer to “Basic Applied Reservoir Simulation” written by Ertekin et al (2001). I begin the discretization process by re-emphasizing that the net rate of change of fluid into a given control volume is zero. The equations determining the flow of water and oil respectively in our reservoir are shown below.

\[
\frac{\partial}{\partial x} \left[ \beta_c k_x A_x \frac{k_{rw}}{\mu_w B_w} \left( \frac{\partial p_w}{\partial x} - \gamma_w \frac{\partial Z}{\partial x} \right) \right] \Delta x \\
+ \frac{\partial}{\partial y} \left[ \beta_c k_y A_y \frac{k_{rw}}{\mu_w B_w} \left( \frac{\partial p_w}{\partial y} - \gamma_w \frac{\partial Z}{\partial y} \right) \right] \Delta y = \frac{V_b}{\alpha_c} \frac{\partial}{\partial t} \left( \phi S_w \right) - q_{wsc} \tag{1}
\]

\[
\frac{\partial}{\partial x} \left[ \beta_c k_x A_x \frac{k_{ro}}{\mu_o B_o} \left( \frac{\partial p_o}{\partial x} - \gamma_o \frac{\partial Z}{\partial x} \right) \right] \Delta x \\
+ \frac{\partial}{\partial y} \left[ \beta_c k_y A_y \frac{k_{ro}}{\mu_o B_o} \left( \frac{\partial p_o}{\partial y} - \gamma_o \frac{\partial Z}{\partial y} \right) \right] \Delta y \\
= \frac{V_b}{\alpha_c} \frac{\partial}{\partial t} \left( \phi S_o \right) - q_{osc} \tag{2}
\]

where

- \( \beta_c \) is a flow term conversion factor
- \( k_x \) is the permeability in the x – direction
- \( k_y \) is the permeability in the y – direction
- \( A_x \) is the cross-sectional area perpendicular to the flow in the x-direction
- \( A_y \) is the cross-sectional area perpendicular to the flow in the y-direction
- \( k_{rw} \) is the water phase relative permeability
- \( k_{ro} \) is the oil phase relative permeability
\( \mu_w \) is the viscosity of water

\( \mu_o \) is the viscosity of oil

\( B_w \) is the formation volume factor of water

\( B_o \) is the formation volume factor of oil

\( p_w \) is the pressure water

\( p_o \) is the pressure of oil

\( \gamma_w \) is the gravity or density of water

\( \gamma_o \) is the gravity or density of oil

\( Z \) is the elevation with respect to a given datum

\( \Delta x \) is an incremental distance used for discretization purposes in the x-direction

\( \Delta y \) is an incremental distance used for discretization purposes in the y-direction

\( V_b \) is the bulk volume of the control volume

\( \phi \) is the porosity

\( S_w \) is the pore volume water saturation

\( S_o \) is the pore volume oil saturation

\( q_{w_{sc}} \) is the source/sink term of water, measured at standard conditions

\( q_{o_{sc}} \) is the source/sink term of oil, measured at standard conditions

The above two equations i.e. (1) and (2) has four unknowns namely pressure of water (\( p_w \)), pressure of oil (\( p_o \)) water saturation (\( S_w \)) and oil saturation (\( S_o \)) so we need to find two other equations to solve our system of equations. The other two equations
come from constraints we need to impose on the system. These are the saturation
constraints and capillary constraints.

At any given time in the control volume, the summation of water and oil saturation
is unity and the oil and water pressures are related via the capillary pressure as shown in
the equations below.

\[ S_w + S_o = 1 \]  \hspace{1cm} (3)

\[ p_{cow} = p_o - p_w \]  \hspace{1cm} (4)

Where

\[ p_{cow} \] is the capillary pressure relating oil and water pressures

In our reservoir model, the effects of gravity and capillary pressure are neglected hence

\[ p_{cow} = 0 \rightarrow p_o = p_w \]  \hspace{1cm} (5)

\[ \frac{\partial Z}{\partial x} = 0 \]  \hspace{1cm} (6)

Using our constraint equations and neglecting gravity and capillary effects we
arrive at the following equations below.

\[ \frac{\partial}{\partial x} \left[ \beta c k_x A_x \frac{k_{rw}}{\mu_w B_w} \left( \frac{\partial p_o}{\partial x} \right) \right] \Delta x + \frac{\partial}{\partial y} \left[ \beta c k_y A_y \frac{k_{rw}}{\mu_w B_w} \left( \frac{\partial p_o}{\partial y} \right) \right] \Delta y \]

\[ = \frac{V_b}{\alpha_c} \frac{\partial}{\partial t} \left( \frac{\phi S_w}{B_w} \right) - q_{wsc} \]  \hspace{1cm} (7)
Equations (7) and (8) are the water and oil equations respectively. The first two terms on the left side of either equation accounts for flow into and out of the control volume with respect to the x and y direction while the two terms on the right side respectively accounts for the accumulation rate of oil and water in the control volume and the injection or production of fluids into or out of the control volume i.e. sources or sinks.

**Discretization of Partial Differential Equations**

In order to solve our system of partial differential equations numerically, a discretization scheme has to be chosen. We have various discretization schemes available. For our simulator, we chose a finite differences scheme. The partial differential equations governing flow of both phases i.e. oil and water are discretized in both space and time as shown in the steps below.

We begin by discretizing the left side of both water and oil equations i.e. equations (7) and (8) in space. After discretizing in space, the end results are the equations below.
To make the equations more compact, we introduce the term transmissibility as shown in equation (10). It is a grouping of the geometrical factor of flow between two adjacent grid blocks and parameters such as viscosity, formation volume factor and relative permeability that are dependent on the saturation and pressure of the flowing fluid.

\[ T_{wx} = \beta_c k_x A_x \frac{k_{rw}}{\mu_w B_w}, \quad T_{wy} = \beta_c k_y A_y \frac{k_{rw}}{\mu_w B_w} \]  

We apply finite difference to \( \frac{\partial p_o}{\partial x} \) and \( \frac{\partial p_o}{\partial y} \) and substitute equations (10) into (9) and we obtain the more compact equation below.

\[
\frac{\partial}{\partial x}\left[\beta_c k_x A_x \frac{k_{rw}}{\mu_w B_w} \left(\frac{\partial p_o}{\partial x}\right)\right] \Delta x + \frac{\partial}{\partial y}\left[\beta_c k_y A_y \frac{k_{rw}}{\mu_w B_w} \left(\frac{\partial p_o}{\partial y}\right)\right] \Delta y = \\
\left[T_{wx,i+\frac{1}{2},j}(p_{o,i+1,j} - p_{o,ij})\right] + \left[T_{wx,i-\frac{1}{2},j}(p_{o,i-1,j} - p_{o,ij})\right]
\]
The oil equation is discretized in a similar way using the transmissibility of oil and applying finite differences accordingly to obtain the equation below:

\[
T_{wy_{i,j+\frac{1}{2}}} (p_{oi,j+1} - p_{oij}) + T_{wy_{i,j-\frac{1}{2}}} (p_{oi,j-1} - p_{oij})
\]

The term on the right side of the partial differential equations i.e. equations (7) and (8) accounting for accumulation can be discretized as shown below.

\[
\frac{\partial}{\partial x} \left[ \beta_c k_x A_x \frac{k_{ro}}{\mu_o B_o} \frac{\partial p_o}{\partial x} \right] \Delta x + \frac{\partial}{\partial y} \left[ \beta_c k_y A_y \frac{k_{ro}}{\mu_o B_o} \frac{\partial p_o}{\partial y} \right] \Delta y
\]

\[
= \left[ T_{ox_{i+\frac{1}{2}j}} (p_{oi+1,j} - p_{oij}) \right] + \left[ T_{ox_{i-\frac{1}{2}j}} (p_{oi-1,j} - p_{oij}) \right]
\]

\[
+ \left[ T_{oy_{i,j+\frac{1}{2}}} (p_{oij+1} - p_{oij}) \right] + \left[ T_{oy_{i,j-\frac{1}{2}}} (p_{oij-1} - p_{oij}) \right]
\]

The above equations can be further expanded in the final form below.
The derivatives of porosity and formation volume factor with respect to pressure can be determined by numerical perturbation as shown in the equations below.

\[
\frac{V_b}{\alpha_c} \frac{\partial}{\partial t} \left( \frac{\phi S_w}{B_w} \right) =
\]

\[
\frac{V_b}{\alpha_c \Delta t} \left[ \left( \frac{\varnothing'}{B_w^n} + \varnothing^{n+1} \left( \frac{1}{B_w} \right)' \right) S_w^n \left( p_o^{n+1} - p_o^n \right) + \left( \frac{\varnothing}{B_w} \right)^{n+1} (S_w^{n+1} - S_w^n) \right]
\]

(15)

\[
\frac{V_b}{\alpha_c} \frac{\partial}{\partial t} \left( \frac{\phi (1-S_w)}{B_o} \right) = \frac{V_b}{\alpha_c \Delta t} \left[ \left( \frac{\varnothing'}{B_o^n} + \varnothing^{n+1} \left( \frac{1}{B_o} \right)' \right) (1 - S_w^n) \left( p_o^{n+1} - p_o^n \right) - \right.
\]

\[
\left. \left( \frac{\varnothing}{B_o} \right)^{n+1} (S_w^{n+1} - S_w^n) \right]
\]

(16)

The derivatives of porosity and formation volume factor with respect to pressure can be determined by numerical perturbation as shown in the equations below.

\[
\varnothing' = \frac{(\varnothing^{n+1} - \varnothing^n)}{(p^{n+1} - p^n)}
\]

(17)

\[
\left( \frac{1}{B_w} \right)' = \frac{\left( \frac{1}{B_w} \right)^{n+1} - \left( \frac{1}{B_w} \right)^n}{(p^{n+1} - p^n)}
\]

(18)

With equations derived thus far, the final form of the partial differential equations for water and oil respectively can be represented with equations (19) and (20).
\[
\left[T_{wx}^{i+\frac{1}{2},j}\left(p_{o_{i+1,j}} - p_{o_{ij}}\right)\right] + \left[T_{wx}^{i-\frac{1}{2},j}\left(p_{o_{i-1,j}} - p_{o_{ij}}\right)\right] + \\
\left[T_{wy}^{i,j+\frac{1}{2}}\left(p_{o_{i,j+1}} - p_{o_{ij}}\right)\right] + \left[T_{wy}^{i,j-\frac{1}{2}}\left(p_{o_{i,j-1}} - p_{o_{ij}}\right)\right]
\]
\[
= \frac{v_b}{\alpha_c\Delta t}\left[\frac{\phi'}{B_w^{n+1}} + \phi^{n+1}\left(\frac{1}{B_w}\right)'\right]S_w^n(p_o^{n+1} - p_o^n) + \left(\frac{\phi}{B_w}\right)^{n+1}\left(S_w^{n+1} - S_w^n\right) - q_{wsc}
\]
\[
\left[T_{ox}^{i+\frac{1}{2},j}\left(p_{o_{i+1,j}} - p_{o_{ij}}\right)\right] + \left[T_{ox}^{i-\frac{1}{2},j}\left(p_{o_{i-1,j}} - p_{o_{ij}}\right)\right] + \\
\left[T_{oy}^{i,j+\frac{1}{2}}\left(p_{o_{ij,j+1}} - p_{o_{ij}}\right)\right] + \left[T_{oy}^{i,j-\frac{1}{2}}\left(p_{o_{ij,j-1}} - p_{o_{ij}}\right)\right]
\]
\[
= \frac{v_b}{\alpha_c\Delta t}\left[\frac{\phi'}{B_o^{n+1}} + \phi^{n+1}\left(\frac{1}{B_o}\right)'\right] \left(1 - S_w^n\right)(p_o^{n+1} - p_o^n) - \left(\frac{\phi}{B_o}\right)^{n+1}\left(S_w^{n+1} - S_w^n\right) - q_{asc}
\]

The source and sink terms represent flow into and out of the reservoir. The general sign convention of flow is defined as positive if mass is flowing into the reservoir and negative if flowing out of the reservoir. In both cases of injection and producer wells, the constraints on the wells are either rate specified or pressure specified in terms of a bottom-hole pressure. If rate specified the rate constraint is applied directly to the algebraic equations but if pressure specified a well model is necessary to couple the well production rate, well bottom-hole pressure and grid block pressure together.
For this model, we made use of the Peaceman’s well index equation to couple the well production rate, well bottom-hole pressure and grid block pressure. The well index, production rate and equivalent wellbore radius equations is shown below

\[
WI = \frac{2\pi \beta_c k k_r h}{\mu B [\ln(r_o/r_w) + s]}
\]

\[
q = WI (p_o - p_{wf})
\]

\[
r_0 = 0.28 \left\{ \left( \frac{k_y}{k_x} \right)^{1/4} (\Delta x)^2 + \left( \frac{k_x}{k_y} \right)^{1/4} (\Delta y)^2 \right\}^{1/2}
\]

where

- \( WI \) is the Peaceman’s well index
- \( \beta_c \) is a flow term conversion factor
- \( k \) is the total permeability
- \( k_r \) is the fluid relative permeability
- \( k_x \) is the permeability in the x - direction
- \( k_y \) is the permeability in the y – direction
- \( \Delta x \) is the grid block dimension relative to flow in the x-direction
- \( \Delta y \) is the grid block dimension relative to flow in the y – direction
- \( h \) is the well grid-block thickness
- \( \mu \) is the fluid viscosity
$B$ is the fluid formation volume factor

$r_o$ is the Peaceman wellbore radius

$r_w$ is the actual wellbore radius

$s$ is the skin factor

$p_o$ is the well grid-block pressure

$p_{wf}$ is the wellbore bottom-hole pressure

$q$ is the fluid flow rate in standard condition

Since $k_x$ is equal to $k_y$ and $\Delta x$ is equal to $\Delta y$, the Peaceman’s wellbore radius equation can be simplified further with the equation below.

$$r_0 = 0.208 \times \Delta x \quad (24)$$

After the partial differential equations have been discretized both in space and time the next step is to choose a suitable formulation to solve our set of equations. We can solve the equations explicitly which is the case where flow terms on the left of our equations are evaluated at the old time steps. The equations can also be solved implicitly i.e. at the new time-steps for pressures and saturations. It is well known in reservoir simulation circles that solving the equations implicitly is preferred over other formulation methods because of its advantages among which is its unconditionally stable nature but for this project, since our main objective is concerned more with establishing the framework for developing plug-ins and extending the functionality of Petrel, less emphasis was placed on the method used to formulate our equations for solving.
explicit and implicit formulations are expressed using equations (25) and (26) below at 
\{n\} and \{n + 1\} respectively.

\begin{align*}
T_{wx, i+\frac{1}{2}, j}^{n,n+1} \left( p_{o,i+1,j} - p_{o,i,j} \right)^{n,n+1} + T_{wx, i-\frac{1}{2}, j}^{n,n+1} \left( p_{o,i-1,j} - p_{o,i,j} \right)^{n,n+1} \\
+ T_{wy, i+\frac{1}{2}, j}^{n,n+1} \left( p_{o,i,j+1} - p_{o,i,j} \right)^{n,n+1} \\
+ T_{wy, i-\frac{1}{2}, j}^{n,n+1} \left( p_{o,i,j-1} - p_{o,i,j} \right)^{n,n+1} \\
= \frac{V_b}{\alpha_c \Delta t} \left[ \frac{\phi'}{B_w^n} + \phi^{n+1} \left( \frac{1}{B_w} \right)' \right] S_{w}^n (p_o^{n+1} - p_o^n) \\
+ \left( \frac{\phi}{B_w^n} \right)^{n+1} (S_{w}^{n+1} - S_{w}^n) - q_{wsc}
\end{align*}
\\( (25) \)

\begin{align*}
T_{ox, i+\frac{1}{2}, j}^{n,n+1} \left( p_{o,i+1,j} - p_{o,i,j} \right)^{n,n+1} + T_{ox, i-\frac{1}{2}, j}^{n,n+1} \left( p_{o,i-1,j} - p_{o,i,j} \right)^{n,n+1} \\
+ T_{oy, i+\frac{1}{2}, j}^{n,n+1} \left( p_{o,i,j+1} - p_{o,i,j} \right)^{n,n+1} \\
+ T_{oy, i-\frac{1}{2}, j}^{n,n+1} \left( p_{o,i,j-1} - p_{o,i,j} \right)^{n,n+1} \\
= \frac{V_b}{\alpha_c \Delta t} \left[ \frac{\phi'}{B_0^n} + \phi^{n+1} \left( \frac{1}{B_0} \right)' \right] (1 - S_{w}^n) (p_o^{n+1} - p_o^n) \\
- \left( \frac{\phi}{B_0^n} \right)^{n+1} (S_{w}^{n+1} - S_{w}^n) - q_{osc}
\end{align*}
\\( (26) \)
The next step is to select a choice of method to solve the set of discretized partial differential equations. Among the choices, we have lagging coefficients, implicit pressure explicit saturation, sequential implicit and fully implicit. For this project we solve the equations using lagging coefficients meaning that the saturations and pressures on the left hand side of the partial differential equations are evaluated at the new time step with the non-linear pressure and saturation dependent terms evaluated at the current time step. The lagging coefficient equations are shown below.

\[
T_{wx_i+\frac{1}{2},j}^n (p_{o_{i+1,j}} - p_{o_{i,j}})^{n+1} + T_{wx_i-\frac{1}{2},j}^n (p_{o_{i-1,j}} - p_{o_{i,j}})^{n+1} \\
+ T_{wy_{i,j+\frac{1}{2}}}^n (p_{o_{i,j+1}} - p_{o_{i,j}})^{n+1} + T_{wy_{i,j-\frac{1}{2}}}^n (p_{o_{i,j-1}} - p_{o_{i,j}})^{n+1} \\
= \frac{V_b}{\alpha_c \Delta t} \left[ \frac{\varrho'}{B_w^n} + \varphi^n \left( \frac{1}{B_w^n} \right) ' \right] S_w^n (p_o^{n+1} - p_o^n) + \left( \frac{\varrho}{B_w^n} \right)^n (S_w^{n+1} - S_w^n) \\
- q_{ws}^{n+1}
\]

\[
T_{ox_i+\frac{1}{2},j}^n (p_{o_{i+1,j}} - p_{o_{i,j}})^{n+1} + T_{ox_i-\frac{1}{2},j}^n (p_{o_{i-1,j}} - p_{o_{i,j}})^{n+1} \\
+ T_{oy_{i,j+\frac{1}{2}}}^n (p_{o_{i,j+1}} - p_{o_{i,j}})^{n+1} + T_{oy_{i,j-\frac{1}{2}}}^n (p_{o_{i,j-1}} - p_{o_{i,j}})^{n+1} \\
= \frac{V_b}{\alpha_c \Delta t} \left[ \frac{\varrho'}{B_o^n} + \varphi^n \left( \frac{1}{B_o^n} \right) ' \right] (1 - S_w^n) (p_o^{n+1} - p_o^n) \\
- \left( \frac{\varrho}{B_o^n} \right)^n (S_w^{n+1} - S_w^n) - q_{os}^{n+1}
\]
To further simplify notations, the following terms below are defined to be substituted into our algebraic equations.

\[
d_{wp} = \frac{V_b}{\alpha_c \Delta t} \left[ \frac{\theta'}{B_w^n} + \theta^{n+1} \left( \frac{1}{B_w} \right)^' \right] S_w^n \tag{29}
\]

\[
d_{ws} = \frac{V_b}{\alpha_c \Delta t} \left( \frac{\theta}{B_w} \right)^{n+1} \tag{30}
\]

\[
d_{op} = \frac{V_b}{\alpha_c \Delta t} \left[ \frac{\theta'}{B_o^n} + \theta^n \left( \frac{1}{B_o} \right)^' \right] (1 - S_w^n) \tag{31}
\]

\[
d_{os} = -\frac{V_b}{\alpha_c \Delta t} \left( \frac{\theta}{B_o} \right)^n \tag{32}
\]

Substituting equations (30) to (32) directly into equations our water and oil equations we obtain the simplified equations below.

\[
T_{wx_{i+\frac{1}{2},j}}^n \left( p_{o_{i+1,j}} - p_{o_{i,j}} \right)^{n+1} + T_{wx_{i-\frac{1}{2},j}}^n \left( p_{o_{i-1,j}} - p_{o_{i,j}} \right)^{n+1} + T_{wy_{i,j+\frac{1}{2}}}^n \left( p_{o_{i,j+1}} - p_{o_{i,j}} \right)^{n+1} + T_{wy_{i,j-\frac{1}{2}}}^n \left( p_{o_{i,j-1}} - p_{o_{i,j}} \right)^{n+1} = d_{wp_i}^n (p_o^{n+1} - p_o^n) + d_{ws_i}^n (S_w^{n+1} - S_w^n) - q_{wsc_i}^{n+1} \tag{33}
\]

\[
T_{ox_{i+\frac{1}{2},j}}^n \left( p_{o_{i+1,j}} - p_{o_{i,j}} \right)^{n+1} + T_{ox_{i-\frac{1}{2},j}}^n \left( p_{o_{i-1,j}} - p_{o_{i,j}} \right)^{n+1} + T_{oy_{i,j+\frac{1}{2}}}^n \left( p_{o_{i,j+1}} - p_{o_{i,j}} \right)^{n+1} + T_{oy_{i,j-\frac{1}{2}}}^n \left( p_{o_{i,j-1}} - p_{o_{i,j}} \right)^{n+1} = d_{op_i}^n (p_o^{n+1} - p_o^n) - d_{os_i}^n (S_w^{n+1} - S_w^n) - q_{osc_i}^{n+1} \tag{34}
\]
The next step is to set up the equations for every grid block applying initial conditions and boundary conditions. The set of equations obtained are re-arranged and placed in a matrix form as shown below.

\[ T^n X^{n+1} = D^n (X^{n+1} - X^n) - Q^{n+1} \]  

(35)

where

- \( T^n \) is the transmissibility matrix evaluated at the current time-step
- \( D^n \) is the accumulation matrix evaluated at the current time-step
- \( Q^{n+1} \) is the source/sink vector evaluated using new time-step pressure but with the well index evaluated at the current time-step
- \( X^{n+1} \) is the state vector of the new time step
- \( X^n \) is the state vector of the current time step

**Structure of Matrices**

To show the typical structure of the matrices to be solved, we make use of a two-dimensional two-phase problem with a 3 by 3 system and make use of the notations \( T_{l_1j_1}, T_{l_1j_w}, T_{l_1j_n}, T_{l_1j_s}, T_{l_1j_c} \) (where \( l \) stands for water(\( w \)) or oil(\( o \))) to represent the contribution to the transmissibility from each surrounding grid block to the center grid block of the 5-point stencil as we loop through all grid blocks. \( T_{w_1j_c} \) is the negative summation of the other applicable components of transmissibility.
Since the transmissibility and accumulation coefficients are obtained at the current time-step equation (35) can be re-arranged in the form below to be solved
For source and sink terms, its representation in the Q matrix is determined by the mode of operation i.e. if it’s an injection or production well. If the well is flow rate specified, then the rate value can be directly entered into the matrix equation but if it is bottom hole pressure specified then the flow terms in the matrix are replace with the equations below for oil and water with the well index calculated at the current time-step.

\[
T^n X^{n+1} = D^n (X^{n+1} - X^n) - Q^{n+1}
\]  \hspace{1cm} (36)

\[
(T^n - D^n) X^{n+1} = -D^n X^n - Q^{n+1}
\]  \hspace{1cm} (37)

The algebraic equations for oil and water are adjusted for grid blocks that have wells with flowing bottom-hole specifications by adding to the transmissibility matrix the negative of the product of the well index and well block pressure and the left side of the equations modified by adding the negative product of the well index and well flowing pressure.

\[
q_{oi} = WI_{oi}^n (p_{oi}^{n+1} - p_{wf}^i)
\]  \hspace{1cm} (38)

\[
q_{oi} = WI_{oi}^n p_{oi}^{n+1} - WI_{oi} p_{wf}^i
\]  \hspace{1cm} (39)

\[
q_{wi} = WI_{wi}^n (p_{wi}^{n+1} - p_{wf}^i)
\]  \hspace{1cm} (40)

\[
q_{wi} = WI_{wi}^n p_{wi}^{n+1} - WI_{wi} p_{wf}^i
\]  \hspace{1cm} (41)

At this point, the matrices representing our system of equations are ready to be solved. The equations can be solved by using a series of solvers either direct or iterative.
solvers. For my Matlab implementation of this simulator, a direct solver was used followed by iterative solvers such as GMRES and bi-conjugate gradient. For my Ocean (C#) implementation because of the choices available to iterative solvers were used amongst which a generalized product bi-conjugate gradient iterative matrix solver was the best.
CHAPTER III

MODEL IMPLEMENTATION IN PETREL

Introduction

In the previous chapter, the partial differential equations that govern the flow of fluid in porous media were described and thereafter, we proceeded to discretize and transform these partial differential equations into a form that could be used to obtain a set of algebraic equations that are solved to provide a unique solution by applying the discretized equations to every grid block in our reservoir.

This project is three-fold, firstly, understanding the partial differential equations that govern fluid flow, secondly, implementing a reservoir simulator with a particular configuration i.e. two-phase and two-dimensional oil-water 5-spot pattern in a programming environment such as Matlab for prototyping and thirdly, implementing the same reservoir model configuration in Petrel with Eclipse 100 and a Custom External Simulator added to the choices of available simulators by using the application programming interface, Ocean which is the primary language used to extend the functionality of Petrel.

The intention of this project is to develop a framework for extending the functionality of Petrel with efficient algorithms in areas of interest such as model order reduction and production optimization. The end product of this project will be used for educational purposes in the classroom environment as a tool to explore the fundamentals of reservoir simulation by integrating the commercial reservoir simulator i.e. Petrel and a custom simulator developed by students in a graduate student reservoir simulation class.
In order to make this exposition clearer, we will build our algorithms based on a pre-defined reservoir model. The following sections of this chapter, shows how the reservoir simulation configuration for the oil-water two dimensional 5-spot pattern is implemented in Petrel beginning by creating a simple grid, creating surfaces and horizons and layers followed by setting up 3-D properties, after which we proceed to add vertical wells, make fluid models, create rock physics functions, create a development strategy and finally define a simulation case to be simulated by a choice of Eclipse formatted simulators in this case, Eclipse 100.

As mentioned earlier, the second part of this project involved the implementation of an in-house simulation configuration which is a two phase two-dimensional 5-spot water flood pattern as shown in Figure 3. Shown below are the inputs in Tables 1, 2 and 3 and a flow chart, figure 4 that describes the process of simulation for this configuration and the responses of the simulator.

![Figure 3: Picture of Implemented Reservoir Model](image-url)
## Model Inputs

<table>
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<th>Input</th>
<th>Value/ Comment</th>
</tr>
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<tbody>
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<td>Number of Grids in X-direction</td>
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<td>Producer 4 location</td>
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Table 1: Inputs of Simulator
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</tr>
<tr>
<td>0.4375</td>
<td>0.0158</td>
<td>0.1985</td>
</tr>
<tr>
<td>0.51</td>
<td>0.05</td>
<td>0.1016</td>
</tr>
<tr>
<td>0.5825</td>
<td>0.1221</td>
<td>0.0429</td>
</tr>
<tr>
<td>0.655</td>
<td>0.2531</td>
<td>0.0127</td>
</tr>
<tr>
<td>0.7275</td>
<td>0.4689</td>
<td>0.0016</td>
</tr>
<tr>
<td>0.8</td>
<td>0.8</td>
<td>0</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>

Table 2: Water and Oil Relative Permeability as a Function of Water Saturation

<table>
<thead>
<tr>
<th>Pressure (psi)</th>
<th>Oil FVF (RB/STB)</th>
<th>Oil Viscosity (cP)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1160.301902</td>
<td>1.12028</td>
<td>1.461437638</td>
</tr>
<tr>
<td>1356.102848</td>
<td>1.117055</td>
<td>1.489130119</td>
</tr>
<tr>
<td>1551.903794</td>
<td>1.113839</td>
<td>1.52165029</td>
</tr>
<tr>
<td>1747.70474</td>
<td>1.110632</td>
<td>1.558580779</td>
</tr>
<tr>
<td>1943.505686</td>
<td>1.107435</td>
<td>1.599605902</td>
</tr>
<tr>
<td>2139.306632</td>
<td>1.104247</td>
<td>1.644480675</td>
</tr>
<tr>
<td>2335.107577</td>
<td>1.101068</td>
<td>1.693010352</td>
</tr>
<tr>
<td>2530.908523</td>
<td>1.097898</td>
<td>1.74503633</td>
</tr>
<tr>
<td>2726.709469</td>
<td>1.094737</td>
<td>1.800426135</td>
</tr>
<tr>
<td>2922.510415</td>
<td>1.091585</td>
<td>1.859066091</td>
</tr>
<tr>
<td>3118.311361</td>
<td>1.088443</td>
<td>1.920855845</td>
</tr>
<tr>
<td>3314.112307</td>
<td>1.085309</td>
<td>1.985704189</td>
</tr>
<tr>
<td>3509.913253</td>
<td>1.082184</td>
<td>2.053525828</td>
</tr>
<tr>
<td>3705.714199</td>
<td>1.079069</td>
<td>2.124238845</td>
</tr>
<tr>
<td>3901.515145</td>
<td>1.075962</td>
<td>2.197762692</td>
</tr>
<tr>
<td>4097.316091</td>
<td>1.072865</td>
<td>2.274016601</td>
</tr>
<tr>
<td>4293.117037</td>
<td>1.069776</td>
<td>2.35291831</td>
</tr>
<tr>
<td>4488.917983</td>
<td>1.066696</td>
<td>2.434383055</td>
</tr>
<tr>
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</tr>
<tr>
<td>4880.519875</td>
<td>1.060563</td>
<td>2.60464548</td>
</tr>
<tr>
<td>5076.320821</td>
<td>1.05751</td>
<td>2.693254824</td>
</tr>
</tbody>
</table>

Table 3: Oil Formation Volume Factor and Oil Viscosity as a Function of Pressure
Water Viscosity was given as 1cp

Water Formation Volume Factor ($B_w$) was calculated with the given expression below

$$B_w = B_{wref} / (1 + x + x^2)$$ \hspace{1cm} (42)

Where

$$x = cw \times (pw - pref)$$ \hspace{1cm} (43)

$B_{wref}$ is unity

cw is Water compressibility

pref is Reference pressure

pw is grid pressure

With the inputs specified above been applied to our algebraic equations describing the flow of oil and water with boundary conditions factored in and a given time to march our simulator in time to, the simulator follows the process of the flow chart below.
Figure 4: The Process Flow Chart for our Simulator Code
Model Construction

It is worth reminding the reader at this point what Petrel is. Petrel is the pre and post processor of Eclipse which is a commercial simulator widely used in the Oil and Gas industry. With Petrel a user has the ability to develop very complicated models that span every domain of reservoir modeling including reservoir engineering, production engineering, well completions, geology, geophysics and a host of other domains that are intermingled to develop an eventual model that will be used for reservoir simulation.

The primary goal of this chapter is to fully introduce the in-house model that is used to develop our framework for extending the features of Petrel as it pertains to the motivation for this project. So far in this thesis, we’ve gone through the motivation for this project, had a description of the partial differential equations that govern the flow of fluids in porous media and introduced our in-house test model for Petrel extension. For the remaining of this chapter, I show how our test model is implemented in Petrel using the inputs in Tables 1, 2 and 3.

In the model implementation, I make use of “preset choices” including dead oil which is an in-built oil and water fluid model in Petrel, consolidated sandstone and saturation functions. These preset choices are actually the sources of our relative permeability, oil viscosity and oil formation volume factor tables. It is important that we go through this section showing the model implementation in Petrel so that the reader can gain a better understanding of the subtle differences of how the custom simulator is developed with ocean which is the application programming interface used for Petrel extension. The extension of Petrel with ocean is covered in the next chapter. Our in-
house two phase two dimensional 5-spot water flood pattern configuration is implemented as follows.

- Open Petrel and begin a new project as shown in Figure 5
- Open the project settings window
- Under the Coordinates and units tab select Field as the unit System. Click OK.

![Figure 5: New Project Settings](image.png)
Next we create a Simple Grid as shown in Figure 6

- On the Processes tab, open the Utilities category and double click on “Make simple grid.”

![Processes Tab Screenshot]

**Figure 6: Selection of Make Simple Grid**

- In the “Make simple grid” window that opens, give it the name 11x11.
- Under the Input data tab as shown in Figure 7, change the top and base limits as shown in the figure below.
- In the Geometry tab as shown in Figure 8, change the Xmin and Ymin values of 1000 and change Xmax and Ymax both to 1352
- Click Apply and OK:
Figure 7: Make Simple Grid (Input Data Tab)
Figure 8: Make a Simple Grid (Geometry Tab)

- On the Models tab, Expand New model and 11x11
- Right-click on Skeleton and select Convert to surface (three surfaces are created on the Input tab) as shown in Figure 9 and 10
Figure 9: Conversion to Surface

Figure 10: Surfaces for 11x11 (Input Pane)
• On the Processes tab, open the Corner point gridding category as shown in Figure 11

![Processes](image)

**Figure 11:** Make Horizon in Processes Tab

• Double click on Make horizons
• Click the Insert button and add two rows in the table as shown in Figure 12
• Select the Top surface on the Input tab, then click the first blue arrow
• Select the Base surface on the Input tab, then click the third blue arrow
• Click Apply and OK
Figure 12: Make Horizon with 11x11

- On the process tab, open Corner point gridding category
- Double click on Layering process
- Change number of layers to 1 as shown in Figure 13
- Click Apply and OK

Figure 13: Layering Process
Create a simple 3D property
- Open the Models tab
- Right-click on Properties and select Calculator as shown in Figure 14

![Figure 14: Property Calculator](image)

- Select Porosity template under Attach new to template
- Enter Porosity=0.2 (no spaces) in the textbox as shown in Figure 15
- Click the ENTER button.
- Click the” x” (top right) button to close the calculator window.

The property format above sets the porosity value to 0.2 in all cells of the grid.
Figure 15: Porosity and Permeability Calculation

- Open a new 3D window (Window > New 3D window)
- In the Models pane open the Properties folder as shown in Figure 16
- Select the radio button on the left of Porosity
- Click on the Show/Hide grid lines icon
Figure 16: Porosity Property in a 3D Window

- Right click on Porosity property
- Select Color table
- Click the Max and Min buttons to scale color
- Click Apply and OK to change display color

Permeability Property

- Select Permeability template under Attach new to template
- Enter Permeability=ran(80,220) in the textbox
- Click the ENTER button.
- Click the” x” (top right) button to close the calculator window.
- Open a 3D window to view “perm”
- Right click on perm
- Select Color table
- Click the Max and Min buttons to scale color
- Click Apply and OK to change display color

The above format of permeability gives random values of permeability between 80md and 220md in each grid block.
Add vertical wells

- Select Insert/New well to bring up Create new well window
- Name the first well P1
- Select Oil as Well Symbol
- Give Well Head X a value of 1016 (X and Y Co-ordinates for all wells are calculated with respect to the chosen reservoir top limit i.e. 1000)
- Give Well Head Y a value of 1016
- Keep KB value as 0
- Check the Specify vertical trace box
- Change the Bottom MD to 950

Repeat the above process for 4 more wells i.e. P2, P3 P4 and INJ1 (all oil producers except INJ1, an injector). Change the position of the wells accordingly

P2
Well head X value of 1336
Well head Y value of 1016

P3
Well head X value of 1016
Well head Y value of 1336

W4
Well head X value of 1336
Well head Y value of 1336

INJ1
- Select Injection water (15) as the Well symbol
- Well Head X value of 1176
- Well Head Y value of 1176
- KB and Bottom MD values the same as the producers

Create PVT Property
- Open the Processes tab
- Open the Simulation category
- Double click on Make fluid model process
- In the Make fluid model window, click on the Use presets button and select Dead oil as shown in Figure 17
- Change the reference pressure to 2800 and accept other default values
Figure 17: Make Fluid Model Window General Tab

- Click on the Initial conditions tab
- Enter the following values as shown in Figure 18 (keep value on other tabs unchanged)
- Click Apply and OK to exit
Figure 18: Make Fluid Model (Initial Condition Tab)

The newly created PVT property is stored in the Fluids folder on the Input tab, the properties can be plotted on a function window as shown in Figure 19 (use Window> New function window)
Create rock physics function

- Open the Processes tab
- Open Simulation category
- Double click on Make rock physics functions process
- Select Create new
- Click on the Saturation tab
- Click on Use Preset button and select Sand as shown in Figure 20
- Click Apply then Ok

Figure 19: Oil Formation Volume Factor and Oil Viscosity for Dead Oil
Figure 20: Make Rock Physics Functions (Saturation Tab)

Switch to Compaction tab

- Click on Use preset and select Consolidated sandstone
- Update the Compaction tab with the values shown in figure 21 below
- Click Apply and OK to exit
Create Development Strategy

- Open the Processes tab
- Open Simulation category
- Double click on Make development strategy process
- On the Input tab select the Wells folder name
- In the Make development strategy window
- Drop the Wells folder from the input pane in the Wells folder tree on the left hand side using the drop arrow as shown in Figure 22
- Change the end from 2031-01-01 to 2012-10-10
Click the Add rules icon to bring up the Add rules window as shown in Figure 23.
Select the Well pressure production control rule (double click or use the Add rule button).
Select P1 and Drop it into Wells.
Choose bottom hole pressure as Control mode and enter a value of 2900 [psi].
Repeat the process for P2, P3 and P4.
For INJ1 Add a Well water injection control rule.
Choose surface rate as Control mode and enter a value of 4000 Stb/d.
Click on Reporting Frequency and change time to days.
Define and run a Simulation case

- Open the Processes tab
- Open Simulation category
- Double click on Define simulation case process
- Enter Eclipse_sim as case name (no spaces allowed in case name)
- Select ECLIPSE 100 as Simulator as shown in Figure 24
- Select Single porosity as Type
- Drop Permeability property in the models pane into PERMX, PERMY and PERMZ
- Drop Porosity property into Porosity(PORO)
Figure 24: Define Simulation Case (Grid)

- Click on the Functions tab
- Select Drainage relative permeability as shown in Figure 25
- Highlight Sand in the Rock physics functions folder in the input pane and drop in the property
Figure 25: Define Simulation Case (Drainage Relative Permeability)

- Switch to Rock Compaction and drop in Consolidated sand from the input as shown in Figure 26
Figure 26: Define Simulation Case (Rock Compaction)

- Switch to Black oil and drop in Initial condition 1 from the input as shown in Figure 27
Figure 27: Define Simulation Case (Initial Condition)

- Switch to Strategy tab
- Use the insert button to add a row to the table
- Select Development strategy1 and drop into Development strategy as shown in Figure 28
Figure 28: Define Simulation Case (Development Strategy)

- Click Export button to export the simulation case
- Click the Run button (Run your project from the C-drive)

Results

- Open a New function window
- On the Cases tab check the box next to Eclipse_sim
- Open Results tab check Well under the identifier folder and select Wells P1, P2, P3 and P4 under the PROD folder
- Select Dynamic Data
• Open the Dynamic results data folder and the Rates folder
• Check the Oil production rate checkbox to plot the oil production rates for all 4 producers.
• Open another function window and repeat the process to plot the Water production rate.

Figure 29: Oil Production Rates for the 4 Producers with Eclipse 100
Figure 30: Water Production Rates for all 4 Producers with Eclipse 100
Figure 31: Water Saturation Map at 100 days with Eclipse 100
Figure 32: Water Saturation Map at 200 days with Eclipse 100

Figure 33: Pressure Map at 100 days with Eclipse 100
Figures 29 to 34 show the responses obtained using Eclipse100 for oil and water production rates up till the end of simulation time, water and pressure saturation maps at 100 and 200 days. As expected, within a considerable amount of time we have water breakthrough and the oil and water production rates increase significantly with the water production rates leveling off and the oil production declining quite steeply after a while. The water saturation and pressure maps in both cases increase with increasing time away from the injector as expected exhibiting the typical performance of a water flood pattern.

In the next chapter, we cover the topic of Petrel extension via Ocean Software Development Kit and show in general terms how Petrel can be extended with respect to
the motivation of this project using our in-house two phase two dimensional oil water 5-spot pattern water flood simulation configuration as the model to establish the framework for Petrel extension.
CHAPTER IV

OCEAN IN PETREL AND CUSTOM SIMULATOR

Ocean in Petrel

Ocean in Petrel is an application programming interface that is used by a software developer to extend the functionality of Petrel, the pre and post processor of Eclipse by creating plug-ins which contains algorithms designed by developers to implement unique objectives. The software developer can be a member of academia, industry or a separate third party that has identified a need to extend the features of Petrel for their personal benefit or with the purpose of making their plug-ins available commercially for interested parties. The software developer is able to create plug-ins by making use of different Ocean templates (Figure 34) that are available in Visual Studio which is the programming environment through which Ocean and Petrel communicate.

The user of Ocean in Petrel has access to common services such as service locator, messaging interface, module lifecycle, data source manager, event transaction manager, domain object hosting, generic data types etc. With the ocean software development kit, the developer has the ability to “unleash their creativity” and extend the user interface of petrel, manipulate data and add to the workflow of model building with the numerous tools and services that give them access to the various domains in Petrel such as seismic domain, well domain, geology, shapes, reservoir model, simulation etc.

The basic building block of a plug-in is the module which is self-contained in assembly. The modules which contain the algorithms to extend the Petrel data user interface are loaded at Petrel start up and unloaded at shut down. The module goes
through five steps in its life cycle namely initialize, integrate, integrate presentation, disintegrate and dispose.

The features of ocean and its capability are far too extensive to be covered in this project so for a more comprehensive coverage of Ocean in Petrel Software Development Kit, I refer you to attend the class thought by the software owners i.e. Schlumberger by registering for a session through the company website “www.ocean.slb.com.”

At this time, I need to remind the reader of the general objective of this project which is to develop a framework for adding our developed algorithms in the form of “processes” to the workflow of Petrel for model building to prepare “cases” for reservoir simulation. For our purposes, there are two primary ways of accomplishing this goal. One is through what is referred to as a “workstep” which has its algorithm self-contained in a module that on execution will implement the code in the module. Depending on what the algorithm is designed to do, it could accept inputs and pass the inputs through the algorithm and produce outputs that could be used for further processing. The other suggested method which is the preferred method is the implementation of a custom simulator which will accept inputs in a certain format to be fed to a custom simulator which is passed through its algorithm and returns responses back to Petrel workspace for visualization. Presently in Petrel we have available to us four simulators i.e. Eclipse 100, Eclipse 300, FrontSim and INTERSECT. Ocean in Petrel gives the user the ability to add a custom simulator to the drop box in the “define simulation case” window of Petrel.

For completion, I will talk about the general flow in developing plugs-in with an example that will give a sense of the workflows and then I will concentrate on the
implementation of the custom simulator. Figure 35 below shows the Petrel user interface with the different forms in which personalized algorithms and their responses could be added. They include processes/worksteps, data models, menus, windows, toolbars, dialogs 2D and 3D renderers etc.

Figure 35: Petrel User Interface (Adapted from Ocean Software Development Kit Fundamental Training Volume 1)
After Ocean and Petrel have been installed, the process of developing plug-ins in Petrel begins with selecting the appropriate Ocean template in Visual Studio which is the programming environment for interaction between Ocean and Petrel. Figure 36 below shows the choices of templates that can be chosen to implement your algorithms.

![Ocean Templates in Visual Studio](image)

**Figure 36: Ocean Templates in Visual Studio**

**How to Make a Plug-in**

Once a choice is made, a wizard is opened which guides the developer through a series of steps to assist them to generate generic blocks of code that are needed to implement their algorithms. As an example, I will implement in the next section a plug-
in that will add a workstep to Petrel to display the name of the active project opened in Petrel.

Making Ocean Plug-ins

- Open up Visual Studio 2010
- Select OceanPlugin as the template as shown in Figure 37
- Give it a name and location
- Click Ok

Figure 37: Choosing Ocean Template
In the wizard that shows after clicking Ok, enter the relevant information as shown below in Figure 38.

Figure 38: Ocean Plug-in Create - Step 1

- Click Next and next again
- In the next steps, fill the information as shown in figure 39, 40 and 41 below
- Click Next and then Finish
Figure 39: Ocean Plug-in Create - Step 3

Figure 40: Ocean Plug-in Create - Step 4
After going through these steps the ocean wizard will create a project in Visual Studio that will have a plug-in as a project and a module with a workstep attached to it. An instance of a plug-in and module are shown in Figures 42 and 43 respectively. The wizard has been designed to do a lot of the setting up for developers leaving the customization based on the desired objectives of the workstep (plug-in) for the developer to fill in. To complete the wizard, on the last page that comes up, check the “Additional Reference Settings” box to add the relevant assemblies to the project solution. These assemblies can be found in the Petrel 2012 folder in the Schlumberger folder in the “Programs” folder.

Figure 41: Ocean Plug-in Create - Step 5
Figure 42: Instance of Plug-in

Figure 43: Instance of Module
For most customization in Petrel, the most important part of customizing modules is the code placed in the “ExecuteSimple ()” Method as shown in Figure 44. It is in this block that the developer places their algorithm for Petrel extension.

- Click on OceanModuleWorkstep.cs in the Solution Explorer window

```csharp
public Executor(Arguments arguments, workflowSimulateContext context)
{
    this.arguments = arguments;
    this.context = context;
}

public override void ExecuteSimple()
{
    // TODO: Implement the workflow logic here.
    IDatasourceManager dataSourceManager;
    SIlo ocn = PetrelProject.CreateProjectInfo(dataSourceManager);
    System.IO.FileInfo fi = pl.ProjectFile;
    if (fi != null)
    {
        arguments.ProjName = fi.Name;
    }
    PetrelLogger.Info("The petrel file name is: "+ arguments.ProjName);
    return;
}
```

**Figure 44: ExecuteSimple Method**

- Build project in Visual Studio
- Open Petrel
- Click on Processes pane (Bottom Left) and open Plug-ins
- Double click on OceanModuleWorkstep and then Apply/OK
Since no project is open you will get the default value i.e. “No project is open” as shown in Figure 45. If a project is opened, then it will display the project name. As simple as this example is, it shows the basics in developing plug-ins by utilizing the wizard to generate most of the code that is used with the relevant parts to be customized according to your objectives. With a workstep, inputs can be passed to the “ExecuteSimple” method for processing to produce outputs that could be used further.
for model building. In the next section, I describe the development of our custom simulator.

**Custom Simulator**

Being able to develop a custom simulator to add to the choices of simulators available is a very useful resource of Ocean. Granted that the present simulators Eclipse 100, 300 etc. are very powerful but in certain instances, a certain level of customization is needed to achieve our objectives. This project began with a certain motivation in mind which was to develop a framework for creating a means to implement custom algorithms in the areas of model order reduction and production optimization. In order to achieve our goal, a test model was used to develop a custom simulator i.e. a two phase two dimensional 5-spot pattern waterflood. For propriety reasons, certain parts of the code implementing the custom simulator cannot be shown here but the general steps to develop the simulator will be explained below with a few figures to give a visual of the process.

The process of developing the custom simulator began with calibrating our test model which was earlier programmed in Matlab (for prototyping) against the same reservoir simulation algorithm re-programmed in C-sharp which is the language of Ocean. After calibration was completed, in collaboration with a Schlumberger software development team, I was able to add a custom simulator to the choices of simulators in the “define simulation case” dialog box of Petrel and develop our framework for custom simulator building.
As displayed in Chapter 3 with the implementation of our test model in Petrel, the prepared “case” is developed by going through a series of actions to complete the description needed to come up with a scenario for reservoir simulation. It begins with making a simple grid, making horizons, layering, creating properties like porosity and permeability, selecting a fluid model, selecting rock physics functions, adding wells and constraining those wells, creating a development strategy and finally defining the case and submitting it to a choice of reservoir simulators (in this case Eclipse 100) for simulation. Figure 46 below shows the process of reservoir simulation in Petrel with regard to our test case.

Figure 46: Reservoir Simulation in Petrel (Slide Courtesy of Schlumberger)
After the case has been defined in “define simulation case” and submitted for reservoir simulation, the case is ran and then on completion the results are imported back into Petrel for visualization. To make a custom simulation, the flow process is not so different. In order to give the developer the opportunity to add their functionality, a series of steps are included in order to customize and format the custom simulator into a well suited form. Figure 47 below captures the additional steps implemented for customization.

Figure 47: Process for Developing a Custom Simulator
The steps of note are the PostExport, PreRun, PreImport and SubmitCommand. For each of these additional steps it’s the place where the developer can add the customized functionality they want. For instance in the PostExport(Case) method, the deck that is exported from the define simulation case is reformatted and modified in a form that the custom simulator expects. The PreRun sets up the environment before the simulation launch. The submit command submits the case to the simulator and the PreImport method formats the results from the simulator in an Eclipse format suited for visualization in Petrel. The reservoir simulation code was developed by the author and the connection with Petrel was developed by a Schlumberger software developer. Figure 48 below shows a snap shot of the custom simulator “Student Simulator” added to the drop list of simulators.

![Image of Define Simulation Case]

Figure 48: Student Simulator in Define Simulation Case
As mentioned, for propriety reasons certain parts of the code cannot be shown but the general framework for connecting the reservoir simulator to Petrel is shown below with a few visuals. I draw the attention of the reader to Figure 49 below that shows the “Student Simulator” i.e. our custom simulator. When this choice is selected from the simulator drop list, an additional tab named “Custom tab” appears.

![Figure 49: Student Simulator and Custom Tab](image)

When the user clicks on the “Custom tab” tab in order to submit their “simulation file they would need to have pre-built (compiled) their reservoir simulation solution in Visual Studio to generate a “dll” file that will be uploaded using the “Browse Simulator”
button shown in the figure above. The custom simulator has been developed such that
certain parts such as the code for the PostExport, PreRun, SubmitCommand and
PreImport methods are sealed but the parts that can be shown are displayed below.

In order to generate the “dll” to be passed to the custom simulator, the user has to
prepare a reservoir simulation file in the format shown in Figure 50 below.

![Image of code snippet]

**Figure 50: Format of the Code to Generate Dll File**

Also specific to generating the dll file the user has to follow the instructions as stated below.

*Please follow the instructions for locating *.dll file.*

---

Program.cs

```csharp
namespace SimDllFile
{
    class SimClass
    {
        public OutputArgs Simulate(InputArgs inputs)
        {
            #region Running Simulator
            Input region
            
            // Computation region: This is the Only region that should be modified by the students. The "Input" will
            #endregion
            OutputArgs Output = RunSimulatorMethod(Inputs, qo_qw, Po_Sw_Values);
            return Output;
        }
        private static Saturation_QoilPressureDependencies ComputeDependencies(double[,] so, double[,] sw, double
        
        private static OutputArgs RunSimulatorMethod(InputArgs Inputs, double[,] qo_qw, double[,] Po_Sw_Values);
        }
        class InputArgs
        class OutputArgs
        class Saturation_QoilPressureDependencies
    }
}
```

---
/// 2. The namespace must contain three classes: "SimClass","InputArgs","OutputArgs"
/// 3. "SimClass" should contain a public method named "Simulate" of the form:
///     public OutputArgs Simulate(InputArgs inputs)
///     This method is the place for writing the simulator's computations. Note that
///     in the following example "RunSimulatorMethod" is a private method that is
called by "Simulate" method. "RunSimulatorMethod"
///     method is NOT a part of presumed structure. The developer can have any
number of private methods to be called from "Simulate" method.
///
/// --Where to locate *.dll file:
/// 1. press F6 to build the *.files. The file that should be loaded by simulator
is "SimDllFile.dll" and is copied at C:\StudentSimulator.
/// or ...Documents\StudentSimulator.
///
/// --Note: before creating *.dll file, with the explained structure, make sure
that your simulator runs, independently. You could test this in a console
application with constant inputs.

The inputs in the “Input region” in the code shown are passed from Petrel to the
reservoir simulator in the format shown in Figure 51 below which is compatible with
Eclipse to be used in the “Computation region” of the code which is where
transmissibility, sources and sink terms and accumulation matrices are calculated, the
set of equations are solved using an iterative solver, the oil and water production rates
for the producers and “lists” containing pressure and water saturation values for every
grid block at every time step to be converted to maps at every time step are stored for
visualization at the end of simulation with the “RunSimulatorMethod” method.
Of note in my program are two methods i.e. Saturation_OilPressureDependencies and RunSimulatorMethod. Both perform two separate functions. The first calculates the water and oil relative permeability as a function of water saturation for every grid block and also the oil viscosity and formation volume factor as a function of pressure all at every time step. This function is basically an interpolation function that accepts the values for saturations and pressures for every grid block as inputs and parses out interpolated values of relative permeability, viscosity and formation volume factors to be used in calculations in the Computation region. The second method i.e. RunSimulatorMethod is the method used to store oil and water production rates, pressure
and saturation values in every grid block at every time step to be passed back to Petrel
workspace for visualization. The block of code below shows the RunSimulationMethod
in its entirety as shown in Figure 52 below which after simulation is completed, stores
the oil and water production rates for all the well, pressure and saturations for every grid
block at every timestep to be imported via a connection into Petrel for visualization.

```csharp
private static OutputArgs RunSimulatorMethod(InputArgs Inputs, double[,] qo_qw, double[,] Po_Sw_Values)
{
    // Production Rates
    List<double> _Producer1_oilratevector = new List<double>(); // for recording Oil Production Rate of Producer 1
    List<double> _Producer1_waterratevector = new List<double>(); // for recording Water Production Rate of Producer 1
    List<double> _Producer2_oilratevector = new List<double>(); // for recording Oil Production Rate of Producer 2
    List<double> _Producer2_waterratevector = new List<double>(); // for recording Water Production Rate of Producer 2
    List<double> _Producer3_oilratevector = new List<double>(); // for recording Oil Production Rate of Producer 3
    List<double> _Producer3_waterratevector = new List<double>(); // for recording Water Production Rate of Producer 3
    List<double> _Producer4_oilratevector = new List<double>(); // for recording Oil Production Rate of Producer 4
    List<double> _Producer4_waterratevector = new List<double>(); // for recording Water Production Rate of Producer 4

    // Pressures and Water Saturations
    List<double[, ,]> _pressuregridvector = new List<double[, ,]>() // for recording Pressure Calculations
    List<double[, ,]> _watersaturationgridvector = new List<double[, ,]>() // for recording Water Saturation Calculations

    //..............for storing pressures and saturations while looping
    //+++++++++++++++++++ looping through all the time steps

    Figure 52: RunSimulationMethod Code
```
int rowshifter = 0;
int pressurecolumnshifter = 0;
int watersaturationcolumnshifter = 1;

for (int i = 0; i < Inputs.ReportingTimes.Count; i++)
{
    // ................ Calculating Oil and Water Production Rate Vectors
    _Producer1_oilratevector.Add(qo_qw[i, 0]);
    _Producer1_waterratevector.Add(qo_qw[i, 1]);
    _Producer2_oilratevector.Add(qo_qw[i, 2]);
    _Producer2_waterratevector.Add(qo_qw[i, 3]);
    _Producer3_oilratevector.Add(qo_qw[i, 4]);
    _Producer3_waterratevector.Add(qo_qw[i, 5]);
    _Producer4_oilratevector.Add(qo_qw[i, 6]);
    _Producer4_waterratevector.Add(qo_qw[i, 7]);

    double[,] PressureGrid = new double[Inputs.Nx, Inputs.Ny, Inputs.Nz];

    for (int ii = 0; ii < Inputs.Nx; ii++)
    {
        for (int j = 0; j < Inputs.Ny; j++)
        {
            for (int k = 0; k < Inputs.Nz; k++)
            {
                PressureGrid[ii, j, k] = Po_Sw_Values[rowshifter, pressurecolumnshifter];
                WaterSaturationGrid[ii, j, k] = Po_Sw_Values[rowshifter, watersaturationcolumnshifter];
            }
            rowshifter = rowshifter + 1;
        }
    }

    rowshifter = 0;
    pressurecolumnshifter = pressurecolumnshifter + 2;
    watersaturationcolumnshifter = watersaturationcolumnshifter + 2;

    // ..........Adding the Grids list
    _pressuregridvector.Add(PressureGrid);
    _watersaturationgridvector.Add(WaterSaturationGrid);
}

Figure 52: Continued
// returning the outputs
        OutputArgs Outputs = new OutputArgs();
        Outputs.Producer1_OilRateVector = _Producer1_oilratevector;
        Outputs.Producer1_WaterRateVector = _Producer1_waterratevector;
        Outputs.Producer2_OilRateVector = _Producer2_oilratevector;
        Outputs.Producer2_WaterRateVector = _Producer2_waterratevector;
        Outputs.Producer3_OilRateVector = _Producer3_oilratevector;
        Outputs.Producer3_WaterRateVector = _Producer3_waterratevector;
        Outputs.Producer4_OilRateVector = _Producer4_oilratevector;
        Outputs.Producer4_WaterRateVector = _Producer4_waterratevector;

        Outputs.PressureGridVector = _pressuregridvector;
        Outputs.WaterSaturationGridVector = _watersaturationridVector;

        return Outputs;
    }
}

Figure 52: Continued

As part of the motivation of this project, the final product will be used in the classroom environment for educational purposes where students will be given the inputs used to develop the case as presented here in this project and develop a reservoir simulator that will be connected via ocean to Petrel to match the responses of the Eclipse100 equivalent simulated case. We ran the Student Simulator with the same inputs and obtained figures 53 to 58 for the responses for oil and water production rates and water and pressure maps.
Results

Figure 53: Oil Production Rate Using Custom Simulator
Figure 54: Water Production Rate Using Custom Simulator

Figure 55: Water saturation at 100 Days with Custom Simulator
Figure 56: Water Saturation at 200 Days with Custom Simulator

Figure 57: Pressure Map at 100 Days with Custom Simulator
The above plots show the responses for oil and water saturation up till the end of simulation time, pressure and water saturation maps at 100 and 200 days using the custom simulator. In a similar fashion as the Eclipse100 simulation run shown in chapter three, the custom simulator exhibits similar responses in performance. The custom simulator in its present form can accept various operating parameter values for permeability, porosity, well constraints (bottom-hole pressures for producers and rates for injectors) etc. Figures displaying the oil and water production rates and the pressure and water saturation maps with the following values are shown

Figure 58: Pressure Map at 200 Days with Custom Simulator
- Permeability (randomly generated with min of 100md and max of 400md)
- Porosity of 0.35
- Bottom-Hole Pressure of 3200 psi
- Injection rate 4,500 STB/day

Figure 59: Oil Production Rates - Custom Simulator
Figure 60: Water Production Rates - Custom Simulator

Figure 61: Water Saturation at 100 Days - Custom Simulator
Figure 62: Water Saturation at 200 Days - Custom Simulator

Figure 63: Pressure at 100 Days - Custom Simulator
Figures 59 to 64 show the responses for oil and water production rates and pressure and water saturation maps at 100 and 200 days with the Custom simulator using the new values of permeability, porosity, bottom-hole pressure and injection rates. As we can see from the plots comparing it to the original inputs the responses especially with the production rates are reasonably higher as expected due to the higher bottom hole pressure, injection rates and the higher range of permeability and porosity in every grid block. Comparison of results will be shown in the next chapter.
CHAPTER V

COMPARISON OF RESULTS AND CONCLUSIONS

In this chapter, I compare the results obtained from using Eclipse100 and the custom simulator for accuracy and error analysis. The analysis is conducted by comparing the results of two cases, one which makes use of a homogenous permeability of 100md and the other which is a heterogeneous permeability case used in developing our framework; both cases of which are five-spot water flood pattern configured.

I compare the results on a well by well basis for oil and water production rates visually and present a table with root mean square error for each well. The error analysis will be followed by interpretations based on the results of the simulations and at the end I make my conclusions and recommendations.

Root Mean Square Error Analysis

Figures 65 to 68 show the oil and water production rates for the homogenous case.
Figure 65: Oil Production Rate of P1 with both Simulators (Homogeneous Case)

Figure 66: Water Production Rate of P1 with both Simulators (Homogeneous Case)
Figure 67: Oil Production Rate of P2 with both Simulators (Homogeneous Case)

Figure 68: Water Production Rate of P2 with both Simulators (Homogeneous Case)
Since this first case is homogeneous, all four producers have the same performance of oil and water production rates so figures for Producers three and four are not shown. Visually it can be seen that the oil and water production rates using Eclipse100 and the Custom Simulator match for all wells. Below is a table showing the root mean square error calculated with the expression below.

\[
\sqrt{\frac{\sum(Q_{\text{eclipse}} - Q_{\text{custom simulator}})^2}{n}} \quad (44)
\]

Where

\(Q_{\text{eclipse}}\) is the production rate of oil or water at a particular time using Eclipse100

\(Q_{\text{custom simulator}}\) is the production rate of oil or water at a particular time using the custom simulator

\(n\) is the number of reported values

<table>
<thead>
<tr>
<th></th>
<th>RMS ERROR AT END OF SIMULATION</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>OIL PRODUCTION RATE (STB/D)</td>
</tr>
<tr>
<td></td>
<td>WATER PRODUCTION RATE (STB/D)</td>
</tr>
<tr>
<td>PRODUCER 1</td>
<td>25</td>
</tr>
<tr>
<td>PRODUCER 2</td>
<td>25</td>
</tr>
<tr>
<td>PRODUCER 3</td>
<td>25</td>
</tr>
<tr>
<td>PRODUCER 4</td>
<td>25</td>
</tr>
</tbody>
</table>

Table 4: Rms Error of Production Rate for all Producers (Homogenous Case)

From Table 4 above, the root mean square error for both oil and water production rate for all producers for this case of homogenous permeability is notable which is rather misleading because visually, the oil and water production rates match very well. I
attribute this root mean square error value to the values reported at different times for the Eclipse100 and Custom Simulator runs which might be as a result of the manner in which values are reported for the Custom Simulator and quite possibly internal computations peculiar to Eclipse that the Custom Simulator doesn’t account for. I analyzed the performance of the heterogenous permeability case for both simulators and obtained Figures 69 to 76 below showing the oil and water production rates.

![P1 Oil production rate](image)

**Figure 69: Oil Production Rate of P1 with both Simulators (Heterogenous Case)**
Figure 70: Water Production Rate of P1 with both Simulators (Heterogenous Case)

Figure 71: Oil Production Rate of P2 with both Simulators (Heterogenous Case)
Figure 72: Water Production Rate of P1 with both Simulators (Heterogenous Case)

Figure 73: Oil Production Rate of P3 with both Simulators (Heterogenous Case)
Figure 74: Water Production Rate of P3 with both Simulators (Heterogenous Case)

Figure 75: Oil Production Rate of P4 with both Simulators (Heterogenous Case)
Visualy, the oil and water production rates for all wells of this heterogeneous case obtained using Eclipse100 and the Custom Simulator are also matched quite well with the exception of the water production rate curves for Producers two and three which has a maximum water production rate point error of five percent. Table 5 below shows the root mean square error of oil and water production rates for all Producers.

<table>
<thead>
<tr>
<th>Producer</th>
<th>RMS Error Oil Production Rate (STB/D)</th>
<th>RMS Error Water Production Rate (STB/D)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PRODUCER 1</td>
<td>22.3</td>
<td>10.3</td>
</tr>
<tr>
<td>PRODUCER 2</td>
<td>28</td>
<td>38</td>
</tr>
<tr>
<td>PRODUCER 3</td>
<td>14</td>
<td>43</td>
</tr>
<tr>
<td>PRODUCER 4</td>
<td>24</td>
<td>9.24</td>
</tr>
</tbody>
</table>

Table 5: Rms Error of Production Rate for all Producers (Heterogenous Case)
Similarly to the homogenous case, the root mean square error for each well comparing the oil and water production rates is misleading since from just visually inspecting the rates, we expect to see a much smaller root mean square error. I attribute this difference again to the manner in which the reporting values are obtained between both simulators and the internal computation done by Eclipse to calculate responses based on its algorithm.

At this point, I have established that the responses using oil and water production rates match reasonably well bearing in mind that in order to understand the difference in reported values, a reasonable amount of time has to be spent understanding the subtle differences between the computations done by Eclipse and how it differs from that of the Custom Simulator.

**Pressure and Time-Step Analysis**

Between both cases i.e. the homogenous and heterogenous cases, we have a reasonable match but to see if we can quantify further the differences, two more situations are investigated. Firstly, a comparison between the pressures in every grid block at random time steps during simulation obtained from Eclipse100 and the Custom Simulator using root mean square error analysis and secondly, a time step reduction to account for the difference in partial difference equation formulation and solving method since Eclipse is a fully implicit simulator and the custom simulator is formulated using lagging coefficient. For this second case, the time step of the custom simulator will be reduced from 0.2 days to 0.001 days and the results will be compared against the
Eclipse100 results for the heterogenous case; specifically the oil and water production rates of Producers two and three which seemed to have the highest discrepancy.

To conduct the pressure analysis, the pressures in every grid block at different time steps capturing the performance of the entire simulation are compared. Figures 77 to 81 below show a comparison of pressures at different time steps.

![Figure 77: Pressures from Eclipse 100 and Custom Simulator on February 15th](image-url)
Figure 78: Pressures from Eclipse 100 and Custom Simulator on April 15\textsuperscript{th}

Figure 79: Pressures from Eclipse 100 and Custom Simulator on June 15\textsuperscript{th}
Figure 80: Pressures from Eclipse 100 and Custom Simulator on August 15th

Figure 81: Pressures from Eclipse 100 and Custom Simulator on October 1st
The Table below shows how the Custom Simulator performs in comparison to Eclipse100 using root means square error analysis with the following expression.

$$\sqrt{\frac{\sum (P_{\text{eclipse}} - P_{\text{custom simulator}})^2}{n}}$$  \hspace{1cm} (45)

Where

$P_{\text{eclipse}}$ is the pressure of oil or water at a particular time using Eclipse100.

$P_{\text{custom simulator}}$ is the pressure of oil or water at a particular time using the custom simulator.

$n$ is the number of reported values.

<table>
<thead>
<tr>
<th>PRESSURE (ROOT MEAN SQUARE ERROR)</th>
</tr>
</thead>
<tbody>
<tr>
<td>February 15th</td>
</tr>
<tr>
<td>----------------</td>
</tr>
<tr>
<td>339 psi</td>
</tr>
</tbody>
</table>

Table 6: Pressure Rms Analysis between Both Simulators (Heterogenous Case)

From the pressure plots above it can be inferred that the custom simulator shows a similar pattern of performance to the Eclipse100 simulator. From inspection, it seems that Eclipse probably makes use of some kind of multiplier which the Custom Simulator doesn’t factor in. The root mean square error analysis in Table 6 above gives a measure of the average difference between the pressures between both simulators at different time steps during simulation. Following the analysis of the second case with a reduced
time step, I will make my final observations, conclusions and recommendations concerning my analysis.

For my last scenario of investigation, I reduced the time step. For a lagging coefficient formulated simulator, the biggest issue dealt with is that of stability. In order to ensure stability, the simulator can only march in time at or below a certain threshold which for this simulator is 0.2 days and so we have this time step constraint. It is expected that if the time step of a lagging coefficient simulator is reduced orders of magnitude lower than the threshold, results obtained will be in closer agreement with a fully implicit simulator such as Eclipse but from the analysis presented thus far it would seem the discrepancy between both Eclipse and the Custom simulator lies more within the internal computations of Eclipse that the Custom Simulator doesn’t account for. For investigating purposes and completeness, the time step was reduced to 0.001 days to see if results are improved. Figures 82 to 85 below show the oil and water production rates of Producers two and three followed by a root mean square error analysis table.
Figure 82: Oil Production Rate of P2 with Custom Simulator (0.001 days)

Figure 83: Water Production Rate of P2 with Custom Simulator (0.001 days)
Figure 84: Oil Production Rate of P3 with Custom Simulator (0.001 days)

Figure 85: Water Production Rate of P3 with Custom Simulator (0.001 day)
From the plots above, it can be seen visually that the oil and water production rates obtained with the lower timestep of 0.001 gives a very similar performance to the 0.2 timestep simulation. It is also seen from table 7 with the root mean square error analysis at the end of simulation that the oil production rate obtained with the custom simulator converges with Eclipse while the water production rate does not. Water break through occurs at approximately 14 days into the simulation which is also the time oil production begins to decline rapidly. It is possible that with lower timesteps, the oil production rates my match even much better but there is no indication that the water production rate will improve any better. Thus far in my analysis, I have attributed the mismatch to internal computations in Eclipse that are not considered in the custom simulator and perhaps the formulation method and have so far seen no indications to convince me otherwise.

<table>
<thead>
<tr>
<th></th>
<th>RMS ERROR AT END OF SIMULATION</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>OIL PRODUCTION RATE (STB/D)</td>
</tr>
<tr>
<td>PRODUCER 2 (0.2 days)</td>
<td>28</td>
</tr>
<tr>
<td>PRODUCER 3 (0.2 days)</td>
<td>14</td>
</tr>
<tr>
<td>PRODUCER 2 (0.001 days)</td>
<td>5</td>
</tr>
<tr>
<td>PRODUCER 3 (0.001 days)</td>
<td>3.2</td>
</tr>
</tbody>
</table>

Table 7: Rms Analysis of Production Rate at 0.001 and 0.2 Timesteps for P2 and P3
Recommendation

With all the analysis done comparing the performance of the Custom Simulator for both homogenous permeability and heterogenous permeability case to the Eclipse run, the mismatch between both simulators is more apparent for the heterogenous case especially for the water production of Producers two and three. Analyzing the distribution of pressures in each grid block at different timesteps as shown in figures 77 to 81 the argument of internal computations in Eclipse is supported because of the pattern exhibited indicating that there probably is a multiplier applied in Eclipse that is responsible for the upward shift in pressures. The objective of this thesis which is to develop a framework for extending Petrel has been achieved. The plug-in its present form is ready to be used in a classroom environment as part of the tools in educating students on reservoir simulation. Matching the performance of Eclipse is definitely an objective of ours but to match Eclipse fully, a considerable amount of time has to be spent in understanding its internal computations. After that study has been conducted, findings can be implemented in future versions of the plug-in. Also it is recommended that the plug-in be upgraded to a fully implicit formulated simulator not only to obtain results much closer to Eclipse but to compare the performance between a lagging coefficient simulator and a fully implicit simulator. In the next section, I present plans for future work and make my final conclusions.

Future Work and Conclusion

Reservoir simulation is continuously developing with the tools and methods for reservoir simulation becoming more efficient and accurate. We began this project with
the goal in mind to ultimately make our contribution to reservoir simulation
development in general by first identifying and working to improve on results obtained
in the past in certain areas of interest. Efficient and robust algorithms for areas such as
model order reduction and production optimization are in progress and with the work
accomplished in this project, when these algorithms are completed, similar custom
simulators will be built to implement the algorithms.

Regarding future work, the extent of this project is very far reaching. We have
accomplished our goal of establishing our framework for extending Petrel via a custom
simulator with Ocean. The test model that we used was a very simple shoe box model.
As I mentioned in my introduction, we currently deal with very complex reservoir
systems that require the use of unstructured grids. This is one area where our plug-in
could be improved to better accommodate more realistic reservoir scenarios. The
configuration that we used was fixed i.e. a five-spot pattern with wells placed in a fixed
position. The code could be altered to allow for more flexibility of well placement and
well constraint. Lastly, Ocean gives the software developer access to every domain in
reservoir modeling. Future plug-ins made might have influences from other domains of
model building such as geology, geophysics etc. The software developer is only limited
by their creativity so the possibilities for improvements are endless. And lastly, in
connection with the result comparison chapter, to model more realistic scenarios using
Eclipse as a basis, studies have to be conducted to factor into the Custom simulator the
internal computations that are representative of real life reservoir performance.
In conclusion, we have taken a first step to accomplish our ultimate goal. With this project, as it is firstly academic, it will be used in a classroom environment. It will be of immense value to students because it will help them in getting a stronger understanding of fundamental and advanced topics in reservoir simulation. The other aspect is its research component which is tied to the future plug-ins to be made in our identified areas of model order reduction and production optimization. The algorithms for these areas are currently developed and when completed, custom simulators will be developed to implement them.
REFERENCES


