Developing a Smart Proxy for Fluidized Bed Using Machine Learning

Amir Ansari

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Developing a Smart Proxy for Fluidized Bed Using Machine Learning

Amir Ansari

Thesis submitted
to the Benjamin M. Statler College of Engineering and Mineral Resources
at West Virginia University

in partial fulfillment of the requirements for the degree of

Masters of Science in
Petroleum and Natural Gas Engineering

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Keywords: Coal Gasification, Fluidized-bed, Multiphase Flow, Artificial Neural Network, Data Mining

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ABSTRACT

Developing a Smart Proxy for Fluidized Bed Using Machine Learning

Amir Ansari

Using fossil fuel which has been grown dramatically during the recent century, causes an increase in greenhouse gas emission. The global warming issue pushes the engineers toward the cleaner type of energy like Hydrogen. Coal gasification is one of the cheapest methods to obtain Hydrogen. Coal gasification is a special case of more general problem called fluidized bed. In order to design and optimize a gasification process, a deep understanding of multiphase flow in a gasifier is needed. MFiX is a commercial multi-phase flow simulator which has been used to simulate the gas and solid transport and reaction in the gasifier using Computational Fluid Dynamics (CFD). Although simulating multiphase flow using commercial CFD software has a lot of flexibilities, it is really time-consuming and some other way could be implemented to reduce the run time. The effort of this project is to develop an alternate method to perform the same analysis but with much lower computational cost. A data-driven approach is used to build a smart proxy by employing the knowledge of Artificial Intelligence (AI) and Data Mining (DM).

In this project, a smart proxy will be developed to study and analyze the fluidized bed problem. This smart proxy is then will be used as a replicate of the CFD solver, with a good accuracy and faster speed. This proxy needs an incredible less amount of time in comparison to the CFD solver with a reasonable error (less than 10%). MATLAB neural network toolbox is used for training.

The goal of this project is to prove the concept of using AI&DM for computational fluid dynamics especially predicting multiphase flow. Multiphase flow has a wide range of application in petroleum industry such as multi-phase flow in the wellbore, surface lines, and hydraulic fracturing such as proppant transport in the hydraulic fracture. This project opens a new way to accelerate the fluid dynamics analysis and reduce its costs.
NOMENCLATURE

d = Particle diameter
\( \vec{f}_G \) = Flow resistance due to the internal porous surfaces
\( \vec{g} \) = Acceleration of gravity

\( I_{gm} \) = interaction force representing the momentum transfer between the gas phase and the \( m^{th} \) solids phase
\( I_{ml} \) = interaction force representing the momentum transfer between the \( m^{th} \) and \( l^{th} \) solids phases

\( J \) = Objective Function

\( P \) = Gas pressure

\( P_s \) = Solid pressure

\( R_{gn} \) = Mass transfer from each of solid phases to the gas phases

\( R_{smn} \) = Mass transfer from each of gas phases to the solid phases

\( \vec{S}_g \) = Gas-phase stress tensor

\( \vec{S}_{sm} \) = Stress tensor of the \( m^{th} \) solid phase

\( u_g \) = Velocity of gas in x direction

\( u_s \) = Velocity of solid in x direction

\( v_g \) = Velocity of gas in y direction

\( \vec{v}_g \) = Gas velocity vector

\( v_s \) = Velocity of solid in y direction

\( \vec{v}_{sm} \) = Solid Velocity Vector

\( w_g \) = Velocity of gas in z direction

\( w_s \) = Velocity of solid in z direction

\( y_{actual} \) = The actual value which is given by CFD simulator

\( y_{predicted} \) = The predicted value which is calculated by ANN

\( \varepsilon_g \) = Gas volume fraction

\( \varepsilon^* \) = Maximum packing volume fraction

\( \varepsilon_{sm} \) = Solid volume fraction

\( \rho' \) = Apparent Solid Density

\( \rho_g \) = Gas density

\( \rho_{sm} \) = Solid density
ACKNOWLEDGMENT

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<td>Artificial Intelligence</td>
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<tr>
<td>ANN</td>
<td>Artificial Neural Network</td>
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<tr>
<td>CFD</td>
<td>Computational Fluid Dynamics</td>
</tr>
<tr>
<td>CMG</td>
<td>Computer Modeling Group</td>
</tr>
<tr>
<td>CSV</td>
<td>Comma Separated Value</td>
</tr>
<tr>
<td>DM</td>
<td>Data Mining</td>
</tr>
<tr>
<td>EIA</td>
<td>Energy Information Administration</td>
</tr>
<tr>
<td>IGCC</td>
<td>Integrated Coal Gasification Combined Cycle</td>
</tr>
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<td>KPI</td>
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<td>NETL</td>
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Chapter 1 Introduction

The global warming is becoming a critical issue nowadays and pushes the engineers more toward the clean and environmentally-friendly fuels. Coal is a major fuel for the power plants. According to the EIA\(^1\) report in April 2016, 33\% of the U.S. electricity was generated using coal, and coal has a huge contribution to greenhouse gas emission. Integrated Gasification Combined Cycle (IGCC) has the potential to reduce the CO\(_2\) emissions of carbon based power plants by converting coal into synthesis gas (also known as syngas) while capturing CO\(_2\) for storage thereby reducing the environmental impact of power generation. Hydrogen gas generated, as the results of the gasification process can be used to power gas turbines in power generation industry \([1]\). Modeling and simulation capabilities reduce the cost and time to market of new technologies like IGCC by reducing costly design and scale-up testing.

1.1. Problem Statement

A coal based power plants (IGCC\(^2\)) are getting popular in power generation facilities. There are about 160 gasification plants in operation in all over the world \([2]\). IGCC consists of several parts; feed system, gasifier, gas clean-up system, and heat exchanger. The heart of an IGCC power plant is a gasifier. Understanding the hydrodynamics inside a gasifier allows for achieving optimum design. Basically, there are three approaches to simulate a process and find out the characteristics of the process; First method is to create a very simple mathematical model and solve the governing equations analytically, second method is to create a mathematical model with more complexities and discretize the domain in time and space and solve the equations numerically. The third approach is to build a prototype (usually in a smaller size) and do some experiments on a smaller scale and perform the upscaling to obtain the actual characteristics in the desired scale.

It is near impossible to capture all the properties of the gasification process with a simple analytical model since analytical models apply simplified assumptions which may eliminate the major characteristics of the problem. It is also extremely challenging to design a small experimental prototype that can mimic the gasification process at very high temperature \([3]\) and high pressure and take detailed measurements for improving the performance of the gasifier.

---

\(^1\) U.S. Energy Information Administration  
\(^2\) Integrated Coal Gasification Combined Cycle
The recommended method is to take the advantage of computational science and using numerical simulation to predict the main features of this process. The complexity of this problem is mainly due to presence of multiple phases (gas and solid) and the energy and mass transfer between these phases. In a complete gasification problem, as many as twenty-two equations should be solved simultaneously including mass conservation (2 equations), momentum equation (6 equations), energy balance (2 equations) and species mass fraction (remaining 12 equations) [4], [5]. Gasification is a transient process with a high degree of non-linearity and chaos that increases the computational cost of the process dramatically. Modeling this process by CFD solver typically is very time-consuming. As an example, modeling of a very simple case without any reaction and mass transfer takes about 4 days on super-computers with several clusters just to simulate 10 seconds of the actual time. Furthermore, by adding more complexities to the system, the run-time will be increasing exponentially.

This is where there is a critical need to develop new data-driven smart proxies that can help engineers to reduce the time needed for simulating complex fluid dynamics problems such as coal gasification. This method takes advantage of machine learning algorithm and artificial intelligence to come up with a powerful tool to predict the behavior of a system with far less computational cost compared to traditional CFD solvers.

1.2. Objective

The commercial CFD software’s which are available in the market such as MFIX, Open-FOAM, and COMSOL need several days to several weeks to simulate a simple fluidization problem. This makes the optimization of any process that requires multiple runs extremely computationally intensive and expensive. Therefore, engineers working in the field of fluid dynamics are looking for new techniques and tools that have the same capability but with faster turnaround time and much lower computational cost. Artificial intelligence and big-data analytics have a proven record in replicating the simulation results of complex problems with huge data sets such as those introduced in oil and gas industry by Mohaghegh and his team. The same techniques Artificial Intelligence (AI) and Data Mining (DM) can be used in a new frame work to develop a new tool that is able to completely replicate numerical simulation of CFD problems with the same accuracy and in much shorter time. The new approach requires huge amount of data, so the pattern
recognition techniques and machine learning algorithms can be used to learn the behavior of the system from the given data.

The main objective of this research is to create a surrogate model that can replicate the CFD numerical simulation results for a non-reacting fluidized bed. The final goal of this project is to deliver a software package containing a system of artificial neural networks which is able to do the same job as CFD simulators does at much faster speed and lower computational cost.

1.3. Chapter Review

This proposal includes four chapters. In chapter one, the problem was defined and the final objective of the research was determined.

In chapter two, an introduction about the gasification will be presented. A brief definition of multiphase flow and its governing equations will be provided. Also, a literature review that has been conducted about using smart proxies in the fluid dynamics problems, will be reviewed.

Chapter three will be discussing the methodology and the machine learning method which is used in this thesis. The artificial neural network with all the required information will be introduced. The network architecture with all input and output system will be discussed.

Results and discussions will be also presented in the fourth chapter. The conclusion will be made in the fifth chapter. And finally, this thesis will end up with recommendations and suggestions for the new works.
Chapter 2 Background

In this chapter, detailed discussions on three main factors of this project will be presented. First, the gasification process will be introduced. Then, the fundamental of fluid dynamic equations and their application in gasification process will be presented. Finally, the application of machine learning and pattern recognition in this project will be introduced.

2.1. Gasification

Gasification is the process by which feedstock such as coal or biomass (dried plants, woods, and farm waste) is converted into synthesis gas (also called syngas). Syngas is mainly composed of CO, H\textsubscript{2} and CO\textsubscript{2} along with contaminants such as H\textsubscript{2}S [6]. Primary species such as H\textsubscript{2} and CO can be used in power generation or chemical processing industries. Figure 2-1 shows the gasifier vessel where the gasification takes place.

![Gasifier with inputs and outputs](image)

Gasifiers are classified into three main categories: fixed bed gasifiers, entrained flow gasifiers and fluidized bed gasifiers. A fluidized bed gasifier can efficiently mix the feedstock (coal) particles, which may be at different stages of gasification. An advantage of a fluidized bed gasifier is the nearly uniform temperature that can be achieved in the reactor. In a fluidized bed gasifier, a mixture
of steam and oxidizer (air) is introduced at the bottom of the gasifier, with coal particles injected along the side of the reactor. After the initial moisture in the coal is released, the coal particles undergo devolatilization, where the volatile gases, such as CO, CO₂, H₂, H₂O, CH₄ to name a few, that are trapped in the coal are released. In the next step, carbon in coal and oxygen react to produce carbon monoxide and heat according to equation (2-1). Carbon monoxide further reacts with oxygen to produce carbon dioxide and heat, according to equation (2-2).

\[
C + \frac{1}{2}O₂ \rightarrow CO \quad (2-1)
\]

\[
CO + \frac{1}{2}O₂ \rightarrow CO₂ \quad (2-2)
\]

The heat of combustion generated is key in maintaining the gasification process, which is an endothermic process. Steam and carbon dioxide gasification takes place, when carbon in coal reacts with H₂O and CO₂ to produce carbon monoxide and hydrogen according to reactions (2-3) and (2-4).

\[
C + H₂O \rightarrow CO + H₂ \quad (2-3)
\]

\[
C + CO₂ \rightarrow 2CO \quad (2-4)
\]

Depending on the gasification goal, CO and H₂ in the syngas can be directed to a water-gas shift reactor, where the ratio of CO to H₂ produced can be altered according to reaction (2-5).

\[
CO + H₂O \leftrightarrow CO₂ + H₂ \quad (2-5)
\]

Gasification process also has its own complexity and concerns. One of the major issues of this process is handling and processing the solids. On the other hand, simulation of fluid dynamics in a gasifier is a tedious task due to multi-phase nature of flow and reaction in a high temperature and pressure.

2.2. Multiphase Flow

Transport in the gasification process consists of two flow regimes; coal transport and smoke transport. This is a challenging problem because of the multi-phase nature of the process and
heterogeneous fluid and particle interactions. Traditionally, the modeling and optimization of multiphase flow is performed by empirical modeling based on the experimental observation without going into the detail of physics and mathematics. The empirical modeling needs large investment for data gathering and data processing. Data gathering is itself a big challenge in gasification vessel since it is hard to measure the flow properties inside the gasifier due to its high temperature. Multiphase computational fluid dynamics does not have these difficulties, it also has a lot of flexibilities to tweak the system characteristics to optimize the process.

Basically, there are two different modeling scheme; Eulerian and Lagrangian. Eulerian refers to bulk fluid simulation that has a fix coordinate system while Lagrangian refers to particle tracking simulation where there is a coordinate assigned to each particle and moving with particle velocity with respect to fixed coordinate.

- Continuum (Eulerian): Select a fix control volume and assume the flow is continuum and use the Navier-Stokes equation by averaging the properties over the volume. Continuum approach needs less computational time but it cannot capture all the complexities, especially in multiphase flow where interaction between particles plays a major role [8].
- Discrete Particle (Lagrangian): Track each particle in the fluid by using Newton’s Law of motion. This method is more straightforward to apply, even in multiphase flow, but the computational costs is high [8].

There are several approaches to modeling multiphase flows. Depending on the application, either the gas phase or the solid phase or both phases can be modeled in an Eulerian or a Lagrangian framework [8]–[10]. Table 2-1 shows different approaches to model multiphase flows.

<table>
<thead>
<tr>
<th>Name</th>
<th>Gas Phase</th>
<th>Solid Phase</th>
<th>Coupling</th>
<th>Scale</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Discrete bubble model</td>
<td>Lagrangian</td>
<td>Eulerian</td>
<td>Drag Closure for bubbles</td>
<td>10 m</td>
</tr>
<tr>
<td>2 Two Fluid Model</td>
<td>Eulerian</td>
<td>Eulerian</td>
<td>Gas-Solid drag closure</td>
<td>1 m</td>
</tr>
<tr>
<td>3 Unresolved Discrete particle model</td>
<td>Eulerian</td>
<td>Lagrangian</td>
<td>Gas-particle drag closure</td>
<td>0.1 m</td>
</tr>
<tr>
<td>4 Resolved Discrete particle model</td>
<td>Eulerian</td>
<td>Lagrangian</td>
<td>Boundary condition at particle surface</td>
<td>0.01 m</td>
</tr>
<tr>
<td>5 Molecular Dynamics</td>
<td>Lagrangian</td>
<td>Lagrangian</td>
<td>Elastic collisions at particle surface</td>
<td>&lt;0.001 m</td>
</tr>
</tbody>
</table>
Coal gasification includes two phases; first phase is gas (oxygen and sometimes water vapor) and the second phase is solids (coal). The governing equations of the motion are partial differential equations. To completely model this process, coupling of those equations for mass transport, energy transport, and momentum equation for two phases should be considered. Solving the coupled partial differential equations analytically is not an easy task, so the numerical method should be utilized to solve the equations.

CFD (computational fluid dynamics) is a combination of numerical analysis, fluid dynamics, and computer science. The governing partial differential equations for the fluid flow could be solved more easily by CFD method which is a tool to replace PDE by a set of linear algebraic equations. There are several commercial CFD simulator available in the market with different capabilities in different aspects. Among those software, MFiX has been chosen for this project to simulate fluid and solid dynamics in gasifier.

Modeling the multi-phase fluid dynamics inside a gasifier is extremely complex. To model the gasification process, not only the hydrodynamics of gas and solid flow have to be modeled by solving for gas and particle transport equations, also the chemical reaction processes such as devolatilization, coal combustion and gasification, homogeneous oxidation and water gas shift reactions have to be modeled as well Often times it’s difficult to make detailed measurements inside a gasifier, due to the high temperature, high pressure and harsh conditions in a gasifier. Simulations, therefore, provides an alternative to designers and engineers to access the performance of a gasifier.

Watanabe and Otaka [3] used CFX-4 CFD software to investigate the effect of air flow rate and coal type on the performance of a gasifier. The purpose of their study was to optimize the performance of the gasifier by changing the air ratio and coal type.

Papadopoulos et al., [7] used a ANSYS-CFX CFD code to develop a high pressure, high temperature model. He used the same concepts and equations as Watanabe and Otaka did.
2.2.1. MFiX

MFiX is an open source multiphase CFD software developed and maintained by NETL\textsuperscript{1}. MFiX is able to handle heat and mass transfer and chemical reactions in fluid-solids systems. It has been used to model the bubbling, circulating fluidized beds, and spouted beds. The output of MFiX consists of transient information on the distribution of volume fractions, pressure, velocity, temperature, and species mass fractions of each phase being considered. MFiX has its own post-processing tool to visualize the results, but it is compatible with multiple software such as Paraview [4], [5].

2.2.2. Governing Equation

In this section, all the equations which MFiX uses to simulate fluid and solid dynamics are reviewed. There will not be any detailed discussion on the derivation of the equations and the purpose of providing these equations is to introduce all the important parameters that later on will be used in the machine learning algorithm as input. Moreover, the step by step algorithm that MFiX uses to solve the problem will be discussed since this will play an important role for the machine learning when implicit prediction is utilizing.

2.2.2.1. Conservation of mass

MFiX can handle one gas phase and multiple solid phases, so there is one scalar equation for conservation of gas and multiple equations for conservation of each solid phase [11].

Equation (2-6) shows the conservation of mass of gas phase

\[
\frac{\partial}{\partial t} (\varepsilon_g \rho_g) + \nabla \cdot (\varepsilon_g \rho_g \vec{v}_g) = \sum_{n=1}^{N_g} R_{gn}
\]  

(2-6)

Where \(\varepsilon_g\) is the gas volume fraction, \(\rho_g\) is the gas density, \(\vec{v}_g\) is the gas velocity vector and \(R_{gn}\) is mass transfer from each of solid phases to the gas phase, this mass transfer could be due to chemical or physical processes such as evaporation and reaction.

\textsuperscript{1} National Energy Technology Laboratory
As mentioned, MFiX can handle multiple solid phases. Equation (2-7) shows the conservation of mass of $m^{th}$ solid phase.

$$\frac{\partial}{\partial t}(\varepsilon_{sm}\rho_{sm}) + \nabla \cdot (\varepsilon_{sm}\rho_{sm}\vec{v}_{sm}) = \sum_{n=1}^{N_{sm}} R_{smn}$$ (2-7)

Where $\varepsilon_{sm}$ is the solid fraction of $m^{th}$ solid phase, $\rho_{sm}$ is the density of $m^{th}$ solid phase, $\vec{v}_{sm}$ is the solid velocity vector and $R_{smn}$ is mass transfer from gas phase to the solid phase.

### 2.2.2.2 Conservation of Momentum

The gas-phase momentum balance is expressed by equation (2-8).

$$\frac{\partial}{\partial t}(\varepsilon_g \rho_g \vec{v}_g) + \nabla \cdot (\varepsilon_g \rho_g \vec{v}_g \vec{v}_g) = \nabla \cdot (\vec{S}_{g}) + \varepsilon_g \rho_g \vec{g} - \sum_{m=1}^{M} I_{gm} + \vec{f}_g$$ (2-8)

Where $\vec{S}_{g}$ is the gas-phase stress tensor, $I_{gm}$ is the interaction force representing the momentum transfer between the gas phase and the $m^{th}$ solids phase, and $\vec{f}_g$ is the flow resistance offered by internal porous surfaces.

The momentum equation for the $m^{th}$ solids phase is

$$\frac{\partial}{\partial t}(\varepsilon_{sm}\rho_{sm}\vec{v}_{sm}) + \nabla \cdot (\varepsilon_{sm}\rho_{sm}\vec{v}_{sm}\vec{v}_{sm}) = \nabla \cdot (\vec{S}^{m}_{sm}) + \varepsilon_{sm}\rho_{sm}\vec{g} - \sum_{l \neq m}^{M} I_{ml} + I_{gm}$$ (2-9)

Where $\vec{S}^{m}_{sm}$ is the stress tensor for the $m^{th}$ solid phase, $I_{gm}$ is the interaction force representing the momentum transfer between the gas phase and the $m^{th}$ solids phase, and $I_{ml}$ is the interaction force representing the momentum transfer between the $m^{th}$ and $l^{th}$ solids phases.

All the important parameters describing the gasification process can be extracted using equations (2-6) to (2-9) and equations described in appendix I. Table 2-2 shows the key parameters involved in the simulation process, this information will be used in the next chapter to define the input parameters for the artificial neural network.
Table 2.2- Important parameters for multiphase flow

<table>
<thead>
<tr>
<th>Key factors of gas-solid system</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas Density ($\rho$)</td>
</tr>
<tr>
<td>Volume fraction ($\varepsilon$)</td>
</tr>
<tr>
<td>Particle diameter ($d$)</td>
</tr>
<tr>
<td>Maximum packing volume fraction ($\varepsilon^*$)</td>
</tr>
<tr>
<td>Velocity vector of gas ($u, v, w$)</td>
</tr>
<tr>
<td>Velocity vector of solid ($u, v, w$)</td>
</tr>
<tr>
<td>Pressure field of gas ($P$)</td>
</tr>
<tr>
<td>Pressure field of solid ($P_s$)</td>
</tr>
<tr>
<td>Time ($t$)</td>
</tr>
<tr>
<td>Location to the boundaries ($x, y, z$)</td>
</tr>
<tr>
<td>Location to the interface ($x, y, z$)</td>
</tr>
</tbody>
</table>

2.2.3. MFiX solution Algorithm

Equations (2-6) to (2-9) form a system of nonlinear partial differential equations. In order to solve this system of PDEs, a step by step iterative algorithm has been developed by MFiX solver. The sequence of solving the equations and calculating the parameters are important for this project in order to establish the best solution scenario. Each solution scenario has its own approach, in some of them (implicit scenarios), the sequence of calculating of parameters is so important. The MFiX solution algorithm might be a good start point for the implicit scenarios. There will be a detailed discussion about different solution scenarios in the next chapter. Figure 2-2 shows the step by step algorithm which is used by MFiX to solve the system of coupled PDE equations.
2.3. Machine Learning

Based on Arthur Samuel (1959) definition, “Machine learning is a field of study that gives computers the ability to learn without being explicitly programmed.”

Machine learning is a process through which computer will learn from data to find a possible pattern in the data set. This process encompasses three main components as follows:

- Learning algorithm
- Data
- A pattern in the data
If these three components are present, a successful learning process can be achieved based on the capability of the learning algorithm. There are two major type of Machine Learning: supervise learning and unsupervised learning [12].

### 2.3.1. Supervised Learning

In supervised learning, some data, is provided as input and output to the learning algorithm, with the goal of finding the relationship between input and output. There are two general types of supervised learning; Regression and Classification. When the output data is in continuous form, regression should be used to find the trend between input and output. This trend could be linear or nonlinear based on the problem characteristics. When for different inputs, there is finite possible output, the classification should be considered. For example, the type of cancer (malignant, benign) could be classified based on the age of the patient and size of the tumor.

### 2.3.2. Unsupervised Learning

In unsupervised learning, there is little or no information about the output. The learning algorithm tries to find the pattern between input data without having the output. This process is named clustering. For example, grouping the customer of a company based on the type of product that they buy daily.

### 2.3.3. Artificial Neural Network

One of the popular machine learning processes is Artificial Neural Network (ANN). The idea of ANN came from the neurons of the brain and the way they are communicating with each other to solve a problem. Each artificial neural network consists of an input layer, one or more hidden layers, and an output layer. The number of output and input layers are chosen based on the problem and the property which is going to be predicted. Figure 2-3 shows a typical ANN with three inputs and two outputs. ANN has one or more hidden layers and each layer has a specific number of neurons [13].

In order to have a well-trained network, proper parameters should be introduced to the network. If improper data are used to train the network there is no guarantee to have a well-trained network that lead to correct predictions, in other words, “Garbage in, Garbage out.” In chapter three, a smart way of picking parameters will be introduced.
The number of hidden layers and the neurons in each hidden layer depends on the complexity of the problem, number of parameters, and number of records. Experience also plays a role in this decision. But generally, there is no solid rule for them. As a rule of thumb, the number of neurons in the first hidden layer shouldn’t be less than the number of input parameters.

2.3.4. Objective function

Regardless of the learning method, each machine learning process needs an optimization procedure that helps the process reduce the error as much as possible. The very common and simple objective function in supervised learning is the summation of all the differences between predicted values by the learning method and the actual values of output. Sometimes the negative errors cancel the positive errors and the total error becomes very small although none of the data points have good predictions, by calculating the square of the differences, the mentioned problem could be resolved. Equation (2-10) shows this objective function [13].

\[
J(\theta_0, \theta_1) = \frac{1}{2m} \sum_{i=1}^{m} (y_{actual} - y_{predicted})^2
\] (2-10)

During the learning process, the learning algorithm tries to assign different weights to each of the lines in Figure 2-3, in a way that the global error of objective function becomes minimum. Also, a
blind validation is done simultaneously to stop the learning process. We will discuss the validation and test in more depth in the next chapter.

2.4. Previous work done in this area

The idea of using AI in petroleum engineering was first introduced by S. Mohaghegh [14]. He took advantage of ANN for predicting the permeability of the formation based on geological well logs. He showed that neural network is capable of making the task of permeability determination automated rather than doing it over and over by log analyst. He also stated that neural network can handle far more complex tasks. He also used ANN for predicting gas storage well performance after hydraulic fracture in the same paper and his later investigations [15].

Alizadehdakhel et al.[16] used ANN to predict the pressure loss of a two-phase flow in the 2 cm diameter tube. In two-phase fluid, separation of the phases may occur because different phases may have different velocities, so the traditional Navier-Stokes equation is not capable of finding the exact pressure drop in different flow condition. The authors used three main property of the fluid (gas velocity number, liquid velocity number, and line slop\(^1\)) as the input of the ANN and only one output which was the average pressure drop. They utilized 8 different networks with a different number of neurons to find out the optimum number neurons. MSE and R-square were used as a criterion to pick the best network design. They also tried to find the most efficient transfer function between Log-Sigmoid, Hyperbolic-Tangent Sigmoid, and linear. The data had come from the experimental setup that they had built. The pressure in upstream and downstream of the pipe was measured and the pressure loss was calculated.

Shahkarami et al. [17] took advantage of ANN to model the pressure and saturation distribution in a reservoir which was used for CO\(_2\) sequestration purpose. This problem required a large number of time steps for simulation of CO\(_2\) injection and storage using commercial software. They ran 10 different cases in CMG (commercial reservoir simulator) and then the results were used as input for ANN. The output of the ANN was selected to be pressure distribution, water saturation, and CO2 mole fraction. 80% of the data coming from CMG simulation were used to train the network, 10% were used for the validation and the rest of the data were used for the test process. They have

\(^1\) Pressure drop in one meter, \(\Delta P/L\) (Pa m\(^{-1}\))
shown that ANN can be used as a powerful tool for multiphase flow simulation in oil and gas industry.

Esmaili and Mohaghegh [18] introduced a new way of using completion and production data with the well logs in order to find out the shale reservoir behavior under certain condition. By understanding the behavior of the shale reservoir, conducting the hydraulic fracture could be much easier. Moreover, this method has the ability to perform the history matching on the production data.

Kalantari-Dehghani et al. [19] coupled reservoir numerical simulator with AI method to develop a shale proxy model that is able to regenerate a numerical simulation results in just a few seconds. They introduced three different well-based tier systems to achieve a comprehensive input data for the ANN. In another research [20], they showed that data-driven proxy models at the hydraulic fracture cluster level could be used separately as a reservoir simulator especially in low permeability reservoir such as shale which has a nonlinear behavior.
Chapter 3 Methodology

In this chapter, the methodology of solving a problem in the field of computational fluid dynamics will be discussed in detail. First, the problem will be defined with all the initial and boundary conditions. Then the modeling process in MFiX (commercial CFD simulator) will be explained. Creating the input of the neural network is the next discussion in this chapter that is the most important step of data training. By knowing all the provided information from previous chapters, the neural network model will be created and the training will be performed.

3.1. Defining the problem

The gasification process is a very complicated problem. In order to demonstrate the usefulness of using ANN to create proxy models for the gasification process, first a proof of concept study has been carried out to show that ANN can be applied to a simple CFD simulation of a flow inside a rectangular fluidized bed.

Figure 3-1 shows the geometry of the problem which is a rectangular fluidized bed with a square cross section. The dimension of the fluidized bed is depicted in Figure 3-1. One-sixth of the fluidized bed is filled with sand which is initially at rest. The sand particles are perfect spheres with the density of 1160 kg/m$^3$. The initial volume fraction of gas (voidage) is 0.42.

The $y$-component of the initial velocity of the air inside the fluidized bed (where sand is located) is 1.43 m/s and the $y$-component of the initial velocity of the air in the freeboard region (above the sand) is 0.6 m/s.

The inlet air velocity is set to 0.6 m/s that is uniformly distributed in the upward direction. The air discharges into atmospheric pressure up on top of the bed.

As air is injected into the bed, initially large gas pockets form that cause sand particles to move upward as a slug. In matter of few seconds, the gas pockets break up, leading to the breakup of sand slugs, which shower back down. At this time, air entering the bed forms bubbles, which move up through the sand bed, leading to the fluidization of the sand particles.
3.2. MFiX

The geometry setup in the previous section, is simulated in MFiX. The output data generated by MFiX is used as the input data to the ANN.

The model has been created and ran successfully. The next step is to get the results from MFiX and organize the data in order to make the data ready for the ANN. Since MFiX reports the results based on the grids, the order and exact location of each grid is extremely important for ANN. The output file of MFiX has an extension of *.vtu for each time-step which needs to be converted to
*.csv files. ParaView is an open source software which can be used for data visualization and format conversion.

3.2.1. Grid system

Several mesh resolutions could be considered depending on the desired accuracy. Table 3-1 shows different grid sizes and the number of cells associated with each grid size. The very fine grid system with 118,098 cells was selected for this study.

<table>
<thead>
<tr>
<th>Grid Classification</th>
<th>Cell size</th>
<th>No. of Cells</th>
<th>No. of Nodes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coarse</td>
<td>8<em>48</em>8 (15 mm)</td>
<td>3,072</td>
<td>3,969</td>
</tr>
<tr>
<td>Medium</td>
<td>12<em>72</em>12 (10 mm)</td>
<td>10,368</td>
<td>12,337</td>
</tr>
<tr>
<td>Fine</td>
<td>18<em>108</em>18 (6.6 mm)</td>
<td>34,992</td>
<td>39,349</td>
</tr>
<tr>
<td>Very Fine</td>
<td>27<em>162</em>27 (4.4 mm)</td>
<td>118,098</td>
<td>127,792</td>
</tr>
</tbody>
</table>

It is important to understand the output files structure of MFiX and understand the format of each file and the order information that is reported. For this purpose, the grid system should be completely known. Figure 3-2 shows the numbering order of the grids. The numbers start from the origin of the coordinates and moves in the y-direction first, as it gets to the last grid in y-direction it moves in the x-direction and goes to the next column. After the first layer numbering is completed, it moves in z-direction to the second, third and all the way to the last layer.

3.3. Artificial Neural Network Setup

The output file of MFiX which was converted to *.csv file is ready to be reorganized to serve as the input for the ANN. Every time-step has one *.csv file containing 9 columns and 118,098 rows. Each column indicates one property and each row corresponds to one cell. Table 3-2 shows those 9 parameters. The input of the ANN is all the data at time-step t while the output will be one or more parameters of time-step t+1. In this approach, the network will learn what the output should be given a set of input data. When the learning process gets done, the deployment process (prediction) will be performed.
Table 3-2- All the useful parameters reported in the MFiX result file

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\varepsilon_g$</td>
<td>Gas volume fraction</td>
</tr>
<tr>
<td>$P$</td>
<td>Gas Pressure</td>
</tr>
<tr>
<td>$P_s$</td>
<td>Solid Pressure</td>
</tr>
<tr>
<td>$u_g$</td>
<td>Velocity of gas in x direction</td>
</tr>
<tr>
<td>$v_g$</td>
<td>Velocity of gas in y direction</td>
</tr>
<tr>
<td>$w_g$</td>
<td>Velocity of gas in z direction</td>
</tr>
<tr>
<td>$u_s$</td>
<td>Velocity of solid in x direction</td>
</tr>
<tr>
<td>$v_s$</td>
<td>Velocity of solid in y direction</td>
</tr>
<tr>
<td>$w_s$</td>
<td>Velocity of solid in z direction</td>
</tr>
</tbody>
</table>

3.3.1. Tier System

In order for the ANN to learn in an effective manner, a tier system has been developed. Each cell is in contact with 28 surrounding cells; 6 of them have the surface contact with the original cell,
12 of them have line contact with the original cell, and 8 of them have point contact with the original cell.

Like any numerical method, the values of each cell has a relation with the value of the surrounding blocks. With that idea in mind, the ANN will not only learn from the 9 parameters (Table 3-2) of the cell, it will also learn from the surrounding cells which are called “Tier”. There are several tiers at the neighbor of each cell and depending on the complexity of the problem, one can use tier 1 (surface contact), tier 2 (line contact), and tier 3 (point contact). Figure 3-3 shows a tier 1 structure, where the main cell is surrounded by its 6 neighboring cells. For this case, the 9 parameters of the original cell and 9 parameters of the tier 1 cells make 63 different parameters, which are the input for the ANN. Depending on the complexity of the problem and spatial and temporal correlations between different tiers and the center cell more or less input parameters might be required.

3.3.2. Input Matrix

It is not enough to consider only the values of each parameter in a center cell and related tiers in the input matrix, but for the network to learn the behavior of the process and perform pattern recognition step, the location of each cell in the geometry is also crucial. Adding the location as an input helps the system understand the spatial correlation between different parameters, as well. On the other hand, wall has a large effect on the flow pattern and the location of wall should be
somehow included into the ANN. To accommodate these ideas, six different distances to the wall confinements (top, bottom, east, west, north, and south) are considered to define the exact location of each cell and parameters associated with each cell. By adding these 6 distances to the previous 63 parameters, a total number of parameters used as input becomes 69. So, the dimension of input matrix is 69 by 118,098 (i.e., number of parameters multiply by the number of cells).

3.3.3. Neural Network Architecture

Each artificial neural network consists of an input layer, one or more hidden layers, and an output layer. The inputs and outputs are chosen based on the nature of the problem and the property which is going to be predicted. In the last section, the number of input parameters were selected to be 69. The output of the ANN could be only one parameter, or it could be more than one parameter. There will be different scenarios to compare different ANN with different number of output parameters. There is no clear guideline on how many hidden layers and neurons are required at each layer and it is basically chosen based on the problem and experience. The only rule of thumb is that, the number of neurons in the first hidden layer shouldn’t be less than the number of input parameters.

For the first try, only one hidden layer with 100 neurons is considered. 69 parameters as input and only one parameter as output were selected. All the required numbers to define an ANN are chosen and shown in Table 3-3.

<table>
<thead>
<tr>
<th>Number of Inputs</th>
<th>69</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of hidden layers</td>
<td>1</td>
</tr>
<tr>
<td>Number of Hidden Neurons</td>
<td>100</td>
</tr>
<tr>
<td>Number of records</td>
<td>118,098</td>
</tr>
<tr>
<td>Number of Output</td>
<td>1</td>
</tr>
</tbody>
</table>

The network characteristics are defined and shown in the Table 3-4. Feed-forward back propagation method is using for the training. The transfer function for hidden layer and the output layer was chosen to be TANSIG which is depicted in the Figure 3-4.
Table 3-4: Neural Network characteristics

<table>
<thead>
<tr>
<th>Network Type</th>
<th>Feed-forward Back propagation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Training Function</td>
<td>Levenberg-Marquardt</td>
</tr>
<tr>
<td>Adaption Learning Function</td>
<td>LEARNGDM</td>
</tr>
<tr>
<td>Performance Function</td>
<td>MSE</td>
</tr>
<tr>
<td>Transfer Function</td>
<td>TANSIG</td>
</tr>
</tbody>
</table>

Figure 3-4: Neural network transfer function (TANSIG)

3.3.4. Data Partitioning

If all the data is used for the training, the network will learn perfectly for the given dataset, but it might not be good to use for new dataset, since the goal of ANN is to predict the same problem but with a new database. This problem is called overfitting. Overfitting occurs when the network learns to mimic the exact data (that was used in the training process) but it is not general enough to predict the new dataset of the same problem. To overcome the overfitting problem only a portion of it is used to train the network. Depending on the nature of the problem, a different percentage can be assigned for the training purpose and remaining data is then used for the validation and test. The validation is a kind of blind test, which is done while training the network. In the test process, the rest of data will be used to check the performance of the network after training. The percentage of the data prioritization used for the preliminary study of this research is shown in Table 3-5. It is important to mention that this partitioning is the preliminary one and a deeper study will be conducted on the percentage of the data in section 3.5.6.3.

Table 3-5: Original Data Partitioning

<table>
<thead>
<tr>
<th>Data</th>
<th>Training</th>
<th>Validation</th>
<th>Test</th>
</tr>
</thead>
<tbody>
<tr>
<td>Percentage of data (%)</td>
<td>70</td>
<td>15</td>
<td>15</td>
</tr>
</tbody>
</table>
3.4. Spatio-Temporal Database

In order to build the smart proxy, a fluidized bed problem has been modeled using MFiX. The result has been used to create a Spatio-Temporal Data base. This provided a comprehensive database for fluidized bed problem.

![Figure 3-5: Spatio-Temporal Database and optimized database](image)

The Spatio-Temporal database is created based on the data from one single time step for the first attempt, and later, more time steps will be added to the database (will be discussed in section 3.5.5). The Spatio-Temporal database includes all the 9 different parameters for each cell and its neighbors from a certain time step, plus the exact time (simulation elapsed time) of that time step and the location of each cell. By this definition, the model ended up having 70 parameters. The Spatio-Temporal database treats each cell as a separate record, so the model has 118,098 records which is equal to the number of cells.

This database should be sent to an optimization process to get an optimized database with the smaller size but the same efficiency.

3.5. Solution Scenarios

Different scenarios are considered to reach the final goal of this project. The term “Different scenarios” refers to have different input and output structures and also using different time-steps for the training, while the training technique is the same in all the scenarios. Depending on what time-steps and how those time-steps are going to be used for the training, different scenarios will be designed which is the main discussion of the following section.

Each scenario has two parts, first is the training process and second is the deployment process. A pair of time-steps is used in the training process. The training process stops based on the criteria that the user determines. This criterion could be the total number of iteration, the total time of
training, or the number of validation failure or a combination of those (In this project, the combination of all the mentioned criteria was used). The learning algorithm is such that the network learns more and more as it goes through each iteration but in order to avoid overfitting or memorization, validation error is always checked. If the validation error increases for a predefined number of iterations, the training stops. Most of the time, validation is the criterion which makes the training stop.

As mentioned in the previous sections, 69 parameters are used as the input for the ANN. Figure 3-6 shows all the 69 input parameters including 6 distances to the boundaries and 9 properties for the orange cell and also 6x9 set of parameters for tier blocks. The network also needs the output to be trained. In this problem, there are total 9 parameters which any of those could be the output of ANN. The output could be one parameter at a time or multiple parameters.

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Figure 3-6- 69 different parameter of ANN

The trained network is then ready for the deployment process. One time-step is given to the trained network and the network will give its prediction for the next time-step. The input of the ANN for each deployment could come from the CFD directly or from the ANN itself. Cascading and non-cascading deployment are defined based on what type of input is used for the network and it will be discussed in detail in the following sections.

3.5.1. Early time versus late time

In this scenario, the 69 inputs come from time-step t and the output is from time-step t+1. The output could be one parameter or multiple parameters, which in this case, only one output is used at the same time (Figure 3-7). The main question here is which time-step should be used for the training since there are multiple time-steps available. For the preliminary runs, one time-step from the early time is used for the training when the motion in the system is like a slug flow and no bubbles are in the fluidized bed. Figure 3-8 shows a pair of the time-steps for training.
For the second try, one pair of time-steps is chosen from the late time when the flow is completely chaotic and bubbles are everywhere in the system. Figure 3-9 shows the input/output pair of time-steps for this scenario.

The reason for choosing these two training (early time and late time) is because there are two different flow regimes at work in these time-steps. Figure 3-10 shows the distribution of solids in the fluidized bed in the early time and late time. This figure shows two complete different motions in the system. The color bar is the gas volume fraction (voidage); all the figures are generated by MATLAB.
The purpose of this analysis is to show that the ANN is capable of capturing all the physics involved in different time-steps. In the next chapter, complete results of this analysis will be presented and discussed in detail.

![Gas volume fraction distribution on the wall; early time (a) versus late time (b)](image)

**Figure 3-10: Gas volume fraction distribution on the wall; early time (a) versus late time (b)**

### 3.5.2. Cascading versus non-cascading

Cascading and non-cascading refer to what kind of input is used for the deployment process. If the input comes from the CFD solver for each deployment stage, it is called non-cascading. If the input of the ANN for each deployment stage comes from the output of previous deployment, it is called cascading.

Although it seems that non-cascading deployment has no benefit because the real input from CFD solver should be available for every stage, it should always be studied in order to prove that the trained network is working properly. Eventually, every parameter should be predicted by cascading method but to accomplish this goal, first non-cascading should be done.
To better understand the difference between these two approaches, two schematic figures are provided. Figure 3-11 shows the non-cascading deployment sequence while Figure 3-12 shows the cascading deployment.

Figure 3-11- The process of non-cascading deployment

Figure 3-12- The process of cascading deployment
The non-cascading and cascading deployment process is going to be completed for both early and late time and the results will be depicted in the next chapter.

3.5.3. Single output versus multiple output

As discussed earlier, ANN can have one output at the same time or multiple outputs. Obviously, having multiple outputs simultaneously increases the training time, furthermore, the network has to fit multiple outputs with the same weights, so the network has less flexibility to learn from data but sometimes it gives the better results especially when there is a correlation between the outputs. Figure 3-13 and Figure 3-14 show the input and output of the ANN when only one output is used and when 3 outputs are used, respectively.

To examine the ability of the ANN when multiple outputs are used, some different cases are considered. Selecting the set of outputs is the most important concern at this point and the main question is what outputs could be used at the same time. It is decided to have three components of gas velocity at the same time as outputs of the ANN because it is more likely for the gas velocity components to have a correlation and the chances are less for the solid pressure and gas velocity to have a correlation.

As another advantage of this approach, it should be stated that having multiple outputs at the same time would reduce the number of neural networks. As mentioned in the last section, there are total
nine different ANN needed for cascading deployment, this number could be reduced to three if each network has three outputs at the same time. The result of this approach is also available in chapter 4.

3.5.4. Explicit versus implicit

Regardless of the training scenario, the training process needs a pair of data; input and output (time-step t and time-step t+1). If all the input data come from time-step t and the output data come from time-step t+1, it is called explicit method, exactly like what is common in CFD solution methods. Figure 3-14 is a demonstration of explicit training. It is also possible to have the combination of data from time-step t and t+1 as input and have time-step t+1 as the output as well. Obviously, the parameters from time-step t+1 that has been used for input will not be used for the output; this approach is called implicit training. Figure 3-15 shows one of the examples of implicit training. The input consists of gas volume fraction, pressures, and gas velocity vector from time-step t in addition to solid velocity vector from time-step t+1. The output is gas velocity vector from time-step t+1.

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This approach is very common in the numerical solution of PDE’s, which increases the converging speed. It is expected to have a lower error when the implicit approach is applied.

3.5.5. Training with multiple time-steps

For all the training until this point of this research, only one pair of time-steps was used. Figure 3-16 shows the input and output pair for the training with single time-step. The trained network for early time-step (when a slugging flow is dominant) is valid for those time-steps that have the same characteristics as slug flow but it is not valid for the entire time range when the bed...
fluidizes. Vice versa, the trained network for the late time-step, when the bed is fully fluidized is not valid for earlier time, when slug flow was present. The question that comes to the mind is “Is there a neural network that can predict different time-steps?” In other words, “Is there a general neural network that can capture different physics involved in the system?”

When the data from early time-step is used to train the neural networks and the training process completes with a good performance, it is definitely a good answer for early time-steps but it might not be a good answer for late time-steps. If the data from two time-steps are used for the training process, the network algorithm will converge to the solution that can mimic the behavior of both time-steps. So, if more time-steps are used in training, the solution will be applicable to more time-steps and the ANN covers wider time range.

Furthermore, there are different physics involved in different time-steps of the simulation, and in order for the network to learn from all the possible behavior, more time-steps should be used in the training process. So, it is decided to use at least three different time-steps with different flow characteristics. One time-step from the early time, one time-step from late time, and one time-step from the time when the bubbles start forming. In order for the ANN to distinguish between different time-steps, another parameter (extra column) is added to the input data that is the exact time of the time-step in seconds. Figure 3-17 shows the input and output pair for the training with three different time-steps. The three time-steps were chosen visually by looking at the gas volume fraction distribution in the fluidized bed. Time-step 200 was chosen because the slugging flow is the dominant flow regime, time-step 1000 was chosen because the bubbles started forming but the flow is still symmetric, and Time-step 4000 was chosen because the bubbles were developed completely and no symmetric motion is in the system and the bed is fluidized. Figure 3-18 depicts these three time-steps.
For the result demonstration, it is not possible to present all the figures, so a different method should be used to show the quality of the ANN. In order to quantify the amount of error in each simulation, RMSE which is the square root of mean square error will be used. Equation (3-1) shows the definition of RMSE. This criterion will be also used when more time-steps is going to be added to the training dataset.

\[
RMSE = \sqrt{\frac{1}{n} \sum_{j=1}^{n} (y_{actual} - y_{predicted})^2}
\]  

(3-1)
Each time-step has 118,098 cells (27x162x27), and also there are 70 parameters, which makes the total number of data point to be equal to 8,266,860. By adding one time-step for training, more than 8 million data points will be added for the analysis. Since our computer resources are limited, the number of data points should be decreased in some fashion if more time-steps are going to be used for the training.

3.5.6. Reducing the size of the system

The size of the input is already very huge and it is not possible to include more than 2 time-steps for the training process because of the memory limitation1. The input data is a matrix which rows and columns represent records and parameters, respectively. There are a couple of ways to reduce the input size; reducing the number of rows (records) or the number of columns (parameters). There is also another way to reduce the computational cost for the training that is using fewer data for training and using more data for validation and test, in other words, by changing the data prioritization. All the mentioned method will be discussed in the next sections.

3.5.6.1. Reducing number of records

There are two ways to reduce the number of records. The first method is removing some cells just randomly from all the locations, and the second method is removing some cells that have less valuable information. For example, the solids are located on the bottom of the fluidized bed, so it is a reasonable idea to concentrate only on the bottom of the fluidized bed for all the solid-related properties since there are no solids in the freeboard portion of the fluidized bed. Figure 3-19 shows the distribution of gas volume fraction at time-step 4000. There are no solids above the blue line, so it is a good assumption to remove all the cells above the blue line. As it is shown in Figure 3-20, it is decided to keep all the cells below the 70 cells’ line and remove the cells with no significant impact in selected parameter.

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1 This Study was done on a Corei7 machine with 32 GB of RAM
Figure 3-19: Distribution of gas volume fraction at time-step 4000

Figure 3-20: The important section of the fluidized bed
3.5.6.2. Reducing number of parameters (KPI1)

The latest input data for the training had 70 parameters and 118,098 records. By concentrating on the lower part of the fluidized bed, the number of records reduced to 51,030, while the number of parameters remains the same. Some of the parameters are going to be eliminated in this section. Reducing the number of parameters is not as straightforward as reducing the number of records and it needs some analysis regarding the prioritization of the parameters. Every parameter has several weights assigned to it to communicate with the hidden layer, as it is depicted in Figure 3-21. If all the weights assigned to one parameter ($w_{11}, w_{12}, ...$) are integrated to one value ($w_1$), that value will represent the total weight and show the priority of that particular parameter when it compares to all the other total weights.

![Network schematic with its weights](attachment:network_schematic.png)

*Figure 3-21: Network schematic with its weights*

After obtaining all the total weights of the parameters, the tornado chart of each ANN could be plotted and the key parameters could be determined.

3.5.6.3. Changing the Data Partitioning

In all the previous sections, data from Table 3-5 were the base for data prioritization. According to the Table 3-5, 70% of the data is used for training, 15% is used for the validation, and the other 15% is used for the test. Changing the data prioritization percentage could reduce the

1 Key Performance Indicator
computational costs. The computational cost of the training process is mainly for the data training, and the validation and test are only a simple multiplication, so by reducing the training percentage, the computational costs could be reduced while the total amount of data remains constant.

3.5.6.4. Reducing number of records using smart sampling

Reducing the number of records was previously accomplished by concentrating on the lower section of the bed. Although using this idea led to having smaller database, still there are some points in the lower section of the bed that has less value. Eliminating less valuable records could reduce database size further. Figure 3-22 shows the distribution of gas volume fraction at time step 4000. The distribution has two peaks. The first peak is the gas volume fraction, when solid is at maximum packing (cells having fully packed solid). The second peak is when the gas volume fraction is either 1 or close to 1 (all air).

By introducing all the data for the training, the model will learn more about these two peaks and will learn less about the transition part that has the gas volume fraction between 0.42 and 1. Since it is desired for the model to learn equally from all the different locations and different events in the fluidized bed, it is decided to reduce the number of records with the value of 1 and 0.42. By this approach, a data with uniform distribution is introduced to the model, and the model will learn equally from all the features in the system. Furthermore, the cells with the value of gas volume fraction between 0.42 and 1, are located at the interface of the gas and solid phases, and the fluid dynamic characteristics are changing significantly in these location, so it is important for the model to learn about transition zone due to the dynamic change.

All the data between these two peaks were kept in the database. And only portions of data from two peaks were selected in a way that the final distribution looks like a uniform distribution. The gas volume fraction distribution is shown in the Figure 3-23. Total number of records reduced from 118,098 to 25,827.

3.6. Summary

In this chapter, the problem was defined with all details and assumptions. Also, different scenarios were introduced in order to achieve the final goal of this project. Different simulations based on the mentioned scenarios or a combination of them were designed and deployed. In the next chapter, the results of these different scenarios will be provided.
Figure 3-22 - Distribution of Gas volume fraction at time step 4000

Figure 3-23 - Distribution of Gas volume fraction at time step 4000 after smart sampling
Chapter 4 Results and Discussion

Different scenarios were introduced in the previous chapter. In this chapter, the results of all the scenarios will be discussed in detail. The results which will be demonstrated are coming from different approaches; early time or late time, single time-step or multiple time-steps for training, cascading or non-cascading deployment, single output or multiple outputs, explicit or implicit method, and reduced order models or complete models. Before proceeding with the results, there will be a short description of how the results are going to be presented.

4.1. Result Demonstration

The current problem is three dimensional in space, with time being the fourth dimension. In order to demonstrate the results, 5 different vertical cross sections were chosen as it is shown in Figure 4-1, all the planes are perpendicular to the z-axis, and the results will be shown for different time-steps.

Each figure has three subplots, the left plot is the result of CFD solver which is coming from MFiX directly, the middle plot is the result of smart proxy which is the output of ANN, and the right plot is the error distribution which is basically the difference between CFD and smart proxy.

4.2. Early time-step, non-cascading, single output, explicit

The simplest case to consider is when one time step from the early time is selected as the input. The ANN had only one output, so 9 separate ANN have been trained for all 9 parameters. The approach was non-cascading and explicit.

Time-steps 100 and 101 were used to train the system, and after the training completed, different time-steps were input to the trained ANN to get the results. All the time-steps from 101 all the way to 120 were input to the ANN and acceptable results were obtained. In the next sections, the results of gas volume fraction and gas pressure are provided for one time-step. The results for the rest of the parameters are in Appendix III.
4.2.1. Gas volume fraction

The results of the smart proxy versus CFD for gas volume fraction are shown in the next figures. As it is shown in Figure 4-2, Smart proxy is able to replicate the MFIX simulation results that show gas volume fraction distribution at time-step 102 for layer one. The maximum prediction error occurred occasionally at some points with the value less than 5% while we see an error near to zero in the rest of the domain. Very similar results have been obtained in other layers (from Figure 4-3 to Figure 4-6).
Figure 4-2: Comparison of CFD and smart proxy results for gas volume fraction of time-step 102 for layer one.

Figure 4-3: Comparison of CFD and smart proxy results for gas volume fraction of time-step 102 for layer two.
Figure 4.4: Comparison of CFD and smart proxy results for gas volume fraction of time-step 102 for layer three

Figure 4.5: Comparison of CFD and smart proxy results for gas volume fraction of time-step 102 for layer four
4.2.2. Gas Pressure

The results of the smart proxy versus CFD for the gas pressure are shown in the next figures. As it is shown in Figure 4-7, Smart proxy is able to replicate the CFD MFIX simulation results that show gas pressure distribution at time-step 102 for layer one. The maximum prediction error occurred occasionally at some points with the value less than 20% while we see an error near to zero in the rest of the domain. Very similar results have been obtained in other layers (from Figure 4-8 to Figure 4-11).
Figure 4-7: Comparison of CFD and smart proxy results for gas pressure of time-step 102 for layer one

Figure 4-8: Comparison of CFD and smart proxy results for gas pressure of time-step 102 for layer two
Figure 4.9: Comparison of CFD and smart proxy results for gas pressure of time-step 102 for layer three

Figure 4.10: Comparison of CFD and smart proxy results for gas pressure of time-step 102 for layer four
4.3. Late time-step, non-cascading, single output, explicit

The first attempt which was using the early time-step for training was successful, now the second attempt is to figure out if the ANN is capable of handling the bubbles in the fluidized bed or not. So, time-steps 4000 and 4002 were picked to train the network. The reason of having time-step 4002 rather than time-step 4001 is that there is no significant movement in the system only in one time-step and the neural network will see more movements by using the pair of 4000-4002 and it can learn better. More variation in data set helps the network to be trained better and being more powerful in prediction.

Similar to the previous scenario, the ANN had only one output, so 9 different ANN trained for all 9 parameters. The approach was non-cascading and explicit. All the time-steps from 4002 all the way to 4040 were input to the ANN and acceptable results were obtained. In the next sections, the results of the gas volume fraction are provided for one time-step.

4.3.1. Gas Volume Fraction

The results of the smart proxy versus CFD for gas volume fraction when the bed is fluidized are shown in the next figures. As it is shown in Figure 4-12 all the way to Figure 4-16, the smart proxy
could be able to mimic the CFD simulation results even when the bubbles are in the system. The figures show the gas volume fraction distribution at time-step 4004 for layer one through five. The maximum prediction error occurred occasionally at some points with the value around 4% while we see an error near to zero in the rest of the domain. Very similar results have been obtained in other layers (from Figure 4-3 to Figure 4-6).

*Figure 4-12- Comparison of CFD and smart proxy results for gas volume fraction of time-step 4004 for layer one*
Figure 4-13: Comparison of CFD and smart proxy results for gas volume fraction of time-step 4004 for layer two.

Figure 4-14: Comparison of CFD and smart proxy results for gas volume fraction of time-step 4004 for layer three.
Figure 4-15: Comparison of CFD and smart proxy results for gas volume fraction of time-step 4004 for layer four

Figure 4-16: Comparison of CFD and smart proxy results for gas volume fraction of time-step 4004 for layer five
4.4. Cascading, single output, explicit

Previous scenarios proved that the ANN is able to mimic the CFD results both when there is moderate change in the dynamics of the multi-phase flow and when the fluid dynamics is chaotic. All the analysis has been shown so far indicated the non-cascading scenario. The next attempt is to apply the cascading approach, with explicit scheme.

4.4.1. Gas volume fraction for early time

To accomplish this scenario, 9 different ANN have been trained by introducing time-step 100 as input and time-step 101 as output. Then for the deployment process, time-step 100 was used as the input of all 9 trained ANN. The outputs of those ANNs were used again to input the next time-step. In the following pages, the results of the cascading approach for gas fraction are shown. All the results are for layer two but different time-steps in order to see the solid motion.

By looking at the figures, error propagation could be seen from each time-step to the next one. This means that the ANN is able to replicate the results of CFD simulation using cascading scheme for only a few time-steps and after a couple of time-steps, the amount of error exceeds the acceptable range. In order to overcome the error propagation, more time-steps should be used for training. This has been practiced in the later scenarios.
Figure 4-17 - Comparison of CFD and smart proxy results for gas volume fraction of time-step 101 for layer two (Cascading)

Figure 4-18 - Comparison of CFD and smart proxy results for gas volume fraction of time-step 102 for layer two (Cascading)
Figure 4-19: Comparison of CFD and smart proxy results for gas volume fraction of time-step 103 for layer two (Cascading)

Figure 4-20: Comparison of CFD and smart proxy results for gas volume fraction of time-step 104 for layer two (Cascading)
Figure 4-21: Comparison of CFD and smart proxy results for gas volume fraction of time-step 105 for layer two (Cascading)

Figure 4-22: Comparison of CFD and smart proxy results for gas volume fraction of time-step 106 for layer two (Cascading)
Figure 4-23: Comparison of CFD and smart proxy results for gas volume fraction of time-step 107 for layer two (Cascading)

Figure 4-24: Comparison of CFD and smart proxy results for gas volume fraction of time-step 108 for layer two (Cascading)
Figure 4-25 - Comparison of CFD and smart proxy results for gas volume fraction of time-step 109 for layer two (Cascading)

Figure 4-26 - Comparison of CFD and smart proxy results for gas volume fraction of time-step 110 for layer two (Cascading)
4.4.2. Gas volume fraction for late time

The same type of analysis as discussed in the previous section has been performed here using time-steps 4000 and 4002 as input and output respectively. Only some of the time-steps were shown here to show the error propagation from each time-step to the next time-steps. These results show that in order to perform the prediction with the cascading approach that is the final goal of this research, further investigation is required. In the next sections, more discussion will be provided in order to solve this issue.

Figure 4-27: Comparison of CFD and smart proxy results for gas volume fraction of time-step 4002 for layer two (Cascading)
Figure 4-28- Comparison of CFD and smart proxy results for gas volume fraction of time-step 4004 for layer two (Cascading)

Figure 4-29- Comparison of CFD and smart proxy results for gas volume fraction of time-step 4006 for layer two (Cascading)
4.5. Early time-step, non-cascading, multiple output, explicit

Based on the discussion in chapter 3, it is sometimes beneficial to have multiple outputs rather than only one output. Three components of gas velocity were selected to be the output of the ANN; the input data is exactly the same as previous scenarios. The inputs came from time-step 100 and the outputs came from time-step 101. The ANN was trained successfully and the time-step 102 was predicted. Figure 4-31 to Figure 4-33 show the results of smart proxy and comparison with CFD simulation results. Fairly good results were obtained. Only the lower section of the fluidized bed is shown in the figures. Gas velocity in the y-direction has the highest error, which is less than 20% at some points, but the rest of the domain has error near zero. The maximum error for the gas velocity in the x-direction and Gas velocity in the z-direction is smaller. In the next section, the implicit results will be presented.
Figure 4.31 - Comparison of CFD and smart proxy results for gas $x$-velocity of time-step 102 for layer four (explicit)

Figure 4.32 - Comparison of CFD and smart proxy results for gas $y$-velocity of time-step 102 for layer one (explicit)
4.6. Early time-step, non-cascading, multiple output, implicit

The final goal of this project is to perform all the simulations using cascading approach and implicitly. In this section, the effect of an implicit solution is examined. All the parameters except solid velocity are coming from time-step 100, three components of solid velocity are coming from time-step 101, and this combination makes the input of the ANN. Also, three components of gas velocity from time-step 101 make the output. Figure 4-34 to Figure 4-36 show the results of this scenario.

To investigate the effect of implicit solution, the results are compared to the results from section 4.5.
By looking at the maximum error, which is the right plot in Figure 4-34, it is concluded that the amount of error was reduced from 12% to less than 9% (comparing Figure 4-31 and Figure 4-34). Also, Figure 4-35 shows the error reduction from 19% to 12% when it is compared to Figure 4-32, and Figure 4-36 shows the error reduction from 5% to 3% (in comparison to Figure 4-33).
Figure 4.35 - Comparison of CFD and smart proxy results for gas \( y \)-velocity of time-step 102 for layer one (implicit)

Figure 4.36 - Comparison of CFD and smart proxy results for gas \( z \)-velocity of time-step 102 for layer two (implicit)
4.7. Using multiple time-steps for training, non-cascading, single output, explicit

As discussed in the previous chapter, the ANN for early time is valid only for early times and probably a short period of time before and after the training time-step. This is also true for the ANN for late time. To overcome that problem, both time-steps should be used for training one ANN. To generalize the ANN even for times between early time and late time, third time-step was also added to train the ANN (Time-steps 200-202, 1000-1002, and 4000-4002 were used). The network has been trained and the deployment process was done by inputting time-step 200 all the way to time-step 4200. The results are presented in terms of RMSE of gas volume fraction.

Figure 4-37 shows the RMSE distribution versus time-steps. It is clear that in the time-steps that we had training data, the amount of error is minimum but in the other time-steps the RMSE increased a little. Also, there are some peaks in the figure and the peaks observed in figure 4-37 illustrate that further learning at additional time steps are required. Probably, there is a kind of new motion in the system in those time-steps that caused ANN did not learn enough to mimic the behavior of those time-steps.

![RMSE when 3 time steps were used](image)

*Figure 4-37- RMSE distribution versus time-step when three time pair of data were used for training*
Figure 4-38 shows time-step 500, which is one of the time-steps with a high value of error. By looking at this time-step closely, it could be understood that there is a specific kind of motion that neural network has not seen so far. Particles are falling down in this time-step and ANN did not learn about this kind of motion.

It is decided to include one more time-step for the training which was time-step 500, and that time-step was chosen based on Figure 4-37. The same scenario was followed but with 4 different time-steps for the training. Figure 4-39 shows the improvement in simulation by adding the fourth time-step for training. The blue curve shows the RMSE before adding the fourth time-step, and the red curve shows the RMSE after adding the fourth time-step.

*Figure 4-38- Comparison of CFD and smart proxy results for gas volume fraction of time-step 500 for layer two*
4.8. Using four time-steps for training, cascading, single output, explicit

The same simulation using 4 time-steps for the training was done with the cascading approach. Unfortunately, the error propagation is still seen in the results after a few time-steps. One of the solutions for this problem is to include more time-steps for the training but because of the memory issue, it is not possible to have more than 4 time-steps unless the size of the system reduces. The next effort is reducing the size of data in order to be able to add some more time-steps for the training.

4.9. Reducing the number of parameters (KPI)

Eliminating the upper part of the fluidized bed did reduce the size of the input data. In order to reduce the data input size even more, some of the parameters should also be eliminated from the training. The ANN has been trained with 69 and 70 parameters up to this point. There is a chance
for the ANN to be able to predict the behavior of the system by fewer parameters. In this way, the size of input can decrease.

To figure out what parameters has the top priority and what parameters has not, the tornado chart of the total weights of the parameters were plotted for each ANN. Since in back propagation method, there is a kind of weighted summation between all the parameters from each layer to the next layer, the total weight could be obtained by averaging all the weights corresponding to a specific parameter. There are two different ways to find the total weights; averaging all the weights by considering their signs, and averaging all the weights by removing the signs. Both approaches were accomplished. Tornado chart for both approaches is provided. Figure 4-40 shows tornado chart when the weights are averaged regularly, and as we expected before, the gas volume fraction of the cell has top priority. Also, Figure 4-41 shows the tornado chart when the weights are averaged after removing the sign.

![Tornado Chart of Neural Network Weights](image)

*Figure 4-40* - parameter prioritization for Gas volume fraction ANN (averaging all the weights)
The 14 less important parameters were removed from the input and the ANN has been trained again with the new 56 parameters based on the two different approaches. Figure 4-42 shows the amount of error when a different number of parameters were used for the training. The blue curve is when all the 70 parameters were used, red curve shows the error when 14 parameters were removed from the training by simply calculating the average and the yellow curve is when 14 parameters were eliminated based on the averaging of the absolute value of the weights. This graph shows that the prioritization of the parameters is more accurate when the weights are averaged without considering the sign. This conclusion is reasonable because some positive and negative weights might cancel each other when the average is calculated by considering the signs, the wrong result may be obtained.

The removed parameters for both approaches are shown in the Table 4-1 and Table 4-2. The parameters are sorted in two different fashions; by parameter or by location.
As discussed, the second method was chosen to perform the KPI analysis for the rest of the parameters.

Another interesting fact about Figure 4-42 is that the error distribution of yellow curve (using 56 parameters) is even less than the blue curve (which was obtained by using 70 parameters). To
explain the reason for this phenomenon, it should be stated that the blue curve is from non-optimized database (Spatio-Temporal database) and the yellow curve is from a reduced database. Reducing the number of parameters does not end up having higher prediction error always, it could help the network to find the relation between the parameters more easily. In other words, there might be some irrelevant parameters in the Spatio-Temporal database which without them the process of fitting could be accomplished by higher accuracy. So, in the optimization process, the less important parameters are eliminated until the prediction error goes far from the original error.

![Comparison between RMSE](image.png)

*Figure 4-42: Comparison of RMSE distribution versus time-step for two different approach of averaging*

The number of parameters was reduced to 56 and still the amount of error is lower than the original simulation. This motivates us to go further and reduce the number of parameters even more until the system breaks. For the second attempt, the number of parameters was reduced to 42 and Figure 4-43 shows the error distribution after removing some other parameters. In this figure, the amount of error increased a little bit in comparison to the when 56 parameters were used but it is still comparable to the original one.
For the third attempt, another seven parameters were removed based on the KPI analysis and the below curve was obtained. Figure 4-44 shows the error distribution when only 35 parameters were used to train the system and it means that almost the same results were achieved by using only half of the data.
The same analysis was done for all the parameters and the size of data for all of them was reduced. Although it was possible to consider a different number of parameter for different ANN, the number of parameters was fixed to be 43 for all the ANN for consistency. But it should be mentioned that these 43 parameters vary from one ANN to another, for example when the gas volume fraction is going to be trained, different parameters are important in comparison to when the gas pressure is training. In the previous section, it was mentioned that in order to be able to perform the cascading deployment, more time-steps should be used in the training process. Because of the memory limitations, it was not possible to add more time-steps in the training but now, when the size of input has been decreased; more time-steps could be used in the training, which is the discussion of the next section.
Table 4-3 shows the size of data before and after size reduction, the data size will be reduced 5 times after using the latest model.

<table>
<thead>
<tr>
<th>Model</th>
<th>Size of input</th>
<th>Total Data Point</th>
</tr>
</thead>
<tbody>
<tr>
<td>Original Model</td>
<td>118,098 by 70</td>
<td>8,148,762</td>
</tr>
<tr>
<td>Latest Model</td>
<td>51,030 by 35</td>
<td>1,786,050</td>
</tr>
</tbody>
</table>

4.10. Using seven time-steps for training, cascading, single output, explicit

The goal of this section is to train a smart ANN that can replicate the CFD completely, it means that the smart package will feed itself instead of feeding by CFD, which is the definition of a cascading deployment. The only input from CFD is one time-step as the initial condition. Fairly good results were obtained with the non-cascading approach by using 4 time-steps. In this section, another three time-steps are going to be added to the training set in order to improve the results. The question is what time-steps should be included in the training set. In order to answer this question, the error distribution of non-cascading result is plotted versus time as it is shown in Figure 4-45. Another three time-steps were selected based on the peaks on the error curve. Time-steps 574, 904, and 1842 were selected.

![Figure 4-45: Comparison between RMSE with and without P₁](image)
All the parameters were trained based on the reduced input size by including 7 time-steps for the input data. The training process was done successfully and the cascading deployment was performed by inputting time-step 4000 to the ANN. The results of the cascading are shown in Figure 4-46 for time-step 4020. This figure shows that after 20 time-steps, the error propagates and reaches to an unacceptable value. This figure shows that the final goal of this project is not achieved yet and more investigations are required.

![Gas Fraction of Layer=1 at time step=4020](image)

*Figure 4-46: Comparison of CFD and smart proxy results for gas volume fraction of time-step 4020 for layer one (Cascading)*

4.11. Changing the data prioritization

As the last method to reduce the input size, the percentage of data for training will be reduced in order to include more time-steps in the training set. The initial percentage for the training was 70%. Three different simulations were conducted with different data prioritization and acceptable results were obtained.
Table 4-4: Data Partitioning in different scenarios

<table>
<thead>
<tr>
<th>Data</th>
<th>Training (%)</th>
<th>Validation (%)</th>
<th>Test (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Original simulation</td>
<td>70</td>
<td>15</td>
<td>15</td>
</tr>
<tr>
<td>First attempt</td>
<td>60</td>
<td>20</td>
<td>20</td>
</tr>
<tr>
<td>Second attempt</td>
<td>40</td>
<td>30</td>
<td>30</td>
</tr>
<tr>
<td>Third attempt</td>
<td>30</td>
<td>35</td>
<td>35</td>
</tr>
</tbody>
</table>

The results of the three attempts are shown in the following figures and all of them show that ANN still is able to learn the pattern by reducing the percentage of training data down to the 30%.

Figure 4-47: CFD and smart proxy results for gas volume fraction of time-step 4004 for layer one by 60% training (non-cascading)
Figure 4-48: CFD and smart proxy results for gas volume fraction of time-step 4004 for layer one by 40% training (non-cascading)

Figure 4-49: CFD and smart proxy results for gas volume fraction of time-step 4004 for layer one by 30% training (non-cascading)
4.12. Smart sampling

The data from time step 4000 was sent in to the Smart Sampling procedure. The deployment started from time step 3950 all the way to time step 4150. The same analysis was done without smart sampling and all the data were used in the training.

Figure 4-50 shows RMSE of smart sampling in comparison to the original model. As it is shown in the figure, by reducing the number of records (only 20% of the records were used) almost the same result obtained.

![Figure 4-50: Comparison of RMSE in different time steps with/without smart sampling](image)

This idea let us include more time steps in the training process in order to reduce the error in the cascading deployment.
Chapter 5 Conclusions and Recommendations

5.1. Conclusions

The original database (Spatio-Temporal database) included 70 parameters and 118,098 records. This database was then sent through an optimization process to get an optimized database with the smaller size but the same efficiency. In the optimization process, the size of Spatio-Temporal database reduced more than 25 times. This optimization was done by different approaches as follows and the exact numbers are shown in the Table 5-1.

1. Reducing the number of parameters (using Key Performance Indicator)
2. Reducing the number of records (focusing on the more important cells using smart sampling)
3. Reducing the percentage of training (by Intelligent Partitioning).

| Table 5-1: comparison between Spatio-Temporal database and optimized database |
|-----------------|------------------|------------------|------------------|
|                  | No. Records      | No. of Parameters | Training Percentage | Total data       |
| Spatio-Temporal Database | 118,098          | 70               | 100               | 8,266,860        |
| Optimized Database     | 25,827           | 43               | 30                | 333,168          |

All the data from time step t were used as the input of the model, and all the data from time step (t+1) were used as the output (since the method is supervised learning, the output is required). A model has been trained by using the original database.

In order to see the efficiency of the model and successfulness of this method, some blind cases (which the model has not seen in the training process) were used for deployment. The results proved that the smart proxy is able to predict all the entire range, including the initiation of the bubbles, bursting of them, bulk flow, and also the non-linear and chaotic motion.

A data-driven smart proxy was developed to mimic the CFD results, with a good accuracy and faster speed. Table 5-2 shows the comparison of run time of these two approaches. This proxy needs an incredible less amount of time in comparison to the CFD solver with a reasonable error (less than 10%). This project proved that the idea of using AI&DM for computational fluid dynamics actually works.
Table 5-2: comparison between speed of run for CFD and Smart proxy

<table>
<thead>
<tr>
<th>Method</th>
<th>Execution Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>CFD</td>
<td>4 seconds simulation = 3 days on 4 CPUs</td>
</tr>
<tr>
<td>Smart Proxy</td>
<td>4 seconds simulation = 180 s = 3 min</td>
</tr>
</tbody>
</table>

5.2. Recommendations and future works

This study showed that the smart proxy is feasible to handle a complex, multi-physics, nonlinear flow and it is worth to spend more time and effort to improve the results.

In order to make the model more general in the whole time ranges, more data from different time steps could be included in the training process, and this is where using the optimized database is beneficial.

Also, there is a need to spend more time on the cascading deployment in order to completely replace the CFD with the smart proxy.
BIBLIOGRAPHY


APPENDIX

Appendix I: MFiX Equations

Fluid-Solids Momentum Transfer

The fluid-solids interaction force is a combination of buoyancy, the drag force, and momentum transfer due to mass transfer.

\[
\vec{I}_{gm} = -\varepsilon_{sm} \nabla P_g - F_{gm} \left( \bar{v}_{sm} - \bar{v}_g \right) + R_{0m} \left[ \varepsilon_{0m} \bar{v}_{sm} + \bar{\varepsilon}_{0m} \bar{v}_g \right]
\]  

(0-1)

Where

\[
\bar{\varepsilon}_{0m} = \begin{cases} 
1 & \text{for } R_{0m} < 0 \\
0 & \text{for } R_{0m} \geq 0 
\end{cases}
\]

\[
\varepsilon_{0m} = 1 - \varepsilon_{0m}
\]

(0-2)

And

\[
F_{gm} = \frac{3}{4} \varepsilon_{sm} \varepsilon_{g} \frac{\rho_g}{V_{rm} d_{pm}} C_{Ds} \left( \frac{Re_m}{V_{rm}} \right) \left| \bar{v}_{sm} - \bar{v}_g \right|
\]

(0-3)

where \( V_{rm} \) is the terminal velocity correlation for the \( m^{th} \) solids phase which is a function of gas volume fraction only, and the \( Re_m \) is Reynolds number of \( m^{th} \) phase.

\[
Re_m = \frac{d_{pm} \left| \bar{v}_{sm} - \bar{v}_g \right| \rho_g}{\mu_g}
\]

(0-4)

Or

\[
F_{gm} = \frac{3\varepsilon_{sm} \varepsilon_{g} \rho_g}{4V_{rm} d_{pm}} \left( 0.63 + 4.8 \sqrt{Re_m} \right) \left| \bar{v}_{sm} - \bar{v}_g \right|
\]

(0-5)
\[ V_{em} = 0.5 \left[ A - 0.06 Re_m + \sqrt{(0.06 Re_m)^2 + 0.12 Re_m (2B - A) + A^2} \right] \] (0-6)

The coefficient A and B are calculated as follows.

\[ A = \varepsilon_s^{4.14} \] (0-7)

\[ B = \begin{cases} 
0.8 \varepsilon_s^{1.28} & \text{if } \varepsilon_s \leq 0.85 \\
\varepsilon_s^{2.65} & \text{if } \varepsilon_s > 0.85 
\end{cases} \] (0-7)

**Solids-Solids Momentum Transfer**

The solids-solids momentum transfer comes from the drag force between different phases and is calculated with the below equation.

\[ \mathbf{\ddot{I}_{ml}} = -F_{sml} \left( \mathbf{\ddot{v}_{sl}} - \mathbf{\ddot{v}_{sm}} \right) + R_{ml} \left[ \varepsilon_{ml} \mathbf{\ddot{v}_{sl}} + \varepsilon_{ml} \mathbf{\ddot{v}_{sm}} \right] \] (0-8)

Where

\[ F_{sml} = \frac{3 \left( 1 + \varepsilon_{lm} \right) \left( \pi / 2 + C_{film} \pi^2 / 8 \right) \varepsilon_{sl} \rho_{sl} \rho_{sm} \rho_{sm} \left( d_{pl} + d_{pm} \right)^2 g_{0lm} | \mathbf{\ddot{v}_{sl}} - \mathbf{\ddot{v}_{sm}} |}{2 \pi \left( \rho_{sl} d_{pl} + \rho_{sm} d_{pm} \right)} \] (0-9)

where \( \varepsilon_{lm} \) and \( C_{film} \) are the coefficient of restitution and coefficient of friction, respectively, between the \( l^{th} \) and \( m^{th} \) solids-phase particles. \( g_{0lm} \) is the radial distribution function at contact.

\[ g_{0pm}^{th} = \frac{1}{\varepsilon_s} \frac{3 \left( \sum_{l=1}^{M} \varepsilon_{sl}/d_{p2} \right) d_{pl} d_{pm}}{\varepsilon_s^2 (d_{pl} + d_{pm})} \] (0-10)
**Fluid-Phase Stress Tensor**

The fluid-phase stress is stated as follows.

\[ S_g = -p_g \bar{I} + \tau_g \quad (0-11) \]

Where \( P_g \) is the gas pressure and also \( \tau_g \) is the viscous stress tensor for Newtonian fluid which is in the below form.

\[ \tau_g = 2 \, \varepsilon_g \, \mu_g \, \bar{D}_g + \varepsilon_g \, \lambda_g \, tr \left( \bar{D}_g \right) \bar{I} \quad (0-12) \]

Where \( \bar{I} \) is identity tensor and \( \bar{D}_g \) is strain tensor which is calculated by the following equation.

\[ \bar{D}_g = \frac{1}{2} \left[ \nabla \bar{v}_g + \left( \nabla \bar{v}_g \right)^T \right] \quad (0-13) \]

**Solids-Phase Stress Tensor**

In MFIX, there are two different theories for calculating the solid-phase stress and those two theories are plastic flow and viscous flow. The stress definition is switched based on the comparison of the gas volume fraction with a critical packing which is \( \varepsilon_g^* \).

\[ S_{sm} = \begin{cases} -p_{sm} \bar{I} + \tau_{sm}^p \text{ if } \varepsilon_g \leq \varepsilon_g^* \\ -p_{sm} \bar{I} + \tau_{sm}^v \text{ if } \varepsilon_g > \varepsilon_g^* \end{cases} \quad (0-14) \]

where \( P_{sm} \) is the pressure and \( \tau_{sm} \) is the viscous stress in the \( m \)th solids phase. The superscript \( p \) stands for the plastic regime and \( v \) for the viscous regime. In fluidized-bed simulations, \( \varepsilon_g^* \) is usually set to the void fraction at minimum fluidization.

In the plastic flow, \( \tau_{sm}^p \) is calculated as follows.

\[ \tau_{s1}^p = 2 \, \mu_{s1}^p \, \bar{D}_{s1} \quad (0-15) \]

In viscous flow, \( \tau_{sm}^v \) is in the below form.

\[ \tau_{sm}^v = 2 \, \mu_{sm}^v \, \bar{D}_{sm} + \lambda_{sm}^v \, tr( \bar{D}_{sm} ) \, \bar{I} \quad (0-16) \]
Appendix II: MFiX Gasification code

# 3-D Rectangular Fluidized Bed
# ---------------------------------------------------------
# Run control
# ---------------------------------------------------------
RUN_NAME = 'FB_S2G1'
DESCRIPTION = 'Bubbling Fluidized Bed Simulation'
RUN_TYPE = 'new'
UNITS = 'si'
TIME  = 0.0
TSTOP = 30.0 'change as needed
DT    = 5.0E-3
DT_MAX = 5.0E-3 !Don't go larger that write_usr output freq
NODESI = 4
NODESJ = 16
NODESK = 4
# ---------------------------------------------------------
# Equations
# ---------------------------------------------------------
ENERGY_EQ = .FALSE.
SPECIES_EQ = .FALSE. .FALSE.
DRAG_TYPE = 'GIDASPOW'
SCHAEFFER = .FALSE.
FRICTION = .FALSE.
# ---------------------------------------------------------
# Geometry Section
# ---------------------------------------------------------
COORDINATES = 'cartesian'
XLENGTH = 0.120 IMAX = 27
YLENGTH = 0.720 JMAX = 162
ZLENGTH = 0.120 KMAX = 27
# ---------------------------------------------------------
# Gas-phase Section
# ---------------------------------------------------------
MU_g0   = 1.9E-5
MW_avg  = 29.0
# Solids-phase Section

RO_s0 = 2000.0 ! kg/m3
D_p0 = 0400.0E-6 ! m
C_c = 0.80
Phi = 30.0
EP_star = 0.42

# Initial Conditions Section

! Bed Freeboard
IC_X_w(1:2) = 0.000 0.000
IC_X_e(1:2) = 0.120 0.120
IC_Y_s(1:2) = 0.000 0.120
IC_Y_n(1:2) = 0.120 0.720
IC_Z_b(1:2) = 0.000 0.000
IC_Z_t(1:2) = 0.120 0.120
IC_EP_g(1:2) = 0.420 1.000
IC_P_g(1:2) = 101.325E3 101.325E3
IC_T_g(1:2) = 300.0 300.0
IC_U_g(1:2) = 0.000 0.000
IC_V_g(1:2) = @(0.6/0.42) 0.600
IC_W_g(1:2) = 0.000 0.000
IC_U_s(1:2,1) = 0.000 0.000
IC_V_s(1:2,1) = 0.000 0.000
IC_W_s(1:2,1) = 0.000 0.000
IC_T_s(1:2,1) = 300.0 300.0

# Boundary Conditions Section

! South North West East Back Front
BC_X_w(1:6) = 0.000 0.000 0.000 0.120 0.000 0.000
BC_X_e(1:6) = 0.120 0.120 0.000 0.120 0.120 0.120
BC_Y_s(1:6) = 0.000 0.720 0.000 0.000 0.000 0.000
BC_Y_n(1:6) = 0.000 0.720 0.720 0.720 0.720 0.720
BC_Z_b(1:6) = 0.000 0.000 0.000 0.000 0.000 0.120
BC_Z_t(1:6) = 0.120 0.120 0.120 0.120 0.120 0.120
BC_TYPE(1:6) = 'MI' 'PO' 'NSW' 'NSW' 'NSW' 'NSW'
BC_EP_g(1:2) = 1.0 1.0
BC_P_g(1:2) = 2*101.325E3
BC_T_g(1:2) = 2*300.0
BC_U_g(1) = 0.0
BC_V_g(1) = 0.6
BC_W_g(1) = 0.0
! BC_ROP_s(1,1) = 0.000
! BC_U_s(1,1) = 0.000
! BC_V_s(1,1) = 0.000
! BC_W_s(1,1) = 0.000
#
# Output Control
#
RES_DT = 0.01
!
! EP_g P_g U_g U_s ROP_s T_g X_g
! P_star V_g V_s T_s1 X_s Theta Scalar
! W_g W_s T_s2
SPX_DT = 0.01 0.01 0.01 0.01 0.01 100. 100. 100.0 100.0
NLOG = 100
full_log = .true.
write_dashboard = .true.
write_vtk_files = .true.
time_dependent_filename = .true.
vtk_dt = 0.05
vtk_varlist(1:1:5) = 1 2 3 4 5
call_usr = .true.
usr_dt(1) = 0.005
ENABLE_DMP_LOG = .F
CHK_BATCHQ_END = .T.
BATCH_WALLCLOCK = 1800d20 ! 30 Days
TERM_BUFFER = 120d0 ! 15 min
Appendix III: Early time-step, non-cascading, single output, explicit

In the below graphs, the results of non-cascading deployment are shown for early time steps for different parameters.

*Figure 0-1* - Comparison of CFD and smart proxy results for gas x-velocity of time-step 102 for layer one (explicit)
Figure 0-2: Comparison of CFD and smart proxy results for gas y-velocity of time-step 102 for layer one (explicit)

Figure 0-3: Comparison of CFD and smart proxy results for solid x-velocity of time-step 102 for layer one (explicit)
Figure 0-4: Comparison of CFD and smart proxy results for solid y-velocity of time-step 102 for layer one (explicit)

Figure 0-5: Comparison of CFD and smart proxy results for gas z-velocity of time-step 102 for layer one (explicit)
Appendix IV: MATLAB Code (Creating Spatio-Temporal database)

%% This program read two different time step as input and output
clc
clear;
close all;
%% Input and Output Time step

Time_Step_in=100;
Time_Step_out=Time_Step_in+1;

%% Defining the geometry
imax = 27; %number of cell in x-direction
kmax = 27; %number of cell in z-direction
jmax = 162; %number of cell in y-direction
Dim_x = 0.12; %length of domain
Dim_y = 0.72; %height of domain
Dim_z = 0.12; %width of domain
Dim_cell = Dim_x/imax; %dimension of cell, the cell is a cube
CellNo=imax*jmax*kmax; %total number of cells

%% Loading the CSV files from time step start to final

data_input = ReadFiles( Time_Step_in,Time_Step_in,CellNo );
data_output = ReadFiles( Time_Step_out,Time_Step_out,CellNo );

%% Initialize the matrices
location = zeros(CellNo,3); % (x,y,z) of each cell
CellID = zeros(CellNo,1); % ID of each cell
tier1 = zeros(CellNo,6); % tier system with of order 1 (bottom,top,west,north,east,south)
DisToBoundary = zeros(CellNo,6); % distance to the boundaries (bottom,top,west,north,east,south)
counter = 1;

%% Filling the geomtry matrices
for k = 1:kmax
    for i = 1:imax
        for j = 1:jmax
            location(counter,:) = [i j k];
            CellID(counter) = (k-1)*jmax*imax+(i-1)*jmax+j;
            %cell ID of tier cells (bottom,top,west,north,east,south)
            tier1(counter,:) = [(k-1)*jmax*imax+(i-1)*jmax+j-1 (k-1)*jmax*imax+(i-1)*jmax+j+1 (k-1)*jmax+1 (k-1)*jmax*imax+(i-2)*jmax+j (k)*jmax+1 (k-1)*jmax*imax+(i-1)*jmax+j];
            %distance of center of each cell to the boundaries (bottom,top,west,north,east,south)
            DisToBoundary(counter,:) = [(j-0.5)*Dim_cell Dim_y-(j-0.5)*Dim_cell (i-0.5)*Dim_cell Dim_z-(k-0.5)*Dim_cell Dim_x-(i-0.5)*Dim_cell (k-0.5)*Dim_cell ];
            %filling zero at for the tiers for the cells at the boundary
            if i==1
            end
        end
    end
end
tier1(counter,3) = 0;
elseif i==imax
    tier1(counter,5) = 0;
end

if j==1
    tier1(counter,1) = 0;
elseif j==jmax
    tier1(counter,2) = 0;
end

if k==1
    tier1(counter,6) = 0;
elseif k==kmax
    tier1(counter,4) = 0;
end

counter=counter+1;
end
end
end

%% Filling the properties of tier1 system

tierData=zeros(CellNo,7*11);
for i = 1:CellNo
    tierData(i,1:11)=data_input(i,:);
    for j = 1:6
        index = tier1(i,j);
        if index==0 %if index iz zero, it means that no cell is there
            t = zeros(1,11);
        else
            t = data_input(index,:);
        end
        tierData(i,11*(j)+1:11*(j+1)) = t;
    end
end

%% Creating the output file

geometry=[CellID location DisToBoundary tier1 tierData data_output(:,:)];

NN_input=[DisToBoundary tierData(:,1:9) tierData(:,12:20) tierData(:,23:31) tierData(:,34:42) tierData(:,45:53) tierData(:,56:64) tierData(:,67:75)];

Fraction=data_output(:,1);
Gas_Pressure=data_output(:,2);
Solid_Pressure=data_output(:,3);

Gas_Velocity=data_output(:,[4 5 6]);
Gas_Velocity_x=data_output(:,4);
Gas_Velocity_y=data_output(:,5);
Gas_Velocity_z=data_output(:,6);

Solid_Velocity=data_output(:,[7 8 9]);
Solid_Velocity_x=data_output(:,7);
Solid_Velocity_y=data_output(:,8);
Solid_Velocity_z=data_output(:,9);

NN_input=NN_input';
Fraction=Fraction';
Gas_Pressure=Gas_Pressure';
Solid_Pressure=Solid_Pressure';
Gas_Velocity=Gas_Velocity';
Gas_Velocity_x=Gas_Velocity_x';
Gas_Velocity_y=Gas_Velocity_y';
Gas_Velocity_z=Gas_Velocity_z';

Solid_Velocity=Solid_Velocity';
Solid_Velocity_x=Solid_Velocity_x';
Solid_Velocity_y=Solid_Velocity_y';
Solid_Velocity_z=Solid_Velocity_z';