Simulation of Proton Exchange Membrane Fuel Cell by using ANSYS Fluent

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Abstract. Proton exchange membrane fuel cells (PEMFC) are attractive alternative source of electricity. The current study involves the computational fluid dynamics simulations of PEMFC under isothermal and non-isothermal conditions to investigate the performance of fuel cell. Effect of pressure and temperature on fuel cell performance is studied under non-isothermal conditions. PEMFC is modeled at 323 K and 1 atm under isothermal conditions whereas under non-isothermal conditions, the simulation is run on 353 K and 3 atm. The results show that the current density increases with increase in operating pressure of PEMFC and vice-versa with operating temperature.

Keywords. Simulation, proton exchange membrane, fuel cell, ANSYS Fluent

1. Introduction
Fuel cell converts chemical energy into electrical energy. It runs continuously until the reactants run out. The ever growing efficiency and very low emissions of fuel cells will ensure their commercial success in coming years [1]. PEMFC’s operate at low temperatures (below 100 °C) which is easy to operate and handle. Recent research work on PEMFC is generally focused on its size, cost, performance and durability. Computer simulations are getting more attention of researchers as low cost option which produces reliable results.

PEMFC performance depends upon various factors like: Physical parameters (i.e. ref. current density, conc.), operating parameters (i.e. operating temperature, pressure), geometrical parameters, advanced parameters like contact resistance. Siegel C. et al reviewed PEM fuel cell simulations performed on different softwares [2]. Researchers [3] validated PEMFC model for investigating convergence criteria. According to authors [4], incomplete and unreliable data is the main issue for obtaining correct results. There is a lot of work going on flow fields to get better performance of PEMFC. Computational model developed for a 5 cm² fuel cell with serpentine and parallel flow fields are considered in experimentation and in modeling [5].

Simulation has been performed by modifying the design of flow field which decrease the pressure drop in PEMFC. Modifications are based on theory of injection engine to reduce pressure drop [6]. A comprehensive non isothermal, 3-D model is established to investigate the performance of PEMFC with straight and serpentine flow fields [7]. Aspect ratio of flow fields also has effect on fuel cell performance [8]. The local transport phenomena and cell performance is performed for parallel and integrated flow fields [9]. Effect of single (1-s), double (2-s) and triple (3-s) serpentine flow fields is also investigated. It is reported that effects of flow fields variations are negligible below 0.7 operating voltage [10]. A stepped flow field is proposed to improve cell performance [11]. Researchers made the comparison between the conventional straight gas flow channel and a novel wave-like channel [12]. Different shapes and lengths of parallel flow field affect cell performance [13]. Water management is one of the main issues in PEM fuel cell [14].

Percolation theory was coupled with 3-D PEMFC model to investigate the effects of water flooding on gas diffusion layer [15]. Using different height and length tapered flow fields due to which velocity increases and water flooding does not produce, it also enhances rib-convection among
adjacent channels [16]. Muthukumar M et.al analyzed the effects of landing to channel (LXC) on the generation of power and current density [17]. Researchers [18] focus on the relation between flow velocity and optimum channel width. PEM permits only proton ions and PEM’s activity depends upon the humidity on surface of membrane [19]. Researchers made the comparison between 3 phenomenological membrane models and the results indicates that springer model and Nguyen and white model over predict drying of the membrane while the fuller and new man model provides the best match with experimental data [20]. In present study the transport equations are used after [7].

**Table 1. PEMFC Dimensions [7].**

| Cell dimensions                  | Units | Values |
|----------------------------------|-------|--------|
| Gas channel length               | mm    | 10     |
| Height of gas channel            | mm    | 1      |
| Width of the gas channel         | mm    | 1      |
| Width of the cell                | mm    | 2      |
| Thickness of catalyst layer      | mm    | 0.014  |
| Thickness of gas diffusion layer | mm    | 0.0254 |
| Thickness of current collector   | mm    | 2.5    |
| Thickness of membrane            | mm    | 0.051  |
| Overall cell height              | mm    | 5.1295 |

**Figure 1.** 3-D view of single channel PEMFC

2. **Methodology**

The simulated 3-D PEMFC consists of single straight channel shown in figure 1. The dimensions are given in table [1]. Boundary conditions and physical parameters used in the simulation are shown in tables 2 and 3 respectively. The model is developed with the following assumptions:

- Flow is laminar.
- Incompressible fluids.
- Steady state system exists.
- Inlet gases follow Ideal gas law.
- The catalyst layer, membrane and gas diffusion layer are Isotropic materials.
• For isothermal modeling temperature is constant.
• For non-isothermal modeling heat flux is constant.

| BC types          | Location                              | Parameters                     | Non-Isothermal | Isothermal | Units   |
|-------------------|---------------------------------------|-------------------------------|----------------|------------|---------|
| **Velocity inlet**| Inlet anode flow channel face         | Velocity inlet                | 2              | 0.3        | m/s     |
|                   |                                       | Mass fraction of H₂          | 0.3            | 0.3        | -       |
|                   |                                       | Mass fraction of H₂O         | 0.7            | 0.7        | -       |
|                   | Inlet Cathode flow channel face       | Velocity inlet                | 2              | 0.5        | m/s     |
|                   |                                       | Mass fraction of O₂          | 0.14           | 0.212      | -       |
|                   |                                       | Mass fraction of H₂O         | 0.2            | 0.079      | -       |
| **Pressure Outlet**| Outlet anode flow channel face        | Anode outlet gas pressure    | 0              | 0          | Pa      |
|                   |                                       | Temperature                  | Default        | 323        | K       |
|                   | Outlet cathode flow channel face      | Cathode outlet gas pressure  | 0              | 0          | Pa      |
|                   |                                       | Temperature                  | Default        | 323        | K       |
| **Wall**          | The terminal and upper anode current collector face | Specified electric potential | 0              | 0          | Volts   |
|                   | (only terminal)                        |                               |                |            |         |
|                   | The terminal and lower cathode current collector face | Specified electric potential | 0.5-0.9       | 0.4-0.9    |         |
|                   | (only terminal)                        |                               |                |            |         |
|                   | All Outer cell faces                  | Thermal condition constant   | -              | 323        | K       |
|                   |                                       | Temperature                  |                |            |         |
|                   |                                       | Heat flux                    | 0              | -          |         |

**Table 2. Boundary conditions (BC).**

**Table 3. Parameters values.**

| Parameters                                      | Isothermal | Non-isothermal | Units    |
|------------------------------------------------|-------------|----------------|----------|
| Cell operating temperature                     | 323         | 353            | K        |
| Cell operating pressure                        | 1           | 3              | Atm      |
| Open-circuit voltage                           | 1.07        | 1.05           | Volt     |
| Anode Reference exchange current density       | 10,000      | 30             | A/m²     |
| Cathode Reference exchange current density     | 20          | 0.004          | A/m²     |
| CL Electric conductivity                       | Default     | 100            | 1/ohm-m  |
| Current collector Electric conductivity         | 4000        | Default        | 1/ohm-m  |
| GDL Electric conductivity                       | 300         | Default        | 1/ohm-m  |
| Anode exchange coefficient                      | 1           | 0.5            | -        |
| Cathode exchange coefficient                    | 1           | 2              | -        |
| Reference concentration of anode               | 1           | 0.04           | -        |
| Reference concentration of cathode             | 1           | 0.00086        | -        |
| CL Porosity                                    | 0.112       | 0.6            | -        |
| GDL Porosity                                   | 0.6         | 0.6            | -        |

ANSYS meshing is used to generate hexahedral mesh with 59200 cells whereas Fluent is used as a solver. Joule heating, electrochemistry and Butler-Volmer are used as model equations for
PEMFC simulation. SIMPLE algorithm is used for solving equations. The least square cell base is chosen under spatial discretization.

3. Results and Discussion
The simulation of single channel PEMFC is performed according to the condition given in tables 2 and 3. Results are presented in figures 2 and 3. The models are validated by using data of Lee Wang et.al. [21] and the results are in good agreement with the experimental data except for the low current density region under non-isothermal conditions and high current density region under isothermal conditions. As the redox reaction in the fuel cell increases, the current density also increases. At high current density, production of water increases in catalyst layer and gas diffusion layer (GDL), due to which effective porosity of catalyst layer and GDL decreases and material resistance increases. The assumption of the simulation that water is being produced in the form of mist does not remain valid due to this phenomenon at high current densities as the large quantities of liquid water fill the pores of catalyst and membrane, thereby increasing transport resistance [5].

Figure 2. Comparison of simulation and experimental results under isothermal conditions.

Figure 3. Comparison of simulation and experimental results under non-isothermal condition.
The most affecting reactant at higher current density is oxygen, as shown in figure 4 (a). The mass fraction of oxygen is very high in cathode channel at 353 K. Under the current collector, oxygen is more depleted in MEA region. This might be due to the limited oxygen diffusivity and also due to the difficulty of water removal in cathode side. As vapor saturation pressure depends on temperature, the relative humidity decreases with the increase in temperature. The membrane might be dehydrated thereby slowing the protonic ions permeation and leads to ohmic losses. Another reason of high mass fraction of oxygen might be slow protonic ions generation in membrane as compared to oxygen diffusion into GDL. On the other hand, it can be clearly seen that at high current density there is low change in hydrogen value as compared to oxygen as shown in figure 4 (b). Because hydrogen (small molecule) has better diffusivity than oxygen and it shows better transport at high current density even if the porosity is low. Usually there is no blockage at porous region of hydrogen side because flooding is not taking place on anode side. Pressure decreases from inlet to outlet as shown in pressure contours in figure 4(c). Temperature increases from inlet to outlet as shown in temperature contours in figure 4(d) due to reduction reaction and ohmic losses.

The current densities over-potential near the CCL/PEM interfaces are so big that the maximum current density generated in cathode region is produced on this interface. At this current density, starvation region is clearly shown in figure 5 under the current collector region; this might be due to oxygen diffusivity problem [22].

![Figure 4](image1.png)

**Figure 4.** Mass fraction distribution under non-isothermal conditions at 0.6 volts for (a) O₂ (b) H₂ (c) Static pressure (d) Temperature.

![Figure 5](image2.png)

**Figure 5.** Current Distribution volts non-isothermal.
Oxygen mass fraction decreases from inlet to outlet because oxygen is consumed in the reaction on its way to the outlet. Current density is greater at the hydrogen inlet as compared to oxygen inlet since the reaction starts at hydrogen inlet as shown in figure 6.

It is shown in figure 4 and figure 6 that no gases are present in MEA region of fuel cell. This means that at anode side, the hydrogen ions enter into the catalyst layer while on cathode side, oxygen ions enter into the catalyst layer instead of gas molecules [23].

**Figure 6.** Mass fraction distribution under isothermal conditions at 0.6 volts for (a) O₂ (b) H₂

Current density value is low at collectors rib because there is depletion of oxygen in those region which is shown in figure 7. Drying of anode side might be major cause of decrease in cell performance. Multi-phase model should be used to observe this phenomenon in detail [5].

**Figure 7.** Current distribution at 0.6 volts (isothermal).
Polarization curves of different pressures at 353 K temperature are shown in figure 8. The pressures of both anode and cathode sides were kept the same. The performance of the fuel cell improves with the increase of pressure. The overall polarization curves shift positively as the pressure increases. With the increase in pressure, partial pressure of the reactant gases increases [21].

Figure 8. Effects of Pressure on voltage and current density at 253 K.

Figure 9. Mass fraction of oxygen at (a) 1 atm (b) 2 atm (c) 3 atm.
The mass fraction of oxygen decreases in PEMFC as the operating pressure is increased as shown in figure 9. The decreased mass fraction is due to increase in reaction rate which is favored by increased operation pressure.

![Figure 10](image1.png)

**Figure 10.** Effect of Temperature on voltage and current density.

![Figure 11](image2.png)

**Figure 11.** Mass Fraction of O₂ at 323K (a), 333K (b) and 353K (c).
Polarization curve at different temperatures are shown in figure 10. Oxygen mass fraction is shown in figure 11. Oxygen mass fraction distribution is often considered the criteria of fuel cell performance because the diffusivity of hydrogen is high as compared to oxygen. As the temperature is increased, humidification decreases and leaves the membrane less hydrated [24]. Oxygen mass fraction is slightly high at 353 K and 333 K as compared to 323K as shown in figure 11.

4. Conclusions
Simulation of single straight channel, proton exchange Membrane fuel cell is carried out under isothermal and non-isothermal conditions. Non-isothermal flow design is co current flow and isothermal model is counter flow design. It is observed that current was being produced at inlet of hydrogen. No oxygen and hydrogen molecules can enter into catalyst layer and membrane. Models predictions are compared with published experimental data which are in good agreement. The deviations observed might be due to single phase assumption in the study. Temperature and pressure are one of very important parameters which play main role in cell performance. The results show that cell performance under non-isothermal conditions is better at higher pressures.

### List of Symbols

| Symbol | Description |
|--------|-------------|
| CL     | Catalyst layer |
| GDL    | Gas diffusion layer |
| CC     | Current collector |
| F      | Faraday constant |
| M      | Molecular weight (kg/mol) |
| ε      | Porosity |
| ρ      | Density |
| u      | Velocity vector |
| \(\bar{\varepsilon}\) | Shear tensor |
| \(\mu\) | Viscosity |
| \(k_i\) | Permeability of GDL and CL |
| \(\bar{\varepsilon}_{eff}\) | Effective stress tensor |
| \(h_k\) | Enthalpy of species k |
| \(Y_k\) | Mass fraction of species |
| T      | Temperature |
| \(T_0\) | Reference temperature |
| \(i_a\) | Transfer current density for anode |
| \(i_c\) | And cathode |
| \(i_c^{ref}\) | Ref. exchange current density per surface area Cathode |
| \(\zeta_a\) | Specific active surface area at anode |
| \(\zeta_c\) | Specific active surface area at cathode |
| \(\iota^{ref}_a\), \(\iota^{ref}_c\) | Ref. exchange current density per surface area of Anode |
| \(\phi_{sel}\) | Electric potential |
| \(\phi_{mem}\) | Protonic potential |
| \(\sigma_{sol}, \sigma_{mem}\) | Electrode conductivity, Ionic conductivity |
| \(\lambda_{eff}\) | Effective thermal conductivity in a porous media |
| \(I_k\) | Diffusional mass flux vector of specie k |
| \(\lambda_S\) | Thermal conductivity of solid porous media |
| \(\eta_{a,c}\) | Over potential at anode and cathode |
| \(\rho_{k,j}\) | Effective diffusional coefficient |
| \(p_0\) | Reference pressure |

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