Numerical Model for Polymer Electrolyte Membrane Fuel Cells with Experimental Application and Validation

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Abstract

The aim of this paper is to present a simple 3D computational model of a Polymer Electrolyte Membrane Fuel Cell (PEMFC) that simulates over time the heat distribution, energy and mass balance of the reactant gas flows in the fuel cell including pressure drop, humidity and liquid water. Although this theoretical model can be adapted to any type of PEMFC, for verification of the model and to present different analysis, it has been adapted to a single cell test fixture. The model parameters were adjusted through a series of experimental tests and the model was experimentally validated for a well defined range of operating conditions: H2/Air as reactants, flow rates of 0.5-1.5 SLPM, dew points and cell temperatures of 30-80°C, currents 0-5 A and with/without water condensation. The model is especially suited for the analysis of liquid water condensation in the reactant channels. A key finding is that the critical current at which liquid water is formed is determined at different flows, temperatures and humidity.

0. Nomenclature

$A_{i,j}$: contact area between nodes i and j (m$^2$)
$\rho_{i}$: thermal capacity of fluid i (J/kgK)
$\alpha$,$\beta$,$\gamma$,$\delta$,$\epsilon$,$\zeta$,$\eta$,$\theta$,$\iota$,$\kappa$,$\lambda$,$\mu$,$\nu$,$\xi$,$\omicron$,$\pi$,$\rho$,$\sigma$,$\tau$,$\upsilon$,$\phi$,$\chi$,$\psi$,$\omega$:

$C_{i}$: thermal capacity of node i (J/kgK)
$D$: channel hydraulic diameter (m)
$F$: Faraday constant 96485 (C/mol)
$g$: heat generated in node i at time k (W/m$^3$)
$h_{i}$: convection coefficient of node i (W/m$^2$K)
$k$: thermal conductivity of the channel hydraulic perimeter (m$^2$/K)
$K$: thermal transfer coefficient between nodes i & j at discrete time k (W/mK)
$K_{w}$: mass flow of fluid xx (kg/s)
$L$: channel length (m)
$m$: mass flow of fluid xx (kg/s)
$m_{w}$: mass flow of gases at average conditions (kg/s)
$M$: molar mass of element xx (g/mol)
$m_{i}$: molar flux of fluid i (mol/s)
$P_{i}$: total pressure of gases (Pa)
$P_{w}$: partial pressure of fluid xx (Pa)
$P_{w}/T_{in}$: saturation pressure of water at temperature $T_{in}$
$\Delta P_{in}$: pressure drop due to major losses (Pa)
$\Delta P_{kin}$: pressure drop due to minor losses (Pa)
$\Delta P_{lk}$: pressure drop due to liquid water inside channel (Pa)
$P_{c}$: channel hydraulic perimeter (m)
$P_{e}$: effective channel hydraulic perimeter with liquid water (m)
$\eta$: fuel cell efficiency

$\eta_{t}$: heat transfer coefficient between gases and in/out channel walls (W)
$\eta_{act}$: heat transfer between gases and active area channel walls (W)
$\eta_{i}$: thermal generation inside the active area due to the reaction (W)
$R_{cont}$: contact thermal resistance between nodes i and j (W/m$^2$/K)
$R_{cont,m-m}$ and $R_{cont,m-c}$: contact thermal resistance between anode and membrane (mem. and cathode respectively) (W/m$^2$/K)
$S$: channel cross section (m$^2$)
$S_{w}$: effective channel cross section with liquid water (m$^2$)
$T_{i}$: temperature of node i for i=1..N, where N equals the number of cubes inside the mesh), at time k (K)
$T_{v}$: temperature in (K)
$V$: volume of the cube represented by node i (m$^3$)
$V_{i}$: vector of nodal temperatures
$\Delta T$: time elapsed between instants k and k+1 (s)
$\Delta t$: time interval from k to k+1 (s)
$\lambda$: thermal conductivity of fluid i of the flow or thermal conductivity of node i (subindex mem for membrane) (W/mK)
$\lambda_{w}$: thermal conductivity of the gases at average conditions (W/mK)
$\mu$: viscosity of the fluid (Pa s)
$\rho$: density of the gas (kg/m$^3$)
$\rho_{f}$: density of the fluid i of the flow or density of node i, what corresponds (kg/m$^3$)
$\rho_{m}$: density of the gas at average conditions (kg/m$^3$)
$\nu$: viscosity of the gas (Pas)
$\omega$: Subindex, represents ambient node
1. Introduction

Polymer electrolyte membrane fuel cells (PEMFC) have tremendous potential as energy conversion devices for a wide range of applications. However there is a great deal to be learned about the various interactions between the different physical phenomena that occur in the fuel cell. There are a vast number of studies in the literature that model the fuel cells with varying degrees of complexity. However the majority do not validate their models with experimental data. Static models, in particular, had an important evolution in the 1990’s. For instance, the isothermal conditions are assumed by many authors such as Springer et al 1991 [6], while others include the thermal distribution (Nguyen 1993) [4]. Spatial variation in only one dimension was considered initially (Springer 1991) [6], while the variation in two and three dimensions was introduced later (Broka, 1997) [13], (Coppo, 2006)[14]. Some works do not model the two phases of water while others do (Bernardi, 1992) [15]. Dynamic models of the physics and control solutions for PEMFC have evolved greatly in the past decade. However, the majority of these dynamic models have a low level of detail. Very few dynamic studies include essential characteristics, such as temperature distribution, the formation of liquid water in different zones of the fuel cell or the pressure drop along the channels. (Shan, 2005) [16] and (Um 2006) [17] present very complete models with heat distribution but their works do not take liquid water into account.

This study presents a dynamic model for PEMFC, which includes the computation of the thermal properties and temperature distribution of the fuel cell and the pressure drop in the reactant gases while taking into account the condensation of water, as well as the resulting effects on the flow and pressure fields. The experimental tests for parameter identification are described. The model is based on the numerical solution of heat transfer problems expressed as various equilibrium differential equations, using numerical iterative methods. In section 2 the model is described, sections 3 and 4 explain the application of the model to a specific single cell and the experimental methodology for the parameter identification and model validation; section 5 collects the principal simulation results and emphasizes the features of the model. The main conclusions are summarized in section 6.

2. Description of the model

The anode and cathode reactants are modeled separately from the stack and interact with it through convection heat transfer in the channels [1]. The pressure drop in the channels is experimentally determined using major and minor losses (for laminar flows) [2]. Water and heat created inside the fuel cell due to the reaction is determined by commonly used equations which are explained in section 2.1.1 and 2.1.2 [3].

The model can be conceived as two sub-models working together, a thermal and a pressure drop. (what is the analytical nature of the thermal equation)

2.1 Thermal model:

The thermal equilibrium equation (thermal generation plus heat transfer equal to energy stored per second) can be approximately solved by creating a mesh of the fuel cell, assuming constant temperature in each cuboid inside the mesh. Iterating with an adequate time interval each cuboid is represented by a node, situated at its midpoint. A correct mesh is crucial to obtain good results. As an example, the mesh used in our case is shown in section 3.2.

In this work, the following finite differences method is used for the simulation of the temperature field:

\[ T_{i}^{n+1} = \frac{\Delta t}{C_i} \left[ \sum_{j \neq i} K_{i}^{j} T_{j}^{n} + (g_{i}^{n} T_{i}^{n}) \right] + \left[ 1 - \frac{\Delta t}{C_i} \sum_{j \neq i} K_{i}^{j} \right] T_{i}^{n} \]  \hspace{1cm} [2.1.2]

where the thermal transfer coefficient between nodes \( K_{i}^{j} \) can be obtained by the following equations: (0 for non adjacent nodes),

\[ K_{i}^{j} = \frac{A_{j}}{\Delta s_i + R_{con, j} + \frac{1}{\lambda_j}} \] \hspace{1cm} [2.1.3]

or

\[ K_{i}^{j} = \frac{A_{j}}{\Delta s_i + R_{con, an-elect} + \frac{1}{\lambda_{an}} + R_{con, cath-elect} + \frac{1}{\lambda_{cath}}} \] \hspace{1cm} [2.1.4]

for adjacent nodes, and
between a surface node and the exterior.

In order to assure stability of the method, the time interval must verified [1]:

\[ \Delta t \leq \frac{C_i}{\sum K_i^j} \]  \[ (2.1.2b) \]

For computation of the temperatures of each node at time \( k \) (\( T(k) \)), it is beneficial to transform the above equations (2.1.2) to the matrix notation. [1]:

\[ T(k+1) = MC \cdot [MK \cdot T(k) + MH \cdot T_{in} + MG] + T(k) \]  \[ (2.1.6) \]

where

\[
MC = \begin{bmatrix}
\frac{N}{C_\infty} & 0 & \cdots & 0 \\
0 & \ddots & \ddots & \vdots \\
\vdots & \ddots & \ddots & 0 \\
0 & \cdots & 0 & \frac{N}{C_\infty}
\end{bmatrix}
\]  \[ (2.1.7a) \]

\[
MK = \begin{bmatrix}
-\sum K_{ii} & K_{i2} & \cdots & K_{in} & K_{i,\infty} \\
K_{i1} & -\sum K_{i1} & \cdots & K_{i,n-1} & K_{i,n} \\
\vdots & \ddots & \ddots & \ddots & \vdots \\
K_{i,n-1} & K_{i,n} & \cdots & -\sum K_{i,n-1} & K_{i,n} \\
K_{i,n} & K_{i,n} & \cdots & K_{i,n} & -\sum K_{i,n}
\end{bmatrix}
\]  \[ (2.1.7b) \]

\[
MH = \begin{bmatrix}
K_{in} & \cdots & K_{in}
\end{bmatrix}
\]  \[ (2.1.7c) \]

And the thermal generation at node \( i \) is:

\[
MG = \begin{bmatrix}
g^{k}_{ch,in/out} & \cdots & g^{k}_{ch,an}
\end{bmatrix}
\]  \[ (2.1.7d) \]

Although it seems to be a quadratic expression, it can be computed as a linear expression due to the matrix MC being diagonal and that MK has a maximum of 7 non-zero values per row and column. This structure reduces significantly the computational time needed by the program.

The channels are included in the matrix representation as isolated nodes with all thermal properties equal to zero (literally holes). The heat transfer between the gases and the channel walls (\( \dot{q}_{ch,in/out}^{k} \) and \( \dot{q}_{ch,an}^{k} \)) is introduced as thermal generation inside the adjacent nodes (to the channels). The thermal generation (\( \dot{q}_{ch}^{k} \)) due to the load inside the active area is also introduced as thermal generation inside the adjacent nodes to the active area and distributed between them. The channels in the active area are in contact their respective gas diffusion layer (3/4 of total surface) and active area (1/4) and the thermal generation due to the reaction is distributed between anode (\( K_{an} \)) and cathode (\( K_{cn} \)), this heat is distributed in the model in the following way:

\[ \sum g^{k}_{i} = \dot{q}_{ch,in/out}^{k} \] for nodes in contact with in/out channels. \[ (2.1.8a) \]

\[ \sum g^{k}_{i} = \frac{3}{4} \dot{q}_{ch,an}^{k} \] for gas diffusion layer nodes in contact with active area channels. \[ (2.1.8b) \]

\[ \sum g^{k}_{i} = \frac{1}{4} \dot{q}_{ch,an}^{k} + \dot{q}_{ch,an}^{k} \] for nodes in active area (distributed between another and cathode). \[ (2.1.8c) \]

2.1.1 Heat transfer (thermal generation) at the active area

The quantity of heat created by the reaction inside the fuel cell can be obtained through its higher heating value efficiency, using the equation:

\[ \dot{q}_{k}^{r} = \left( \frac{1}{\eta} - 1 \right) V \cdot I \text{ where } \eta = \frac{V}{1.482} \]  \[ (2.1.8) \]

The heat is assumed to be generated on the cathode active area, while the heat generation from the anode reaction is neglected [12]. In the literature it is shown that the calculated heat generation due to the reaction is distributed between the anode, membrane and cathode for fully humidified gas streams in a proportion of 3%, 4%, and 93% respectively [11].
2.1.2 Heat transfer at the channels and active area due reactant flows

The heat transfer between the reactant gases and the channel walls, is calculated as convection heat exchange inside circular channels for a laminar flow \[1\].

It is assumed that both reactants and vapor behave as ideal gases, their partial pressures and/or vapor mass flow can be determined by Dalton’s Law

\[
\frac{m_{\text{vap}}}{m_{\text{H}_{2}O}} = \frac{P_{\text{vap}}}{P_{\text{H}_{2}O}} \quad \text{and relative humidity is} \quad H_r = \frac{P_{\text{vap}}}{P_{\text{sat}}(T_{\text{in}})} \leq 1. \quad [2.1.9]
\]

Reactant consumption and water generation can be obtained by the equations

\[
\begin{align*}
\dot{m}_{\text{H}_{2}\text{O, out}} &= \dot{m}_{\text{H}_{2}\text{O, in}} - \frac{M_{\text{H}_{2}O}}{2} \cdot \frac{I}{F} \\
\dot{m}_{\text{O}_{2}\text{, out}} &= \dot{m}_{\text{O}_{2}\text{, in}} - \frac{M_{\text{O}_{2}}}{4} \cdot \frac{I}{F} \\
\dot{m}_{\text{H}_{2}\text{O, out, an}} &= \dot{m}_{\text{H}_{2}\text{O, in, an}} + K_{\text{wtd1}} \cdot \frac{M_{\text{H}_{2}O}}{2} \cdot \frac{I}{F} \\
\dot{m}_{\text{H}_{2}\text{O, out, cut}} &= \dot{m}_{\text{H}_{2}\text{O, in, cut}} + K_{\text{wtd2}} \cdot \frac{M_{\text{H}_{2}O}}{2} \cdot \frac{I}{F}
\end{align*}
\]

as stated in \[3\], \(K_{\text{wtd1}}+K_{\text{wtd2}}=1\) and can be determined by the equations shown in \[4\].

Gas mixture properties are obtained from the data in NASA database \[5\], evaluated at average conditions inside the channel and mixed according to the equations:

\[
\begin{align*}
\dot{c}_{\text{p}} &= \sum c_{p,i} \cdot \frac{m_i}{m} \\
\lambda &= \sum \lambda_i \cdot \frac{m_i}{m}
\end{align*}
\]

\[ [2.1.14, 2.1.15] \]

The heat exchanged between gases and channel can be obtained by an iterative method, explained at \[1\] with equations:

\[
h_m = 3.66 \cdot \frac{\lambda}{D} \quad \text{(for laminar flow)} \quad [2.1.18]
\]

\[
T_{\text{out}} = T_{\text{ch}} + (T_{\text{in}} - T_{\text{ch}}) \cdot \exp\left(-\frac{h_m \cdot L \cdot \text{Per}}{m_m \cdot \text{c}_{\text{p, in}}}ight) \quad [2.1.19]
\]

\[
\dot{q} = -h_m \cdot L \cdot \text{Per} \cdot \text{LMTD} \quad \text{where} \quad \text{LMTD} = \frac{T_{\text{out}} - T_{\text{in}}}{\log\left(\frac{T_{\text{ch}} - T_{\text{in}}}{T_{\text{ch}} - T_{\text{out}}}\right)} \quad [2.2.20]
\]

To this convective heat transfer, the heat created/absorbed due to the water phase change is added; it can be obtained by the difference between liquid water in and out, multiplied by the evaporation heat constant of water (\(h_v\)). Then, heat transfer between channel and the gases (\(\dot{q}_{\text{ch, in/out}}^k\) and \(\dot{q}_{\text{ch, an}}^k\)) is obtained as the sum of both.

2.2 Pressure drop model

The pressure drop in the channel is influenced by the frictional losses in the channel and by the changing channel cross section due to the liquid water. In order to model pressure drop inside the fuel cell, two terms are required: the pressure drop produced by the mass flow and a term incorporating liquid water content inside the fuel cell channels.

2.2.1 Pressure drop related to mass flow

The pressure drop inside a channel can be modeled by calculating the major and minor losses, where the first equation depend linearly with mass flow and are modeled by:

\[
\Delta P_{\text{in}} = -\frac{57}{\text{Re}} \cdot \frac{L}{D} \cdot \frac{\rho_m \cdot v^2}{2} = -\frac{57}{32} \cdot \frac{\mu_m \cdot L \cdot \text{Per}^2}{S} \cdot m_m = -K_{\text{in}} \cdot \frac{\mu_m \cdot m_m}{\rho_m} \quad [2.2.1]
\]

and where the second equation depends quadratically to the mass flow and are modeled by

\[
\Delta P_{\text{ing}} = K_{\text{ing}} \cdot \frac{1}{\rho_m} \cdot m_m^2 \quad [2.2.2]
\]

An explanation of these equations can be found in \[2\]. Roughly, major losses are associated to pressure drop inside a straight channel, where the minor losses are produced in turns, entrances, and exits which are dependent on empirical data of a given geometry.
Properties of the gas mixtures can be obtained by the equations shown in 2.1 and the following:

\[ \rho = \sum \rho_i \]  

\[ \mu = \frac{\mu_i}{1 + \psi_i \frac{M_i}{M}} + \frac{\mu_g}{1 + \psi_g \frac{M_g}{M}} \]  

where \[ \psi_i = \frac{\sqrt{2}}{4} \left( 1 + \left( \frac{\mu_i}{\mu_g} \frac{\bar{n}_i}{\bar{n}_g} \right)^{0.3} \right) \left( 1 + \frac{\bar{n}_2}{\bar{n}_1} \right)^{-0.5} \]  

2.2.2 Pressure drop related to liquid water content inside the fuel cell

When there is liquid water droplets present inside the channels, a greater pressure drop is detected in the fuel cell [10]. This extra pressure drop shall be added to equation 2.2.1. We believe that this loss is directly associated to a narrowing of the effective cross section of the channels. The amount of restriction is related to the pressure drop induced in the channel due to the flow. The higher the pressure drop the greater the force that is placed on the upstream side of the water droplet. This force needs to exceed the surface contact resistance of the droplet and the channel walls for the droplet to be removed. The dynamic associated to the buildup of water droplets in the channel is not taken into account in this model; however a steady state constant is determined experimentally for the specific geometry and surface finish of the gas channels.

This effect, at least for the single cell, can be modeled as a special major loss with the equation:

\[ \Delta P_{tw} = -K_w m \dot{m}_w \]  

2.2.3 Combined pressure drop in the case of liquid water inside the fuel cell

According to the previous section, the pressure drop is determined by:

\[ \Delta P = \Delta P_{in} + \Delta P_{sl} + \Delta P_w = -(K_{in} \frac{\mu_m}{\rho_m} - K_w) \dot{m}_m - K_{sl} \frac{1}{\rho_m} \dot{m}_w^2 \]  

From equation [2.2.4] and comparing it with equation [2.2.1] it is possible to obtain an approximate equation that models the effective cross section when liquid water appears. It is:

\[ K_{in} \frac{\mu_m}{\rho_m} - K_w = \frac{57}{32} \frac{\mu_m}{\rho_m} L \frac{P_{tw}}{S_{tw}^3} \]  

[2.2.5]

If it is assumed that channel section at active area remains square, even if liquid water is accumulated, then the following equation is obtained.

\[ S_{tw}^2 = \frac{2}{57} \frac{L \left( K_{in} - K_w \frac{\rho_m}{\mu_m} \right)}{K_{in} - K_w} \frac{P_{tw}}{\dot{m}_m^2} \]  

[2.2.6]

The effective cross section has been found experimentally and is relatively constant for a wide range of operating conditions. Nevertheless, this may be true in the active area of the single cell because the experiments were done with very large stoichiometries, however in a multi-cell stack it will require a more complex equation.
3. Application of the model to a single cell test fixture

3.1 Single cell test fixture description

The single cell test fixture modeled in this paper is an ElectoChem® (model # EFC-05EFC-05-02-02) 5 cm² active area fuel cell. It is equipped with a Nafion® 115 membrane with 1 mgPt/cm² catalyst loading and Toray® carbon fiber paper (Type TGP-H-060) gas diffusion layers, serpentine/straight channel flow field with groups of three channels that come back together at each turn in a uniting manifold. There are five straight sections in the flow field which makes it similar to a five pass serpentine configuration, as shown in figure 3.1.1.

![Image of fuel cell channels configuration](image.png)

The dimensions of the channel are 0.78mm wide, 0.78mm deep, and the space between each channel is 0.78mm. The length of the straight section of the channel is 23.25mm. The gases enter the flow field through two 1.6mm diameter holes and exit through the same size holes. The flow fields are machined into POCO® graphite plates that are 19.1mm thick. The Teflon® fitting which connect the fuel cell to the test station are also screwed into the graphite plates. The current collector bus plates are gold plated copper for enhanced surface conductivity. Resistive square planner 60W heaters with adhesive are in the center of each bus plate with the dimensions of 50mm wide.

<table>
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<tr>
<th>Dimensions</th>
<th>Material</th>
<th>x (mm)</th>
<th>y (mm)</th>
<th>z (mm)</th>
<th>volume (m³)</th>
<th>convective area (m²)</th>
<th>contact area (m²)</th>
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</tr>
</tbody>
</table>

Table 3.1.1 Dimensional properties of the cell

3.2 Model assumptions

The following assumptions are made in order to build a reliable, fast and precise model.

1. One global convection coefficient for the whole cell: With the experimental setup it is impossible to obtain different coefficients for each surface, error commited is low.
2. Physical properties of the materials do not depend on temperature: Temperature range is slim (40-80°C).
3. Calculated properties (c_p, λ, ρ, μ) of dry air do not change due to oxygen depletion; due to large stochiometry.
4. Properties of the flow inside the active area are determined for the average mass flow between entrance and exit.
5. Ideal gases are assumed.
6. It is assumed that K_wd₁=0.3 and K_wd₂=0.7, in fact these constants have little effect in the simulations due to the high stochiometries.
7. Channels in active area modeled as one 3-parallel long straight tube: Reduces precision and computational time. For a small single cell this assumption can be made. For larger fuel cells or in order to obtain more precise results, the channel can be modeled with shorter consecutive tubes.

8. No heat transfer by radiation: Low temperature, non lineal equation (thermal model is lineal), the determined convection coefficient includes radiation at a given temperature.

3.3 Fuel cell mesh

The mesh applied in the model of the single cell is shown in Fig 3.2.1. It divides the different zones of the cell with the corresponding materials. Smaller cuboids are placed in the active area for higher precision.

Fig. 3.2.1 Single cell mesh

3.3 Properties of the materials

Most of the constants were obtained from material suppliers, fuel cell supplier or different reference books.

<table>
<thead>
<tr>
<th>Material</th>
<th>(\rho) (kg/m(^3))</th>
<th>(c_p) (J/(K·Kg))</th>
<th>(\lambda) (W/(m·K))</th>
</tr>
</thead>
<tbody>
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<td>-</td>
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</tr>
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</table>

Table. 3.2.2 Thermal properties of the cell

4. Experimental methodology

In this section we present the single cell test fixture, the experimental setups, and the LabView hardware and software used for data acquisition.

4.1 Experimental Setup

The test station consists of two reactant (anode and cathode) gas subsystems. Each subsystem contains a Bronkhorst® mass flow controller, membrane based humidification with dew point sensors for control, inlet line heaters to prevent condensation, absolute pressure transducers at the inlet, differential pressure transducers between the inlet and outlet of each reactant, and back pressure regulator at the outlet of the fuel cell to control system pressure. The mass flow controllers are each calibrated for the specific gas (Hydrogen for the anode and synthetic air for the cathode).
There are 8 temperature sensors that come from the fuel cell by way of K Type thermal couples. Four of the measurements are of the graphite plates (two in each plate) with one connected to a RedLion® PID controller model #T4810105, which controls the temperature of the fuel cell. The cooling of the cell is attained mainly by natural convection. The other four temperature measurements are of the reactants inlets and outlets. The inlet temperature measurements are close to the outlet of the gas line heaters but still out side of the cell. The outlet temperatures are measured inside the fuel cell in the outlet manifold. All the measurement and control are made in real time through LabView® which is explained in more detail in the following section.

Data acquisition and control system consists of two computers define the data acquisition and control system. The first one is responsible for the user interface and setting the operating conditions. This is done by means of a graphical interface developed by our laboratory through LabView. The second computer runs under a Real Time Operating System (RTOS) and works in deterministic mode offering consistent and stable functionality, implementation of controllers and stores the data. Inside RTOS computer there areCompactRIO cards with input/output modules that contain configurable signal conditioning, isolation and screw terminals to provide direct connections to our sensors and actuators.

4.2 Experiments development for parameter identification and verification

In order to adjust some unknown parameters, to be known: global convective coefficient, internal contact resistances and pressure drop constants, and to verify the model, several experiments were performed, and simulated with the model. These experiments where designed based on the data acquisition components and the fuel cell active area. All parameters were adjusted following a “minimum global error” strategy.

The first set of experiments were designed to obtain the thermal constants (global convective coefficient and internal contact resistances) and check the thermal model (fig. 4.2.1 and 4.2.3), all of them were done without reactant flows. These experiments are divided in six groups:

- Adjust the current of the heaters in order to reach a thermal equilibrium inside the cell. Used to calculate and verify the convection heat transfer parameter of the cell.
- Heat the cell with both heaters (no equilibrium is reached). Used to calculate and verify convection parameter and heat capacity of the cell.
- Cool the cell (no refrigeration system, no equilibrium is reached). Used to verify all the previously stated parameters.
- Heat the cell with only one heater and adjust its current to reach thermal equilibrium inside the cell. Used to calculate and verify the convection parameter and internal contact resistances.
- Heat the cell with only one heater, at maximum current (no equilibrium is reached). Used to verify all the previously stated parameters under different conditions.
- Follow a temperature profile turning on and off the heaters. Used to verify all the previously stated parameters.

The adjusted parameters are:

<table>
<thead>
<tr>
<th></th>
<th>$R_{ij}$ contact resistance (W·m²/K)</th>
<th>$h$ convectivity (W/(m²·K))</th>
</tr>
</thead>
<tbody>
<tr>
<td>heating sides</td>
<td>-</td>
<td>12</td>
</tr>
<tr>
<td>heaters</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>copper plates</td>
<td>Copper - Graphit 0</td>
<td>12</td>
</tr>
<tr>
<td>graphite plates</td>
<td>Graphit - Diffusion layer 0.00045</td>
<td>12</td>
</tr>
<tr>
<td>diffusion layer</td>
<td>Dif. layer Anode - MEM 0.00045</td>
<td>-</td>
</tr>
<tr>
<td>gasket</td>
<td>Dif. layer Cathode - MEM 0.00045</td>
<td>-</td>
</tr>
<tr>
<td>membrane</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>lab. support</td>
<td>-</td>
<td>9.23</td>
</tr>
</tbody>
</table>

Table. 4.2.1 Thermal properties of the cell (adjusted)

The second set of experiments were designed to check pressure drop inside the cell (table 4.2.2), all of them were done with H2/Air, at different flow rates (0.5-1.5 SLPM), at different dew point and cell temperatures (30-80ºC), different currents (0-5 A) and with/without water condensation. It was impossible to work with low flow rates due to the extremely low pressure drops. Most of the experiments were done with flow rate 1.5 SLPM on both sides.
The obtained constants are the following ones:

<table>
<thead>
<tr>
<th></th>
<th>H2</th>
<th>AIR</th>
</tr>
</thead>
<tbody>
<tr>
<td>$K_{in}$ (s/m)</td>
<td>$3.7\cdot10^{12}$</td>
<td>$9\cdot10^{12}$</td>
</tr>
<tr>
<td>$K_{vis}$ (Pa·s²/m³)</td>
<td>$1.8\cdot10^{12}$</td>
<td>$1.9\cdot10^{12}$</td>
</tr>
<tr>
<td>$K_w$ (Pa·s/kg)</td>
<td>$2.8\cdot10^{7}$</td>
<td>$7.5\cdot10^{7}$</td>
</tr>
</tbody>
</table>

Table. 4.2.2 Pressure drop constants (adjusted)

The comparison between acquired data and simulated with the Matlab model is shown in the next two tables.

<table>
<thead>
<tr>
<th></th>
<th>2 heaters to thermal equilibrium</th>
<th>2 heaters full power</th>
<th>2 heaters 1/4 power</th>
<th>1 heater thermal equilibrium</th>
<th>1 heater full power</th>
<th>no heaters</th>
</tr>
</thead>
<tbody>
<tr>
<td>$P_{heaters_real}$ (W)</td>
<td>-15,12</td>
<td>-16,38</td>
<td>-100,28</td>
<td>-28,50</td>
<td>-28,50</td>
<td>-16,38</td>
</tr>
<tr>
<td>$P_{conv_mod}$ (W)</td>
<td>15,90</td>
<td>18,02</td>
<td>14,03</td>
<td>23,21</td>
<td>12,11</td>
<td>14,67</td>
</tr>
<tr>
<td>$P_{stored_mod}$ (W)</td>
<td>0,00</td>
<td>0,00</td>
<td>95,32</td>
<td>67,28</td>
<td>13,29</td>
<td>12,37</td>
</tr>
<tr>
<td>Error (W)</td>
<td>0,78</td>
<td>1,64</td>
<td>9,07</td>
<td>-9,79</td>
<td>-3,10</td>
<td>-1,46</td>
</tr>
</tbody>
</table>

Table. 4.2.3 Heat balance for several experiments

Table. 4.2.4 Pressure drop comparison of experimental data with simulated data

5. Selected results for the single cell test fixture

This section presents selected results for the 2D temperature distribution snapshot as well as the temperature evolution in the cell.

5.1 Temperature distribution inside the fuel cell at a fixed time

Figures 5.1.1 through 5.1.3 show the temperature distribution across the fuel cell and the change in the inlet and outlet reactant temperatures at a given time where every rectangle represents the temperature of a cuboid. This corresponds to a small fuel cell where the majority of the heat comes from two contact resistances, one on each side. It is critical to determine the thermal distribution because of its relationship with the phase change of water.
5.2 Evolution of fuel cell temperature over time

The model simulates the evolution over time of every cuboid temperature and temperature of the reactant. This allows for the simulation of non-equilibrium temperature pattern in the fuel cell. Figure 5.2 shows the temperature evolution through time, at a fixed location in the fuel cell while the heaters are cycled on and off.

5.3 Figure 5.3 shows relative humidity of the gases and liquid water content, through the anode and cathode. These illustrations are important in order to determine if and where water is condensing inside the fuel cell, which is a key factor in fuel cell efficiency and stability. In this case; on the anode side the hydrogen reaches its saturation temperature at the first node entering the fuel cell and starts condensing, however on the cathode side the air reaches its saturation temperature at the third node.
**5.3** Relative humidity and liquid water content of the reactant gases through the cell

**5.4** Figure 5.4 shows the current at which water condenses inside the active area (critical current), for different temperatures and flow rates. Any point above each line will result in condensation in the flow field at the given conditions. Raising the current while all the other parameters remain constant will eventually lead to water condensation inside the active area. This model is designed to determine this critical maximum current.

**5.5** Figure 5.5 shows the pressure drop in the channels, for a fixed flow rate and the difference in temperature between the fuel cell and dew point while modifying the current. The discontinuity corresponds to the critical current. The simulation shows the sensitivity of condensation of water with respect to fuel cell and dew point temperatures, current and flow rates. A one degree change in the temperature difference greatly affects the critical current. Increasing overall temperature increases the capacity of the gases to remove water in the vapor form which in turn increases the critical current.
Figures 5.6 show the pressure drop for the reactant gases with a fixed stoichiometry and temperature vs. modifying the current. Increasing the current implies increasing the flow rate; which provokes a greater pressure drop inside the cell. It can be seen that both dew point temperature and stoichiometry have a great effect on the pressure drop. This is due to the quantity of water added to the cell. NOT QUIT SURE ABOUT THIS ONE

Fig. 5.5 Pressure drop against current, fixed mass flow

Fig. 5.6 Pressure drop against current, fixed stoichiometry (analog for AIR)
6. Conclusions
In this work a four-dimensional model of heat distribution and pressure drop for PEMFC was developed and experimentally tested. It includes a simple two phase flow pressure drop model which is representative of the true effects of liquid water in the flow field. The model is especially suited for the analysis of liquid water condensation in the reactant channels.

The model was applied to a single cell test fixture and some of the parameters were adjusted through a series of experimental tests. The model was validated using experiments designed for this purpose. The identified parameters are contact resistances, convective heat transfer coefficients and pressure drop constants.

Several simulations were performed and analyzed. The results show that there is a step jump in the pressure drop when the critical current is reached. This determines the maximum current that should be applied to the fuel cell to avoid flooding of the catalyst layers and diffusion layers. Critical currents and the effects of liquid water and water vapor on the pressure drop of the fuel cell were evaluated for a range of operating conditions.

The model shows that the current at which water condenses inside the reactant channels (or critical current) depends strongly on the flow rate and even more on the temperature difference between the fuel cell and the reactant dew point. The model also shows that lower overall temperature gives a smaller critical current.

7. Future work
Improvements in future versions of the model are:
- Apply the model to a fuel cell stack.
- Introduce more complex equations to model the pressure drop inside the cell. To achieve it, it is necessary to apply the model to a fuel cell (not single cell) with higher pressure drops and lower stoichiometry in order to have a greater range of possible experiments. For the operating range of the single cell in the lab, it has been checked that this simple model is quite enough.
- Introduce more complex equations to model water movements inside the active area. Once again, that will be necessary when modeling a fuel cell stack. As a reference, these equations can be found at [4].
- Introduce an electrical model, it is to say, a sub-model which determines cell voltage at working conditions. Some of the future objective of research that will be conducted with more lab experiments and the help of the model are:
  - Determine the quantity of liquid water that remains inside the cell dynamically and in steady state.
  - Determine, or verify the water movements inside the active area. It is related to the figure 5.5.1.
  - Create a more universal and precise approximation of section reduction due to liquid water formation inside the cell.

8. Acknowledgments
This work has been funded partially by the projects CICYT DPI2007-62966 and CICYT DPI2004-06871-C02-01, of the Spanish Government.

9. Model & Contact
The implementation in Matlab/Simulink of the model for the single cell used in our laboratory can be found at the following address:
http://wikiri.upc.es/wiki/

10. References