Modeling and parametric study of solid oxide fuel cell performance

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ABSTRACT

Renewable energies are in the news these days and are seen as a solution to endorse energy independence and diminish greenhouse releases. One possible new renewable source that is emerging as a promising technology is the fuel cell. The ‘F.C’ is an electrochemical system that transfers the chemical power of a redox response into electrical energy with simultaneous production of water and heat. In fact, there are numerous kinds of fuel cells; in our work we are interested in studying the solid oxide fuel cell (SOFC). In the context of this work, a modeling tool has been implemented for SOFCs to the analogy between the electrical, thermal and chemical domains; this way of proceeding constitutes a simple, evolutionary and efficient tool. Using this modeling, a simulation was carried out in order to obtain the dissimilar characteristics and the impact of the studied parameters on the performance of the SOFC fuel cell.

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1. INTRODUCTION

In recent decades, solid oxide fuel cell (SOFC) systems have attracted a lot of attention, owing to their potential uses as stationary power producers and mobility (ground, marine, air). The great energy conversion efficiency and minimal hazardous emission levels of SOFCs make them appealing (only the CO2 released by the hydrogen production process is a concern). Modularity, fuel flexibility, and minimal noise are among the other benefits [1]-[3]. Furthermore, high operating temperatures afford good features, such as potential utilize of SOFCs in extremely proficient cogeneration applications. Solid oxide fuel cells are also appropriate for internally reforming the fuel (e.g. natural-gas, propane, methanol, gasoline, and diesel), as a result making it doable to stay away from the acceptance of both highly sophisticated, expensive outside reformer and to remove the complexities from fuel storage [4]. The giant confronts to promote the dispersion of SOFC–based energy conversion systems are mostly connected to manufacture costs and permanence. The accomplishments of these objectives shall definitely contribute to upholding the technology and lastly beginning a mass production stage. Above and beyond expenses and performance, long-standing steadiness is a crucial condition for the marketable application of the SOFC technology [5], [6].

The hydrogen vector, namely fuel cells and electrolysis (to make hydrogen), also offers promising potential for these large-scale storage applications, where conventional batteries play a key role. As a result, the storage solution will be multi-system, and the electrical study of the association of many sources of energy - hybridization - will take on new significance in this context. Solid oxide fuel cells (SOFC) are electrochemical devices that allow chemical energy to be converted into electrical energy. The premise is straightforward: water and electricity are produced from a fuel like hydrogen and an oxidant like oxygen.
Apart from the fact that these systems are environmentally friendly since they do not emit greenhouse gases, they also show the potential of hydrogen as an energy vector for both the production of electricity and the storage of energy in chemical form. The applications primarily pertain to decentralized energy generation for residential and urban regions with electrical powers ranging from a few kW to a few hundred kW over hundreds of hours [7], [8]

2. OPERATING MODEL OF SOFC CELL

Cell is a set of an anode, an electrolyte, a cathode and the interconnections. The anode is supplied with hydrogen or gas mixture (fuel) and the cathode is supplied with air (oxidizer). The operating principle of the SOFC is hinged on the following mechanism: oxygen is dissociated at the cathode into $O_2^-$, then the cation migrates through the ionic conductive electrolyte at high temperature (from 700 °C) and will combine at the anode with hydrogen to form water and release electrons that go via an outsider electrical circuit [8]-[10]. The main characteristic of SOFC is therefore their operating temperature (600 to 1,000 °C), which is mandatory to obtain a adequate ionic dynamism of the ceramic electrolyte. This temperature has a double advantage. Firstly, it allows the direct use of hydrocarbons, which can be easily reformed without the need for a noble metal catalyst. Secondly, it produces a high level of heat that can be easily used in cogeneration, with an efficiency of up to 80%.

But it also has a disadvantage; the heating is long and complicates any use with short and repetitive cycles (as in the case of transport). For these reasons, the technology lends it itself particularly well to decentralized electricity production and cogeneration (areas covering powers ranging from 1 KW to several tens of MW) [11]-[13]. Thanks to its high efficiency and its potential capacity to operate directly with liquid hydrocarbons, it will also find an outlet in naval propulsion. The development of this type of cell, because of its high operating warmth and the resolution of thermo-mechanical problems of material resistance, is quite complex. One of the particularities of the SOFC is its solid electrolyte, usually zirconium (Zr2) doped with a mole of 8 to 10% ytterbium (Y3+), which acts as a conductor for the oxygen anion ($O_2^-$).

The following Figure 1 shows the operating principle of the SOFC fuel cell.

![Figure 1. Picture of SOFCs](image)

3. POLARIZATIONS

Polarizations, or overvoltage’s, are voltage losses as consequence of drawbacks in the structure of the F.C polyanionic cellulose (PAC) and materials microstructure. The ohmic confrontation of ‘oxyg’ ions passing through the electrolyte (i.R $\Omega$), electro_chemicals activated hurdles at the ‘ano’ and ‘cath’, and terminaly focus polarizations in view the incapacity of gas’s to distribute at high velocities through the porous anode and cathode [10]. The cell voltage can be calculated employing in (1):

$$\text{Volt} = E_0 - R \times i - \eta_{\text{cathode}} - \eta_{\text{anode}}$$ (1)

In SOFC, it is frequently essential to spotlight on Ohmic and attentiveness biases because high operating heats background little’s activation polarizations. Nonetheless, like the lowest boundary of the S.O.F.C functioning hotness is came close to (~873,15 kelvin), these polarizations turn out to be significant. Research by Huo [14] The above equation is utilized to find out the S.O.F.C voltage. This method showed results which gives aexcellent accord with scrupulous investigational Information and deprived agreement for
experimental working parameters other than the originals. Additionally, most of the equations employed need the addition of multiple factors that are hard or unfeasible in order to identify. This that any process of optimizing the S.O.F.C operating factors and the choosing very complicated geometry configuration. Because these circumstances, a few other equations have been suggested [15].

\[
E_{\text{SOFC}} = \frac{E_{\text{max}} - E_{\text{max}} \cdot \eta_f \cdot r}{r} (1 - \eta_f) + 1
\]

Where:
- \(E_{\text{SOFC}}\) : Cell voltage
- \(E_{\text{max}}\) : The Nernst equation predicts the maximum voltage
- \(i_{\text{max}}\) : Current density at its maximum (for a given fuel flow)
- \(\eta_f\) : Factor of fuel usage [15], [16]
- \(r\) : The electrolyte's particular ionic resistance
- \(r\) : The electrolyte's particular electrical resistance

4. THE LOSSES SOFC MODEL

There are three main forms of polarization overvoltage's: activation, ohmic and concentration. A constant offset also contributes to the total polarization, resulting from small overvoltages like contact resistor, inside-current and leakage. According to [17], the shift was approximated to be 0.07 Volts. The additions of the different polarizations result in a voltage sag from the optimal Nernst potential to the actual working value. As the interconnector and the electrodes are iso-polar, the array voltage is constant over the cell and can be expressed in (3):

\[
V_{\text{SOFC}} = E_{\text{Nernst}} - V_{\text{act}} - V_{\text{ohm}} - V_{\text{con}} - V_{\text{offset}}
\]

The total energy derived from the S.O.F.C is determined as shown in (4):

\[
E_{\text{cel}} = V_{\text{SOFC}} \cdot I_{\text{SOFC}}
\]

4.1. Activation polarization

`Activ` polarization means the power obstacle being diminished in order to enable the electrochemical feedback taking part at the interface of the electrodes [18]. This quantity of power unavoidably leads to a noteworthy overvoltage, that is typically modelled by the non-linear equation recognized as butler-wolmer formula [19]:

\[
V_{\text{act}} = \frac{R_i J_i}{a(t_i)^{2/3}} \left( \frac{J_i}{a(t_i)} \right) (\sinh^{-1} \left( \frac{J_i}{a(t_i)} \right))
\]

where \(a\) is the charge transmitt coefficient and \(J_0\) is the switch present mass.

4.2. ‘Ohm’ polarization

‘Ohm’ polarization primarily relies on the ionic momentum of electrodes and the electronic momentum of an electrolyte. Such over voltages are assessed finalizing the content of every part of the S.O.F.C, as (6):

\[
V_{\text{ohm}_k} = \frac{l_k}{\sigma_k(t_i)} J_i
\]

\[
V_{\text{ohm}} = \sum V_{\text{ohm}_k} k = [\text{ano}, \text{cath}, \text{el}]
\]

The conductances are approximated with, suggestions in the review of literature [17] for 2nd edition ceramic ‘S.O.F.C’ stacks, in which the ‘ano’, ‘cath’ and ‘el’ cover materials are nickel cermets, strontium-doped lanthanum manganese and yttrium oxide cured zirconia, correspondingly:

\[
\sigma_{an} = 10^3
\]

\[
\sigma_{ca}(T_s) = c_{-3}(T_s)^2 - c_{-2}(T_s)^2 - c_{-3}
\]

\[
\sigma_{el}(T_s) = c_4(T_s^4 - 273.15)^2 + c_{-3}(T_s - 273.15) + c_6
\]
4.3. Concentration polarization

As the relative stresses of hydrogen and oxygen at the ‘ano’ and ‘cath’, respectively, drop as fuel is depleted. The rate of depletion is determined by the cell’s average current density. When a result, as a current thickness rises, the relative stresses fall, and a deficiency number re-agents is eventually carried to the electrodes. Until the voltage is decreased to \([20, 21]\), considerable losses occur. ‘Ano’ and ‘cath’ restrict currents are the values at which this phenomena occurs. Concentration polarization is the term for the voltage loss that occurs at high current densities and can be calculated as (8).

\[
V_{\text{Con}}^i = -\frac{R T_i}{2 F} \left[ \frac{1}{2} \ln \left( 1 - \frac{J_i}{J_{cs}} \right) + \ln \left( 1 - \frac{J_i}{J_{as}} \right) - \ln \left( 1 + \frac{\phi_{H_2}}{\phi_{O_2} \cdot I_{as}} \right) \right]
\]

The ‘ano’ and ‘cath’ limiting currents are determined as a function of the species scattering coefficients, using the formula given in [22].

5. MODELING OF THE SOFC

The purpose of this paper is to give the model adopted to account for the physical phenomena in the cell. The first part of the paper is dedicated to the modeling principles applied to the chemical, thermal and electrical domains. The second part concerns the application of the modeling to the S.O.F.C Cell. The chemical § thermal modeling of the ‘F.C’ is based on their formal analogy with the electrical domain. The temperature is similar to the voltage and the heat flow is similar to the current. Two heat sources within the cell are considered: endothermic or exothermic heat sources produced by chemical reactions and Ohmic heat sources. Furthermore, pressure is similar to voltage and flow is similar to current. The application of this analogy offers the possibility of representing physical laws within a system with electrical circuits. The use of such a representation allows the use of different techniques in the electrical domain: putting chemical and thermal models in series or in parallel, realization of the regulation of chemical or thermal quantities with the help of methods used in automation (ex: pole compensation), ease and flexibility of integration of developed models in a global model (battery coupling or battery associated with the converter) The electrical model translates the electrochemical phenomena of each cell. The electrical model of the SOFC is based on the reversible Nernst voltage and the different voltage drops.

5.1. Electro-chemical system

We will choose a static SOFC system based on thermodynamics and electrochemical reactions to determine the electrical characteristics of the fuel cell. In a second step we vary the parameters and observe the variation [23]:

\[
\Delta G = \Delta H - T \Delta s
\]  

(9)

\[
\Delta H = \Delta U + \Delta PV
\]  

(10)

\[
\Delta U = \omega + \phi
\]  

(11)

\(\Delta V\): Growth in size, \(\Delta U\): Internal energy changes, \(\omega\): labour, \(\phi\): thermal.

\[
\Delta G = \omega + \phi + P \Delta V - T \Delta s
\]  

(12)

\[
Q = T \Delta s
\]  

(13)

\[
\Delta G = \omega + P \Delta V
\]  

(14)

\(-P \Delta V\), thus:

\[
\Delta G = -n \Gamma \epsilon - P \Delta V + P \Delta V
\]  

(15)

As a result, it was simply:

\[
\Delta G = -n \Gamma \epsilon
\]  

(16)

With:

\(\epsilon\): electromotive strength of the stack, \(\Gamma\): Constant of Faraday, \(n\): number of electrons transported.
\[
\Delta G_r = G_{H_2O} - G_{H_2} - \frac{1}{2} G_{O_2} 
\]

\[
\Delta G_r = G_r - NRT \log \frac{P_{O_2}^\frac{1}{2}}{P_{H_2O}} 
\]

\[
P_{O_2}, P_{H_2}, P_{H_2O} 
\]

\[
E = E^0 + \frac{RT}{nF} \log \frac{P_{O_2}^\frac{1}{2}}{P_{H_2O}} 
\]

5.2. Real potential of the stack

The survoltages happened because of the shedding of non-reversible load, also named, which are polarization of ‘activ’, ‘Ohm’ polarization § ‘conc’. High-temperature fuel cells are able to generate enough heat to power the vaporiser, to power the steam generator. This will produce a hydrogen-rich reformate which is then which is then supplied to the cell.

\[
V = E - \text{losses} 
\]

5.3. Capacity and quality requirements

When the temperature of the battery drops, the variance of free energy increases, hence the standard voltage of a stack is in relation to the change in its standard free energy [24].

For water produced in a liquid state at 25°C, it is:

\[
\Delta G_r = -237,190 \text{ joul/Mol.} \]

\[
\Delta H_r = -285,840 \text{ joul/Mol.} \]

The Rated stack Voltage is:

\[
E^0 = \frac{-\Delta G}{nF} = 122.9 \text{ V} 
\]

\[
\Delta G_r = -228,590 \text{ j/Mol.} \]

\[
\Delta H_r = -241,830/\text{mol} \rightarrow E^0 = 11.8 \text{ V.} \]

However at 1000°C [25].

\[
\Delta H_{1000°C} = \Delta H_{25°C} + \int_{25}^{1000} \Delta C_p \text{d}t 
\]

\[
\Delta S_{1000°C} = \Delta S_{25°C} + \int_{25}^{1000} \frac{C_p}{T} \text{d}t 
\]

\[
\Delta G_{1000°C} = \Delta H_{1000°C} - T\Delta S_{1000°C} 
\]

In this scenario, the start signal voltage is: \(V_0 = 0.935\text{Volt}\).

5.4. Performance of the cell

The chemical reaction is followed by a change in entropy, which leaves a free enthalpy, which may be transformed into work, in this case electrical energy, with the remainder being converted into heat during the chemical reaction in the stack. The report of the free enthalpy on the standard enthalpy of the reaction of water formation corresponds to the performance thermodynamics for a reversible transformation:

\[
\eta = \frac{\Delta G^0}{\Delta H^0_{max}} 
\]

This performance ranges from 83 percent for liquid water creation to 95 percent for water formation in gaseous forms under typical conditions, but above 10^3 °C, the theoretical performance is:

\[
\eta_{th} = 0.7337% 
\]

a. The voltaic performance [24]-[26]:

Of the in (24) one can define a voltaic performance to performance by voltage:

\[
\eta_{V} = \frac{V_e(t)}{\eta} 
\]
b. The electrical performance:

It is motivating to describe the notion of electrical performance:

$$\Gamma_{\text{electrique}} = \Gamma E \frac{\Delta g_0 V_c(I)}{\Delta n^0 F^0} \frac{1}{V_c} \frac{E}{E_0} \frac{1}{E_{\text{max}}} \frac{1}{V_c(I)} \frac{V_c}{E_0} (26)$$

$$E = \frac{\Delta g_0}{n^0 F^0} \frac{1}{V_c} \frac{E}{E_0} \frac{1}{E_{\text{max}}} \frac{1}{V_c(I)} \frac{V_c}{E_0} (27)$$

It is then possible to directly measure this electrical performance experimentally using a voltage measurement, or to plot the electrical performance to T Constant by drawing the characteristic V(I) returns. Because this performance is highly dependent on temperature, it is necessary to recalculate E_{\text{max}} for each temperature [27]:

$$\Gamma_{\text{Fonctionnement}} = \frac{V_c(I)}{V_c(0)} \frac{V_c(I)}{V_c(0)}$$

6. SIMULATION OF THE SOFC

Simulation and mathematical models shall undoubtedly help in the development of a variety of power generators. But they are even more important in the development of fuel cells. Simulation of these cells allows you to focus on experimental research and improve the accuracy of interpolation and extrapolation of the results. In addition, mathematical models serve as a valuable tool for the design and optimization of fuel cell systems. Simulations are run on different systems with different software packages [28]-[30]. The following Table 1 shows the simulation parameters.

| Table 1. The SOFC model values [30] |
|-----------------------------------|
| **Settings**                     | **Data** | **Available units** |
| Nominal power                    | 15,000   | Watt               |
| Number of Stack                  | 384.00   |                     |
| ‘Ano’ forces                     | 2,280.006| Torr               |
| ‘Cath’ forces                    | 2,280.006| Torr               |
| fuel airflow                     | 0.0012   | Mol/s              |
| combustible Water rate           | 0.00001  | Mol/s              |
| combustible Hydrogen rate        | 0.00009  | Mol/s              |
| T fuel                           | 799.85   | °C                 |
| T air                            | 799.85   | °C                 |
| Current Density                  | 60.00    | A/cm²              |
| V_{\text{out}}                   | 440.00   | Volt               |

7. RESULTS AND DISCUSSION

In our study by using a simulation model with block diagrams under simulink Matlab to obtain the electrical and electrochemical characteristics of the SOFC type fuel cell, and then we varied the parameters of the effect of temperature and the effect of gas pressure, the results are the following:

In order to verify the performance of the fuel cell generation system under normal operating conditions, the previously established model was simulated in the MATLAB/Simulink software environment. MATLAB/Simulink software environment. For this simulation, the model was planned With the parameter values extracted from the literature together with experimental data, the results are presented in the following and the interpretation is just after the presentation of the figures.

7.1. Polarization curve

A polarization curve for the SOFC fuel cell is shown in Figures 2 and 3. The characteristic begins with a no-load voltage at zero current, as can be shown. A dramatic reduction in voltage is noted when the cell current increases. The loss of activation voltage and parasites produced by the pace of the electrochemical process create this reduction in cell voltage. The non-linear activation losses contribute to the V(i) curve’s downward tendency at low current densities. The activation losses remain the most significant in absolute terms. The Ohmic loss generated by the electrical resistances of the cell causes an almost linear drop in cell voltage and an increase in current above specific values [31]-[33].
7.2. Temperature effect

In this part we have varied the temperature for several values (T=600 °C, 700 °C, 800 °C, 900 °C, 1000 °C) and we fix the partial pressure. Figures 4 and 5 show figures at various temperatures typical of SOFCs. The activation voltage drop dominates the voltage drop in the low current region (Figure 6). As the load current increases, the resistance voltage drop increases rapidly and is the main cause of the SOFC voltage drop (Figure 7). At certain load currents, the centralized voltage drop in the SOFC reduces the output voltage of the fuel cell Figure 8. Figure 4 also demonstrates the effect of temperature on SOFC-VI characteristics. The SOFC output voltage is high at low temperatures in the low current region and high at high temperatures in the high current region. From this figure, we can see that rising the cell temperature has a beneficial effect on the cell voltage. This is explained by the high ionic conductivity of the cells (electrolytes and electrodes) at high temperatures (such as T=900 °C, T=1000 °C) Figure 4. Therefore, it is needed to raise the temperature to ensure proper cell function [34], [35]. Tables 2 show the power values according to the influence of temperature:

![Figure 2. Current-voltage characteristic of a fuel cell SOFC](image1)

![Figure 3. Current-power characteristic of a fuel cell SOFC](image2)

![Figure 4. Current-voltage characteristics for different temperatures values](image3)

![Figure 5. Current-power characteristics for different temperatures values](image4)

![Figure 6. Activation losses characteristics for different temperatures values](image5)

![Figure 7. Ohmic losses characteristics for different temperatures values](image6)
Table 2. Influence of temperature on power

| T (°C) | Power (Watt) | Ohmic Losses (Volt) | Activation Losses (Volt) | Concentration Losses (Volt) |
|--------|--------------|---------------------|--------------------------|-----------------------------|
| 600    | 15,120       | 10.38               | 0.6345                   | 0.001938                    |
| 700    | 15.25        | 12.58               | /                        | /                           |
| 800    | 15.32        | 15.8                | /                        | /                           |
| 900    | 15.35        | 20.8                | /                        | /                           |
| 1000   | 15.22        | 29.15               | /                        | /                           |

7.3. Pressure effect
a. Variation of H2 pressure

In this case the temperature is fixed at T=1000°C and the partial pressure of the H2 gas is varied for the following values (PH2=0.01 Atm, 0.02A tm, 0.03 Atm) the results are the following. The polarization, activation, Ohmic resistance, and concentration-to-voltage curves at various pressures at 800 °C are shown in Figures 9, 10, 11, 12, and 13, respectively. In the event of a flooding failure, the pressure effect corresponds to the hydrogen and oxygen measured at the fuel cell inlet.
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b. Variation of O₂ pressure

In this case the temperature is fixed at T=1000 °C and the partial pressure of the O₂ gas is varied for the following values (PO₂=0.01 Atm, 0.02 Atm, 0.03 Atm) the results are the following. As shown in Figures 14, 15, 16, 17 and 18, as the pressure of oxygen increases, the activation loss decreases and the increasing current density decreases. In addition, increasing pressure reduces concentration loss and increases current density.

![Figure 13. Concentration losses characteristic for different pressures H₂ values](image1)

![Figure 14. Current-voltage characteristics for different pressures O₂ values](image2)

![Figure 15. Current-power characteristics for different pressures O₂ values](image3)

![Figure 16. Activation losses characteristics for different pressures O₂ values](image4)

![Figure 17. Ohmic losses characteristics for different pressures O₂ values](image5)

c. Variation of H₂O pressure

In this case the temperature is fixed at T=1000°C and the partial pressure of the H₂O gas is varied for the following values (PH₂O=0.1 Atm, 0.2 Atm, 0.3 Atm) the results are the following. As shown in Figures 19, 20, 21, 22, and 23. In general, increasing the inlet pressure increases the fuel cell voltage and power density.
as shown in Figure 19. In normal operation of the cell, this pressure is maintained at 1 atmosphere. During the flood condition, the pressure at the inlet will automatically increase, causing cell failure. Therefore, reliable operation of the cell requires efficient monitoring of the cell. Tables 2 to 5 show the power values according to the influence of the pressure for H2O, O2, and H2. From the table, we can see that the value of power rises with the increasing temperature, and the parameter affecting this increase is Ohmic loss. After changing the pressure, you can see that the ohm loss and activation loss, and the concentration remain constant. The result was interpreted after the illustration of all figures.

![Figure 18. Concentration losses characteristics for different pressures O2 values](image1)

![Figure 19. Current-voltage characteristics](image2)

![Figure 20. Current-power characteristics for different pressures H2O values](image3)

![Figure 21. Activation losses characteristics for different pressures H2O values](image4)

![Figure 22. Ohmic losses characteristics for different pressures H2O values](image5)

![Figure 23. Concentration losses characteristics for different pressures H2O values](image6)
Influences of electrolyte type and thickness on solid oxide fuel cell hybrid system

Two-dimensional computational fluid dynamics simulation of anode side flow field in anode-supported SOFC,

https://tel.archives-ouvertes.fr/tel-00608632.

As explained above for a summary or overview, the shaping of all the physically applicable properties in a fuel stack system will necessitates agile tools that can integrate multiple physical domains into the same model. Equation-based language and causal modeling capabilities enable physics models. This is very practical when you want to reuse a component in a variety of settings for a range of objectives.

8. CONCLUSION

In the context of this work, a macroscopic modelling tool has been implemented to the analogy between the electrical and thermal domains. This method constitutes a simple, scalable and efficient modelling tool. Once the component library has been completed, we will have a complete tool for studying the analogy between the electrical and thermal. Finally, we look at cogeneration systems using solid oxide fuel cells (SOFC).

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