



Article Numerical Modeling and Simulation of the Solid Oxide Cell Stacks and Metal Interconnect Oxidation with OpenFOAM

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Abstract: Solid oxide cells are capable of efficiently converting various chemical energy carriers to electricity and vice versa. The urgent challenge nowadays is the faster degradation rate compared with other fuel cell/electrolyzer technologies. To understand the degradation mechanisms, simulation of a solid oxide cell is helpful. Since most previous research developed models using commercial software, such as COMSOL and ANSYS Fluent, a gap for knowledge transfer is being gradually formed between academia and industry due to licensing issues. This paper introduces a multiphysics model, developed by a computational code, openFuelCell2. The code is implemented with an open-source library, OpenFOAM. It accounts for momentum transfer, mass transfer, electrochemical reactions and metal interconnect oxidation. The model can precisely predict I–V curves under different temperatures, fuel humidity and operation modes. Comparison between OpenFOAM and COMSOL simulations shows good agreement. The metal interconnect oxidation is modeled, which can predict the thickness of the oxide scale under different protective coatings. Simulations are conducted by assuming an ultra-thin film resistance on the rib surface. It is revealed that coatings fabricated by atmospheric plasma spraying can efficiently prevent metal interconnect oxidation, with a contribution of only 0.53 % to the total degradation rate.

Keywords: solid oxide cell; multiphysics modeling; OpenFOAM; openFuelCell2; metal interconnect oxidation

1. Introduction

The transition to a renewable energy system where solar and wind energy prevail leads to a challenge for the grid balance. Applying solid oxide cells (SOCs) as energy storage and conversion devices may be part of the solution to these problems. Their operation at high temperatures provides several advantages, such as high efficiencies, the possibility of sector coupling on the heat side, the capability of being reversibly operated, and the tolerance to various fuels. In order to increase the lifetime of SOCs, their degradation behavior should be studied, as it obstructs the long-term operation. Significant degradation mechanisms, such as Ni agglomeration and Cr poisoning, are related to the local distribution of the overpotential and chemical species. However, measuring local values for these during the operation of SOCs is not only a technical challenge but also a potentially dangerous task, as gas leakages, burns and electric shocks could happen. In contrast, simulation can be an efficient and safe method to visualize the local physical values. Therefore, a three-dimension model that can capture the physical processes in an SOC is necessary.

Although there have been a number of scientific findings due to simulation in the past decade [1–8], simulations are usually steady-state simulations, making it impossible to trace the degradation over time. It can be seen that many numerical investigations of SOCs were carried out with commercial software, for example, DETCHEM, ANSYS Fluent, COMSOL,



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Copyright: © 2023 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). etc. Mathematical models used to describe the multiphysical and multiscale transport processes in the SOCs have been incorporated into a 'black-box'. It is usually impossible to visualize the implementation of these models. In addition, the knowledge transfer may be limited within the research community due to the license issues. In this work, the authors employed a computational code, openFuelCell2 [9,10], which was implemented with an open-source library, OpenFOAM [11]. The code witnessed reliable performance in lowtemperature electrochemical applications, for example, the proton exchange membrane fuel cells [12–15]. This work aims to extends the capability of openFuelCell2 to hightemperature applications, i.e., SOFC and SOEC. The comparison between the simulation results with openFuelCell2 and COMSOL is conducted for the basic steady-state model. A degradation model to describe the performance degradation with the evolution of time due to metal interconnect (MIC) oxide scale growth is additionally proposed and implemented into openFuelCell2. The effects of different protective coatings on MIC oxidation are discussed. Numerical simulations are performed for the F10 SOC stack design [16] in Forschungszentrum Jülich GmbH (FZJ), the detailed description of which is given in the next section.

2. Geometry of the Model

It is helpful to introduce the general information of the F10 SOC stack design in FZJ first. The simplified geometry of the repeating unit of the SOC stack is shown at the left-hand side of Figure 1. The standard design is an fuel-electrode supported cell (FESC), where a thick support layer at the fuel side, made of Ni/8YSZ (8 mol% yttriastabilized zirconia), provides necessary mechanical strength. The repeating unit consists of roughly 1000 µm thick Ni-mesh as the distributor and contacting element at the fuel side, a 500 µm thick support layer Ni/8YSZ [17], a 7 µm thick fuel electrode Ni/8YSZ [17], a 10 μ m thick 8YSZ electrolyte [17], a 45 μ m thick air electrode La_{0.58}Sr_{0.4}Co_{0.2}Fe_{0.8}O_{3- δ} (LSCF) [17], and a 100 μ m thick contact layer LSCF (or LCC12, La_{0.97}Mn_{0.4}Co_{0.3}Cu_{0.3}O_{3-\delta}). A barrier layer (GDC, gadolinia-doped ceria) is added between the electrolyte and the air electrode to prevent interfacial reaction. Furthermore, there is a protective coating on the MIC to alleviate issues of metal oxidation and Cr poisoning. In FZJ, the protective coating on the MIC surface is usually made of $MnCo_{1.9}Fe_{0.1}O_4$ (MCF) applied by atmospheric plasma spraying (APS). Another widely used coating is made of MnO_x and is added onto MIC via wet powder spraying (WPS). The drawback of WPS, in contrast to APS, is that it can only generate a porous layer that cannot efficiently prevent metal oxidation and chromium poisoning. Consequently, stacks with APS coatings have better performance than stacks with WPS coatings [18]. The MIC is made by either Crofer 22 APU or intermediate temperature metal (ITM). For sealing the stack glass, ceramic sealants are used on all bonding surfaces.

The numerical simulation of the whole geometry usually means a rather large calculation effort. With the purpose of accelerating the computation while capturing the main features of the repeating unit of a real SOC stack, the following assumptions are made:

- 1. The barrier layer and protective coating are ignored in the geometry of the model. Their effects on the resistance are considered in the electronic conductivity of the air electrode and air-side contact layer.
- 2. The metal frame is ignored in the geometry of the model.

Therefore, the computational domain in this study refers to a representative threedimensional slice of one channel in a real SOC stack as shown on the right-hand side of Figure 1.



Figure 1. Simplification of the geometry of a standard F10 stack [16]. The left figure is a representive geometry of the active volume. The right figure is the cross-section geometry used in the simulations. The barrier layer and protective coating are not considered in the model. Not to scale.

3. Governing Equations of the Basic Model

The SOC stack is a complex electrochemical device that contains different physics processes, which are coupled with each other. This interplay strongly affects the performance of the SOC stacks. Therefore, suitable mathematical models should be implemented; otherwise, the results are questionable. Although it is recommended to involve as many details as possible, several plausible simplifications can be made to alleviate the difficulty in multiphysics modeling:

- 1. Properties of materials, such as permeability, porosity and electrical conductivity, are homogeneous and isotropic.
- 2. All fluid is in the laminar flow regime.
- 3. Heat transfer is ignored. In other words, isothermal assumption is applied in the model. The assumption holds well for stacks listed in this work, as the temperature difference is $10 \sim 15$ °C according to thermocouple measurements.
- 4. The quality of the inlet gas is perfect, meaning the air only has oxygen and nitrogen, and the fuel only has hydrogen and steam.

In this section, models of major physical processes, including electrochemical reaction, momentum transfer and mass transport, are present. A summary of the models used in this work and corresponding variables to be solved is shown in Table 1.

Tabi	le 1.	Summar	v of r	nain	model	s used	in thi	s work.
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Physical Phenomenon	Domain or Surface	Main Equation	Variables to Be Solved
Electrochemical reaction	Electrodes	Chang–Jaffe kinetic	Electronic and ionic potential
Momentum transfer	Channels	Navier-Stokes equations	Pressure and velocity
	Porous medium	Darcy–Forchheimer equation	Pressure and velocity
Mass transport	Channels and porous medium	Fick's law	Molar fractions of species
MIC oxidation	Rib surface	Parabolic law	Electronic potential difference due to the oxide scale

3.1. Electrochemical Reaction

Electrochemical reaction is a key part in modeling SOCs due to its direct relation to I–V curves and strong coupling with other physics. Commonly, it is simulated by the Butler–Volmer equation in Equation (1) [19]:

$$i = i_0 \left[\exp\left(\frac{\beta_a n_e F}{RT} \eta\right) - \exp\left(\frac{-\beta_c n_e F}{RT} \eta\right) \right]$$
(1)

where *i* is the current density, i_0 is the exchange current density, β_a (β_c) is the anodic (cathodic) charge transfer coefficient, n_e is the number of transferred electrons, F is the

Faraday constant, R is the gas constant, *T* is the temperature, and η is the overpotential. In spite of the popularity of Equation (1), Mogensen et al. [20] raised several arguments against using Equation (1) for SOCs, such as the reaction rate probably not being limited by the charge transfer. Similarly, Adler et al. [21] found that, except for charge transfer, the surface diffusion of chemical species and dissociative chemisorption were also important for reaction rates. As illustrated by Maria et al. [22], it could be that the high operation temperature of SOC makes adsorption, diffusion and other processes more rate-limiting than the charge transfer. Additionally, in most cases, linear I–V curves instead of exponentially changed I–V curves are observed from the measurement in FZJ.

In light of the above reasons, in this work, a linear expression of electrochemical kinetics is used, shown in Equations (2) and (3) [22,23]:

$$i_{\rm v,fuel} = \frac{\eta_{\rm fuel}}{R_{\rm fuel} \exp(\frac{-E_{\rm a,fuel}}{RT}) a_{\rm H_2}^{0.1} a_{\rm H_2}^{-0.33}}$$
(2)

$$i_{\rm v,air} = \frac{\eta_{\rm air}}{R_{\rm air} \exp(\frac{-E_{\rm a,air}}{RT})a_{\rm O_2}^{-0.25}}$$
(3)

where i_v is the volumetric reaction current density, η is the overpotential, R is a pre-factor, E_a is the energy barrier, and a is the activity of the reactant. The overpotential can be calculated by [24]

$$\eta_{\rm air} = \phi_{\rm ele, \ air} - \phi_{\rm io, \ air} - \phi_{\rm eq, \ air} \tag{4}$$

$$\eta_{\text{fuel}} = \phi_{\text{ele, fuel}} - \phi_{\text{io, fuel}} - \phi_{\text{eq, fuel}} \tag{5}$$

where ϕ_{io} is the ionic potential, ϕ_{ele} is the electromotive potential and ϕ_{eq} is the equilibrium potential defined as [22]

$$\phi_{\text{eq, air}} = \frac{\text{RT}}{4\text{F}} \ln(Y_{\text{O}_2}) \tag{6}$$

$$\phi_{\rm eq, \ fuel} = -\frac{\Delta G_{\rm water \ splitting}}{2F} - \frac{RT}{2F} \ln\left(\frac{Y_{\rm H_2}}{Y_{\rm H_2O}}\right) \tag{7}$$

where Y_{O_2} , Y_{H_2} and Y_{H_2O} are molar fractions of oxygen, hydrogen and water, respectively. $\Delta G_{\text{water splitting}}$ is the Gibbs free energy change of water splitting.

Two types of current exist in SOC: ionic current i_{io} and electronic current i_{ele} . The current and potential obey Ohm's law:

$$i_{\rm io} = -\sigma^{\rm io} \nabla \phi_{\rm io} \tag{8}$$

$$i_{\rm ele} = -\sigma^{\rm ele} \nabla \phi_{\rm ele} \tag{9}$$

where σ^{io} and σ^{ele} are the ionic conductivity and electronic conductivity that can be found in Table 2. In a porous and composite medium, the conductivity needs to be corrected by considering the effect of the solid volume fraction χ of and tortuosity τ of different materials. If the number of different material properties in the medium is *n*, the effective conductivity is [25]

$$\sigma^{\text{ele,eff}} = \sum_{i=1}^{n} \sigma_i^{\text{ele}} \frac{\chi_i}{\tau_i} \tag{10}$$

$$\sigma^{\text{io,eff}} = \sum_{i=1}^{n} \sigma_i^{\text{io}} \frac{\chi_i}{\tau_i} \tag{11}$$

The conservation of current is guaranteed by

$$\nabla i_{\rm io} = S_{\rm current} \tag{12}$$

$$\nabla i_{\rm ele} = -S_{\rm current} \tag{13}$$

where S_{current} is a current source defined as the following:

$$S_{\text{current}} = \begin{cases} i_{\text{v}} & \text{in electrochemical active regions} \\ 0 & \text{in electrochemical inactive regions} \end{cases}$$
(14)

and i_v is given in Equations (2) and (3).

The electronic and ionic conductivity is present in Table 2. Other parameters, such as the porosity and volume factions of the porous medium, are given in Table A1.

Table 2. Conductivity of material.

Component	$\sigma^{ m ele}$ (S/m) [22,26,27]	σ^{io} (S/m) [22]
Ni 8YSZ LSCF LCC12 ¹ Crofer 22 APU	$3.27 \times 10^{6} - 1065.3T$ 22,591-1.6 × 10 ⁶ exp(-6024/T) 22,591-1.6 × 10 ⁶ exp(-6024/T) 9.09 × 10 ⁵	$6.25 \times 10^4 \exp(-10,300/T)$ $5.5 \times 10^9 \exp(-9050/R/T)$ $5.5 \times 10^9 \exp(-9050/R/T)$

¹ Assume LCC12 has the same electronic and ionic conductivity as LSCF.

3.2. Momentum Transfer

Momentum transfer of fluid happens in gas channels and porous medium. In gas channels, it is a common practice to apply Navier–Stokes (NS) equations to describe the momentum transfer as shown in Equation (15) [28]:

$$\frac{\partial(\rho_{\rm f}\vec{U})}{\partial t} + \rho_{\rm f}(\vec{U}\cdot\nabla)\vec{U} = -\nabla(P\vec{I}) + \rho_{\rm f}\vec{g} + \nabla\left[\mu\left(\nabla\vec{U} + (\nabla\vec{U})^{\rm T}\right) - \frac{2\mu}{3}(\nabla\cdot\vec{U})\vec{I}\right]$$
(15)

where ρ_f is the density of the fluid, *P* is the pressure, \vec{U} is the velocity, μ is the viscosity, and \vec{I} is the identity matrix.

In a porous medium, momentum transfer can be simulated by several models. The easiest one is Darcy's law, considering that the velocity of fluid should linearly respond to the pressure gradient:

$$\vec{U} = -\frac{\kappa}{\mu} \nabla P \tag{16}$$

where κ is the permeability of the porous medium. In spite of its conciseness, it should be kept in mind that it is an empirical relation that originated from the experimental data of water passing through sands at low velocity [29]. Therefore, there are some limitations to Equation (16). For example, Darcy's law neglects viscous forces. Assuming that Darcy's law is implemented in a porous medium and the NS equation is used in gas channels, two calculation domains need to be defined because Equation (15) has second-order differential terms, while Equation (16) misses it. This inherent conflict leads to trouble in setting boundary conditions at the interface between channels and the porous medium [30].

Provided by the drawbacks in Darcy's law, it is recommended to use the Darcy's modified NS equation, which is able to be implemented in both gas channels and porous media, as given in Equation (17) [31,32]:

$$\frac{\partial(\rho_{\rm f}\vec{U}/\varepsilon)}{\partial t} + \rho_{\rm f}(\frac{\vec{U}}{\varepsilon}\cdot\nabla)\frac{\vec{U}}{\varepsilon} = -\nabla(P\vec{I}) + \rho_{\rm f}\vec{g} + \nabla\left[\frac{\mu}{\varepsilon}\left(\nabla\vec{U} + (\nabla\vec{U})^{\rm T}\right) - \frac{2\mu}{3\varepsilon}(\nabla\cdot\vec{U})\vec{I}\right] - \frac{\mu}{\kappa}\vec{U}$$
(17)

where ε is the porosity of the porous medium. In gas channels where permeability approaches infinity and porosity becomes one, Equation (17) can automatically become

Equation (15), which avoids setting another boundary condition at the interface between the gas channels and porous medium. The continuity equation is

$$\frac{\partial(\epsilon\rho_{\rm f})}{\partial t} + \nabla(\rho_{\rm f}\vec{U}) = S_{\rm mass} \tag{18}$$

where S_{mass} is the mass source due to reactions.

The data of porosity and permeability are given in Table A1.

3.3. Mass Transport

Mass transport is governed by Equation (19):

$$\frac{\partial(cY_i)}{\partial t} + \nabla(c\vec{U}Y_i) + \nabla\vec{N}_i = S_{\text{mole},i}$$
(19)

where *c* is the molar density, Y_i is the molar fraction, N_i is the molar diffusion flux of species *i* and $S_{\text{mole},i}$ is the source term of moles of chemical specie *i* due to electrochemical reactions, and it is described by the following Equation (20):

$$S_{\text{mole},i} = \begin{cases} \frac{i_v}{n_e F} & \text{in electrochemical active regions} \\ 0 & \text{in electrochemical inactive regions} \end{cases}$$
(20)

where i_v is the reaction current density, and n_e is the number of electrons transferred in electrochemical reactions. To solve Equation (19), N_i needs to be calculated. Several diffusion models can be chosen. For instance, Fick's law can be used, which is the most popular model due to its simplicity, and it has been successfully implemented in SOC numerical modeling [33,34]. The mathematical description is shown in Equation (21)

$$\vec{N}_i = -D^{\rm B}_{ii} \nabla(cY_i) \tag{21}$$

where D_{ij}^{B} is the binary diffusion coefficient for species *i* and *j*. For a multi-component system, it is more plausible to apply the Stefan–Maxwell equation to describe the transport phenomena, which is the extension of Fick's law. The Stefan–Maxwell equation is given in Equation (22)

$$\sum_{i=1,j\neq i}^{n} \frac{Y_j \dot{N_i} - Y_i \dot{N_j}}{D_{ij}^{\text{B,eff}}} = -\frac{P}{RT} \nabla Y_i - \frac{Y_i}{RT} \nabla P$$
(22)

where $D_{ij}^{\text{B,eff}}$ is the effective binary diffusion coefficient. It should be noted that both Fick's law and the Stefan–Maxwell equation are factually applied for the open space instead of the porous medium [35]. To simulate the behavior in the porous medium, effective diffusion coefficients are introduced, which are related to the porosity and tortuosity of the porous medium. In the model, Equation (23) is used to define the effective diffusion coefficient [25]:

$$D_{ij}^{\text{B,eff}} = \frac{\varepsilon_k}{\tau_k} D_{ij}^{\text{B}}$$
(23)

where ε is the porosity and τ is the tortuosity. *k* is the component, such as the fuel electrode and the air electrode. The tortuosity is calculated through following relationship:

$$\tau_k = (1 - \varepsilon_k)^{\frac{2 - D_s}{2 - D_s}} \tag{24}$$

where D_s is usually chosen as 1.2 for SOC systems [36]. The binary diffusion coefficient D_{ij}^{B} is evaluated by the Chapman–Enskog theory [37]:

$$D_{ij}^{\rm B} = D_{ji}^{\rm B} = \frac{3.16 \times 10^{-8} T^{1.75}}{P(v_i^{1/3} + v_i^{1/3})^2} (\frac{1}{M_i} + \frac{1}{M_j})^{1/2}$$
(25)

where *v* is the diffusion volume and *M* is the molar mass.

Fick's law and the Stefan–Maxwell model function well regarding mass transfer in a porous medium, but they will fail if the interaction between the species and solids cannot be ignored. In this case, Knudsen diffusion should be taken into consideration in addition to molecular diffusion. One model that includes Knudsen diffusion is the dusty gas model as given in Equation (26):

$$\sum_{j=1,j\neq i}^{n} \frac{Y_j \vec{N}_i - Y_i \vec{N}_j}{D_{ij}^{\text{B,eff}}} + \frac{N_i}{D_i^{\text{K,eff}}} = -\frac{P}{RT} \nabla Y_i - \frac{Y_i}{RT} \left(1 + \frac{\kappa P}{\mu D_i^{\text{K,eff}}}\right) \nabla P$$
(26)

where κ is the permeability, $D_i^{\text{K,eff}}$ is the effective Knudsen diffusion coefficient for the species *i*, and it can be calculated through Equation (27) [25]:

$$D_i^{\text{K,eff}} = \frac{\varepsilon_k}{\tau_k} D_i^{\text{K}} = \frac{d_p \varepsilon_k}{3\tau_k} \sqrt{\frac{8\text{RT}}{\pi M_i}}$$
(27)

where d_p is the diameter of the pore size.

In brief, three models, Fick's law, the Stefan–Maxwell model and the dusty gas model, can be chosen to simulate the mass transport. In this paper, Fick's law is chosen after several considerations. First, it is easiest to achieve numerically, as it has a simpler form than the others. Secondly, Fick's law actually can approximate the other two models well, particularly when the number of components in the system is less than four [38]. Thirdly, to improve the accuracy, Fick's law can be modified to include the Knudsen diffusion [22,39,40] by employing the Bosanquet formula as shown in Equation (28):

$$\frac{1}{D_{ij}^{\text{eff}}} = \frac{1}{D_{ij}^{\text{B,eff}}} + \frac{1}{D_{ij}^{\text{K,eff}}}$$
(28)

where $D_{ij}^{K,eff}$ is a mean Knudsen diffusion coefficient that is obtained by replacing M_i with M_{ij} in Equation (27), and M_{ij} is defined as

$$\frac{1}{M_{ij}} = \left(\frac{1}{M_i} + \frac{1}{M_j}\right) / 2 \tag{29}$$

Given the above, Fick's law (Equations (21), (27)–(29)) is applied in the model.

4. Governing Equations of the Degradation Model of MIC Oxidation

The typical material used for interconnects of SOC stacks are metals (e.g., Crofer 22 APU). It provides strong mechanical support and high electronic conductivity. However, it always suffers from oxidation [41], leading to the formation of the oxide scale. During the growth of the oxide scale, the resistance at the interface increases, which serves as one part of the total degradation. The model in this work only considers the MIC oxidation at the air side. On the fuel side, MIC oxidation is believed to have little influence on the performance of SOCs. The Ni mesh results in good contact with the fuel support layer, providing spot welding is used to connect MIC and Ni mesh, which provides a strong and reliable contact.

4.1. Thickness of the Oxide Scale on a Clean Surface

To simulate the growth of the oxide scale, it is a common practice to employ the parabolic law [42,43] given in Equation (30)

$$d^2 = \Xi \cdot t \tag{30}$$

where *d* is the thickness of the oxide scale, *t* is time, and Ξ is the parabolic constant. The "constant" means that its value does not depend on the thickness of the oxide scale.

Before applying Equation (30), it is necessary to prove that, for SOCs, the growth of the Cr_2O_3 fulfills the parabolic format. It was pointed out that the growth of Cr_2O_3 obeys the parabolic law when the temperature is above 623 K [44], which is lower than the operating temperature of SOCs. There are works in the literature [42,45], where the parabolic law was used to ensure good curve fitting. In brief, it is reasonable to assume that Equation (30) is satisfied during the whole operation of SOCs.

The next step is to obtain the parabolic constant Ξ . The typical method is Wagner's theory, which relates ions' tracer diffusion coefficients to the parabolic constant. According to Wagner's theory, the parabolic constant can be written as follows:

$$\Xi = \int_{\substack{Y_{O_2}^{\text{oxide}-\text{gas}}\\ Q_2}} \int \left[\left(Z_{Cr^{3+}} / Z_{O^{2-}} \right) D_{Cr^{3+}}^{\text{eff}} + D_{O^{2-}}^{\text{eff}} \right] d \ln Y_{O_2}$$
(31)

where Z is the absolute value of the valency, D^{eff} is the effective tracer diffusion coefficients, which should include the bulk diffusion and grain-boundary diffusion [46,47], the superscript oxide–gas and metal–oxide refer to the interface of oxide–gas and the interface of metal–oxide, respectively. Although Equation (31) is widely used for the simulation of alloy oxidation, it is not suitable for the model of SOCs due to the following reasons:

- In order to obtain tracer diffusion coefficients, the concentration of defects (e.g., V^{'''}_{Cr}) that serve as the path for the ions needs to be known. However, there remain debates [47–49] as to which defects dominate inside Cr₂O₃.
- 2. No literature about tracer diffusion coefficients inside the oxide scale on Crofer 22 APU can be found. If the data of a different alloy are used (e.g., Ni-Cr alloy [49]), the parabolic constant obtained from Equation (31) will possibly deviate from the experimental findings.

In light of the difficulties in applying Equation (31), an empirical relation is used and introduced in the following section, which was proposed by FZJ [42].

4.2. Thickness of the Oxide Scale on the Surface Covered by a Porous Coating

It should be noted that Equation (30) only works for a clean surface without coatings. However, the MIC is usually protected by WPS MnO_x or APS MCF. Considering the thickness of oxide scale is proportional to the mole of oxygen atoms adsorbed on the MIC, the following equation is obtained:

$$d_{\text{poro}} \propto n_{\text{O, poro}} = \iint \dot{n}_{\text{O, poro}} dA_{\text{poro}} dt$$
 (32)

$$= \iint \dot{n}_{\rm O,\ poro} d(\varepsilon_{\rm coating} A) dt \tag{33}$$

$$=\varepsilon_{\text{coating}} \iint \dot{n}_{\text{O, poro}} dAdt \tag{34}$$

$$=\varepsilon_{\text{coating}} \iint \dot{n}_{\text{O}} dA dt \tag{35}$$

$$=\varepsilon_{\text{coating}}n_{\text{O}}$$
 (36)

which means

$$d_{\text{poro}} = \varepsilon_{\text{coating}} d \tag{37}$$

where d_{poro} is the thickness of the oxide scale under porous coatings (e.g., the protective coating), $n_{\text{O}, \text{ poro}}$ is the mole of oxygen atoms adsorbed on the MIC surface covered by coatings, $\dot{n}_{\text{O}, \text{ poro}}$ is the adsorption rate of oxygen atoms per area over the MIC surface covered by coatings, A_{poro} is the free surface area under a porous coating that can be reached by oxygen gas, A is the total surface area of MIC, $\varepsilon_{\text{coating}}$ is the porosity of the porous coatings, \dot{n}_{O} is the adsorption rate of oxygen atoms over the MIC surface without coatings,

and n_0 is the mole of oxygen atoms adsorbed on the MIC surface without coatings. Two underlying assumptions are used for derivation of Equation (37). First, the homogenization assumption is applied in Equation (33). It is usually hard for CFD models to capture micro-structure values. Secondly, it is assumed in Equation (35) that the adsorption rate over the MIC surface is not affected by the coating. The most evident difference of physical phenomena between the cases with and without coating is the gas transport of oxygen, it means the oxygen partial pressure affects insignificantly on the adsorption rate. If the porosity of the coating is higher than 0.3, this assumption is reasonable, as it was found that oxygen could freely reach the surface [45]. At present, there is no direct evidence to support the second assumption regarding the application of a dense coating. However, it is thought that the rate-limiting step during metal oxidation is usually the diffusion of ions instead of gas transport over the metal surface [43]. Therefore, the second assumption is generally valid for the growth of the oxide scale under a dense coating.

In brief, Equations (30) and (37) are used to simulate the growth of the oxide scale on MIC in SOCs.

4.3. Voltage Drop Due to MIC Oxidation

To simulate the voltage drop due to MIC oxidation, an internal boundary condition is needed. In the model, this boundary is called the ultra-thin film resistance. The effect of the oxide scale is represented by an internal surface boundary condition at the rib, as shown in Figure 2. The mesh of the rib surface is split into two sub-surfaces, and Ohm's law is applied to each of them as follows:

$$-\vec{n}_{\rm up} \cdot \vec{i}_{\rm ele} = -\frac{\sigma_{\rm oxide \ scale}}{d_{\rm poro}} \left(\phi_{\rm ele, \ up} - \phi_{\rm ele, \ low}\right) \tag{38}$$

$$-\vec{n}_{\text{low}} \cdot \vec{i}_{\text{ele}} = -\frac{\sigma_{\text{oxide scale}}}{d_{\text{poro}}} \left(\phi_{\text{ele, low}} - \phi_{\text{ele, up}} \right)$$
(39)

where \vec{n} is the unit surface normal vector, \vec{t}_{ele} is the electronic current density, $\sigma_{oxide \ scale}$ is the conductivity of the oxide scale, and ϕ_{ele} is the electronic potential. The subscripts "up" and "low" refer to the upper surface (with higher electronic potential) and lower surface (with lower electronic potential), respectively. The directions of the above vectors can be found in Figure 2. The difference between $\phi_{ele, up}$ and $\phi_{ele, low}$ arises from the resistance of the growing oxide scale, which can be obtained through

$$\Delta \phi_{\rm ele} = \phi_{\rm ele, up} - \phi_{\rm ele, low} = |\vec{i}_{\rm ele} \cdot \frac{d_{\rm poro}}{\sigma_{\rm oxide \ scale}}| \tag{40}$$

where $\Delta \phi_{ele}$ is the voltage degradation due to the MIC oxidation.



Figure 2. Schematic diagram of the ultra-thin film resistance model. To more clearly illustrate the model, only the contact layer and MIC are shown in the figure.

4.4. Parameters

According to the experimental data [42], the parabolic constant of Crofer 22 APU (700 \sim 900 °C) is summarized in Equation (41):

$$\Xi_{\rm Crofer} = 5.5075 \times 10^{-9} \exp\left(\frac{-1.9992 \times 10^5}{RT}\right)$$
(41)

It was found that under 800 °C, the growth rate of the oxide scale on ITM was half of that on Crofer 22 APU [50]. Assuming this is valid for the temperature range of 700 \sim 900 °C, the parabolic constant for ITM is,

$$\Xi_{\rm ITM} = 0.25 \times \Xi_{\rm Crofer} = 1.3769 \times 10^{-9} \exp\left(\frac{-1.9992 \times 10^5}{RT}\right)$$
(42)

The conductivity of the oxide scale on Crofer 22 APU, $\sigma_{\text{oxide scale, Crofer}}$, is summarized as follows with the help of experimental data [43]:

$$\sigma_{\text{oxide scale, Crofer}} = \frac{1}{T} \cdot 10^{-3228/T + 6.7661}$$
(43)

The available conductivity data of the oxide scale on ITM are very limited. It was found that the increasing rate of ASR on ITM during oxidation was about 3.5 times as much as that on Crofer 22 APU [51], even if the thickness was half. The explanation was that if the coating is not a spinel phase, the oxide scale generated on ITM is porous, while that on Crofer 22 APU is quite dense [51]. ITM was usually combined with MnO_x coating (not a spinel phase) in FZJ SOC stacks design, it is assumed that the conductivity of the oxide scale on ITM, $\sigma_{\text{oxide scale, ITM}}$, is

$$\sigma_{\text{oxide scale, ITM}} = \frac{1}{7} \sigma_{\text{oxide scale, Crofer}} = \frac{1}{7T} \cdot 10^{-3228/T + 6.7661}$$
(44)

The porosity of WPS coating is 0.45 [52], while the porosity of APS coating is 0.03 [53].

5. Numerical Procedure

The mesh in the model is presented in Figure 3, where, in total, 74,800 elements are made. Mesh refinement is performed at the interface to capture electrochemical reactions. In other areas, mesh is coarser.



Figure 3. The mesh of the model.

The basic model is steady state, while MIC oxidation is a time-dependent model. Discretization of the time derivative is achieved by the backward Euler method. To increase the convergence of the time-dependent simulation, the basic model is solved first. The time-dependent simulation of MIC oxidation is carried out with the initial values being results of the basic model. The time step for the time-dependent simulation is 10 h. The discretization of the gradient terms and divergence terms is performed with the Gauss linear method; the discretization of all Laplacian terms is Gauss linear, except the conservation of the current, which is discretized by the Gauss harmonic. In addition, a SIMPLE solution algorithm [54] is employed to solve the Navier–Stokes equation. The coupling between each physics is summarized in Figure 4, where the arrows mean the passed variables to the other physics. For example, to solve the governing equation of the mass transport, pressure and velocity

The solver and discretization used in COMSOL and OpenFOAM are different. The former employs a coupled solution manner, which solves all variables simultaneously. During time intervals, a direct solver, which solves the inverse of the matrix, is used. In contrast, the segregated solver is applied in OpenFOAM that subdivides the numerical calculation into several steps. For each iteration, the iterative solver is applied. Moreover, COMSOL implements the finite element method, while OpenFOAM utilizes the finite volume method. Despite the above differences in the numerical calculation, it is expected that both software will give similar results when it comes to the same model with the same boundary conditions and parameters. All the simulations are conducted on a single core of a computer with CPU i7-9700K and 32 GB RAM.

fields are needed, which are taken from the solution of the momentum transfer.



Figure 4. A schematic diagram showing how each of the physics is coupled with each other.

6. Experiment

SOC stacks were operated in a constant current mode with the gases supplied in a counter-flow regime. The characterization and operation of SOC stacks were carried out in an oven test rig. Inside the oven, there were electric heating elements to maintain the temperature of stacks through heat radiation. Deionized water with a resistivity of 10 M Ω cm [55] was supplied to the electrically heated steam generator. Hydrogen with a minimal purity of 99.9% was used for the inlet fuel. Dehumidified air with a dew point below -40 °C was sent as the inlet air. The mass flow of inlet air and inlet fuel was controlled by the mass flow controllers (MFCs). Mica sealants between the test bench interface and the stack in combination with a compressive force of 1 kN were applied to prevent gas leakage. Table 3 shows details of the MFCs, operation conditions and stack

designs of several stacks used for the validation in this work. The temperature values in Table 3 were measured by thermocouples that were placed roughly 10 mm deep into the intermediate interconnector (details of the locations can be found in reference [56]).

Table 3. Experimental details of SOC stacks.

		F1002-97	F1002-95	F1004-67	F1004-115
Instrument	MFC	Bronkhorst, F-201CV	Bronkhorst, F-201CV	Bronkhorst, F-201CV	Brooks, 0254
Stack design	Contact layer MIC Protective coating	LCC12 ITM WPS MnO _x	LCC12 Crofer 22 APU WPS MnO _x	LSCF Crofer 22 APU APS MCF	LSCF Crofer 22 APU APS MCF
Operation conditions	Temperature Fuel mass flow Molar ratio in fuel Air mass flow Molar ratio in air Current density Time	$\begin{array}{c} 720\ ^{\circ}\text{C}\\ 1.18\times10^{-7}\ \text{kg/s}\\ \text{H}_2/\text{H}_2\text{O}=79/21\\ 1.9\times10^{-6}\ \text{kg/s}\\ \text{O}_2/\text{N}_2=21/79\\ 0.5\ \text{A/cm}^2\\ 100\ \text{kh} \end{array}$	$\begin{array}{c} 710\ ^{\circ}\text{C}\\ 1.15\times10^{-7}\ \text{kg/s}\\ \text{H}_2/\text{H}_2\text{O}=80/20\\ 1.18\times10^{-6}\ \text{kg/s}\\ \text{O}_2/\text{N}_2=21/79\\ 0.5\ \text{A/cm}^2\\ 17\ \text{kh} \end{array}$	$\begin{array}{c} 730\ ^{\circ}\text{C}\\ 1.15\times10^{-7}\ \text{kg/s}\\ \text{H}_2/\text{H}_2\text{O}=80/20\\ 1.18\times10^{-6}\ \text{kg/s}\\ \text{O}_2/\text{N}_2=21/79\\ 0.5\ \text{A/cm}^2\\ 25\ \text{kh} \end{array}$	
Conditions of I-V curves characterization	Temperature Fuel mass flow Molar ratio in fuel Air mass flow Molar ratio in air Current density	$\begin{array}{c} 700{\sim}800\ ^{\circ}\text{C} \\ 1.52\times10^{-7}\ kg/s \\ H_2/H_2O = 88/12 \\ 2.38\times10^{-6}\ kg/s \\ O_2/N_2 = 21/79 \\ 0{\sim}0.8\ \text{A/cm}^2 \end{array}$		$\begin{array}{c} 700{\sim}800\ ^{\circ}\text{C} \\ 1.85 \times 10^{-7}\ \text{kg/s} \\ \text{H}_2/\text{H}_2\text{O} = 80/20 \\ 1.79 \times 10^{-6}\ \text{kg/s} \\ \text{O}_2/\text{N}_2 = 21/79 \\ 0{\sim}0.8\ \text{A/cm}^2 \end{array}$	$\begin{array}{c} 700{\sim}800\ ^{\circ}\text{C} \\ 5.53 \times 10^{-7}\ \text{kg/s} \\ \text{H}_2/\text{H}_2\text{O} = 50/50 \\ 2.83 \times 10^{-6}\ \text{kg/s} \\ \text{O}_2/\text{N}_2 = 21/79 \\ -0.5{\sim}0.5\ \text{A/cm}^2 \end{array}$

F1004-115 consists of a 400 μ m thick support layer, 10 μ m thick fuel electrode and a 6 μ m thick electrolyte. The geometry of the rest of the parts of F1004-115 and other stacks can be found in Figure 1.

7. Validation

7.1. Validation of I–V Curves of the Basic Model

The validation of the OpenFOAM simulation has two parts: (1) The first shows a comparison between the OpenFOAM simulation and the COMSOL simulation. The same mesh, geometry, model, boundary conditions and parameters are used in both simulations. Although OpenFOAM and COMSOL use different discretization methods and solvers, their results are expected to be close. (2) The second shows a comparison between the OpenFOAM simulation and the experimental results. The comparison is carried out under different operation modes (e.g., fuel cell and electrolysis modes) and various operation conditions (e.g., different operation temperature and humidity of inlet fuel).

The validation by I–V curves is shown in Figure 5. Figure 5a,b present the I–V curves only in the fuel cell mode. With varied humidity and temperature, very good matches between the OpenFOAM simulation and the experimental data can be found, where the maximum voltage deviation is only 4.5%. Figure 5c presents the I–V curves for reversible SOCs. The negative current density means the electrolysis operation, while the positive current density indicates the fuel cell operation. There is good agreements between OpenFOAM and the experimental measurement providing the maximum voltage deviation is just 4.1%. The OpenFOAM simulation closely approximates the COMSOL simulation in all cases shown in Figure 5, which proves an additional validity of the numerical model in OpenFOAM. The deviations between the simulated and experimentally measured I–V curves could be due to the fluctuating temperature (5~10 °C) during the I–V curve characterizations.



(c) I–V curves of F1004-115

Figure 5. Validations of I–V curves for (**a**) F1002-97, 12% humidified fuel, SOFC, (**b**) F1004-67, 20% humidified fuel, SOFC and (**c**) F1004-115, 50% humidified fuel, SOFC and SOEC.

7.2. Validation of MIC Degradation Model

The core equation of the MIC oxidation model is Equation (37), which describes the thickness growth of the oxide scale under a porous coating. In the following, examples are given to demonstrate the validity of Equation (37).

7.2.1. Result from Persson: Crofer 22 APU with and without (La,Sr)MnO₃ (LSM) Coating

Persson [45] conducted a series of oxidation tests on Crofer 22 APU. The experimental data were taken from thermogravimetric analysis (TGA), which made it possible to track the growth of the oxide scale during operation. Assuming that Equation (37) provides a good approximation of the experimental results, we would expect the ratio of the thickness of the oxide scale with coating to the thickness of the oxide scale without coating to be close to the porosity of the coating, which is 0.3. According to the results in Figure 6, after 1000 hours of operation, the ratio is close to 0.3, which confirms the validity of the model. However, at the beginning of the operation, the ratio is higher than 0.3. This could result from the sintering process of the coating, as Persson used spray guns to add coatings on the MIC at room temperature and then started the oxidation tests directly [45]. However, in general, it can be claimed that the model is capable of predicting an accurate thickness of the oxide scale.

7.2.2. Result from FZJ: F1002-95 and F1002-97

Two stacks designed by FZJ are used for validation, F1002-97 and F1002-95. The former used ITM as the MIC and WPS MnO_x as the protective coating, while the latter employed Crofer 22 APU as the MIC and WPS MnO_x as the protective coating. Details of the stacks can be found in Table 3.



Figure 6. Validation of MIC oxidation model. The experimental data are taken from TGA results of Persson [45]. Crofer 22 APU was used as the MIC. One stack was operated without coatings, while another stack was coated with 15 µm LSM, whose porosity was measured as 0.3 after 4000 h of operation [45].

Unlike the TGA data in Figure 6, the validation data in this section are manually obtained from SEM figures. It should be kept in mind that the model predicted an average thickness. As shown in Figure 7 where average thickness is compared, good agreement is observed, although there exist slight deviations between the simulation and the experimentally measured data. The deviations can be due to the limited choices of parameters (e.g., the porosity of the protective coating and properties of ITM) and the errors when processing data from SEM images.



Figure 7. Validations of MIC oxidation model. The experimental data (21 data points for F1002-97 and 29 data points for F1002-95) are obtained from a scanning electron microscope (SEM) image of FZJ SOC stacks after they were shut down [57,58].

8. Results and Discussions

8.1. Spatial Distribution of Mole Fraction and Overpotential

Distribution of mole fraction and overpotential can affect the local degradation. Here, only analysis of the SOEC is given.

The spatial distribution of the mole fraction and overpotential are shown in Figure 8 and Figure 9, respectively. The simulation works for the electrolysis operation of F1004-115 under -0.5 A/cm^2 , fuel inlet composition $H_2/H_2O = 50:50$ and air inlet composition $O_2/N_2 = 21:79$. In the simulation, the flow field is the counter flow, where fuel flows toward the x-coordinate, while air flows against the x-coordinate.





(c) Mole fraction of oxygen at AEL/ELEC (OpenFOAM)



(d) Mole fraction of oxygen at AEL/ELEC (COMSOL)

Figure 8. Spatial distribution of mole fractions in the x-y plane of the fuel electrode and the air electrode. The simulation is the electrolysis operation of F1004-115, where fuel flows toward the x-coordinate direction and air flows against the x-coordinate. The geometry is scaled by the transformation vector (x, y, z) = (1, 20, 1).

Figure 8 shows the mole fraction distribution in the electrolysis mode of F1004-115, at the interface between the fuel electrode and the electrolyte (FEL/ELEC) and the interface between the air electrode and the electrolyte (AEL/ELEC). From Figure 8a, it is found that along the x-coordinate, which is the fuel flow direction, Y_{H_2} gradually increases due to the electrochemical reaction. Similarly, as indicated by Figure 8c, Y_{O_2} increases along the flow direction (against the x-coordinate). In addition, it should be noted that O_2 tends to segregate at the rib zones. As a comparison, on the fuel side, H_2 is more uniformly distributed. The main reason behind this phenomenon is that a gas channel is used for air distribution, while the Ni mesh is used for sending the fuel. Unlike the Ni mesh, the air channel does not directly contact the rib zones. Hence, O_2 transportation at the rib zones mainly relies on the diffusion process. The diffusion coefficient of O_2 is not high, as it is one order of magnitude lower than that of hydrogen and water. Consequently, the lack of enough of a driving force of transportation leads to O_2 segregation at the rib zones. When it comes to the mole fraction, the OpenFOAM simulation is close to the COMSOL simulation.

There is a very slight difference between Figure 8a,b, which can be caused by different solvers applied in OpenFOAM and COMSOL.



(c) Activation overpotential of the air electrode (OpenFOAM)

(d) Activation overpotential of the air electrode (COMSOL)

Figure 9. Spatial distribution of overpotential on the outer surface of the fuel electrode and the air electrode in the x-z plane (see geometry in Figure 1). The simulation is the electrolysis operation of F1004-115, where fuel flows toward the x-coordinate direction and air flows against the x-coordinate. The geometry is scaled by the transformation vector (x, y, z) = (0.05, 1, 60).

Figure 9 shows the overpotential distribution in the electrolysis mode of F1004-115, where η_{fuel} is negative and η_{air} is positive. According to Figure 9a,b, both overpotentials η_{fuel} and η_{air} reach the maximum absolute values close to the electrolyte as expected because the overpotential is the driving force for electrochemical reactions, which majorly happen near the electrolyte interface. Compared to the COMSOL simulation (right column), the evolution tendency is the same, while the values are slightly different.

8.2. Voltage Degradation Due to MIC Oxidation in F1002-97

The details of the stack F1002-97 can be found in Table 3. Figure 10 shows stack voltage evolution predicted by simulation at four time points. The inserted figures show the distributions of the electronic potential in the contact layer and the MIC. At time = 0 h, when no oxide scale exists, the electronic potential is uniformly distributed because the electronic conductivity of materials is high. With time going on, the oxide scale gradually grows, leading to degradation. It can be seen that the ultra-thin film resistance leads to a sudden potential drop across the rib in the inserted figures. Figure 11 gives the evolution of the voltage of the whole lifetime and the share of degradation rate only due to MIC oxidation. Because of the unstable water supply, the voltage measured in experiments fluctuate all the time. With the help of linear fitting, the degradation rate is determined to be 7.3 mV/kh before 40 kh and 1.6 mV/kh after 40 kh. Figure 11a indicates that, overall, the MIC oxidation contributed ~11% of the total degradation, which is not negligible. In addition, as shown in Figure 11b, its influence could be high in the first 5 kh, contributing 20–31% to the total degradation.



Figure 10. Simulation of the stack voltage evolution of F1002-97 with time. Only MIC oxidation is considered to be the degradation. The inserted figures are the distribution of the electronic potential across the contact layer and the MIC.



Figure 11. (a) Voltage evolution from experiments and simulation. The red texts refer to the degradation rates: (b) Ratio of the degradation rate due to MIC oxidation to the total degradation rate. Comparison of the experimental data and simulation results of F1002-97: (a) shows the voltage evolution only due to MIC oxidation and (b) exhibits the share of voltage degradation rate caused by MIC oxidation. A sudden increase is observed in (b) because the total degradation rate decreases from 7.3 mV/kh to 1.6 mV/kh at 40 kh.

8.3. Voltage Degradation Due to MIC Oxidation in F1004-67

Crofer 22 APU was used as the MIC, and APS MCF was applied as the protective coating for the stack F1004-67. It was operated at 730 °C and 0.5 A/cm² with a degradation rate of 3 mV/kh over 25 kh operation. With a dense coating, the simulation predicts that the voltage decrease at a rate of 0.016 mv/kh due to MIC oxidation, which only contributes \sim 0.53% to the total degradation. Therefore, degradation due to the MIC

oxidation can be safely neglected. The degradation mainly arise from other processes, such as Ni agglomeration and Cr poisoning.

9. Conclusions and Outlook

A three-dimensional multiphysics SOC model is built by using a computational code, openFuelCell2. The code is constructed inside an open-source library, OpenFOAM. The model is proven to be able to carry out not only steady-state but also time-dependent simulations. The model is intensively validated for the steady-state simulation to precisely predict I–V curves under different operation temperatures, fuel humidities and operation modes with a maximum error below 4.5%. It is indicated that the model is able to predict the SOC performance, as the spatial distribution of overpotential and mole fractions simulated by OpenFOAM and COMSOL are very close. A time-dependent model of MIC oxidation is proposed. Different experiments validated that the model can predict accurate thicknesses of the oxide scale. According to the simulation, it is found that if MCF protective coating fabricated by APS is applied, the degradation due to the MIC oxidation can be generally neglected since it only contributes ~0.53% to the total degradation rate. In contrast, if WPS MnO_x protective coating is used, the MIC oxidation cannot be ignored, especially at the beginning of the operation.

This work establishes a framework for further sophisticated modeling studies. For example, although an empirical formulation is currently used to calculate the parabolic constant, models accounting for more detailed physics can be proposed and implemented. More degradation mechanisms, such as Ni agglomeration and Cr poisoning, will be integrated in the model in the near future to reveal how the performance of the SOC stack evolves with time and how to optimize the stack design.

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Nomenclature

Abbreviations

8YSZ	8 mol% yttria-stabilized zirconia
AEL/ELEC	The interface between the air electrode and the electrolyte
APS	Atmospheric plasma spraying
FEL/ELEC	The interface between the fuel electrode and the electrolyte
FESC	Fuel electrode-supported cell (FESC)
FZJ	Forschungszentrum Jülich GmbH
GDC	Gadolinia-doped ceria
ITM	Intermediate temperature metal
LCC12	$La_{0.97}Mn_{0.4}Co_{0.3}Cu_{0.3}O_{3-\delta}$
LSCF	$La_{0.58}Sr_{0.4}Co_{0.2}Fe_{0.8}O_{3-\delta}$
MCF	$MnCo_{1.9}Fe_{0.1}O_{3-\delta}$

MFC	Mass flow controller	
MIC	Metal interconnect	
NS	Navier–Stokes	
PVD	Physical vapor deposition	
SOC	Solid oxide cell	
SP	Screen printing	
WPS	Wet powder spraying	
Greek symbols		
X	Volume fraction	[1]
κ	Permeability	[m ²]
$\phi_{ m ele}$	Electronic potential	[V]
$\phi_{\rm io}$	Ionic Potential	[V]
σ_i^{ele}	Electronic conductivity of phase <i>i</i>	[S/m]
σ_i^{io}	Ionic conductivity of phase <i>i</i>	[S/m]
$\sigma_{\text{oxide scale}}$	Electronic conductivity of the oxide scale	[S/m]
τ	Tortuosity	[1]
υ	Diffusion volume	[m ³ /mol]
ε	Porosity	[1]
Ξ	Parabolic constant	[m ² /s]
Roman symbols		
Ũ	Velocity	[m/s]
С	Molar density	[mol/m ³]
d	Thickness of the oxide scale	[m]
D_{ii}^{B}	Binary diffusion coefficient	[m ² /s]
$D_{ij}^{\mathbf{K}}$	Knudsen diffusion coefficient	$[m^2/s]$
i _{ele}	Electronic current density	$[A/m^2]$
i _{io}	Ionic current density	$[A/m^2]$
i _v	Reaction current density	$[A/m^3]$
М	Molar mass	[g/mol]
Ν	Molar diffusion flux	$[mol/(m^2 s)]$
Р	Pressure	[Pa]
S _{mass}	Source term of mass	$[g/(m^3 s)]$
S _{mole}	Source term of mole	$[mol/(m^3 s)]$
Т	Temperature	[K]
Y	Molar fraction	[1]
R	Gas constant	[(J mol)/K]

Appendix A

Table A1. Other physical parameters used in the model.

Parameter	Value	Unit	Reference
Mean pore diameter of the fuel electrode $(d_{p,fel})$	641	nm	[59]
Mean pore diameter of the air electrode $(d_{p,ael})$	306	nm	[59]
Permeability in the fuel electrode (κ_{fuel})	1×10^{-13}	m ²	
Permeability in the air electrode (κ_{air})	1×10^{-13}	m ²	
Hydrogen diffusion volume (v_{H_2})	6.12×10^{-6}	m ³ /mol	[60]
Water diffusion volume $(v_{H_{2}O})^2$	1.31×10^{-5}	m ³ /mol	[60]
Nitrogen diffusion volume (v_{N_2})	1.79×10^{-5}	m ³ /mol	[22]
Oxygen diffusion volume (v_{O_2})	1.66×10^{-5}	m ³ /mol	[22]
Volume fraction of Ni in solid part of fuel electrode (χ_{Ni})	0.4	1	[25]
Volume fraction of YSZ in solid part of fuel electrode (χ_{YSZ})	$1-\chi_{\rm Ni}$	1	[25]
Volume fraction of LSCF in solid part of air electrode (χ_{LSCF})	1	1	[57]
Porosity of the Ni mesh ($\varepsilon_{Ni mesh}$)	0.8	1	
Porosity of the fuel electrode (ε_{fel})	0.2	1	57
Porosity of the fuel electrode support layer ($\varepsilon_{\text{felsup}}$)	0.415	1	[25]
Porosity of the air electrode (ε_{ael})	0.45	1	[25]
Porosity of the contact layer (ε_{ocl})	0.45	1	
Porosity of the WPS coating ($\varepsilon_{\text{coating,WPS}}$)	0.45	1	[52]
Porosity of the APS coating ($\varepsilon_{\text{coating APS}}$)	0.03	1	[53]
Tortuosity of LSCF ($\tau_{\rm LSCF}$)	2.2	1	[25]
Tortuosity of Ni (τ_{Ni})	4.64	1	251
Tortuosity of YSZ (τ_{YSZ})	2.18	1	[25]

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