

Cite this: *Sustainable Energy Fuels*,
2023, 7, 280

Multiscale modelling potentialities for solid oxide fuel cell performance and degradation analysis

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Solid oxide fuel cells are electrochemical devices that are able to directly convert the chemical energy of fed fuels to electricity as well as to provide heat through exhausted gases allowing a higher energy efficiency compared to traditional thermal engines. However, the state-of-the-art materials show a drastic performance drop after too few working hours because of irreversible microstructural changes. Here the main issue consists of improving cell durability by optimising its structure and operative conditions. Modelling can significantly support this target, permitting a better understanding of different phenomena and providing information that are difficult to directly measure. However, degradation simulation is a quite challenging task due to the complexity of the studied systems, where different phenomena overlap, as well as due to the numerous data requested on both electrochemical and microstructural features. Depending on the available cell information and the analysis detail level, a multiscale modelling approach is a promising solution for providing effective results with reduced computational efforts. Based on a macroscale characterization, for example, semi-empirical degradation functions can be directly derived from electrochemical impedance spectra and area-specific resistance variations without knowing anything on the microstructure in order to estimate global cell performance and durability through a lumped-parameter model. Whereas, when aiming at the identification of an aged element specific behaviour, detailed formulations have to be introduced for each mechanism following a microscale approach. In such cases, a local-level modelling is fundamental in view of uneven distributions of properties on the cell plane which influence locally the degradation process development and resulting performance.

Received 15th August 2022
Accepted 22nd November 2022

DOI: 10.1039/d2se01118b

rsc.li/sustainable-energy

1. Introduction

Solid Oxide Fuel Cells (SOFCs) are electrochemical devices commonly applied in gas to power applications.¹ The electrochemical exothermic oxidation of hydrogen permits direct electricity generation with minimum environmental issues realizing water as the primary product. Modularity, performance independence from the system scale, reduced noise and absence of mobile parts are other significant advantages compared to conventional power generators.² Different from other available fuel cell designs, high working temperatures favour the use of SOFCs as co-generation units, which are mainly applied at the residential level providing ~ 0.75 – 1.5 kW_{el} and ~ 1 kW as the maximum thermal power.³ There have been also some attempts of use in the mobile sector as auxiliary power units or for vehicle propulsion combined with batteries.^{4–6} Despite promising results in terms of nominal performance, SOFCs are still characterised by an insufficient lifetime reaching 40 000–45 000 h as the best still rare durability in stationary applications.⁷ Whereas the commercial target

requires a double value at least to be competitive with other solutions.⁸ This limit derives from significant microstructural variations which involve electrodes as well as interconnects and, to a lesser extent, the electrolyte.⁹ Indeed, the state-of-the-art anode consists of a Ni-YSZ (Yttria Stabilized Zirconia) cermet, where the hydrogen reacts at the Triple Phase Boundary (TPB) sites, namely the active zones characterised by the contemporaneous presence of an electron conductor (*i.e.*, the metal), an ionic conductor (*i.e.*, the ceramic) and the gas phase to favour reactant transport. Ni permits the achievement of excellent performance due to its good catalytic activity and electron conductivity, on the other hand this metal becomes unstable under long-lasting operation with observed agglomeration and migration.¹⁰ Indeed, high working temperatures favour the particle growth reducing the active area and thus provoking a significant performance reduction (~ 18 – 41% of the total losses).¹¹ Under a stressful fuel utilization, Ni can be oxidised by the produced steam and then reduced again when the nominal operation is restored, which provokes crack formation because of the volume changes between these two states.¹² Moreover, Ni migrates away from the electrolyte interface further varying the effective active area.¹³ At the cathode, perovskites are commonly applied as a pure layer or mixed with high ionic conductor

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materials in order to increase their performance by extending TPB sites and enhancing ion migration. LSCF-CGO (Lanthanum Strontium Cobalt Ferrite-Cerium Gadolinium Oxide) or LSM-YSZ (Lanthanum Strontium Manganite-Yttria Stabilized Zirconia) are the main available solutions at the industrial level.¹⁴ However, both these electrodes easily react with other cell components forming high resistive phases,¹⁵ without the introduction of barrier layers. For instance, LSCF demixing consists of a heavy accumulation of strontium and cobalt, in less amount, near the electrolyte interface creating insulating zirconate deposits.¹⁶ Moreover, secondary phase formation also derives from the interactions with ferritic stainless-steel based interconnects, which release Cr volatile species covering the catalytically active sites.¹⁷ In addition to the release of Cr, interconnects provoke fracture development and electrical resistance increase due to the thickening of the oxide scale used to prevent steel corrosion.^{18,19} Less frequently, YSZ electrolyte can progressively change its lattice from the initial cubic phase to the tetragonal one reducing ion conductivity.²⁰

In view of these several degradation mechanisms, further experimental and theoretical studies are still required in order to have a better insight of long-lasting cell operation and to project effective mitigation strategies.

As underlined in several experimental tests, cell degradation is strictly correlated to applied working conditions and the initial material structure. Temperature has been pointed out as one of the main sources, since an increase of $\sim 100^\circ$ can result in an acceleration of 1.5–2 times.²¹ Gas composition, and above all, the humidity level significantly influence cell durability, with an observed increase of anodic Ni instability at high steam content.²² System performance has also a relevant dependence on the initial material structure. For instance, the presence of protective layers among electrodes, electrolyte and the interconnect reduces the risk of secondary phase formation.²¹ Moreover, larger Ni particles in the anodic cermet are more prone to agglomeration resulting in a lower catalytic activity.²³ Due to the degradation variability being a function of specific test features, quite different results have been presented in the literature for long-lasting cell operations. For instance, in ref. 24, the Ni-YSZ/YSZ/LSM-YSZ system was tested for 6000 h at 1123 K and 0.3 A cm^{-2} reporting a degradation rate of 1.4 V\% kh^{-1} . After 5000 h at 1023 K and 0.2 A cm^{-2} , a lower degradation rate was reported resulting in 0.75 V\% kh^{-1} .²⁵ Whereas the Ni-YSZ/YSZ/LSCF-CGO system showed a more stable performance with an overall value of 0.5 V\% kh^{-1} after 100 000 h of operation working at 973 K and 0.5 A cm^{-2} .²⁶

Referring to modelling activity, a multiscale analysis well fits the challenge to study SOFCs at nominal and aged states. However, significant computation efforts have to be made in order to merge different simulation levels. For the nominal cell behaviour, some studies have been recently published in the literature, where a multiscale approach has been followed starting from atomic level models of specific transport phenomena and reaction paths to heat and material transport macro models, as well as from 3D material reconstruction to cell voltage prediction.^{27–30} Looking at degradation analysis, different modelling approaches have been proposed which

include both formulations of single mechanisms and general expressions for voltage time evolution. Referring to a specific phenomenon, the easiest alternative consists of determining semi-empirical correlations which represent the time changes of cell-specific features or microstructural variables. For instance, the Ni particle size increase or directly the resulting TPB density reduction were modelled through a power-law formulation to represent agglomeration mechanisms.^{11,31} The decrease of electrolyte conductivity due to phase changes was forecast by introducing an empirical temperature-dependent corrective coefficient.³² Otherwise, theoretical studies have formulated physically based kinetics to predict degradation mechanisms. In ref. 33, the oxide scale growth on the interconnects was computed considering the oxidation reaction at the metal interface as well as the oxygen diffusion through the formed oxide. In ref. 34 the anodic Ni reduction and oxidation kinetics were evaluated in the cell working temperature range to understand if they are characterised by a reaction or a diffusion controlled path. In view of the fundamental role of microstructural features in the electrode catalytic activity, geometrical models have been developed to predict the specific surface area and TPB density for composite electrodes, such as Ni-YSZ and LSCF-CGO.^{35,36} These were considered as a random distribution of interconnected spheres evolving during cell operation to reach lower energy-level states. Considering the simulation of global cell degradation, some empirical correlations have been formulated which express the total resistance variation measured through electrochemical impedance spectra as a further polarization loss.³⁷ In addition, the time evolution of different resistances has been determined by applying a proper equivalent circuit model which suggests the needed corrective terms in the kinetic formulation.³⁸ However, multiscale simulation codes that are able to merge all aspects, from the single mechanisms to global cell degradation, and involving simultaneously empirical correlations as well as microstructure-dependent kinetic formulations to be applied depending on the available experimental observations, are still rare in the literature.^{39–41}

In such a scenario, the authors propose SIMFC (SIMulation of Fuel Cells) as an effective multiscale simulation code which combines the different levels of analysis in a single tool for predicting cell performance and durability. SIMFC is an in-home Fortran code for high-temperature cell operation including MCFCs (Molten Carbonate Fuel Cells) and SOFCs.^{42,43} Based on the conservation equation resolution for anodic and cathodic sides, it permits global performance prediction by a lumped-parameter modelling approach for a preliminary analysis of the system. Solving the same formulae for each sub-element discretizing the cell plane in a local-level modelling approach, the global cell behaviour is evaluated as a function of local physicochemical parameter distributions.⁴⁴ Focusing on SOFC durability, specific degradation functions derived empirically for a macroscale approach and dependent on the local microstructure for a microscale approach were introduced into the kinetic formulations. In addition to being one of the rare comprehensive tools for solid oxide cell modelling, it is also characterised by a high computational velocity reaching



Table 1 SIMFC kinetic multiscale formulation^a44,46,48

Term	Macroscale formulation	Microscale formulation	Material properties
Ohmic overpotential	$\eta_{\text{ohm}} = R_{\text{ohm}}j$	$\eta_{\text{ohm}} = \sum \frac{d_i}{\sigma_i^{\text{eff}}} J$	$\sigma_i^{\text{eff}} = \sigma_{i,0}(1-\varphi)^2 \left(\frac{\psi_i - \psi_i'}{1 - \psi_i'(1-\varphi)} \right)^2$
Activation overpotential	$\eta_{\text{act}} = \frac{2RT}{zF} \sinh^{-1} \left[\frac{J}{2J_0} \right]$	$\gamma = \gamma' l_{\text{TPB}} P_{\text{el}} P_{\text{cd}}$	$\psi_i' = \frac{1.764r_i}{(Z - 1.764)r_j + 1.764r_i}$
Diffusion overpotential	where $J_0 = \gamma \prod y_i^{\alpha_i} \exp \left(\frac{-E_{\text{act}}}{RT} \right)$ $\eta_{\text{diff}} = \frac{RT}{zF} \ln \left[\prod \left(\frac{p_{i,\text{bulk}}}{p_{i,\text{TPB}}} \right)^{2\alpha_i} \right]$	$p_{i,\text{TPB,an}} = p_{i,\text{bulk,an}} \pm \frac{JRT}{zF} \left(\frac{d_{\text{ac,cm}}}{3D_{i,\text{ac}}^{\text{eff}}} + \frac{d_{\text{sup,an}}}{D_{i,\text{sup}}^{\text{eff}}} \right)$ $p_{i,\text{TPB,cat}} = \frac{p}{\varphi_{\text{cat}}} - \left(\frac{p}{\varphi_{\text{cat}}} - p_{i,\text{bulk,cat}} \right) \exp \left(\frac{\varphi_{\text{cat}} RT d_{\text{cat}} J}{z F p D_{i,\text{cat}}^{\text{eff}}} \right)$	$l_{\text{TPB}} = \frac{\left(1 + \frac{r_{\text{cd}}^2}{r_{\text{el}}^2} \right) \psi_{\text{el}} r_{\text{cd}} Z}{3(1-\varphi) \psi_{\text{cd}} \sin \beta \min(r_{\text{cd}}, r_{\text{el}})} \frac{1}{4r_{\text{cd}}^3} \frac{\psi_{\text{el}} r_{\text{cd}} + \psi_{\text{cd}} r_{\text{el}}}{\psi_{\text{el}} r_{\text{cd}} + \psi_{\text{cd}} r_{\text{el}}}$ $D_i^{\text{eff}} = \frac{\varphi}{\xi} \left(\frac{1}{D_{\text{mix},i}^{\text{M}}} + \frac{1}{D_i^{\text{K}}} \right)^{-1}$ $D_{\text{mix},i}^{\text{M}} = \sum_{j \neq i} \left(\frac{y_j}{D_{i-j}^{\text{M}}} \right)^{-1} (1 - y_i)$ and $D_i^{\text{K}} = \frac{2r_p}{3} \sqrt{\frac{8000RT}{\pi M_i}}$ $r_p = \frac{2}{3(1-\varphi)} \frac{r_{\text{el}} r_{\text{cd}}}{\psi_{\text{el}} r_{\text{cd}} + \psi_{\text{cd}} r_{\text{el}}}$

^a Nomenclature for electrochemical kinetics. d : thickness (an: anode, cat: cathode), D^{eff} : effective diffusion coefficient (ac: active layer, sup: support), D^{K} : Knudsen diffusion coefficient, D^{M} : molecular diffusion coefficient according to Chapman–Enskog theory, E_{act} : activation energy, E_{eq} : reversible voltage, F : Faraday constant, f : current density, J_0 : exchange current density, l_{TPB} : triple phase boundary density, M : molecular weight in g mol^{-1} , P : percolation network probability, p : gas partial pressure, R : ideal gas constant, r_{cd} : electron conductive particle radius, r_{el} : ion conductive particle radius, r_p : pore radius, R_{ohm} : ohmic resistance, T : temperature, y : molar fraction, V_{sofc} : cell voltage under load, Z : coordination number, z : charge number, α : kinetic order, β : contact angle, γ : kinetic constant, η : overpotential, φ : porosity, φ_{cat} : combination of cathodic diffusion coefficients, ξ : tortuosity, σ : conductivity, σ_0 : pure material conductivity, φ : porosity, ψ : phase fraction (ed: electronic conductive, el: ionic conductive), and ψ' : threshold phase fraction.

convergency usually under one minute. This is a significant improvement compared to reference multiscale models, mainly based on computational fluid dynamic software.³⁹ All potentialities of SIMFC as a simulation tool are described in the following chapters, also reporting the results of two specific case studies as examples.

2. Nominal performance modelling

SIMFC is a physically based model which evaluates the cell behaviour solving conservation equations in terms of material, energy and momentum balances as described in authors' previous studies.^{43,45} Its core consists of ad hoc electrochemical kinetics to compute the cell voltage derived from Nernst, Ohm and Butler–Volmer equations (eqn (1)), described in detail elsewhere.^{44–46}

$$V_{\text{SOFC}} = E_{\text{eq}} - \sum \eta$$

$$= E^0(T) + \frac{RT}{zF} \ln \left(\frac{p_{\text{H}_2} p_{\text{O}_2}^{0.5}}{p_{\text{H}_2\text{O}}} \right) - \eta_{\text{ohm}} - \eta_{\text{act}} - \eta_{\text{diff}} \quad (1)$$

Considering overpotential formulations reported in Table 1, the main kinetic parameters are derived semi-empirically for a macroscale analysis, whereas in a microscale study they are expressed as a function of material features following the percolation theory to estimate available catalytic sites, conductivities and gas diffusion paths. In such a model, the electrodes are assumed as a random packing of ideal spheres consisting of electronic and ionic conductive phases.⁴⁷ Here the ohmic term (*i.e.*, charge migration resistance) results in the combination of different layer contributions, where the effective conductivity is computed knowing the pure material value, particle radius and phase fraction. However, according to this formulation the pure ionic–electronic transport resistances are just evaluated, neglecting the effects due to cell setup and contact resistances. The activation overpotential depends on the availability of catalytically active sites or, in other terms, the triple phase boundary density. Finally, the diffusion overpotential for gas mass transport considers the diffusion coefficients and transport paths in view of the particle distribution. Table 1 summarises the proposed multiscale formulation for overpotential terms applied to different cell layers referring to the nominal behaviour; introduced degradation functions are discussed in detail in the following sections.

As results, SIMFC computes the voltage dependence on working conditions as the global cell behaviour index and maps of main physicochemical parameters on the cell plane as local results. SOFC performance is estimated referring to the electric efficiency $\varepsilon_{\text{SOFC}}$ (*i.e.*, the ratio between the produced power P_{SOFC} and the power corresponding to the fed hydrogen flow rate N_{H_2} in terms of its Lower Heating Value LHV_{H_2}) and the Degradation Rate DR_{SOFC} (*i.e.*, the voltage or resistance/overpotential variation over time t normalised to its initial value), as shown in eqn (2) and (3), respectively.

$$\varepsilon_{\text{SOFC}}(\%) = \frac{P_{\text{SOFC}}}{N_{\text{H}_2} \text{LHV}_{\text{H}_2}} = \frac{V_{\text{SOFC}} J A}{N_{\text{H}_2} \text{LHV}_{\text{H}_2}} \quad (2)$$

$$\text{DR}_{\text{SOFC}}(\%) = \left(\frac{x_t - x_0}{x_0} \right) \frac{1000}{t} \quad (3)$$

where A is the cell active area. Here, eqn (2) represents the energy conversion efficiency from fuel chemical energy to electricity through the electrochemical process, whereas eqn (3) provides a preliminary quantification of performance degradation assuming a linear variation of voltage or resistance/overpotential resulting in a negative value for the first case (*i.e.*, voltage reduction) and a positive one for the second (*i.e.*, resistance/overpotential increase).

3. Macro-level cell degradation modelling

3.1 Methodology

At a macroscale level without knowing the cell microstructure, the performance degradation can be modelled through the introduction of further time-dependent overpotential terms η_{deg} distinguishing between the increase of ohmic and polarization contributions (eqn (4)). These degradation functions are determined based only on electrochemical characterization measurements, such as electrochemical impedance spectra and characteristic curves (Fig. 1). Here the attending voltage drop can be predicted during cell operation; nevertheless, these equations are quite specific for just the studied system.

$$V_{\text{SOFC}} = E_{\text{eq}} - \sum \eta - \eta_{\text{deg,ohm}} - \eta_{\text{deg,pol}} \quad (4)$$

In detail, the ohmic evolution derives from the time profile of internal resistance (*i.e.*, the first intercept of the real axis in electrochemical impedance spectra) independently from measured conditions, at the open circuit voltage or under load, in view of the direct proportionality between the ohmic resistance and the current density. Whereas the degradation function of the polarization term requires the J - V curve profile knowledge because of its more complex dependences summing activation and diffusion overpotentials (Table 1). Knowing the ohmic resistance every time, $\eta_{\text{deg,pol}}$ can be obtained by the punctual variation of the differential resistance R_d , defined as the derivative of the voltage with respect to the current density.⁴⁹ Indeed, the voltage reduction means an R_d increase under fuel cell operation. Note the R_d profile is usually distinguished in three domains as a function of applied loads. The initial rapid decrease at low current densities represents the region where the activation loss dominates, the second with a plateau is due to charge transport resistances which are constant terms independent of load according to Ohm's law, and the last one shows gas diffusion limitations resulting in a drastic increase at high current densities.

3.2 Case study

This approach was validated based on reference data of an anode supported cell tested for 9000 hours (~ 375 days) in fuel



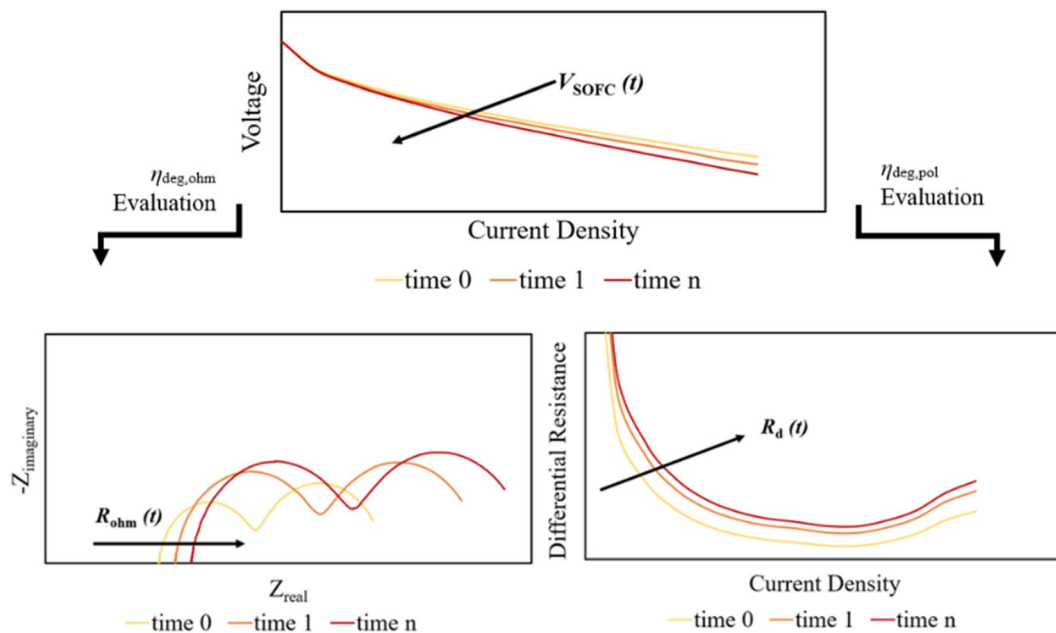


Fig. 1 Macroscale evaluation of SOFC voltage degradation based on electrochemical characterization under operation.

cell mode under 0.5 A cm^{-2} to avoid the fuel starvation (H_2 utilization equal to 60%).⁵⁰ The 9 cm^2 cell consisted of a Ni-YSZ anode ($\sim 260 \mu\text{m}$), a YSZ electrolyte ($\sim 8 \mu\text{m}$), a CGO barrier layer ($\sim 5 \mu\text{m}$) and an LSCF-CGO cathode ($\sim 50 \mu\text{m}$). Electrochemical characterization included continuous voltage monitoring during the operation, electrochemical impedance spectra and characteristic curves at regular time steps after the first 4000 testing hours, allowing for the determination of the discussed

empirical degradation functions. Assuming an isothermal behaviour, 0D-level SIMFC was applied as a simulation tool in view of the small-scale cell and missing microstructural data. Table 2 reports the tested working conditions of the cell, the kinetic parameters with reference ranges and the derived empirical degradation functions for ohmic and polarization contributions dependent on current density, J , in A cm^{-2} and time, t , in hours. Note the ohmic term follows the linear proportionality consistent with Ohm's law. Whereas the polarization overpotential degradation was computed punctually assuming 0.05 A cm^{-2} as the current density step ΔJ in differential resistance calculation.

The obtained good match between simulated and experimental voltages validated the proposed approach, suitable as a preliminary study of the cell durability without performing detailed microstructural analysis but only based on common measurements carried out during operation. Both voltage time evolution and characteristic curves show an averaged relative error lower than 2% (Fig. 2). A bit lower accuracy with a still acceptable relative error of 3% characterises OCV conditions ($\sim 1.2 \text{ V}$) which might be due to leakage issues reducing the experimental value. The voltage had a degradation rate of -0.7 V\% kh^{-1} , reducing from 0.813 V to 0.763 V . This trend showed a more significant reduction in the first 2000 hours (DR_{SOFC} higher than -1 V\% kh^{-1}), followed by a quite stable operation and then a gradual continuous decrease in the final testing hours. Looking at different kinetic terms, at the nominal state the ohmic loss had the main weight among different overpotentials representing 57% vs. 42% of the activation loss and only 1% of the diffusion loss. Under operation the cell underwent a drastic increase of ohmic contribution, which had a difference of $\sim 37 \text{ mV}$ after 5000 and of $\sim 49 \text{ mV}$ after 9000 h at the reference working condition of 0.5 A cm^{-2} , representing the

Table 2 Operating conditions, kinetic parameters and degradation functions for a 9000-hour working Ni-YSZ/YSZ/LSCF-CGO cell

Working condition	Anode	Cathode
T [K]	1023	1023
p [atm]	1	1
Flow rate [Nl h^{-1}]	3.16	22.6
Feed composition	Pure hydrogen	Dry air
Kinetic parameter	Anode	Cathode
E_{act} [kJ mol^{-1}]	100 100–120 as ref. 51	110 110–160 as ref. 51
Kinetic order α [—]	0.1 for H_2 0.1–2 as ref. 52 0.44 for H_2O –0.5–1 as ref. 52	0.20 for O_2 ref. 53
γ [A cm^{-2}]	6×10^4 5.5×10^4 – 5.5×10^6 as ref. 54	1×10^6 7×10^4 – 7×10^6 as ref. 54
Degradation function		
$\eta_{\text{deg,ohm}}$ [V]		$(10^{-9}t^2 + 2 \times 10^{-5})J$
$\eta_{\text{deg,pol}}$ [V]		$(2J^2 + 3J + 1) \times 10^{-5}t\Delta J$



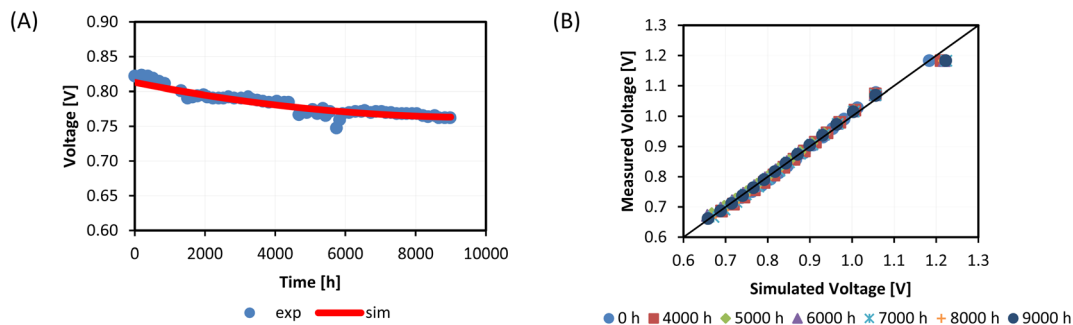


Fig. 2 Comparison between experimental and simulated voltages looking at the durability test (A) and J - V curve measurements at different operation hours (B) for a 9000-h working Ni-YSZ/YSZ/LSCF-CGO cell.

66% of the total losses at the test end. Polarization contribution varied only about some millivolts. This cell aging caused an efficiency reduction of 8% after 9000 h, starting from an initial value of 38% in view of imposed quite low hydrogen utilization.

4. Micro-level cell degradation modelling

4.1 Methodology

Knowing microstructural parameter variations, a more detailed analysis can be performed trying to distinguish the role of different degradation mechanisms in the global cell performance worsening. In this framework, a local-level model is fundamental to evaluate the specific structure changes as a function of local working conditions, such as temperature and gas composition, which have a significant influence on the system behaviour.⁴⁸ Here, starting from the time evolution profiles of microstructural features, the simulation can estimate the overpotential changes and the resulting cell voltage

decrease as schematized in Fig. 3. Referring to the most severe degradation processes, a local-level simulation was carried to evaluate the performance failure of an anode supported cell due to anodic Ni agglomeration and Cr poisoning at cathodic active sites in the case of poorly working interconnect barrier layers.⁹ Following both experimental and theoretical observations,^{23,55} the agglomeration of the Ni-YSZ electrode was modelled using a power law type formulation (eqn (5)).

$$r_{\text{Ni}}^n(t) - r_{\text{Ni}}^n(t_0) = k(T) \times (1 - \exp^{-\lambda t}) \quad (5)$$

where r_{Ni} is the Ni radius within the cermet, t is the observation time, and n , k and λ are the empirical parameters (k depends on the temperature).

Knowing Ni radius time evolution, the consequences on cell behaviour can be evaluated by applying the microscale kinetics formulation presented in Table 1, which expresses the exchange current density J_0 , the material conductivity σ and gas diffusion coefficients D^{eff} as functions of the particle size. As observed in reference studies,^{23,35,40,56,57} the radius increase causes the

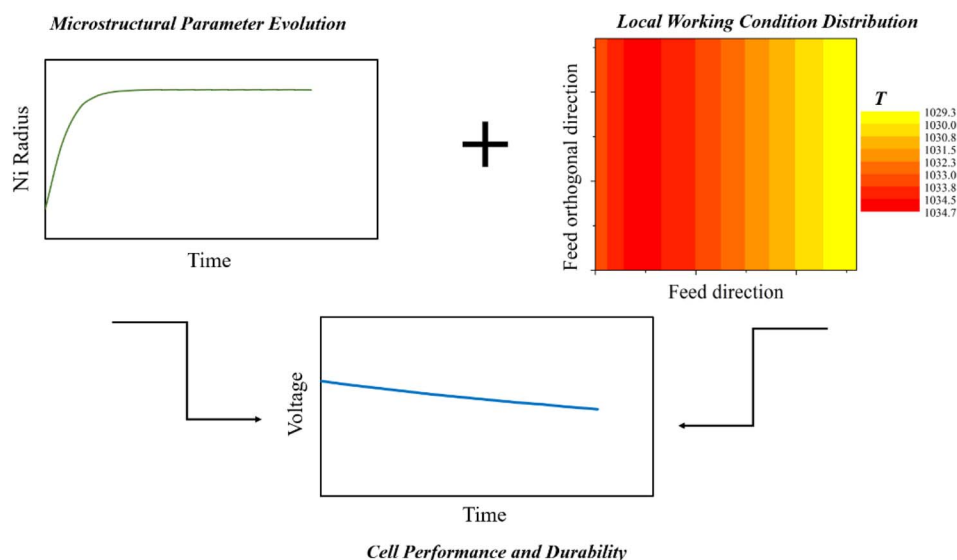


Fig. 3 Microscale evaluation of SOFC voltage degradation based on microstructural parameter time evolution and local working condition maps.



reduction of the particle surface area, which means a lower contact section between both metal–ceramic particles (*i.e.*, lower catalytic activity and TPB density) and metal–metal, ceramic–ceramic particles (*i.e.*, lower electron and ion conductivity, respectively). Moreover, larger particles influence material porosity also by increasing the pore dimension.

Referring to cathodic degradation processes, LSM–YSZ poisoning is mainly due to electrochemical reactions of gaseous chromium-based compounds which deposit on the active sites. Considering $\text{CrO}_2(\text{OH})_2$ as the vapour phase species with the highest partial pressure in equilibrium with the formed Cr_2O_3 solid phase, $p_{\text{CrO}_2(\text{OH})_2}$ was computed according to an empirical correlation showing an Arrhenius-type dependence on the temperature (eqn (6)). Assuming Cr oxidation as a single electrochemically driven process, the current density J_{Cr} required for this reaction was evaluated by applying the Butler–Volmer equation and it allowed to determine the time evolution profile of cathodic TPB density (eqn (7)), as presented in ref. 58. Since the charge transfer associated with Cr deposition is significantly smaller than the oxygen evolution value, J_{Cr} can be neglected in the global overpotential calculation.

$$p_{\text{CrO}_2(\text{OH})_2}(\text{atm}) = 0.0226 p_{\text{H}_2\text{O}}^{0.992} \exp\left(\frac{-67000}{RT}\right) \quad (6)$$

$$\begin{aligned} \ln\left(\frac{l_{\text{TPBcat}}(t)}{l_{\text{TPBcat}}(t_0)}\right) &= -\int_0^t K_{\text{Cr}_2\text{O}_3} J_{\text{Cr}} dt \\ &= -\int_0^t K'_{\text{Cr}_2\text{O}_3} \nu_{\text{CrO}_2(\text{OH})_2}^{0.5} \nu_{\text{H}_2\text{O}}^{0.5} \sinh\left(\frac{F\eta_{\text{actcat}}}{2RT}\right) dt \end{aligned} \quad (7)$$

where K and K' are constants dependent on Cr_2O_3 physical properties. If Ni agglomeration can also occur without applying a load, Cr deposition is driven by the local cathodic overpotential. Moreover, in this second process the electrode conductivity is less influenced by not varying the particle size and thereby the contact area between them. Nevertheless, the points where LSM and YSZ should interact with the gas mixture are covered preventing oxygen diffusion into the catalytic surface and thereby blocking its electrochemical reduction. In other terms, TPB density decrease occurs as pointed out in ref. [32, 59–61].

4.2 Case study

Advantages of following a microscale approach were underlined in a devoted case study, analysing a commercial size anode supported cell with 80 cm^2 active area. A current density of 0.4 A cm^{-2} was imposed resulting in 70% hydrogen utilization and 16% oxygen utilization. Considering a cell composed of a Ni/YSZ anode ($\sim 250 \mu\text{m}$), a YSZ electrolyte ($\sim 8 \mu\text{m}$) and an LSM–YSZ oxygen electrode ($\sim 50 \mu\text{m}$), Ni particle agglomeration and Cr poisoning were modelled following the methodology proposed in the previous section. Working conditions, electrode microstructure, kinetics parameters and degradation coefficients are reported in Table 3 (note that anodic kinetic values are equal to previous case study values using the same electrocatalytic material). In view of a larger size, the cell was

Table 3 Operating conditions, kinetic and microstructural parameters and degradation model coefficients for 1000-h working Ni–YSZ/YSZ/LSM–YSZ cell simulation

Working condition	Anode	Cathode
T [K]	973–1123	973–1123
p [atm]	1	1
Flow rate [Nl h^{-1}]	20	200
Feed composition	96/4% vol $\text{H}_2/\text{H}_2\text{O}$ fed mixture	Air with 0.1% vol of steam
Kinetic parameter	Anode	Cathode
E_{act} [kJ mol^{-1}]	100 100–120 as ref. 51	120 110–160 as ref. 51
α [–]	0.10 for H_2 0.44 for H_2O	0.20 for O_2 ref. 53 –0.5–1 as ref. 52
γ' [mA]	0.26	0.9
Microstructural parameter	Anode	Cathode
r [μm]	Ni: 0.4 – YSZ: 0.3 Ref. 63	LSM: 0.6 – YSZ: 0.4 Ref. 58
Ψ [–]	Ni: 0.53 – YSZ: 0.47 Ref. 63	LSM: 0.52 – YSZ: 0.48 Ref. 58
φ [–]	0.15 (active layer) – 0.3 (support) Ref. 63	0.3 Ref. 58
ξ [–]	4 Ref. 64	3 Ref. 64
Degradation model coefficient		
n [–]		5 as ref. 55
$k(T)$ [m^5]		4.3×10^{-30} $T(K)$ ref. 56
λ [h^{-1}]		0.001 ref. 56
$K'_{\text{Cr}_2\text{O}_3}$ [h^{-1}]		70 ref. 58

not assumed isothermal; the temperature local gradient was computed on the cell plane to underline the influence on degradation through a sensitivity analysis in a common working range. Note that modelled results referred to an ideal case where only Ni-agglomeration and Cr deposition were considered, here these values could not be validated on reference experimental tests where other mechanisms are also observed, such as interconnect oxidation, electrolyte phase transition, secondary phase formation, crack development and layer delamination.^{9,20,62} However, their addition went beyond the aim of this work which wanted to show the relevance of the variable local control in having an effective prediction of commercial-scale cells as well as of the simulation being a support for experimental activities where the isolation of single processes is quite difficult when a full cell is tested.

Assuming a 1000-h working cell with a gas inlet temperature of 1023 K at both sides, the simulation allowed the estimation of performance worsening as a function of the Ni particle and Cr deposition increase. In such conditions, the voltage decreased



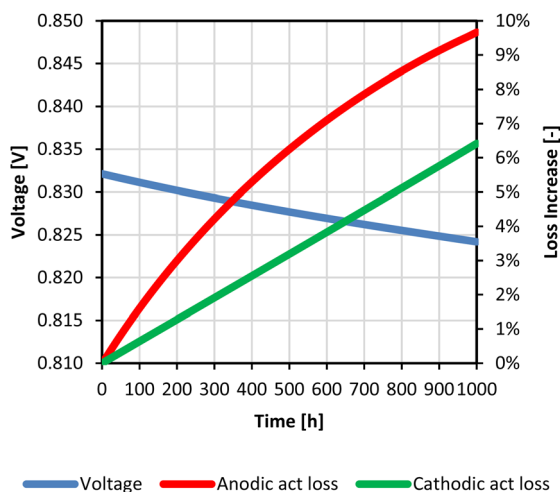


Fig. 4 Voltage and activation overpotential variations during 1000 h of working of the Ni-YSZ/YSZ/LSM-YSZ cell at 1023 K as the gas inlet temperature.

from 0.832 V as the nominal performance value to 0.824 V as the aged status. At both electrodes the main effects were detected looking at the activation overpotential (Fig. 4). Indeed, the particle agglomeration caused $\sim 7\%$ increase of Ni size, which meant a reduction of catalytic sites (*i.e.*, anodic TPB density decrease of $\sim 10\%$) and a resulting $\sim 10\%$ increase of anodic activation loss. Whereas the consequences on anode conductivity and gas diffusion were less evident. The growth of particles reduced the contact area among them,⁴⁷ increasing the ohmic resistance of $\sim 18\%$; however its value was quite low with respect to electrolyte contribution still guaranteeing an effective continuous network for the charge migration within the anode. Ni agglomeration also influenced the pore size and here the gas transport, yet the effects on diffusion overpotential in the studied time interval were negligible. Looking at the cathodic side, Cr poisoning provoked a TPB density decrease of $\sim 11\%$ resulting in $\sim 6\%$ increase of cathodic activation overpotential. In the proposed modelling no specific functions were introduced to evaluate the Cr deposition effect on other cathodic

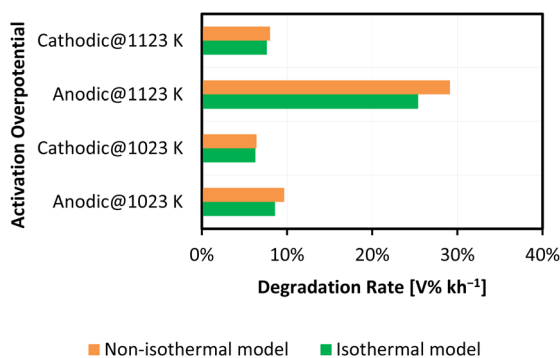


Fig. 5 Degradation rate of activation overpotential because of anodic and cathodic TPB density reduction assuming an isothermal and a non-isothermal Ni-YSZ/YSZ/LSM-YSZ cell working for 1000 h.

losses, however, these resulted quite negligible compared to the activation one looking at previous experimental observations.⁵⁸ In more detail, according to SIMFC local analysis a higher anodic TPB density decrease characterized the inlet section where the temperature was higher due to the faster reaction rate, resulting in more significant Ni particle sizes. Whereas the cathodic term showed an opposite trend since the worst performance occurred at the outlet due to lower temperatures.

The proposed local approach provided not only additional information useful for the local control of cell operation, but also more effective results of global performance for industrial scale systems. Neglecting the temperature gradient on the cell plane underestimated the activation overpotential degradation rates as shown in Fig. 5, where main differences between an isothermal and a non-isothermal cell behaviour are reported for 1023 K and 1123 K anodic and cathodic gas inlet temperatures as examples. According to experimental observations,⁹ the temperature is one of the main degradation source as confirmed by model results where $\sim 10\text{--}15^\circ$ increase on the cell plane due to the exothermic reaction was sufficient to have a non-negligible higher performance worsening. For instance, on feeding gas at 1123 K, $DR_{\eta_{act}}$ changed from 25 V\% kh^{-1} to 29 V\% kh^{-1} for the anodic side and from 9 V\% kh^{-1} to 10 V\% kh^{-1} for the cathodic side, comparing isothermal and non-isothermal results. At each temperature, more significant anodic variations confirmed the Ni agglomeration process to be more sensitive to temperature than Cr deposition, as discussed in more detail below.

In view of the significant role of temperature in degradation processes, a devoted analysis was performed to underline the changes of the Ni-YSZ/YSZ/LSM-YSZ cell behaviour in the common working range. As Table 4 reports, better nominal voltages occurred at high gas inlet temperatures due to lower polarization losses; indeed both catalytic activity and charge migration are thermoactivated mechanisms. Global voltage $DR_{V_{SOFC}}$ after 1000 h of operation was also slower (remember a negative value means voltage decrease): it was halved varying from 973 K to 1123 K because of better initial performances.

Looking at the local performance time evolution, further information was obtained. Fig. 6 shows the maps of the fuel cell produced power P_{SOFC} , the electric efficiency ε_{SOFC} and the voltage degradation rate $DR_{V_{SOFC}}$ computed every 100 h

Table 4 Ni-YSZ/YSZ/LSM-YSZ cell performance and the voltage degradation rate after 1000 h of working at variable gas inlet temperatures

Gas inlet temperature [K]	Voltage @ 0 h [V]	$DR_{V_{SOFC}}$ @ 1000 h [V\% kh^{-1}]
973	0.794	-1.07
998	0.815	-1.02
1023	0.832	-0.95
1048	0.845	-0.86
1073	0.854	-0.75
1098	0.860	-0.62
1123	0.861	-0.50



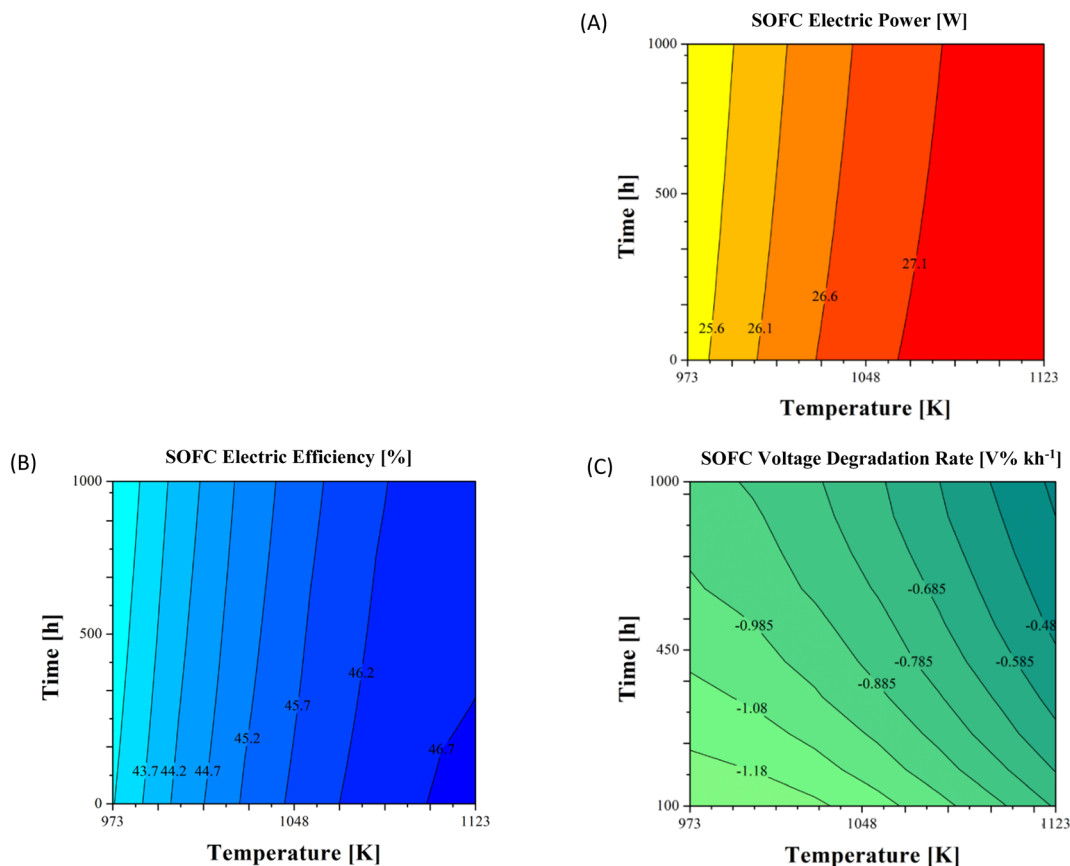


Fig. 6 Ni–YSZ/YSZ/LSM–YSZ cell performance time evolution in terms of the produced power (A), the electric efficiency (B) and the voltage degradation rate (C) at variable inlet gas temperatures assuming 1000 h operation.

operation. As observed, increasing the inlet temperature allowed for a bit higher electricity production in terms of some watts due to lower overpotentials every time. During operation the degradation caused a gradual power decrease, showing a more evident variation at 973 K compared to 1123 K: ~ 271 mW vs. ~ 137 mW (Fig. 6A). This lower value derived from the major weight of cathodic overpotential compared to the anodic one using LSM–YSZ as the electrocatalyst. Indeed, the oxygen reduction is limited at the TPB sites showing a higher resistance compared to the Ni–YSZ cermet, different from LSCF and LSCF–CGO where oxygen reduction can also occur directly on the perovskite surface without contacts with the ionic conductor.^{14,65} If on one hand a higher temperature favoured Ni agglomeration in view of a thermally activated surface diffusion as the driving mechanism,⁵⁵ on the other hand it slowed down Cr poisoning since the applied cathodic overpotential decrease disadvantages its deposition reaction.⁶⁶ Here, a lower global degradation rate and higher power were observed at 1123 K.

According to this trend, the electric efficiency reached its maximum at the 1123 K inlet temperature (*i.e.*, $\sim 47\%$ at the nominal state) and, however, a quite negligible variation was reported during 1000 h ageing (Fig. 6B), requiring more hours before a visible change of the efficiency.⁷ Also at lower

temperatures, $\varepsilon_{\text{SOFC}}$ never dropped below $\sim 42.5\%$ which is a still accepted performance.

Finally, considering the DR_{SOFC} time evolution (Fig. 6C), a decreasing trend of absolute values occurred at each temperature condition. After the first 100 h operation the lowest value of $-1.28 \text{ V\% kh}^{-1}$ was obtained at 973 K, whereas it changed until $-0.93 \text{ V\% kh}^{-1}$ in the final 100 hours. This trend was observed also at higher temperatures ($-1.12 \text{ V\% kh}^{-1}$ vs. $-0.70 \text{ V\% kh}^{-1}$ and $-0.76 \text{ V\% kh}^{-1}$ vs. $-0.35 \text{ V\% kh}^{-1}$ at gas inlet temperatures of 1048 K and 1123 K, respectively). The slowdown was correlated to the asymptotic growth of the Ni particle size which was characterized by a rapid increase within the first working hours and then stabilized to an asymptotic value due to the inhibiting effect of the YSZ backbone, since the YSZ percentage was sufficiently high to guarantee a good percolated network.⁶⁷ This stabilization for long-term operation was also observed for Ni–YSZ based cells in ref. 68. Moreover, the degradation rate value reduction was faster at higher working temperatures: $\sim 54\%$ at 1123 K vs. $\sim 31\%$ at 998 K. As observed, the anodic agglomeration had a major influence at higher temperatures due to the slowdown of the Cr deposition mechanism.



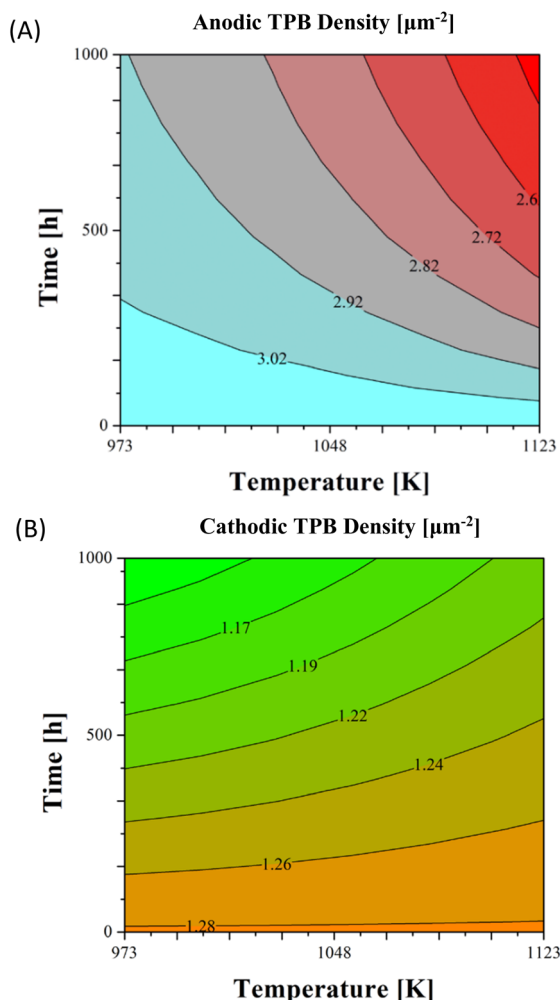


Fig. 7 Ni-YSZ/YSZ/LSM-YSZ electrode performance time evolution in terms of TPB density for anodic (A) and cathodic (B) sides at variable gas inlet temperatures assuming 1000 h operation.

Looking now at the specific electrode performance to better understand previously discussed results on the global cell behaviour, both Ni agglomeration and Cr poisoning provoked TPB density decrease: in the first case for the reduction of the

contact area between anodic particles, in the second for Cr deposition on the cathodic active sites. Nevertheless, different dependences on temperature and time evolution were underlined (Fig. 7). Indeed, the Ni particle size growth is a strongly thermoactivated process driven by vacancy diffusion from larger into smaller particles: here an increase of just $\sim 4\%$ was computed after 1000 h at 973 K, whereas it reached $\sim 17\%$ at 1123 K. This caused a more significant l_{TPB} decrease at higher temperatures, changing from the reference structure of $3.11 \mu\text{m}^{-2}$ to $2.4 \mu\text{m}^{-2}$ at 1123 K compared to $2.9 \mu\text{m}^{-2}$ at 973 K (Fig. 7A). Similar trends were reported for the active layer of anode supported cells in ref. 23, where TPB density reduction differed more than $1.5 \mu\text{m}^{-2}$ from 1023 K to 1123 K in performed durability tests. Consequently, the anodic activation overpotential variation was more highlighted by increasing working temperatures overcoming already $\sim 20\%$ after 500 h and $\sim 29\%$ after 1000 h. Whereas it never exceeded the 10% in the range between 973 K and 1023 K (Fig. 8A). In view of Ni asymptotic growth as previously discussed (eqn (5)), the degradation was faster in the initial working hours, whereas it slowed down after 600–700 h. Looking at the cathode behaviour, the opposite trend is visible: TPB density reduction occurred mainly at low temperatures decreasing of $\sim 0.16 \mu\text{m}^{-2}$ at 973 K vs. $\sim 0.08 \mu\text{m}^{-2}$ at 1123 K (Fig. 7B). In theory, Cr volatilization is favoured at temperature rise resulting in higher $\text{CrO}_2(\text{OH})_2$ partial pressures (eqn (6)), for instance 7.4×10^{-9} atm at 998 K vs. 1.6×10^{-8} atm at 1123 K. Devoted tests on stainless steel interconnects confirmed a higher evaporated Cr amount at 1123 K with respect to 923 K.⁶⁹ Nevertheless, simultaneously a higher temperature increases the exchange current density (Table 1) and here the local cathodic activation loss decreases (eqn (7)), becoming the predominant contribution for Cr deposition process, as confirmed in previous references for the LSM-YSZ electrode.^{60,66} Looking at the cathodic activation overpotential variation (Fig. 8B), a more homogeneous map was obtained due to a lower discussed thermal dependence. Moreover, a quite proportional increase is visible along 1000 h operation with higher variations in the final hours in agreement with trends observed experimentally.^{70,71}

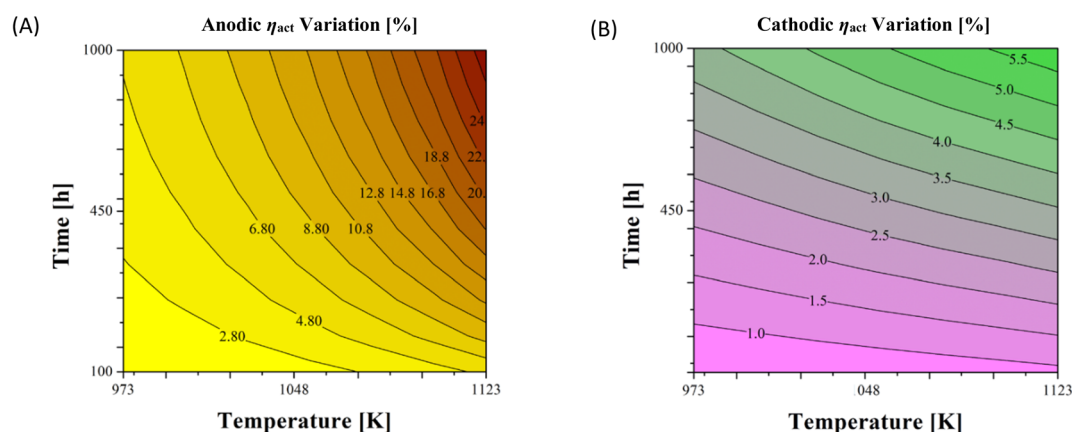


Fig. 8 Ni-YSZ/YSZ/LSM-YSZ electrode performance time evolution in terms of activation overpotential variation for anodic (A) and cathodic (B) sides at variable gas inlet temperatures assuming 1000 h operation.



5. Conclusions

A multiscale approach results in an effective solution to study solid oxide fuel cell performance and durability for both experimental and, above all, simulation activities. Indeed, it allows the choice of the most proper model dimension as a function of the investigation goal as well as of the system scale, moving from few centimetres in laboratory tests to the industrial size. Moreover, depending on the cell characterization available information, different levels of study can be applied into electrochemical kinetics formulation and degradation functions.

In this framework, in-home built SIMFC (SIMulation of Fuel Cells) was proposed as an effective tool resulting in one of the few available models comprehensive of all discussed features. Its potentialities were presented with reference to devoted case studies providing clear examples of its possible uses. For a preliminary analysis of cell durability, empirical correlations as a function of the applied load and time were derived by looking at the electrochemical impedance spectrum and J - V curve variations. Here, the weight of different overpotentials was evaluated by identifying the main sources of the global performance drop, as shown in the proposed case study on a 9000-h working anode supported cell. Whereas, knowing microstructural features, models specific for each degradation mechanism were applied and then correlated to kinetic parameters such as TPB density, conductivity and diffusivity coefficient. This more detailed approach is needful for industrial scale applications in view of working-condition-significant local gradients which have a relevant influence on degradation development. Considering temperature-dependent processes such as Ni agglomeration and Cr deposition reducing TPB density above all, neglecting local gradients predicted wrong activation losses of aged samples. This was more evident for the anodic contribution (error of $\sim 4\%$ after 1000 h operation) since a high temperature promotes the particle growth. Whereas Cr deposition is less sensitive to temperature changes, achieving an activation overpotential variation of ~ 5 – 7% at most.

Note that, after a proper tuning of the specific studied system, the proposed macroscale model approach is already ready to simulate the global cell behaviour and durability. Whereas in the microscale model approach two of the main degradation causes (*i.e.*, Ni agglomeration and Cr deposition) were only considered as examples. Following steps could focus on including further degradation mechanisms, after specific *ex situ* experimental campaigns where the behaviour of single layers is underlined. Indeed, in view of model robustness the computational time would not vary. A more challenging task consists of formulating degradation functions to predict domino effects, namely the correlations among different processes that can enhance or inhibit each other. Since the literature is quite scarce, an ad hoc design of the experiment would be projected, including both electrochemical and microstructural characterization of the cell to have appropriate information for modelling activities.

Author contributions

Bosio: conceptualization, methodology, software, and writing – review & editing; Bianchi: conceptualization, methodology, software, validation, visualization, writing – original draft, and writing – review & editing.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This research has received funding from the Fuel Cells and Hydrogen 2 Joint Undertaking (now Clean Hydrogen Partnership) under Grant Agreement no. 825027 (AD ASTRA project). This Joint Undertaking received support from the European Union's Horizon 2020 Research and Innovation program and Hydrogen Europe and Hydrogen Europe Research. The authors thank Daria Vladikova and her group for providing data on the 9000 h durability test used in the macroscale approach discussion.

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