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Short communication

Evaluation of proton conducting BCY10-based anode supported cells by co-pressing method: Up-scaling, performances and durability



J. Dailly ^{a,*}, M. Marrony ^a, G. Taillades ^b, M. Taillades-Jacquin ^b, A. Grimaud ^c, F. Mauvy ^c, E. Louradour ^d, J. Salmi ^e

- ^a EDF-EIFER, Emmy-Noether-Strasse 11, 76131 Karlsruhe, Germany
- ^b CNRS, Université de Montpellier, AIME, 87, Av. du Dr Schweitzer, 33608 Montpellier, France
- ^cCNRS, Université de Bordeaux, ICMCB, 87, Av. du Dr Schweitzer, 33608 Pessac, France
- ^d Céramiques Techniques Industrielles CTI, 382 Avenue du Moulinas, 30340 Salindres, France
- ^e Marion Technologies, Parc Technologique Delta Sud, 09340 Verniolle, France

HIGHLIGHTS

- Commercial powders produced at kilogram scale have been analysed and used.
- The maximum power density of around 180 mW cm⁻² is obtained at 600 °C on both sizes.
- The co-pressing method is only efficient at the laboratory scale.

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ABSTRACT

One of the main target of the French ANR CONDOR (2009–2011) project coordinated by EDF-EIFER was the elaboration of proton conducting $BaCe_{0,9}Y_{0,1}O_{3-\delta}$ -based cells using processes suitable at industrial level. Thus, commercial powders produced at kilogram scale (anode, electrolyte and cathode) have been analysed and used. Then, elaboration of protonic ceramic half-cells by co-pressing method from 25 mm to 80 mm in diameter has been assessed in order to establish the suitability of this shaping method regarding industrial requirements. According to results, the co-pressing method is relevant for the elaboration of cells up to 40 mm in diameter (laboratory experiment). A sintering step was carried out to densify the electrolyte layer to get a gas-tight membrane. Nickelate-based cathodes were screen-printed and electrochemical performances of the final assembly have been measured. Electrical power up to 180 mW cm⁻² at 600 °C has been obtained on both 25 and 40 mm cells.

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1. Introduction

The high energy conversion efficiency and the low impact to the environment are part of great advantages of Solid Oxide Fuel Cells (SOFCs), which are promising devices for electricity generation. They convert directly the chemical energy from the fuel to electricity. However, most of technical issues of SOFCs remain the high operating temperatures (>700 $^{\circ}$ C), leading to accelerated ageing of materials. During the last ten years, Protonic Ceramic Fuel Cell has considerably attracted attention thanks to the lower temperature range (400–600 $^{\circ}$ C) and the interest of a better energy efficiency

with the non-dilution of the fuel since the water is produced to the air electrode side.

Since Iwahara published in the 80's his work on protonic perovskites as electrolyte for fuel cells [1–4], many proton conducting oxide structures have been studied, like perovskites [5–11], based-perovskites (like $Ba_3Ca_{1,18}Nb_{1,82}O_9$ or $Baln_{0,8}Ti_{0,2}O_{2,6}\square_{0,4}$) [12–15], ortho-niobates [16–18] and others candidates [19–22].

Now, most of protons conducting ceramic cell researches are devoted to the optimization of their performances and reliability. This aspect request notably the strict reduction of ohmic resistance of each interface layers by promoting very thin and dense electrolyte layer [23–26], a well-architecture anode functional layer to increase the triple-phase boundary (TPB) for hydrogen electrode reactions [27,28] and a good chemical compatibility and microstructure for air electrodes side [29–32].

^{*} Corresponding author. Tel.: +49 721 61 05 13 52; fax: +49 721 6105 1332. E-mail address: dailly@eifer.org (J. Dailly).

But, until now, most of electrical performances related in the literature do refer to lab-scale size cell below 10 mm of diameter. Thus, no real investigations of efficient up scaling routes have been led until now in order to fit to the industrial requirements for the introduction of cells in stacks. This aspect appears fundamental to approach the stack manufacturing then the system integration key steps research in a reasonable term.

Here proposed is the assessment of co-pressing methods for an up-scaling of PCFC anode- supported cell manufacture. Microstructure and electrical performances of such cells are investigated.

2. Experimental

2.1. Powders

All samples were prepared using commercial powders (Marion Technologies®) produced at kilogram scale, listed below:

- Electrolytes: $BaCe_{0,9}Y_{0,1}O_{3-\delta}$ (BCY10) - Anode (cermet): $NiO-BaCe_{0,9}Y_{0,1}O_{3-\delta}$

- Cathode: $Pr_2NiO_4 + \delta$

2.2. Cells elaboration

The half cells were made by co-pressing of powders (at AIME laboratory and Céramiques Techniques Industrielles CTI^{\otimes}). In a first time, cermet spray-dried powder (NiO-BCY10) was pressed (100 MPa) into discs (-25-40-80 mm in diameter, 2 mm in thickness). Then, a given weight of electrolyte powder (BCY10) was sieved on the top of the anode discs in order to reach a homogeneous and reproducible thickness for all the samples, and then copressed at 150 MPa. The green half cells were sintered at 1350 °C for 10 h on a bed of anode powder in order to limit chemical pollution and geometrical deformation.

The cathode was made by screen-printing (at ICMCB and EIFER laboratories). The slurry was prepared using cathode-powder and 6 wt.% ethylcellulose in Terpineol. Two layers were screen-printed on half-cells, dried at 100 $^{\circ}$ C in a drying box and were then fired at 1150 $^{\circ}$ C for 2 h in air.

2.3. Characterization of the samples

Crystallographic phases were determined at room temperature by X-ray diffraction (INEL brand), in the $\theta/2\theta$ mode, using the K α 1Cu radiation. The phase identification was done using the software WPA.

The size distributions of powders dispersed in water are determined by dynamic laser light scattering using a Mastersizer 2000 apparatus (Malvern brand) using the Mastersizer 2000 5.12 F software.

In order to appreciate their TEC difference, dilatometric measurement were carried out on sintered material samples with a Netzsch DIL 402 PC from room temperature to 1000 $^{\circ}$ C with a heating rate of 5 $^{\circ}$ C min $^{-1}$.

The cell morphologies are evidenced by electron microscopy in the SEM mode in a field-emission scanning electron microscope.

2.4. Electrochemical test conditions

Single cells were tested in fuel cell testing systems at 600 °C with non-humidified hydrogen as fuel and compressed air as oxidant. Electrical contacts to the cell were made using platinum wires welded to the platinum meshes used as current collectors. AC impedance spectroscopy was obtained over the frequency range

from 0.01 to $10^5\ \mathrm{Hz}$ under open-circuit conditions and under polarization.

3. Results and discussion

3.1. Characterization of powders

As shown in Figs. 1 and 2, the as-prepared powders of BaCe_{0,9}Y_{0,1}O_{3- δ} and Pr₂NiO_{4+ δ} exhibit a well-developed crystallization and all the peaks can be well indexed as pure perovskite and Ruddlesden–Popper phases, respectively. It could be clearly seen that there is no evidence of the formation of other substances. All the XRD measurements made on the synthesized-phases are in agreement with the corresponding JCPDS card (n° 82–2372 for BaCe_{0,9}Y_{0,1}O_{3- δ}, n° 44–1159 for NiO, n° 34–1113 for Pr₂NiO_{4+ δ}, respectively). The average crystallite sizes are listed in Table 1.

Dilatometric experiments were carried out on sintered samples in order to measure their thermal expansion coefficient (TEC), and evaluate the potential mechanical stresses during co-firing. Values are summarized in Table 2: even if the difference between TEC values are rather different, no layer delamination has been observed after cell elaboration, meaning that membrane and substrates are likely to compensate this gap.

3.2. Up-scaling of cells

In the first stage, NiO–BaCe $_{0.9}Y_{0.1}O_{3-\delta}/BaCe_{0.9}Y_{0.1}O_{3-\delta}/Pr_2NiO_4+\delta$ assemblies were prepared at lab-scale (i.e. 25 mm in diameter, Fig. 3). Microscopic observations of the cells before testing show a fully densified electrolyte (density higher than 95%, nearly 50 μ m thick) which is well stuck on a homogeneous biphasic anode (Fig. 4). Figs. 5 and 6 also confirm that good planarity and crack-free assemblies can be obtained by simple manual deposition.

The up-scaling of cell fabrication consisted in improving the size of samples while using the same co-pressing method (Fig. 5). The microstructure remained the same (Figs. 4 and 6) and planar and crack-free assemblies have been also obtained.

The repartition of each element of 40 mm cells has been checked by Energy-dispersive X-ray spectroscopy mapping (Fig. 7). All elements are well dispersed and there is no evidence of nickel in the electrolyte. All the observation on 40 mm cells shows that the first up-scaling step was a success.

Up-scaling the size up to 80 mm in diameter brings more mechanical and technical problems, the most critical step being the deposition of the electrolyte: the bigger the cell is, the less the thickness of the electrolyte is homogeneous. Experience proved

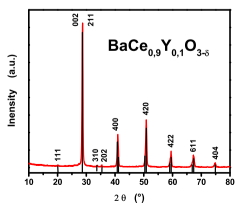


Fig. 1. XRD pattern for the $BaCe_{0,9}Y_{0,1}O_{3-\delta}$ powder.

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