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Implications of Cation Interdiffusion between Double Perovskite Cathode and Proton-Conducting Electrolyte for Performance of Solid Oxide Fuel Cells

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Abstract: Chemical compatibility and cation interdiffusion between the double perovskite cobaltites $RBaCo_2O_{6-\delta}$ (R = Gd, Pr) and proton-conducting electrolyte $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ were studied. Chemical interaction was found to occur already at 1100 °C as a result of the partial dissolution of $RBaCo_2O_{6-\delta}$ (R = Gd, Pr) in $BaZr_{0.8}Y_{0.2}O_{3-\delta}$. Analysis of the element distribution along the cross sections of diffusion couples $RBaCo_2O_{6-\delta}$ (R = Gd, Pr) | $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ showed strong interdiffusion of cations, with cobalt being the most mobile one. Its diffusion depth in the electrolyte reaches up to several hundreds of micrometers. The addition of NiO as a sintering aid to $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ promotes cation diffusion especially through the grain boundary mechanism, increasing the diffusion depth of Co. The possible implications of cation interdiffusion on the performance of proton-conducting SOFCs are discussed based on the results obtained.

Keywords: solid oxide fuel cells; cation interdiffusion; double perovskites; cathode materials; proton-conducting solid electrolyte

1. Introduction

Solid oxide fuel cells (SOFCs) are solid-state electrochemical devices that convert the chemical energy of the fuel (hydrogen, natural gas and other hydrocarbons) directly into electrical energy and possess high energy conversion efficiency and low environmental impact [1-3]. At the same time, a number of problems impede large-scale applications of SOFCs, among them high-temperature degradation processes, chemical compatibility of cell components and the high cost of materials [1-4]. The lowering of the operating temperature is the general way to promote the commercial application of SOFCs because it broadens the range of compatible construction materials, prolongs their lifetime and, consequently, reduces the cost of energy production. In this respect, proton-conducting solid oxide fuel cells (H-SOFCs) are generally considered to be better suited for practical applications due to lower operating temperatures and the absence of fuel contamination. In turn, conclusions on the chemical compatibility of materials are usually made based only on a costly and time-consuming trial-and-error method: heating a physical mixture of different powders at high temperature for several hours followed by analyzing its phase composition using X-ray diffraction (XRD). Such an approach may lead to contradictory results because of the kinetic limitations of solid-state interaction and non-zero detection limits of conventional XRD. In this respect, thermodynamic analysis of materials' compatibility is completely free from such limitations and, probably, represents the only way of unambiguous interpretation of compounds' stability under particular conditions.

The choice of thermodynamically stable and chemically compatible materials is a necessary but not sufficient condition for successful operation of SOFCs, because solid-state materials exposed to a thermodynamic potential gradient (gradient of temperature,



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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/). chemical potential of elements, electrical potential) are subjected to kinetic degradation governed by diffusion processes [5]. The applied gradients induce directed fluxes of the mobile components and lead to three basic degradation phenomena of materials: (i) kinetic demixing, (ii) kinetic decomposition and (iii) morphological instability [5]. Thus, in SOFCs the different chemical natures of components and high operating temperatures (600–1000 °C) inevitably lead to cation interdiffusion and continuous kinetic degradation of materials and, therefore, determine the lifetime of the devices [4].

Obviously, understanding the diffusion processes at the electrode/electrolyte interfaces in SOFCs is of critical importance for their successful large-scale commercial application [4]. Unfortunately, this important information is mostly lacking, and only a limited number of studies of cation diffusion phenomena in perovskite oxides may be found in the literature. Thus, Ca- and Sr-doped LaCrO₃, and Sr- and Mg-doped LaGaO₃ were investigated by tracer diffusion method [6–8]. Cation interdiffusion in the systems LaFeO₃, LaCoO₃, La_{0.9}Sr_{0.1}NiO₄, La₂Ni_{0.8}Cu_{0.2}O₄ and La₂₇W₅O_{55.5} was studied by diffusion couple method [9–13]. The kinetic demixing process was studied only for $La_{0.5}Sr_{0.5}Fe_{0.5}Co_{0.5}O_{3-\delta}$ and $La_{0.3}Sr_{0.7}CoO_{3-\delta}$ [14,15]. Comprehensive investigation was carried out for $La_2NiO_{4+\delta}$. a Ruddlesden–Popper phase, including determination of the self-diffusion coefficient of Ni²⁺ [16], chemical tracer diffusion [17], interdiffusion coefficients for the A- and B-site cations [18], as well as kinetic decomposition of La_2NiO_{4+d} under an oxygen chemical potential gradient [19]. As for the interdiffusion between the components of SOFCs, to the best of our knowledge this has been studied only for LSCF|GDC|YSZ interfaces, where Gd-doped ceria (GDC) acts as a buffer layer between the $La_{1-x}Sr_xCo_{1-y}Fe_yO_{3-\delta}$ (LSCF) cathode and yttria-stabilized zirconia (YSZ) electrolyte [20,21]. In turn, for protonconducting SOFCs, such studies have not yet been performed.

The thermodynamic analysis of the stability of the double perovskite $PrBaCo_2O_{6-\delta}$ under different conditions has recently been carried out by us [22]. As a result, its instability against chemical interaction with CeO_2 was found. The diffusion couple $(PrBaCo_2O_{6-\delta}(PBC)-SDC)$ experiment showed interdiffusion of Pr and Sm and chemical interaction between PBC and SDC with the formation of barium cerate as a product, in full agreement with thermodynamic calculations. Both phenomena were shown to affect significantly both thermal expansion and total conductivity of the composite electrodes and lead to higher than expected TECs of the composites. Thus, the potential problems with cation interdiffusion have already been outlined [22]. Formation of barium cerate on the border between PBC and SDC was also shown to cause a significant drop in the composites' total conductivity and should be detrimental for the long-term stability of the SOFC cell. One can expect that formation of BaCO₃ due to the chemical reaction of PBC with CO_2 from the ambient air, as predicted by thermodynamic calculations, will lead to lowering of the SOFC's performance as well. The PBC double perovskite was shown to be more suitable as an electrode for proton-conducting solid oxide fuel cells because of its thermodynamic stability against chemical interaction with barium-cerateand zirconate-based electrolytes [22].

In addition, the influence of the sintering aids such as, for example, widely employed NiO on the electrode–electrolyte compatibility and interdiffusion is completely unknown.

Therefore, the priority aims of the present work were (i) to investigate the chemical compatibility of double perovskite cathodes $RBaCo_2O_{6-\delta}$ (R = Gd, Pr) with proton-conducting solid electrolyte $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ +NiO (0, 1 wt.%) and (ii) to study interdiffusion processes at the interfaces $RBaCo_2O_{6-\delta}$ (R = Gd, Pr) | $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ +NiO (0, 1 wt.%).

2. Experimental Procedure

Powder samples of the double perovskites $PrBaCo_2O_{6-\delta}$ (PBC) and $GdBaCo_2O_{6-\delta}$ (GBC) and the solid electrolyte $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ (BZY0.2) were prepared by means of a glycerol–nitrate method using Gd_2O_3 , Pr_6O_{11} , Y_2O_3 , $BaCO_3$, Co and $Zr(OH)_2CO_3 \cdot xH_2O$ as starting materials. The purity of all the materials used was 99.99% (wt.).

 $Zr(OH)_2CO_3 \cdot xH_2O$ was preliminarily dissolved in concentrated HNO₃ (purity 99.99%), and the Zr concentration in the resulting solution was determined by the gravimetric method. The as-obtained solution with known concentration of Zr was used as a precursor for the solid electrolyte synthesis.

For preparation of $BaZr_{0.8}Y_{0.2}O_{3-\delta}$, the stoichiometric quantities of $BaCO_3$ and Y_2O_3 were dissolved in concentrated HNO₃. This solution was mixed with the $Zr(OH)_2CO_3 \cdot xH_2O$ solution described above, and a stoichiometric amount of glycerol (99% purity) was added as a complexing agent and a fuel. The required weight of glycerol was calculated according to full reduction of the nitrates to molecular N₂. The as-obtained solution was heated continuously at 100 °C till complete water evaporation and pyrolysis of the dried precursor. The resulting ash was subsequently calcined at 1100 °C for 24 h in air and then pressed into pellets (10 mm in diameter at 150 MPa) and sintered at 1550 °C for 12 h in air. A composite material of $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ with 1 wt.% of NiO as a sintering aid was made by physical mixing of the components in an agate mortar in an ethanol medium.

For synthesis of the $RBaCo_2O_{6-\delta}$ (R = Gd, Pr) powders, a stoichiometric mixture of the appropriate starting materials was dissolved in concentrated HNO₃, and glycerol (99% purity) was added to the solution. The weight of glycerol was calculated as described above in the case of $BaZr_{0.8}Y_{0.2}O_{3-\delta}$. As-prepared solutions were heated continuously at 100 °C till complete water evaporation and pyrolysis of the dried precursor. To yield the $RBaCo_2O_{6-\delta}$ (R = Gd, Pr) powders, the product of pyrolysis was subsequently calcined at 900, 1000 and 1100 °C for 12 h in air followed by thorough regrinding in an ethanol medium in an agate mortar after each annealing step.

The phase composition of the as-synthesized samples was studied by means of X-ray diffraction (XRD) with an XRD 7000 diffractometer (Shimadzu, Kyoto, Japan) using Cu K α radiation (λ = 1.5418 Å). The phase identification was performed using the PDF–2 2021 database [23] and Match! software [24]. The XRD patterns demonstrating phase purity of all the as-prepared oxides are presented in Figure S1.

The refinement of the lattice parameters was performed using the Le Bail method as implemented in Rietica 4.0 software [25]. The refined unit cell parameters of the starting double perovskite and electrolyte powders are given in Table S1.

The chemical compatibility of $RBaCo_2O_{6-\delta}$ (R = Gd, Pr) and $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ was studied by annealing the corresponding powder mixtures (50:50 wt.%) at different temperatures in the range 1100–1250 °C for 12 h in air with subsequent phase identification by the XRD.

Ceramic pellets (Ø10 mm × 3 mm) for cation interdiffusion experiments were prepared by dry pressing at 150 MPa and sintering for 12 h in air at 1240, 1250 and 1550 °C for PrBaCo₂O_{6- δ}, GdBaCo₂O_{6- δ} and BaZr_{0.8}Y_{0.2}O_{3- δ}+NiO (0, 1 wt.%), respectively. Density of the ceramic samples was estimated by the Archimedes method using water as a medium. The relative densities of the as-prepared pellets were 95% for GdBaCo₂O_{6- δ}, 97% for PrBaCo₂O_{6- δ}, 92% for BaZr_{0.8}Y_{0.2}O_{3- δ} and 95% for BaZr_{0.8}Y_{0.2}O_{3- δ}+NiO (1 wt.%). One surface of each pellet was polished down to 1 µm using diamond suspension. Then it was purified ultrasonically by means of an IL 10 ultrasonic generator (INLAB, Saint Petersburg, Russia) in an ethanol medium and dried at 70 °C in a drying oven in air atmosphere.

Cation interdiffusion was studied by the diffusion couple method. The following diffusion couples were studied: $GdBaCo_2O_{6-\delta}-BaZr_{0.8}Y_{0.2}O_{3-\delta}$, $GdBaCo_2O_{6-\delta}-BaZr_{0.8}Y_{0.2}O_{3-\delta}+NiO$ (1 wt.%), $PrBaCo_2O_{6-\delta}-BaZr_{0.8}Y_{0.2}O_{3-\delta}$ and $PrBaCo_2O_{6-\delta}-BaZr_{0.8}Y_{0.2}O_{3-\delta}+NiO$ (1 wt.%). Before annealing, each diffusion couple was pressed in a special device and fixed with a clamping mechanism. For each system, two independent experiments were performed, both at 1100 °C for 20 h and at 1200 °C for 48 h in air. After annealing, diffusion couples were placed in a silicon form, fixed with epoxy resin and cut perpendicular to the reaction interface. Then cross section of the reaction zone was polished down to 1 μ m using diamond suspension, purified ultrasonically in distilled water and dried at 70 °C in a drying oven in air atmosphere. The microstructure and element distribution across the couples' sections were examined by means of scanning electron microscopy (SEM) using a VEGA

3 microscope (Tescan, Brno, Czech Republic) equipped with an Ultim Max 40 (Oxford Instruments, Abingdon, UK) detector for energy dispersive X-ray spectroscopy (EDX).

Additionally, diffusion couples with porous layers of $RBaCo_2O_{6-\delta}$ (R = Gd, Pr) on top of the dense ceramic pellet of $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ were studied to mimic the behavior of SOFCs' cathodes. These diffusion couples were prepared by screen-printing the cathode paste on the polished surface of the ceramic electrolyte pellet. The cathode paste was obtained by thoroughly mixing the powder of double perovskite cobaltite with a solution of polyvinyl butyral (5 wt.%) in ethanol. The as-prepared diffusion couple was annealed at 1100 °C for 6 h in air. The treatment of the reaction zone and subsequent investigation by SEM were performed as described above for the other diffusion couples.

3. Results and Discussion

3.1. Chemical Compatibility

The results of chemical compatibility experiments are shown in Figures 1 and S2. At a first glance, XRD patterns of the sample mixtures look very similar, at least, at temperatures lower than 1100 °C. Such XRD patterns can be readily found in the current literature along with a typical conclusion that they indicate the absence of chemical interaction between the components of the mixture investigated. However, the XRD pattern of the GdBaCo₂O_{6- δ} + $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ mixture calcined at the highest studied temperature (see Figure 1) clearly indicates the formation of a new compound isostructural to Gd_2O_3 (possibly doped by Y_2O_3). Furthermore, the intensities of the peaks corresponding to the components of the mixtures as well as peaks' positions (and, hence, lattice parameters) depend on the calcination temperature as seen in Figure 1c,d. For example, the intensities of the peaks corresponding to double perovskites $RBaCo_2O_{6-\delta}$ (R = Gd, Pr) strongly decrease with calcination temperature. At the same time, XRD peaks of $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ slightly increase in intensity and shift towards high $2\Theta^\circ$ angles. Similar changes can be observed in the XRD patterns of the mixtures containing NiO; see Figure S2. All these evidences undoubtedly indicate in favor of chemical interaction between double perovskites and a solid electrolyte which occurs already at 1100 °C and likely at lower temperatures. The XRD results seem to suggest at least partial dissolution of $RBaCo_2O_{6-\delta}$ (R = Gd, Pr) in $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ oxide. Therefore, the similarity of the XRD patterns of the starting mixture (denoted as 25 °C in Figure 1) and the one calcined at 1100 °C cannot be used as evidence of the absence of chemical interaction.

The refined unit cell parameters of $RBaCo_2O_{6-\delta}$ (R = Gd, Pr) and $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ after annealing the powder mixtures (50:50 wt.%) are shown in Figure 2.

As seen in Figure 2a,b, lattice parameters of $RBaCo_2O_{6-\delta}$ (R = Gd, Pr) oxides, first slightly increase but then decrease with increasing calcination temperature. This can be explained by the combined effect of two factors: (i) variation of the $RBaCo_2O_{6-\delta}$ oxygen content, $6-\delta$, in a wide range [26,27] with temperature and the sample's prehistory and (ii) partial substitution of the smaller Y³⁺ cation for larger Gd³⁺ and Pr³⁺ [28] in $RBaCo_2O_{6-\delta}$ due to cation interdiffusion between components of the sample mixture.

As for $BaZr_{0.8}Y_{0.2}O_{3-\delta}$, its unit cell parameter, as seen in Figure 2c, decreases with calcination temperature and more strongly in contact with $GdBaCo_2O_{6-\delta}$ oxide. This can be, at least qualitatively, explained assuming cobalt dissolution in $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ leads to lattice contraction as shown in [29], whereas dissolution of Gd and Pr, on the contrary, facilitates the lattice expansion of barium zirconate due to differences in the cation radii [28].

3.2. Analysis of Diffusion Couples

The results of SEM and EDX analysis of the cross sections of diffusion couples $RBaCo_2O_{6-\delta}$ (R = Gd, Pr)– $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ +NiO (0, 1 wt.%) after annealing are presented in Figures 3–5. They support the conclusions drawn above and based on the results of chemical compatibility study by XRD. As seen in Figures 3–5, cation interdiffusion can be observed, with Co being the most mobile cation demonstrating pronounced diffusion into the electrolyte pellet for all the couples studied.



Figure 1. XRD patterns of the powder mixtures $RBaCo_2O_{6-\delta}-BaZr_{0.8}Y_{0.2}O_{3-\delta}$ (50:50 wt.%) annealed at different temperatures for 12 h in air: (**a**,**c**) R = Gd; (**b**,**d**) R = Pr.

The analysis of the cross section of the GdBaCo₂O_{6- δ}-BaZr_{0.8}Y_{0.2}O_{3- δ} diffusion couple annealed at 1200 °C for 48 h in air, as seen in Figure 3a, reveals the formation of Gd₂O₃ on the surface of the GdBaCo₂O_{6- δ} pellet along the interface of the diffusion couple. This is obviously in agreement with the XRD results discussed above. In addition, slight interdiffusion (depth ~15 µm) of Gd³⁺ ions in the BaZr_{0.8}Y_{0.2}O_{3- δ} electrolyte and Y³⁺ into the GdBaCo₂O_{6- δ} phase was detected. The diffusion of Co ions occurs mainly through the bulk of the electrolyte. The diffusion depth reaches 300 µm. Cobalt concentration in the electrolyte pellet is practically constant at a distance of up to 30 µm from the interface with GdBaCo₂O_{6- δ}. The average chemical composition of this electrolyte layer was estimated as Ba_{1.02}Zr_{0.76}Y_{0.16}Co_{0.06}O_{3- δ}.

In the case of the PrBaCo₂O_{6-δ}-BaZr_{0.8}Y_{0.2}O_{3-δ} couple annealed at 1200 °C for 48 h in air, significant diffusion of both Co and Pr into the BaZr_{0.8}Y_{0.2}O_{3-δ} pellet was found, as shown in Figure 4a. Their diffusion occurs both along the grain boundaries and through the bulk of BaZr_{0.8}Y_{0.2}O_{3-δ}. The former mechanism is dominating. The diffusion depth of Co reaches 560 µm. The analysis of the grain boundary phase's chemical composition gives near the interface PrBaCo₂O_{6-δ} | BaZr_{0.8}Y_{0.2}O_{3-δ} the following averaged composition: Ba_{1.02}Co_{0.72}Y_{0.21}Pr_{0.09}O_{3-δ}. However, the composition of the grains' interior may be expressed as Ba_{0.99}Zr_{0.69}Y_{0.16}Co_{0.13}Pr_{0.03}O_{3-δ}.



Figure 2. The unit cell parameters vs. annealing temperature of powder mixtures $RBaCo_2O_{6-\delta}$ (R = Gd, Pr)– $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ (50:50 wt.%): (**a**) $GdBaCo_2O_{6-\delta}$, (**b**) $PrBaCo_2O_{6-\delta}$ and (**c**) $BaZr_{0.8}Y_{0.2}O_{3-\delta}$.

The addition of NiO as a sintering aid to the BaZr_{0.8}Y_{0.2}O_{3- δ} solid electrolyte strongly promotes interdiffusion especially along the grain boundaries, as can be observed in Figures 3b, 4b and S3. As a result, for example, the depth of Co diffusion from PrBaCo₂O_{6- δ} to BaZr_{0.8}Y_{0.2}O_{3- δ} is almost doubled, up to 1000 µm when 1 wt.% of NiO is added to the electrolyte.

The diffusion couple GdBaCo₂O_{6- δ}-BaZr_{0.8}Y_{0.2}O_{3- δ}+NiO (1 wt.%) annealed at 1200 °C for 48 h in air does not show the formation of Gd₂O₃ as a reaction product, contrary to the couple without NiO. Instead, diffusion of Gd³⁺ into the electrolyte phase occurs mainly through the grain boundary mechanism, as seen in Figure 3b. Similar behavior is evident for diffusion of Co ions. As a result, the formation of Ba_{1.03}Co_{0.81}Ni_{0.03}Gd_{0.13}O_{3- δ} oxide was detected on the grain boundaries of the electrolyte pellet near the interface with double perovskite, whereas the composition of the electrolyte grains' bulk corresponds to Ba_{1.02}Zr_{0.78}Y_{0.16}Co_{0.04}O_{3- δ}. In addition, the diffusion of Ni²⁺ ions from the electrolyte pellet to the GdBaCo₂O_{6- δ} phase was observed.





Figure 3. SEM images and concentration maps of reaction zones for diffusion couples after annealing at 1200 °C for 48 h in air: (a) GdBaCo₂O_{6- δ}-BaZr_{0.8}Y_{0.2}O_{3- δ} and (b) GdBaCo₂O_{6- δ}-BaZr_{0.8}Y_{0.2}O_{3- δ}+NiO (1 wt.%); BSE (back scattered electrons) image and element maps were obtained near the phase boundary of the diffusion couple. The red arrow marks the interface between the pellets.



Figure 4. SEM images and concentration maps of reaction zones for diffusion couples after annealing at 1200 °C for 48 h in air: (**a**) PrBaCo₂O_{6- δ}-BaZr_{0.8}Y_{0.2}O_{3- δ}; BSE image and element maps obtained near the phase boundary of diffusion couple and (**b**) PrBaCo₂O_{6- δ}-BaZr_{0.8}Y_{0.2}O_{3- δ}+NiO (1 wt.%); BSE image and element maps obtained in the area of Co diffusion maximum in the depth of about 500 µm from the phase boundary. The corresponding images near the phase boundary of the diffusion couple may be observed in Figure S3. The red arrow marks the interface between the pellets.



Figure 5. Concentration profiles of diffusion couples after annealing at 1200 °C for 48 h in air: (a) GdBaCo₂O_{6- δ}-BaZr_{0.8}Y_{0.2}O_{3- δ}, (b) PrBaCo₂O_{6- δ}-BaZr_{0.8}Y_{0.2}O_{3- δ}, (c) GdBaCo₂O_{6- δ}-BaZr_{0.8}Y_{0.2}O_{3- δ}, (c) GdBaCo₂O_{6- δ}-BaZr_{0.8}Y_{0.2}O_{3- δ}+NiO (1 wt.%) and (d) PrBaCo₂O_{6- δ}-BaZr_{0.8}Y_{0.2}O_{3- δ}+NiO (1 wt.%).

The largest diffusion depth, reaching 1000 μ m, was found in the PrBaCo₂O_{6-δ}– BaZr_{0.8}Y_{0.2}O_{3-δ}+NiO (1 wt.%) couple annealed at 1200 °C for 48 h in air. Again, the diffusion of Pr and Co occurs through both grain boundaries and bulk of the electrolyte phase with the former mechanism being the dominating one. Interestingly, it seems in this case the cation diffusion in the electrolyte pellet so strongly depends on the presence of NiO that near the interface, where the concentration of Ni is relatively lower than that in the bulk of the electrolyte pellet due to nickel diffusion into the double perovskite pellet, a layer depleted in Co and Pr is formed. The maximum of Co concentration is, thus, shifted deep into the electrolyte pellet at a distance of 400–540 μ m from the interface with double perovskite. In this area, two types of Co-containing compounds were observed. The one with an approximate composition Y_{0.70}Pr_{0.24}Ba_{0.94}Co_{1.76}Ni_{0.12}O_{6-δ} was detected on the grain boundaries (Figure 4b). The other is a mixed cobalt–nickel oxide with Co to Ni ratio 2.04:0.9, which forms particles with a size of about 2–5 μ m, as seen in Figure 4b. The composition of the bulk of the electrolyte's grains in the area of maximum cobalt content corresponds on average to Ba_{1.01}Zr_{0.80}Y_{0.16}Co_{0.03}O_{3-δ}.

The averaged (over the cross section area) concentration profiles illustrating the abovementioned distribution of different species in the diffusion zone of the $RBaCo_2O_{6-\delta}$ (R = Gd, Pr) | $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ +NiO (0, 1 wt.%) couples annealed at 1200 °C for 48 h in air are shown in Figure 5.

Decreasing the annealing temperature of all the diffusion couples discussed above to 1100 °C does not change the general view of the diffusion zone except that the diffusion depth roughly becomes two times less than that at 1200 $^{\circ}$ C. In the case of porous $RBaCo_2O_{6-\delta}$ (*R* = Gd, Pr) double perovskite layers annealed on the surface of the pure $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ electrolyte pellet at 1100 °C for 6 h in air, i.e., under conditions which can be used to produce a cathode layer in the SOFC, the diffusion depth was found to be equal to ~50 µm irrespective of the nature of rare-earth element. Taking into account the typical thickness of the electrolyte layer of the order of $10-50 \ \mu\text{m}$, the results of this work give a clear indication that for the time necessary for the preparation of the cathode layer, cobalt from the double perovskite seems to be able to reach the SOFC's anode, diffusing through the electrolyte layer. Since the leading diffusion mechanism is the grain boundary one, all the electrolyte grains in that case would be covered with Co-rich compounds. This would potentially lead to electrolyte breakage due to cobalt cations' reduction by hydrogen from the anode compartment. Hence, the cation interdiffusion puts strong technological restrictions on the use of cobaltites in proton-conducting SOFCs with $BaZr_{1-x}Y_xO_{3-\delta}$ -based electrolytes. Methods must be found to prevent or at least reduce the diffusion of cobalt into the electrolyte by, for example, using some buffer layer; alternatively, other cathode materials that are less prone to similar diffusion problems should be identified.

In any case, the results of our work convincingly point to the crucial necessity of performing similar studies with other potential cathode and electrolyte materials of protonconducting SOFCs. Furthermore, the NiO-sintering aid, as discussed above, seems to promote the grain boundary diffusion in the electrolyte layer, hence, the quantity of NiO should be limited and the influence of other sintering aids on diffusion in the system electrode | electrolyte should be studied.

4. Conclusions

Chemical compatibility and cation interdiffusion between the double perovskites $RBaCo_2O_{6-\delta}$ (R = Gd, Pr) and proton-conducting electrolyte $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ were studied by XRD and diffusion couples methods. Both techniques gave consistent results indicating that chemical interaction between the double perovskites and the solid electrolyte occurs already at 1100 °C and, likely, at lower temperatures. The partial dissolution of $RBaCo_2O_{6-\delta}$ (R = Gd, Pr) in $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ oxide was found. Element distribution maps obtained through the cross sections of diffusion couples showed strong interdiffusion of cations, with cobalt being the most mobile cation possessing in some instances the diffusion depth in the electrolyte up to several hundreds of micrometers. In the case of the GdBaCo₂O_{6- δ} | BaZr_{0.8}Y_{0.2}O_{3- δ} couple without the NiO-sintering aid, Co diffusion occurs through the bulk of the electrolyte phase, whereas a layer of Gd_2O_3 forms on the interface with double perovskite. Yet, in the presence of NiO, both Gd and Co diffuse from GdBaCo₂O_{6- δ} inside the electrolyte pellet with the grain boundary mechanism as the main one. In turn, when $PrBaCo_2O_{6-\delta}$ is in contact with $BaZr_{0.8}Y_{0.2}O_{3-\delta}$, the diffusion of both Pr and Co in the electrolyte pellet takes place irrespective of NiO addition. The diffusion occurs through both bulk and grain boundary mechanisms, with the latter as the leading one. The diffusion depth in the presence of the NiO-sintering aid is almost doubled, pointing to the necessity to control the content of NiO in the electrolyte as well as to search for other sintering aids. The observed strong diffusion of cobalt through the $BaZr_{0.8}Y_{0.2}O_{3-\delta}$ layer reveals a potential problem for the long-term stability of SOFCs and, thus, must be understood and controlled as much as possible.

Supplementary Materials: The following supporting information can be downloaded at: https: //www.mdpi.com/article/10.3390/en16072980/s1. Figure S1. XRD patterns of the as-prepared oxides: (a) GdBaCo₂O_{6- δ}, (b) PrBaCo₂O_{6- δ}, (c) BaZr_{0.8}Y_{0.2}O_{3- δ}; Table S1. Refined unit cell parameters of the as-prepared oxides; Figure S2. XRD patterns of the powder mixtures (50:50 wt.%) RBaCo₂O_{6- δ}-BaZr_{0.8}Y_{0.2}O_{3- δ} +NiO (1 wt.%) annealed at different temperatures for 12 h in air: (a) *R* = Gd, (b) *R* = Pr; Figure S3. BSE image and concentration maps near the phase boundary of the $\label{eq:prback} PrBaCo_2O_{6-\delta}-BaZr_{0.8}Y_{0.2}O_{3-\delta}+NiO~(1~wt.\%)~diffusion~couple~after~annealing~at~1200~^\circC~for~48~h~in~air.$

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