Samples of $\text{Ba}_{1-x}\text{Sr}_x\text{Ce}_{0.9}\text{Y}_{0.01}\text{O}_{3-\delta}$, $0 < x < 0.1$, with Improved Chemical Stability in $\text{CO}_2$-$\text{H}_2$
Gas-Involving Atmospheres as Potential Electrolytes for a Proton Ceramic Fuel Cell

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1. Introduction

One promising group of ceramic proton conductors is composed of perovskite ($\text{ABO}_3$)-based oxides. The $\text{Y}_2\text{O}_3$-doped zirconates $\text{BaZr}_{1-x}\text{Y}_x\text{O}_{3-\delta}$ or cerates $\text{BaCe}_{1-x}\text{Y}_x\text{O}_{3-\delta}$, where $0 < x < 0.3$,
are currently considered to be suitable materials for ceramic proton-conducting fuel cells operating at reduced temperatures. Ceramic proton-conducting fuel cells (PCFCs) and solid oxide fuel cells (SOFCs) are being intensively studied with the aim of constructing power sources which can be operated within a temperature range of 400–700 °C [1,2]. Ceramic proton conductors can be applied to the construction of other electrochemical devices important for hydrogen infrastructures. Classic electrochemical devices for this technology include solid oxide electrolyzers, hydrogen or hydrocarbon sensors, hydrogen units for gas purification processes, and reactors for the hydrogenation of compounds [3–5].

Generally, Y2O3-doped cerates exhibit higher levels of ionic conductivity than BaZr1−xYxO3−δ. The limited chemical stability of BaCe0.9Y0.1O3−δ-based materials in H2O and CO2 gas atmospheres is a crucial factor limiting their practical utilisation in fuel cells supplied by rich hydrogen processing gases. Ba(OH)2, BaCO3, and CeO2 have been observed as secondary phases appearing as a result of the decomposition of yttrium-doped BaCeO3 in reaction with CO2 [6–10]. Multicomponent solid solutions in which ceria was partially replaced by zirconia in Ba(Ce1−xZrx)Y0.9O3−δ exhibited greater resistance to corrosion from CO2 attacks. These materials were successfully applied in IT-SOFCs in which power density involving BaCeO3-based proton-conducting membranes varied from 30 to 1400 mW/cm² [11–16].

In the near future, increased use of hydrogen is expected in the fuel-energy sector as well as in various areas of transport. Hydrogen is most often produced from fossil fuels by means of various technologies in which feed streams involving hydrogen, prior to their entry into the PSA cleaning unit, include some impurities, such as CO, CO2, N2, Ar, and CH4. The level of CO2 may range from 1.5 to 10 vol.%. Some efforts have also been made to obtain biohydrogen by means of various biological, biotechnological, and renewable energy technologies [17–20].

The elaboration of either new chemical compositions of ceramic proton-conducting materials, obtained by various processing methods, which are highly tolerant to contamination in hydrogen, or of ways to improve the chemical stability of BaCeO3-based electrolytes in a CO2 gas atmosphere, is required for the continued development of hydrogen technologies. It was found that the addition of Ba3(PO4)2 BaWO4 to BaCe0.9Y0.1O3−δ improved resistance to CO2 corrosion; however, a deterioration in electrical properties was observed instead [21–23].

Structural modification via partial substitution of the hosting A ion in ABO3 by other ions with the same valence but different ionic radii may also be a way to improve protonic conductivity as well as to reduce chemical reactivity with impurities such as H2O or CO2 in processing gases.

ABO3 oxides exhibit greater tilts when the A site is characterized by a smaller ion or when the B site is doped with increasing concentrations of M3+ (Y, Sm, Gd) dopant [24–26]. In the previous paper [27] authors found that Ba0.95Ca0.05Ce0.9Y0.1O3−δ exhibited improved corrosion resistance in CO2–H2 gas atmosphere compared to BaCe0.9Y0.1O3−δ. It was also reported that strontium-modified Ba0.9Sr0.1Ce0.8Y0.2O3−δ was characterized by improved chemical stability in H2O gas atmospheres compared to BaCe0.9Y0.1O3−δ. The variation in the electrical conductivity of these materials has not been clearly explained [28,29].

The aim of the present study was to determine the effect of the concentration of SrO in Ba1−xSrxCe0.9Y0.1O3−δ, where 0 < x < 0.1, on its electrical and electrochemical properties as well as on its chemical resistance to attacks of CO2 corrosion. Another particular goal was the acquisition of a modified ceramic proton conductor characterized by improved chemical stability in a CO2 gas atmosphere and by electrical conductivity values comparable to the level of BaCeO3-based samples.

2. Experimental

2.1. The Preparation of Powder and Sintered Samples of Ba1−xSrxCe0.9Y0.1O3−δ, 0 < x < 0.1

Powders from SBCY, i.e., Ba1−xSrxCe0.9Y0.1O3−δ, 0 < x < 0.1, were synthesized via solid-state reaction. The starting reagents were barium carbonate and strontium carbonate (Avantor Performance Materials Poland S.A, Gliwice, Poland, 99.999%), yttrium (III) oxide (Sigma-Aldrich,
St. Louis, MO, USA, 99.99%), and cerium(IV) oxide (Acros Organics, Geel, Belgium 99.9%). Stoichiometric amounts of reagents corresponding to the chemical composition of series of samples of BaCe$_{0.9}$Y$_{0.1}$O$_{3–\delta}$ (BCY), Ba$_{0.95}$Sr$_{0.05}$Ce$_{0.9}$Y$_{0.1}$O$_{3–\delta}$ (2SBCY), Ba$_{0.9}Sr$0.05Ce$_{0.9}$Y$_{0.1}$O$_{3–\delta}$ (5SBCY) and Ba$_{0.9}$Sr$_{0.1}$Ce$_{0.9}$Y$_{0.1}$O$_{3–\delta}$ (10SBCY) were weighed and homogenised. In the SBCY series of samples, the concentration of strontium varied from 2 to 10 mol.%. The powdered chemical reagents were homogenized and milled in a planetary ball mill (Retsch PM 100) in dry ethanol (Avantor Performance Materials Poland S.A, 99.8%). Grinding balls (diameter ~5 mm) made from tetragonal yttria-zirconia (TZP) solid solutions were used in this operation. The milling process elaborated for the 5SBCY sample was applied.

A small portion of mixed reagents prepared for the SBCY-series powders was calcined within a temperature range of 850–1150 °C for 2 h. The monophase orthorhombic phase for all investigated compositions of the SBCY series was found at a temperature of 1150 °C. Subsequently, a 200 g portion of powder was synthesized at 1200 °C for 2 h. The SBCY powders were additionally ground in dry ethanol. The total milling time for monophase SBCY powder samples was 6 h. The details of the homogenisation procedure and milling conditions for the monophase samples was described in the previous paper [30]. The BSCY disc samples were formed via the isostatic pressing method, using a National Forge isostatic cold press, EPSI Engineered Pressure Systems International N.V. (Temse, Belgium). The samples were pressed in a mould with a diameter of 12 under 0.5 MPa. Once pressed, the samples were placed in plastic foil, subjected to outgassing, and pressed under 250 MPa. The samples were sintered at 1550 °C for 2 h in air.

2.2. Analytical Methods Used to Evaluate the Physicochemical Properties of the Series of SBCY Samples

X-ray diffraction (XRD) measurements were made using a PANalytical Empyrean diffractometer (Malvern, United Kingdom; CuK$_\alpha$; $\lambda$ = 0.15418 nm) equipped with a high-temperature measurement oven in order to determine the phase compositions of Ba$_{1-x}$Sr$_x$Ce$_{0.9}$Y$_{0.1}$O$_{3–\delta}$ powders and sintered samples. XRD patterns were collected in the heating-cooling cycle. The temperatures of XRD measurement varied within a range of 25–800 °C; a ramp of 10 °C min$^{-1}$ was applied in the heating-cooling cycle. Variations in the cell volumes of SBCY samples were calculated based on cell parameters, determined by means of the Rietveld method. In order to find the impact of the incorporation of strontium resulting in Ba$_{1-x}$Sr$_x$Ce$_{0.9}$Y$_{0.1}$O$_{3–\delta}$ on variations in its structure compared to the BaCe$_{0.9}$Y$_{0.1}$O$_{3–\delta}$ sample, the Goldschmidt tolerance factor (t), specific free volume (SFV), and global instability index (GII) were calculated using the SPuDS software package (2.18.06.19, Michael Lufaso, Jacksonville, FL, USA, dedicated to analysis of the structure of ABO$_3$-based oxides [27,31,32].

Raman measurements were also carried out for Ba$_{1-x}$Sr$_x$Ce$_{0.9}$Y$_{0.1}$O$_{3–\delta}$ samples. A Horiba Jobin Yvon LabRAM HR micro Raman spectrometer (Higashi-kanda, Chiyoda-ku, Tokyo, Japan) equipped with a CCD detector was used in this investigation. An excitation wavelength of 532 nm was used, with a beam intensity of approximately 10 mW. Acquisition time was set at 30 s [27].

Observations of cross sections as well as surfaces of all SBCY sintered samples were carried out by means of scanning electron microscopy. A Nova NanoSEM 200 (FEI, Eindhoven, Netherlands) coupled with an EDS system was used in these investigations. Comparative observations were made, both of initial SBCY powders, sintered pellets and of samples that had undergone an additional corrosion resistance test in a CO$_2$ gas atmosphere [33]. The thermograviometry (TG) method was used to determine mass variation for the series Ba$_{1-x}$Sr$_x$Ce$_{0.9}$Y$_{0.1}$O$_{3–\delta}$ in pure CO$_2$ gas. A thermobalance (Rubotherm DynTHERM 1100-40 MP-G analyser, TA Instruments, Bochum, Germany) was used in these investigations. TG curves were recorded within a temperature range of 25–1100 °C, at a pressure of 0.1 MPa (abs). Prior to these measurements, all SBCY samples were ground to powder; the powdered SBCY samples were then placed in an alumina crucible. The gas flow was 100 mL min$^{-1}$, the temperature ramp 10 °C min$^{-1}$. The final heating temperature of 1100 °C was maintained for 20 min. Tests of the corrosion resistance of SBCY sintered samples to CO$_2$ attack were carried out in a quartz reactor placed in an electric resistance furnace. The SBCY electrolytes were heated to 550 °C and
maintained at that temperature for 100 h. A gas mixture containing 2.6 or 5 vol.% of CO₂ in Ar was used in this experiment. The chemical resistance of BCY and SBCY samples in H2O-Ar was examined in analogous conditions. In these experiments, argon was passed first through a scrubber with water, then into the quartz reactor. The SBCY samples used for these experiments were of the same dimensions and thickness as those used in the investigations of electrical properties. Following the experiments, samples were subjected to X-ray investigations and SEM microstructural observations as well as electrical conductivity measurements, performed in both air and hydrogen gas atmospheres [27].

2.3. Electrical and Electrochemical Investigations of Ba₁₋ₓSrₓCe₀.9Y₀.1O₃₋δ Samples

SBCY cylindrical samples with two Pt electrodes prepared using Pt paste (Gwent Electronic Materials Ltd., Taipei, Taiwan) were investigated by means of electrochemical impedance spectroscopy (EIS) in the range 10–10¹⁰ Hz with a density of 54 points per decade and an amplitude of 0.1 VRMS in a two-electrode setup [34,35]. EIS measurements were performed in air from room temperature to 730 °C. Electrical conductivity measurements for SBCY samples were also performed in wet 5 vol.% H₂/Ar gas atmospheres or in wet air. The AC four-probe method was applied. The experiments were carried out within a temperature range of 400–700 °C.

The values of electromotive force (EMF) of the elaborated concentration cells (1):

\[ H_2 (I), Pt | SBCY | Pt, H_2 (II) \]  

(1)

were used to determine the ionic transport number. Pure humidified hydrogen and a humidified mixture of 10% H₂ in Ar (flow ~30 mL/min) were used as feeding streams. The experimental procedure was similar to that described in paper [36].

3. Results

All of the XRD diffraction patterns recorded for all of the SBCY powders and sintered samples (Figure 1a,b) at RT reflected the BCY monophase orthorhombic Pnma space group. In the case of SBCY samples with increased SrO content in Ba₁₋ₓSrₓCe₀.9Y₀.1O₃₋δ (Figure 1c), the diffraction peaks shifted towards greater diffraction angles in the recorded XRD patterns. A similar phenomenon was also mentioned by Wang et al. [28]. This suggests that decreases in the cell parameters of SBCY series samples should also be expected. The impact of temperature on possible phase composition changes was also investigated within a temperature range of 25–800 °C.

![Figure 1. XRD patterns collected for Ba₁₋ₓSrₓCe₀.9Y₀.1O₃₋δ (0 < x < 0.1) (SBCY) powders (a), sintered samples (b), and diffraction peak (211) shifts of SBCY samples toward greater diffraction angles (c) vs. chemical composition.](image-url)
The variation in the cell volume of $\text{Ba}_{1-x}\text{Sr}_x\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ samples vs. chemical composition is shown in Figure 2a. A gradual decrease was observed in cell volume vs. increase in SrO content in the $\text{Ba}_{1-x}\text{Sr}_x\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ samples. These results are also the consequences of previously observed shifts in diffraction peaks, in the case of SBCY samples, to angles with greater values in XRD diffraction patterns. Figure 2b presents the representative XRD patterns recorded for a 5SBCY sample during the heating-cooling cycle. No precipitation of impurities as additional phases or phase transitions of the orthorhombic phase of SBCY were found within this temperature range.

![Figure 2a](image)

**Figure 2.** Variation in the cell volume of $\text{Ba}_{1-x}\text{Sr}_x\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ vs chemical composition at RT (a); XRD patterns collected for $\text{Ba}_{0.95}\text{Sr}_{0.05}\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ (5SBCY) during the heating-cooling cycle within the temperature range 25–800 °C (b) variation in $\text{BaCe}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ (BCY) and 5SBCY cell volume vs temperature (c).

Figure 2c presents the dependence of the cell volumes of the BCY and 5SBCY samples during heating-cooling cycles; linear increases were noted.

The values of the cell volume of 5SBCY, lower than those recorded for BCY, are in close agreement with data reflecting comparison of the ionic radius of the Sr$^{2+}$ ion (r = 135 pm) with the greater radius of the Ba$^{2+}$ ion (r = 149 pm) it replaced.

Hayashi et al. [37] studied the correlation between structural parameters such as tolerance factor t and specific free volume SFV and variations in values of ionic oxide conductivity for a wide spectrum of ABO$_3$-based perovskite oxide-ion conductors. The influence of SFV on oxide-ion migration may have an indirect impact on proton migration as well as on oxide-proton conductivity in ABO$_3$-based materials. The authors of the cited paper found that, in order to obtain maximum ionic conductivity as a result of the balance between SFV and t, the optimum value of t was approximately 0.96. It was shown that high values of oxide ion conductivity could be expected for perovskite materials with SFV values higher than 0.4. Neither the structural parameters of $\text{Ba}_{1-x}\text{Sr}_x\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ samples nor t, SFV, GII, nor their impact on the samples’ physicochemical properties had been analyzed previously. Figure 3 presents the dependence of calculated values of such parameters as t, SFV, and GII vs chemical composition for $\text{Ba}_{1-x}\text{Sr}_x\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ samples.

The highest value of t was found for the $\text{BaCe}_{0.9}\text{Y}_{0.1}\text{O}_3$ sample. The partial replacement of BaO with SrO in $\text{Ba}_{1-x}\text{Sr}_x\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ led to a gradual decrease in the tolerance factor vs increased SrO content. This finding is also in close agreement with data reflecting the comparison of the ionic radius of the Sr$^{2+}$ ion (r = 135 pm) with the greater radius of the Ba$^{2+}$ ion (r = 149 pm) which it replaced. On the other hand, introduction of SrO resulting in $\text{Ba}_{1-x}\text{Sr}_x\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ led to higher values of SFV. According to data analyzed in the subject literature, an increase in the SFV coefficient is desirable for an improvement in ionic migration. The global instability index (GII) value is typically < 0.1 valence units (vu) for unstrained structures of ABO$_3$-based oxides, and as high as 0.2 vu in structures with induced strains. A GII value of approximately 0.2 vu has been found to be critical for several structures typically designated as unstable. The substitution of SrO for BaO in the range 0 < x < 0.1 in
Ba_{1-x}Sr_xCe_{0.9}Y_{0.1}O_{3-\delta} caused a small increase in GII, the values of which were lower than the expected crucial values.

Figure 3. Dependence of the t tolerance factor, specific free volume SFV, and global instability index GII vs strontium content in Ba_{1-x}Sr_xCe_{0.9}Y_{0.1}O_{3-\delta} samples.

Possible variations in the ABO$_3$ structure in sintered SBCY samples were also investigated, using Raman spectroscopy as complementary to X-ray diffraction analysis. The interpretation of the Raman spectra recorded for BCY and SBCY samples within a range of 400–50 cm$^{-1}$ is presented in Figure 4a–d. It was found that the obtained SBCY samples indicated the same orthorhombic phase, with the Pnma space group.

Figure 4. Raman spectra recorded for BCY (a), Ba$_{0.98}$Sr$_{0.02}$Ce$_{0.9}$Y$_{0.1}$O$_{3-\delta}$ (2SBCY) (b), 5SBCY (c), and Ba$_{0.9}$Sr$_{0.1}$Ce$_{0.9}$Y$_{0.1}$O$_{3-\delta}$ (10SBCY) (d) samples before and after additional heating (h) in 5 vol.% H$_2$O-Ar at 550 °C for 100 h.

Due to the similarity of the collected spectra, it can be unequivocally stated that the substitution of SrO for BaO resulting in Ba$_{1-x}$Sr$_x$Ce$_{0.9}$Y$_{0.1}$O$_{3-\delta}$ was accomplished within the same crystallographic structure. No additional chemical bands in the range 300–100 cm$^{-1}$, which would correspond to variations in ABO$_3$ crystal structure, were found in the SBCY series of samples. These findings enabled us to conclude that the introduced Sr$^{2+}$ ions were placed in the same positions as the original Ba$^{2+}$ ions in Ba$_{1-x}$Sr$_x$Ce$_{0.9}$Y$_{0.1}$O$_{3-\delta}$, 0 < x < 0.1. The crystallographic structure of SBCY sample series was the same as that of BaCe$_{0.9}$Y$_{0.1}$O$_{3-\delta}$. The Raman studies performed for SBCY samples were complementary to observations, which were made during the XRD investigations.

A similar situation was observed for a series of samples of CaO-modified BaCe$_{0.9}$Y$_{0.1}$O$_{3-\delta}$ resulting in Ba$_{0.95}$Ca$_{0.05}$Ce$_{0.9}$Y$_{0.1}$O$_{3-\delta}$ (5CBCY), whose electrolytic properties were described in the
paper [27]. As a result of the analysis of Raman spectra (Figure 4b,c), it can be stated that samples 2SBCY-h and 5SBCY-h, additionally heated in a H₂O-Ar gas atmosphere, were characterized by greater concentrations of point defects compared to the same samples (2SBCY; 5SBCY) prior to the heating experiments. This may also be attributable to the adsorption of H₂O from the gas atmosphere and interaction with oxygen vacancies, as evidenced by the significantly lower half-width of the bands on the spectra of the 2SBCY-h and 5SBCY-h samples. This was especially visible in the case of the 2SBCY-h sample [38–40].

Sintered pellets were prepared from SBCY grounded powders, which were also characterized. Based on XRD investigations (Figure 1a), crystalline size d_{(211)} was calculated for all powders from the SBCY series. It was found that the crystalline size d_{(211)} estimated from XRD patterns varied within a range of ~97 to ~106 nm. No significant impact of strontium content in Ba\(^{1–x}\)Sr\(^x\)Ce\(^{0.9}\)Y\(^{0.1}\)O\(_3–δ\) on the regular variation of crystalline sizes was observed. The SEM/EDS method was also used to investigate ground SBCY powders. The representative SEM recorded for 5SBCY powder is presented in Figure 5a. Based on SEM observation, the particle size falls within the range ~80 to ~400 nm. The agglomerate form of particles was observed in the investigated powders. Figure 5b presents results of chemical analysis performed on the SBCY sample, confirming its established chemical composition.

![Figure 5](image_url)

**Figure 5.** The representative SEM image recorded for 5SBCY powder (a) and SEM/EDS chemical analysis (b) performed for this sample.

Figure 6a–d presents the microstructures of the representative surfaces of sintered BCY, 2SBCY, 5SBCY, and 10SBCY samples. All of the sintered SBCY samples were characterized by fairly uniform microstructures and similar levels of closed porosity.

In the case of sintered BCY samples, the average grain size was ~4 μm. The introduction of SrO into Ba\(^{1–x}\)Sr\(^x\)(Ce\(^{0.9}\)Y\(^{0.1}\))O\(_3–δ\) solid solutions led to a slight gradual decrease in average grain size, to 2–3 μm.

Figure 7a presents relative mass (m/m₀) variation vs chemical composition recorded for all SBCY powdered samples. The TG curves were recorded in pure CO₂.
In the case of sintered BCY samples, the average grain size was ~4 \( \mu \)m. The introduction of SrO into \( \text{Ba}_{1-x}\text{Sr}_x(\text{Ce}^{0.9}\text{Y}^{0.1})\text{O}_{3-\delta} \) solid solutions led to a slight gradual decrease in average grain size, to 2-3 \( \mu \)m.

Figure 7a presents relative mass (\( m/m_o \)) variation vs chemical composition recorded for all SBCY powdered samples. The TG curves were recorded in pure CO\(_2\).

**Figure 6.** The microstructure of the representative surface of sintered (a) BCY, (b) 2SBCY, (c) 5SBCY, and (d) 10SBCY samples.

**Figure 7.** TG curves recorded for \( \text{Ba}_{1-x}\text{Sr}_x\text{Ce}^{0.9}\text{Y}^{0.1}\text{O}_{3-\delta} \) samples in a pure CO\(_2\) gas atmosphere, at temperatures ranging from 25 to –1100 °C (a); XRD pattern recorded for SBCY samples after heating in 5 vol.% CO\(_2\) in an Ar gas atmosphere at 550 °C for 100 h (b).

The greatest increase in relative mass (\( m/m_o \)) was determined in the case of the BCY sample. Sawant et al. [41] investigated the chemical stability of \( \text{BaCeO}_3 \) as well as of \( \text{BaCe}_{0.9}\text{Y}^{0.1}\text{O}_{3-\delta} \) in a CO\(_2\) gas atmosphere, using the thermogravimetric method within a limited temperature range of
100–900 °C. In these conditions, weight gains were observed above ~450 °C, continuing up to 900 °C due to the following reaction:

\[ \text{BaCeO}_3 + \text{CO}_2 \rightarrow \text{BaCO}_3 + \text{CeO}_2 \] (2)

It was also pointed out that BaCe_{0.9}Y_{0.1}O_{3-δ} exhibited a higher level of chemical stability than BaCeO_{3}. In the case of BCY, lesser weight gain was observed in the same conditions. The TG curves we obtained for BCY (Figure 6a) are close to the results of the cited paper [42].

At temperatures from 450 to 900 °C, a continuous increase in mass was observed. No peak of decomposition of carbonate compounds was found on the thermogravimetric curve (m/m₀ vs temperature) in these conditions. The first mass decrease on the TG curve for BCY was recorded at 970 °C [43]. This may correspond to the thermal decomposition of BaCO_3. The introduction of SrO into Ba_{1-x}Sr_xCe_{0.9}Y_{0.1}O_{3-δ} caused an increase in the maximum temperature decomposition of carbonate compounds in the TG curve. In the case of SBCY, increase in the temperature of carbonate decomposition was accompanied by increased SrO content in Ba_{1-x}Sr_xCe_{0.9}Y_{0.1}O_{3-δ} within the temperature range 995–1015 °C. The considerable decrease in mass following attainment of the maximum temperature might be identified as the release of CO_2 due to the reaction of SrCO_3/BaCO_3 with solid solutions of CeO_2 and Y_2O_3, according to reverse reactions (2) and (3).

\[ \text{Ba}_{1-x}\text{Sr}_x\text{Ce}_{0.9}Y_{0.1}\text{O}_{3(\delta)} + \text{CO}_2(\delta) \rightarrow (1-x)\text{BaCO}_3(\delta) + x\text{SrCO}_3(\delta) + \text{Ce}_{0.9}Y_{0.1}\text{O}_{1.95(\delta)} \] (3)

Two factors were decisive in terms of the probable origin of the faster reaction kinetics of BaCe_{0.9}Y_{0.1}O_{3-δ} (BCY). One was the basicity of the Ba^{2+} ion, which, being greater than that of the Sr^{2+} ion, led to the greater reactivity of the former, leading in turn to greater chemical reactivity against CO_2 with acidic properties at the surface of the particle. The second factor may be connected with the more rapid diffusion kinetics in the ABO₃ crystal structure of Ba^{2+} cations in BaCe_{0.9}Y_{0.1}O_{3-δ} compared to Sr^{2+} in Ba_{1-x}Sr_xCe_{0.9}Y_{0.1}O_{3-δ}. Thermogravimetric analysis enabled the observation that the presence of SrO in the Ba_{1-x}Sr_xCe_{0.9}Y_{0.1}O_{3-δ} samples caused considerably smaller relative mass changes compared to the BCY sample. In the case of the BCY sample, the increase in relative mass within the investigated temperature range could be attributed to CO_2 absorption at the surface of the particles and (via bulk diffusion) into the particles. These findings suggested the occurrence of a superficial reaction as well as bulk diffusion, given the bulk diffusion rate of Ba^{2+} for BaCe_{0.9}Y_{0.1}O_{3-δ}, which is higher than that of Sr^{2+} for Ba_{1-x}Sr_xCe_{0.9}Y_{0.1}O_{3-δ}. This may also be correlated with the structural parameter t and SFV. The least variation in mass was observed for the sample with strontium, where x = 0.02. A larger amount of SrO introduced into Ba_{1-x}Sr_xCe_{0.9}Y_{0.1}O_{3-δ} caused a lesser reduction in mass. These results also indicated that the smaller deviation from the original BCY structure may have been responsible for the more rapid kinetics of Ba^{2+} diffusion [44].

The chemical stability of the Ba_{1-x}Sr_xCe_{0.9}Y_{0.1}O_{3-δ} sintered sample was also tested in gas atmospheres involving 2.6 or 5 vol.% CO_2 in Ar at 550 °C for 100 h. The XRD diffraction patterns recorded for Ba_{1-x}Sr_xCe_{0.9}Y_{0.1}O_{3-δ} samples following additional heating in a 5% Ar-CO_2 gas atmosphere (Figure 7b) indicated that all samples exhibited precipitation of BaCO_3 and ceria solid solution. The results obtained from XRD measurements were correlated with data obtained via TG analysis in CO_2. A quantitative phase analysis performed using the Rietveld method clearly indicated that the amount of precipitation was greater for BCY than for SBCY. The amount of BaCO_3 for the BCY sample was estimated at about 61%. In the case of SBCY samples, the amount of BaCO_3 was estimated at 30%–33%. These data clearly indicated that the incorporation of SrO into Ba_{1-x}Sr_xCe_{0.9}Y_{0.1}O_{3-δ} inhibited the corrosion of samples in CO_2 gas atmospheres.

Wang et al. [28] investigated variation in the electrical conductivity of Ba_{1-x}Sr_xCe_{0.8}Y_{0.2}O_{3-δ} in wet hydrogen within the temperature range 450–800 °C by means of AC impedance spectroscopy conducted within a frequency range of 0.1 Hz–1 MHz. It was found that electrical conductivity slowly decreased along with increased SrO content in Ba_{1-x}Sr_xCe_{0.8}Y_{0.2}O_{3-δ} samples. Greater activation
energy of the electrical conductivity process was observed in samples with higher concentrations of SrO in the investigated solid solutions.

Hung et al. [29] also found that the total electrical conductivity of $\text{Ba}_{1-x}\text{Sr}_x\text{Ce}_{0.8}\text{Y}_{0.2}\text{O}_{3-\delta}$ samples decreased slightly along with increasing SrO content in air within the temperature range 400–800 °C. Systematic investigations concerning the impact of either grain boundary conductivity or bulk conductivity in $\text{Ba}_{1-x}\text{Sr}_x\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ samples are lacking. Data concerning variation in electrical conductivity in hydrogen or humidity-involving atmospheres also require detailed clarification regarding variation in structure and microstructure. The ionic conductivity of BaCeO$_3$-based electrolytes is usually measured in air, hydrogen, and steam gas atmospheres using the electrochemical impedance spectroscopy (EIS) method, which is limited to frequencies of 1–10 MHz [45–47].

This frequency limit does not reflect all of the phenomena occurring in bulk transport properties at temperatures higher than 450 °C, although this temperature range is crucial for LT- and IT-SOFCs. In an IT-SOFC with a BaCeO$_3$-based electrolyte, several processes take place in the bulk, including proton transport, oxygen-ion transport, and, possibly, electronic conductivity. These relaxations at elevated temperatures are shifted well above 10 MHz; therefore, ultrabroadband frequency equipment is necessary to investigate and resolve them [34,35].

Ultrabroadband frequency electrochemical impedance spectroscopy within the range 1–10 GHz had previously been applied to the study of the electrical conductivity of $\text{BaCe}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ and $\text{Ba}_{0.95}\text{Ca}_{0.05}\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ samples [27]. In the present study we continued the application of this method in order to study the electrical properties of SBCY sintered samples. All data were normalised to the sample geometry. The shape of the impedance in the complex plane plots measured in the two-electrode setup of both $\text{Ba}_{0.95}\text{Sr}_{0.05}\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ (2SBCY) and $\text{Ba}_{0.95}\text{Sr}_{0.05}\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ (5SBCY) consisted of semicircles: one separated high-frequency (HF), up to two overlapped medium-frequency (MF), and one separated low-frequency (LF).

Figure 8a–c presents examples of the impedance spectra for 2SBCY and 5SBCY measured at different temperatures.

The bulk and grain boundary ionic conductivity of SBCY sintered samples was determined using equivalent electrical circuits (EECs) consisting of up to four sub-circuits connected in series ($R_i$, CPE$_i$, where $i = 1–4$). In each sub-circuit, a capacitor was replaced by a constant phase element (CPE) whose impedance is expressed by the following formula:

$$Z_{\text{CPE}} = \frac{1}{2\pi f_0 C_0} \left( \frac{f_0}{f} \right)^{\alpha}$$

(4)

where $j$ is an imaginary unit, $f$ is frequency, $f_0$ is reference frequency, $C_0$ is capacitance at reference frequency (1 kHz), and $\alpha$ is a coefficient usually assuming a value between 0.5–1.0, close to 1 for an ideal capacitor and close to 0.5 for a diffusion process.

The HF semicircle ($R_1$, CPE$_1$) was ascribed to grain interior (bulk) processes. The fitted constant phase element (CPE$_1$) obtained from the fitting procedure fell within a range from 2 to $18 \times 10^{-10}$ F m$^{-1}$. The first MF semicircle was characterised by capacitances ranging from $3 \times 10^{-4}$ to $8 \times 10^{-8}$ F m$^{-1}$, ascribed to the ionic conductivity of grain boundary regions. The corresponding semicircle was substantially overlapped by the second MF semicircle; thus, the obtained values of grain boundary conductivities are characterised by considerable errors. The other MF and LF semicircles were ascribed to processes occurring in electrodes.
Figure 8. Complex impedance plane plots measured in air (a) at 287 °C for 2SBCY ceramics and (b) at 507 °C for 5SBCY ceramics; (c) an example of fit for 2SBCY ceramics measured at 407 °C; crosses indicate measured data, circles indicate the fitted model; numbers above red symbols denote logarithms of frequency.

Figure 8c gives an example comparing the fitted EEC model to measured data. The evolution of the various relaxation processes mentioned above can be inferred from the frequency dependencies of an imaginary impedance part (Figure 9).

The frequency where the maximum of $Z''$ is observed, called the relaxation frequency, is a good indication for the relaxation dispersion of $Z'$ (shown in Figure 9a). The relaxation frequency of the charge carrier in the grain interior changes from about 0.2 MHz at 87 °C to about 2 GHz at 707 °C. Consequently, the electrical resistance of grain interiors is characterised by lower values along with increases in temperature. Similarly, the increase in the first MF relaxation frequency, from about 100 Hz to about 40 MHz, was observed over the temperature range shown in Figure 9b. This relaxation frequency was associated with charge carriers in the grain boundaries of electrolyte ceramics. The separation of two MF relaxations was possible only by means of data fitting. The $Z''$ maximum of the low-frequency
relaxation process was observed only at 707 °C. The bulk conductivity dependence vs 1000/T is presented in Figure 10a.

![Figure 9. Spectra of (a) real \(Z'\) and (b) imaginary \(Z''\) parts for a 2SBCY sintered pellet. Arrows indicate (approximately) the shift of the relaxation frequency of several processes with changes in temperature.](image)

![Figure 10. Variation in grain interior \(\sigma_B\) (a) and grain boundary \(\sigma_{CB}\) (b) conductivities of the SBCY sample series vs temperature.](image)

The introduction of strontium resulting in \(\text{Ba}_{1-x}\text{Sr}_x\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}\) samples, where \(0 < x < 0.1\), led to variation in the electrical conductivity and activation energy of grain interiors along with SrO content in air within the temperature range 250–1000 K. For all SBCY sintered samples, two areas of variation of bulk conductivity could be distinguished in the plot (\(\sigma_B - 1000/T\)). The first region could be observed within the low temperature range, 298–600 K, in which higher activation energy \(E_a\) was observed. In the temperature range 600–1000 K, lower activation energy of the bulk conductivity of SBCY sinters was observed. Similar dependences of \(\sigma_B\) vs temperature were also observed for \(\text{BaCe}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}\) and \(\text{Ba}_{0.95}\text{Ca}_{0.05}\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}\) sintered samples and for \(\text{BaCe}_{1-x}\text{M}_x\text{O}_{3-\delta}\), where \(M = \text{Y}, \text{La}, \text{Gd}\) and \(x = 0.1–0.2\).

Analysis of electrical conductivity studies vs temperature or partial oxygen pressure indicated that, at temperatures higher than 600 K, oxide-hole conductivity dominates at high oxygen partial pressure (above \(pO_2 = 10^{-5}\) atm) [48–50]. Based on analysis of Figure 10a, it can be stated that the 2SBCY sintered sample was characterized by bulk ionic conductivity slightly higher than that of the BCY sample in this condition. Comparative analysis of the chemical resistance vs \(\text{CO}_2\) attack and bulk conductivity of the 2SBCY sample in a \(\text{CO}_2\) gas atmosphere indicated that the introduction of SrO into the \(\text{Ba}_{1-x}\text{Sr}_x\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}\) samples (where \(x = 0.02\)) led to both an improvement in chemical resistance in a \(\text{CO}_2\) gas atmosphere and an increase in bulk conductivity in comparison to the BCY sample.
A continued increase in the share of strontium, i.e., for $x > 0.02$ in the $\text{Ba}_{1-x}\text{Sr}_x\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ samples, led to a gradual decrease in bulk conductivity in air.

Grain boundary conductivity is also an important factor in determining the total electrical conductivity of ceramic material. The high electrical resistance of intergranular areas in a sintered ceramic is very often a factor preventing the acquisition of high values of ionic conductivity in proton-conducting electrolytes containing $\text{BaZr}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ (BZ) or $\text{BaCe}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ (BCY) [51,52]. Lindman presented an analysis of the potential for reducing the value of grain boundaries for BZY and BCY. The results of the considerations presented in the cited paper suggest that $\text{BaCe}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ [53] exhibits a greater tendency towards modification of grain boundaries aimed at reducing their electrical resistance. The variation in grain boundary ionic conductivity vs $1000/T$ for sintered $\text{Ba}_{1-x}\text{Sr}_x\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ samples is presented in Figure 10b. Based on the data presented therein, it can be observed that the gradual substitution of SrO for BaO in $\text{Ba}_{1-x}\text{Sr}_x\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ samples, where $0 < x < 0.1$, led to slight changes in grain boundary conductivity in relation to the BCY sample. Improved grain boundary conductivity was observed for the 2SBCY sample. It is also widely known that the introduction of small amounts of SrO or CaO into a CeO$_2$-Gd$_2$O$_3$-SrO solution enable the acquisition of greater electrical conductivity as a result of the potential for cleansing the grain boundary area of resistive non-organic compounds [54,55]. In the case of $\text{Ba}_{1-x}\text{Sr}_x\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ samples, where $x > 0.02$, with higher SrO concentrations, a deterioration in grain boundary conductivity was observed.

The total electrical conductivity $\sigma$ of the sintered $\text{Ba}_{1-x}\text{Sr}_x\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ samples within the temperature range 400–800 °C was also measured using AC electrochemical impedance spectroscopy in 5 vol.% H$_2$ in Ar. Figure 11a presents variations in $\sigma$ vs temperature for $\text{Ba}_{1-x}\text{Sr}_x\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ sintered samples.

It can be concluded that the sintered 5SBCY sample was characterised by the highest value of electrical conductivity and the lowest value of activation energy in an atmosphere of humidified hydrogen. In the case of $\text{Ba}_{1-x}\text{Sr}_x\text{Ce}_{0.9}\text{Y}_{0.1}\text{O}_{3-\delta}$ samples, the partial replacement of BaO with SrO led to a small improvement in total conductivity $\sigma$ in an atmosphere of humidified hydrogen gas. The $t_{\text{ion}}$ transference number was also estimated from EMF measurement of the steam/hydrogen concentration cell (1) (Figure 11b). In these conditions, the transference number determined for the BCY sample fell within the range 0.90–0.95. The 5SBCY and 25BCY sintered samples exhibited values of the $t_{\text{ion}}$ transference number similar to that of the BCY electrolyte. These results confirmed that the investigated SBCY samples exhibited the same level of ionic conductivity as the BCY ceramic membrane.

The electrical conductivity of the SBCY samples was also measured following additional heat treatment at 550 °C for 100 h in a CO$_2$/Ar gas atmosphere involving 2.6 or 5 vol.% CO$_2$.

Figure 12a presents the impedance spectra ($\mathcal{Z}''$–$\mathcal{Z}'$) recorded for the 5SBCY sample before and after additional heating in a 5 vol.% CO$_2$ gas atmosphere at 390 °C. As can be seen, increases in the
electrical resistance of grain interiors and grain boundary regions of the 5SBCY sample were observed following additional heat treatment in a CO₂ gas atmosphere. Figure 12b shows the variation in σ vs 1000/T determined for the 5SBCY sample and presents the electrical conductivity values obtained for samples before and after additional exposure in 5 or 2.6 vol.% CO₂ in Ar. All measurements were performed via the EIS method. A slight decrease in electrical conductivity was recorded for 5SBCY following additional tests performed in a gas atmosphere involving 2.6 or 5 vol.% CO₂.

The impact of the presence of H₂O in an Ar gas atmosphere on the total electrical conductivity of the 5SBCY sample was also studied. In Figure 13, the electrical conductivity values recorded before and after additional heating in a 5 vol.% CO₂ gas atmosphere at 390 °C (a); variation in σ vs temperature recorded for 5SBCY before and after additional exposure in 5 or 2.6 vol.% CO₂ (b).

The comparative analysis of electrical conductivity data for the 5SBCY electrolyte before and after test enabled us to state that exposure of the sample in an H₂O gas atmosphere led to a slight improvement in electrical conductivity. This finding may be directly connected with the incorporation of water in the 5SBCY sample.
4. Conclusions

All of the investigated Ba$_{1-x}$Sr$_x$Ce$_{0.9}$Y$_{0.1}$O$_{3-\delta}$ samples were monophase. The orthorhombic (Pnma space group) structure was identified exclusively using the XRD diffraction method. In the case of SBCY samples with increased SrO content in Ba$_{1-x}$Sr$_x$Ce$_{0.9}$Y$_{0.1}$O$_{3-\delta}$ the diffraction peaks shifted towards greater diffraction angles in the recorded XRD patterns; consequently, a decrease was found in the cell volume of Ba$_{1-x}$Sr$_x$Ce$_{0.9}$Y$_{0.1}$O$_{3-\delta}$ samples when compared to BaCe$_{0.9}$Y$_{0.1}$O$_{3-\delta}$. Raman studies confirmed that the introduced Sr$^{2+}$ ions were located in the same position in the crystallographic structure in Ba$_{0.98}$Sr$_{0.02}$Ce$_{0.9}$Y$_{0.1}$O$_{3-\delta}$, Ba$_{0.95}$Sr$_{0.05}$Ce$_{0.9}$Y$_{0.1}$O$_{3-\delta}$, and Ba$_{0.90}$Sr$_{0.10}$Ce$_{0.9}$Y$_{0.1}$O$_{3-\delta}$ as the original Ba$^{2+}$ ions in BaCe$_{0.9}$Y$_{0.1}$O$_{3-\delta}$. The increase in the concentration of SrO in Ba$_{1-x}$Sr$_x$Ce$_{0.9}$Y$_{0.1}$O$_{3-\delta}$, where 0 < x < 0.1, also led to a decrease in tolerance factor as well as increases in specific free volume and global instability factor. The analyzed structural parameters were responsible for variation in chemical resistance in CO$_2$ and electrical conductivity in air, hydrogen, or wet-gas-involving atmospheres. It was found that Ba$_{1-x}$Sr$_x$Ce$_{0.9}$Y$_{0.1}$O$_{3-\delta}$, where 0.02 < x < 0.05, exhibited improved electrical conductivity in air as well as in hydrogen or humidity-involving gas atmospheres. Superior chemical stability in CO$_2$-involving gas atmospheres was also observed in the case of these samples in comparison to BaCe$_{0.9}$Y$_{0.1}$O$_{3-\delta}$. The values of the $t_\text{ton}$ transference number determined in hydrogen concentration galvanic cells for Ba$_{0.98}$Sr$_{0.02}$Ce$_{0.9}$Y$_{0.1}$O$_{3-\delta}$ and Ba$_{0.95}$Sr$_{0.05}$Ce$_{0.9}$Y$_{0.1}$O$_{3-\delta}$ are comparable to data obtained for BaCe$_{0.9}$Y$_{0.1}$O$_{3-\delta}$. The composition of Ba$_{0.98}$Sr$_{0.02}$Ce$_{0.9}$Y$_{0.1}$O$_{3-\delta}$ or Ba$_{0.95}$Sr$_{0.05}$Ce$_{0.9}$Y$_{0.1}$O$_{3-\delta}$ mentioned above exhibited not only the same level of electrical conductivity as BaCe$_{0.9}$Y$_{0.1}$O$_{3-\delta}$, but also improved chemical resistance in 2.6–5.0 vol.% CO$_2$ gas atmospheres. The variations in electrical conductivity recorded for the Ba$_{0.95}$Sr$_{0.05}$Ce$_{0.9}$Y$_{0.1}$O$_{3-\delta}$ sample following exposure in gas atmospheres with close to 5 vol.% of CO$_2$ were smaller than those for BaCe$_{0.9}$Y$_{0.1}$O$_{3-\delta}$ and the previously investigated Ba$_{0.95}$Ca$_{0.05}$Ce$_{0.9}$Y$_{0.1}$O$_{3-\delta}$. These materials appear to be more suitable ceramic proton-conducting membranes for the construction of ceramic fuel cells operating in hydrogen gas atmospheres involving contamination of CO$_2$ below 2 vol.%.

Author Contributions: M.D.: Conceptualization of the whole paper, methodology of all experimental parts, interpretation results, writing original draft, preparing answer for reviewers, writing final version of paper; B.L.: SBCY samples preparation, SPUDS analysis, electrical and electrochemical investigations of samples in humidity or H$_2$ involving gas atmosphere, technical editing of original and final version of manuscript; R.L.: XRD investigations and interpretation all XRD results, answers for review comments concerning the XRD methodology; S.D.: investigations of SBCY samples by broadband impedance spectroscopy method, visualisation of data, T.S.: interpretation of results, broadband impedance spectroscopy methodology, consultation of final interpretation of all electrical investigations and results, some correction of final manuscript; A.K.: investigations of SBCY samples by broadband impedance method, supervision for all measurements, software development, electrochemical setup, modelling data of EIS results; M.M.: final consultation, modelling analysis of EIS data, EIS methodology, software, visualisation, writing the EIS part in original and final manuscript; M.S.: Raman Spectroscopy investigations, interpretation of these results, consulting and writing some part of text in original and final manuscript according to this method; A.R.-K.: SEM microscopy measurements, investigations, consulting the results, P.G.: thermogravimetry investigations, methodology, consulting the interpretation of TG results. All authors have read and agreed to the published version of the manuscript.

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