Surface and Bulk Oxygen Kinetics of BaCo$_{0.4}$Fe$_{0.4}$Zr$_{0.2-x}$YxO$_{3-\delta}$ Triple Conducting Electrode Materials

Jack H. Duffy 1,2, Yuqing Meng 1, Harry W. Abernathy 2 and Kyle S. Brinkman 1,2,*

Abstract: Triple ionic-electronic conductors have received much attention as electrode materials. In this work, the bulk characteristics of oxygen diffusion and surface exchange were determined for the triple-conducting BaCo$_{0.4}$Fe$_{0.4}$Zr$_{0.2-x}$YxO$_{3-\delta}$ suite of samples. Y substitution increased the overall size of the lattice due to dopant ionic radius and the concomitant formation of oxygen vacancies. Oxygen permeation measurements exhibited a three-fold decrease in oxygen permeation flux with increasing Y substitution. The DC total conductivity exhibited a similar decrease with increasing Y substitution. These relatively small changes are coupled with an order of magnitude increase in surface exchange rates from Zr-doped to Y-doped samples as observed by conductivity relaxation experiments. The results indicate that Y-doping inhibits bulk O$_2^-$ conduction while improving the oxygen reduction surface reaction, suggesting better electrode performance for proton-conducting systems with greater Y substitution.

Keywords: triple ionic-electronic conductor; oxygen permeation membrane; electrical conductivity relaxation

1. Introduction

Mixed ionic-electronic conductors (MIECs) are utilized in a wide range of applications in intermediate to high temperature systems, from protonic ceramic fuel cells (PCFCs) to oxygen separation and catalysis [1–3]. Recently, a subset of MIECs known as triple ionic-electronic conductors (TIECs), predominantly electron hole conductors, but also proton and oxygen ion conductors, have emerged as a promising class of materials, especially as cathodes for PCFCs due to their excellent oxygen reduction reaction (ORR) activity and ease of synthesis [4–6]. The conduction of these three species is particularly useful to extend the range of active sites from the electrolyte-electrode interface to the entire surface of the electrode [7].

Despite their strong performance metrics in PCFC applications, bulk properties of TIECs are relatively unknown, especially surface interactions and bulk conductivity of protons and oxygen ions. Historically, measurements for cathode materials focused on characterization in its microstructural, porous architecture, often using electrical impedance spectroscopy (EIS) and power density measurements. These methods provide useful information about ORR kinetics across samples [8], while changes in processes in samples of the same material can be successfully identified [9]. However, it may be difficult to deconvolute these data in different materials as the result of either microstructural effects or material effects. It is therefore pertinent to understand how a material behaves in its dense, nonporous form to understand further how it behaves in functional devices.

Recent work has pushed toward better characterization of bulk TIEC properties, further probing the mechanisms which enable strong surface kinetics and high ionic conductivity in these materials. Electrical conductivity relaxation (ECR) is a prominent method...
which allows for relatively simple measurements to probe the oxygen reduction kinetics of cathode materials [10–14]. ECR has also recently been used to probe the kinetics of other surface behaviors, such as proton uptake and interactions with water [15–17]. Measuring ionic conductivity in TIECs, however, is more challenging, as most known materials in this class are predominantly electron conductors, due to their high doping concentration of multivalent elements, such as Co, Fe, Ni, and Mn [18–20]. High electronic conductivity renders traditional methods such as EIS, often used for predominantly ionic conductors, ineffective. Instead, permeation is a widely used technique to probe the ionic conductivity in dominant electronic conductors such as MIECs and TIECs [21–23]. Zhou et al. recently utilized ECR and permeation methods to report a new cathode material Sr₂Co₀.₅ Nb₀.₅ Fe₀.₅ O₆−δ (SSNCF) with both performance data and bulk property data compared to a well-known material Ba₀.₅Sr₀.₅Co₀.₈ Fe₀.₂ O₃−δ (BSCF) [24]. Despite recent developments, there remains a large literature gap in bulk property data for many claimed triple-conductors.

Among these TIECs, BaCo₄Fe₄Zr₀.₁Y₀.₁O₃−δ (BCFZY0.1) is one of the most widely studied state-of-the-art single-phase cathodes. Aside from strong BCFZY0.1 performance data on PCFCs and SOFCs, its fundamental surface kinetics [15,16,25] and bulk conductivity are well-studied. Multiple variants of BCFZY0.1 have been synthesized and characterized to compare with the base system. A-site-deficient BCFZY0.1 materials have been synthesized, showing improved hydrogen evolution reaction (HER), oxygen diffusion, and hydration parameters from its original structure [26,27]. A B-site deficient variant, BaCo₄Fe₄Zr₀.₁Y₀.₁O₃−δ (BCFZY0.95), was also synthesized, exhibiting improved proton conduction and ORR kinetics [28,29]. Liang et al. created a Ni-substituted variant, BaCo₄Fe₄Zr₀.₁Y₀.₁Ni₀.₀5O₃−δ (BCFZYN), with improved ORR kinetics and ionic conductivity [30]. These previous results show that BCFZY0.1 can be tuned through doping and deficiency strategies with some measured success.

BCFZY0.1 was initially derived from BaCo₄Fe₄Zr₀.₂O₅−δ (BCFZ) [31], as yttrium was substituted into the materials system to impart more proton conductivity and improve ORR kinetics [1,32,33]. The specific composition was chosen based on EIS measurement of the area specific resistance of symmetric cells comprised of different Zr to Y ratios [1]. While the resulting composition was a better performing cathode than BCFZ, the underlying properties of Y substitution in BCFZ remained unstudied.

To better understand how to design TIEC materials for electrodes, including single-phase and multiphase systems, it is paramount to understand the tunability of state-of-the-art, model materials, such as BCFZY0.1. In this work, the oxygen surface kinetics and bulk conductivity are studied for the BaCo₄Fe₄Zr₀.₁₂−₂X₃O₅−δ (BCFZYₓ, X = 0, 0.05, 0.1, 0.15, 0.2) materials system. It is shown that a tradeoff occurs as Y is substituted into the system, with a small decrease in bulk oxygen ion conductivity and an order-of-magnitude increase in oxygen surface exchange kinetics with increasing Y concentration. This work begins to probe the effect of yttrium in the BCFZYₓ system and lays the foundation for future work to understand the important factors which make high performance electrode materials.

2. Materials and Methods

2.1. Preparation of BCFZYₓ Precursor Powders

All compositions of BCFZYₓ precursor powders were prepared by a modified Pechini (citric-acid) method [1]. Stoichiometric amounts of Ba(NO₃)₂ (Alfa Aesar, 99+%), Co(NO₃)₂·6H₂O (Alfa Aesar, >98.0%), Fe(NO₃)₃·9H₂O (Alfa Aesar, >98%), ZrO(NO₃)₂ solution (Sigma-Aldrich, ≥99%, 35 wt.% in dilute nitric acid), and Y(NO₃)₃·6H₂O (Alfa Aesar, 99.9%) were dissolved in deionized water in appropriate molar ratios. The obtained solution is mixed with citric acid (Sigma-Aldrich, ≥99.5%) and EDTA (Alfa Aesar, 99.4+%) with a citric acid to EDTA to metal-ion ratio of 1.5:1:5.1. The solution was then adjusted to reach a desired pH value of 9 by adding NH₄OH (Alfa Aesar, 28.0–30.0% NH₃) and heated at 80
°C to obtain a dark purple gel through water evaporation. The gel was dried at 150 °C for 48 h to obtain the primary powder.

2.2. Fabrication of Dense BCFZYx pellets for Characterization

For preparation of the 1 mm thick membranes, the primary BCFZYx powders were calcined at 1000 °C for 8 h in air. After manually grinding, the powders were mixed with binder (5% polyvinyl alcohol in water) and uniaxially pressed into pellets of 15 mm diameter under 300 MPa for 2 min. The pellets were buried in calcined powder and subsequently sintered at 1270 °C for 8 h. All sintered samples had relative densities greater than 95%. The sintered pellets were subsequently polished to approximately 1 mm thickness using progressively finer sanding paper grits to achieve a polished, shiny surface to reduce surface effects during characterization.

The BCFZY0.1 membrane was also tested with a porous surface coating of the same composition. To make this surface coating, the BCFZY0.1 primary powder was pre-calcined at 600 °C for 5 h followed by ball milling in 1-butanol (Fisher Chemical, ≥99.4%) for 7 days. The precursor powder was obtained by drying the ball-milled powder at 500 °C for 5 h. Surface coating paste was prepared by mixing the precursor powder with dispersant (20 wt% solspers 28000 in terpinol solution) and binder (5 wt% Heraeus V-600 in terpinol solution) in a 15:3:1 weight ratio by manual grinding in a mortar and pestle for 45 min. For the surface coated BCFZY0.1 membrane, the BCFZY0.1 paste was screen printed on both sides of the pellet. The whole structure was fired at 900 °C to form the porous BCFZY0.1 | dense BCFZY0.1 | porous BCFZY0.1 membrane.

2.3. Characterization

The prepared, sintered membranes were crushed into powders and analyzed by X-ray diffraction (XRD; Ultima IV, Rigaku Americas Corporation, The Woodlands, TX, USA) using Cu/α radiation (λ = 1.54108 Å), with a scan range from 20–90° and a step size of 0.02°. The microstructures of the membranes and coating layer were analyzed using scanning electron microscopy (SEM; SU6600, Hitachi High-Tech, Tokyo, Japan).

For oxygen permeation measurements, the BCFZYx membranes were sealed on one side with an alumina tube (O.D. = 0.5 in., I.D. = 0.375 in.) by a ceramic paste (Ceramabond 552, Aremco), followed by additional sealing on the sidewalls of the membrane to reduce sidewall permeation and ensure low leakage. A schematic of the experimental setup is provided in Figure S1. The sealed membrane was heated to 100 °C for 2 h and then 250 °C for 2 h to ensure proper curing of the bond before heating to 550 °C at a ramp rate of 1 °C·min⁻¹. The feed side remained unsealed and was exposed to ambient air inside the furnace. The permeate side was supplied with ultra-high purity helium (He; 99.999%, Airgas) at a rate of 50 mL·min⁻¹. The gas concentrations of O₂ and N₂ were measured on the membrane permeate side using a gas chromatograph (GC; Micro GC Fusion, INFICON, Bad Ragaz, Switzerland), with the concentration of N₂ used to calculate the physical leakage of air from the feed side to the permeate side. The oxygen permeation flux was subsequently corrected using the total measured oxygen on the permeate side minus the calculated physical leakage of oxygen. Measured leakage was less than 3% for all samples.

Electrical conductivity was measured for all BCFZYx samples using a DC four-point probe method with a digital multimeter (Keithley 2001 Series, Tektronix, Inc., Beaverton, OR, USA) on bar samples fabricated from the sintered pellets. Each pellet was polished to create approximately 12 mm x 5 mm x 1 mm dense bars for measurement. Conductivity was measured both in dry air and in humidified air passed through a room temperature (20 °C) bubbler. In addition, this setup was utilized in an ECR method to determine the surface exchange coefficient, \( k_{\text{chem}} \), and the diffusion coefficient, \( D_{\text{chem}} \). Gas composition was abruptly changed from \( pO_2 \) of 0.1 to 0.21 at a flow rate of 225 mL·min⁻¹, and the electrical conductivity changed continuously during the gas switching. The ECR measurement was also performed in both dry and humidified air.
3. Results

3.1. Structure and Composition

Figure 1a displays the x-ray diffraction pattern for a crushed, sintered pellet of BaCo_{0.4}Fe_{0.4}Zr_{0.15}Y_{0.05}O_{3-δ} (BCFZY0.05) with Rietveld refinement. All compositions, including BCFZY0.05, show a cubic perovskite (Pm̅3m) phase without any detectable impurities after sintering at 1270 °C for 8 h, and can be seen in Figure S3. The (110) peak is analyzed in Figure 1b, indicating an increased lattice parameter with more incorporated Y. The Y substitution should cause an increase in oxygen vacancies from the original BCFZ material, as written in Kröger–Vink notation as follows:

\[ Y_2O_3 + O_0^\delta + 2Zr^{4+}_X \leftrightarrow 2Y^{4+}_Z + V^{\ast}_O + 2ZrO_2. \]  

(1)

Using Rietveld refinement for the measured samples, the calculated lattice parameter for each sample is shown in Figure 1c. Generally, the lattice parameter increased with increased Y substitution, as expected with a larger Y^{3+} cation substitution for the Zr^{4+} cation, consistent with the (110) peak shift from Figure 1b. The lattice parameter was used to calculate parameters which may affect the overall oxygen mobility in the structure, including lattice free volume, critical radius. These data are included in Table S1.

Sample compositions observed under SEM determined the microstructural differences across compositions. The probed samples show that the average cross-sectional grain size generally increases with Y substitution from 3 μm (BCFZ) to 5 μm (BCFZY0.05) to 10 μm (BCFZY0.1) to 18 μm for BaCo_{0.4}Fe_{0.4}Zr_{0.05}Y_{0.15}O_{3-δ} (BCFZY0.15) and BaCo_{0.4}Fe_{0.4}Y_{0.2}O_{3-δ} (BCFY), indicating greater sinterability with greater Y substitution. The grain size is within the same order of magnitude, meaning that there should be no nanostructural effects from grain boundary diffusion of oxygen. In addition, samples were probed using energy-dispersive x-ray spectroscopy (EDX) to confirm the doping of Zr and Y to the system. In all materials, the normalized ratio of Zr:Y was found to be the same as the desired ratio (shown in Table S2).

3.2. Oxygen Permeation Properties

A representative figure of the oxygen permeation flux as a function of measurement time is shown in Figure 2 for BCFZY0.05 at temperatures of 600 to 800 °C. After ramping, the temperature is held for 30 min to ensure stabilization of any temperature hysteresis before performing the measurement. Each measurement lasted approximately one hour, with individual gas samples taken in 15-min intervals. The flux remains stable throughout
the duration of each temperature. No secondary phase formation or microstructural changes were observed after the permeation measurements as confirmed by SEM/EDX analysis, indicating good material stability in these conditions. In all samples, the final three measurement points at each temperature were averaged as the representative permeation flux for further figures and discussion.

Figure 2. Oxygen permeation flux as a function of measurement time at increasing temperatures for BCFZY0.05, showing stable flux over each one-hour measurement.

Figure 3a displays the oxygen permeation flux as a function of membrane temperature for all BCFZYs compositions, normalized to 1 mm thickness. Error bars were calculated from measurement propagation error and the standard deviation of the three representative data points. It is observed that as Y substitution increases, the oxygen permeation flux decreases. At 800 °C, as shown in Figure 3b, BCFZ exhibits a flux of 0.485 mL·min⁻¹·cm⁻², markedly similar to previously reported literature [34], and continues to decrease with Y substitution until it reaches a plateau for the BCFZY0.1 and BCFZY0.15 compositions, followed by a continued decrease in flux to 0.198 mL·min⁻¹·cm⁻² for BCFY. The general trend of decreasing flux with increasing Y is somewhat unexpected, as it was hypothesized that increasing oxygen vacancies in the material would result in more pathways for oxygen diffusion. Based on the structural data from XRD, the estimated free volume and average metal-oxygen bond energy would also suggest that increasing Y concentration would cause an increase in oxygen mobility. Rather, it is hypothesized that Y substitution distorts the lattice from BCFZ due to its much greater ionic size than Zr, Fe, and Co, thus decreasing perovskite symmetry. The decreasing crystal symmetry may cause larger vacancy-lattice interactions and may help explain the trend for oxygen mobility in this materials system [35].
Figure 3. Oxygen permeation flux normalized for 1 mm BCFZY\textsubscript{x} pellets as a function of (a) temperature, and (b) \textit{Y} stoichiometry at 800 °C.

For the BCFZY0.1 composition, three pellets were measured for oxygen permeation flux at thicknesses of 1 mm, 0.54 mm, and 0.37 mm. As shown in Figure 4a, the oxygen permeation increased with decreasing thickness across all temperatures, consistent with the modified Wagner equation below [36]:

\[
J_{O_2} = \frac{RT}{L_c} \frac{\ln \sigma_e \ln \sigma_i}{\sigma_e + \sigma_i} d \ln P_{O_2},
\]

where \(J_{O_2}\) is the oxygen permeation flux, \(L_c\) is the material characteristic thickness, \(L\) is the membrane thickness, \(T\) is the temperature, \(R\) and \(F\) are the gas and Faraday constants, respectively, and \(\sigma_e\) and \(\sigma_i\) are the electronic and ionic conductivities, respectively. Each thickness also shows a characteristic leveling-off of oxygen flux at higher temperatures. This is due to the increase in \(L_c\) with temperature, resulting in a material with more surface limited performance [36].

Figure 4. Oxygen permeation flux for the BCFZY0.1 sample (a) as a function of temperature, under varying thicknesses, and (b) as a function of thickness at 600 °C.
A similar phenomenon is seen in Figure 4b, which displays the flux as a function of membrane thickness at 600 °C. The theoretical bulk-controlled and mixed-controlled oxygen permeation is plotted using Equation (2) with experimentally determined values, along with the experimental oxygen permeation for bare and coated samples. It is noted that the experimental \( J \) vs \( L^{-1} \) plot is non-linear, with thinner bare membranes leveling and exhibiting fluxes further from the theoretical value. This divergence suggests that the membranes become more limited by surface processes as the thickness decreases.

By adding the surface coating of the same composition at approximately the same thickness, the permeation flux significantly increases. The full surface coating dataset is available in Figure S4. The addition of the surface coating increases the surface exchange kinetics by increasing the availability of surface sites for oxygen reduction, thereby increasing the oxygen permeation. The surface coated membrane also aligns closely with the theoretical bulk-controlled prediction, affirming that this membrane has less surface limitation than the bare sample at similar thickness. The respective increases in flux due to surface coating and decreasing thickness suggest that the bare BCFZY0.1 membranes are controlled by a mixture of surface and bulk processes for the measured thickness range.

3.3. Conductivity

The total conductivity vs temperature plot for all BCFZY\(_x\) compositions is shown in Figure 5a for dry atmosphere. At 600 °C, the conductivity ranges from 2.71 S∙cm\(^{-1}\) for BCFZ to 1.25 S∙cm\(^{-1}\) for BCFY, decreasing with increasing Y substitution. This decrease in total conductivity can be somewhat attributed to the increase in lattice constant with Y substitution, as the Y\(^{3+}\) ion is larger than the Zr\(^{4+}\) ion. The BCFZY\(_x\) materials also show significantly lower electronic conductivity compared to other triple-conducting cathode materials such as SSNCF [24], BSCF [37,38], and PrBa0.5Sr0.5Co1.5Fe0.5O\(_{4+\delta}\) (PBSCF) [5,39].

In wet (2.3% H\(_2\)O) atmosphere, the total conductivity uniformly decreases by about 0.02 S∙cm\(^{-1}\) across all samples and temperatures, as seen in Figure 5b for the BCFY composition. This difference in conductivity is attributable to Equation (3), where in redox-active materials, an introduction of humidity leads to a reduction in the hole concentration and conductivity of all compositions at constant oxygen partial pressures [40]:

\[
H_2O + 2O_\Delta^\delta + 2h^* \leftrightarrow 2OH^- + \frac{1}{2}O_2, \tag{3}
\]

where \(O_\Delta^\delta\) and \(OH^-\) represents oxygen and hydroxide ions, respectively, on lattice sites, and \(h^*\) denotes electron holes. In addition, in both wet and dry atmospheres, two characteristic zones of conductivity appear. Below 475 °C, the activation energy \(E_a = 0.11\) eV, while above 475 °C, \(E_a = 0.04\) eV. This temperature-driven distinction is consistent with previous measurements of BCFZY0.1 [1].
From the oxygen permeation measurements, the oxygen ion conductivity can be estimated for all species, as shown in Figure S5. It has been previously shown that BCFZY0.1 exhibits the p-type carrier regime where $\sigma_{\text{electronic}} \gg \sigma_{\text{ionic}}$ [25]. Given the similarities in structure, total conductivity, and oxygen permeation, it is reasonable to assume that all BCFZYx compositions follow these p-type regime characteristics. Using this p-type assumption and assuming the membranes are bulk limited ($L_C/L$ approaches zero) at 1mm thickness, Equation (2) can be applied to solve for $\sigma_{\text{ion}}$ using the measured oxygen permeation flux as follows [41]:

$$\sigma_{\text{ion}} = \frac{16F^2L\cdot J_{O_2}}{R \cdot T \cdot \ln \left( \frac{P_l}{P_h} \right)}$$

where $P_l$ and $P_h$ are the low and high oxygen partial pressures across the membrane. Because the estimates were calculated from the oxygen permeation measurements and the Wagner equation, the same general trend and shape is seen in both the permeation and conductivity data. Relatively small changes in conductivity were observed, as conductivity remained well within an order of magnitude at all temperatures, with approximately a factor of three separating the lowest ionic conductor (BCFY) and the highest ionic conductor (BCFZ) at 800 °C. BCFZY0.1 also exhibited similar conductivity to that in previously reported literature [30]. In air, the ionic transport in relation to electronic transport (tion) remains approximately constant across all samples from 550 to 700 °C, owing to the relative changes in electronic and ionic conductivity across samples.

### 3.4. Electrical Conductivity Relaxation

To understand the surface effect of oxygen transport in BCFZYx, each composition’s conductivity was measured under rapid gas switching. At high temperatures, oxygen exchange occurs in perovskites according to the following:

$$\frac{1}{2}O_2 + V_o^{*} \leftrightarrow O_o^* + 2h^*,$$

where $V_o^*$ represents oxygen vacancies. From Equation (5), the electronic conductivity of each composition is dependent on the oxygen partial pressure. Following a rapid oxidation from 0.1 to 0.21 atm, the conductivity response of each composition is represented in Figure 6a by fitted, normalized conductivity as a function of time at 600 °C. BCFZY0.1
exhibits the fastest response time at around 300 s, increasing in response time as the composition reaches the BCFZ (around 600 s) and BCFY (around 500 s) endmembers. The same gas switch was also performed in humidified atmospheres to simulate PCFC conditions. As shown in Figure 6b for BCFZY0.05, BCFZY0.1, and BCFZY0.15, humidifying air increased the relaxation response time. This increase in response time is likely the result of competition between Equation (5) and the following equation below, as both utilize oxygen vacancies for oxygen and water adsorption, respectively [25]:

\[
H_2O + V^*_O + O^*_O \leftrightarrow 2O^*H^+.
\] (6)

The resultant normalized conductivity curves were fitted using the diffusion equations to determine \(k_{chem}\) and \(D_{chem}\), specifically using the ECR tool developed by the National Energy Technology Laboratory (NETL) [42]. In Figure 6c and Figure 6d, the calculated surface exchange coefficient \((k_{chem})\) and diffusion coefficient \((D_{chem})\), respectively, are displayed as functions of composition. From these data, it is evident that \(k_{chem}\) increases with increasing Y substitution, which agrees with the expectation that an increase in Y substitution increases the oxygen vacancy concentration, resulting in more active sites for surface exchange and reduction via Equation (3). The surface exchange coefficient \(k\) decreases with the introduction of water to the atmosphere (Figure S6), exhibited by the increase in relaxation time. The diffusion coefficient, \(D_{chem}\), does not appear to have any clear trend; \(k_{chem}\) has an overall order-of-magnitude increase over the range of Y substitution, while \(D_{chem}\) only changes by about a factor of 2 from its lowest (BCFY) to its highest (BCFZY0.1) values. BCFZY0.1 has the fastest response time of all samples, which may be attributed to the best combination of bulk diffusion with surface exchange.
These data, coupled with XRD and oxygen permeation data, which showed a decrease in oxygen permeation at 600 °C, suggest that increased oxygen vacancies from Y substitution changes local structuring in the material, which benefits surface reactions, but inhibits bulk oxygen diffusion. The weaker increase in bulk oxygen transport properties is thus impacted by the interplay between a combination of increased oxygen vacancies with the impact on the oxygen vacancy mobility from structural changes. Techniques such as neutron scattering may be used to probe local structure to confirm this hypothesis, presenting a clear path toward understanding these conflicting phenomena.

Using $k_{\text{chem}}$ and $D_{\text{chem}}$ obtained from ECR measurements, the characteristic thickness, $L_c$, can be estimated for all compositions at 600 °C. Defined as the ratio of $D$ to $k$ [21], $L_c$ quantitatively expresses the relative control of transport via bulk diffusion or surface exchange. In Figure 7a, the characteristic thickness decreases with increasing Y substitution, from 209 μm for BCFZ to 21 μm for BCFY. This decrease in $L_c$ signifies that the materials become more bulk controlled as Y is substituted in the material system as a result of faster surface kinetics. BCFY is identified as a strong candidate for thin-film membranes, due to its low characteristic thickness, and for an infiltration material in fuel cell cathodes because of its superior ORR kinetics.

**Figure 7.** (a) The characteristic thickness, $L_c$, at 600 °C, calculated from ECR measurements for each BCFZYx composition. (b) Oxygen ion conductivity estimated using the Wagner equation with and without a correction for the characteristic thickness.

With experimentally determined $L_c$, Equation (2) can be utilized further to estimate the oxygen ion conductivity more accurately at this temperature. Figure 7b displays the oxygen ion conductivity for all BCFZYx samples with and without the correction from characteristic thickness. For compositions with low $L_c$, such as BCFY, there is little deviation (about 4%) between the bulk-controlled assumption and the mixed-controlled assumption. However, for compositions with higher $L_c$ such as BCFZ, the estimates can deviate by as much as 40%. This deviation highlights the importance of $L_c$ to improve the accuracy for permeation as a method of estimating oxygen ion conductivity for triple conducting materials.

**4. Conclusions**

A suite of BCFZYx samples with different ratios of Zr:Y were probed for oxygen surface kinetics and bulk conductivity. Increasing Y concentrations in this material system
increased the lattice parameter of the perovskite structure. In addition, Y substitution resulted in the following across the entire system at 600 °C: (i) an order-of-magnitude increase in oxygen surface exchange coefficient, (ii) a decrease in oxygen permeation by a factor of three, and (iii) a decrease in electronic conductivity by a factor of two. Bulk oxygen ionic conductivity was also estimated for the range of samples, noting the importance of the characteristic thickness on the accuracy of this estimation. It was also determined through the surface coating and varying thickness measurements that the BCFZY0.1 membranes between 1 mm and 0.37 mm were controlled by a mixture of surface and bulk processes. Further research is required on the proton-conducting nature of these materials to better understand underlying reasons for their superior performance as cathode materials.

Supplementary Materials: The following are available online at www.mdpi.com/article/10.3390/membranes11100766/s1, Figure S1: Oxygen permeation experimental setup, Figure S2: SEM images for BCFZYs, Figure S3: BCFZYs X-Ray diffraction, Table S1: Estimated and calculated structural data for BCFZYs, Table S2: Stoichiometry of BCFZYs calculated from EDX data, Figure S4: Oxygen permeation flux for bare and surface-coated membranes, Figure S5: Oxygen conductivity as a function of temperature, Figure S6: $k_{chem}$ in wet and dry atmospheres, and Figure S7: Arrhenius plot of oxygen-ion conductivity for BCFZYs.

Author Contributions: J.H.D. performed the experiments; J.H.D. and Y.M. performed the data analysis; J.H.D., Y.M., and K.S.B. conceived and designed the experiments; Y.M., H.W.A., and K.S.B. supervised the study and provided scientific discussions; J.H.D. prepared the original draft manuscript. All authors edited and reviewed the manuscript. All authors have read and agreed to the published version of this manuscript.

Funding: This work was financially supported by the National Energy Technology Laboratory (NETL) and Oak Ridge Institute for Science and Education (ORISE). K.S.B. was supported in part by an appointment to the NETL Research Participation Program, sponsored by the U.S. Department of Energy and administered by the Oak Ridge Institute for Science and Education.

Institutional Review Board Statement: Not applicable.

Informed Consent Statement: Not applicable.

Data Availability Statement: All data are contained within the article and its Supplementary Materials.

Conflicts of Interest: The authors declare no conflict of interest.

References
8. Endler-Schuck, C.; Joos, J.; Niedrig, C.; Weber, A.; Ivers-Tiffée, E. The chemical oxygen surface exchange and bulk diffusion coefficient determined by impedance spectroscopy of porous La0.58Sr0.4Co0.2Fe0.8O3-δ (LSCF) cathodes. Solid State Ionics 2015, 269, 67–79, doi:10.1016/j.ssi.2014.11.018.
Membranes 2021, 11, 766


16. Chen, Y.; Hong, T.; Wang, P.; Brinkman, K.; Tong, J.; Cheng, J. Investigate the proton uptake process of proton/oxygen ion/hole triple conductor BaCo0.4Fe0.4Zr0.1Y0.1O3-δ by electrical conductivity relaxation. J. Power Sources 2019, 440, 227122, doi:10.1016/j.jpowsour.2019.227122.


23. Balaguer, M.; Solis, C.; Serra, J.M. Study of the transport properties of the mixed ionic electronic conductor Ce0.8-TbO0.2+Co (x = 0.1, 0.2) and evaluation as oxygen-transport membrane. Chem. Mater. 2011, 23, 2333–2343, doi:10.1021/cm103881w.


26. Ren, R.; Wang, Z.; Xu, C.; Sun, W.; Qiao, J.; Rooney, D.W.; Sun, K. Tuning the defects of the triple conducting oxide BaCo 0.4 Fe 0.4 Zr 0.1 Y 0.1 O 3-δ perovskite toward enhanced cathode activity of protonic ceramic fuel cells. J. Mater. Chem. A 2019, 7, 18365–18372, doi:10.1039/c9ta03359z.


31. Shang, M.; Tong, J.; O’Hayre, R. A promising cathode for intermediate temperature protonic ceramic fuel cells: BaCo0.4Fe0.4Zr0.1O3-δ. RSC Adv. 2013, 15769–15775, doi:10.1039/c3ra41828f.


37. Chen, D.; Shao, Z. Surface exchange and bulk diffusion properties of Ba0.5Sr 0.5Co0.8Fe0.2O3-δ mixed conductor. *Int. J. Hydrogen Energy* 2011, 36, 6948–6956, doi:10.1016/j.ijhydene.2011.02.087.


40. Poetzsch, D.; Merkle, R.; Maier, J. Proton uptake in the H+-SOFC cathode material Ba 0.5 Sr 0.5 Fe 0.8 Zn 0.2 O 3−δ: Transition from hydration to hydrogenation with increasing oxygen partial pressure. *Faraday Discuss.* 2015, 182, 129–143, doi:10.1039/C5FD00013K.
