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Short communication

# High-performance anode-supported Solid Oxide Fuel Cells based on $Ba(Zr_{0.1}Ce_{0.7}Y_{0.2})O_{3-\delta}$ (BZCY) fabricated by a modified co-pressing process

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#### 1. Introduction

Oxide proton conductors are promising electrolytes for low-temperature SOFCs because of their relatively high ionic conductivities at low temperatures [1–8]. To date, electrolytesupported cells were commonly used due to difficulties associated with fabrication of anode-supported cells, leading to much lower performance in comparison to anode-supported cells using yttriastabilized zirconia (YSZ) or gadolinia doped ceria (GDC) as the electrolyte [1,9]. For example, isostatic pressing at 216 MPa followed by sintering at 1600 °C for 10 h was required to form dense BaCe<sub>0.8</sub>Gd<sub>0.2</sub>O<sub>3-d</sub> pellets of ~1 mm thick [10]. These electrolytesupported cells with La<sub>0.6</sub>Sr<sub>0.4</sub>COO<sub>3- $\delta$ </sub> cathodes demonstrated a peak power density of ~5.5 mW cm<sup>-2</sup> at 700 °C under typical fuel cell operating conditions.

A reactive sintering process was recently developed for fabrication of thin and dense proton conducting electrolyte films on Ni-based anodes at relatively low temperatures (~1400 °C) [11–14]. The reactions among precursors (BaCO<sub>3</sub>, CeO<sub>2</sub>, and Y<sub>2</sub>O<sub>3</sub>) in the anode and electrolyte layers facilitated densification of the electrolyte films. However, it is noted that the conductivities of these electrolyte films are far lower than the anticipated values. For example, a 25  $\mu$ m thick Ba(Zr<sub>0.1</sub>Ce<sub>0.7</sub>Y<sub>0.2</sub>)O<sub>3- $\delta$ </sub> (BZCY) electrolyte showed an area-specific Ohmic resistance of ~0.5  $\Omega$  cm<sup>2</sup> at 700 °C [12], corresponding to conductivity of 0.005 S cm<sup>-1</sup>; however, the

### ABSTRACT

A modified co-pressing process was developed to fabricate anode-supported dense and uniform  $Ba(Zr_{0.1}Ce_{0.7}Y_{0.2})O_{3-\delta}$  (BZCY) electrolyte films (~20 µm thick) from BZCY powders with different characteristics; the powders derived from a glycine nitrate process was used for the anode whereas the powders from solid state reaction for the electrolyte. The BZCY electrolyte films sintered at 1350 °C for 6 h reached a conductivity of ~0.025 S cm<sup>-1</sup> at 700 °C, similar to that of BZCY pellet sintered at 1550 °C for 10 h. Further, a test cell based on such an anode-supported BZCY electrolyte demonstrated peak power densities of ~780 and ~490 mW cm<sup>-2</sup> at 700 and 600 °C, respectively.

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conductivity of a properly sintered BZCY pellet at 700 °C should be ~0.021 S cm<sup>-1</sup> [4]. The large discrepancy in conductivity is due probably to the non-stoichiometry in BZCY resulting from complicated synthesis reactions at 1400 °C, especially Ba evaporation from carbonate during sintering [15,16]. It was reported that BaCO<sub>3</sub> experienced maximum mass loss at ~1200 °C and the resultant BaO is extremely volatile at this temperature [17]. In contrast, the Ba loss in BaCeO<sub>3</sub> is relatively small even at ~1500 °C [13]. Electrical conductivity of cerates and zirconates depends sensitively on stoichoimetry; small deviation from desired composition might dramatically reduce conductivity, sinterability, and chemical stability in atmospheres containing H<sub>2</sub>O and CO<sub>2</sub> [18,19]. Therefore, it is necessary to develop a process for fabrication of BZCY films supported by anode while maintaining the desired compositions and properties.

In this communication, we report a modified co-pressing, cofiring process for fabrication of anode-supported BZCY electrolyte films with high electrical conductivity (~0.025 S cm<sup>-1</sup> at 700 °C) at relatively low sintering temperature (1350 °C for 6 h) using BZCY powders derived from a glycine nitrate process for the anode whereas the powders from solid state reaction for the electrolyte. This unique fabrication process produced uniform, homogeneous, and dense BZCY films of ~20  $\mu$ m supported by anode; single cells based on these anode-supported BZCY films demonstrated much higher power densities.

#### 2. Experimental

Two different methods were used to prepare  $Ba(Zr_{0.1}Ce_{0.7}Y_{0.2})O_{3-\delta}$  powders. In the solid state reaction (SSR) method, stoichiometric amounts of high-purity barium carbonate,

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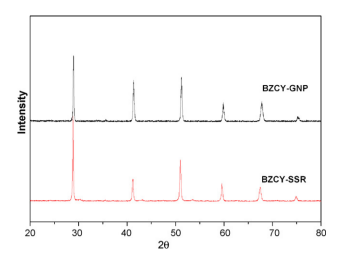
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zirconium oxide, cerium oxide, and yttrium oxide powders (all from Aldrich Chemicals) were mixed using ball milling with stabilized zirconia media in ethanol for 48 h. The resultant mixture was dried at 60°C for 24 h, followed by calcination at 1100°C for 10 h. The ball milling and calcination under same conditions were repeated twice to obtain pure phase. In the glycine nitrate process (GNP), suitable amounts of  $Ba(NO_3)_2$ ,  $Ce(NO_3)_3$ ,  $Y(NO_3)_3$ , and ZrO(NO<sub>3</sub>)<sub>2</sub> (all from Aldrich Chemicals) were dissolved in distilled water (0.1 M) and mixed with glycine (molar ratio of  $NO_3^-$  to glycine = 1.5:1). Then the solution was heated on a hot plate to vaporize water, converted to gel, and finally ignited to flame. The resultant yellow ash was then fired at 900 °C for 2 h to obtain BZCY powders. In subsequent discussions, BZCY-SSR stands for BZCY powders prepared by a solid state reaction process whereas BZCY-GNP for those derived from a glycine nitrate process.

A co-pressing and co-firing process was then used to prepare anode-supported electrolyte bilayers. Mixed powders of NiO, BZCY–GNP, and rice starch (from Aldrich Chemicals) with a weight ratio of 65:35:10 was pre-pressed at 100 MPa in a stainless steel mold as the substrate. Subsequently the BZCY powders from SSR were added to the surface of green anode through a screen (500 meshes) to guide uniform distribution of BZCY powders on the anode. Without a screen, it is difficult to uniformly and fully cover the anode when the amount of BZCY electrolyte powders was relatively small to make the electrolyte layer sufficiently thin. The anode-supported electrolyte bilayers were uniaxially co-pressed at 250 MPa, followed by sintering at 1350 °C for 6 h.

Composite cathode consisting of Sm<sub>0.5</sub>Sr<sub>0.5</sub>CoO<sub>3- $\delta$ </sub> (SSC, derived from GNP as describe elsewhere [3]) and BZCY (weight ratio of 7:3) were prepared as follows. The mixture of SSC and BZCY was mixed with V006 and acetone (weight ratio of 1:1:1) to form a cathode slurry, which was brush-painted on the BZCY electrolyte. The composite cathode was then fired at 1000 °C for 3 h. All the cells in this report utilized same cathode; the only difference is the various combinations of BZCY powders in anode and electrolyte. For simplicity, cells were labeled with the process method used for fabrication of the BZCY powders. For example, GNP/SSR denotes the cell that used BZCY powder derived from GNP for the anode and BZCY powder from SSR for the electrolyte.

The phase of BZCY was examined by XRD (PW-1800 system,  $2\theta = 20-80^{\circ}$ ). The morphologies of the cell components were observed using a Hitachi S-800 scanning electron microscope. All standard electrochemical experiments were performed using a Solartron 1286 electrochemical interface and a Solartron 1255 HF frequency response analyzer.



**Fig. 1.** X-ray diffraction patterns of the BZCY powder synthesized by glycine nitrate process and solid state reaction.

#### 3. Results and discussion

Fig. 1 shows typical X-ray diffraction patterns of the BZCY powders prepared by solid state reaction and glycine nitrate process, suggesting that single phase was obtained using both methods. This is required to ensure high conductivity and stability. Shown in Fig. 2 are the typical morphologies of BZCY powders by the two methods. The BZCY powder prepared by the SSR process has relatively high filling density; each individual particle is relatively dense and dry pressing of the powder may result in high packing density of green body. In contrast, the BZCY powder derived from a GNP process are highly porous; the pore sizes ranged from tens of nanometers to sub-micrometers. The low filling density of the GNP powders would result in large amount of shrinkage during firing. When used in the anode support, the BZCY powder derived from a GNP process would lead to accelerated densification of the electrolyte during firing: it may help to increase triple phase boundary length of the anode as well

Other combinations of BZCY powders were also examined under identical co-pressing and co-firing conditions. Fig. 3(a) shows the cross-sectional morphology of a cell using BZCY–SSR powders in both anode and electrolyte. Obviously, some pinholes were observable in the BZCY electrolyte, due primarily to insufficient shrinkage ( $\sim$ 12% in diameter) after sintering. Higher sintering temperature will yield denser electrolyte, but may cause deactivation of the anode. Fig. 3(b) shows the microstructures of an anode-

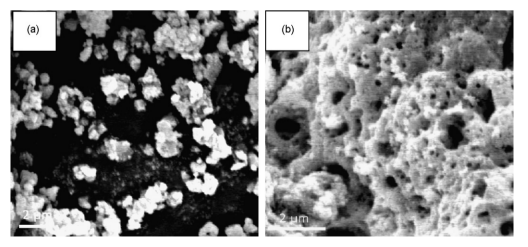


Fig. 2. Morphologies of BZCY powders prepared by (a) a solid state reaction and (b) a glycine nitrate process.

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