Direct foamed and nano-catalyst impregnated solid-oxide fuel cell (SOFC) cathodes

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A binder system containing polyurethane precursors was used to in situ foam (direct foam) a (La$_{0.6}$Sr$_{0.4}$)$_{0.98}$ (Co$_{0.2}$ Fe$_{0.8}$)O$_3$ (LSCF) cathode composition upon a yttrium-stabilized zirconia (YSZ) electrolyte coated with a porous ~10 µm thick cathode active layer. The YSZ electrolyte was ~110 µm in thickness, and a full cell was created by application of a Ni((Ce$_{0.8}$Ga$_{0.2}$)O$_3$) cermet at the baseline anode. Cells possessing the foamed LSCF cathode were compared to cells constructed via standard methods in terms of resultant microstructure, electrochemical performance, and introceptive character. The foamed cathode tended to possess a high level of tortuous porosity which was ellipsoidal and interconnected in character. Both the standard and foamed cathode structures were subjected to an infiltration process, and the resultant microstructure was examined. The impregnation efficiency of the foamed cathode was at least ~10% greater per deposition than that of an unfoamed porous LSCF cathode. The SOFC with the Pt nano-catalyst impregnated foamed cathode demonstrated a maximum power density of 593 mW/cm$^2$ utilizing wet H$_2$ fuel, which is 52% higher than a SOFC with the baseline Pt-impregnated LSCF cathode (~390 mW/cm$^2$) at 800 °C. The cathode compositional and microstructural alterations obtainable by foaming led to the elevated power performance, which was shown to be quite high relative to standard SOFCs with a thick YSZ electrolyte.

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1. Introduction

Solid oxide fuel cells (SOFCs) are attractive systems for the efficient electrochemical conversion to electricity of the chemical energy stored in gas and liquid fuels such as hydrogen, carbon monoxide, methane, coal syngas, and liquid hydrocarbon fuels. One issue that limits the commercialization of SOFC systems is related to the long-term degradation of the fuel cells, which can be correlated to the high operation temperature. In order to lower the operation temperature, and potentially increase cell life, the oxygen reduction reaction (ORR) at the SOFC cathodes must be improved. A recent strategy used to enhance oxygen reduction kinetics of traditional (La,Sr)MnO$_3$ (LSM) and LSCF cathodes is through the liquid impregnation of a nano-catalyst within the final sintered cathode microstructure [1]. The inclusion of the nano-catalyst extends the overall triple-phase boundary (TPB) area and alters the oxygen reduction mechanism by providing alternative adsorption and ion/electron spillover sites. A few reports indicated up to a two times improvement in fuel cell performance for Cu and Ce oxide nano-catalyst within the cathode microstructure [2–5]. The nano-catalyst is typically added by dropping or dip-coating a liquid based solution or dispersion of the nano-catalyst, which leaves a dispersion of the particle distributed over the pre-sintered cathode microstructure. A low temperature-firing step is usually required to bond the nano-catalyst to the pre-sintered cathode backbone microstructure.

Previous literature demonstrates the effectiveness of the nano-catalyst impregnation in SOFC cathodes for improving oxygen reduction reaction [1,5–11]; unfortunately, the impregnation process may be time and labor intensive to incorporate the required nano-catalyst content within the open porosity. A well connected deposition of nano-catalyst into an electrode is typically only achieved by repetitive impregnation steps, which leads to more costly processing [11]. In addition, as the open cathode porosity is filled with the precipitated salts, the ability to infiltrate the porous structure to the active cathode (near the electrolyte interface) becomes more of an issue [12]. In this work, a novel in situ direct foaming process was utilized to form open and interconnected porosity throughout the cathode current collector. This porosity provided a microstructure to retain high intrinsic cathode performance, while at the same time, providing an open
porous network to permit the efficient deposition of nanocatalyst to the active cathode area.

2. Experimental

The electrolyte-support membranes were fabricated from 8 mol% YSZ powder (Daichi Kigen Kagaku Kogoyo Co., LTD, Japan) by a tape-casting, lamination process, and sintering (∼110 µm final thickness). A ~3 µm thick Ce0.9Gd0.1O2 (GDC) buffer layer was incorporated between the electrolyte and both electrodes by screen-printing and firing onto the electrolyte at 1350 °C for 1 h. The GDC used in this study was synthesized using a conventional co-precipitation method [12]. A 50 vol% NiO (HPGNO, Novamet, Wyckcoff, NJ)-50 vol% GDC composite was mixed with an ink vehicle (J2M, 63/2, Johnson Matthey, UK). The ink was screen-printed and sintered at 1350 °C for 2 h to a thickness of ∼50 µm. The LSCF powder was synthesized by a solid-state process and attrition-milled to an average surface area of 5 m²/g. A 50 vol% LSCF-50 vol% GDC composite was formed as an active cathode composition and was printed on the cathode side of the cell (~10 µm thickness). For the baseline samples, a pure LSCF ink was printed over the active area and sintered to 1150 °C for 1 h. The alternative cathode architecture was fabricated by printing a LSCF ink containing polyurethane precursors, which were similar to precursors previous demonstrated for direct foaming filled polyurethanes [13,14]. The ink was composed of LSCF powder and 8:4:1 polymer precursor composition (isocyanate:polyol:surfactant). The precursor materials used in this work were polymethylene isocyanate (Volanate M220, Dow Chemicals), polyethylene glycol (PEG200, Aldrich) and polyoxyethylene sorbitan monooleate (Tween 80 Fluka). These polymer chemicals), polyethylene glycol (PEG200, Aldrich) and polyoxymethylene isocyanate (Volanate M220, Dow (isoscyanate:polyol:surfactant). The precursor materials used in this work were polymethylene isocyanate (Volanate M220, Dow Chemicals), polyethylene glycol (PEG200, Aldrich) and polyoxymethylene sorbitan monooleate (Tween 80 Fluka). These polymer components were added to the J2M ink vehicle and printed over the active layer within an argon filled glove box. On exposure to ambient atmosphere (relative humidity 45–55%), the polymerization reaction was initiated, and the CO₂-blowing reaction resulted in the foaming of the printed LSCF structure. The LSCF/polymer thin film gel was then fired in a similar manner as the baseline (unfoamed) cathode microstructure (1150 °C for 1 h).

An aqueous platinum catalyst (H₂PtCl₆-6H₂O, Alfa Aesar-Pemion) solution was prepared at a 0.1 M concentration, and this solution was used for the impregnation process. This solution was impregnated into the cathode with micro-pipette (10–100 µL, Eppendorf International). The calcination of the impregnated cathode was performed at 850 °C for 1 h. The weight change of the impregnated structure was monitored until the catalyst level reached 5 wt% of cathode initial weight. The scanning electron microscope (SEM) imaging and energy-dispersive spectroscopy (EDS) analysis of the half-cells with the above-mentioned compositions were performed using a JEOL 7600 SEM microscope.

The SOFCs were mounted on an alumina tube fixture with a pair of 9 × 9 mm platinum mesh and wire as the current collectors for both electrodes. Six light dots of LSCF and Ni metal inks were used to secure the interconnect leads on the cathode and anode, respectively. The cell was then heated to 800 °C at 1–2 °C/min. under 60 sccm of argon gas within the anode chamber and ambient air within the cathode chamber. After the cell reached 800 °C, the anode atmosphere was slowly transitioned to 100 sccm of moist H₂ (3% H₂O). Cell power curves (V–I–P curves) data was collected using Solatron SI-1287 interface and an SI-1252 frequency response analyzer for the electrochemical impedance spectroscopy (EIS).

3. Results and discussion

Initial direct foaming experiments were completed with a low LSCF solids loading within the organic carrier. Fig. 1a displays the SEM micrograph of a cathode generated from a 20 vol% LSCF loading in the ceramic-polymer precursor suspension. The average density for this sample measured by image analysis of SEM micrographs (using National Institute of Health (NIH) ImageJ) was ~14% theoretical density (86% porosity) with > 20 µm pore size. In a subsequent specimen, the LSCF solids loading was increased to 70 vol%, while still maintaining the same precursor composition stated above. Fig. 1b shows a different resultant microstructure with an array of elongated or interconnected porous channels and a mixture of fine porosity between the LSCF particles. A polished cross-section of the foamed LSCF cathode on the substrate was analyzed using ImageJ with a subprogram developed by Impoco et al. [15]. The average porosity level was found to be 46.5% calculated from nine images at various magnifications (500–2000 ×) across the film thickness. The calculated two-dimensional mean pore perimeter length was 17.7 ± 7.54 µm, with an average two-dimensional pore area of ~15.9 µm² and a circularity factor of 6.9. The circularity factor was calculated by dividing the pore perimeter value by the Ferret’s diameter, with a circularity factor farther from π (3.14) indicating a departure from an isotropic pore shape towards an interconnected and elongated channel shape. For comparison, an unfoamed LSCF cathode fired at 1150 °C onto the YSZ electrolyte (using no polyurethane precursor mixture) displayed an average porosity level of 43.1% with a two-dimensional pore area of 7.9 µm² and a circularity factor of 3.1. Interestingly, the average pore size for both samples was ~4 µm.

The microstructures of the in situ foamed and baseline cathodes were then impregnated with nano-catalyst. The nano-Pt composition was chosen for its high stability and low reactivity.
with the LSCF composition; therefore, the processing benefits can be directly evaluated without misinterpretation due to potential chemical interaction or surface modification of the LSCF cathode backbone (as is the case for nano-Ag or -Cu oxide incorporation) [16–18]. The amount of catalyst per infiltration step was monitored. The primary calcinations were performed after the 1st and 2nd infiltration steps at 450 °C for 1 h, which resulted in the production of Pt/PtOₓ and some carbonaceous content. Final calcination was completed after further infiltration cycles (at 850 °C for 1 h). Fig. 2 displays the accumulative normalize weight after each impregnation/calcination step. The catalyst weight was normalized as weight per volume to account for slight differences in thickness between the foamed and baseline cathodes. The sudden decrease in weight after the 3rd step reflects the removal of the carbonaceous content from all steps to that point. The foamed cathode showed the ability to accept a higher catalyst solution volume (and thus, higher solid nano-catalyst amount) after every step with similar porosity level (~40–45% theoretical).

The average amount of the precursor solution added per impregnation step was 2.45 ± 0.25 g/cm³ over 5 impregnation steps, whereas for the baseline was 0.845 ± 0.12 g/cm³. The results are well correlated with the above-discussed microstructural characteristics, where the interconnected and high porosity of foamed cathode assisted in providing open channels for efficient impregnation.

The voltage–current–power (V–I–P) performance data for the button cell SOFCs were tested with foamed and unfoamed LSCF cathodes using moist H₂ fuel at 800 °C. The SOFCs were mounted on an alumina tube and interconnected with Pt leads. The area specific resistance (ASR) values were calculated from the slope of the linear portion for the voltage–current density data. The performance of the SOFC with in situ foamed cathode showed a significant improvement over the baseline (unfoamed) cell. The cell with the foamed cathode displayed an average maximum power density of 514 mW/cm² (ASR = 0.52 Ω cm²), which is 43% higher than the cells with the baseline cathode (360 mW/cm², ASR = 0.73 Ω cm²), as shown in Fig. 4. For reference, the cell's electrolyte should contribute ~0.30 Ω cm² to the total ASR for the stated thickness at 800 °C. The same cathode microstructures were impregnated with the Pt precursor using the five-step

![Fig. 2. Normalized weight of nano-Pt catalyst incorporated into the microstructure of the baseline and foamed cathode after each impregnation step.](image)

![Fig. 3. SEM micrograph of the impregnated (a) baseline (unfoamed) and (b) foamed LSCF cathode at a distance of ~2–5 μm from the active cathode layer.](image)

![Fig. 4. V–I–P performance curves of SOFC button cells that possessed a LSCF cathode that was unfoamed, foamed, unfoamed/Pt-impregnated, and foamed/Pt-impregnated. The SOFCs were measured at 800 °C with wet-H₂ fuel.](image)
impregnation and thermal processing method. It must be re-stated that the foamed sample at this point contained nearly twice the loading of the baseline sample due to the restricted ability of this structure to accept the catalyst impregnation solution (Fig. 3). The cell with the impregnated foamed cathode displayed a maximum power density of 593 mW/cm² (ASR ≈ 0.46 Ω cm²), which is 14% higher in power than the un-impregnated foamed cathode (514 mW/cm²). This performance increase is twice that demonstrated recently by other researchers for Pt-enrichment of LSCF and LSM cathodes (~7–8%) at 800 °C [8,18,19]. The SOFC baseline cell with a Pt-impregnated LSCF cathode displayed a maximum power density of ~390 mW/cm² (ASR ≈ 0.71 Ω cm²) at 800 °C. This accounts for ~7.7% higher power performance value over that of an un-impregnated baseline cell (~360 mW/cm²), as shown in Fig. 4. This improved performance can be attributed to the higher content of the Pt loading within the electrode and the homogeneous dispersion of fine catalyst particles deep within the cathode microstructure. Again, it must be stated that the loading was achieved with a higher degree of processing efficiency, leading potentially to a lower number of impregnation steps in the future to achieve a desired loading level.

4. Conclusion

The use of an in situ foaming process was used for the first time to fabricate porous LSCF cathode architectures with a mixed pore size range and a tortuous porous microstructure. The average pore area and circularity factors were twice as high as that demonstrated for an unfoamed LSCF cathode. The microstructure formed by this foaming process resulted in an electrode with a much lower cathodic polarization and an increase in maximum power performance by ~43%. The open and interconnected cathode pore structure also provided a higher level of efficiency in the impregnation of the cathode, which resulted in increased oxygen reduction kinetics. Future optimization of the polyurethane precursor composition and content, and the use of alternative surfactants within the cathode inks may permit distinct control of the blowing reaction. These alterations may potentially allow for the microstructural design of the cathode structure, specifically designed for increased TPB population, pore distribution/gradient, oxygen mass flow, and nano-catalyst incorporation.

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