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Performance and electrochemical analysis of solid oxide fuel cells based on LSCF-YSZ nano-electrode

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Abstract: NiO-YSZ /YSZ/ LSCF-YSZ tri-layer structure SOFC has been fabricated by tape casting and infiltration methods. Subsequently, polarization curves and electrochemical impedance spectra measurement were carried out to evaluate cell performance at 850-700\textdegree C with varied steam content in hydrogen supplied to anode. At 800\textdegree C, open circuit voltage for 4\% steam humidified hydrogen has reached 1.069V, along with power density of 831mW cm\textsuperscript{-2} at 0.7V. Furthermore, 500h of long-term durability test has been performed at 750\textdegree C in hydrogen atmosphere, as a consequence that a voltage degradation rate of 1.7%/kh has been measured.

Keywords: LSCF-YSZ, SOFC, electrochemical analysis, DRT

I Introduction

Solid oxide fuel cell (SOFC) has attracted increased attention as energy-converting
device, which can directly convert chemical energy to electricity. [1] Because of its high electronic and oxygen ion conductivities at low temperatures, the strontium- and cobalt-doped lanthanum ferrites, La$_x$Sr$_{1-x}$Co$_y$Fe$_{1-y}$O$_{3-\delta}$ (LSCF), have been widely used as cathode materials for intermediate and low temperature SOFCs. Yttria-stabilized zirconia (YSZ) is the state-of-the-art electrolyte material for SOFC. Nickel oxide (NiO)-YSZ as anode material in combination with YSZ electrolyte and LSCF cathode has been investigated by many researchers [2-6].

To decrease the sintering temperature of the cell, lots of works have been focused on infiltration method, which is used to fabricate the cathode of SOFC. Liu et al. prepared a tri-layer YSZ structured SOFC infiltrated with LSCF and Pd-SDC respectively acting as cathode and anode, which shows good power density of 935mW cm$^{-2}$ at 800°C [3]. Fan et al. also prepared LSCF-YSZ nano-electrode with infiltration for reversible SOFCs [7], showing high catalytic activity towards oxygen reduction/oxidation reaction. However, analysis of electrochemical impedance spectra of infiltrated cells is rarely reported [8]. What’s more, long-term stability of the electrode fabricated by infiltration method is a critical issue. It has been reported that coarsening of infiltrated nano-particles and solid reactions between the infiltrated and scaffold material are the potential causes for the degradation [9, 10]. Meanwhile, long-term stability results of infiltration electrodes are rarely published.

Distributions of relaxation times (DRT) is a technique for deconvolution of a single impedance spectrum into a distribution of relaxation times, invented by Ivers-Tiffée and coworkers [11]. According to DRT method, different electrode processes in SOFC
can be identified. The impedance $Z$ for angular velocity $\omega$ can be decomposed as

$$Z(\omega) = R_0 + Z_{pol}(\omega)$$

$$= R_0 + R_{pol} \int_0^\infty \frac{\gamma(\tau)}{1 + j\omega\tau} d\tau = R_0 + R_{pol} \int_0^\infty \frac{G(\tau)}{1 + j\omega\tau} d\ln \tau$$

in which $R_0$ is ohmic resistance irrelevant to frequency, $Z_{pol}(\omega)$ is the polarization part of total impedance. By using DRT technique, different physical and chemical procedures are shown as different peaks with characteristic frequencies, which can be distinguished with the change of test conditions.

In this study, nano-structural LSCF-YSZ electrode for planar SOFC was fabricated with infiltration method. After that, analysis of electrochemical impedance spectra was carried out using DRT method to evaluate the performance of infiltrated cells. Long-term stability test was also implemented with humidified hydrogen at 750°C for 500h. EIS data was fitted to an equivalent circuit model by the Complex Nonlinear Least Square (CNLS) method. Finally, scanning electron microscope (SEM) photos of test cells before and after long-term stability test were given.

II Experimental

2.1 Preparation of nano-structured NiO-YSZ/YSZ/LSCF-YSZ cells

The procedure for preparation of tested SOFC is mentioned in ref. [12]. The cell was fabricated by tape casting and infiltration methods. The NiO-YSZ/YSZ/porous YSZ substrate was firstly made by tape casting and hot isostatic pressing technology. YSZ (8 mol% yttria stabilized zirconia, Tosoh Company, Japan), NiO powder and graphite (Furunda Zirconium Material Co. Ltd., China) as pore former were mixed in a
weight ratio of 50:50:10 and ball milled with ethanol-butanone solvent, castor oil dispersant, dibutyl phthalate (DBP) plasticizer, and polyvinyl butyral (PVB) binder in a two-stage milling process mentioned in [7]. For preparation of the YSZ electrolyte slurry, almost the same method was used, except for no NiO and graphite pore former. And the porous YSZ layer as cathode was also similar with 50wt% graphite to form backbones. After tape casting and hot isostatic pressing mentioned in [3], the tri-layer structure cell substrate was co-sintered in air at 1300°C for 10h to get the YSZ electrolyte layer densified. After that, LSCF \((\text{La}_{0.6}\text{Sr}_{0.4}\text{Co}_{0.2}\text{Fe}_{0.8}\text{O}_{3-\delta})\) precursor as the solution of infiltration was prepared by dissolving stoichiometric amounts of \(\text{La(NO}_3\text{)}_3\cdot6\text{H}_2\text{O}, \text{Sr(NO}_3\text{)}_2, \text{Co(NO}_3\text{)}_2\cdot6\text{H}_2\text{O}\) and \(\text{Fe(NO}_3\text{)}_3\cdot9\text{H}_2\text{O}\) in deionized water and ethanol. [13] The infiltration process was executed by infiltrating solution into pre-sintered YSZ backbone. After infiltration and dried in air at room temperature, the substrate was sintered at 450°C for 2h to let nitrate decompose to oxide. This procedure was repeated for multiple times until the loading amount of LSCF reaching 30~50 wt% and finally the cells were sintered at 900°C for 2h to obtain nano-structured cathode.

2.2 Procedures of cell characterization, electrochemical measurement and long-term test in hydrogen atmosphere

The prepared NiO-YSZ/YSZ/LSCF-YSZ cells were cut with an effective working area of 4×4 cm². The cells were tested in a SOFC test setup mentioned in [14]. Gold and double-layer nickel meshes were used as current collectors on the oxygen and fuel side, respectively.
Before testing, NiO-YSZ anode was reduced at 850°C in safety hydrogen (9% hydrogen, 91% nitrogen) for 2h. The electrochemical measurement procedures of the cell were executed at temperatures from 850°C to 700°C with a decrement of 50°C, during which 24L/h hydrogen with 4%, 20% and 50% steam content supplied to NiO-YSZ anode, and 140 L/h air supplied to LSCF-YSZ cathode for each temperature. To protect test cells, the lowest voltage applied in i-V test was limited to 700mV. Electrochemical impedance spectroscopy (EIS) of test cells was measured at OCV by using Solatron 1260 frequency analyzer and external shunt in series with the cell. 12 points per decade was recorded from 96,850 Hz to 0.08 Hz. Analysis of impedance data via DRT method was carried out using Ravdav [15]. 500h long-term stability test was examined at 750°C by introducing 24L/h hydrogen with 4% steam content to anode and 140 L/h air to cathode, at the current density of 0.25 A cm\(^2\). EIS data was collected

Fig. 1 Performance of test cell at 800°C with different fuel components supplied to anode side
during test at OCV with an AC amplitude of 20 mV.

Fig. 2 Performance of test cell at different temperatures with 20% steam hydrogen supplied to anode side

III Results and discussion

3.1 Cell characterization of polarization curves

Fig. 1 shows the cell voltage and power density change as function of current density at 800°C with different steam contents of hydrogen to anode side. From Fig. 1, it can be seen that peak power densities of the cell were about 831 mW cm\(^{-2}\), 738 mW cm\(^{-2}\) and 544 mW cm\(^{-2}\) at 0.7V for hydrogen with 4%, 20% and 50% steam supplied to anode, respectively. Open circuit voltage for 4% steam humidified hydrogen reached 1.069V and maximum current density reached 1.2 A cm\(^{-2}\), indicating densified electrolyte and relatively low resistance. Moreover, the activation loss for 4% steam humidified hydrogen is apparently higher.
Fig. 2 shows i-V curves at different temperatures from 850°C to 700°C with 20% steam humidified hydrogen. The open circuit voltages reached 0.985V, 0.994V, 1.004V and 1.016V at 850°C, 800°C, 750°C and 700°C, respectively. Maximum power density of 898mW cm² was measured at 850°C. Resistance of test cell increased as expected with the decrease of temperature with ohmic resistance ($R_s$) of 0.08, 0.11, 0.15, 0.22 Ω cm² and polarization resistance ($R_p$) of 0.22, 0.28, 0.39, 0.64 Ω cm² at 850°C, 800°C, 750°C and 700°C, respectively, which can be deduced from Fig. 4.

3.2 Electrochemical impedance spectra measurement

Fig. 3 and Fig. 4 show the EIS results related to Fig. 1 and Fig. 2, respectively. From Fig. 3, it is clear that steam content influenced polarization resistance significantly, whereas ohmic resistance remained constant. Both high frequency and low frequency arcs in spectra were varied with the change of steam content, indicating that the fuel composition has an effect on almost all physical and chemical procedures in tri-layer structure fuel cells tested, especially for low frequency arc. On the other
hand, temperature exerted an influence on both ohmic and polarization resistance as expected. Almost no influence of temperature can be seen from low frequency arc in spectra, with significantly rise of high frequency arc as the temperature decreased. This illustrated different temperature reaction of physical and chemical processes.

DRT analysis of EIS data in Fig. 3 and Fig. 4 are shown in Fig. 5 and Fig. 6. Five peaks can be observed in DRT curves, namely Peak I to Peak V starting from high frequency. According to results by other researchers, Peak I to V corresponded to ionic transport [16], Triple Phase Boundary (TPB) reaction [17], resistance of O²⁻ diffusivity [17] in cathode, diffusion [18] and conversion [19] in anode, respectively. According to Fig. 5, Peak I remained constant when steam content changed in anode side, indicating ionic transport on cathode side, which is different from the results in [18]. All the other 4 peaks are related to the change of fuel compositions in anode side. And their characteristic frequencies all increased with the increase of steam content in hydrogen. For Fig. 6, two groups can be distinguished clearly. Peak I and II are
temperature-dependent peaks, and the heights of peaks increased with the decrease of temperature. The characteristic frequency of Peak II decreased at the same time, while that of Peak I remained unchanged. Peak III, IV and V are temperature-independent peaks, which means their peak heights and characteristic frequencies remained unchanged with temperature decrease, indicating that the related processes in test cells are temperature-independent. The peaks and their characteristic frequency ranges are listed in Table 1.

![Fig. 6 DRT analysis of EIS data of test cell at different temperatures with 20% steam hydrogen supplied to anode side](image)

<table>
<thead>
<tr>
<th>Peak No.</th>
<th>I</th>
<th>II</th>
<th>III</th>
<th>IV</th>
<th>V</th>
</tr>
</thead>
<tbody>
<tr>
<td>characteristic frequency range (Hz)</td>
<td>10k~50k</td>
<td>1k~10k</td>
<td>100~200</td>
<td>10~40</td>
<td>1~4</td>
</tr>
</tbody>
</table>

Table 1 characteristic frequency range of peaks in DRT analysis of test cells
3.3 Long-term test in hydrogen atmosphere

500h test with 4% steam humidified hydrogen as fuel at 750°C was performed after cell electrochemical characterization. The current density was set to be 0.25 A cm\(^{-2}\). I-V result is shown in Fig. 7. Cell voltage dropped from 0.915V at the beginning to 0.907V at the end of the test, indicating a voltage degradation rate of 1.7%/kh, close to similar tests [20]. EIS results during long-term test are presented in Fig. 8. As it can be seen, ohmic resistance increased slightly from 0.17Ω cm\(^2\) to 0.18Ω cm\(^2\), while polarization resistance increased from 0.31Ω cm\(^2\) to 0.33Ω cm\(^2\), mainly in high frequency arc.
DRT analysis of EIS data is presented in Fig. 9. As shown in the graph, increase of impedance is mainly due to the increase of Peak II, while Peak V showed a slightly tendency of decrease during test. The characteristic frequency of Peak II also decreased slightly with time, which by far has not been noticed before. Previous research has mentioned relation between the long-term performance degradation and the coarsening of microstructure of test cell, which leads to reduction of TPB [7, 21-23]. These corresponded to the phenomenon of Peak II, which was possibly the potential cause of degradation of cell performance.

Each resistance was fitted by the CNLS method using an equivalent circuit of Fig. 10. Resistances and constant phase elements were combined to simulate electric behavior of Peak I, II, IV and V, while a Gerischer element was used to simulate behavior of $O^2-$ diffusion in cathode, which was represented by Peak III.
Fig. 11 Fitting results of impedance data of long-term stability test

Fitting results are shown in Fig. 11. We can see that the resistance of Peak II (R₃) contributed most to the increase of impedance of test cell. It increased from 0.202Ω cm² at 0h to 0.224Ω cm² at 500h. Ohmic resistance (R₁) also increased with time from 0.146Ω cm² to 0.154Ω cm². The resistances of other peaks were all below 0.1Ω cm² and their contribution to the change of total impedance can be ignored. Changing rates of every resistance are listed in Table 2. It has to be mentioned that the changing rate for R₂, R₄, R₅, R₆ are more likely to be affected by random data error because of relatively smaller resistance value.

Table 2 changing rates of resistance in the equivalent circuit in long-term test

<table>
<thead>
<tr>
<th>resistance</th>
<th>R₁</th>
<th>R₂</th>
<th>R₃</th>
<th>R₄</th>
<th>R₅</th>
<th>R₆</th>
</tr>
</thead>
<tbody>
<tr>
<td>changing rate (%/kh)</td>
<td>9.6</td>
<td>-2.4</td>
<td>24.8</td>
<td>-25.7</td>
<td>-6.76</td>
<td>-10.2</td>
</tr>
</tbody>
</table>

SEM photos of testing cell before and after long-term stability test are shown in Fig. 12. In the middle is the dense YSZ electrolyte layer, which is about 10mm in thickness. Above that is the LSCF/YSZ cathode layer and below is the YSZ/NiO anode.
layer. The vertical cracks in the electrolyte layer in Fig. 12b) was caused by cell dismounting. By comparison, no obvious change in cell structure can be observed, while the change of micro-structure needs to be investigated further due to lack of high resolution equipment.

Fig. 12 SEM photos of testing cell for long-term test
a) before test b) after test

IV Conclusion

Cell characterization, electrochemical measurement and 500h durability test were executed for NiO-YSZ/YSZ/LSCF-YSZ tri-layer structure SOFC. Power density of 831mW cm² at 0.7V at 800°C was reached. Five different reaction processes have been distinguished by DRT analysis. 1.7%/kh degradation rate was measured for 500 durability test, and coarsening of microstructure was suspected to be the potential cause for degradation.

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