Performance Investigation of a Nickel Cermet Anode Modified with Copper, Alkaline Earth Metal Oxide, Boron, and Perovskite for Direct Methane Fuel Cell

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Abstract

The metallic copper, alkaline earth metal oxide, boron, and perovskite were incorporated on the surface of a Ni-cermet anode, and the performance of the modified Solid Oxide Fuel Cell (SOFC) anode was evaluated. The cell performance was analyzed by voltage-current characteristics (V-I curve) and H2-CH4 step reactions (P-t curve) in a potentiostatic mode. Besides, we also determined if a metallic phase or high electronic conductivity of the anode is important for a cell to perform well when H2 is used as a fuel, whereas both conductivity and anti-coking capability are critical while using CH4 as a fuel. The results showed that the anodes containing magnesium oxide (MgO), lanthanum strontium titanate (La0.4Sr0.4TiO3-δ), and boron were relatively resistant to the degradation in the CH4 environment when compared with others. The underlying mechanism varied mainly with electronic and structural promotion by the dopants as well as their material compatibility with the Ni-cermet substrate. These findings were evidenced and supported by surface analysis as well as in-situ infrared and mass spectroscopic studies too.

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1. Introduction

The conversion of methane (CH$_4$) into valuable chemicals or electric power is of continuous interest in both academia and industry because of surging global energy demands. Methane is the predominant component of natural gas, while to date, the widely adopted route for the conversion is still the conventional steam methane reforming (SMR) over nickel-based catalysts [1]. Due to the endothermic nature, the SMR reaction is highly energy-intensive, along with the need for large capital investment. As a result, a more economically feasible way is pursued to reduce the cost and increase energy efficiencies. Solid oxide fuel cells (SOFCs) have received much attention in the past decades for the conversion of hydrocarbons such as methane and biogas to electric power with high efficiencies. SOFCs are operated at high temperatures, where modest reaction kinetics could be achieved using nonprecious metal catalysts [2]. Nevertheless, irreversible deactivation remains the major challenge over the state-of-the-art Ni cermet anodes because of carbon deposition and sulfur poisoning when methane is used as a fuel.

Several approaches have been investigated to circumvent the degradation of Ni cermet anodes for the utilization of hydrocarbons in SOFCs [3-6]. Application of steam reforming reaction to the Ni cermet anode inside SOFCs was first proposed. However, the development of internal steam reforming was found to be hindered since a high ratio of steam/methane (H/C) was required to depress carbon accumulation. The high proportion of steam leads to dilution of fuel with low energy efficiency and adds to the complexity of heat management. In the past decade, various perovskites materials such as (LaSr)(CrMn)O$_3$ [5], (LaSr)TiO$_3$ [7], BaZrCeYYbO$_3$ [8], and Sr$_2$Mg$_{1-x}$Mn$_x$MoO [9] have been considered as alternative anode materials with great interest. These ceramic anode materials did not only show improved tolerance to carbon deposition in CH$_4$ but also presented good stability when subjected to redox cycles. However, cell performance with these anodes of Ni-free metal oxide was lower than those demonstrated by conventional Ni-cermet anode supported ones. The drawbacks were attributed to lack of active sites, low conductivity, material mismatch, and formation of undesirable phases [10, 11].

Surface modification or the addition of a protective top layer on the Ni-cermet anode is an effective method to avoid coking in Direct Methane Fuel Cells. Several groups and we have reported that adding a functional layer sustains a stable electrochemical performance of the Ni-cermet anode in CH$_4$ [12-16]. The layer either functions as a catalyst to decompose CH$_4$ or serves as a reforming zone for the conversion of oxidation products. In addition, the insertion efficacy of alkaline earth metal oxide on the Ni-cermet anode has also been demonstrated for the reduction in coking [17-20]. The inclusion of basic oxides increases the Lewis basicity of the anode and allows chemisorption of oxidation products, including H$_2$O and CO$_2$, with subsequent reactions and removal of deposited carbon [6, 18, 21, 22]. The content of basic oxide and the possible phase reaction in the anode composite have shown pronounced effects on the activity of methane oxidation and coking resistance. Nonetheless, the addition of basic oxides such as CaO, MgO, or BaO was effective in some studies, while the effects were not obvious in others [17, 19, 23]. Moreover, the addition of...
alkaline earth metal oxide to the anode is done through an impregnation method. Although the uniform distribution of the oxide particles can be achieved in the interstitial Ni sites by the wet method, the procedures could be tedious with multiple heating cycles.

On the other hand, the incorporation of rare-earth metal oxides and boron into a Ni-based catalyst resulted in a reduced coke formation [24-26]. Rare-earth metal oxides have been claimed to stabilize Ni particles on a catalyst support, acting as a structural promoter to improve the metal dispersion and to suppress the aggregation of Ni grains. The influence of boron on the activity and acidity of a catalyst has been known, mainly arising from better metal dispersion through structure control and metal-boron electronic interaction [27-29]. Nevertheless, no application of any boron promoter to a SOFC anode has been reported in the literature so far. In the present work, we modified the surface of a Ni-cermet anode with copper, alkaline earth metal oxide, boron, and perovskite via a solid-state reaction. The idea of using solid-state synthesis was motivated by the need to develop a simple way that can be adapted to current cell manufacturing methods such as tape-casting or die-pressing. The cell performance with these modified anodes was investigated by voltage-current characteristics and power density versus time in a potentiostatic mode. In situ surface analysis of these anodes was conducted by diffuse reflectance infrared spectroscopy (DRIFTS) to probe the mechanism responsible for coking resistance relevant to the electrode process in detail. The results obtained here gave a complementary study with those in literature and provided valuable information to design an effective anode material for fuel cell applications.

2. Experimental

2.1 Material Synthesis

All electrochemical tests were carried out on an electrolyte-supported SOFC. The cell was composed of electrolyte support (yttrium stabilized zirconia, 8 mol.% Y2O3 doped in ZrO2), a cathode (Lanthanum strontium manganite, LSM), a cathode interlayer (LSM/YSZ, 50/50 wt.%), an anode interlayer (NiO/YSZ, 50/50 wt.%), and an anode (modified NiO/YSZ, 70/30 wt.%). The composition of the modified NiO/YSZ anodes and their preparation is described below. The electrolyte-supported cell was fabricated by die pressing. The electrolyte (~ 4.5 g), anode interlayer (~ 2.5 g), and anode (~ 0.8 g) were pressed (3000 psi) together in a die to form a half-cell, which was then sintered and calcined at 1400 °C for four hours. The cathode slurry consisting of LSM/YSZ and pure LSM was then screen printed on the electrolyte side of the sintered half-cell and calcined at 1200 °C for two hours. The obtained bottom cell was circular in shape with a diameter of about 2 cm.

2.2 Modification and Characterization of NiO/YSZ Anode

Four types of modified NiO/YSZ anodes were prepared by incorporating metal, alkaline earth metal oxide, inorganic element, and perovskite. The metal modified anode was prepared by sputtering copper on the surface of the NiO/YSZ at a deposition rate of 3.5 Å per second (Plasma treatment system, KD-SPUTTR). Two different sputtering times of 20 and 60 min were applied to obtain an estimated thickness of 420 and 1260 nm, respectively. The other three materials were introduced by mixing an additive with NiO/YSZ in the anode layer (the amount was fixed at 1 wt.% of the anode layer unless otherwise specified), followed by pressing and calcination to alter surface properties. The following additives were mixed with NiO/YSZ i) alkaline earth metal oxide: calcium
hydroxide (Ca(OH)$_2$), calcium oxide (CaO), and magnesium oxide (MgO), ii) inorganic element: boric acid (H$_3$BO$_3$), and boron powder (B), and iii) perovskite: lanthanum strontium titanate (LST, La0.4Sr0.4TiO$_3$−γ). The LST powder was synthesized via a solid-state reaction [30, 31], whereas the boric acid was incorporated via a wet impregnation method. Briefly, boric acid was first dissolved in dilute nitric acid, added dropwise to the NiO/YSZ matrix, and then annealed in air at 500 ºC. The steps were repeated to achieve a loading of 1 wt.%. Surface structure and morphology analysis were conducted by a scanning electron microscope (SN 4000).

2.3 Electrical Performance

Figure 1 shows the testing apparatus used in the study. The cell was assembled with two silver foils (as a current collector) and attached to one end of a ceramic tube by silver conductive paste (Alfa Aesar). The area of the layer of silver conductive paste on the cathode was about 1 cm$^2$ and used as an active area for performance normalization. The cell was then placed in a furnace at room temperature and heated in a He (purity 99%) stream. As the temperature arrived at 750 ºC, fuel gas (50 vol.% H$_2$ and CH$_4$ diluted in He) was introduced from the bottom of the tube to the anode, while the cathode was left exposed to air. The electrical performance of each cell was measured by its voltage-current (V-I) curve using a potentiostat (VersaSTAT3–400). The comparison between the performances in H$_2$ and CH$_4$ was obtained by operating the cell in a potentiostatic mode and recording the output of the electric power.

![Schematic of a SOFC testing apparatus.](image)

Figure 1 Schematic of a SOFC testing apparatus.

2.4 In Situ Infrared Spectroscopy Studies

Diffuse Reflectance Infrared Fourier Transform Spectroscopy (DRIFTs) was used to investigate the reaction mechanism and intermediates of CH$_4$ oxidation on the modified NiO/YSZ anode. The IR spectra were collected by an IR spectrometer (Thermo Nicolet iS10) attached with a diffuse reflection accessory (the Praying Mantis™, Harrick). About 0.1 g of anode powder obtained after crushing and grinding the sintered anode disk was loaded into a high-temperature reaction chamber covered with a dome. The reactant gas containing CH$_4$ (10 sccm) and O$_2$ (2 sccm), both diluted with helium (50 sccm), was used without further purification. The composition of the reaction effluent from the DRIFTs reactor was monitored by a mass spectrometer (MS, Hiden Analytical, HPR 20). The
mass/electron ratios (m/e) in the MS were selected for H$_2$ (2), He (4), CH$_4$ (15), H$_2$O (18), CO (28), O$_2$ (32), and CO$_2$ (44). The MS response for H$_2$ and CO was obtained by subtracting the CH$_4$ fragment intensity from the total MS response at m/e = 2, and subtracting the CO$_2$ fragment intensity from the total MS response at m/e = 28, respectively.

3. Results and Discussion

3.1 Cell Performance in H$_2$ and CH$_4$

The voltage and power density versus current density (V-I-P plot) characteristics curves of the four modified anodes in H$_2$ at 800 °C are shown in Figure 2. The copper metal modified NiO/YSZ anode showed the best performance with a current density of 240 mA cm$^{-2}$ and a power density of 50 mW cm$^{-2}$. The cell performance improved as the time of coating was increased from 20 to 60 min. The alkaline earth metal oxide modified anode also showed an improved cell performance, except the 5 wt.% MgO-Ni/YSZ one. For inorganic element modified anodes, the one impregnated with boric acid gave a higher current and power density, while an inferior performance was observed for the B-modified anode. Besides, the anode containing LST performed the worst among all in H$_2$.

![V-I-P plots of a SOFC with different modified anodes in H$_2$ at 800 °C.](image)

Figure 2 V-I-P plots of a SOFC with different modified anodes in H$_2$ at 800 °C.

Figure 3 presents power density versus time (P-t curve) characteristics of different anodes when the fuel was switched from H$_2$ to CH$_4$ in a potentiostatic mode. The results showed that the cells modified with 5 wt.% MgO, LST, and B were relatively resistant to degradation in CH$_4$. However, the one with Cu, CaO/Ca(OH)$_2$, and H$_3$BO$_3$ showed no improvement in comparison with the pristine NiO/YSZ. In particular, the power density of the 5 wt.% MgO and LST modified cells was found to increase drastically upon exposure to CH$_4$. The enhancement in the cell performance probably resulted from the rapid reaction kinetics on the electrode in CH$_4$ than that in H$_2$. The activity of the anode and the cell performance were correlated and further studied with in situ IR spectroscopy.
Figure 3 Power density vs. time (P-t curve) curves of the modified anodes in a potentiostatic mode (at constant potential = 0.7 V unless specified), gas environment: 0-30 min in H₂/He and 30-120 min in CH₄/He.

3.2 In-Situ IR and Mass Spectroscopy

Figure 4 depicts the DRIFTs spectra when the modified anodes were reduced in H₂ at 700 °C. The results showed that the reducibility of the anode followed an order of pristine NiO/YSZ > boron > LST > 5wt.% MgO modified NiO/YSZ (from high to low). The extent of reduction was indicated by the intensity of the Ni-O bond assigned with a band at 667 cm⁻¹, which shifted down with a more reduced anode material as monitored during the reduction. The reducibility and the cell performance in H₂ were found to be well correlated (Figure 2 and 4), indicating that high electrical performance was obtained for a more metallic or reduced anode. Figure 5 further illustrates the correlation with the MgO modified cell, demonstrating that the power density of the rarely reduced anode was low in H₂, while the performance was enhanced when it was exposed to CH₄. The reduction on the anode was manifested in CH₄ due to the depletion of the Ni-O bond as the corresponding Ni-O band shifted continuously. However, it again shifted upward after introducing O₂ flow over the CH₄ exposed anode. Reversible band shifting confirmed the position of the Ni-O band as the redox reaction of Ni-NiO occurred.
Figure 4 DRIFTs spectra of modified anodes on reduction in H\textsubscript{2} at 700 °C.

Figure 5 a) P-t curve of 5 wt.% MgO-NiO/YSZ anode and b) DRIFTs spectra of the anode subjected to various gas atmospheres, including H\textsubscript{2}, CH\textsubscript{4}, and O\textsubscript{2} at 700 °C.

The MS profiles of the eluting gaseous species produced on the modified anode upon exposure to CH\textsubscript{4} are shown in Figure 6. The results showed that the syngas, i.e., H\textsubscript{2} and CO were the major gaseous products formed on all the anode materials. The H\textsubscript{2} production followed a consistently increasing trend with the electrochemical performance in CH\textsubscript{4} (Figure 3 vs. Figure 6). Among all, the 5 wt.% MgO modified anode gave the best cell performance along with stable syngas production. The molar ratio of H\textsubscript{2} to CO in the syngas was about 2 to 1, implying that the primary reaction engaged partial oxidation of methane. Besides, the rapid decrease in the concentration of CO\textsubscript{2} and H\textsubscript{2}O suggested the involvement of a secondary reaction, consuming these by-products.
Figure 6 MS profiles of gaseous products formed on the pristine and modified NiO/YSZ anodes exposed to CH₄ at 700 °C.

The above experimental results demonstrated that the cell performance of the modified anodes was strongly influenced by the type of additives and the resultant properties. The V-I-P results showed that electrical conductivity played a major role in the cell performance in H₂. The cell with the Cu-modified anode exhibited the highest power density, followed by MgO, CaO, B, and LST modified anodes. Compared with metallic Cu, basic metal oxide exhibited low electrical conductivity, thus, leading to lower cell performance. Electrical conductivity investigated for various basic oxide modified Ni/SDC cermets followed an order of CaO > SrO > MgO > BaO > La₂O₃ [20]. On the contrary, in the current study, the MgO modified cell performed slightly better than the CaO modified one. This was probably because the comparison was made for different electrolytes (SDC vs. YSZ), and an inconsistent amount of dopant was used. Additionally, alkaline earth metal oxides such as CaO were reported to dissolve easily in the lattice of fluorite structures, thus, impeding the oxide ion conductivity of the anode. Therefore, the amount of loading needs optimization for different systems due to the diverse results reported in the literature [17, 20].

Apart from Cu and basic metal oxides, the addition of B or LST to the anode showed no improvement in cell performance in H₂. Boron has been shown promoting Ni-based catalysts for steam reforming of hydrocarbons; however, doping of boron precursors might incur a problem of poisoning or material incompatibility in the electrode [32]. The YSZ electrolyte is reported to react with volatile boron species above 600 °C to form YBO₃, disintegrating the electrode structure. This explains the reason behind using the wet method to introduce boric acid rather than solid-state reaction adopted for other compounds studied in this work. In fact, trials with the solid-state reaction using boric acid have been failed. Besides, the addition of LST also suffered from the material mismatch with the NiO/YSZ anode, although it was not very drastic. Moreover, the LST possesses low oxide ion conductivity and poor catalytic activity toward H₂ oxidation due to large polarization resistance [33, 34].

Despite the poor activity of 5 wt.% MgO, B, and LST modified anodes in H₂, these modified cells performed better than their counterparts in CH₄. The improved performance of the 5 wt.% MgO modified anode was attributed to increased electrochemical activity and coking tolerance of the Ni metal. Since CH₄ exhibits higher reducing strength than H₂, the anode (MgO-NiO) could be partially transformed to a metallic Ni phase (or NiₓMg₁₋ₓO) [35, 36]. The Ni phase would then form a
percolating layer on the YSZ electrolyte through the anode matrix and may improve the electrode conductivity. Besides, the Ni metal provides the active sites to dissociate CH₄, thus, facilitating subsequent oxidation reaction. The presence of MgO altered the phase of Ni from metallic to cationic and made the anode reluctant to reduction as observed in the DRIFTs experiment, thereby alleviating deactivation of the Ni active site by coking. It was also noticed that a lot of carbon deposit was found around the MgO modified anode and near current collectors after the test in comparison with the CaO modified one. The inclusion of CaO, a basic metal oxide, in the Ni-cermet anode has been demonstrated to reduce the deactivation in CH₄ too. However, no such improvement was observed in the current study [20]. The likely reason was explained based on different preparation methods. The anode made by the wet impregnation might give a homogeneous distribution of precursors than that by the solid-state reaction. Furthermore, the amount of MgO in the anode was critical for cell performance since the addition of 5 wt.% MgO sustained the Ni cermet anode in CH₄ while the one with 1 wt.% MgO was not effective. Nevertheless, high MgO content would give a poor performance in H₂ due to a reduced electrical conductivity [19, 36]. In contrast to the electronic promotion of MgO and boron, LST seemed to act as a structure promotor, and their additions greatly changed the morphologies of the Ni cermet anode, as shown in Figure 7. In the case of NiO/YSZ in combination with LST, a barrier layer was formed to cover the proximity of Ni cermet, thus, retarding the diffusion of CH₄ and the succeeding activation. Slowing down the bond scission of methane would promote the oxidation of deposited carbon. Also, the effect of promotion was not evident in the composite anode of boron and Ni-cermet, possibly due to the material mismatch between boron species and YSZ electrolyte.

Figure 7 SEM images of the surface of pristine and modified Ni cermet anodes before/after cell test in CH₄.
4. Conclusions

In summary, we modified the surface of a Ni-cermet anode with copper, alkaline earth metal oxide, boron, and perovskite and investigated the cell performance in SOFCs. The results indicated that the copper modified anode gave the best performance in H₂, while the one incorporated with inorganic compounds such as MgO, LST, and boron behaved relatively stable in CH₄. The cell performance of the modified anodes was greatly affected by the interaction between the dopant and the Ni cermet, as well as the material compatibility with the substrate. The promotion of Ni-cermet anodes by MgO worked mainly via electronic interaction with Ni, while in LST and boron, it did through structure control. Meanwhile, the modification with CaO was not effective, probably due to the non-uniform distribution of precursors resulted from the solid-state reaction. Our studies provided valuable information to compare the effect of different dopants on the Ni anode and to obtain the underlying mechanism for coking resistance by an in situ surface analysis. This will allow designing an effective material capable of withstanding methane and other hydrocarbons for fuel cells and other potential energy-related applications.

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Author Contributions

Andrew C. Chien – Project management, experimental design, manuscript writing; Eric Y. Lin – Cell fabrication, electrical measurement, surface analysis; Nicole J. Ye – IR and MS analysis

Competing Interests

The authors have declared that no competing interests exist.

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