Research article

Enhancement of hydrogen production using Ni catalysts supported by Gd-doped ceria

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ABSTRACT

A redox cycle between Ce⁴⁺ and Ce³⁺ is an elementary step in water gas shift (WGS) mechanism. By facilitating the redox cycle between +4 and +3 of cerium, a formation of oxygen vacancy can be enhanced. It is considered to be a dominating factor in developing the WGS performance and the stability of ceria in this work. We have facilitated the redox cycle in CeO₂ to enrich the WGS activity. The WGS reaction was carried out on Ni catalyst supported by Gd-doped ceria (GDC) from Daiichi. Ni and Re were added onto GDC by impregnation method to examine the role of Re addition on surface, structural and reducibility, which affected upon their catalytic activities. Rhenium has an influence on increasing the water gas shift performance of Ni/GDC catalysts because it facilitates the redox process at the surface of ceria, disperses Ni particles and enhances oxygen vacancy formation. The results indicate that the water gas shift activity of 1%Re₄%Ni/GDC is higher than that of 5%Ni/GDC. The dispersion of active site on the surface of catalyst results in an increase of CO molecule adsorption and acceleration of the redox cycle between Ce⁴⁺ and Ce³⁺ of ceria support via oxygen vacancy generation. Therefore, using a combination of these two effects can enhance the WGS performance.

1. Introduction

Many chemical industries and the fuel cell industry utilize hydrogen to provide a pollution-free combustion and high energy density [1, 2, 3]. Hydrogen is a promising fuel for mobile and residential fuel cells. It is a sustainable fuel resource and emission free during the hydrogen reaction. Hence, hydrogen has received a widespread attention to reduce our dependence on fossil fuel, which has created environmental pollution. Currently, a production of hydrogen by natural gas steam reforming is still the predominant way in major industries [4]. Reforming process generates a synthesis gas, which contains H₂ and CO. However, CO must be eliminated because it deactivates Pt electrode in a low-temperature fuel cell. Therefore, a water gas shift (WGS) reaction has been applied to obtain hydrogen rich gas reformates.

Ceria based materials are widely used for many applications because of its high oxygen storage capacity and good reducibility [5, 6, 7, 8]. Excellent redox property of ceria provides an increase in the oxygen transport ability at ceria surface. In addition, a strong interaction between ceria support with active metal particles results in enhancing the metal dispersion and preventing the ceria support agglomeration. Oxygen vacancies within the catalyst can be introduced from the redox cycle between Ce⁴⁺ and Ce³⁺ of ceria [9, 10, 11, 12]. This provides sites for hydrogen dissociation, which is the rate-limiting step for water gas shift reaction. Therefore, increasing oxygen vacancy is a key factor in enhancing the water gas shift activity of CeO₂ based catalysts. Many researchers have tried to increase the oxygen vacancies in ceria-based materials to enrich the WGS activity. An addition of Zr to ceria was found to enhance the surface area, redox property and amount of oxygen vacancies [13, 14]. Some previous study revealed that Nd and La showed positive impact on Cu dispersion, redox properties and oxygen vacancy formation in Cu/CeO₂ catalysts [15]. Furthermore, it was found that an addition of Ba could enhance the sintering resistance, metal dispersion and the oxygen vacancy of ceria materials [16].

In this work, we have focused on accelerating the redox cycle between Ce⁴⁺ and Ce³⁺ of ceria for the bimetallic ReNi catalyst on Gd-doped ceria support to enhance the oxygen vacancies. The oxygen vacancy is the main factor to facilitate the redox cycle. In addition, an increase of the active metal of supported Ni catalysts alerts CO adsorption on the catalyst.

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surface, which leads to increasing rate of WGS reaction. Effects of rhenium addition on CeO$_2$ structure and reducibility were investigated using BET surface area, X-ray diffraction, H$_2$-temperature programmed reduction, chemisorption and X-ray photoelectron spectroscopy analyses to explain the relationship between the predominant factors (i.e. metal dispersion, metal area, and reducibility) and water gas shift activity.

2. Experimental procedure

2.1. Catalysts preparation

Impregnation method was used to prepare Ni/GDC catalyst. Nickel (II) nitrate hexahydrate (Alfa Aesar) was dissolved in a minimal amount of deionized (DI) water. The metal nitrate solution was added to 10% gadolinium doped ceria support (supplied by Daiichi).

NiRe/GDC catalyst was prepared using co-impregnation method. Ni(NO$_3$)$_2$.6H$_2$O (Alfa Aesar) and NH$_4$ReO$_4$ (Sigma-Aldrich) were dissolved with DI water. The mixture solution of metal nitrate was added to GDC support (Daiichi). All samples were dried at 110 °C for 12 h and calcined at 650 °C for 8 h.

2.2. Catalyst characterization

The BET surface area of the specimens was performed using Quantachrome NOVA 1200e. Prior to the measurement, the catalysts were heated under vacuum at 300 °C for 6 h. The BET surface areas of all catalysts were estimated by using N$_2$ adsorption-desorption isotherms at 77 K in the relative pressure range of 0.05–0.3.

The X'Pert Pro diffractometer (PANalytical) operating at 0.02° step and 0.5 s per step over a 20 range of 20°–80° with the current of 40 mA and 40 kV was employed for X-ray powder diffraction measurements. The X-ray diffractograms were collected using nickel-filtered Cu K$_\alpha$ radiation. The full width at half maximum of the strongest (111) reflection was considered for the calculation of ceria crystallite size using Scherrer’s equation.

A catalyst analyzer BELCAT-B was used for H$_2$-Temperature Programmed Reduction (H$_2$-TPR) experiment. The samples were treated under high purity He at 120 °C for 30 min prior to a TPR measurement. A mixed gas of 5%H$_2$–Ar was employed for TPR experiments operating from 50 °C to 1000 °C at the rate of 10 °C min$^{-1}$. The H$_2$ consumption during the reduction was determined using a thermal conductivity detector (TCD).

A mixed gas of 10%CO–He was introduced to the specimen for the measurement of the metal surface area from the total gas chemisorption. The metal surface area and the weight of the specimen were used for the calculation of metal dispersion. All specimens were reduced with H$_2$ at 400 °C for an hour. After cooling with He, carbon monoxide chemisorption pulse was analysed under the flow of carbon monoxide in helium carrier gas at 50 °C at the rate of 30 ml min$^{-1}$. The flow of carbon monoxide out of the reactor was monitored using a TCD.

Amount of adsorption per 1 g of sample $V_m$ in cm$^3$ g$^{-1}$

$$V_m = \frac{V_{chem}}{m}$$

(1)

Metal dispersion (percentage of metal surface exposure) D (%)

$$D = \frac{V_{chem}}{26519 \times SF \times MW \times 100/c}$$

(2)

where $V_m$ is an amount of adsorption in cm$^3$, $m$ is the weight of specimen in g, MW is the atomic weight of metal in g mol$^{-1}$, and SF is the stoichiometry factor.

Metal weight (c) in g:

$$c = m \times p/100$$

(3)

$Am$ is metal surface area (surface area of metal per 1 g of sample) in m$^2$ g$^{-1}$

$$Am = \frac{V_{chem}}{26519 \times SF \times 6.02 \times 10^{23} \times 10^{-18}/m}$$

(4)

where $p$ is the weight percentage of supported metal content in wt%, and $\sigma_m$ is the crosssectioned area of a metal in atom/nm$^2$.

The Raman spectra of Ni and NiRe catalysts were performed using Perkin Elmer System 2000 FT-IR/FT-Raman. The spectra were collected using Ar ion laser irradiation in the range of 200–2000 cm$^{-1}$ with an output power of 10 mW and wavelength of 532 nm.

The chemical composition of specimen surface was investigated using an X-ray photoelectron spectrometer (XPS, AXIS ULTRA DLD, Kratos analytical). The catalysts were reduced ex-situ under a flow of 5%H$_2$/N$_2$ at 400 °C for 1 h before the analysis. The base pressure in the XPS analysis chamber was about 5 × 10$^{-9}$ torr. The specimens were excited with X-ray hybrid mode of 700 × 300 μm spot area with a monochromatic Al K$_\alpha$$_2$ radiation at 1.4 keV. X-ray anode runs at 15kV 10mA 150 W. The photoelectrons are detected with a hemispherical analyzer positioned at an angle of 90° with respect to the normal to the sample surface. The spectra are calibrated using the C1s line (BE = 285 eV).

2.3. Water gas shift activity

Water gas shift performance was tested using a fixed bed reactor with 4 mm inner diameter. The catalysts (~150 mg) were loaded into a stainless steel tube reactor between two layers of quartz wool. The reactor was placed inside a tube furnace. Prior to the water gas shift activity measurement, the prepared catalyst was reduced by heating in 5% H$_2$/N$_2$ balanced gas from room temperature to 400 °C and held for 1 h. After reducing the catalyst, the temperature was lowered to 100 °C. Activity test of Ni catalysts was performed in the temperature range of 100–500 °C. H$_2$O was fed through a pre-heater using a syringe pump. The feed gas contained 5% CO, 10% H$_2$O and 85% N$_2$. Preliminary measurements were operated to consider proper conditions from which internal and external mass transfer effects were not dominant. When considering the effect of external mass transfer, the particle size diameter of the catalysts was between 100-200 μm in all conditions. Furthermore, the total flow rate was kept constant at 100 ml min$^{-1}$ in all experiments. The outlet gasses were analysed using an on-line Shimadzu GC-14B gas chromatography equipped with TCD. The column utilized in the gas chromatography was a Unibead C. The WGS activities were calculated using Eq. (5):

$$\% CO conversion = \frac{CO_{in} - CO_{out}}{CO_{in}} \times 100$$

(5)

The WGS rates were tested in separate experiments using 20 mg of catalysts and the feed gas mixture of 5% CO, 10% H$_2$O and 85% N$_2$ where the CO conversion was less than 30%.

Rate = (F × X) / m

(6)

where Rate is the WGS rate (mol g$^{-1}$ s$^{-1}$), F is the molar flow rate of CO (mol s$^{-1}$), m is the weight of the samples (g), and X is the carbon monoxide conversion.

3. Results and discussion

3.1. Standard characterization of Ni catalysts

Figure 1 presents X-ray diffraction patterns of Ni catalysts. XRD spectra of all catalysts correspond to ceria in fluorite-type cubic crystal structure. The lattice constant of ceria in the catalysts was calculated from 111 diffraction peak broadening. The ceria lattice parameters of GDC (Daiichi) and Ni catalysts were found in the range of 0.5421–0.5424 nm. Nickel and rhenium ion could not be inserted into the lattice of CeO$_2$ due to the nature of impregnation synthesis. There are two diffraction peaks at 20 = 37.3° and 43.4° which correspond to NiO in supported Ni catalysts. The NiO diffraction peaks are very weak suggesting that there is
very small amount of NiO. The results of BET surface areas and crystallite size are summarized in Table 1. An addition of nickel and rhenium onto GDC (Daiichi) by impregnation method leads to an increase in crystallite size and a reduction of the surface area. This result is probably due to CeO2 crystallites agglomeration after calcination at high temperature. CO chemisorption analysis was operated to study a dispersion of nickel over Ni/GDC materials. The reduced ceria gives some signals to the CO titration. Indeed, Ce3+ species can also react with CO during the investigation. Thus, the CO-chemisorption was measured on doped ceria without an addition of nickel, used as a blank test. From the results of chemisorption analysis shown in Table 1, it is found that an addition of rhenium to Ni/GDC (Daiichi) enhances a dispersion of Ni metal on the GDC support. Metallic Ni was believed to provide active sites for the WGS reaction [17]. The effect of Re addition on metallic Ni surface area was probed, which revealed that ReNi/GDC catalyst had the highest metallic surface area of 42 m2/g. Generally, catalysts with higher metallic surface area provide an increase in catalytic activity, as more of their surface active sites are exposed to reactants [18, 19].

3.2. H2-TPR of Ni catalysts

TPR profiles of Ni catalysts are shown in Figure 2. The reduction peaks of GDC support appear at 510 °C and 800 °C, which are assigned to the reduction of the support and the bulk CeO2, respectively. TPR profile of 1%Re/GDC is also plotted for a comparison. The reduction of Rhenium oxide occurs at high temperature (about 400 °C), which indicates that rhenium species alone is difficult to reduce. On the contrary, a combination of Ni and Re species enhances their reducibility. For Ni/GDC, there are three reduction peaks corresponding to the reduction of nickel oxide, surface region of CeO2 and the bulk. An addition of Ni onto Gd-doped ceria from Daiichi shifts the surface reduction toward lower temperatures. H2-TPR profile of bimetallic NiRe on Gd-doped ceria from Daiichi is also compared. It is clearly seen that Re addition onto Ni/GDC drastically shifts the reduction peak of surface oxygen to lower temperature. From the comparison between Ni/GDC and ReNi/GDC catalysts, it was found that an addition of Re increased H2 consumption (Table 2), indicating an improvement of reducibility of Ni catalyst. These results imply that rhenium helps Ni in reducing ceria and leads to the enrichment of Ce3+ at the ceria surface, which gives rise to oxygen vacancies. This simplifies the electron movement at the surface, hence, enhancing easier surface reduction [11].

3.3. Raman spectroscopy

Defect structures of supported Ni catalysts were investigated using Raman spectroscopy. Figure 3 illustrates the Raman spectra of NiRe/GDC, Ni/GDC and GDC support. A Raman peak at ~460 cm⁻¹ is the symmetrical stretching of the oxygen atoms around Ce ions (F2g mode) of the cubic fluorite crystal structure of CeO2. The intensity of this peak decreases when addition of Ni or NiRe onto GDC support. In addition, there is another broad peak of NiRe/GDC and Ni/GDC starting from 520 to 650 cm⁻¹. It is a defect-induced mode band, which is due to the presence of surface

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**Table 1.** Surface areas, crystallite size and %dispersion of Ni catalysts.

| Catalyst       | S_BET (m²/g) | Crystallite size (nm) | Lattice parameter (nm) | %dispersion | S_0 (m²/m³O) | Average particle size (nm) |
|----------------|--------------|-----------------------|------------------------|-------------|--------------|---------------------------|
| GDC            | 76           | 12.0                  | 0.5424                 | 1.8         | -            | -                         |
| 5%Ni/GDC      | 45           | 15.1                  | 0.5423                 | 4.9         | 32.5         | 20.7                      |
| 1%Re5%Ni/GDC  | 42           | 16.1                  | 0.5421                 | 6.5         | 42.0         | 14.9                      |

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* Estimated from N2 adsorption at ~196 °C.
* Calculated from 111 diffraction peak broadening.
* Estimated from CO-chemisorption.

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**Table 2.** WGS rate and H2 consumption of Ni-based catalysts.

| Catalyst       | Rate at 350 °C (mmol/kg.s) | E_a (kJ/mol) | H2-TPR Peak position (°C) | H2 consumption (mmol/g) |
|----------------|----------------------------|--------------|---------------------------|-------------------------|
| 5%Ni/GDC      | 58                         | 69 ± 0.6     | 240, 321                  | 1.26, 16.41             |
| 1%Re5%Ni/GDC  | 77                         | 58 ± 0.4     | 205, 244, 280             | 0.66, 0.74, 16.47       |

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Figure 1. XRD patterns of supported Ni catalysts, (a) GDC, (b) 5%Ni/GDC and (c) 1%Re5%Ni/GDC.

Figure 2. H2-TPR profiles of supported Ni catalysts, (a) GDC, (b) 1%Re/GDC, (c) 5%Ni/GDC and (d) 1%Re5%Ni/GDC.

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defects or a different charge state of doping cations [20]. These defects are related to the evolution of oxygen vacancies.

3.4. XPS analysis

The relationship between chemical species on the surface region of the catalysts and reducibility were studied using XPS analysis. The results are summarized in Table 3. The Ni 2p3/2 XPS spectra of Ni/GDC and ReNi/GDC catalysts were deconvoluted as shown in Figure 4. The prepared catalyst was reduced under a flow of 5%H2/N2 at 400 °C for 1 h before performing the XPS analysis. After the reduction, Ni was in the metallic state whereas the different nickel species co-existed due to the interaction with GDC. The peak around 853 eV is attributed to Ni0 whereas the following two peaks at higher binding energy are assigned to surface Ni2+ [21]. Their relative ratios are summarized in Table 3. Ni0 was suggested to be the dominant active species in accelerating the reactants [22]. For Re addition onto Ni/GDC, it was found that XPS spectrum of surface Ni2+ shift to lower Ni binding energy values (856.35), which indicates that Re appears to hinder the strong interaction between Ni and GDC. Thus, metallic nickel formation is favored [23]. The enhancement of Ni0 content of ReNi/GDC implies a superior WGS activity of Ni catalyst with Re addition.

3.5. WGS activity of Ni catalysts

Figure 5 shows the WGS activity of Ni catalysts. The WGS activity of GDC (Daiichi) support was also plotted for a comparison. It was found that 5%Ni/GDC (Daiichi) is an active catalyst for a water gas shift reaction. The activity of 5%Ni/GDC (Daiichi) starts at about 200 °C and ascends to reach the maximum of 95% conversion at 350 °C. The conversion above 350 °C reaches equilibrium and then slightly decreases. Moreover, it can be seen that ReNi/GDC exhibits higher CO conversion rate compared to 5%Ni/GDC catalysts. The bimetallic catalyst ReNi/GDC becomes active above 200 °C and reaches the maximum of 99% conversion rapidly at around 350 °C. The combination of H2-TPR and CO chemisorption results suggests that ReNi/GDC has an excellent redox property, the smallest particle size and the highest nickel dispersion on GDC support, which contributes to the highest water gas shift activity [24, 25].

Table 3. XPS analysis for quantitative measurement of Ni-based catalysts.

| Catalyst       | BE Ni 2p3/2 (eV) | Ni0 (%) |
|----------------|------------------|---------|
| 5%Ni/GDC       | 853.07           | 28.2    |
| 854.92         | 856.62           |         |
| 1%Re5%Ni/GDC   | 853.11           | 34.1    |
| 5%Ni/GDC (Daiichi) | 854.91         |         |
| 856.35         |                  |         |

Figure 6. WGS catalytic activity of supported Ni catalysts.

4. Conclusions

The role of Re on the water gas shift activity was investigated. The best activities were found for ReNi on GDC from Daiichi. The catalytic activity of ReNi/GDC (Daiichi) rapidly increased to attain the maximum of 99% conversion at 350 °C. An addition of Re into Ni/GDC provided a high metal activity. Further studies are required to understand the detailed mechanism of this catalytic activity.
The authors report that ReNi/GDC is a promising catalyst for a hydrogen production application by water gas shift reaction. The high WGS activity of ReNi/GDC was mainly related to enhanced reducibility and the active Ni\(^0\) species. An increase of the surface area of metallic Ni improved the reactant CO adsorption and enhanced the water gas shift rate. It is concluded that ReNi/GDC is a promising catalyst for a hydrogen production application by water gas shift reaction.

Declarations

**Author contribution statement**

Opas Tojira: Analyzed and interpreted the data; Wrote the paper.
Jessica Gina Lomonaco: Performed the experiments.
Thanathon Sesuk: Performed the experiments; Contributed reagents, materials, analysis tools or data.
Soomitra Chojarjouchul: Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data; Wrote the paper.
Pannipa Tepamatr: Conceived and designed the experiments; Performed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data; Wrote the paper.

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**Data availability statement**

Data will be made available on request.

**Declaration of interests statement**

The authors declare no conflict of interest.

**Additional information**

No additional information is available for this paper.

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