Reverse Water–Gas Shift Chemical Looping Using a Core–Shell Structured Perovskite Oxygen Carrier

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Abstract: Reverse water–gas shift chemical looping (RWGS-CL) offers a promising means of converting the greenhouse gas of CO 2 to CO because of its relatively low operating temperatures and high CO selectivity without any side product. This paper introduces a core–shell structured oxygen carrier for RWGS-CL. The prepared oxygen carrier consists of a metal oxide core and perovskite shell, which was confirmed by inductively coupled plasma mass spectroscopy (ICP-MS), XPS, and high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) measurements. The perovskite-structured shell of the prepared oxygen carrier facilitates the formation and consumption of oxygen defects in the metal oxide core during H 2 -CO 2 redox looping cycles. As a result, amounts of CO produced per unit weight of the core–shell structured oxygen carriers were higher than that of a simple perovskite oxygen carrier. Of the metal oxide cores tested, CeO 2, NiO, Co 3 O 4, and Co 3 O 4-NiO, La 0.75 Sr 0.25 FeO 3-encapsulated Co 3 O 4-NiO was found to be the most promising oxygen carrier for RWGS-CL, because it was most productive in terms of CO production and exhibited long-term stability.

Keywords: reverse water–gas shift chemical looping (RWGS-CL); oxygen carrier; metal oxide; perovskite; core–shell

1. Introduction

Due to the increased use of fossil fuels, atmospheric CO 2 concentrations have steadily increased, which is the major cause of global warming [1–3]. According to a report issued in 2015 by the PBL Netherlands Environmental Assessment Agency, global CO 2 emission from fossil fuel combustion was about 35.7 Gt, which at the time represented a 1.4% annual increase [4]. To replace fossil-based energy resources, various types of renewable energies have been researched for the past few decades [5–7]. However, none of these energy sources are economically feasible as compared with low cost fossil fuels, including shale gas [8]. Accordingly, the use of fossil fuels in combination with an efficient CO 2 capture system offers a feasible near-term solution until low cost renewable energy resources become available. Carbon capture and storage (CCS) systems have been devised to capture and store CO 2 generated by fossil fuel combustion [9,10], but these technologies have CO 2 storage capacity limitations [11] and the leakage of CO 2 from storage systems appears to make permanent storage unattainable [12].

CO 2 reduction to CO has been raised as an attractive alternative to CCS [13–17]. Instead of burying supercritical CO 2 in depleted oil or gas reservoirs, CO 2 could be converted into CO, which is a valuable feedstock for the production of chemical products such as methanol, olefins, and liquid...
fuels [18,19]. More specifically, CO produced from CO$_2$ could be used as syngas, a raw material for gas to liquid processes. Photocatalytic process and solar thermochemical splitting have been considered as potential means of reducing CO$_2$ to CO [20]. However, the efficiency of photocatalytic CO$_2$ conversion is still too low to be commercialized into the industrial scale due to the relatively poor activity and stability of photocatalysts [21]. While the solar thermochemical approach is much more efficient at CO$_2$ conversion than photocatalytic processes, it requires high operating temperatures (>1000 °C), implying an energy-intensive process. Additionally, researchers have found it difficult to identify cost-effective materials that are stable at temperatures above 1000 °C [22]. The hydrogenation of CO$_2$ to produce CO, that is, the reverse water–gas shift (RWGS) reaction (Equation (1)), was devised as an alternative to prepare future society for a time when cost-effective, solar energy-based CO$_2$ reducing processes become available. Using H$_2$, CO$_2$ can be converted to CO at lower temperatures than those required by solar thermochemical processes, and thus energy consumption to produce a unit mole of CO can be decreased [23,24]. In addition, when CO is produced at low temperatures, CO production can be easily sequenced with Fischer–Tropsch synthesis (FTS)-based fuel production [25].

$$\text{CO}_2 + \text{H}_2 \leftrightarrow \text{CO} + \text{H}_2\text{O} \quad (1)$$

Reverse water–gas shift chemical looping (RWGS-CL) is a two-step CO$_2$ hydrogenation process where H$_2$ oxidation and CO$_2$ reduction occur in two separate reactors. By using H$_2$ for CO$_2$ reduction, the operating temperature (500–700 °C) of RWGS-CL becomes lower than those required at solar thermochemical processes [22,26]. Furthermore, it can generate CO without any byproducts (e.g., free of CH$_4$ produced by CO$_2$ methanation), since H$_2$ is not directly contacted with CO$_2$. As a result, syngas can be produced at an energy efficiency 54% greater than that of normal RWGS processing [27]. RWGS-CL is a two-step reduction/oxidation process. During the reduction phase, the oxygen carrier is reduced by injecting hydrogen to form oxygen defects (Equation (2)), and because hydrogen, a strong reducing agent, is used, processing can be carried out at lower temperatures than those required by solar thermochemical processes involving the reduction of a metal oxide. During the oxidation phase, oxygen defects produced during the reduction phase act as active sites for CO$_2$ reduction to produce CO (Equation (3)). A schematic diagram of the RWGS-CL process is provided in Figure 1.

Reduction step: \[ \text{MeO}_x + y\text{H}_2 \rightarrow \text{MeO}_{x-y} + y\text{H}_2\text{O} \quad (2) \]

Oxidation step: \[ \text{MeO}_{x-y} + y\text{CO}_2 \rightarrow \text{MeO}_x + y\text{CO} \quad (3) \]

![Figure 1](image_url)  

**Figure 1.** Schematic of the reverse water–gas shift chemical looping (RWGS-CL) process.

Reducible metal oxides and perovskites have been widely used as oxygen carriers for RWGS-CL. Metal oxides are attractive oxygen carriers for commercial-scale processes due to their accessibilities and
high oxygen storage capacities. However, because of their high tendencies to be sintered at relatively low temperature, the deactivation of the particle easily occurs and the performance of the oxygen carrier cannot be maintained in the long-term experiment [28,29]. ABO$_{3}$-structured perovskite [30,31] presents both high thermal stability and selectivity for converting CO$_{2}$ into CO. However, the oxygen storage capacities of ABO$_{3}$ perovskites are generally lower than those of metal oxides, and the maximum amount of CO production per unit weight of ABO$_{3}$ perovskites is also lower [32].

Core–shell structured particles with a metal oxide core and a perovskite shell might provide a solution as they offer the advantages of metal oxides and perovskite materials. Since perovskite has high lattice oxygen and electron conductivity, a large amount of lattice oxygen can be transferred through the perovskite shell to the metal oxide core. In addition, the sintering of metal oxide particles can be effectively avoided by encapsulating metal oxide with the thermostable perovskite shell [33]. For these reasons, core–shell structured oxygen carriers have been used for chemical looping processes to optimize redox activity, oxygen storage capacity, and stability [34,35]. However, such core–shell oxygen carriers have rarely been used for RWGS-CL.

In the present study, we synthesized metal oxide core–perovskite shell particles and investigated their potentials as oxygen carriers for RWGS-CL. In detail, La$_{0.75}$Sr$_{0.25}$FeO$_{3}$ (LSF; a perovskite) and MeO$_{x}$@LSF (MeO$_{x}$: CeO$_{2}$, NiO, Co$_{3}$O$_{4}$, and Co$_{3}$O$_{4}$-NiO) particles were synthesized and tested. Under repeated redox cycles, core–shell particles have demonstrated their higher CO productivities than a simple perovskite particle. The core–shell structure of these particles was investigated by inductively coupled plasma mass spectroscopy (ICP-MS), X-ray photoelectron spectroscopy (XPS), and high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM).

Temperature-programmed processes were used to quantify oxygen mobility in particles during the redox cycle.

2. Experimental Section

2.1. Oxygen Carrier Preparation

A modified Pechini method was used to prepare core–shell structured oxygen carriers [35]. In detail, stoichiometric amounts of CeO$_{2}$, Co$_{3}$O$_{4}$, and NiO nanoparticles (<50 nm, Sigma-Aldrich, St. Louis, MO, USA) were dispersed in 60 vol% aqueous ethanol and sonicated for 5 min. After settling the dispersion for 6 h, the top phase was separated to obtain nanoparticle suspension. To coat the metal oxide nanoparticles with perovskite, stoichiometric amounts of La(NO$_{3}$)$_{3}$·6H$_{2}$O (99.9%, Alfa Aesar, Haverhill, MA, USA), SrCl$_{2}$·6H$_{2}$O (99%, Sigma-Aldrich), and Fe(NO$_{3}$)$_{3}$·9H$_{2}$O (≥98%, Sigma-Aldrich) were dissolved in deionized water. After heating the solution to 50 °C and stirring for 30 min, citric acid (CA, ≥99.5%, Sigma-Aldrich) was added by 3 times of the total moles of metal cations and stirred for another 30 min. The prepared nanoparticle suspension and perovskite solution were then mixed (molar ratio of metal oxide to perovskite = 1:1) and stirred at 50 °C for 30 min. Ethylene glycol (EG, 99.5%, Samchun Pure Chemical Co., Seoul, Korea) was then added (molar ratio of EG:CA = 2:1) and stirred at 80 °C until a gel formed. This gel was then dried in a 130 °C oven without stirring. To remove volatile species, the dried sample was loaded into a furnace and heated to 450 °C and maintained for 4 h with 80 mL/min air, and then annealed by raising the temperature to 900 °C (800 °C for Co$_{3}$O$_{4}$-containing samples due to the decomposition of Co$_{3}$O$_{4}$ to CoO at 900 °C) and maintaining this temperature for 6 h with the same air flow. The procedure used to prepare a simple perovskite oxygen carrier was as described above, except that the addition of metal oxide nanoparticles was omitted.

2.2. Characterization of Oxygen Carriers

High-resolution powder X-ray diffraction (XRD) measurements were carried out to identify the crystal phases of the prepared oxygen carriers. XRD spectra were recorded on a Rigaku SmartLab X-ray diffractometer (KAIST Analysis Center for Research Advancement) with Cu Kα radiation (λ = 1.5406 Å)
at 45 kV and 200 mA. Samples were scanned over the 2θ range of 20–80° at a scanning rate of 10°/min, and the signals obtained were processed using the PDXL2 program. The metal compositions of the synthesized oxygen carriers were measured by inductively coupled plasma mass spectroscopy (ICP-MS, Agilent ICP-MS 7700S, KAIST Analysis Center for Research Advancement). Surface metal atomic ratios were obtained by X-ray photoelectron spectroscopy (XPS) with a Thermo VG scientific Sigma Probe XPS system (KAIST Analysis Center for Research Advancement) and monochromatic Al Kα radiation at 5.0 µA and 4 kV. To confirm the core–shell structure of the synthesized oxygen carriers, high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) images were obtained using an FEI Talos F200X 200 kV TEM instrument (KAIST Analysis Center for Research Advancement).

2.3. RWGS-CL Experiments

RWGS-CL experiments were performed in a fixed-bed quartz reactor (i.d. = 7 mm). Of the fresh oxygen carrier 100 mg was placed on quartz wool and packed into the reactor, which was then placed in an electric furnace and heated to 600 °C at a ramping rate of 5 °C/min. RWGS-CL experiments were performed in four steps. (1) A reduction step, H₂ with N₂ as a carrier gas (H₂:N₂ = 1:9) was injected into the reactor at a total flow rate of 50 mL/min for 20 min to form oxygen defects in the oxygen carrier. (2) A purge step, which involved purging the reactor with N₂ at 45 mL/min for 10 min. (3) An oxidation step, CO₂ with N₂ as a carrier gas at a total flow rate of 50 mL/min (CO₂:N₂ = 1:9) was injected for 20 min into the reactor to reoxidize the reduced oxygen carrier and generate CO. (4) The purge step mentioned in (3) was then repeated. During experiments, this redox cycle was repeated 5 times, though the long-term stability test was performed for 20 cycles. Product streams were analyzed by an online gas analyzer. H₂ levels were analyzed by a thermal conductivity detector (ZAF-4, Fuji Electric Systems), and the levels of other gases were detected by an infrared detector (ZRI-6, Fuji Electric Systems).

2.4. Temperature-Programmed Processes

Temperature-programmed processes were conducted to access oxygen mobility, which is an ability to form oxygen defects when exposed to hydrogen and the consumption of these defects by CO₂ sourced oxygen during the reduction of CO₂. For H₂ temperature-programmed reduction (H₂-TPR) measurements, 100 mg of oxygen carrier was packed into the fixed-bed quartz reactor and a H₂/N₂ carrier gas (H₂ and N₂ flow rates of 10 and 40 mL/min, respectively) was injected into the reactor at room temperature. Then the reactor temperature was raised to 900 °C at a 5 °C/min in the electric furnace. Unreacted H₂ was detected through the thermal conductivity detector. For CO₂ temperature-programmed oxidation (CO₂-TPO) measurements, 100 mg of oxygen carrier was sealed in the same quartz reactor, pretreated with H₂ at 600 °C, cooled to room temperature, and a CO₂/N₂ carrier gas (CO₂ and N₂ flow rates 10 and 40 mL/min, respectively) was introduced into the reactor. The reactor was then heated to 900 °C at a ramping rate of 5 °C/min with the same furnace system, and the generated CO was detected using the infrared detector.

3. Results and Discussion

3.1. Structure of the Oxygen Carriers

The XRD patterns of synthesized La_{0.75}Sr_{0.25}FeO₃ (LSF), CeO₂@LSF, NiO@LSF, Co₃O₄@LSF, and Co₃O₄-NiO@LSF are shown in Figure 2. The crystal phase of LSF matched the PDF card of La_{0.8}Sr_{0.2}FeO₃ (PDF #: 00-035-1480), indicating the formation of a fully oxidized orthorhombic perovskite structure. Other core–shell structured samples also showed the same perovskite phase with each metal oxide phases, i.e., CeO₂ (PDF #: 01-080-8533), NiO (PDF #: 00-047-1049), Co₃O₄ (PDF #: 00-009-0418), and NiCo₂O₄ (PDF #: 01-073-1702). These observations indicated pure oxygen carriers with the intended phases had been synthesized successfully.
ICP-MS and XPS were carried out to determine the atomic compositions of metal cations in the bulk phase and surface of prepared oxygen carriers, respectively. ICP-MS and XPS data are summarized in Table 1, in which cation compositions of metal oxide and perovskite are referred to ‘cations of metal oxide’ and ‘cations of perovskite’. For example, in CeO$_2$@LSF, ‘cations of metal oxide’ only considers the composition of Ce cation while ‘cations of perovskite’ includes the remaining cations, which consists of the perovskite phase, La, Sr, and Fe. The cation compositions of metal oxide as determined by the ICP-MS matched theoretical value in the bulk phase particle, i.e., 0.33 for Ce in CeO$_2$@LSF, 0.33 for Ni in NiO@LSF, 0.6 for Co in Co$_3$O$_4$@LSF, and 0.5 for Co+Ni in Co$_3$O$_4$-NiO@LSF.

While the result of ICP-MS represented the bulk phase composition, XPS analysis showed the cation composition of the surface side. When ICP-MS and XPS data were compared, molar compositions of metal oxides determined by XPS were around half of the bulk composition determined by ICP-MS. This indicated that metal oxide phases were largely confined to cores and the perovskite structures were mainly presented on the particle surfaces, as has been previously reported for the same preparative procedure [32,35]. HAADF-STEM mapping in Figures 3 and 4 confirmed the core–shell structure of particles with a metal oxide core and perovskite shell. In the STEM mapping, metal oxide cores with 20–40 nm were clearly encapsulated by perovskite shells.
Figure 3. High-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) mapping images of the prepared oxygen carriers. CeO$_2$@LSF (a), NiO@LSF (b), Co$_3$O$_4$@LSF (c), and Co$_3$O$_4$-NiO@LSF (d). Green = La, Sr, and Fe; (Red = each core metal (Co in (d)), Yellow = Ni in (d)).

Figure 4. HAADF-STEM mapping image of La$_{0.75}$Sr$_{0.25}$FeO$_3$.

3.2. RWGS-CL Performance of the Oxygen Carriers

To observe the effect of core–shell structured particles, the amounts of CO produced by the prepared oxygen carriers during the oxidation step are shown in Figure 5. As shown in the figure, NiO@LSF, Co$_3$O$_4$@LSF, and Co$_3$O$_4$-NiO@LSF produced more CO than LSF, whereas LSF and CeO$_2$@LSF produced similar amounts (0.35–0.55 mmol/gOC), which could be due to the low oxygen storage capacity of CeO$_2$. Unlike other metal oxide cores, CeO$_2$ cannot be fully reduced at the operating condition then the amount of reducible oxygen of CeO$_2$@LSF is lower than others. On the other hand, Co$_3$O$_4$ and NiO-based oxygen carriers recorded about 3-fold higher production amounts of CO after the 2nd cycle (1.15 mmol/gOC for NiO@LSF, 1.45 mmol/gOC for Co$_3$O$_4$@LSF, and 1.6 mmol/gOC for Co$_3$O$_4$-NiO@LSF). For successful RWGS-CL cycling, particles should have a high ability to form oxygen defective active sites when exposed to hydrogen atmosphere and these oxygen-defective sites should be properly consumed by CO$_2$ sourced oxygen during the CO$_2$ oxidation step. Therefore, it can
be inferred that the NiO@LSF, Co$_3$O$_4$@LSF, and Co$_3$O$_4$-NiO@LSF have the higher oxygen mobility than simple perovskite catalysts.

3.3. Temperature-Programmed Processes

In order to get more insights, temperature-programmed processes were performed on the prepared oxygen carriers. Firstly, H$_2$-TPR measurements were obtained for LSF and other core–shell structured oxygen carriers to observe temperatures at which oxygen defects are first formed during the hydrogen reduction step. Additionally, the amount of the oxygen defect formed by H$_2$ was measured.

Amounts of H$_2$ consumed as determined by H$_2$-TPR measurements are presented in Figure 6a and Table 2. The results obtained showed that overall hydrogen consumptions by LSF and CeO$_2$@LSF were lower than those of NiO@LSF, Co$_3$O$_4$@LSF, and Co$_3$O$_4$-NiO@LSF as expected from Figure 5. Furthermore, H$_2$ was mainly consumed at a temperature higher than 600 °C for LSF and CeO$_2$@LSF. On the other hand, the other three carriers consumed much more hydrogen even at temperatures lower than 600 °C, which means that these oxygen carriers have a strong tendency to generate an oxygen defect when reacted with H$_2$. Co$_3$O$_4$@LSF consumed the most hydrogen at temperatures lower than 600 °C and was followed by Co$_3$O$_4$-NiO@LSF. However, hydrogen consumptions at temperatures up to 600 °C did not fully follow the amounts of CO generated as described in Figure 5. For instance, Co$_3$O$_4$-NiO@LSF presented the highest amount of generated CO during redox cycles although it consumed less hydrogen than Co$_3$O$_4$@LSF at temperatures lower than 600 °C. This is because the consumption of the oxygen defect by CO$_2$ should also be considered to explain the amount of CO production of the oxygen carrier.

![Figure 5. Amounts of CO produced by the oxygen carriers during the oxidation step.](image)

![Figure 6. Results of H$_2$-TPR (a) and CO$_2$-TPO (b) measurements.](image)
Table 2. Results of \( \text{H}_2 \)-TPR and \( \text{CO}_2 \)-TPO measurements.

| Oxygen Carrier       | Consumed \( \text{H}_2 \) (mmol/gOC) | Generated \( \text{CO} \) (mmol/gOC) |
|----------------------|--------------------------------------|-------------------------------------|
|                      | Until 600 °C  | Until 900 °C  | Until 600 °C  | Until 900 °C  |
| LSF                  | 1.75         | 4.72         | 0.28         | 0.29         |
| CeO\(_2\)@LSF       | 0.47         | 1.86         | 0.11         | 0.11         |
| NiO@LSF             | 3.79         | 6.14         | 1.27         | 1.58         |
| Co\(_3\)O\(_4\)@LSF| 8.18         | 9.06         | 1.48         | 6.26         |
| Co\(_3\)O\(_4\)-NiO@LSF | 5.68       | 7.42         | 1.89         | 5.71         |

To observe oxygen defect consumptions during the \( \text{CO}_2 \) oxidation step, \( \text{CO}_2 \)-TPO measurements were carried out on reduced oxygen carriers. Amounts of \( \text{CO} \) generation during \( \text{CO}_2 \)-TPO measurements are presented in Figure 6b and Table 2. Based on considerations of RWGS-CL at an operating temperature of 600 °C, \( \text{CO} \) formation up to 600 °C was explained better with the trend of RWGS-CL in Figure 5. Co\(_3\)O\(_4\)-NiO@LSF generated the highest amount of \( \text{CO} \) at temperatures up to 600 °C in \( \text{CO}_2 \)-TPO measurements, which explained why it produced the most \( \text{CO} \) in the isothermal experiment in Figure 5. Although Co\(_3\)O\(_4\)@LSF showed the highest amount of \( \text{CO} \) production at temperatures up to 900 °C, more than two-thirds of this \( \text{CO} \) was generated between 600 and 900 °C. Co\(_3\)O\(_4\)-NiO@LSF was chosen as the optimum metal oxide core for RWGS-CL.

3.4. Long-Term Stability Test

Figure 7 portrays the result of the long-term stability test for the Co\(_3\)O\(_4\)-NiO@LSF oxygen carrier. As shown by the figure, the amount of \( \text{CO} \) produced by Co\(_3\)O\(_4\)-NiO@LSF gradually decreased until the 10th cycle and then remained constant from the 10th to the 20th cycles. More specifically, the amounts of \( \text{CO} \) generated during the 2nd, 10th, and 20th cycles were 1.68, 1.37, and 1.33 mmol/gOC, respectively. According to the result, Co\(_3\)O\(_4\)-NiO@LSF maintained 97.6% of \( \text{CO} \) production at the 20th cycle as that observed at the 10th cycle, which indicated that the productivity of \( \text{CO} \) on this sample was stabilized after the 10th cycle. Furthermore, even after 20 redox cycles, Co\(_3\)O\(_4\)-NiO@LSF showed clearly higher performance than the values seen during five redox cycles of LSF, CeO\(_2\)@LSF, and NiO@LSF (Figure 5). For Co\(_3\)O\(_4\)@LSF (Figure 8), amounts of \( \text{CO} \) generated at the 2nd, 10th, and 20th cycles were 1.49, 1.23, and 0.92 mmol/gOC, respectively, which presents the gradual decrease of the activity of oxygen carrier. This is because Co\(_3\)O\(_4\)@LSF requires a higher temperature than 600 °C to be fully oxidized by \( \text{CO}_2 \) (Figure 6b). From the long-term stability test, it was found that the Co\(_3\)O\(_4\)-NiO@LSF sample showed not only the highest \( \text{CO} \) productivity among all samples but also the highest stability during the multiple redox cycles.

Table 3 shows the comparison between the core–shell structured oxygen carrier with others in terms of \( \text{CO} \) production. Most previous research reported lower \( \text{CO} \) production than the core–shell structured oxygen carrier. Although one research that used Co-based perovskite showed a higher \( \text{CO} \) production [36], the operating temperature of Co-based perovskite (850 °C) is much higher than the present research (600 °C). From this comparison, it was confirmed that Co\(_3\)O\(_4\)-NiO@LSF was sufficiently competitive for \( \text{CO} \) production at the relatively low temperature (600 °C).
Figure 7. Cyclic performance of Co$_3$O$_4$-NiO@LSF during the long-term stability test.

Figure 8. Cyclic performance of Co$_3$O$_4$@LSF during the long-term stability test.

Table 3. Comparison for CO productions with various oxygen carriers.

| Oxygen Carrier                | Reference | Temp. (°C) | Produced CO (mmol/gOC) |
|-------------------------------|-----------|------------|------------------------|
| La$_{0.75}$Sr$_{0.25}$CoO$_3$ | [36]      | 850        | 4.03                   |
| Fe$_2$O$_3$-Ce$_{0.5}$Zr$_{0.5}$O$_2$ | [28] | 800        | 1.00 *                 |
| Fe$_2$O$_3$-Al$_2$O$_3$       | [37]      | 750        | 0.84 *                 |
| CeO$_2$-Fe$_2$O$_3$          | [38]      | 600        | 1.01 *                 |
| Co$_3$O$_4$-NiO@LSF         | This study| 600        | 1.33                   |

* Estimated from experimental results.

4. Conclusions

This paper investigated the potential use of metal oxide core–perovskite shell oxygen carriers in a reverse water–gas shift chemical looping (RWGS-CL) process. La$_{0.75}$Sr$_{0.25}$FeO$_3$ (LSF) and MeO$_x$@LSF (MeO$_x$: CeO$_2$, NiO, Co$_3$O$_4$, and Co$_3$O$_4$-NiO) oxygen carriers were prepared and subjected to RWGS-CL experiments. Cyclic RWGS-CL experiments showed that NiO@LSF, Co$_3$O$_4$@LSF, and Co$_3$O$_4$-NiO@LSF generated more CO than LSF. CeO$_2$@LSF showed a similar CO amount with LSF due to the relatively
low reducible oxygen capacity of CeO$_2$ (0.35–0.55 mmol/gOC) as compared with those of Co$_3$O$_4$ and NiO-based oxygen carriers (1.15–1.6 mmol/gOC). Temperature-programmed redox reactions were conducted to determine the oxygen capacity and mobility in the reduction and oxidation steps. These analyses revealed that the core–shell structured oxygen carriers effectively formed oxygen defects and consumed at relatively low temperature. Co$_3$O$_4$-NiO@LSF was chosen as the optimum oxygen carrier because it possessed the advantages of Co$_3$O$_4$@LSF and NiO@LSF. Moreover, the amount of produced CO was stably maintained for 20 cycles on Co$_3$O$_4$-NiO@LSF.

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