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Feasibility Study of an Iron-Based Composite Added with Al₂O₃/ZrO₂ as an Oxygen Carrier in the Chemical Looping Applications

Ching-Ti Kao *, Cheng-Hsien Shen and Heng-Wen Hsu

Green Energy and Environment Research Laboratories, Industrial Technology Research Institute (ITRI) 360, Gao Fa 2nd Rd., Guiren Dist., Tainan City 711010, Taiwan; CHShen@itri.org.tw (C.-H.S.); hsuhw@itri.org.tw (H.-W.H.)
* Correspondence: chingt423@itri.org.tw

Abstract: The chemical looping process is a promising approach for carbon capture. Oxygen carriers play the crucial role of carrying oxygen between oxidation and reduction reactors. In this study, iron-based composites, added with alumina and zirconia, were used as the oxygen carriers. The feasibility study of these composites for chemical looping applications was then evaluated by measuring their properties, including mechanical properties, relative density, microstructures, crystal structure, and their capacity of oxygen. The results suggest that the addition of zirconia led the decrease of the bulk relative density and thus had a negative effect to both crush strength and attrition. Crush strength declined from 57 kgf to 26 kgf when using zirconia, replacing alumina, in an iron-based composite as the inner material. In addition, the phases in oxidizing and reducing reaction were also revealed. The formation of the spinel phase (Fe₆Al₂O₁₄) was the major factor that altered the capacity of oxygen. It inhibited Fe₂O₃’s ability to be completely reduced to Fe and thus decrease the capacity of oxygen. The value was therefore decreased from 9.7% to 6.2% after 50 redox cycles in alumina addition composite. On the other hand, for the zirconia addition, all of the Fe₂O₃ could transform to Fe, which provided 8.5% of oxygen capacity after 50 redox cycles. A dense layer which was identified as the Fe₂O₃ in the bulk surface was observed in the samples reacted with 50 redox cycles. The proposed mechanism of the formation of Fe₂O₃ layer and its corresponding kinetic analysis was also revealed in this study.

Keywords: chemical looping; oxygen carrier; microstructure; crystal structure

1. Introduction

1.1. Background for Carbon Reduction

Most developed countries have been focusing on carbon-reducing technologies for many years to achieve carbon neutrality by 2050. Carbon capture and storage is the most promising approach to achieve this objective [1].

Between 2010 and 2018, more than 70% of the power generated in Taiwan involved fossil fuels. Moreover, nearly 90,000 kt CO₂ is emitted every year [2]. According to statistics released by the International Energy Agency in 2018, global CO₂ emissions totaled 4.42 t CO₂; those of Taiwan were 10.83 t CO₂ per capita—twice the global average [3]. Therefore, lowering CO₂ emissions in Taiwan is imperative.

1.2. Chemical Looping Process (CLP)

The chemical looping process is a highly promising method for reducing CO₂ emissions because of its inherent gas separation property. It also does not require additional facilities to separate CO₂ from flue gas. The schematic of chemical looping is displayed in Figure 1. In the chemical looping process, metal oxide (MeₓOᵧ), usually used as the oxygen carrier, passes through the reduction reactor and reacts with fuel, such as coal, natural
gas, biomass, and so on. High-purity CO₂ is then produced and can be used directly or stored. The oxygen-depleted particle (MeₓOₙ, y > z) can circulate to the oxidation reactor and reacts with air; subsequently, oxygen within the oxygen carrier can be recovered again. Some further applications have been proposed by using this concept.

**Figure 1.** Schematic of the chemical looping process.

### 1.2.1. Chemical Looping Combustion (CLC)

Instead of air, the oxygen contained by the oxygen carrier is applied to combust with fuel in the chemical looping combustion process. Nitrogen can be avoided through this combustion process. For example, only CO₂ is produced from a reducer with the use of coal as the fuel, which is illustrated in Figure 2. The exhaust gas (usually composed with N₂ and low concentration of O₂) from the oxidation reactor will have a high temperature which could be introduced to a gas turbine to produce electricity [4]. The heat value is altered by the composition of the fuel and the material of the oxygen carrier. While all of the combustion reactions are exothermic reactions, the reaction between the fuel and oxygen carrier could be endothermic or exothermic [5]. In most cases, the oxygen carrier should also act as the heat carrier which can conduct the heat from the oxidation reactor to provide to heat to the reduction reactor.

**Figure 2.** Schematic of the chemical looping combustion, chemical looping oxygen uncoupling, and chemical looping air separation.

### 1.2.2. Chemical Looping Oxygen Uncoupling (CLOU)

Since the oxygen carriers are usually solid, the reaction rate with the use of gaseous or liquid fuel such as natural gas is still acceptable for the particle use. However, in the case of the solid fuel such as coal, the reaction only happens in the interface of solid to solid, which leads to a slow reaction rate. Chemical looping oxygen uncoupling (CLOU) was therefore proposed for improving this problem. Chemical looping oxygen uncoupling was first proposed by Lyngfelt and Mattiso in 2009 [6]. The differences between CLC and CLOU...
are mainly in the material of the metal oxide (usually are Cu or Ni). By manipulating the
temperature and oxygen partial pressure, the oxygen can release from the oxygen carrier
spontaneously. The solid fuel can not only react with the oxygen carrier but also with
gaseous oxygen, which is illustrated in Figure 2. The reaction rate of the CLOU was almost
50 times faster than that in the CLC [7].

1.2.3. Chemical Looping Air Separation (CLAS)

Chemical looping air separation (CLAS) is a further application from chemical loop-
ing oxygen uncoupling (CLOU), presented in Figure 2. By introducing the inner gas or
vacuuming the reduction reactor, the oxygen partial pressure can be reduced. Pure oxygen
can be released and collected from the oxygen carrier subsequently [8].

Compared to the conventional methods for oxygen production such as cryogenic air
separation (CAS) and pressure swing adsorption (PSA), CLAS has the specifications of
a lower energy penalty and higher outlet temperature [9]. According to the simulation
results revealed from Newcastle University, the oxygen produced by the CLAS applied to
combustion can only have one-tenth in the energy consumption than that of the CAS [10,11].
In addition, integrating the CLAS process and Oxy-fuel combustion or solid oxide fuel cell
was also a novel application. With the use of higher-temperature oxygen produced from
CLAS, it can avoid the gas preheating process, which can reduce the energy consumption.

1.2.4. Syngas Chemical Looping (SCL) and Chemical Looping Hydrogen Generation
(CLHG)

Through controlling the oxidation state of the metal oxide and the oxygen pressure,
the composition of exhaust gas can be manipulated. For the syngas chemical looping
process (SCL), CO₂, syngas (CO + H₂) can be obtained from the reduction reactor by the
incomplete reaction between the fossil fuel and oxygen carrier [12]. This process can further
combine with the integrated gasification combined cycle (IGCC), which might provide
higher thermal efficiency [13].

Chemical looping hydrogen generation (CLHG) is another potential application of the
chemical looping process. The oxygen carrier developed in this study tends to be applied
in the CLHG process. An additional reactor, which is called a steam reactor, is needed for
hydrogen generation. The steam reactor is usually set up after the reduction process and
before the oxidation process [14]. Compared to other chemical looping applications, metal
oxide using in the CLHG should be deeply reduced, which can provide higher reactivity
for hydrogen generation. Taking iron-based oxygen carriers as an example, iron (Fe) can be
oxidized by steam to be wüstite (FeO) or magnetite (Fe₃O₄). Hydrogen can be produced
through Equation (1) and Equation (2). Afterwards, the reduced oxygen carrier will be
transported to an oxidation reactor to react with air. The phase of iron can be completely
oxidized again.

\[
Fe + H₂O \rightarrow FeO + H₂ \quad (1)
\]

\[
3FeO + H₂O \rightarrow Fe₃O₄ + H₂ \quad (2)
\]

Compared to the conventional method for hydrogen generation of steam methane
reforming (SMR), CLHG has the advantage of lower energy consumption [15]. For example,
the CLHG process can reach higher efficiency due to the preference in its thermodynam-
ics [16]. In addition, it also provides more effective heat usage than SMR process due to the
lower demand for gas cooling [15].

In recent years, CLHG has been not only considered as a single technology but also
as one of the units which can integrate combustion and hydrogen generation, as has been
proposed. For example, CLHG can provide high temperature H₂ to solid oxide fuel cell
(SOFC) processes directly as the fuel which can avoid the energy penalty from the gas
pre-heating process [17,18]. On the other hand, it can also be combined with a supercritical
water gasification system (SCWG) to deal with biomass or waste [19]. Moreover, the concept
of introducing CLHG to chemical production, such as ammonia, was also proposed [20].
On the contrary, the H₂ produced by the CLHG technique is not suitable for storage, such as in a high-pressure cylinder. Additional processes including gas cooling, purification, and pressurization are needed, and thus raise the cost of H₂ production.

1.3. Selection for Oxygen Carrier

Some transition-metal oxides such as iron oxides, nickel oxides, and copper oxides are attractive oxygen carrier candidates. Among them, iron oxides have lower cost, a higher melting point, and a smaller environmental impact than other candidates [5]. In addition, iron-based materials have been revealed as the more suitable reactant for use in CLHG applications because its phase conversion temperature is close to the operating temperature for CLHG [16,21].

Several benefits can be achieved by adding an inner support in the matrix material. The addition of an inner support can disperse the matrix material and minimize the sintering effect between each particle. Some ceramic materials such as alumina, zirconia, and silica are promising inner support candidates for iron oxides [22,23].

Among these ceramic materials, alumina catches the most attention as the inner support for iron oxides because of its remarkable mechanical properties. Several studies for the composite between iron oxide and alumina have been discussed [24,25]. However, some drawbacks for the iron-based with alumina have been revealed. The ability to resist sintering for this iron-based composite was not sufficient for chemical looping combustion or hydrogen generation [26]. The reactivity for this composite will gradually decline under long-term operation.

In addition, some of the studies suggested that alumina was not a good candidate as the inner support material for use in CLHG due to the carbon decomposition effect. This might be caused by the lack of oxygen vacancy for alumina [27]. Another challenge for the iron oxide and alumina composite was the formation of the spinel phase (Fe₃Al₂O₆). The high stability of this spinel structure limits the transformation of iron which might reduce the capacity of oxygen for this composite [24]. However, Fe₃Al₂O₆ was generally regarded as the phase which could benefit chemical looping combustion (CLC). The formation of Fe₃Al₂O₆ could prevent the sintering between each particle [28]. Moreover, it could be reversed to Fe₂O₃ and Al₂O₃ in an oxidizing reaction and then be involved in the reaction again. The role for Fe₃Al₂O₆ applied in CLC or CLHG is still not clear.

In this work, an iron-based composite with alumina and zirconia was selected as the oxygen carrier in chemical looping applications, especially for hydrogen generation. Introducing zirconia into the composite can provide some advantages such as a relatively high Tammann point (ZrO₂ ≈ 1358 °C, Al₂O₃ ≈ 1036 °C, SiO₂ ≈ 855 °C at 1 atm), which can increase resistance to the sintering effect [7]. In addition, either iron oxides or alumina did not have any interaction with zirconia, which meant no second phase could be found in the operating temperature around 900 °C, presented in the phase diagram [29,30].

Further, some researchers suggested that the addition of Zr can enhance Fe₂O₃’s ability to deeply reduce to FeO which could promote the efficiency of hydrogen generation [31]. Furthermore, Zirconia has a similar thermal expansion coefficient to iron oxides compared to alumina (Al₂O₃ ≈ 7.2 × 10⁻⁶ K⁻¹, ZrO₂ ≈ 10.5 × 10⁻⁶ K⁻¹, and Fe₂O₃, Fe₃O₄, FeO and Fe ≈ 8–15 × 10⁻⁶ K⁻¹), which could help to reduce the thermal stress during the redox reaction [32,33]. In this study, the composites of Fe₂O₃, Al₂O₃, and ZrO₂ were synthesized in order to discover the better oxygen carrier for chemical looping hydrogen generation techniques. The role of Fe₃Al₂O₆ for CLHG will also be revealed.

By manipulating the ratio between ZrO₂ and Al₂O₃, which serve as the inner material, a series of properties for iron-based composites, including relative density, microstructures, capacity of oxygen, and mechanical properties, were examined. The iron’s phases, which existed in the oxidizing reaction and reducing reaction, were also measured in order to clarify how they altered the oxygen capacity. The feasibility of the iron-based composites with added alumina and zirconia for chemical looping applications was then evaluated.
2. Materials and Methods

2.1. Samples Preparation

The starting powder of iron oxide (Fe$_2$O$_3$, purity > 99%, Himag Co., Ltd., Gloucestershire, UK) mixed with various proportions of alumina (Al$_2$O$_3$, purity > 99%, Showa, Gyoda, Japan) and zirconia (ZrO$_2$, purity > 99%, HonWay Materials Ltd., Kaohsiung, Taiwan) was prepared through a mechanical mixing process. Ten percent by weight of starch was also added as the pore former to each composite. The powder mixing was performed by a wet process in which dilute water was added as the solvent. The mixed powder was further dried in an oven at 80 °C for 1 day. According to our previous study, the mixing ratio for matrix/support of 40/60 wt% exhibited the best efficiency for dispersion [34]. The composition between matrix and support material was then determined in this study.

Granulation and sieving processes were then conducted by granulator. The dried powder was granulated by the mesh of 1000, 800, 500, 270, and 180 µm, respectively. The particle range of 180 to 270 µm, which displayed the best powder flowability, was selected for the following uniaxial pressing process. A green compact (diameter: 3 mm; height: approximately 2.8 mm) was pressed through uniaxial pressing. The denotations and compositions of the composites used in this study are presented in Table 1. According to our previous study, the selected sintering temperature of 1200 °C for the iron-based materials could serve moderate porosity and strength for a moving bed system [34]. The sintering process was conducted in a muffle furnace at a heating and cooling rate of 5 °C/min, and the dwell time at highest temperature was 1 h in air.

Table 1. Denotations and compositions of the composites in the present study.

| Composition | Fe$_2$O$_3$ Content (wt%) | Al$_2$O$_3$ Content (wt%) | ZrO$_2$ Content (wt%) |
|-------------|---------------------------|---------------------------|-----------------------|
| Fe$_{40}$Al$_{60}$ | 40 | 60 | 0 |
| Fe$_{40}$Al$_{55}$Zr$_5$ | 40 | 55 | 5 |
| Fe$_{40}$Al$_{50}$Zr$_{10}$ | 40 | 50 | 10 |
| Fe$_{40}$Al$_{40}$Zr$_{20}$ | 40 | 40 | 20 |
| Fe$_{40}$Zr$_{60}$ | 40 | 0 | 60 |

2.2. Phase Analysis

The phase of the composites was analyzed with the X-ray diffraction (XRD, D2 Phaser, Bruker, USA) technique. Narrow scan for XRD was adopted for distinguishing the phases, including Fe, Fe$_2$O$_3$, m-ZrO$_2$, t-ZrO$_2$, Al$_2$O$_3$ and FeAl$_2$O$_4$. The specific scanning ranges were selected from 28° to 38° and 28° to 46°. The interval of each step was 0.02°, and the collection time lasted for 1 s. A better signal to noise ratio for XRD results could therefore be achieved through narrow scan measurement.

2.3. Mechanical Properties Measurement

The relative density of the as-sintered samples was measured through Archimedes’ method. The theoretical density of each composite was estimated using the lever rule. Ten pieces of the sintered samples were measured. At least 50 bulks were measured, and the average density was obtained. The microstructures of the samples were observed through scanning electron microscopy (SEM, SUS8010, Hitachi, Japan). Elemental analysis of the samples was performed through energy-dispersive X-ray spectroscopy (EDS, X-max, HORIBA, Japan) coupled with SEM.

The crushing strength of the as-sintered samples was measured using a digital force gauge (FGP-100, Shimpo, Japan). The samples were heated to 110 °C in an oven for 1 h before the test to eliminate the influence of water. The sample was placed in the radial side and then subjected to a vertical compressive force until it was broken. At least 10 samples were measured, and the average crushing strength was obtained. Attrition was measured...
using an abrasion machine, following the ASTM D4058-96 standard. The formula for attrition is as follows:

\[
\text{Attrition (\%)} = \frac{W_{\text{pan}}}{W_{20} + W_{\text{pan}}} \times 100\%
\]  

where \(W_{\text{pan}}\) is the weight of the sample that remained in a screen mesh of 20, and \(W_{20}\) is the weight of the sample that was sieved through a screen mesh of 20.

2.4. Measurement for the Capacity of Oxygen

The weight change for the sample during the redox reaction was measured through thermogravimetric analysis (SETSYS Evolution TGA, Setaram, France), which can provide information on iron’s phase changing during reduced and oxidized reaction.

The heating and cooling ramp rate of the TGA measurement was 10 °C/min, and the operation temperature for the redox reaction was 900 °C. One redox cycle included an oxidation reaction (lasting 10 min), purging (lasting 5 min), a reduction reaction (lasting 40 min), and a purging (lasting 5 min) process. The flow rate of air and the mixed H\(_2\)/N\(_2\) gas (20/80 in vol%) for oxidation and reduction was 200 mL/min, and the flow rate of nitrogen used as the purging gas before each oxidation and reduction process was 200 mL/min.

The capacity of oxygen can be calculated by comparing the weight difference of sample, which was in oxidizing and in reducing state. In this study, the content of Fe\(_2\)O\(_3\), regarded as the reactant, was fixed in consistency for all composites. This value can also be known for the performance of the oxygen carrier used in the chemical looping process [5].

\[
\text{Capacity of oxygen} \% = 100 \times \frac{\text{Weight}_{\text{oxidizing}} - \text{Weight}_{\text{reducing}}}{\text{Initial weight of sample}}
\]  

3. Results and Discussion

3.1. Characterization of the As-Sintered Composites

The images of the as-sintered samples are displayed in Figure 3a. The samples were deep brick red and turned darker with an increase in the zirconia content. The dimensions of the as-sintered samples are listed in Table 2. All of the samples had a similar diameter and height after sintering at 1200 °C. The images of the as-sintered samples after 50 redox reaction cycles are depicted in Figure 3b.
Table 2. Summary of relative densities and dimensions of the as-sintered samples.

| ZrO$_2$ Content (wt%) | Relative Density (%) | Porosity (%) | Dimension
|------------------------|----------------------|--------------|-------------
|                        |                      |              | Height (mm) | Diameter (mm) |
| 0                      | 66.42 ± 1.96         | 33.58        | 2.55 ± 0.02 | 2.85 ± 0.03   |
| 5                      | 62.32 ± 0.99         | 37.68        | 2.65 ± 0.04 | 2.85 ± 0.02   |
| 10                     | 62.43 ± 5.49         | 37.57        | 2.68 ± 0.03 | 2.87 ± 0.03   |
| 20                     | 60.68 ± 1.45         | 39.32        | 2.58 ± 0.02 | 2.85 ± 0.01   |
| 60                     | 54.45 ± 5.81         | 45.55        | 2.38 ± 0.04 | 2.77 ± 0.02   |

As demonstrated in Table 2, except for the Fe$_{40}$Zr$_{60}$, all of the as-sintered samples presented a similar relative density. Several factors can influence the relative density; the phase transformation process of zirconia might be the major reason in the present case. Zirconia will undergo phase transformation from its tetragonal to monoclinic phase during the cooling process, for which the transformation temperature is around 1100 °C. This phase transformation typically produces approximately 5% to 8% volume expansion [35], thus forming some cracks inside the bulk. Moreover, the higher densification temperature needed for ZrO$_2$ (1500–1600 °C) than that of Al$_2$O$_3$ (1400–1500 °C) was another reason for the lower density of ZrO$_2$-added samples [36,37].

XRD analysis was used for the phase identification in this study. In the phase diagram of iron oxides and zirconia, there is no second formed within the operating temperature in this study. In case of the phase diagram of iron oxides and alumina, FeAl$_2$O$_4$ is one of the most common phases existing in iron oxide and alumina composites [38]. The formation of FeAl$_2$O$_4$ might decrease the reactivity for the chemical looping process because it limits the phase transformation of iron [5]. The main peaks for FeAl$_2$O$_4$ were located at 31.1° and 36.6° in the XRD pattern, which was revealed in previous study [39], and cannot be detected in the as-sintered samples, presented in Figure 4.

For the increase of zirconia content, the peaks which are identified as the monoclinic phase (m-ZrO$_2$) located at 28.2° and 31.4° gradually appear. Furthermore, the intensity
of the tetragonal phase (t-ZrO$_2$) peaks, which represent the high temperature phase of zirconia, was too weak to be found in Figure 4. The results of XRD suggest that most of the zirconia have undergone the phase transformation during the cooling process, which led some cracks produced within the samples. This might be the reason that the zirconia addition sample has the lowest relative density, which is listed in Table 2.

### 3.2. Mechanical Properties

The crushing strength and attrition, two of the most critical factors, of the as-sintered samples were measured to evaluate their lifetime in chemical looping. For either a fluidized bed or moving bed reactor in chemical looping, the material should be sufficiently strong to withstand the impacts of several looping cycles.

The value of crush strength can be used as the index for estimation of how well a particle withstands impacts during the looping system. In addition, the value of attrition can serve as the index of the resistance of attrition between each particle.

The influence of relative density on the crushing strength and attrition of the as-sintered samples is displayed in Figure 5. The result suggests that both crushing strength and attrition were highly dependent and exhibited a linear relationship with relative density, for which the R-squared values were 0.945 and 0.983, respectively.

![Figure 5. Relationship among the relative density, crushing strength, and attrition of the as-sintered samples.](image)

The addition of ZrO$_2$ in Fe$_2$O$_3$ had a negative effect on the mechanical properties. Typically, with the increase of zirconia content, crushing strength decreased and attrition increased. This is because most of the mechanical properties, especially strength, are highly dependent on the relative density [40]. The difference of relative density for as-sintered samples was thus the major influence upon both crushing strength and attrition.

### 3.3. Capacity of Oxygen for the Composites

Three individual samples for each iron-based composition were measured by TGA analysis in order to ensure reproducibility. The setting parameter of TGA analysis in this study was referred to the Ellingham diagram, which can estimate the equilibrium phase in certain thermodynamic conditions [41]. According to the application for CLHG, Fe$_2$O$_3$ should be reduced to Fe or FeO. The performed atmosphere for TGA analysis was therefore set in air (oxidation) and 20% H$_2$ with N$_2$ compensation (reduction) at 900 °C.

TGA could be regarded as the small scale of a fixed bed reactor. Each sample was set in the alumina crucible and hung inside the furnace. The sample was reduced by reduction gas, and the weight of the sample was then decreased due to the oxygen was
released. Afterward, nitrogen was used as the purge gas, which removed the reduction gas remaining from the furnace. Air was subsequently switched to the furnace, so that the sample could capture oxygen from the air to oxidize and then increase its weight. The definition of the capacity of oxygen is given in Section 2.4. The capacity of oxygen for the as-sintered samples with 50 redox cycles is displayed in Figure 6. All of the reacted samples showed a drastic decline for their capacity of oxygen in the first 10 redox cycles and became more stable after the 15th cycle.

Figure 6. Capacity of oxygen for the as-sintered samples with 50 redox reaction cycles.

The samples at the 15th cycle in the oxidizing and reducing reaction were chosen for TGA analysis. The reason for the decrease of the capacity of oxygen could then be determined. The cooling process was carried out at the relatively fast ramp of 10 °C/min in order to maintain its phase from measurement temperature. Through the analysis of XRD, the phase which existed in reducing and oxidizing reaction can be obtained, presented in Figure 7. The results of XRD measurement are also summarized in Table 3.

The composition of phase is similar between the sample in the oxidizing reaction and the as-sintered one regardless of the addition of alumina or the zirconia composites. However, for the composites with an alumina addition in reducing reaction, a spinel phase FeAl$_2$O$_4$ was formed. FeAl$_2$O$_4$ was mainly composited of FeO and Al$_2$O$_3$. The formation of FeAl$_2$O$_4$ restricted Fe$_2$O$_3$ completely reduced to Fe which decreased the capacity of oxygen. On the other hand, all of the Fe$_2$O$_3$ can fully convert to Fe in the zirconia addition sample due to the lack of second phase formation. The conversion to Fe from Fe$_2$O$_3$ can promote the efficiency of chemical looping hydrogen generation, as presented in Equation (1). The increase of the capacity of oxygen can also provide more oxygen to react with fuel, which can generate more heat during oxidation reaction to achieve the thermal equilibrium for the chemical looping system. In terms of thermodynamics, the addition of zirconia is more suitable than that of alumina in iron-based composite for chemical looping hydrogen generation.
Figure 7. XRD pattern for the as-sintered samples reacted at 15th cycle in (a) oxidizing and (b) reducing reaction.
Table 3. Summary of the phase composition for the iron-based composites with alumina and zirconia addition.

| Composites   | As-Sintered       | Oxidation                        | Reduction                        |
|--------------|-------------------|----------------------------------|----------------------------------|
| Fe_{40}Al_{60} | Fe_{2}O_{3}, Al_{2}O_{3} | Fe_{2}O_{3}, Al_{2}O_{3}        | FeAl_{2}O_{4}, Al_{2}O_{3}, Fe    |
| Fe_{40}Al_{55}Zr_{5} | Fe_{2}O_{3}, Al_{2}O_{3}, m-ZrO_{2} | Fe_{2}O_{3}, Al_{2}O_{3}, m-ZrO_{2}, t-ZrO_{2} | FeAl_{2}O_{4}, Al_{2}O_{3}, Fe, m-ZrO_{2} |
| Fe_{40}Al_{50}Zr_{10} | Fe_{2}O_{3}, Al_{2}O_{3}, m-ZrO_{2} | Fe_{2}O_{3}, Al_{2}O_{3}, m-ZrO_{2}, t-ZrO_{2} | FeAl_{2}O_{4}, Al_{2}O_{3}, Fe, m-ZrO_{2} |
| Fe_{40}Al_{40}Zr_{20} | Fe_{2}O_{3}, m-ZrO_{2}, t-ZrO_{2} | Fe_{2}O_{3}, Al_{2}O_{3}, m-ZrO_{2} | FeAl_{2}O_{4}, Al_{2}O_{3}, Fe, m-ZrO_{2} |
| Fe_{40}Zr_{60} | Fe_{2}O_{3}, m-ZrO_{2}, t-ZrO_{2} | Fe_{2}O_{3}, Al_{2}O_{3}, m-ZrO_{2} | Fe, m-ZrO_{2} |

3.4. Influence of the Microstructure on Oxygen Carrier

All of the samples for SEM observation were mounted in epoxy and then ground and polished to obtain the cross-sectional view from the surface to the bulk. The corresponding microstructures of as-sintered and the samples with 50 redox cycles are depicted in Figure 8a,b, respectively. The bright particles can be identified as Fe_{2}O_{3} (M = 159.69 g/mol) because they have a higher molecular weight than Al_{2}O_{3} (M = 101.96 g/mol) and ZrO_{2} (M = 123.218 g/mol), which was verified through EDS measurements (not shown in the present study).

Cracks can be observed in the Fe_{40}Al_{40}Zr_{20} and Fe_{40}Zr_{60} samples, presented in Figure 8a, and this echoes the phase transformation of zirconia. The further analysis of this layer was conducted by using EDS mapping, and was identified as the Fe_{2}O_{3}, presented in Figure 8c. Moreover, its thickness increased along with the zirconia addition, given in Figure 8b. This layer could reach almost 5 μm in thickness for the Fe_{40}Zr_{60} sample.

This Fe_{2}O_{3} layer acted as the barrier that inhibited the fuel gas reacting with the inner material. The capacity of oxygen thus decreased with the redox reaction in the samples of Fe_{40}Al_{40}Zr_{20} and Fe_{40}Zr_{60}. In case of the Fe_{40}Al_{40}Zr_{20}, due to the influence from both the formation of FeAl_{2}O_{4} and the dense layer of Fe_{2}O_{3} formed on the surface, it exhibited the lowest capacity of oxygen after 50 redox reactions, presented in Figure 7. On the other hand, for the sample of Fe_{40}Zr_{60}, the Fe_{2}O_{3} layer was peeled off from the bulk because its thickness was too thick. The fresh surface appeared again and could react to the inner material again, which could maintain its reactivity.

![Figure 8. Continuation](image)
Figure 8. Cross-sectional view for (a) as-sintered samples and (b) after 50 redox reaction cycles at 900 °C, and (c) EDS mapping result for Fe$_2$O$_3$ layer formed in surface of the specimen.

More detailed analysis of the dense Fe$_2$O$_3$ layer formed on the surface was needed. In this study, the proposed mechanism of the formation for Fe$_2$O$_3$ layer and its corresponding kinetic analysis is given in Figures 9 and 10.

3.5. Kinetic Analysis for Redox Reaction

The kinetic analysis given in Figure 9 for redox reaction was through the thermal-gravity measurement. Each redox reaction exhibited a similar result within the 50 cycles, presented in Figure 9a. Two distinct peaks can be found during the reduction reaction, and only one peak during oxidation. The further analysis of single redox at 10th cycle reaction is displayed in Figure 9b. Two distinct peaks can be found during the reduction reaction and only one peak during oxidation.

During the oxidation reaction, iron on the surface was oxidized by oxygen to Fe$_2$O$_3$. Meanwhile, the volume of Fe$_2$O$_3$ particles expanded, which is presented in Figure 10. Afterward, each Fe$_2$O$_3$ particles were closer together, which led them to easily sinter together; the dense layer was then formed on the surface. This dense layer acted as a barrier which inhibited the reaction between the inner material and fuel gas. The corresponding kinetics curve for the oxidizing reaction was exhibited in Figure 9b. Only one characteristic peak formed in the oxidizing reaction with the fastest reaction rate which was the special feature for the surface reaction. In addition, no other peak could be observed in the
The proposed schematic diagram for the reducing reaction is given in Figure 10. \( \text{Fe}_2\text{O}_3 \) reacted with the reducing gas, and then was reduced to Fe. Because the volume of particles shrank, the interspace between each particle was formed. Afterward, reducing gas could further react with the inner material. Two characteristic peaks can be obtained in the kinetic analysis diagram during reducing reaction, presented in Figure 9b. The first peak exhibited the faster reaction rate and was the initial reaction; it was the reaction between surface particles and the reducing gas.

![Figure 9](image_url)

**Figure 9.** (a) Reaction kinetic analysis for redox reaction and (b) further analysis for single redox reaction.
The second peak with the feature of a slower reaction rate and longer reaction time period could be regarded as the inner reaction. According to Figure 10, the reduction reaction that happened in the inner material should rely on the interspace which was formed through particle shrinkage. The reaction was thus needed to take more time to occur layer by layer. In case of the Fe\textsubscript{40}Al\textsubscript{60}, the shrinkage of particle volume from Fe\textsubscript{2}O\textsubscript{3} to FeAl\textsubscript{2}O\textsubscript{4} was not as much compared to Fe\textsubscript{2}O\textsubscript{3} to Fe. The width of the interspace could be assumed to be smaller than that of Fe\textsubscript{40}Zr\textsubscript{60}, in which all of the Fe\textsubscript{2}O\textsubscript{3} could be completely reduced to Fe. The rate for inner reaction during reduction was therefore slower than that of Fe\textsubscript{40}Zr\textsubscript{60}.

### 3.6. Formation of the Fe\textsubscript{2}O\textsubscript{3} Layer

The phase separation effect can result in the driving force of the formation of the Fe\textsubscript{2}O\textsubscript{3} layer. A previous study using an Fe-Ni-O composite revealed phase separation similar to that in the present study; the Kirkendall effect was introduced to explain this phenomenon [42]. Outward and inward diffusion was produced because of the different diffusion coefficients of the Fe and Ni atoms. The core–shell structure (Ni core and Fe shell) was then formed.

For the microstructural images presented Figure 8, the phase separation effect derived from outward and inward diffusion indeed affected the composite in this study. Moreover, the effect of phase separation was enhanced significantly in zirconia addition samples than that of the alumina addition. This is caused by different phases of iron.

In the case of the FeZr composite, all of Fe\textsubscript{2}O\textsubscript{3} was completely reduced to Fe during the reducing reaction, listed in Table 3. Because Fe has a lower Tamman point (764 °C) than the operating temperature (900 °C) conducted in the reducing reaction, Fe will then segregate to the surface by several redox reactions. On the other hand, for the FeAl composite, only part of Fe\textsubscript{2}O\textsubscript{3} can only be reduced to FeO, and then reacts with alumina to become the spinel structure of FeAl\textsubscript{2}O\textsubscript{4}. The higher Tamman point of FeAl\textsubscript{2}O\textsubscript{4} (890 °C) than that of Fe (764 °C) could restrict the Fe segregated to the surface. The phase separation effect was therefore less influenced in the FeAl composite than that in the FeZr.

### 4. Conclusions

This feasibility study evaluated iron-based composites added with Al\textsubscript{2}O\textsubscript{3} and ZrO\textsubscript{2} as oxygen carriers for chemical looping applications. Their corresponding microstructures before and after 50 redox reaction cycles were also analyzed. As for the mechanical properties, both crushing strength and attrition were influenced by the relative density of the bulk. For the measurement of the capacity of oxygen, all of the as-sintered samples presented a drastic decrease within the first 10 redox cycles. An Fe\textsubscript{2}O\textsubscript{3} layer that formed on the surface acted as a barrier to inhibit further redox reactions with the inner layer material.

In this study, the phase in oxidizing and reducing reaction was revealed and analyzed. Both the phases of FeAl\textsubscript{2}O\textsubscript{4} and Fe were formed in the alumina addition samples, but...
only Fe was produced in the zirconia addition sample during the reduction reaction. The formation of FeAl$_2$O$_4$ inhibited Fe$_2$O$_3$ complete reduction to Fe, which might decrease the efficiency if it is used for chemical looping hydrogen generation.

A dense layer which was identified as Fe$_2$O$_3$ in the bulk surface was found in the samples with 50 redox cycles. It acted as the barrier which inhibited the reaction between the inner materials and fuel gas. The capacity of oxygen for zirconia addition composites was then influenced.

Through the kinetic analysis during oxidizing and reducing reaction, the formation of an Fe$_2$O$_3$ layer was revealed in this study. Zirconia as the additive for iron-based composite was more suitable for chemical looping hydrogen generation than that of alumina. However, the cracks which derived from zirconia’s phase transformation should be avoided if it is applied in either a fluidized bed or moving bed reactor.

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